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About the book

Science has evolved over thousands of years. It began out of curiosity about how the world around us works as well as a need to know how to make thing work better. From water management to space travel, science is essential for success.

The evolution of science is layered: Early science depended mostly on critical observation, thus the early scientist was considered a philosopher who created a theory. Soon experiments were designed to test these theories. This process is successful when a new principle is established, leading to a deeper and reproducible experimental observation. This typically takes years. A key component of science is the making and testing of models, that are designed to quantitatively evaluate the results of experiments, in a mathematical framework. Good science is observation and experimentation. Great science is the art of making models that explain experimental results. This always results in a deeper question, suggesting new experiments. Each generation has its geniuses. One of these was Galileo, who was a philosopher, experimentalist, and mathematician.

An understanding of physics requires a knowledge of mathematics. The converse is not true. By definition, pure mathematics contains no physics. Yet historically, mathematics has a rich history filled with physical applications. Mathematics was developed by individuals with the intent of making things work. As an engineer, I see these creators of early mathematics as budding engineers. This book is an attempt to tell the story of the development of mathematical physics as viewed by an engineer.

There are two distinct ways to learn mathematics: by learning definitions and relationships, or by associating each mathematical concept with its physical counterpart. Students of physics and engineering best learn mathematics based on the underlying physical concepts. Students of pure mathematics are taught via definitions of abstract structures, not from the history of mathematical physics. These two teaching methods result in very different understandings of the material.

There is a deep common thread between physics and mathematics: the chronological development, or the history of mathematics. This is because much of mathematics was developed to solve physical problems. Most early mathematics evolved from attempts to understand the world, with the goal of navigating it. Pure mathematics followed as generalizations of these physical concepts.

Around 1638 Galileo stated that, based on his experiments with balls rolling down inclined planes and pendulums, the height of a falling object is given by

\[ h(t) = \frac{1}{2} G t^2, \]

where \( t \) is time and \( G \) is a constant. This formula leads to a constant acceleration \( a(t) \) of the object since

\[ a(t) = \frac{d^2}{dt^2} h(t) = G \]

is independent of time. It follows that the force on a body is proportional to its acceleration \( a \), defined as \( G \) – namely, \( F = a \equiv G \). Thus \( G \) must be the object’s mass, which must be a constant. If the object has a constant forward velocity, then the object will have a parabolic trajectory. The relative mass may be measured using a balance scale. I believe Galileo understood all this.

Years later, following up on the observations from Galileo’s study of pendulums and falling objects, Newton showed that differential equations were necessary to explain gravity and that the force of gravity is proportional to the masses of the two objects divided by the square of the distance between them:

\[ \frac{d^2}{dt^2} r(t) = G \frac{mM}{r^2(t)}. \]

To find \( r(t) \) we must integrate this equation. For an object at height \( h(t) \) above the surface of the earth, \( r(t) = R_e + h(t) \approx R_e \), where \( R_e \) is the radius of earth. In this case, the force is effectively constant, since \( h \ll R_e \). Newton’s equation says the acceleration is constant,

\[ \frac{d^2 h(t)}{dt^2} = G \frac{mM}{R_e^2}, \]
but different from Galileo’s $G$ (a simple mass). Yet it seems clear that the physics behind Newton’s formula for the acceleration $a(t)$ of two large masses (sun and earth, or earth and moon) and Galileo’s physics for balls rolling down inclined planes are the same.\footnote{https://physicstoday.scitation.org/do/10.1063/PT.6.3.20191002a/full/} The difference is that Newton’s proportionality constant is a significant generalization of Galileo’s. But, other than the constant, which defines the acceleration, the two formulas are the same.

This is not a typical mathematics book; rather, it is about the relationship of math and physics, presented roughly in chronological order via their history. To teach mathematical physics in an orderly way, our treatment requires a step backward in terms of the mathematics, but a step forward in terms of the physics. Historically speaking, mathematics was created by individuals such as Galileo who, by modern standards, may be viewed as engineers. This book contains the basic information that well-informed engineers needs to know, as best I can provide.

Let the reader beware that engineering and physics texts do not intend to be rigorous, in the mathematical sense. In some ways, mathematics cleans up the mess by proving theorems, which frequently start with speculations in physics and even engineering. The cleanup is a slow, tedious process. Just because something seems obvious based on the known physical facts does not make it a fundamental theorem of mathematics.

Although while there are similarities between this book and that of Graham et al. (1994), the differences are notable. First, Graham’s Concrete Mathematics presents an impossible standard to be measured against. Second, it is clearly a math book, brilliantly written and targeted at computer science students. This book is not just a math book, it is a mathematical physics text, which depends much underlying math. I would like to believe there are similarities in (1) the broad range of topics, (2) the in-depth discussion, and (3) the use of historical context.

**Organization:** As discussed in Sec. 1.2.2 and Fig. 1.6 (p. 21) the book is divided into three mathematical themes, called streams, presented as five chapters: Introduction, Number Systems, Algebraic Equations, Scalar Calculus, and Vector Calculus. Appendices are used to isolate complex self-contained topics and large Tables, such as those for Laplace Transforms.

Chapter 2, Number Systems, introduces two key concepts, the greatest common divisor (GCD) and the continued fraction (CFA). When we deal with simple electrical networks composed of inductors, resistors, and capacitors (Fig. 3.8, page 110), or mechanical networks consisting of masses, dashpots, and springs, or their equivalent, pendulums, as used by Galileo in his studies of gravity (Figs. 1.3, page 16 and 3.11, page 120), the system may be modeled as a Brune impedance, defined as the ratio of polynomials of the Laplace frequency $s = \sigma + \omega j$ (See Sec. 3.2.2, page 79 and Sec. 4.4.2, page 143). Of special importance is the development of ordinary differential equations (Sec. 3.4.2, and Eq. 3.4.2.4) which under generalized symmetry conditions, called postulates (Sec. 3.9.1, page 121), characterize Brune impedances (Brune, 1931a).

Using the CFA (Sec. 2.5.4), we can generalize the Brune impedance. This generalization results in a transmission line, that describes wave propagation in horns, dealt with in Chapters 4 and 5 (Cauer et al., 1958; Cauer, 1958). This topic is both physically and mathematically important (Cauer, 1932).

The material is delivered in numbered sections (e.g., Sec. 1.1) spread out over a semester of 15 weeks, three lectures per week, with a three-lecture time-out for administrative duties. Eleven problem sets are provided for weekly assignments.

Many students have rated these assignments as the most important part of the course. There is a built-in interplay between these assignments and the lectures. There are four exams, one at the end of each of the three sections plus the final. The first exam is in class, two others and the final are evening exams. When a student returns the exam, the full solution is provided while the exam is fresh in their mind, resulting in a teaching moment. It is my personal belief that, in principle, students can see the exam in advance of taking it, since each exam is entirely based on the assignments.

**Author’s personal statement**

An expert is someone who has made all possible mistakes in a small field. I don’t know if I would be called an expert, but I certainly have made my share of mistakes. I openly state that I love making mistakes because I learn so much from them. One might call that the “expert’s corollary.”

This book has been written out of my love for the topic of mathematical physics, a topic that provides many insights, that lead to a deep understanding of important physical concepts. Over the years I have developed a physical sense of math along with a related mathematical sense of physics. While doing my research,\footnote{https://auditorymodels.org/index.php/Main/Publications} I believe that math can be physics, and physics math. I have come across what I feel are certain conceptual holes that need
filling, and I sense many deep relationships between math and physics that remain unidentified. What we presently teach is not wrong, but it is missing these relationships. What is lacking is an intuition for how math “works.” Good scientists “listen” to their data. In the same way, we need to start listening to the language of mathematics. We need to let mathematics guide us toward our engineering goals.

As summarized in Fig. 1, this marriage of math, engineering, and physics (MEP) helps us make progress in understanding the physical world. We must turn to mathematics and physics when trying to understand the universe. My views follow from a lifelong attempt to understand human communication – that is, the perception and decoding of human speech sounds. This research arose from my 32 years at Bell Labs in the Acoustics Research Department. There such lifelong pursuits were not only possible but were openly encouraged. The idea was that if you are successful at something, take it as far as you can, but, on the other hand, you should not do something well that’s not worth doing. People got fired for the latter. I should have left for a university after a mere 20 years at Bell Labs, but the job was just too cushy.

In this text it is my goal to clarify conceptual errors while telling the story of physics and mathematics. My views have been inspired by classic works, as documented in the Bibliography. This book was inspired by my reading of Stillwell (2002) through his Chapter 21. Somewhere in Chapter 22 I switched to the third edition (Stillwell, 2010), at which point I realized I had much more to master. It became clear that by teaching this material to first-year engineers, I could absorb the advanced material at a reasonable pace. This book soon followed.

Summary

This is foremost a math book, but not the typical math book. First, this book is for the engineering minded, for those who need to understand math to do engineering, to learn how things work. In that sense the book is more about physics and engineering than mathematics. Math skills are essential for making progress in building things, be it pyramids or computers, as clearly shown by the great civilizations of the Chinese, Egyptians, Mesopotamians, Greeks, and Romans.

Second, this is a book about the math that developed to explain physics, to allow people to engineer complex things. To sail around the world, one needs to know how to navigate. This requires a model of the planets and stars. You can only know where you are on earth once you understand where earth is relative to the sun, planets, Milky Way, and distant stars. The answer to such a cosmic question depends strongly on who you ask. Who is qualified to answer such a question? It is best answered by those who study mathematics applied to the physical world. The utility and accuracy of that answer depend critically on the depth of understanding of the physics of the cosmic clock.

The English astronomer Edmond Halley (1656–1742) asked Isaac Newton (1643–1727) for the equation that describes the orbit of the planets. Halley was obviously interested in comets. Newton immediately answered, “an ellipse.” It is said that Halley was stunned by the response (Stillwell, 2010, p. 176), as this was what had been experimentally observed by Kepler (ca. 1619) and he knew that Newton must have some deeper insight. Both were eventually knighted.

---

3 MEP is a focused alternative to STEM.
4 I started around December 1970, fresh out of graduate school, and retired on December 5, 2002.
When Halley asked Newton to explain how he knew, Newton responded, “I calculated it.” But when challenged to show the calculation, Newton was unable to reproduce it. This open challenge eventually led to Newton’s grand treatise, *Philosophiae Naturalis Principia Mathematica* (July 5, 1687). It had a humble beginning, as a letter to Halley explaining how to calculate the orbits of the planets. To do this Newton needed mathematics, a tool he had mastered. It is widely accepted that Newton and Gottfried Leibniz invented calculus. But the early record shows that perhaps Bhāskara II (1114–1185 CE) had mastered the art well before Newton.\(^5\)

*Third,* the main goal of this book is to teach mathematics to motivated engineers, in a way that it can be understood, mastered, and remembered. How can this impossible goal be achieved? The answer is to fill in the gaps with *Who did what, and when?* Compared with the math, the historical record is easily mastered.

To be an expert in a field, one must know its history. This includes who the people were, what they did, and the credibility of their story. Do you believe the Pope or Galileo on the roles of the sun and the earth? The observables provided by science are clearly on Galileo’s side. Who were those first engineers? They are names we all know: Archimedes, Pythagoras, Leonardo da Vinci, Galileo, Newton, and so on. All of these individuals had mastered mathematics. This book presents the tools taught to every engineer. Rather than memorizing complex formulas, make the relationships “obvious” by mastering each simple underlying concept.

*Fourth,* when most educators look at this book, their immediate reactions are: *Each lecture is a topic we spend a week on (in our math/physics/engineering class) and You have too much material crammed into one semester.*

The first sentence is correct, the second is not. Tracking the students who have taken the course, looking at their grades, and interviewing them personally demonstrate that the material presented here is appropriate for one semester.\(^6\)

To write this book I had to master the language of mathematics. I had already mastered the language of engineering and a good part of physics. One of my secondary goals was to build this scientific Tower of Babel by unifying the terminology and removing the jargon.

**Acknowledgments**

I would like to acknowledge John Stillwell for his brilliant and constructive historical summary of mathematics as well as my close friend and long-time (40 years) colleague Steve Levinson, who somehow drew me into this project without my even knowing it. Next, my brilliant graduate student Sarah Robinson was constantly at my side, first repairing blunders in my first-draft homeworks and then grading these and the exams and tutoring the students. Without her, I would never have survived the first semester the material I taught. Her proofreading skills are amazing. Thank you, Sarah, for your infinite help. Without Kevin Pitts this work never could have been started, as he provided early funding when the project was a germ of an idea. Matt Ando’s (math) and Michael Stone’s (physics) encouragement was psychologically important in helping me think I might actually write a book. Finally, I would like to thank John D’Angelo for his highly critical comments, in response to my thousands of silly math questions. When it comes to the heavy hitting, John was always there to provide a brilliant explanation that I could easily translate into engineerese (matheering?) (i.e., engineer language).

My delightful friend Robert Fossum Emeritus Professor of Mathematics from the University of Illinois, kindly pointed out flawed mathematical terminology. James (Jamie) Hutchinson’s precise use of the English language dramatically raised the bar on my more than occasionally casual writing style. To each of you, thank you!

Finally I would like to thank my wife Sheau Feng Jeng, (Patricia Allen), for her unbelievable support and love. She delivered constant peace of mind, without which this project could never have been started, much less finished. Many others, including many students, played important roles, but given their large numbers, sadly they must remain anonymous.

–Jont Allen, Mahomet, IL, May, 12, 2019

\(^5\)[https://www-history.mcs.st-and.ac.uk/Projects/Pearce/Chapters/Ch8_5.html](https://www-history.mcs.st-and.ac.uk/Projects/Pearce/Chapters/Ch8_5.html)

\(^6\)[https://www.istem.illinois.edu/news/jont.allen.html](https://www.istem.illinois.edu/news/jont.allen.html)
Chapter 1

Introduction

Much of early mathematics before 1600 BCE involved the love of art and music, the sensations of light and sound. Our psychological senses of color and pitch are determined by the frequencies (i.e., wavelengths) of light and sound. The Chinese and later the Pythagoreans are well known for their early contributions to music theory. We are largely ignorant of exactly what Chinese scholars knew. The best record of early mathematics comes from Euclid, who lived in the third century BCE, after Pythagoras. Thus we can trace early mathematics back to the Pythagoreans in the sixth century (580–500 BCE), who focused on the Pythagorean theorem and early music theory.

Pythagoras strongly believed that “all is number,” meaning that every number, and every mathematical and physical concept, could be explained by integral (integer) relationships, mostly based on either ratios or the Pythagorean theorem. It is likely that his belief was based on Chinese mathematics from thousands of years earlier. It is also believed that his ideas about the importance of integers followed from music theory. The musical notes (pitches) obey natural integral ratio relationships based on the octave (a factor of two in frequency). The western 12-tone scale breaks the octave into 12 ratios called semitones. Today this has been rationalized to be the 12th root of 2, which is approximately equal to $18/17 \approx 1.06$ or 0.0833 [octave]. Our innate sense of frequency ratios comes from the physiology of the auditory organ (the cochlea), with a fixed distance along the organ of Corti, the sensory organ of the inner ear.

As acknowledged by Stillwell (2010, p. 16), the Pythagorean view is still relevant today:

> With the digital computer, digital audio, and digital video coding everything, at least approximately, into sequences of whole numbers, we are closer than ever to a world in which “all is number.”

1.1 Early science and mathematics

Although early Asian mathematics has been lost, it clearly defined its course for at least several millennia. The first recorded mathematics was from the Chinese (5000–1200 BCE) and the Egyptians (3300 BCE). Some of the best early records were left by the people of Mesopotamia (Iraq, 1800 BCE). While the first 5000 years of math are not well documented, the basic timeline is clear, as shown in Fig. 1.1.

Thanks to Euclid, and later Diophantus (ca. 250 CE), we have some basic (but vague) understanding of Chinese mathematics. For example, Euclid’s formula (Eq. 2.5.5.6, p. 50) provides a method for computing Pythagorean triplets, a formula believed to be due to the Chinese.²

Chinese bells and stringed musical instruments were exquisitely developed with tonal quality, as documented by ancient physical artifacts (Fletcher and Rossing, 2008). In fact this development was so rich that one must ask why the Chinese failed to initiate the Industrial Revolution. Specifically, why did European innovation eventually dominate when it was the Chinese who were responsible for such extensive early invention?

It could have been for internal politics, but perhaps our best insight into the scientific history of China may have come from an American chemist and scholar from Cambridge, England. Joseph Needham learned Chinese from a colleague, whom he later married, and ended up researching early Chinese science and technology for the U.S. government (Winchester, 2009).

According to Lin (1995), the Needham question is:

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¹See Fig. 2.8, p. 51.
²One might reasonably view Euclid’s role as that of a mathematical messenger.
CHAPTER 1. INTRODUCTION

Why did modern science, the mathematization of hypotheses about Nature, with all its implications for advanced technology, take its meteoric rise only in the West at the time of Galileo[, but] had not developed in Chinese civilization or Indian civilization?

As discussed by Lin (1995); Apte (2009), Needham cites the many developments in China:

*Gunpowder, the magnetic compass, and paper and printing, which Francis Bacon considered as the three most important inventions facilitating the West’s transformation from the Dark Ages to the modern world, were invented in China.*

Needham’s works attribute significant weight to the impact of Confucianism and Taoism on the pace of Chinese scientific discovery, and emphasize what it describes as the “diffusionist” approach of Chinese science as opposed to a perceived independent inventiveness in the western world. Needham held that the notion that the Chinese script had inhibited scientific thought was “grossly overrated.” (Grosswiler, 2004)

Lin (1995) focused on military applications, missing the importance of nonmilitary contributions. A large fraction of mathematics was developed to better understand the solar system, acoustics, musical instruments, and the theory of sound and light. Eventually the universe became a popular topic, as it still is today.

Regarding the Needham question, I suspect the answer is now clear. In the end, China withdrew from its several earlier expansions because of internal politics (Menzies, 2004, 2008).

### History of Mathematics to the 16th Century CE

<table>
<thead>
<tr>
<th>Century</th>
<th>Event</th>
</tr>
</thead>
<tbody>
<tr>
<td>20th</td>
<td>Chinese (primes; quadratic equation; Euclidean algorithm (GCD))</td>
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<tr>
<td>18th</td>
<td>Babylonians (Mesopotamia/Iraq) (quadratic solution)</td>
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<tr>
<td>6th</td>
<td>Thales of Miletus (first Greek geometry) (624)</td>
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<tr>
<td>5th</td>
<td>Pythagoras and the Pythagorean “tribe” (570)</td>
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<tr>
<td>4th</td>
<td>Euclid; Archimedes</td>
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<tr>
<td>3rd</td>
<td>Eratosthenes (276-194)</td>
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<tr>
<td>3rd</td>
<td>Diophantus (ca. 250)</td>
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<tr>
<td>4th</td>
<td>Library of Alexandria destroyed by fire (391)</td>
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<tr>
<td>7th</td>
<td>Brahmagupta (negative numbers; quadratic equation) (598-670)</td>
</tr>
<tr>
<td>10th</td>
<td>al-Khwarizmi (algebra) (830); Hasan Ibn al-Haytham (Alhazen) (965-1040)</td>
</tr>
<tr>
<td>14th</td>
<td>Bhaskara (calculus) (1114-1183)</td>
</tr>
<tr>
<td>15th</td>
<td>Leonardo da Vinci (1452-1519); Michelangelo (1475-1564); Copernicus (1473-1543)</td>
</tr>
<tr>
<td>16th</td>
<td>Tartaglia (cubic solution); Bombelli (1526-1572); Galileo Galilei (1564-1642)</td>
</tr>
</tbody>
</table>

### 1.1.1 The Pythagorean theorem

Thanks to Euclid’s *Elements* (written ca. 323 BCE) we have a historical record tracing the progress in geometry as established by the Pythagorean theorem, which states that for any right triangle having sides of lengths \((a, b, c) \in \mathbb{R}\) that are either positive real numbers or, more interesting, integers \(c > [a, b] \in \mathbb{N}\) such that \(a + b > c\),

\[
c^2 = a^2 + b^2. \tag{1.1.1.1}
\]

Early integer solutions were likely found by trial and error rather than by an algorithm.

If \(a, b, c\) are lengths, then \(a^2, b^2, c^2\) are each the area of a square. Equation 1.1.1.1 says that the area \(a^2\) plus the area \(b^2\) equals the area \(c^2\). Today a simple way to prove this is to compute the magnitude of the complex number \(c = a + bj\), which forces the right angle

\[
|c|^2 = (a + bj)(a - bj) = a^2 + b^2. \tag{1.1.1.2}
\]

However, complex arithmetic was not an option for the Greek mathematicians, since complex numbers and algebra had yet to be discovered.

Almost 700 years after Euclid’s *Elements*, the Library of Alexandria was destroyed by fire (391 CE), taking with it much of the accumulated Greek knowledge. As a result, one of the best technical records remaining is
1.1. EARLY SCIENCE AND MATHEMATICS

Euclid’s *Elements*, along with some sparse mathematics due to Archimedes (ca. 300 BCE) on geometrical series, the volume of a sphere, the area of a parabola, and elementary hydrostatics. In about 1572 a copy Diophantus’s *Arithmetic* was discovered by Bombelli in the Vatican library (Burton, 1985; Stillwell, 2010, p. 51). This book became an inspiration for Galileo, Descartes, Fermat, and Newton.

**Early number theory:** Well before Pythagoras, the Babylonians (ca. 1,800 BCE) had tables of triplets of integers \([a,b,c]\) that obey Eq. 1.1.1.1, such as \([3,4,5]\). However, the triplets from the Babylonians were larger numbers, the largest being \(a = 12,709,\text{and} \, c = 18,541\). A stone tablet (Plimpton-322) dating back to 1800 BCE was found with integers for \([a,c]\). Given such sets of two numbers, which determined a third positive integer \(b = 13,500\) such that \(b = \sqrt{c^2 - a^2}\), this table is more than convincing that the Babylonians were well aware of Pythagorean triplets (PTs), but less convincing that they had access to Euclid’s formula, a formula for PTs (Eq. 2.5.5.6, p. 50).

It seems likely that Euclid’s *Elements* was largely the source of the fruitful era of the Greek mathematician Diophantus (215–285) (see Fig. 1.1), who developed the field of discrete mathematics, now known as *Diophantine analysis*. The term means that the solution, not the equation, is integer. The work of Diophantus was followed by fundamental changes in mathematics, possibly leading to the development of algebra but at least including these discoveries:

1. Negative numbers
2. Quadratic equations (Brahmagupta, 7th century)
3. Algebra (al-Khwarizmi, 9th century)
4. Complex arithmetic (Bombelli, 15th century)

These discoveries overlapped with the European Middle Ages (also known as the Dark Ages). Although Europe went “dark,” presumably European intellectuals did not stop working during these many centuries.3

1.1.2 What is science?

Science is a process that quantifies hypotheses to build truths.4 It has evolved from early Greek philosophers, Plato and Aristotle, into a method that uses statistical tests to either validate or prove wrong the null hypothesis. Scientists use the term *null hypothesis* to describe the supposition that there is no difference between two intervention groups, or no effect of a treatment on some measured outcome. The measure of the likelihood that an outcome occurred by chance is called the *p*-value. From the *p*-value we can have some confidence that the null hypothesis is either true (the treatment causes no difference between two groups) or false (the probability *p* of a difference is greater than chance). The *p*-value is the present standard of scientific truth, but it is not iron clad and must be used with care. For example, not all experimental questions may be reduced to a single binary test. Does the sun rotate around the moon or around the earth? There is no test of this question, as it is nonsense. To even say that the earth rotates around the sun is, in some sense, nonsense because all the planets are involved in the orbital motion.

Yet science works quite well. We have learned many deep secrets regarding the universe over the last 5000 years.

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3It would be interesting to search the archives of the monasteries, where the records were kept, to determine exactly what happened during this religious blackout.

1.1.3 What is mathematics?

It seems strange when people complain that they “can’t learn math” but then claim to be good at languages. Before high school students tend to confuse arithmetic with math. One does not need to be good at arithmetic to be good at math (but it doesn’t hurt).

Math is a language with symbols taken from various languages—not so different from other languages. Today’s mathematics is a written language with an emphasis on symbols and glyphs, primarily Greek letters, obviously due to the popularity of Euclid’s Elements. The specific evolution of these symbols is interesting (Mazur, 2014). Each symbol is assigned a meaning appropriate for the problem being described. These symbols are then assembled to make sentences. In the Chinese language, the spoken and written versions are different across dialects. Similarly, mathematical sentences may be read out loud in any language (dialect), but the symbols (like Chinese characters) are universal.

Learning languages is an advanced social skill. However, the social outcomes of learning a language and learning math are very different. Learning a new language is fun because it opens doors to other cultures. Math is different due to the rigor of the grammar (rules of the language) as well as the way it is taught (e.g., not as a language). A third difference between math and language is that math evolved from physics and has important technical applications.

As with any language, the more mathematics you learn, the easier it is to understand, because mathematics is built from the bottom up. It’s a continuous set of concepts, much like the construction of a house. If you try to learn calculus and differential equations but skip simple number theory, the lessons will be more difficult to understand. You will end up memorizing instead of understanding, and as a result you will likely soon forget it. When you truly understand something, it can never be forgotten. A nice example is the solution to a quadratic equation: If you learn how to complete the square (see p. 64), you will never forget the quadratic formula.

Mathematical topics need to be learned in order, just as in the case of building the house. You can’t build a house if you don’t know about screws or cement (plaster). Likewise in mathematics, you can’t learn to integrate if you have failed to understand the difference between integers, complex numbers, polynomials, and their roots.

A short list of topics in mathematics includes numbers (N, Z, Q, ℤ, ℚ, ℂ), algebra, derivatives, antiderivatives (i.e., integration), differential equations, vectors and the spaces they define, matrices, matrix algebra, eigenvalues and eigenvectors, solutions of systems of equations, and matrix differential equations and their eigensolutions. Learning is about understanding, not memorizing.

The rules of mathematics are formally defined by algebra. For example, the sentence \( a = b \) means that the number \( a \) has the same value as the number \( b \). The sentence is read as “\( a \) equals \( b \).” The numbers are nouns and the equal sign says they are equivalent; it plays the role of a verb, or action symbol. Following the rules of algebra, this sentence may be rewritten as \( a - b = 0 \). Here the symbols for minus and equal indicate two types of actions (verbs).

Sentences can become arbitrarily complex, such as the definition of the integral of a function or a differential equation. But in each case, the mathematical sentence is written down, may be read out loud, has a well-defined meaning, and may be manipulated into equivalent forms following the rules of algebra and calculus. This language of mathematics is powerful, with deep consequences, first known as algorithms but eventually as theorems.

The writer of an equation should always translate (explicitly summarize the meaning of the expression), so the reader will not miss the main point. This is a simple matter of clear writing.

Just as math is a language, so language may be thought of as mathematics. To properly write correct English it is necessary to understand the construction of the sentence. It is important to identify the subject, verb, object, and various types of modifying phrases. For example, look up the interesting distinction between that and which. Thus, like math, language has rules. Most individuals use language that “sounds right,” but if you’re learning English as a second language, you must understand the rules, which are arguably easier to master than foreign speech sounds.

Context can be critical, and the most important context for mathematics is physics. Without a physical problem to solve, there can be no engineering mathematics. People needed to navigate the earth and weigh things, which required an understanding of gravity. Many questions about gravity were deep, such as Where is the center of the universe? But church dogma goes only so far. Mathematics along with a heavy dose of physics finally answered this huge question. Someone needed to perfect the telescope, put satellites into space, and view the cosmos. Without mathematics none of this would have happened.

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5“it looks like Greek to me.”
6https://en.oxforddictionaries.com/usage/that-or-which
7Actually this answer is simple: Ask the Pope and he will tell you. (I apologize for this inappropriate joke.)
1.2. MODERN MATHEMATICS

Figure 1.2: Timeline covering the two centuries from 1596CE to 1855CE, including the development of modern theories of analytic geometry, calculus, differential equations, and linear algebra. Newton was born about one year after Galileo died and thus was heavily influenced by his many discoveries. The vertical red lines indicate mentor-student relationships. Note the significant overlap among Newton, the Bernoullis, and Euler, a nucleation point for modern mathematics. Gauss had the advantage of input from Newton, Euler, d’Alembert, and Lagrange. Lagrange had a key role in the development of linear algebra. Likely Cauchy had a significant contemporary influence on Gauss as well. Finally, note that Fig. 1.1 ends with Bombelli while this figure begins with him. He famously discovered a copy of Diophantus’s book in the Vatican library. This was the same book that Fermat wrote in, with the margin too small to hold the “proof” of his “last theorem.”

1.1.4 Early physics as mathematics: Back to Pythagoras

We have established that math is the language of science. There is an additional answer to the question What is mathematics? The answer, the creation of algorithms and theorems, comes from studying its history, beginning with the earliest records. This chronological view starts, of course, with the study of numbers. First there is the taxonomy of numbers. It took thousands of years to realize that numbers are more than the counting numbers \( \mathbb{N} \), to create a symbol for nothing (i.e., zero), and to invent negative numbers. With the invention of the abacus, a memory aid for manipulating complex sets of real integers, one could do very detailed calculations. But this required the discovery of algorithms (procedures) to add, subtract, multiply (many additions of the same number), and divide (many subtractions of the same number), such as the Euclidean algorithm for the greatest common divisor (GCD). Eventually it became clear to the experts (early mathematicians) that there were natural rules to be discovered; thus books (e.g., Euclid’s *Elements*) were written to summarize this knowledge.

The role of mathematics is to summarize algorithms (i.e., sets of rules) and formalize an idea as a theorem. Pythagoras and his followers, the Pythagoreans, believed that there was a fundamental relationship between mathematics and the physical world. The Asian civilizations were the first to capitalize on the relationship between science and mathematics, to use mathematics to design things for profit. This may have been the beginning of capitalizing technology (i.e., engineering), based on the relationship between physics and math. This influenced commerce in many ways—map making, tools, implements of war (the wheel, gunpowder), art (music), water transport, sanitation, secure communication, food, and so on. Of course, the Chinese were among the first to master many of these technologies.

Why is Eq. 1.1.1.1 called a theorem? Theorems require a proof. What exactly needs to be proved? We do not need to prove that \((a, b, c)\) obeys this relationship, since this condition is observed. We do not need to prove that \(a^2\) is the area of a square, as this is the definition of an area. What needs to be proved is that the relationship \(c^2 = a^2 + b^2\) holds if, and only if, the angle between the two shorter sides is 90°. The Pythagorean theorem (Eq. 1.1.1.1) did not begin with Euclid or Pythagoras; rather they appreciated its importance and documented its proof.

In the end the Pythagoreans, who instilled fear in their neighborhood, were burned out, and murdered, likely the fate of mixing technology with politics:

> Whether the complete rule of number (integers) is wise remains to be seen. It is said that when the Pythagoreans tried to extend their influence into politics they met with popular resistance. Pythagoras fled, but he was murdered in nearby Mesopotamia in 497 BCE. Stillwell (2010, p. 16)

1.2 Modern mathematics

Modern mathematics (what we practice today) was born in the 15th and 16th centuries in the minds of Leonardo da Vinci, Bombelli, Galileo, Descartes, Fermat, and many others (Burton, 1985). Many of these early masters were, like the Pythagoreans, extremely secretive about how they solved problems. This soon changed with Galileo, Mersenne, Descartes, and Newton, which caused mathematics to blossom. Developments during this time may seem hectic and disconnected, but this is a wrong impression. The developments were dependent on new technologies, such as the telescope (optics) and more accurate time and frequency measurements, due to Galileo’s...
studies of the pendulum, and a better understanding of the relationship $f \lambda = c_0$, among frequency $f$, wavelength $\lambda$, and wave speed $c_0$.

### 1.2.1 Science meets mathematics

**Early studies of vision and hearing:** Since light and sound (music) played such a key role in the development of the early science, it was important to fully understand the mechanism of our perception of light and sound. There are many outstanding examples where physiology impacted mathematics. Leonardo da Vinci (1452–1519) is well known for his early studies of human anatomy, the knowledge of which was key when it came to drawing and painting the human form.

**Galileo:** In 1589 Galileo Galilei famously conceptualized an experiment in which he suggested dropping two different masses from the Leaning Tower of Pisa. He suggested that both must take the same time to hit the ground.

![Figure 1.3: Depiction of Galileo’s argument (from his unpublished book of 1638) as to why objects of different masses (i.e., weights) must fall with the same velocity, contrary to what Archimedes had proposed in about 250 BCE.](image)

Conceptually this is a mathematically sophisticated experiment, driven by a mathematical argument in which Galileo considered the two masses to be connected by an elastic cord (a spring) or rolling down a frictionless inclined plane (see Fig. 1.3). His studies resulted in the concept of conservation of energy, one of the cornerstones of physical theory since that time.

Being joined with an elastic cord, the masses become one. If the velocity were proportional to the mass, as Archimedes believed, the sum of the two masses would necessarily fall even faster. This results in a logical fallacy: How can two masses fall faster than either mass alone? This also violates the concept of conservation of energy, as the total energy of the two masses would be greater than that of the parts. In fact, Galileo’s argument may have been the first time that the principle of conservation of energy was clearly stated.

It seems likely that Galileo was attracted to this model of two masses connected by a spring because he was also interested in planetary motion, which consists of masses (sun, earth, moon) also mutually attracted by gravity (i.e., the spring).

Galileo performed related experiments on pendulums, where he varied the length $l$, mass $m$, and angle $\theta$ of the swing. By measuring the period he was able to formulate precise relationships between the variables. This experiment also measured the force exerted by gravity, so the experiments were related, but in very different ways. The pendulum served as the ideal clock, as it needed very little energy to keep it going, due to its very low friction (energy loss).

In a related experiment, Galileo measured the length of a day by counting the number of swings of the pendulum in 24 hours, measured precisely by the daily period of a star as it crossed the tip of a church steeple. The number of seconds in a day is $24 \cdot 60 \cdot 60 = 86,400 = 2^4 \cdot 3^2 \cdot 5^2 \cdot 7$ [s/day]. Since 86,400 is the product of the first three primes, it is highly composite and thus may be expressed in many equivalent ways. For example, the day can be divided evenly into 2, 3, or 5 parts and remain the same in terms of the number of seconds that transpire. Factoring the number of days in a year ($365 = 5 \cdot 73$) is not useful, since it may not be decomposed into many small parts.

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8The term period refers to the duration in seconds of a periodic function. For example, the periods of the moon and the sun are 28 days and one year, respectively.
primes. Galileo also extended work on the relationship between the wavelength and frequency of a sound wave in musical instruments. On top of these impressive accomplishments, Galileo greatly improved the telescope, which he needed for his observations of the planets and their moons.

Many of Galileo’s contributions resulted in new mathematics, leading to Newton’s discovery of the wave equation (1687), followed 60 years later by its one-dimensional general solution by d’Alembert (1747).

**Mersenne**: Marin Mersenne (1588–1648) contributed to our understanding of the relationship between the wavelength and the dimensions of musical instruments and is said to be the first to measure the speed of sound. At first Mersenne strongly rejected Galileo’s views, partially due to errors in Galileo’s reports of his results. But once Mersenne saw the significance of Galileo’s conclusion, he became Galileo’s strongest advocate, helping to spread the word (Palmerino, 1999).

Consider the development of an important theorem of nature. Are data more like bread or wine? The answer depends on the data. Galileo’s original experiments on pendulums and rolling masses down slopes were flawed by inaccurate data. This is likely because he didn’t have good clocks. But he soon solved that problem and the data became more accurate. We don’t know if Mersenne repeated Galileo’s experiments and then appreciated his theory, or if he communicated with Galileo. But the final resolution was that the early data were like bread (it rots), but when the experimental method was improved with a better clock, the corrected data were like wine (it improves with age). Galileo claimed that the time for the mass to drop a fixed distance was exactly proportional to the square of the time. This expression when integrated led to $F = ma$. One follows from the other. If the mass varies, then you get Newton’s second law of motion (Eq. 3.1.0.2, p. 62).

Mersenne was also a decent mathematician, inventing in 1644 the Mersenne prime (MP) $p_m$ of the form

$$p_m = 2^{p_k} - 1,$$

where $p_k (k < m)$ denotes the $k$th prime (see p. 24). As of December 2018, 51 MPs are known. The first MP is $3 = 2^2 - 1$, and the largest known prime is a MP. Note that $p_{31} = 2^{75} - 1$ is the MP of the MP $p_7$.

**Newton**: With the closure of Cambridge University due to the plague of 1665, Issac Newton returned home to Woolsthorpe-by-Colsterworth (95 miles north of London) to work by himself for over a year. It was during this solitary time that he did his most creative work.

Exploring our physiological senses required a scientific understanding of the physical processes of vision and hearing, first considered by Newton (1643–1727), but researched later in much greater detail, by Helmholtz (Stillwell, 2010, p. 261). While Newton (1642–1726) may be best known for his studies on light and gravity, he was the first to predict the speed of sound. However, his theory was in error by $\sqrt{c_p/c_v} = \sqrt{1.4} = 1.183$. This famous problem would not be corrected for 129 years, awaiting the formulation of thermodynamics and the equipartition theorem by Laplace in 1816 (Britannica, 2004).

Just 11 years prior to Newton’s 1687 *Principia*, there was a basic understanding that sound and light traveled at very different speeds, due to the experiments of Ole Rømer (Feynman, 1968, 2019, *Google for Feynman videos*).

*Ole Rømer first demonstrated in 1676 that light travels at a finite speed (as opposed to instantaneously) by studying the apparent motion of Jupiter’s moon Io. In 1865, James Clerk Maxwell proposed that light was an electromagnetic wave, and therefore traveled at the speed $c_v$ appearing in his theory of electromagnetism. (Wikipedia: Speed of Light, 2019)*

The idea behind Rømer’s discovery was that due to the large distance between Earth and Io, there was a difference between the period of the moon when Jupiter was closest to Earth and when it was farthest from Earth. This difference in distance caused a delay or advance in the observed eclipse of Io as it went behind Jupiter, delayed by the difference in time due to the difference in distance. This is like watching a video of a clock’s motion. When the video is delayed or slowed down, the time will be inaccurate (it will indicate an earlier time).

**The amazing Bernoulli family**: The first individual who seems to have openly recognized the importance of mathematics, enough to actually teach it, was Jacob Bernoulli (Fig. 1.4). Jacob worked on what is now viewed as the standard package of analytic “circular” (i.e., periodic) functions: $\sin(x)$, $\cos(x)$, $\exp(x)$, $\log(x)$. Eventually the full details were developed (for real variables) by Euler (p. 81).
Figure 1.4: Above left: Jacob (1655–1705) and right: Johann (1667–1748) Bernoulli, both painted by their portrait painter brother, Nicolaus. Below left: Leonhard Euler (1707–1783) and right: Jean le Rond d’Alembert (1717–1783). Euler was blind in his right eye, hence the left portrait view.
From Fig. 1.5 (p. 19) we may conclude that Jacob (1654–1705), the oldest brother, would have been strongly influenced by Newton.\textsuperscript{13} Newton would have been influenced by Fermat, Descartes, and Galileo, who died one year before Newton was born\textsuperscript{14} (White, 1999). Because the calendar was modified during Newton’s lifetime, his birth date is no longer given as Christmas 1642 (Stillwell, 2010, p. 175).

Jacob Bernoulli, like all successful mathematicians of the day, was largely self-taught. Yet Jacob was in a new category of mathematicians because he was an effective teacher. Jacob taught his sibling Johann, who then taught his sibling Daniel. But most important, Johann taught Euler, the most prolific (thus influential) of all mathematicians, including Gauss. This teaching resulted in an explosion of new ideas and understanding. It is most significant that all four mathematicians published their methods and findings. Much later, Jacob studied with students of Descartes\textsuperscript{15} (Stillwell, 2010, pp. 268–69).

**Euler:** Leonhard Euler’s mathematical talent went far beyond that of the Bernoulli family (Burton, 1985). Another special strength of Euler was his large number of publications. First he would master a topic, and then he would publish. Once the tools of mathematics were openly published, largely by Euler, mathematics grew exponentially.\textsuperscript{16} His papers continued to appear long after his death (Calinger, 2015). It is also interesting that Euler was a contemporary of Mozart (see Fig. 1.5).

**d’Alembert:** Another individual of that time who also published extensively was Jean la Rond d’Alembert (Fig. 1.4). Some of the most innovative ideas were first proposed by d’Alembert. Unfortunately, and perhaps somewhat unfairly, his rigor was criticized by Euler and later by Gauss (Stillwell, 2010).

**Gauss:** Figures 1.2 and 1.4 show timelines of the most famous mathematicians. This was one of the most creative times in mathematics. Carl Friedrich Gauss was born at the end of Euler’s long and productive life. I suspect that Gauss owed a great debt to Euler; surely he must have been a scholar of Euler. One of Gauss’s most important achievements may have been his contribution to solving the open question about the density of prime numbers and his use of least-squares.\textsuperscript{17}

**Cauchy:** Augustin-Louis Cauchy (1789–1857) was the son of a well-to-do family but had the misfortune of being born during the time of of the French Revolution, which perhaps originated with the Seven Years’ War, around 1756. Today the French still celebrate Bastille Day (July 14, 1789), which is viewed as a celebration of the revolution. The French Revolution left Cauchy with a lifelong scorn for French politics that deeply influenced his life. But Cauchy had an unmatched intellect for mathematics. His most obvious achievement was complex analysis, for which he proved many key theorems.

<table>
<thead>
<tr>
<th>1525</th>
<th>1564</th>
<th>1650</th>
<th>1750</th>
<th>1800</th>
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<tr>
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<td>Euler</td>
<td>Daniel</td>
<td>Gauss</td>
<td>Helmholtz</td>
<td></td>
</tr>
<tr>
<td>Galileo</td>
<td>Jacob</td>
<td>d’Alembert</td>
<td></td>
<td>Rayleigh</td>
<td>Kirchhoff</td>
<td></td>
</tr>
<tr>
<td>Fermat</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Figure 1.5:** Timeline for the 16th through 20th centuries covering Bombelli to Einstein. As noted in the caption for Fig. 1.2, it seems likely that Bombelli’s discovery of Diophantus’s book *Arithmetic* in the Vatican library triggered many of the ideas presented by Galileo, Descartes, and Fermat, followed by others (i.e., Newton). Thus Bombelli’s discovery might be considered a magic moment in mathematics. The vertical red lines indicate mentor–student relationships. For orientation Mozart is indicated along the bottom and Napoleon at the top. Napoleon hired Fourier, Lagrange, and Laplace to help with his many bloody military campaigns. See Figs. 1.1, 1.2, and 3.1 for additional timelines.

\textsuperscript{13}For a similar timeline see https://www.famousscientists.org/joseph-louis-lagrange/

\textsuperscript{14}https://www-history.mcs.st-andrews.ac.uk/Biographies/Newton.html

\textsuperscript{15}It seems clear that Descartes was also a teacher.

\textsuperscript{16}There are at least three useful exponential scales: factors of 2, factors of \( e \approx 2.7 \), and factors of 10. The octave and decibel use factors of 2 (6 [dB]) and 10 (20 [dB]). Information theory uses factors of 2 (1 [bit]), 4 (2 [bits]). Circuit theory uses all three scales.

\textsuperscript{17}https://www-history.mcs.st-andrews.ac.uk/Biographies/Gauss.html
Helmholtz: Hermann von Helmholtz (1821–1894) was educated and experienced as a military surgeon. He also mastered classical music, acoustics, physiology, vision, hearing (Helmholtz, 1863b), and, most important of all, mathematics. Gustav Kirchhoff frequently expanded on Helmholtz’s contributions. It is reported that Lord Rayleigh learned German so he could read Helmholtz’s great works.

Helmholtz’s theories of music and perception of sound are fundamental scientific contributions (Helmholtz, 1863a). His best known mathematical contribution is known as the fundamental theorem of vector calculus, or simply “Helmholtz theorem” (p. 207).

The history during this time is complex. For example, in 1850 Lord Kelvin wrote a letter to George Stokes, suggesting that Stokes try to prove what is today known as “Stokes’s theorem.” As a result, Stokes posted a reward (Smith’s Prize), searching for a proof of “Lord Kelvin’s theorem,” which was finally achieved by Hermann Hankel (1839–1873). Many new concepts were being proved and appreciated over this productive period. Maxwell had published his famous equations and reformatted in modern vector notation by Oliver Heaviside, J. Willard Gibbs, and Heinrich Hertz. Fig. 1.5 should put to rest the idea that one’s best work is done in the early years. Many of these scientists were fully productive into old age. Those who were not died early due to poor health or accidents.

Lord Kelvin: Lord Kelvin (William Thompson, 1824–1907) was one of the first true engineer-scientists, equally acknowledged as a mathematical physicist, well known for his interdisciplinary research, and knighted by Queen Victoria in 1866. Lord Kelvin coined the term thermodynamics, a science more fully developed by Maxwell (the same Maxwell of electrodynamics).

James Clerk Maxwell (1831-1879) In 1869 a Cambridge senate committee was formed to create the Cambridge Physics Laboratory and “the founding of a special Professorship.” The Chancellor of Cambridge was the seventh Duke of Devonshire and a distant relative of Henry Cavendish, his family name. Thus the new laboratory became known as the Cavendish.

There was, naturally, much speculation about the choice of the new Professor of Experimental Physics. [Lord] Kelvin was the most likely candidate, but on being approached in private, refused in order to stay in Glasgow. Another likely candidate was Lord Rayleigh, a brilliant mathematician and physicist who had left Cambridge to work in his private laboratory at his country seat in Essex. When the appointment was eventually announced, the reaction was, if anything, one of disappointment. The new Professor, James Clerk Maxwell, was relatively unknown.

He was a much respected mathematician, but he had not since made any great name for himself – his major and astounding books on Electricity and Kinetic Theory had yet to be published. Moreover, the six years before his appointment had been spent in isolation at his Scottish home. His appointment was announced on March 8th 1871, and in spite of the initial disappointment, his inaugural lecture was looked forward to by his likely students as much as by the rest of the Cambridge scientists.

When, a few days later, Maxwell began his first course with a lecture on Heat, his students had the delight of seeing the lecture room packed with their tutors, lecturers, professors and all the important personages of the University. Thinking that this was his first public appearance they sat, in their formal academic dress, while Maxwell, “with a perceptible twinkle in his eye,” gravely expounded the difference between Fahrenheit and Centigrade, and the principle of the air thermometer.

It was felt afterwards that Maxwell had done it on purpose, perhaps out of modesty, perhaps out of his later well-known sense of humour, or perhaps because he knew of the still considerable opposition his new laboratory had to face. As he had written to his friend Lord Rayleigh, “if we succeed too well, and corrupt the minds of youth till they observe vibrations and deflections and become Senior Ops. instead of Wranglers, we may bring the whole University and all the parents about our ears.”

Lord Rayleigh (William Strutt): Lord Rayleigh (1842–1919) wrote a classic text (1896) that is widely read even today by those who study acoustics. In 1904 he received the Nobel Prize in Physics for his investigations of the densities of the most important gases and for his discovery of argon in connection with these studies.

---

19Lord Kelvin was one of a half dozen interdisciplinary mathematical physicists, all working about the same time, who made a fundamental change in our scientific understanding. Others include Helmholtz, Stokes, Green, Heaviside, Rayleigh, and Maxwell.
20Thermodynamics is another topic that warrants an analysis along historical lines (Kuhn, 1978).
21https://www.phy.cam.ac.uk/history/years/firstten
1.2. MODERN MATHEMATICS

1.2.2 Three Streams from the Pythagorean theorem

From the outset of his presentation, Stillwell (2010, p. 1) defines “three great streams of mathematical thought: Numbers, Geometry and Infinity” that flow from the Pythagorean theorem, as summarized in Fig. 1.6. This is a useful concept, based on reasoning not as obvious as one might think. Many factors are in play here. One of these is the strongly held opinion of Pythagoras that all mathematics should be based on integers. The rest are tied up in the long, necessarily complex history of mathematics, as best summarized by the fundamental theorems (see Sec. 2.3.1, p. 35), each of which is discussed in detail in a later chapter.

1. Numbers
   • $\mathbb{N}$ counting numbers, $\mathbb{Q}$ rationals, $\mathbb{P}$ primes (6th century BCE)
   • $\mathbb{Z}$ common integers, $\mathbb{I}$ irrationals (5th century BCE)
   • zero $\in \mathbb{Z}$ (7th century CE)

2. Geometry (e.g., lines, circles, spheres, toroids, other shapes)
   • Composition of polynomials (Descartes, Fermat),
   • Euclid’s geometry and algebra $\Rightarrow$ analytic geometry (17th century CE)
   • Fundamental theorem of algebra (18th century CE)

3. Infinity ($\infty \to$ sets)
   • Taylor series, functions, calculus (Newton, Leibniz) (17th and 18th century CE)
   • $\mathbb{R}$ real, $\mathbb{C}$ complex (19th century CE)
   • Set theory (20th century CE)

Figure 1.6: Three streams that follow from the Pythagorean theorem: numbers, geometry, and infinity.

As shown in Fig. 1.6, Stillwell’s concept of three streams, following from the Pythagorean theorem, is the organizing principle behind this book.

Ch. 1: The Introduction is a historical survey of pre-college mathematical physics, presented in terms of the three main Pythagorean streams (stream 1–stream 3), leading to the book’s five chapters. Stream 3 is split into 3A and 3B.

Ch. 2: Number systems (p. 23) Some important ideas from number theory, starting with prime numbers, complex numbers, vectors, and matrices. Five classic number theory problems are discussed: the Euclidean algorithm (GCD), continued fractions (CFA), Euclid’s formula, Pell’s equation, and the Fibonacci difference equation. The general solution of these problems leads to the concept of the eigenfunction analysis, which is introduced in Sec. 2.5.7 (p. 53).

Ch. 3: Algebraic equations (p. 61) Algebra and its development, as we know it today. The chapter presents the theory of real and complex equations and functions of real and complex variables. Newton’s method for finding complex roots of polynomials, poles vs. zeros, and the Gauss-Lucas theorem (bounds on the root locations of the derivative of a polynomial). Complex impedance $Z(s)$ of complex frequency $s = \sigma + \omega j$ is covered with some care, developing the topic is needed for engineering mathematics.

While the algebra of real and complex functions is identical, the calculus is fundamentally different. This leads to the concepts of complex analytic functions, complex Taylor series, and the Cauchy-Riemann conditions. These ideas are fundamental when dealing with impedance functions that describe the linear relationships between force and flow in the complex frequency domain (i.e., complex impedance).

Ch. 4: Scalar calculus (Stream 3A) (p. 133) Ordinary differential equations and integral theorems of simple physical systems (mass-springs, inductors-capacitors, heat dynamics), solutions to scalar differential equations that have constant coefficients, colorized mappings of complex analytic functions, multivalued functions, Cauchy’s theorems, and inverse Laplace transforms.

Ch. 5: Vector calculus (Stream 3B) (p. 169) Vector partial differential equations, as well as gradient, divergence, and curl differential operators, Stokes’s and Green’s theorems, and Maxwell’s equations.
Chapter 2

Stream 1: Number Systems

Number theory (the study of numbers) was a starting point for many key ideas. For example, in Euclid’s geometrical constructions the Pythagorean theorem for real $[a, b, c]$ was accepted as true, but the emphasis in the early analysis was on integer constructions, such as Euclid’s formula for Pythagorean triplets (Eq. 2.5.5.6, p. 50).

As we shall see, the derivation of the formula for Pythagorean triplets is the first of a rich body of mathematical constructions—such as solutions of Pell’s equation (p. 52), and recursive difference equations, such as solutions of the Fibonacci recursion formula $f_{n+1} = f_n + f_{n-1}$ (see p. 53)—that goes back at least to the Chinese (2000 BCE). These are early pre-limit forms of calculus, best analyzed using an eigenfunction (e.g., eigenmatrix) expansion, a geometrical concept from linear algebra, as an orthogonal set of normalized unit-length vectors (see Appendix B.3, p. 237).

It is hard to imagine that anyone who uses an abacus would not appreciate the concept of zero and negative numbers. It does not take much imagination to go from counting numbers $\mathbb{N}$ to the set of all integers $\mathbb{Z}$ including zero. On an abacus, subtraction is obviously the inverse of addition. Subtraction to obtain zero abacus beads is no different than subtraction from zero, which gives negative beads. To assume that the Romans, who first developed counting sticks, or the Chinese, who then deployed the concept using beads, did not understand negative numbers is impossible.

However, understanding the concept of zero (and negative numbers) is not the same as having a symbolic notation. The Roman number system has no such symbols. The first recorded use of a symbol for zero is said to be by Brahmagupta in 628 CE.3

However, this is likely wrong, given the notation developed by the Mayan civilization that existed from 2000 BCE to 900 CE.3 There is speculation that the Mayans cut down so much of the Amazon jungle that it eventually resulted in global warming, possibly leading to their demise.

The definition of zero depends on the concept of subtraction, which formally requires the creation of algebra (ca. 830 CE; see Fig. 1.1, p. 13). But apparently it took more than 600 years from the time Roman numerals were put into use, without any symbol for zero, to the time the symbol for zero is first documented. Likely this delay was more about the political situation, such as government rulings, than mathematics.

The concept that caused much more difficulty was $\infty$, or infinity, first proposed by Bernhard Riemann in 1851 with the development of the extended plane, which mapped the plane to a sphere (see Fig. 3.15, p. 127). His construction made it clear that the point at $\infty$ is simply another point on the open complex plane, since rotating the sphere (extended plane) moves the point at $\infty$ to a finite point on the plane, thereby closing the complex plane.

2.1 The taxonomy of numbers: $\mathbb{P}, \mathbb{N}, \mathbb{Z}, \mathbb{Q}, \mathbb{F}, \mathbb{I}, \mathbb{R}, \mathbb{C}$

Once symbols for zero and negative numbers were accepted, progress could be made. To fully understand numbers, a transparent notation was required. First one must identify the different classes (genus) of numbers, providing a notation that defines each of these classes along with their relationships. It is logical to start with the most basic counting numbers, which we indicate with the double-bold symbol $\mathbb{N}$. For easy access, double-bold symbols and set-theory symbols—such as, $\{\cdot\}, \cup, \cap, \in, \notin, \perp$, and so on—are summarized in Appendix A.

---

1 Heisenberg, an inventor of the matrix algebra form of quantum mechanics, learned mathematics by studying Pell’s equation (p. 52) by eigenvector and recursive analysis methods. https://www.aip.org/history-programs/niels-bohr-library/oral-histories/4661-1


3 https://www.storyofmathematics.com/mayan.html
Counting numbers \( \mathbb{N} \): These are known as the natural numbers \( \mathbb{N} = \{1, 2, 3, \ldots \} \) and denoted by the double-bold symbol \( \mathbb{N} \). For clarity we shall refer to the natural numbers as counting numbers, since natural, which means integer, is vague. The mathematical sentence “\( 2 \in \mathbb{N} \)” is read as “\( 2 \) is a member of the set of counting numbers.” The word set is defined as the collection of any objects that share a specific property. Typically the set may be defined either as a sentence or by example.

Primes \( \mathbb{P} \): A number is prime (\( \pi_n \in \mathbb{P} \)) if its only factors are 1 and itself. The set of primes \( \mathbb{P} \) is a subset of the counting numbers (\( \mathbb{P} \subseteq \mathbb{N} \)). A somewhat amazing fact, well known to the earliest mathematicians, is that every integer may be written as a unique product of primes. A second key idea is that the density of primes \( \rho_p(N) \sim 1 / \log(N) \); that is, \( \rho_p(N) \) is inversely proportional to the log of \( N \), an observation first quantified by Gauss (Goldstein, 1973). A third is that there is a prime between every integer \( N \geq 2 \) and \( 2N \).

We shall use the convenient notation \( \pi_n \) for the prime numbers, indexed by \( n \in \mathbb{N} \). The first 12 primes are \( \{n|1 \leq n \leq 12\} = \{\pi_n|2, 3, 5, 7, 11, 13, 17, 19, 23, 29, 31, 37\} \). Since \( 4 = 2^2 \) and \( 6 = 2 \cdot 3 \) may be factored, \( 4, 6 \not\in \mathbb{P} \) (read as “\( 4 \) and \( 6 \) are not in the set of primes”). Given this definition, multiples of a prime—that is, \( [2, 3, 4, 5, \ldots] \cdot \pi_k \) of any prime \( \pi_k \)—cannot be prime. It follows that all primes except 2 must be odd and every integer \( N \) is unique in its prime factorization.

Exercise #1
Write the first 10 to 20 integers in prime-factored form.

Solution: \( 1, 2, 3, 2^2, 5, 2 \cdot 3, 7, 2^3, 2 \cdot 5, 11, 3 \cdot 2^2, 13, 2 \cdot 7, 3 \cdot 5, 2^4, 17, 2 \cdot 3^2, 19, 2^2 \cdot 5 \)

Exercise #2
Write the integers 2 to 20 in terms of \( \pi_n \). Here is a table to assist you:

<table>
<thead>
<tr>
<th>( n )</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>\ldots</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \pi_n )</td>
<td>2</td>
<td>( \pi_1 )</td>
<td>3</td>
<td>( \pi_2 )</td>
<td>5</td>
<td>( \pi_3 )</td>
<td>7</td>
<td>( \pi_4 )</td>
<td>11</td>
<td>( \pi_5 )</td>
<td>13</td>
<td>( \pi_6 )</td>
</tr>
</tbody>
</table>

Solution:

<table>
<thead>
<tr>
<th>( n )</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
<th>13</th>
<th>14</th>
<th>\ldots</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \Pi \pi_n )</td>
<td>2</td>
<td>( \pi_1 )</td>
<td>3</td>
<td>( \pi_1 \pi_2 )</td>
<td>5</td>
<td>( \pi_1 \pi_3 )</td>
<td>7</td>
<td>( \pi_1 \pi_4 )</td>
<td>11</td>
<td>( \pi_1 \pi_5 )</td>
<td>13</td>
<td>( \pi_1 \pi_6 )</td>
<td>17</td>
<td>( \pi_1 \pi_7 )</td>
<td>19</td>
</tr>
</tbody>
</table>

Coprimes are two relatively prime numbers that have no common (i.e, prime) factors. For example, \( 21 = 3 \cdot 7 \) and \( 10 = 2 \cdot 5 \) are coprime, whereas \( 4 = 2 \cdot 2 \) and \( 6 = 2 \cdot 3 \), which have 2 as a common factor, are not. By definition all unique pairs of primes are coprime. We shall use the notation \( m \perp n \) to indicate that \( m \) and \( n \) are coprime. The ratio of two coprimes is reduced, as it has no factors to cancel. The ratio of two numbers that are not coprime may always be reduced by canceling the common factors. This is called the reduced form, or an irreducible fraction. When we do numerical work, for computational accuracy it is always beneficial to work with coprimes. Generalizing this idea, we could define tripri
times as three numbers with no common factor, such as \( \{\pi_3, \pi_9, \pi_2\} \).

The fundamental theorem of arithmetic states that each integer may be uniquely expressed as a unique product of primes. The prime number theorem estimates the mean density of primes over \( N \).

Integers \( \mathbb{Z} \): The integers include positive and negative counting numbers and zero. Notionally we might indicate this using set notation as \( \mathbb{Z} = \mathbb{N} \cup \{0\} \cup \mathbb{N} \). We read this as: “The integers are in the set composed of the negative natural numbers \( -\mathbb{N} \), zero, and \( \mathbb{N} \).”

Rational numbers \( \mathbb{Q} \): These are defined as numbers formed from the ratio of two integers. Given two numbers \( n, d \in \mathbb{N} \), we have \( n/d \in \mathbb{Q} \). Since \( d \) may be 1, it follows that the rationals include the counting numbers as a subset. For example, the rational number \( 3/1 \in \mathbb{N} \).

The main utility of rational numbers is that they can efficiently approximate any number on the real line, to any precision. For example, the rational approximation \( \pi \approx 22/7 \) has a relative error of \( \approx 0.04\% \) (see p. 48).

Fractional number \( \mathbb{F} \): A fractional number \( \mathbb{F} \) is defined as the ratio of signed coprimes. If \( n, d \in \pm \mathbb{P} \), then \( n/d \in \mathbb{F} \). Given this definition, \( \mathbb{F} \subseteq \mathbb{Q} = \mathbb{Z} \cup \mathbb{F} \). Because of the powerful approximating power of rational numbers, the fractional set \( \mathbb{F} \) has special utility. For example, \( \pi \approx 22/7, 1/\pi \approx 7/22 \) (to 0.04\%), \( e \approx 19/7 \) to 0.15\%, and \( \sqrt{2} \approx 7/5 \) to 1\%.
2.1. THE TAXONOMY OF NUMBERS: \( \mathbb{P}, \mathbb{N}, \mathbb{Z}, \mathbb{Q}, \mathbb{F}, \mathbb{I}, \mathbb{R}, \mathbb{C} \)

**Irrational numbers** \( \mathbb{I} \): Every real number that is not rational is irrational (\( \mathbb{Q} \perp \mathbb{I} \)). Irrational numbers include \( \pi, e \), and the square roots of primes. These are decimal numbers that never repeat, thus requiring infinite precision in their representation. Such numbers cannot be represented on a computer, as they would require an infinite number of bits (precision).

The rationals \( \mathbb{Q} \) and irrationals \( \mathbb{I} \) split the reals (\( \mathbb{R} = \mathbb{Q} \cup \mathbb{I} \), \( \mathbb{Q} \perp \mathbb{I} \)); thus each is a subset of the reals (\( \mathbb{Q} \subset \mathbb{I} \), \( \mathbb{I} \subset \mathbb{R} \)). This relationship is analogous to that of the integers \( \mathbb{Z} \) and fractionals \( \mathbb{Q} \), which split the rationals (\( \mathbb{Q} = \mathbb{Z} \cup \mathbb{Q}, \mathbb{Q} \subset \mathbb{Z} \)) (thus each is a subset of the rationals (\( \mathbb{Z} \subset \mathbb{Q} \), \( \mathbb{F} \subset \mathbb{Q} \))).

Irrational numbers \( \mathbb{I} \) were famously problematic for the Pythagoreans, who incorrectly theorized that all numbers were rational. Like \( \infty \), irrational numbers required mastering a new and difficult concept before they could be understood: It was essential to understand the factorization of counting numbers into primes (i.e., the fundamental theorem of arithmetic) before the concept of irrationals could be sorted out. Irrational numbers could only be understood once limits were mastered.

As shown in Fig. 2.6 (p. 49), fractionals can approximate any irrational number with arbitrary accuracy. Integers are also important, but for a very different reason. All numerical computing today is done with \( \mathbb{Q} = \mathbb{F} \cup \mathbb{Z} \). Indexing uses integers \( \mathbb{Z} \), while the rest of computing (flow dynamics, differential equations, etc.) is done with fractionals \( \mathbb{F} \) (i.e., IEEE-754). Computer scientists are trained on these topics, and computer engineers need to be at least conversant with them.

**Real numbers** \( \mathbb{R} \): Reals are the union of rational and irrational numbers–namely, \( \mathbb{R} = \mathbb{Q} \cup \mathbb{I} = \mathbb{Z} \cup \mathbb{F} \cup \mathbb{I} \). Lengths in Euclidean geometry are reals. Many people assume that IEEE 754 floating-point numbers (ca. 1985) are real (i.e., \( \in \mathbb{R} \)). In fact, they are rational (\( \mathbb{Q} = \{ \mathbb{F} \cup \mathbb{Z} \} \)) approximations to real numbers, designed to have a very large dynamic range. The hallmark of fractional numbers (\( \mathbb{F} \)) is their power in making highly accurate approximations of any real number.

Using Euclid’s compass and ruler methods, one can make the length of a line proportionally shorter or longer or (approximately) the same. A line may be made be twice as long, or an angle can be bisected. However, the concept of an integer length in Euclid’s geometry was not defined.\(^4\) Nor can one construct an imaginary or complex line, as all lines are assumed to have real lengths. The development of analytic geometry was an analytic extension of Euclid’s simple (but important) geometrical methods.

Real numbers were first fully accepted only after set theory was developed by Georg Cantor in 1874 (Stillwell, 2010, p. 461). At first blush, this seems amazing, given how widely accepted real numbers are today. In some sense they were accepted by the Greeks as lengths of real lines.

**Complex numbers** \( \mathbb{C} \): Complex numbers are best defined as ordered pairs of real numbers.\(^5\) For example, if \( a, b \in \mathbb{R} \) and \( j = -1 = \sqrt{-1} \), then \( c = a + bj \in \mathbb{C} \). The word complex, as used here, does not mean that the numbers are complicated or difficult. They are also known as imaginary numbers, but this does not mean the numbers disappear. Complex numbers are quite special in engineering mathematics as roots of polynomials. The most obvious example is the quadratic formula for the roots of polynomials of degree 2 that have coefficients \( \in \mathbb{C} \). All real numbers have a natural order on the real line. Complex numbers do not have a natural order. For example, \( j > 1 \) makes no sense.

Today the common way to write a complex number is using the notation \( z = a + bj \in \mathbb{C} \), where \( a, b \in \mathbb{R} \). Here \( 1j = \sqrt{-1} \). We also define \( 1j = -1j \) to account for the two possible signs of the square root. Accordingly \( 1j^2 = 1j^2 = -1 \).

Cartesian multiplication of complex numbers follows the basic rules of real algebra—for example, the rules for multiplying two monomials and polynomials. Multiplication of two first-degree polynomials (i.e., monomials) gives

\[
(a + bx)(c + dx) = ac + (ad + bc)x + bdx^2.
\]

If we substitute \( 1j \) for \( x \) and use the definition \( 1j^2 = -1 \), we obtain the Cartesian product of two complex numbers:

\[
(a + bj)(c + dj) = ac - bd + (ad + bc)j.
\]

Thus multiplication and division of complex numbers obey the usual rules of algebra.

However, there is a critical extension: Cartesian multiplication holds only when the angles sum to less than \( \pm \pi \)—namely, the range of the complex plane. When the angles add to more that \( \pm \pi \), one must use polar coordinates, where the angles add for angles beyond \( \pm \pi \) (Boas, 1987, p. 8). This is particularly striking for the \( LT \) transform of a delay (see Appendix C.3, p. 242).

\(^4\) As best I know.

\(^5\) A polynomial \( a + bx \) and a 2-vector \( [a, b]^T \) are also examples of ordered pairs.
Complex numbers can be challenging and may provide unexpected results. For example, it is not obvious that \( \sqrt{3 + 4j} = \pm(2 + j) \).

**Exercise #3**
Verify that \( \sqrt{3 + 4j} = \pm(2 + j) \).

**Solution:** Squaring the left side gives \( 3 + 4j \). Squaring the right side gives \((2 + j)^2 = 4 - j^2 + 4j = 3 + 4j \). Thus the two are equal. □

**Exercise #4**
What is special about the above example?

**Solution:** Note this is a \( \{3, 4, 5 \} \) triangle. Can you find another example like this one? Namely, how does one find integers that obey Eq. 1.1.1.1 (p. 12)? □

An alternative to Cartesian multiplication of complex numbers is to work in polar coordinates. The polar form of the complex number \( z = a + bj \) is written in terms of its magnitude \( \rho = \sqrt{a^2 + b^2} \) and angle \( \theta = \angle z = \tan^{-1} z = \arctan z \) as

\[
z = \rho e^{i\theta} = \rho(\cos \theta + j \sin \theta).
\]

From the definition of the complex natural log function, we have

\[
\ln z = \ln \rho e^{i\theta} = \ln \rho + i\theta,
\]

which is important, even critical, in engineering calculations. When the angles of two complex numbers are greater than \( \pm\pi \), one must use polar coordinates. It follows that as computing the phase, this is much different than the single- and double-argument \( \angle \theta = \arctan(z) \) function.

The polar representation makes clear the utility of a complex number: Its magnitude scales while its angle \( \Theta \) rotates. The property of scaling and rotating is what makes complex numbers useful in engineering calculations. This is especially obvious when dealing with impedances, which have complex roots with very special properties, as discussed on page 142.

**Matrix representation:** An alternative way to represent complex numbers is in terms of \( 2 \times 2 \) matrices. This relationship is defined by the mapping from a complex number to a \( 2 \times 2 \) matrix:

\[
a + bj \leftrightarrow \begin{bmatrix} a & -b \\ b & a \end{bmatrix}, \quad 1 \leftrightarrow \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix}, \quad 1j \leftrightarrow \begin{bmatrix} 0 & -1 \\ 1 & 0 \end{bmatrix}, \quad e^{i\theta} \leftrightarrow \begin{bmatrix} \cos(\theta) & -\sin(\theta) \\ \sin(\theta) & \cos(\theta) \end{bmatrix}.
\]

The conjugate of \( a + bj \) is then defined as \( a - bj \leftrightarrow \begin{bmatrix} a & b \\ -b & a \end{bmatrix} \). By taking the inverse of the \( 2 \times 2 \) matrix (assuming \(|a + bj| \neq 0\)), we can define the ratio of one complex number by another. Until you try out this representation, it may not seem obvious or even possible.

This representation proves that \( 1j \) is not necessary when defining a complex number. What \( 1j \) can do is to conceptually simplify the algebra. It is worth your time to become familiar with the matrix representation, to clarify any possible confusions you might have about multiplication and division of complex numbers. This matrix representation can save you time, heartache, and messy algebra. Once you have learned how to multiply two matrices, it’s a lot simpler than doing the complex algebra. In many cases we will leave the results of our analysis in matrix form, to avoid the algebra altogether.\(^6\) Thus both representations are important. More on this topic may be found on page 23.

**History of complex numbers:** It is notable that complex numbers were not accepted until 1851 even though Bombelli introduced them in the 16th century. One might have thought that the solution of the quadratic, known to the Chinese, would have settled this question. It seems that complex integers (Gaussian integers) were accepted before nonintegral complex numbers. Perhaps this was because real numbers (\( \mathbb{R} \)) were not accepted (i.e., proved to exist, thus mathematically defined) until the development of real analysis in the late 19th century, thus providing a proper definition of the real numbers.

\(^6\) Sometimes we let the computer do the final algebra, numerically, as \( 2 \times 2 \) matrix multiplications.
2.1. THE TAXONOMY OF NUMBERS: P, N, Z, Q, F, I, R, C

Exercise #5
Using Matlab/Octave, verify that

\[
\frac{a + bj}{c + dj} = \frac{ac + bd + (bc - ad)j}{c^2 + d^2} \quad \leftrightarrow \quad \left[ \begin{array}{cc} a & -b \\ b & a \end{array} \right] \left[ \begin{array}{cc} c & -d \\ d & c \end{array} \right]^{-1} = \left[ \begin{array}{cc} a & -b \\ b & a \end{array} \right] \left[ \begin{array}{cc} c & d \\ -d & c \end{array} \right] \frac{1}{c^2 + d^2}.
\]

Solution: The typical solution may use numerical examples. A better solution is to use a symbolic code:

\[
\text{syms a b c d A B}
A = \left[ \begin{array}{cc} a & -b \\ b & a \end{array} \right];
B = \left[ \begin{array}{cc} c & -d \\ d & c \end{array} \right];
C = A \cdot \text{inv}(B)
\]

2.1.1 Numerical taxonomy

A simplified taxonomy of numbers is given by the mathematical sentence

\[
\pi_k, P \subset N \subset Z \subset Q \subset R \subset C.
\]

This sentence says:

1. Every prime number \(\pi_k\) is in the set of primes \(P\),
2. which is a subset of the set of counting numbers \(N\),
3. which is a subset of the set of integers \(Z = -N \cup \{0\} \cup N\),
4. which is a subset of the set of rationals \(Q\) (ratios of signed counting numbers \(\pm N\)),
5. which is a subset of the set of reals \(R\),
6. which is a subset of the set of complex numbers \(C\).

The rationals \(Q\) may be further decomposed into the fractionals \(F\) and the integers \(Z\) \((Q = F \cup Z)\), and the reals \(R\) into the rationals \(Q\) and the irrationals \(I\) \((R = Q \cup I)\). This classification nicely defines all the numbers (scalars) used in engineering and physics.

The taxonomy structure may be summarized with the single compact sentence, starting with the prime numbers \(\pi_k\) and ending with complex numbers \(C\):

\[
\pi_k \in P \subset N \subset Z \subset (Z \cup F = Q) \subset (Q \cup I = R) \subset C.
\]

As discussed in Appendix A (p. 221), all numbers may be viewed as complex; that is, every real number is complex if we take the imaginary part to be zero (Boas, 1987). For example, \(2 \in P \subset C\). Likewise, every purely imaginary number (e.g., \(0 + 1j\)) is complex with zero real part.

Finally, note that complex numbers \(C\), much like vectors, do not have rank order, which means that one complex number cannot be larger or smaller than another. It makes no sense to say that \(j > 1\) or \(j = 1\) (Boas, 1987). The real and imaginary parts, and the magnitude and phase, have order. Order seems restricted to \(R\). If time \(t\) were complex, there could be no yesterday and tomorrow.\(^7\)

2.1.2 Applications of integers

The most relevant question at this point is why are integers important? First, we count with them so that we can keep track of “how much.” But there is much more to numbers than counting: We use integers for any application where absolute accuracy is essential, such as in banking transactions (making change), the precise computing of dates (Stilwell, 2010, p. 70) and locations (“I’ll meet you at 34th and Vine at noon on Jan. 1, 2034”), and the construction of roads or buildings out of bricks (objects built from a unit size).

To navigate we need to know how to predict the tides and the location of the moon and sun. Integers are important precisely because they are precise: Once a month there is a full moon, easily recognizable. The next day it’s slightly less than full. If we could represent our position as integers in time and space, we would know exactly where we are at all times. But such an integral representation of our position or time is not possible.

The Pythagoreans claimed that everything was Integer. From a practical point of view, it seems they were right. Today all computers compute floating-point numbers as fractionals. However, in theory the Pythagoreans were wrong. The error (difference) is a matter of precision.

\(^7\)One can usefully define \(\xi = x + 1j \cdot ct\) to be complex \((x, t \in R)\), with \(x\) in meters [m], \(t\) in seconds [s], and the speed of light \(c_0\) [m/s].
Numerical Representations of \( \mathbb{I}, \mathbb{R}, \mathbb{C} \): When doing numerical work, one must consider how to compute within the set of reals (i.e., which contain irrationals). There can be no irrational number representation on a computer. The international standard of computation, IEEE floating point numbers, is based on rational approximation. The mantissa and the exponent are both integers, having sign and magnitude. The size of each integer depends on the precision of the number being represented. An IEEE floating-point number is rational because it has a binary (integer) mantissa, multiplied by 2 raised to the power of a binary (integer) exponent. For example, \( \pi \approx \pm 2^{a}b^c \) with \( a, b, c \in \mathbb{Z} \). In summary, IEEE floating-point rational numbers cannot be irrational because irrational representations would require an infinite number of bits.

True floating-point numbers contain irrational numbers, which must be approximated by fractional numbers. This leads to the concept of fractional representation, which requires the definition of the mantissa, base, and exponent, where both the mantissa and the exponent are signed. Numerical results must not depend on the base. One could dramatically improve the resolution of the numerical representation by the use of the fundamental theorem of arithmetic (see p. 35). For example, one could factor the exponent into its primes and then represent the number as \( 2^a3^b7^c \) (\( a, b, c \in \mathbb{Z} \)), etc. Such a representation would improve the resolution of the representation. But even so, the irrational numbers would be approximate. For example, base ten is natural using this representation, since \( 10^n = 2^n5^n \).

Thus

\[
\pi \cdot 10^5 \approx 314,159.27 \ldots = 3 \cdot 2^55^5 + 1 \cdot 2^45^4 + 4 \cdot 2^35^3 + \cdots + 9 \cdot 2^05^0 + 2 \cdot 2^05^{-1} \ldots.
\]

Exercise #6
If we work in base 2 and use the approximation \( \pi \approx 22/7 \), then according to the Matlab/Octave \texttt{dec2bin()} routine, show that the binary representation of \( \pi_2 \cdot 2^{17} \) is

\[
\pi_2 \cdot 2^{17} \approx 411,940_{16} = 64,924_{10} = 1,100,100,100,100,100,100_2.
\]

Solution: First we note that this must be an approximation, since \( \pi \in \mathbb{I} \), which cannot have an exact representation \( \in \mathbb{F} \). We are asking for the fractional \( \in \mathbb{F} \) approximation to \( \pi \):

\[
\hat{\pi}_2 = 22/7 = 3 + 1/7 = [3; 7], \tag{2.1.2.3}
\]

where \( \text{int64} (\text{fix}(2^{17} \cdot 22/7)) = 411,940 \) and \( \text{dec2hex} (\text{int64} (\text{fix}(2^{17} \cdot 22/7))) = 64,924 \), and where 1 and 0 are multipliers of powers of 2, which are then added together:

\[
411,940_{10} = 2^{18} + 2^{17} + 2^{14} + 2^{11} + 2^8 + 2^5 + 2^2.
\]

Computers keep track of the decimal point using the exponent, which in Exercise #6 is the factor \( 2^{17} = 131,072_{10} \). The concept of the decimal point is replaced by an integer that has the desired precision, and a scale factor of any base (radix). This scale factor may be thought of as moving the decimal point to the right (larger number) or left (smaller number). The mantissa fine-tunes the value about a scale factor (the exponent). In all cases the number actually is an integer. Negative numbers are represented by an extra sign bit.

Exercise #7
Using Matlab/Octave and base 16 (i.e., hexadecimal) numbers, with \( \pi_2 = 22/7 \), find (a) \( \pi_2 \cdot 10^5 \) and (b) \( \pi_2 \cdot 2^{17} \).

1. \( \pi_2 \cdot 10^5 \)

Solution: (a) Using the command \texttt{dec2hex(fix(22/7*10^5))} we get \texttt{4cbad}_{16}, since \( 22/7 \times 10^5 = 314,285.7 \ldots \) and \texttt{hex2dec('4cbad')} = 314,285. (b) \( 2^{18} \cdot 11_{16}/7_{16} \)

2. \( \pi_2 \cdot 2^{17} \) Solution: \( 2^{18} \cdot 11_{16}/7_{16} \).

Exercise #8
Write the first 11 primes, base 16.

Solution: The Octave/Matlab command \texttt{dec2hex()} provides the answer:

\footnote{IEEE 754: \url{https://www.h-schmidt.net/FloatConverter/IEEE754.html}}
\footnote{Base 10 is the accepted standard simply because we have 10 fingers that we count with.}
2.1. THE TAXONOMY OF NUMBERS: P, N, Z, Q, F, I, R, C

<table>
<thead>
<tr>
<th>(n)</th>
<th>dec</th>
<th>(\pi_n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\pi_n)</td>
<td>dec</td>
<td>2 3 5 7 11 13 17 19 23 29 31</td>
</tr>
<tr>
<td>(\pi_n)</td>
<td>hex</td>
<td>2 3 5 7 0B 0D 11 13 17 1D 1F</td>
</tr>
</tbody>
</table>

Exercise #9

\(x = 2^{17} \times 22/7\), using IEEE754 double precision:\(^{10}\)

\[
x = 411,940.5625_{10}
\]

\[
= 2^{18} \times 1,198.372
\]

\[
= 0,10010,00,110010,010010,010010,010010_2
\]

\[
= 0x48c92,492_{16}.
\]

The exponent is \(2^{18}\) and the mantissa is 4,793,490\(_{16}\). Here the commas in the binary (0,1) string are to help visualize the quasiperiodic nature of the bitstream. The numbers are stored in a 32-bit format, with 1 bit for the sign, 8 bits for the exponent, and 23 bits for the mantissa.

Perhaps a more instructive number is

\[
x = 4,793,490.0
\]

\[
= 0,100,1010,100,100,100,100,100,100,100_2
\]

\[
= 0x4a924,924_{16},
\]

which has a repeating binary bit pattern of \((100)\)_2, broken by the scale factor 0x4a. Even more symmetrical is

\[
x = 0x24,924,924_{16}
\]

\[
= 00,100,100,100,100,100,100,100,100,100_2
\]

\[
= 6.344,131,191,146,900 \times 10^{-17}.
\]

In this example the repeating pattern is clear in the hex representation as a repeating \((942)_2\), as represented by the double brackets, with the subscript indicating the period—in this case, three digits. As before, the commas are to help with readability and have no other meaning.

The representation of numbers is not unique. For example, irrational complex numbers have approximate rational representations (i.e., \(\pi \approx 22/7\)). A better example is complex numbers \(z \in \mathbb{C}\), which have many representations, as a pair of reals (i.e., \(z = (x, y)\)), or by Euler’s formula, and matrices \((\theta \in \mathbb{R})\):

\[
e^\theta = \cos \theta + j \sin \theta \leftrightarrow \begin{bmatrix} \cos \theta & -\sin \theta \\ \sin \theta & \cos \theta \end{bmatrix}.
\]

At a higher level, differentiable functions (analytic functions) may be represented by a single-valued Taylor series expansion (see p. 76), limited by its region of convergence (RoC).

Pythagoreans and Integers: The integer is the cornerstone of the Pythagorean doctrine—so much so that it caused a fracture within the Pythagoreans when it was discovered that not all numbers are rational. One famous proof of such irrational numbers comes from the spiral of Theodorus, as shown in Fig. 2.1, where the short radius of each triangle has length \(b_n = \sqrt{n}\) with \(n \in \mathbb{N}\), and the long radius (the hypotenuse) is \(c_n = \sqrt{1 + b_n^2} = \sqrt{1 + n}\). This figure may be constructed using a compass and ruler by maintaining right triangles.

Public-key security: An important application of prime numbers is public-key encryption, which is essential for internet security applications (e.g., online banking). Most people assume encryption is done by a personal login and passwords. Passwords are fundamentally insecure for many reasons. Decryption depends on factoring large integers, formed from products of primes having thousands of bits.\(^{11}\) The security is based on the relative ease of multiplying large primes, along with the virtual impossibility of factoring their products.

When a computation is easy in one direction, but its inverse is impossible, it is called a trap-door function. We shall explore trapdoor functions in Appendix 2. If everyone were to switch from passwords to public-key encryption, the internet would be much more secure.\(^{12}\)

\(^{10}\)https://www.h-schmidt.net/FloatConverter/IEEE754.html

\(^{11}\)It would seem that public-key encryption could work by having two numbers with a common prime, and then using the Euclidean algorithm, the greatest common divisor (GCD) could be worked out. One of the integers could be the public-key and the second could be the private key.

\(^{12}\)https://fas.org/irp/agency/dod/jason/cyber.pdf
Figure 2.1: The spiral of Theodorus, made from contiguous right triangles having lengths \( a = 1, b_n = \sqrt{n}, \) with \( n \in \mathbb{N} \), and \( c_n = \sqrt{n+1} \). In this way, each value of \( c_n^2 = b_n^2 + 1 \in \mathbb{N} \). This sequence of triangles generate the set \( \{ \sqrt{n} \}, n \in \mathbb{N} \), and is easily generated using a compass and a ruler. (Adapted from https://en.wikipedia.org/wiki/spiral_of_Theodorus.)

**Puzzles:** Another application of integers is imaginative problems that use integers. An example is the classic Chinese four stone problem: Find the weight of four stones that can be used with a scale to weigh anything (e.g., salt, gold) between 0, 1, 2, \ldots, 40 \text{ [gm]} (Fig. 8, p. 39 in Assignment AE-2). As with the other problems, the answer is not as interesting as the method, since the problem may be easily recast into a related one. This type of problem can be found in airline magazines as amusement on a long flight. This puzzle is best cast as a linear algebra problem, with integer solutions. Again, once you know the trick, it is “easy.”

---

13Whenever someone tells you something is “easy,” you should immediately appreciate that it is very hard, but once you learn a concept, the difficulty evaporates.
2.2 Problems NS-1

Topic of this homework:
Introduction to Matlab/Octave (see the Matlab or Octave tutorial for help)
Deliverables: Report with charts and answers to questions. Hint: Use L\LaTeX.\textsuperscript{14}

Plotting complex quantities in Octave/Matlab

Problem # 1: Consider the functions \( f(s) = s^2 + 6s + 25 \) and \( g(s) = s^2 + 6s + 5 \).

- 1.1: Find the zeros of functions \( f(s) \) and \( g(s) \) using the command \texttt{roots()}

- 1.2: Show the roots of \( f(s) \) as red circles and of \( g(s) \) as blue plus signs.
The \( x \)-axis should display the real part of each root, and the \( y \)-axis should display the imaginary part. Use \texttt{hold on} and \texttt{grid on} when plotting the roots.

- 1.3 Give your figure the title “Complex Roots of \( f(s) \) and \( g(s) \).” Label the \( x \)- and \( y \)-axes “Real Part” and “Imaginary Part.” Hint: Use \texttt{xlabel}, \texttt{ylabel}, \texttt{ylim([-10 10])}, and \texttt{xlim([-10 10])} to expand the axes.

Problem # 2: Consider the function \( h(t) = e^{j2\pi ft} \) for \( f = 5 \) and \( t = \{0:0.01:2\} \).

- 2.1: Use \texttt{subplot} to show the real and imaginary parts of \( h(t) \).
Make two graphs in one figure. Label the \( x \)-axes “Time (s)” and the \( y \)-axes “Real Part” and “Imaginary Part.”

- 2.2: Use \texttt{subplot} to plot the magnitude and phase parts of \( h(t) \).
Use the command \texttt{angle} or \texttt{unwrap(angle())} to plot the phase. Label the \( x \)-axes “Time (s)” and the \( y \)-axes “Magnitude” and “Phase (radians).”

Prime numbers, infinity, and special functions in Octave/Matlab

Problem # 3: Prime numbers, infinity, and special functions.

- 3.1: Use the Matlab/Octave function \texttt{factor} to find the prime factors of 123, 248, 1767, and 999,999.

- 3.2: Use the Matlab/Octave function \texttt{isprime} to determine whether 2, 3, and 4 are prime numbers. What does the function \texttt{isprime} return when a number is prime or not prime? Why?

- 3.3: Use the Matlab/Octave function \texttt{primes} to generate prime numbers between 1 and \( 10^6 \). Save them in a vector \( x \). Plot this result using the command \texttt{hist(x)}.

\textsuperscript{14}https://www.overleaf.com
3.4: Now try \([n, \text{bincenters}] = \text{hist}(x)\). Use \text{length}(n)\) to find the number of bins.

3.5: Set the number of bins to 100 by using an extra input argument to the function \text{hist}. Show the resulting figure, give it a title, and label the axes. Hint: \text{help hist} and \text{doc hist}.

**Problem # 4: Inf, NaN, and logarithms in Octave/Matlab.**

- 4.1: Try \(1/0\) and \(0/0\) in the Octave/Matlab command window. What are the results? What do these “numbers” mean in Octave/Matlab?

- 4.2: Try \(\log(0), \log_{10}(0),\) and \(\log_{2}(0)\) in the command window. In Matlab/Octave, the natural logarithm \(\ln(\cdot)\) is computed using the function \(\log\). Functions \(\log_{10}\) and \(\log_{2}\) are computed using \(\log_{10}\) and \(\log_{2}\).

- 4.3: Try \(\log(1)\) in the command window. What do you expect for \(\log_{10}(1)\) and \(\log_{2}(1)\)?

- 4.4: Try \(\log(-1)\) in the command window. What do you expect for \(\log_{10}(-1)\) and \(\log_{2}(-1)\)?

- 4.5: Explain that Matlab/Octave arrives at the answer in problem 4.4. Hint: \(-1 = e^{i\pi}\).

- 4.6: Try \(\log(\exp(j*\sqrt{\pi}))\) (i.e., \(\log e^{j\sqrt{\pi}}\)) in the command window. What do you expect?

- 4.7: What does inverse mean in this context? What is the inverse of \(\ln f(x)\)?

- 4.8: What is a decibel? (Look up decibels on the internet.)

**Problem # 5: Very large primes on Intel computers.** Find the largest prime number that can be stored on an Intel 64-bit computer, which we call \(\pi_{\text{max}}\). Hint: As explained in the Matlab/Octave command \text{help flintmax}, the largest positive integer is \(2^{53}\); however, the largest integer that can be factored is \(2^{32} = \sqrt{2^{64}}\). Explain the logic of your answer. Hint: \text{help isprime()}. 

**Problem # 6: We are interested in primes that are greater than \(\pi_{\text{max}}\). How can you find them on an Intel computer (i.e., one using IEEE floating point)?** Hint: Consider a sieve that contains only odd numbers, starting from 3 (not 2). Since every prime number greater than 2 is odd, there is no reason to check the even numbers. \(n_{\text{odd}} \in \mathbb{N}/2\) contain all the primes other than 2.

**Problem # 7: The following identity is interesting. Can you find a proof?**
1 = 1^2
1 + 3 = 2^2
1 + 3 + 5 = 3^2
1 + 3 + 5 + 7 = 4^2
1 + 3 + 5 + 7 + 9 = 5^2
\vdots
\sum_{n=0}^{N-1} 2n + 1 = N^2.
2.3 The role of physics in mathematics

Integers arose naturally in art, music, and science. Examples include the relationships between musical notes, the natural eigenmodes (tones) of strings and other musical instruments. These relationships were so common that Pythagoras believed that to explain the physical world (the universe), one needed to understand integers. As discussed on page 11, “all is number” was a seductive song.

As we will discuss on page 61, it is best to view the relationships among acoustics, music, and mathematics as historical, since these topics inspired the development of mathematics. Today integers play a key role in quantum mechanics, again based on eigenmodes, but in this case, eigenmodes follow from solutions the of Schrödinger equation, with the roots of the characteristic equation being purely imaginary. If there were a real part (i.e., damping), the modes would not be integers.

As discussed by Vincent Salmon (1946, p. 201), Schrödinger’s equation follows directly from the Webster horn equation. While Philip Morse (1948, p. 281) (a student of Arnold Sommerfeld) fails to make the direct link, he comes close to the same view when he shows that the real part of the horn resistance goes exactly to zero below a cutoff frequency. He also discusses the trapped modes inside musical instruments created by the horn flare. One may assume Morse read Salmon’s paper on horns, since he cites Salmon (Morse, 1948, footnote 1, p. 271).

Engineers are so accustomed to working with real (or complex) numbers that they rarely acknowledge the distinction between real (i.e., irrational) and fractional numbers. Integers arise in many contexts. One cannot master computer programming without understanding integer, hexadecimal, octal, and binary representations, since all numbers in a computer are represented in numerical computations in terms of rationals \( \mathbb{Q} = \mathbb{Z} \cup \mathbb{F} \).\(^{15}\)

The primary reason integers are so important is their absolute precision. Every integer \( n \in \mathbb{Z} \) is unique\(^{16}\) and has the indexing property, which is essential for making lists that are ordered, so that one can quickly look things up. The alphabet also has this property (e.g., a book’s index).

Because of the integer’s absolute precision, the digital computer quickly overtook the analog computer once it was practical to make logic circuits that were fast. From 1946 the first digital computer was thought to be the University of Pennsylvania’s ENIAC. We now know that the code-breaking effort in Bletchley Park, England, under the guidance of Alan Turing, created the first digital computer (the Colossus), which was used to break the World War II German Enigma code. Due to the high secrecy of this war effort, the credit was only acknowledged in the 1970s when the project was finally declassified.

There is zero possibility of analog computing displacing digital computing because of the importance of precision (and speed). But even with binary representation, there is a nonzero probability of error—for example, on a hard drive—due to physical noise. To deal with this, error-correcting codes have been developed, reducing the error by many orders of magnitude. Today error correction is a science, and billions of dollars are invested to increase the density of bits per area to increasingly larger factors. A few years ago the terabyte drive was unheard of; today it is standard. In a few years petabyte drives will certainly become available. It is hard to comprehend how these will be used by individuals, but they are essential for online (cloud) computing.

2.3.1 The three streams of mathematics

Modern mathematics is built on a hierarchical construct of fundamental theorems, as summarized in the following boxed material (p. 35). The importance of such theorems cannot be overemphasized. Gauss’s and Stokes’s laws play a major role in understanding and manipulating Maxwell’s equations. Every engineering student needs to fully appreciate the significance of these key theorems. If necessary, memorize them. But that will not do over the long run, as each and every theorem must be fully understood. Fortunately most students already know several of these theorems but perhaps not by name. In such cases, it is a matter of mastering the vocabulary.

\(^{15}\)See Appendix A (p. 221) for a review of mathematical notation.

\(^{16}\)Check out the history of \( 1729 = 1^3 + 12^3 = 9^3 + 10^3 \).
The three streams of mathematics

1. Number systems: Stream 1
   - Arithmetic
   - Prime numbers

2. Geometry: Stream 2
   - Algebra

3. Calculus: Stream 3 (Flanders, 1973)
   - Leibniz $\mathbb{R}^1$
   - Complex $\mathbb{C} \subset \mathbb{R}^2$
   - Vectors $\mathbb{R}^3, \mathbb{R}^n, \mathbb{R}^\infty$
     - Gauss’s law (divergence theorem)
     - Stokes’s law (curl theorem, or Green’s theorem)
     - Vector calculus (Helmholtz’s theorem)

These theorems are naturally organized and may be thought of in terms of Stillwell’s three streams. For Stream 1 we have the fundamental theorem of arithmetic and the prime number theorem. For Stream 2 there is the fundamental theorem of algebra, and for Stream 3 there are a host of theorems on calculus, ordered by their dimensionality. Some of these theorems seem trivial (e.g., the fundamental theorem of arithmetic). Others are more challenging, such as the fundamental theorem of vector calculus and Green’s theorem.

Complexity should not be confused with importance. Each of these theorems, as stated, is fundamental. Taken as a whole, they are a powerful way of summarizing mathematics.

2.3.2 Stream 1: Prime number theorems

There are two easily described fundamental theorems about primes:

1. The fundamental theorem of arithmetic: This states that every integer $n \in \mathbb{Z}$ may be uniquely factored into prime numbers. This raises the question of the meaning of factor (split into a product). The product of two integers $m, n \in \mathbb{Z}$ is $mn = \sum m \ b + \sum n \ a$. For example, $2 \times 3 = 2 + 2 + 2 = 3 + 3$.

2. The prime number theorem: One would like to know how many primes there are. That is easy: $|\mathbb{P}| = \infty$ (the size of the set of primes is infinite). A better way of asking this question is What is the average density of primes in the limit as $n \to \infty$? This question was answered, for all practical purposes, by Gauss, who in his free time computed the first three million primes by hand. He discovered that, to a good approximation, the primes are equally likely on a log scale. This is nicely summarized by the limerick:

Chebyshev said, and I say it again: There is always a prime between $n$ and $2n$.

This is attributed to the mathematician Pafnuty Chebyshev and nicely sums up the prime number theorem (Stillwell, 2010, p. 585).

When the ratio of two frequencies (pitches) is 2, the relationship is called an octave. With a slight stretch of terminology, we could say there is at least one prime per octave. An interesting extension of this observation is: as $n \to \infty$ what is the limiting density in primes per fraction of an octave? More specifically let $\pi < \pi_n < \pi_{k+1}$, then what is the limit of $\log_2 \pi_{n+1}/\pi_{n-1}$?

In modern western music the octave is further divided into 12 ratios called semitones equal to $\frac{12}{\sqrt{2}}$. Twelve semitones is an octave. In the end, it is a question of the density of primes on a log-log (i.e., ratio) scale. One might wonder about the maximum number of primes per octave as a function of $N$, or ask for the fractions of an octave (factors of 2) for $\pi_k$ as $k$ becomes large? The maximum value of $R_k < 0.5$ thus Chebyshev’s bound of 2 is quite conservative, by a factor of 3 (0.6/2=0.3). As $k \to \infty$ the bound is exponentially tightened. The results of this calculation are show in Fig. 2.2. For reference when $k = 9592$, $\pi_k = 99991$, $\pi_{k+1} = 99989$, $\pi_{k+1} = 100003$, thus $1 - P(k+1)/P(k-1) = 1.4e - 4$. 

2.3.3 Stream 2: Fundamental theorem of algebra

This theorem states that every polynomial in $x$ of degree $N$,

$$P_N(x) = \sum_{k=0}^{N} a_k x^k,$$

has at least one root (see p. 89). When a common root is factored out, the degree of the polynomial is reduced by 1. Applied recursively, a polynomial of degree $N$ has $N$ roots. Note there are $N + 1$ coefficients (i.e., $[a_N, a_{N-1}, \ldots, a_0]$). If we are interested in only the roots of $P_N(x)$, it is best to define $a_N = 1$, which defines the monic polynomial. If the roots are fractional numbers, this might be possible. However, if the roots are irrational numbers (likely), a perfect factorization is at least unlikely if not impossible.

2.3.4 Stream 3: Fundamental theorems of calculus

In Sec. 5.6 and Sec. 5.6.6 we will deal with each of the theorems for Stream 3. We consider the several fundamental theorems of integration, starting with Leibniz’s formula for integration on the real line ($\mathbb{R}$), then progressing to complex integration in the complex plane ($\mathbb{C}$) (Cauchy’s theorem), which is required for computing the inverse Laplace transform. Gauss’s and Stokes’s laws for $\mathbb{R}^2$ require closed and open surfaces, respectively. One cannot manipulate Maxwell’s equations, fluid flow, or acoustics without understanding these theorems. Any problem that deals with the wave equation in more than one dimension requires an understanding of these theorems. They are the basis of the derivation of the Kirchhoff voltage and current laws.

Finally we define three vector operations based on the gradient operator,

$$\nabla \equiv \hat{x} \frac{\partial}{\partial x} + \hat{y} \frac{\partial}{\partial y} + \hat{z} \frac{\partial}{\partial z},$$

pronounced “del” (preferred) or “nabla,” which are the gradient $\nabla()$, divergence $\nabla \cdot ()$, and curl $\nabla \times ()$.

Second-order operators such as the scalar Laplacian $\nabla \cdot \nabla () = \nabla^2 ()$ (God) may be constructed from first-order operators. The most important of these is the vector Laplacian $\nabla^2 ()$, which is required when working with Maxwell’s wave equations.

The first three operations are defined in terms of integral operations on a surface in one, two, or three dimensions by taking the limit as that surface, or the volume contained within, goes to zero. These three differential operators are essential to fully understand Maxwell’s equations, the crown jewel of mathematical physics. Hence mathematics plays a key role in physics, as does physics in math.

2.3.5 Other key mathematical theorems

In addition to the widely recognized fundamental theorems for the three streams, there are a number of equally important theorems that have not yet been labeled “fundamental.”

---

17 It is not clear what it takes to reach this more official sounding category.
The widely recognized Cauchy integral theorem is an excellent example, since it is a steppingstone to Green’s theorem and the fundamental theorem of complex calculus. In Sec. 4.5 (see p. 151) we clarify the contributions of each of these special theorems.

Once these fundamental theorems of integration (Stream 3) have been mastered, the student is ready for the complex frequency domain, which takes us back to Stream 2 and the complex frequency plane \((s = \sigma + \omega j \in \mathbb{C})\). While the Fourier and \(\mathcal{L}T\)’s are taught in mathematics courses, the concept of complex frequency is rarely mentioned. The complex frequency domain (see p. 120) and causality are fundamentally related (see pp. 152–162), and are critical for the analysis of signals and systems, especially when dealing with the concept of impedance (see p. 142).

Without the concept of time and frequency, we cannot develop an intuition for the Fourier and Laplace transforms, especially within the context of engineering and mathematical physics. The Fourier transform covers signals, while the Laplace transform describes systems. Separating these two concepts, based on their representations as Fourier and Laplace transforms, is an important starting place for understanding physics and the role of mathematics. However, these methods, by themselves, do not provide the insight into physical systems that we need to be productive or, better, creative with these tools. We need to master the tools of differential equations and then partial differential equations to fully appreciate the world that they describe. Electrical and mechanical networks composed of inductors, capacitors, and resistors are isomorphic to mechanical systems composed of masses, springs, and dash-pots. Newton’s laws are analogous to those of Kirchhoff, which are the rules needed to analyze simple physical systems composed of linear (and nonlinear) subcomponents. When lumped-element systems are taken to the limit in several dimensions, we obtain Maxwell’s partial differential equations, the laws of continuum mechanics, and beyond.

The ultimate goal of this text is to make you aware of and productive in using these tools. This material can be best absorbed by treating it chronologically through history, so you can see how this body of knowledge came into existence, through the minds and hands of Galileo, Newton, Maxwell, and Einstein. Perhaps one day you too can stand on the shoulders of the giants who went before you.
2.4 Problems NS-2

Topic of this homework:
Prime numbers, greatest common divisors, the continued fraction algorithm
   Deliverables: Answers to questions

Prime numbers

Problem # 1: Every integer may be written as a product of primes.

– 1.1: Write the numbers 1,000,000, 1,000,004, and 999,999 in the form \(N = \prod_k \pi_k^{\beta_k}\).
   Hint: Use Matlab/Octave to find the prime factors.

– 1.2: Give a generalized formula for the natural logarithm of a number \(\ln(N)\) in terms of its primes \(\pi_k\) and their multiplicities \(\beta_k\). Express your answer as a sum of terms.

Problem # 2: Using the computer

– 2.1: Explain why the following brief Matlab/Octave program returns the prime numbers \(\pi_k\) between 1 and 100.

\[
n=2:100; \quad k = \text{isprime}(n); \quad n(k)
\]

– 2.2: How many primes are there between 2 and \(N = 100\)?

Problem # 3: Prime numbers may be identified using a sieve (see 2.3).

– 3.1: By hand, complete the sieve of Eratosthenes for \(n = 1, \ldots, 49\). Circle each prime \(p\), then cross out each number that is a multiple of \(p\).

– 3.2: What is the largest number you need to consider before only primes remain?

– 3.3: Generalize: For \(n = 1, \ldots, N\), what is the largest number you need to consider before only the primes remain?

– 3.4: Write each of these numbers as a product of primes: 22, 30, 34, 43, 44, 48, 49.

– 3.5: Find the largest prime \(\pi_k \leq 100\). Do not use Matlab/Octave other than to check your answer. Hint: Write the numbers starting with 100 and count backward: 100, 99, 98, 97, \ldots. Cross off the even numbers, leaving 99, 97, 95, \ldots. Pull out a factor (only one is necessary to show that it is not prime).
3.6: Find the largest prime $\pi_k \leq 1000$. Do not use Matlab/Octave other than to check your answer.

3.7: Explain why $\pi_k^{-s} = e^{-s\ln \pi_k}$.

### Greatest common divisors

Consider using the Euclidean algorithm to find the greatest common divisor (GCD; the largest common prime factor) of two numbers. Note that this algorithm may be performed using one of two methods:

<table>
<thead>
<tr>
<th>Method</th>
<th>Division</th>
<th>Subtraction</th>
</tr>
</thead>
<tbody>
<tr>
<td>On each iteration...</td>
<td>$a_{i+1} = b_i$</td>
<td>$a_{i+1} = \max(a_i, b_i) - \min(a_i, b_i)$</td>
</tr>
<tr>
<td></td>
<td>$b_{i+1} = a_i - b_i \cdot \text{floor}(a_i/b_i)$</td>
<td>$b_{i+1} = \min(a_i, b_i)$</td>
</tr>
<tr>
<td>Terminates when...</td>
<td>$b = 0$ (GCD $= a$)</td>
<td>$b = 0$ (GCD $= a$)</td>
</tr>
</tbody>
</table>

The division method (Eq. 2.1, Sec. 2.1.2, Ch. 2) is preferred because the subtraction method is much slower.

**Problem #4: Understanding the Euclidean algorithm (GCD)**

4.1: Use the Octave/Matlab command `factor` to find the prime factors of $a = 85$ and $b = 15$.

4.2: What is the greatest common prime factor of $a = 85$ and $b = 15$?

4.3: By hand, perform the Euclidean algorithm for $a = 85$ and $b = 15$.

4.4: By hand, perform the Euclidean algorithm for $a = 75$ and $b = 25$. Is the result a prime number?

4.5: Consider the first step of the GCD division algorithm when $a < b$ (e.g., $a = 25$ and $b = 75$). What happens to $a$ and $b$ in the first step? Does it matter if you begin the algorithm with $a < b$ rather than $b < a$?

4.6: Describe in your own words how the GCD algorithm works. Try the algorithm using numbers that have already been divided into factors (e.g., $a = 5 \cdot 3$ and $b = 7 \cdot 3$).

4.7: Find the GCD of $2 \cdot \pi_{25}$ and $3 \cdot \pi_{25}$.

**Problem #5: Coprimes**

5.1: Define the term coprime.

5.2: How can the Euclidean algorithm be used to identify coprimes?

5.3: Give at least one application of the Euclidean algorithm.

5.4: Write a Matlab function, `function x = my_gcd(a, b)`, that uses the Euclidean algorithm to find the GCD of any two inputs $a$ and $b$. Test your function on the $(a, b)$ combinations from the previous problem. Include a printout (or hand write) your algorithm to turn in.

**Hints and advice:**
• Don’t give your variables the same names as Matlab functions! Since \texttt{gcd} is an existing Matlab/Octave function, if you use it as a variable or function name, you won’t be able to use \texttt{gcd} to check your \texttt{gcd()} function. Try clear all to recover from this problem.

• Try using a “while” loop for this exercise (see Matlab documentation for help).

• You may need to use some temporary variables for \(a\) and \(b\) in order to perform the algorithm.

### Algebraic generalization of the GCD (Euclidean) algorithm

**Problem #6:** In this problem we are looking for integer solutions \((m, n) \in \mathbb{Z}\) to the equations

\[ ma + nb = \gcd(a, b) \text{ and } ma + nb = 0 \text{ given positive integers } (a, b) \in \mathbb{Z}^+. \]

Note that this requires that either \(m\) or \(n\) be negative. These solutions may be found using the Euclidean algorithm only if \((a, b)\) are coprime \((a \perp b)\). Note that integer (whole number) polynomial relations such as these are known as Diophantine equations. The above equations are linear Diophantine equations, possibly the simplest form of such relations.

**Example:** \(\gcd(2, 3) = 1\): For \((a, b) = (2, 3)\), the result is

\[
\begin{bmatrix}
1 \\
0
\end{bmatrix} = \begin{bmatrix}
0 & 1 \\
1 & -2
\end{bmatrix} \begin{bmatrix}
0 & 1 \\
1 & 0
\end{bmatrix} \begin{bmatrix}
2 \\
3
\end{bmatrix} = \begin{bmatrix}
-1 & 1 \\
3 & -2
\end{bmatrix} \begin{bmatrix}
2 \\
3
\end{bmatrix}.
\]

Thus from the above equation we find the solution \((m, n)\) to the integer equation

\[ 2m + 3n = \gcd(2, 3) = 1; \]

namely, \((m, n) = (-1, 1)\) (i.e., \(-2 + 3 = 1\)). There is also a second solution \((3, -2)\) (i.e., \(3 \cdot 2 - 2 \cdot 3 = 0\)) that represents the terminating condition. Thus these two solutions are a pair and the solution exists only if \((a, b)\) are coprime \((a \perp b)\).

**Subtraction method:** This method is more complicated than the division algorithm because at each stage we must check whether \(a < b\). Define

\[
\begin{bmatrix}
a_i \\
b_i
\end{bmatrix} = Q \begin{bmatrix}
a_{i+1} \\
b_{i+1}
\end{bmatrix}, \quad Q = \begin{bmatrix}
1 & -1 \\
0 & 1
\end{bmatrix}, \quad S = \begin{bmatrix}
0 & 1 \\
1 & 0
\end{bmatrix},
\]

where \(Q\) sets \(a_{i+1} = a_i - b_i\) and \(b_{i+1} = b_i\) assuming \(a_i > b_i\), and \(S\) is a swap matrix that swaps \(a_i\) and \(b_i\) if \(a_i < b_i\). Using these matrices, we implement the algorithm by assigning

\[
\begin{bmatrix}
a_{i+1} \\
b_{i+1}
\end{bmatrix} = Q \begin{bmatrix}
a_i \\
b_i
\end{bmatrix} \text{ for } a_i > b_i, \quad \begin{bmatrix}
a_{i+1} \\
b_{i+1}
\end{bmatrix} = QS \begin{bmatrix}
a_i \\
b_i
\end{bmatrix} \text{ for } a_i < b_i.
\]

The result of this method is a cascade of \(Q\) and \(S\) matrices. For \((a, b) = (2, 3)\), the result is

\[
\begin{bmatrix}
1 \\
1
\end{bmatrix} = \begin{bmatrix}
1 & -1 \\
0 & 1
\end{bmatrix} \begin{bmatrix}
1 \\
0
\end{bmatrix} = \begin{bmatrix}
1 \\
0
\end{bmatrix} = \begin{bmatrix}
2 \\
3
\end{bmatrix} = \begin{bmatrix}
-1 & 1 \\
0 & 1
\end{bmatrix} \begin{bmatrix}
2 \\
3
\end{bmatrix}.
\]

Thus we find two solutions \((m, n)\) to the integer equation \(2m + 3n = \gcd(2, 3) = 1\).

- 6.1: By inspection, find at least one integer pair \((m, n)\) that satisfies \(12m + 15n = 3\).

- 6.2: Using matrix methods for the Euclidean algorithm, find integer pairs \((m, n)\) that satisfy \(12m + 15n = 3\) and \(12m + 15n = 0\). Show your work!!

- 6.3: Does the equation \(12m + 15n = 1\) have integer solutions for \(n\) and \(m\)? Why or why not?

**Problem #7: Matrix approach:**

It can be difficult to keep track of the \(a\)’s and \(b\)’s when the algorithm has many steps. We need an alternative way to run the Euclidean algorithm using matrix algebra. Matrix methods provide a more transparent approach to the operations on \((a, b)\). Thus the Euclidean algorithm can be classified in terms of standard matrix operations. Write out the indirect matrix approach discussed at the end of Sec. 2.5.3 (Eq. 2.5.3.3).
Continued fractions

**Problem #8:** Here we explore the continued fraction algorithm (CFA), discussed in Sec. 2.5.4. In its simplest form, the CFA starts with a real number, which we denote as $\alpha \in \mathbb{R}$. Let us work with an irrational real number, $\pi \in \mathbb{I}$, as an example because its CFA representation will be infinitely long. We can represent the CFA coefficients $\alpha$ as a vector of integers $n_k$, $k = 1, 2, \ldots, \infty$:

$$\alpha = [n_1; n_2, n_3, n_4, \ldots] = n_1 + \frac{1}{n_2 + \frac{1}{n_3 + \frac{1}{n_4 + \cdots}}}.$$

As discussed in Sec. 2.5.3 (p. 44), the CFA is recursive, with three steps per iteration. For $\alpha_1 = \pi$, $n_1 = 3$, $r_1 = \pi - 3$, and $\alpha_2 \equiv 1/r_1$,

$$\alpha_2 = \frac{1}{0.1416} = 7.0625\ldots$$

$$\alpha_1 = n_1 + \frac{1}{\alpha_2} = n_1 + \frac{1}{n_2 + \frac{1}{\alpha_3}} = \cdots.$$

In terms of a Matlab/Octave script,

```matlab
alpha0 = pi;
K=10;
n=zeros(1,K); alpha=zeros(1,K);
alpha(1)=alpha0;
for k=2:K %k=1 to K
n(k)=round(alpha(k-1));
%n(k)=fix(alpha(k-1));
alpha(k)= 1/(alpha(k-1)-n(k));
%disp([fix(k), round(n(k)), alpha(k)]); pause(1)
end
disp([n; alpha]);
%Now compare this to matlab’s rat() function
rat(alpha0,1e-20)
```

- **8.1:** By hand (you may use Matlab/Octave as a calculator), find the first three values of $n_k$ for $\alpha = e^\pi$.

- **8.2:** For the preceding question, what is the error (remainder) when you truncate the continued fraction after $n_1, \ldots, n_3$? Give the absolute value of the error and the percentage error relative to the original $\alpha$.

- **8.3:** Use the Matlab/Octave program provided to find the first 10 values of $n_k$ for $\alpha = e^\pi$, and verify your result using the Matlab/Octave command `rat()`.

- **8.4:** Discuss the similarities and differences between the Euclidean algorithm and the CFA.

- **8.5:** Extra Credit: Show that the CFA is the inverse operation of the GCD (i.e., the CFA is the GCD run in reverse). (Hint: see Sec. 2.5.3.)

**Continued fraction algorithm (CFA)** (8 pts)

**Problem #9:** CFA of ratios of large primes
9.1: Starting from the primes below $10^6$, form the CFA of $\pi_j/\pi_k$ with $j = 78498$ and $k < j$.

9.2: Look at other ratios of prime numbers and look for a pattern in the CFA of the ratios of large primes. What is the most obvious conclusion?

9.3: (4pts) Expand $23/7$ as a continued fraction. Express your answer in bracket notation (e.g., $\pi = [3., 7, 16, \cdots]$). Show your work.

9.4: (2pts) Can $\sqrt{2}$ be represented as a finite continued fraction? Why or why not?

9.5: (2pts) What is the CFA for $\sqrt{2} - 1$?

Hint: $\sqrt{2} + 1 = \frac{1}{\sqrt{2} - 1} = [2; 2, 2, \cdots]$.

9.6: Find the CFA for $1 + \sqrt{3}$

9.7: Show that

$$\frac{1}{1 - \sqrt{a}} = a^{1/2} + a^{3/2} + a^{7/2} + a^{17/2} + a^{31/2} + a^5 + a^8 + a^9 + a^{12} + a + 1 = 1 - a^6$$

Use symbolic analysis to show this, then explain.

2.5 Applications of prime numbers

If someone asked you for a theory of counting numbers, I suspect you would laugh and start counting. It sounds like either a stupid question or a bad joke. Yet integers are a rich topic, so the question is not even slightly dumb. It is somewhat amazing that even birds and bees can count. While I doubt birds and bees can recognize primes, cicadas and other insects crawl out of the ground only in prime-number cycles (e.g., 13- or 17-year cycles). If you have ever witnessed such an event (I have), you will never forget it. Somehow they know. Finally, there is an analytic function, first introduced by Euler based on his analysis of the sieve, now known as the Riemann zeta function $\zeta(s)$, that is complex analytic, with its poles at the logs of the prime numbers. The properties of this function are truly amazing, even fun. Many of the questions and answers about primes go back to at least the early Chinese (ca. 1500 BCE).

2.5.1 The importance of prime numbers

Each prime perfectly predicts multiples of that prime, but there seems to be no regularity in predicting primes. It follows that prime numbers are the key to the theory of numbers because of the fundamental theorem of arithmetic (FTA).

It is likely that the first insight into the counting numbers started with the sieve shown in Fig. 2.3. A recursive sieve method for finding primes was first devised by the Greek Eratosthenes (O’Neill, 2009). A sieve answers the question How can one identify the prime numbers? The answer comes from looking for irregular patterns in the counting numbers.

For example, starting from $\pi_1 = 2$, we strike out all even numbers $2(2, 3, 4, 5, 6, \ldots)$ but not 2. By definition, the multiples are products of the target prime (2 in our example) and every other integer ($n \geq 2$). In this way all the even numbers are removed in this first iteration. The next remaining integer (3 in our example) is identified as the second prime $\pi_2$. Then all the multiples of $\pi_2 = 3$ are removed. The next remaining number is $\pi_3 = 5$, so all multiples of $\pi_3 = 5$ are removed (i.e., $10, 15, 20, \ldots$). This process is repeated until all the numbers of the list have been either canceled or identified as prime.
1. Write \( N - 1 \) integers \( n \), starting from 2: \( n \in \{2, 3, \ldots, N\} \) (e.g., \( N = 4, n \in \{2, 3, 4\} \)). Note that the first element \( \pi_1 = 2 \) is the first prime. Cross out all multiples of \( \pi_1 \); that is, cross out \( n \cdot \pi_1 = 4, 6, 8, 10, \ldots, 50 \), or all \( n \) such that \( \text{mod}(n, \pi_1) = 0 \).

2. Let \( k = 2 \) and note that \( \pi_2 = 3 \). Cross out \( n\pi_2 \cdot (2, 3, 4, 5, 6, 7, \ldots, 45) \), that is, all \( n \) such that \( \text{mod}(n, \pi_2) = 0 \).

3. Let \( k = 3, \pi_3 = 5 \). Cross out \( n\pi_3 \cdot (25, 35) \) (mod \( n, 5 \) = 0).

4. Finally let \( k = 4, \pi_4 = 7 \) (mod \( n, 7 \) = 0). Cross out \( n\pi_4 \): (49). Thus there are 15 primes less than \( N = 50 \): \( \pi_k = 2, 3, 5, 7, 11, 13, 17, 19, 23, 29, 31, 37, 41, 43, 47 \) (highlighted in red). Above 2, all end in odd numbers, and above 5, all end with 1, 3, 7, or 9.

Figure 2.3: Sieve of Eratosthenes for \( N = 50 \).
As the word *sieve* implies, this process takes a heavy toll on the integers, rapidly pruning the nonprimes. In four iterations of the sieve algorithm, all the primes less than $N = 50$ are identified in red. The final set of primes is displayed in step 4 of Fig. 2.3.

Once a prime greater than $N/2$ has been identified (25 in the example), the recursion stops, since twice that prime is greater than $N$, the maximum number under consideration. Thus once $\sqrt{49}$ has been reached, all the primes have been identified (this follows from the fact that the next prime $\pi_n$ is multiplied by an integer $n = 1, \ldots, N$).

When we use a computer, memory efficiency and speed are the main considerations. There are various schemes for making the sieve more efficient. For example, the recursion $n\pi_k = (n - 1)\pi_k + \pi_k$ will speed up the process by replacing the multiplication with an addition of $\pi_k$.

### 2.5.2 Two fundamental theorems of primes

Early theories of numbers revealed two fundamental theorems (see Sec. 2.3.3, Sec. 2.3.4). The first of these is the fundamental theorem of arithmetic, which says that every integer $n \in \mathbb{N}$ greater than 1 may be uniquely factored into a product of primes:

$$n = \prod_{k=1}^{K} \pi_k^{\beta_k},$$

where $k = 1, \ldots, K$ indexes the integer’s $K$ prime factors $\pi_k \in \mathbb{P}$. Typically prime factors appear more than once—for example, $25 = 5^2$. To make the notation compact we define the *multiplicity* $\beta_k$ of each prime factor $\pi_k$. For example, $2312 = 2^3 \cdot 17^2 = \pi_1^3 \pi_2^2$ (i.e., $\pi_1 = 2, \beta_1 = 3; \pi_2 = 17, \pi_7 = 2$) and $2313 = 3^2 \cdot 257 = \pi_3^2 \pi_55$ (i.e., $\pi_3 = 3, \beta_3 = 2; \pi_55 = 257, \beta_{55} = 1$). Our demonstration of this is empirical, using the Matlab/Octave *factor*(N) routine, which factors $N$.\(^{18}\)

What seems amazing is the unique nature of this theorem. Each counting number is uniquely represented as a product of primes. No two integers can share the same factorization. Once you multiply the factors out, the result is unique ($N$). Note that it’s easy to multiply integers (e.g., primes) but expensive to factor them. And factoring the product of three primes is significantly more difficult than factoring two.

Factoring is much more expensive than division. This is not due to the higher cost of division over multiplication, which is less than a factor of 2.\(^{19}\) Dividing the product of two primes, given one, is trivial, slightly more expensive that multiplying. Factoring the product of two primes is nearly impossible, as one needs to know what to divide by. Factoring means dividing by some integer and obtaining another integer with remainder zero.

This brings us to the prime number theorem (PNT). The security problem is the reason these two theorems are so important: (1) Every integer has a unique representation as a product of primes, and (2) the density of primes is large (see the discussions on p. 35). Thus security reduces to the “needle in the haystack problem” due to the cost of a search.

One could factor a product of primes $N = \pi_k \pi_l$ by doing $M$ divisions, where $M$ is the number of primes less than $N$. This assumes the primes less than $N$ are known. However, most integers are not a simple product of two primes.

But the utility of using prime factorization has to do with their density. If we were simply looking up a few numbers from a short list of primes, it would be easy to factor them. But given that their density is logarithmic ($\gg 1$ per octave), factoring comes at a very high computational cost.

### 2.5.3 Greatest common divisor (Euclidean algorithm)

The Euclidean algorithm is a systematic method to find the largest common integer factor $k$ between two integers $n$ and $m$, denoted $k = \gcd(n, m)$, where $n, m, k \in \mathbb{N}$ (Graham et al., 1994). For example, $15 = \gcd(30, 105)$ since, when factored $(30, 105) = (2 \cdot 3 \cdot 5, 7 \cdot 3 \cdot 5) = 3 \cdot 5 \cdot (2, 7) = 15 \cdot (2, 7)$. The Euclidean algorithm was known to the Chinese (i.e., not discovered by Euclid) (Stillwell, 2010, p. 41). Two integers are said to be *coprime* if their GCD is 1 (i.e., they have no common factor).

The Euclidean algorithm is best explained by a trivial example: Consider the two numbers 6 and 9. At each step the smaller number (6) is subtracted from the larger (9) and the smaller number and the difference (the remainder) are saved. This process continues until the two resulting numbers are equal, which is the GCD. For our example, $9 - 6 = 3$, leaving the smaller number 6 and the difference 3. Repeating this, we get $6 - 3 = 3$, leaving the smaller number 3 and the difference 3. Since these two numbers are the same, we are done; thus

\(^{18}\)If you wish to be a mathematician, you need to learn how to prove theorems. If you’re a physicist, you are happy that someone else has already proved them, so that you can use the result.

\(^{19}\)https://streamcomputing.eu/blog/2012-07-16/how-expensive-is-an-operation-on-a-cpu/
2.5. APPLICATIONS OF PRIME NUMBERS

Examples of the GCD: \( l = \gcd(m, n) \)

- Examples \((m, n, l \in \mathbb{Z})\):
  - \(5 = \gcd(13 \cdot 5, 11 \cdot 5)\). The GCD is the common factor 5.
  - \((13 \cdot 10, 11 \cdot 10) = 10 \gcd(130, 110) = 10 = 2 \cdot 5\) is not prime
  - \(\gcd(1234, 1024) = 2\), since \(1234 = 2 \cdot 617, 1024 = 2^{10}\)
  - \(\gcd(\pi_k \pi_m, \pi_k \pi_n) = \pi_k\)
  - \(l = \gcd(m, n)\) is the part that cancels in the fraction \(m/n \in F\)
  - \(m/\gcd(m, n) \in \mathbb{Z}\)

- Coprimes \((m \perp n)\) are numbers that have no distinct common factors; that is, \(\gcd(m, n) = 1\)
  - The GCD of two primes is always 1: \(\gcd(13, 11) = 1, \gcd(\pi_m, \pi_n) = 1 (m \neq n)\)
  - If \(m \perp n\), then \(\gcd(m, n) = 1\).
  - If \(\gcd(m, n) = 1\), then \(m \perp n\).

\(3 = \gcd(9, 6)\). We can verify this result by factoring [e.g., \((9, 6) = 3(3, 2)\)]. The value may also be numerically verified using the Matlab/Octave GCD command \(\gcd(6, 9)\), which returns 3.

\[3 = \gcd(9, 6)\]

Figure 2.4: The Euclidean algorithm recursively subtracts \(n\) from \(m\) until the remainder \(m - kn\) is less than either \(n\) or zero. Note that this recursion is the same as \(\text{mod}(m, n)\). Thus the GCD recursively computes \(\text{mod}(m, n)\) until the remainder \(\text{rem}(m, n)\) is less than \(n\), which is called the GCD’s turning point. It then swaps \(m\) and \(n\), so that \(n < m\). This repeats until it terminates on the GCD. Due to its simplicity this is called the direct method for finding the GCD. For the case depicted here, the value of \(k\) renders the remainder \(m - 6n < n\). If one more step were taken beyond the turning point \((k = 7)\), the remainder would become negative. Thus the turning point satisfies the linear relationship \(m - \alpha n = 0\) with \(\alpha \in \mathbb{R}\).

Direct matrix method: The GCD may be written as a matrix recursion given the starting vector \((m_0, n_0)^T\). The recursion is then

\[
\begin{bmatrix}
    m_{k+1} \\
    n_{k+1}
\end{bmatrix} = \begin{bmatrix}
    1 & -1 \\
    0 & 1
\end{bmatrix} \begin{bmatrix}
    m_k \\
    n_k
\end{bmatrix}.
\]

This recursion continues until \(m_{k+1} < n_{k+1}\), at which point \(m\) and \(n\) must be swapped. The process is repeated until \(m_k = n_k\), which equals the GCD. We call this the direct method (see Fig. 2.4). The direct method is inefficient because it recursively subtracts \(n_k\) many times until the resulting \(m_k\) is less than \(n_k\). It also must test for \(m \leq n\) after each subtraction and then swap them if \(m_k < n_k\). If they are equal, we are done.

The GCD’s turning point may be defined using the linear interpolation \(m - \alpha n = 0, \alpha \in \mathbb{R}\), where the solid line crosses the abscissa in Fig. 2.4. If, for example, \(l = 6 + 43/97 \approx 6.443298 \ldots\), then \(6 = \lfloor m/n \rfloor < n\) and \(\alpha \in \mathbb{F} \subset \mathbb{R}\). This is nonlinear (truncation) arithmetic, which is a natural property of the GCD. The \(\text{floor()}\) functions finds the turning point, where we swap the two numbers, since by definition, \(m > n\). In this example, \(6 = \lfloor l \rfloor\).
Exercise #10
Show that
\[
\begin{bmatrix}
1 & -1 \\
0 & 1 \\
\end{bmatrix}^n = \begin{bmatrix}
1 & -n \\
0 & 1 \\
\end{bmatrix}.
\]

**Solution:** To prove this let \( n = 2 \) and then 3. Each recursive multiplication adds 1 to the upper right corner.

**Why is the GCD important?** The utility of the GCD algorithm arises directly from the fundamental difficulty in factoring large integers. Computing the GCD using the Euclidean algorithm costs less than factoring when finding the coprime factors, which is extremely expensive. The utility surfaces when the two numbers are composed of very large primes.

When two integers have no common factors, they are said to be **coprime** and their GCD is 1. The ratio of two integers that are coprime is automatically in reduced form (they have no common factors). For example, \( \frac{4}{2} \in \mathbb{Q} \) is not reduced, since \( 2 = \gcd(4, 2) \) (with a zero remainder). Canceling out the common factor 2 gives the reduced form \( \frac{2}{1} \in \mathbb{F} \). If the GCD were 10^9 digits, it is obvious that any common factor would need to be be removed, thus greatly simplifying further computation. This will make a huge difference when using IEEE754.

The floor function and the GCD are related in an important way, as discussed next.

**Indirect matrix method:** A much more efficient method uses the floor() function, which is called division with rounding, or simply the indirect method. Specifically the GCD may be written in one step as
\[
\begin{bmatrix}
m \\
n \\
\end{bmatrix}_{k+1} = \begin{bmatrix}
0 & 1 \\
1 & -\left\lfloor \frac{m}{n} \right\rfloor \\
\end{bmatrix} \begin{bmatrix}
m \\
n \\
\end{bmatrix}_k. 
\]

This matrix is Eq. 2.5.3.2 to the power \( \left\lfloor \frac{m}{n} \right\rfloor \), followed by swapping the inputs (the smaller must always be on the bottom).

**The GCD and multiplication:** Multiplication is simply recursive addition, and finding the GCD takes advantage of this fact. For example, \( 3 \ast 2 = 3 + 3 = 2 + 2 + 2 \). Since division is the inverse of multiplication, it must be recursive subtraction.

**The GCD and long division:** When we learn how to divide a smaller number into a larger one, we must learn how to use the GCD. For example, suppose we wish to compute \( 6 \div 110 \). We start by finding out how many times 6 goes into 11. Since \( 6 \times 2 = 12 \), which is larger than 11, the answer is 1. We then subtract 6 from 11 to find the remainder 5. This is, of course, the floor function (e.g., \( \left\lfloor \frac{11}{6} \right\rfloor = 1, [11/5] = 2 \)).

**Example:** Start with the two integers \([873, 582]\). In factored form these are \([\pi 25 \cdot 3^2, \pi 25 \cdot 3 \cdot 2]\). Given the factors, we see that the largest common factor is \( \pi 25 \cdot 3 = 291 \) (\( \pi 25 = 97 \)). When we take the ratio of the two numbers, this common factor cancels:
\[
\frac{873}{582} = \frac{\pi 25 \cdot 3}{\pi 25 \cdot 2} = \frac{3}{2} = 1.5.
\]

Of course if we divide 582 into 873 we numerically obtain the answer 1.5 \( \in \mathbb{F} \).

**Exercise #11**
What does it mean to reach the turning point when using the Euclidean algorithm?

**Solution:** When \( m/n - \left\lfloor m/n \right\rfloor < n \), we have reached a turning point. When the remainder is zero (i.e., \( m/n - \left\lfloor m/n \right\rfloor = 0 \)), we have reached the GCD.

**Exercise #12**
Show that in Matlab/Octave \( \text{rat}(873/582) = 1+1/(-2) \) gives the wrong answer (\( \text{rats}(837/582)=3/2 \)).

**Hint:** Factor the two numbers and cancel out the GCD.

**Solution:** Since
\[
\text{factor}(873) = 3 \cdot 3 \cdot 97 \text{ and factor}(582) = 2 \cdot 3 \cdot 97,
\]
the GCD is $3 \cdot 97$. Thus $3/2 = 1 + 1/2$ is the correct answer. But due to rounding methods, it is not $3/2$. As an example, in Matlab/Octave $\text{rat}(3/2) = 2+1/(-2)$. Matlab’s $\text{rat}()$ command uses rounding rather than the floor function, which explains the difference. When the $\text{rat}()$ function produces negative numbers, rounding is employed.

**Exercise #13**
Divide 10 into 99. The floor function ($\text{floor}(99/10)$) must be used, followed by the remainder function ($\text{rem}(99,10)$).

**Solution:** When we divide a smaller number into a larger one, we must first find the floor followed by the remainder. For example, $99/10 = 9 + 9/10$ has a floor of 9 and a remainder of 9/10.

**Graphical description of the GCD:** The Euclidean algorithm is best viewed graphically. In Fig. 2.4 we show what is happening as one approaches the turning point, at which point the two numbers must be swapped to keep the difference positive, which is addressed by the upper row of Eq. 2.5.3.3.

Here is a simple Matlab/Octave code to find $l=\text{gcd}(m,n)$ based on the Stillwell (2010) definition:

```matlab
function k = \gcd(m,n)
while m ~=0
A=m; B=n;
 m=max(A,B); n=min(A,B); %m>n
m=m-n;
endwhile %m=n
k=A;
```

This program loops until $m = 0$.

**Coprimes:** When the GCD of two integers is 1, the only common factor is 1. This is of key importance when trying to find common factors between the two integers. When $1=\text{gcd}(m,n)$, the two integers are said to be **coprime** or **relatively prime**, which is frequently written as $m \perp n$. By definition, the largest common factor of coprimes is 1. But since 1 is not a prime ($\pi_1 = 2$), the two integers have no common primes. It can be shown (Stillwell, 2010, pp. 41–2) that when $a \perp b$, there exist $m,n \in \mathbb{Z}$ such that

$$am + bn = \text{gcd}(a,b) = 1.$$ 

This equation may be related to the addition of two fractions that have coprime numerators ($a \perp b$). For example,

$$\frac{a}{m} + \frac{b}{n} = \frac{am + bm}{mn}.$$ 

It is not obvious that this is simply $1/mn$.

**Exercise #14**
Show that

$$\begin{bmatrix} 0 & 1 \\ 1 & \frac{m}{n} \end{bmatrix} = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix} \begin{bmatrix} 1 & -1 \\ 0 & 1 \end{bmatrix}.$$ 

**Solution:** This exercise uses the results of an earlier Exercise # 10, times the row swap matrix.

**2.5.4 Continued fraction algorithm**

As shown in Fig. 2.5, the continued fraction algorithm (CFA) starts from a single real decimal number $x_o \in \mathbb{R}$ and recursively expands it as a fraction $x \in \mathbb{F}$ (Graham et al., 1994). Thus the CFA may be used for forming rational approximations to any real number. For example, $\pi \approx 22/7$, an excellent approximation well known to Chinese mathematicians.

The Euclidean algorithm (i.e., GCD), on the other hand, operates on a pair of integers $m, n \in \mathbb{N}$ and returns their greatest common divisor $k \in \mathbb{N}$, such that $m/k, n/k \in \mathbb{F}$ are coprime, thus reducing the ratio to its irreducible form (i.e., $m/k \perp n/k$). Note this is done without factoring $m$ and $n$. 
Despite this seemingly irreconcilable difference between the GCD and CFA, the two are closely related so close that Gauss called the Euclidean algorithm for finding the GCD the continued fraction algorithm (CFA) (Stillwell, 2010, p. 48). At first glance it is not clear why Gauss would be so “confused.” One is forced to assume that Gauss had some deeper insight into this relationship. If so, it would be valuable to understand that insight. Since Eq. 2.5.3.3 may be inverted, the process may be reversed, which is closely related to the CFA as discussed in Fig. 2.5. This might be the basis behind Gauss’s insight.

**Definition of the CFA**

1. Start with \( n = 0 \) and the positive input target \( x_0 \in \mathbb{R}^+ \). 
2. Rounding: Let \( m_n = \lfloor x_n \rfloor \in \mathbb{N} \).
3. The input vector is then \([m_n, x_n]^T\).
4. Remainder: \( r_n = x_n - m_n \) \((-0.5 \leq r_n \leq 0.5)\)
5. Reciprocate:

\[
x_{n+1} = \begin{cases} 
1/r_n, & n \leftarrow n + 1; \text{go to step 2} \\
0, & \text{terminate} 
\end{cases} \quad \begin{align*}
n &\neq 0 \\
r &\neq 0 \quad x_2 = 1/0.14159 = 7.06 \ldots \\
r &\neq 0 \quad r_0 = \pi - 3 \approx 0.1416 \\
Output: & [m_n, x_{n+1}]^T = [3, 7.06]^T 
\end{align*}
\]

**Notation:** Writing out all the fractions can become tedious. For example, expanding \( e = 2.7183 \ldots \) using the Matlab/Octave command `rat(exp(1))` gives the approximation

\[
\exp(1) = 3 + 1/(-4 + 1/(2 + 1/(5 + 1/(-2 + 1/(-7)))))) - o\left(1.75 \times 10^{-6}\right) \\
= [3; -4, 2, 5, -2, -7] - o(1.75 \times 10^{-6}).
\]

Here we use a compact bracket notation, \( \hat{e}_6 \approx [3; -4, 2, 5, -2, -7] \), where \( o() \) indicates the error of the CFA expansion.

Since entries are negative, we deduce that rounding arithmetic is being used by Matlab/Octave (but this is not documented). Note that the leading integer part \( m_0 \) may be indicated by an optional semicolon.\(^{20}\) If the steps are carried further, the values of \( m_n \in \mathbb{Z} \) give increasingly more accurate rational approximations. The five rounding schemes are discussed in Appendix A.1.5.

**Exercise #15**

Let \( x_0 \equiv \pi \approx 3.14159 \ldots \) As shown in Fig. 2.6, \( a_0 = 3 \), \( r_0 = 0.14159 \), \( x_1 = 7.065 \approx 1/r_0 \), and \( a_1 = 7 \). If we were to stop here, we would have

\[
\hat{x}_2 = 3 + \frac{1}{7 + 0.0625} \ldots = 3 + \frac{1}{7} = \frac{22}{7}. 
\]

(2.5.4.4)

This approximation of \( \hat{x}_2 = 22/7 \) has a relative error of 0.04%:

\[
\frac{22/7 - \pi}{\pi} \approx 4 \times 10^{-4}.
\]

**Exercise #16**

For a second level of approximation we continue by reciprocating the remainder \( 1/0.0625 \approx 15.9966 \), which rounds to 16, giving a negative remainder of \( \approx -1/300 \):

\[
\hat{x}_3 \approx 3 + 1/(7 + 1/16) = 3 + 16/(7 \cdot 16 + 1) = 3 + 16/113 = 355/113.
\]

\(^{20}\)Unfortunately Matlab/Octave does not support the bracket notation.
With rounding, the remainder is $-0.0034$, resulting in a much more rapid convergence. If floor rounding is used ($15.9966 = 15 - 0.9966$) the remainder is positive and close to 1, resulting in a much less accurate rational approximation for the same number of terms. It follows that there can be a dramatic difference depending on the rounding scheme, which, for clarity, should be specified, not inferred.

**Rational approximation examples**

$$\hat{\pi}_2 = \frac{22}{7} = [3; 7] \approx \hat{\pi}_2 + o(1.3 \times 10^{-3})$$

$$\hat{\pi}_3 = \frac{355}{113} = [3; 7, 16] \approx \hat{\pi}_3 - o(2.7 \times 10^{-7})$$

$$\hat{\pi}_4 = \frac{104,348}{33,215} = [3; 7, 16, -249] \approx \hat{\pi}_4 + o(3.3 \times 10^{-10})$$

Figure 2.6: The expansion of $\pi$ to various orders, using the CFA, along with the order of the error of each rational approximation, with rounding. For example, $\hat{\pi}_2 = \frac{22}{7}$ has an absolute error ($|\frac{22}{7} - \pi|$) of about 0.13%.

**Exercise #17**

Find the CFA using the floor function, to the 12th order.

**Solution:** $\hat{\pi}_{12} = [3; 7, 15, 1, 292, 1, 1, 2, 1, 3, 1]$. Octave/Matlab will give a different answer due to the use of rounding rather than floor.

**Exercise #18**

Matlab/Octave’s `rat(pi,1e-20)` gives

$$3 + 1/(7 + 1/(16 + 1/(-294 + 1/(3 + 1/(-4 + 1/(5 + 1/(-15 + 1/(-3))))))))$$

In bracket notation,

$$\hat{\pi}_9 = [3; 7, 16, -294, 3, -4, 5, -15, 03].$$

Because the sign changes, it is clear that Matlab/Octave uses rounding rather than the floor function.

**Exercise #19**

Based on the several examples given above, which rounding scheme is the most accurate? Explain why.

**Solution:** Rounding results in a smaller remainder at each iteration and thus results in a smaller net error and faster convergence. Using the floor truncation always gives positive coefficients, which could have some applications.

When the CFA is applied and the expansion terminates ($r_n = 0$), the target is rational. When the expansion does not terminate (which is not always easy to determine, as the remainder may be ill-conditioned due to small numerical rounding errors), the number is irrational. Thus the CFA has important theoretical applications with irrational numbers. You may explore this using Matlab’s `rats(pi)` command.

In addition to these five basic rounding schemes, there are two other important $\mathbb{R} \rightarrow \mathbb{N}$ functions (i.e., mappings) that will be needed later: `mod(x,y)` and `rem(x,y)` with $x, y \in \mathbb{R}$. The base-10 numbers may be generated from the counting numbers using $y=\text{mod}(x,10)$.

**Exercise #20**

1. Show how to generate a base-10 real number $y \in \mathbb{R}$ from the counting numbers $\mathbb{N}$ using the $m=\text{mod}(n,10) + k10$ with $n, k \in \mathbb{N}$.

**Solution:** Every time $n$ reaches a multiple of 10, $m$ is reset to 0 and the next digit to the left is increased by 1 by adding 1 to $k$, generating the digit pair $km$. Thus the `mod()` function forms the underlying theory behind decimal notation.

2. How would you generate binary numbers (base 2) using the $\text{mod}(x,b)$ function? **Solution:** Use the same method as in part (a), but with $b = 2$. 


3. How would you generate hexadecimal numbers (base 16) using the mod\((x,b)\) function? **Solution:** Use the same method as in part (a), but with \(b = 16\).

4. Write the first 19 numbers in hex notation, starting from zero. **Solution:** 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, A, B, C, D, E, F, 10, 11, 12. Recall that 10\(_{16}\) = 16\(_{10}\), thus 12\(_{16}\) = 18\(_{10}\), resulting in a total of 19 numbers if we include 0.

5. What is FF\(_{16}\) in decimal notation? **Solution:** hex2dec\(\left(\text{‘ff’}\right) = 255\(_{10}\)\)

Symmetry: A continued fraction expansion can have a high degree of recursive symmetry. For example, consider the CFA
\[
R_1 \equiv \frac{1 + \sqrt{5}}{2} = 1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \ddots}}} = 1.618033988749895 \ldots
\] (2.5.4.5)

Here \(a_n\) in the CFA is always 1 \((R_1 \equiv [1; 1, \ldots])\), thus the sequence cannot terminate, proving that \(\sqrt{5} \in \mathbb{R}\). A related example is \(R_2 \equiv \text{rat}\left(1 + \sqrt{2}\right)\), which gives \(R_2 = [2; 2, 2, 2, \ldots]\).

When we expand a target irrational number \((x_0 \in \mathbb{R}\) and the CFA is truncated, the resulting rational fraction approximates the irrational target. For the example above, if we truncate at three coefficients \(([1; 1, 1])\), we obtain
\[
1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \ddots}}}}}}}}}} = 1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \ddots}}}}}}}}}} = 1.5 = \frac{3}{2} + 0.118 \ldots
\]

Truncation after six steps gives
\[
[1., 1, 1, 1, 1, 1, 1] = 13/8 \approx 1.6250 = \frac{1 + \sqrt{5}}{2} + 0.0070 \ldots
\]

Because all the coefficients are 1, this example converges very slowly. When the coefficients are large (i.e., the remainder is small), the convergence will be faster. The expansion of \(\pi\) is an example of faster convergence.

**In summary:** Every rational number \(m/n \in \mathbb{R}\), with \(m > n > 1\), may be uniquely expanded as a continued fraction, with coefficients \(a_n\) determined using the CFA. When the target number is irrational \((x_0 \in \mathbb{Q}\), the CFA does not terminate; each step produces a more accurate rational approximation, converging in the limit as \(n \to \infty\).

Thus the CFA expansion is an algorithm that can, in theory, determine when the target is rational, but with an important caveat: One must determine whether the expansion terminates. This may not be obvious. The fraction \(1/3 = 0.33333 \ldots\) is an example of such a target, where the CFA terminates yet the fraction repeats. It must be that
\[
1/3 = 3 \times 10^{-1} + 3 \times 10^{-2} + 3 \times 10^{-3} + \ldots
\]

Here \(3 \cdot 3 = 9\). As a second example,\(^21\)
\[
1/7 = 0.142857, 142857, 142857, 142857 \ldots = 14, 2857 \times 10^{-6} + 142, 857 \times 10^{-12} + \ldots
\]

There are several notations for repeating decimals, such as \(1/7 = 0.142857\) and \(1/7 = 0.1(142857)\). Note that 142, 857 = 999, 999/7. Related identities include 1/11 = 0.090909... and 11 \times 0.090909 = 999, 999. When the sequence of digits repeats, the sequence is predictable and it must be rational. But it is impossible to be sure that it repeats because the length of the repeat can be arbitrarily long.

**Exercise #21**

Discuss the relationship between the CFA and the transmission line modeling method on page 110.

**Solution:** The solution is detailed in Appendix E.

### 2.5.5 Pythagorean triplets (Euclid’s formula)

Euclid’s formula is a method for finding three integer lengths \([a, b, c] \in \mathbb{N}\) that satisfy Eq. 1.1.1.1 (p. 12). It is important to ask Which set are the lengths \([a, b, c]\) drawn from? There is a huge difference, both practical and theoretical, if they are from the real numbers \(\mathbb{R}\) or from the counting numbers \(\mathbb{N}\). Given \(p, q \in \mathbb{N}\) with \(p > q\), the three lengths \([a, b, c] \in \mathbb{N}\) of Eq. 1.1.1.1 are given by
\[
a = p^2 - q^2, \quad b = 2pq, \quad c = p^2 + q^2.
\] (2.5.5.6)
2.5. APPLICATIONS OF PRIME NUMBERS

This result may be directly verified, since

\[ (p^2 + q^2)^2 = (p^2 - q^2)^2 + (2pq)^2 \]

or

\[ p^4 + q^4 + 2pq^2 = p^4 + q^4 - 2pq^2 + 4p^2q^2. \]

Thus, Eqs. 2.5.5.6 are easily proved once given. Deriving Euclid’s formula (See AE-2, problem #2) is obviously much more difficult, and is similar to the proof of Pell’s equation.

A well-known example is the right triangle depicted in Fig. 2.7, defined by the integer lengths \([3, 4, 5]\) that have angles \([0.54, 0.65, \pi/2]\) [rad], which satisfies Eq. 1.1.1.1. As quantified by Euclid’s formula (Eqs. 2.5.5.6), there are an infinite number of Pythagorean triplets (PTs). Furthermore, the seemingly simple triangle that has angles of \([30, 60, 90] \in N\) [deg] (i.e., \([\pi/6, \pi/3, \pi/2] \in I\) [rad]) has one irrational (i) length \((1, \sqrt{3}, 2)\).

<table>
<thead>
<tr>
<th>(a)</th>
<th>(c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>119</td>
<td>169</td>
</tr>
<tr>
<td>3367</td>
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<td>106</td>
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</tbody>
</table>

The set from which the lengths \([a, b, c]\) are drawn was not missed by the early Asians and was documented by the Greeks. Any equation whose solution is based on integers is called a Diophantine equation, named for the Greek mathematician Diophantus of Alexandria (ca. 250 CE) (see Fig. 1.1, p. 13).

A stone tablet from the 19th century BCE with the numbers engraved on it, as shown in Fig. 2.8, was discovered in Mesopotamia, and cataloged in 1922 by George Plimpton. These numbers are \(a\) and \(c\) pairs from PTs \([a, b, c]\).

Given this discovery, it is clear that the Pythagoreans were following those who came long before them. Recently a second stone, dating between 350 and 50 BCE, has been reported, that indicates early calculations on the orbit of Jupiter’s moons.\(^{23}\)

\(^{21}\)Taking the Fourier transform of the target number, represented as a sequence, could help to identify an underlying periodic component. The number 1/7 ↔ \([1, 4, 2, 8, 5, 7]_8\) has a 50 (dB) notch at 0.8π [rad] due to its six-digit periodicity, carried to 15 digits (Matlab/Octave maximum precision), Hamming windowed, and zero padded to 1024 samples.


Table 2.1: Table of Pythagorean triplets computed from Euclid’s formula, Eq. 2.5.6.6, for various \([p, q]\). The last three columns are the first, fourth, and penultimate values of Plimpton-322, along with their corresponding \([p, q]\). In all cases \(c^2 = a^2 + b^2\) and \(p = q + l\), where \(l = \sqrt{c-b} \in \mathbb{N}\).

<table>
<thead>
<tr>
<th>(q)</th>
<th>(l)</th>
<th>(a)</th>
<th>(b)</th>
<th>(c)</th>
<th>(p)</th>
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<th>(2)</th>
<th>(3)</th>
<th>(4)</th>
<th>(5)</th>
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<td>5</td>
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<td>45</td>
<td>169</td>
<td>18541</td>
<td>3229</td>
</tr>
</tbody>
</table>

It is of interest that PTs play a role on atomic physics, as discussed in Appendix H.

2.5.6 Pell’s equation

Pell’s equation,

\[
x_n^2 - N y_n^2 = (x_n - \sqrt{N} y_n)(x_n + \sqrt{N} y_n) = 1,
\]

(2.5.6.7)

with nonsquare \(N \in \mathbb{N}\) specified and \(x, y \in \mathbb{N}\) unknown, has a venerable history in both physics (see p. 23) and mathematics. Given its factored form, it is obvious that every solution \(x_n, y_n\) has the asymptotic property

\[
\left| \frac{x_n}{y_n} \right| \rightarrow \pm \sqrt{N}.
\]

(2.5.6.8)

It is believed that Pell’s equation is directly related to the Pythagorean theorem, since both are simple binomials that have integer coefficients (Stillwell, 2010, p. 48), with Pell’s equation being the hyperbolic version of Eq. 1.1.1.1. For example, with \(N = 2\), a solution is \(x = 17, y = 12\) (i.e., \(17^2 - 2 \cdot 12^2 = 1\)).

A \(2 \times 2\) matrix recursion algorithm, likely due to the Chinese and used by the Pythagoreans to investigate \(\sqrt{N}\), is

\[
\begin{bmatrix}
  x \\ y
\end{bmatrix}_{n+1} = \begin{bmatrix}
  1 & N \\
  1 & 1
\end{bmatrix} \begin{bmatrix}
  x \\ y
\end{bmatrix}_n,
\]

(2.5.6.9)

where we indicate the index outside the vectors.

Starting with the trivial solution \([x_0, y_0]^T = [1, 0]^T\) (i.e., \(x_0^2 - N y_0^2 = 1\)), additional solutions of Pell’s equations are determined, having the property \(x_n/y_n \rightarrow \sqrt{N} \in \mathbb{P}\), motivated by Euclid’s formula for Pythagorean triplets (Stillwell, 2010, p. 44). Note that Eq. 2.5.6.9 is a \(2 \times 2\) linear matrix composition method (see p. 91), since the output of one matrix multiplication is the input to the next.

Asian solutions: The first solution of Pell’s equation was published in about 628 CE by Brahmagupta, who independently discovered the equation (Stillwell, 2010, p. 46). Brahmagupta’s novel solution also used the composition method, but in a different way from Eq. 2.5.6.9. Then in 1150 CE, Bhaskara II independently obtained solutions using Eq. 2.5.6.9 (Stillwell, 2010, p.69). This is the composition method we shall explore here, as summarized in Appendix B, Table B.1.

The best way to see how this recursion results in solutions to Pell’s equation is by example. Initializing the recursion with the trivial solution \([x_0, y_0]^T = [1, 0]^T\) gives

\[
\begin{bmatrix}
  x_1 \\ y_1
\end{bmatrix} = \begin{bmatrix}
  1 & 2 \\
  1 & 1
\end{bmatrix}_0 \begin{bmatrix}
  1 \\ 0
\end{bmatrix} = 1^2 - 2 \cdot 1^2 = -1
\]

\[
\begin{bmatrix}
  x_2 \\ y_2
\end{bmatrix} = \begin{bmatrix}
  3 & 2 \\
  2 & 1
\end{bmatrix}_1 \begin{bmatrix}
  1 \\ 0
\end{bmatrix} = 3^2 - 2 \cdot 2^2 = 1
\]

\[
\begin{bmatrix}
  x_3 \\ y_3
\end{bmatrix} = \begin{bmatrix}
  7 & 3 \\
  5 & 2
\end{bmatrix}_2 \begin{bmatrix}
  1 \\ 0
\end{bmatrix} = (7)^2 - 2 \cdot (5)^2 = -1
\]

\[
\begin{bmatrix}
  x_4 \\ y_4
\end{bmatrix} = \begin{bmatrix}
  17 & 7 \\
  12 & 5
\end{bmatrix}_3 \begin{bmatrix}
  1 \\ 0
\end{bmatrix} = 17^2 - 2 \cdot 12^2 = 1
\]

\[
\begin{bmatrix}
  x_5 \\ y_5
\end{bmatrix} = \begin{bmatrix}
  41 & 17 \\
  29 & 12
\end{bmatrix}_4 \begin{bmatrix}
  1 \\ 0
\end{bmatrix} = (41)^2 - 2 \cdot (29)^2 = -1
\]
Thus the recursion results in a modified version of Pell’s equation,

$$x_n^2 - 2y_n^2 = (-1)^n,$$  \hspace{1cm} (2.5.6.10)

where only even values of \( n \) are solutions. This sign change had no effect on the Pythagoreans’ goal, since they cared about only the ratio \( y_n/x_n \rightarrow \pm \sqrt{2} \).

**Modified recursion:** We may restore the solution of Pell’s equation for \( N = 2 \) using a slightly modified linear matrix recursion. To fix the \((-1)^n\) problem, we multiply the 2 \( \times \) 2 matrix by \( 1_j = \sqrt{-1} \), which gives

\[
\begin{align*}
\begin{bmatrix} x \\ y \end{bmatrix}_1 &= J \begin{bmatrix} 1 \\ 0 \end{bmatrix} = J \begin{bmatrix} 1 & 0 \\ 1 & 1 \end{bmatrix} \begin{bmatrix} 1 \\ 0 \end{bmatrix} = 1 \\
\begin{bmatrix} x \\ y \end{bmatrix}_2 &= J^2 \begin{bmatrix} 3 \\ 2 \end{bmatrix} = J \begin{bmatrix} 1 & 0 \\ 1 & 1 \end{bmatrix} J \begin{bmatrix} 1 \\ 0 \end{bmatrix} = 3 \\
\begin{bmatrix} x \\ y \end{bmatrix}_3 &= J^3 \begin{bmatrix} 7 \\ 5 \end{bmatrix} = J \begin{bmatrix} 1 & 0 \\ 1 & 1 \end{bmatrix} J^2 \begin{bmatrix} 3 \\ 2 \end{bmatrix} = 7 \\
\begin{bmatrix} x \\ y \end{bmatrix}_4 &= J^4 \begin{bmatrix} 17 \\ 12 \end{bmatrix} = J \begin{bmatrix} 1 & 0 \\ 1 & 1 \end{bmatrix} J^3 \begin{bmatrix} 7 \\ 5 \end{bmatrix} = 17 \\
\begin{bmatrix} x \\ y \end{bmatrix}_5 &= J^5 \begin{bmatrix} 41 \\ 29 \end{bmatrix} = J \begin{bmatrix} 1 & 0 \\ 1 & 1 \end{bmatrix} J^4 \begin{bmatrix} 17 \\ 12 \end{bmatrix} = 41.
\end{align*}
\]

**Solution to Pell’s equation:** By multiplying the matrix by \( 1_j \), all the solutions (\( x_k \in \mathbb{C} \)) to Pell’s equation are determined. The \( 1_j \) factor corrects the alternation in sign, so every iteration yields a solution. For \( N = 2, n = 0 \) (the initial solution), \([x_0, y_0]\) is \([1, 0]\), \([x_1, y_1]\) is \([1, 1]\), and \([x_2, y_2]\) is \([-3, 2]\). These are easily checked using this recursion.

The solution for \( N = 3 \) is given in Appendix B.2.1 Table. B.1 (page 236) shows that every output of this slightly modified matrix recursion gives solutions to Pell’s equation: \([1, 0], [1, 1], [4, 2], [10, 6], \ldots, [76, 44], \ldots\).

At each iteration, the ratio \( x_n/y_n \) approaches \( \sqrt{2} \) with increasing accuracy, coupling it to the CFA, which may also be used to find approximations to \( \sqrt{N} \). The value of \( 41/29 \approx \sqrt{2} \), with a relative error of <0.03%.

### 2.5.7 Fibonacci sequence

Another classic problem, also formulated by the Chinese, is the Fibonacci sequence, generated by the relationship

\[ f_{n+1} = f_n + f_{n-1}. \] \hspace{1cm} (2.5.7.11)

Here the next number \( f_{n+1} \in \mathbb{N} \) is the sum of the previous two. If we start from \([0, 1]\), this linear recursion equation leads to the Fibonacci sequence \( f_n = [0, 1, 1, 2, 3, 5, 8, 13, 21, 34, \ldots] \). Alternatively, if we define \( y_{n+1} = x_n \), then Eq. 2.5.7.11 may be compactly represented by a 2 \( \times \) 2 companion matrix recursion (see the Fibonacci exercises NS-3).

\[
\begin{bmatrix} x \\ y \end{bmatrix}_{n+1} = \begin{bmatrix} 1 & 1 \\ 1 & 0 \end{bmatrix} \begin{bmatrix} x \\ y \end{bmatrix}_n,
\]

which has eigenvalues \((1 \pm \sqrt{5})/2\).

The correspondence of Eqs. 2.5.7.11 and 2.5.7.12 is easily verified. Starting with \([x, y]_0^T = [0, 1]^T\), we obtain for the first few steps:

\[
\begin{align*}
\begin{bmatrix} 1 \\ 0 \end{bmatrix}_1 &= \begin{bmatrix} 1 & 1 \\ 1 & 0 \end{bmatrix} \begin{bmatrix} 1 \\ 0 \end{bmatrix}_0, & \begin{bmatrix} 1 \\ 1 \end{bmatrix}_1 &= \begin{bmatrix} 1 & 1 \\ 1 & 0 \end{bmatrix} \begin{bmatrix} 1 \\ 1 \end{bmatrix}_0, & \begin{bmatrix} 2 \\ 1 \end{bmatrix}_2 &= \begin{bmatrix} 1 & 1 \\ 1 & 0 \end{bmatrix} \begin{bmatrix} 1 \\ 1 \end{bmatrix}_1, & \begin{bmatrix} 3 \\ 2 \end{bmatrix}_3 &= \begin{bmatrix} 1 & 1 \\ 1 & 0 \end{bmatrix} \begin{bmatrix} 2 \\ 1 \end{bmatrix}_2, & \cdots
\end{align*}
\]

From the above, \( x_n = [0, 1, 1, 2, 3, 5, \ldots] \) is the Fibonacci sequence, since the next \( x_n \) is the sum of the previous two, and the next \( y_n \) is \( x_n \).

**Exercise #22**

Use the Octave/Matlab command `compan(c)` to find the companion matrix of the polynomial coefficients defined by Eq. 2.5.7.11.

**Solution:** Using Matlab/Octave: \( f=[1, -1, -1]; C=compan(f); \) returning

\[
C = \begin{bmatrix} 1 & 1 \\ 1 & 0 \end{bmatrix}
\] \hspace{1cm} (2.5.7.13)
Exercise #23
Find the eigenvalues of matrix \( C \).

Solution: The characteristic equation is

\[
\det \begin{bmatrix} 1 - \lambda & 1 \\ 1 & -\lambda \end{bmatrix} = 0
\]

or \( \lambda^2 - \lambda - 1 = (\lambda - 1/2)^2 - 1/4 - 1 = 0 \), which has roots \( \lambda_{\pm} = (1 \pm \sqrt{5})/2 \approx \{1.618, -0.618\} \).

The mean-Fibonacci sequence: Suppose that the Fibonacci sequence recursion is replaced by the mean of the last two values–namely, let

\[
f_{n+1} = \frac{f_n + f_{n-1}}{2}.
\]

(2.5.7.14)

This seems like a small change. But how does the solution differ? To answer this question it is helpful to look at the corresponding \( 2 \times 2 \) matrix.

Exercise #24
Find the \( 2 \times 2 \) matrix corresponding to Eq. 2.5.7.14. The \( 2 \times 2 \) matrix may be found using the companion matrix method (see p. 71).

Solution: Using Matlab/Octave code, we have

```matlab
f=[1, -1/2, -1/2];
C=companion(f);
```

which returns

\[
C = \frac{1}{2} \begin{bmatrix} 1 & 1 \\ 2 & 0 \end{bmatrix}.
\]

(2.5.7.15)

Exercise #25
Find the steady-state solution for the mean-Fibonacci, starting from \( [1, 0]^T \). State the nature of both solutions.

Solution: By inspection one steady-state solution is \( [1, 1]^T \) or \( f_n = 1^n \). To find the full solution, we need to find the two eigenvalues, defined by

\[
\det \begin{bmatrix} 1/2 - \lambda & 1/2 \\ 1 & -\lambda \end{bmatrix} = \lambda^2 - \lambda/2 - 1/2 = (\lambda - 1/4)^2 - (1/4)^2 - 1/2 = 0.
\]

Thus \( \lambda_{\pm} = (1 \pm 3)/4 = \{1, -0.5\} \). The first solution converges to 1 while second solution is \((-1/2)^n\), which changes sign at each time step and quickly converges to zero. The full solution is given by \( E\Lambda^n E^{-1}[1, 0]^T \) (see Appendix B, p. 233).

■
2.5. APPLICATIONS OF PRIME NUMBERS

Relationships to digital signal processing: Today we recognize Eq. 2.5.7.11 as a discrete difference equation, which is a pre-limit (pre-Stream 3) recursive form of a differential equation. The $2 \times 2$ matrix form of Eq. 2.5.7.11 is an early precursor to 17th- and 18th-century developments in linear algebra. Thus the Greeks’ recursive solution for the $\sqrt{2}$ and Bhaskara’s solution of Pell’s equation are early precursors to discrete-time signal processing as well as to calculus.

There are strong similarities between Pell’s equation and the Pythagorean theorem. As we shall see in Sec. 2, Pell’s equation is related to the geometry of a hyperbola, just as the Pythagorean equation is related to the geometry of a circle. We shall show, as one might assume, that there is a Euclid’s formula for the case of Pell’s equations, since these are all conic sections with closely related conic geometry. As we have seen, the solutions involve $\sqrt{-1}$. The derivation is a trivial extension of that for Euclid’s formula for Pythagorean triplets. The early solution of Brahmagupta was not related to this simple formula.
2.6 Problems NS-3

**Topic of this homework:** Pythagorean triplets, Pell’s equation, Fibonacci sequence

**Deliverables:** Answers to problems

**Pythagorean triplets**

**Problem #1:** Euclid’s formula for the Pythagorean triplets \( a, b, c \) is \( a = p^2 - q^2, \) \( b = 2pq, \) and \( c = p^2 + q^2. \)

- 1.1: What condition(s) must hold for \( p \) and \( q \) such that \( a, b, \) and \( c \) are always positive and nonzero?

- 1.2: Solve for \( p \) and \( q \) in terms of \( a, b, \) and \( c. \)

**Problem #2:** The ancient Babylonians (ca. 2000 BCE) cryptically recorded \((a, c)\) pairs of numbers on a clay tablet, archeologically denoted Plimpton-322 (see 2.8).

- 2.1: Find \( p \) and \( q \) for the first five pairs of \( a \) and \( c \) shown here from Plimpton-322.

\[
\begin{array}{c|c}
 a & c \\
\hline
119 & 169 \\
3367 & 4825 \\
4601 & 6649 \\
12709 & 18541 \\
65 & 97
\end{array}
\]

Find a formula for \( a \) in terms of \( p \) and \( q. \)

- 2.2: Based on Euclid’s formula, show that \( c > (a, b). \)

- 2.3: What happens when \( c = a? \)

- 2.4: Is \( b + c \) a perfect square? Discuss.

**Pell’s equation:**

**Problem #3:** Pell’s equation is one of the most historic (i.e., important) equations of Greek number theory because it was used to show that \( \sqrt{2} \in \mathbb{Q}. \) We seek integer solutions of

\[ x^2 - Ny^2 = 1. \]

As shown on page 52, the solutions \( x_n, y_n \) for the case of \( N = 2 \) are given by the linear \( 2 \times 2 \) matrix recursion

\[
\begin{bmatrix}
x_{n+1} \\
y_{n+1}
\end{bmatrix} = M
\begin{bmatrix}
x_n \\
y_n
\end{bmatrix}
\]

where

\[
M = \begin{bmatrix} 1 & 2 \\ 1 & 1 \end{bmatrix}
\]
with \([x_0, y_0]^T = [1, 0]^T\) and \(1_j = \sqrt{-1} = e^{j\pi/2}\). It follows that the general solution to Pell’s equation for \(N = 2\) is
\[
\begin{bmatrix}
x_n \\
y_n
\end{bmatrix} = (e^{j\pi/2})^n \begin{bmatrix} 1 & 2 \\ 1 & 1
\end{bmatrix} \begin{bmatrix} x_0 \\
y_0
\end{bmatrix}.
\]

To calculate solutions to Pell’s equation using the matrix equation above, we must calculate
\[
A^n = e^{j\pi n/2} \begin{bmatrix} 1 & 2 \\ 1 & 1
\end{bmatrix} = e^{j\pi n/2} \begin{bmatrix} 1 & 2 \\ 1 & 1
\end{bmatrix} \begin{bmatrix} 1 & 2 \\ 1 & 1
\end{bmatrix} \cdots \begin{bmatrix} 1 & 2 \\ 1 & 1
\end{bmatrix},
\]
which becomes tedious for \(n > 2\).

Diagonalization of a matrix (eigenvalue/eigenvector decomposition):

As derived in Appendix B, the most efficient way to compute \(A^n\) is to diagonalize the matrix \(A\) by finding its eigenvalues and eigenvectors.

The eigenvalues \(\lambda_k\) and eigenvectors \(\vec{e}_k\) of a square matrix \(A\) are related by
\[
A\vec{e}_k = \lambda_k \vec{e}_k, \quad (\text{NS-3 2.1})
\]
such that multiplying an eigenvector \(\vec{e}_k\) of \(A\) by the matrix \(A\) is the same as multiplying by a scalar, \(\lambda_k \in \mathbb{C}\) (the corresponding eigenvalue). The complete eigenvalue problem may be written as
\[
AE = \Lambda E.
\]
If \(A\) is a \(2 \times 2\) matrix,\(^{24}\) the matrices \(E\) and \(\Lambda\) (of eigenvectors and eigenvalues, respectively) are
\[
E = \begin{bmatrix} \vec{e}_1 & \vec{e}_2 \end{bmatrix} \quad \text{and} \quad \Lambda = \begin{bmatrix} \lambda_1 & 0 \\ 0 & \lambda_2 \end{bmatrix}.
\]
Thus the matrix equation \(AE = \begin{bmatrix} A\vec{e}_1 & A\vec{e}_2 \end{bmatrix} = \begin{bmatrix} \lambda_1 \vec{e}_1 & \lambda_2 \vec{e}_2 \end{bmatrix} = E \Lambda\) contains Eq. NS-3 1.1 for each eigenvalue-eigenvector pair.

The diagonalization of the matrix \(A\) refers to the fact that the matrix of eigenvalues, \(\Lambda\), has nonzero elements only on the diagonal. The key result is found by postmultiplication of the eigenvalue matrix by \(E^{-1}\), giving
\[
AEE^{-1} = A = E\Lambda E^{-1}. \quad (\text{NS-3 2.2})
\]
If we now take powers of \(A\), the \(n\)th power of \(A\) is
\[
A^n = (A\Lambda E^{-1})^n = E\Lambda E^{-1} E\Lambda E^{-1} \cdots E\Lambda E^{-1} = E\Lambda^n E^{-1}. \quad (\text{NS-3 2.3})
\]
This is a very powerful result because the \(n\)th power of a diagonal matrix is extremely easy to calculate:
\[
\Lambda^n = \begin{bmatrix} \lambda_1^n & 0 \\ 0 & \lambda_2^n \end{bmatrix}.
\]
Thus, from Eq. NS-3 1.3 we can calculate \(A^n\) using only two matrix multiplications:
\[
A^n = E\Lambda^n E^{-1}.
\]

Finding the eigenvalues:

The eigenvalues \(\lambda_k\) are determined from Eq. NS-3 1.1, by factoring out \(\vec{e}_k\):
\[
A\vec{e}_k = \lambda_k \vec{e}_k \quad (A - \lambda_k I)\vec{e}_k = 0.
\]
Matrix \(I = [1, 0; 0, 1]^T\) is the identity matrix, having the dimensions of \(A\), with elements \(\delta_{ij}\) (i.e., diagonal elements \(\delta_{11,22} = 1\) and off-diagonal elements \(\delta_{12,21} = 0\)).

\(^{24}\)These concepts may be easily extended to higher dimensions.
The vector \( \vec{e}_k \) is not zero, yet when operated on by \( A - \lambda I \), the result must be zero. The only way this can happen is if the operator is degenerate (has no solution)—that is,

\[
\det(A - \lambda I) = \det \begin{bmatrix} a_{11} - \lambda & a_{12} \\ a_{21} & a_{22} - \lambda \end{bmatrix} = 0. \tag{NS-3 2.4}
\]

This means that the two equations have the same roots (the equation is degenerate).

This determinant equation results in a second-degree polynomial in \( \lambda \):

\[
(a_{11} - \lambda)(a_{22} - \lambda) - a_{12}a_{21} = 0,
\]

the roots of which are the eigenvalues of the matrix \( A \).

**Finding the eigenvectors:**

An eigenvector \( \vec{e}_k \) can be found for each eigenvalue \( \lambda_k \) from Eq. NS-3 1.1,

\[
(A - \lambda_k I)\vec{e}_k = \vec{0}.
\]

The left side of the above equation becomes a column vector, where each element is an equation in the elements of \( \vec{e}_k \), set equal to 0 on the right side. These equations are always degenerate, since the determinant is zero. Thus the two equations have the same slope.

Solving for the eigenvectors is often confusing because they have arbitrary magnitudes, \( ||\vec{e}_k|| = \sqrt{\vec{e}_k \cdot \vec{e}_k} = \sqrt{\vec{e}_{k,1}^2 + \vec{e}_{k,2}^2} = d \). From Eq. NS-3 1.1, we can determine only the relative magnitudes and signs of the elements of \( \vec{e}_k \), so we have to choose a magnitude \( d \). It is common practice to normalize each eigenvector to have unit magnitude (\( d = 1 \)).

– 3.1: Find the companion matrix and thus the matrix \( A \) that has the same eigenvalues as Pell’s equation. Hint: Use Matlab’s function \([E, \text{Lambda}] = \text{eig}(A)\) to check your results!

– 3.2: Solutions to Pell’s equation were used by the Pythagoreans to explore the value of \( \sqrt{2} \). Explain why Pell’s equation is relevant to \( \sqrt{2} \).

– 3.3: Find the first three values of \((x_n, y_n)^T\) by hand and show that they satisfy Pell’s equation for \( N = 2 \). By hand, find the eigenvalues \( \lambda_{\pm} \) of the \( 2 \times 2 \) Pell’s equation matrix

\[
A = \begin{bmatrix} 1 & 2 \\ 1 & 1 \end{bmatrix}.
\]

– 3.4: By hand, show that the matrix of eigenvectors, \( E \), is

\[
E = \begin{bmatrix} \vec{e}_+ & \vec{e}_- \end{bmatrix} = \frac{1}{\sqrt{3}} \begin{bmatrix} -\sqrt{2} & 1 \\ 1 & \sqrt{2} \end{bmatrix}.
\]

– 3.5: Using the eigenvalues and eigenvectors you found for \( A \), verify that

\[
E^{-1}AE = \Lambda \equiv \begin{bmatrix} \lambda_+ & 0 \\ 0 & \lambda_- \end{bmatrix}
\]

– 3.6: Now that you have diagonalized \( A \) (Equation NS-3 1.3), use your results for \( E \) and \( \Lambda \) to solve for the \( n = 10 \) solution \((x_{10}, y_{10})^T\) to Pell’s equation with \( N = 2 \).

**The Fibonacci sequence**

The Fibonacci sequence is famous in mathematics and has been observed to play a role in the mathematics of genetics. Let \( x_n \) represent the Fibonacci sequence,

\[
x_{n+1} = x_n + x_{n-1}, \tag{NS-3 2.5}
\]
where the current input sample \( x_n \) is equal to the sum of the previous two inputs. This is a “discrete time” recurrence relationship. To solve for \( x_n \), we require some initial conditions. In this exercise, let us define \( x_0 = 1 \) and \( x_{n<0} = 0 \). This leads to the Fibonacci sequence \{1, 1, 2, 3, 5, 8, 13, \ldots\} for \( n = 0, 1, 2, 3, \ldots \).

Equation NS-3 1.5 is equivalent to the \( 2 \times 2 \) matrix equations

\[
\begin{bmatrix} x_n \\ y_n \end{bmatrix} = A \begin{bmatrix} x_{n-1} \\ y_{n-1} \end{bmatrix}, \quad A = \begin{bmatrix} 1 & 1 \\ 1 & 0 \end{bmatrix}.
\] (NS-3 2.6)

**Problem # 4:** Here we seek the general formula for \( x_n \). Like Pell’s equation, Eq. NS-3 1.5 has a recursive, eigenanalysis solution. To find it we must recast \( x_n \) as a \( 2 \times 2 \) matrix relationship and then proceed as we did for the Pell case.

- **4.1:** By example, show that the Fibonacci sequence \( x_n \) as described above may be generated by

\[
\begin{bmatrix} x_n \\ y_n \end{bmatrix} = A^n \begin{bmatrix} x_0 \\ y_0 \end{bmatrix}, \quad A = \begin{bmatrix} 1 & 1 \\ 1 & 0 \end{bmatrix}.
\] (NS-3 2.7)

- **4.2:** What is the relationship between \( y_n \) and \( x_n \)?

- **4.3:** Write a Matlab/Octave program to compute \( x_n \) using the matrix equation above. Test your code using the first few values of the sequence. Using your program, what is \( x_{40} \)? Note: Consider using the eigenanalysis of \( A \), described by Eq. NS-3 1.3 (p. 16).

- **4.4:** Using the eigenanalysis of the matrix \( A \) (and a lot of algebra), show that it is possible to obtain the general formula for the Fibonacci sequence

\[
x_n = \frac{1}{\sqrt{5}} \left[ \left( \frac{1 + \sqrt{5}}{2} \right)^{n+1} - \left( \frac{1 - \sqrt{5}}{2} \right)^{n+1} \right].
\] (NS-3 2.8)

- **4.5:** What are the eigenvalues \( \lambda_\pm \) of the matrix \( A \)?

- **4.6:** How is the formula for \( x_n \) related to these eigenvalues? Hint: Find the eigenvectors.

- **4.7:** What happens to each of the two terms

\[
[(1 \pm \sqrt{5})/2]^{n+1}?
\]

- **4.8:** What happens to the ratio \( x_{n+1}/x_n \)?

**Problem # 5:** Replace the Fibonacci sequence \( x_n \) such that the value \( x_n \) is the average of the previous two values in the sequence.

- **5.1:** What matrix \( A \) is used to calculate this sequence?

- **5.2:** Modify your computer program to calculate the new sequence \( x_n \). What happens as \( n \to \infty \)?

- **5.3:** What are the eigenvalues of your new \( A \)? How do they relate to the behavior of \( x_n \) as \( n \to \infty \)? Hint: You can expect the closed-form expression for \( x_n \) to be similar to Eq. NS-3 1.8.
– 5.4: What matrix $A$ is used to calculate this sequence?

– 5.5: Modify your computer program to calculate the new sequence $x_n$. What happens as $n \to \infty$?

– 5.6: What are the eigenvalues of your new $A$? How do they relate to the behavior of $x_n$ as $n \to \infty$? Hint: You can expect the closed-form expression for $x_n$ to be similar to Eq. NS-3 1.8.

**Problem # 6:** Consider the expression

$$\sum_{n=1}^{N} f_n^2 = f_N f_{N+1}.$$  

– 6.1: Find a formula for $f_n$ that satisfies this relationship. Hint: It only holds for the Fibonacci recursion formula.

**CFA as a matrix recursion**

**Problem # 7:** The CFA may be written as a matrix recursion. For this we adopt a special notation, unlike other matrix notations,\(^{25}\) with $k \in \mathbb{N}$:

$$\begin{bmatrix} n \\ x_{k+1} \end{bmatrix} = \begin{bmatrix} 0 & [x_k] \\ 0 & x_k - [x_k] \end{bmatrix} \begin{bmatrix} n \\ x_k \end{bmatrix}.$$  

This equation says that $n_{k+1} = [x_k]$ and $x_{k+1} = 1/(x_k - [x_k])$. It does not mean that $n_{k+1} = [x_k] x_k$, as would be implied by standard matrix notation. The lower equation says that $r_k = x_k - [x_k]$ is the remainder—namely, $x_k = [x - k] + r_k$ (Octave/Matlab’s `rem(x, floor(x))` function), also known as `mod(x, y)`.

– 7.1: Start with $n_0 = 0 \in \mathbb{N}$, $x_0 \in \mathbb{I}$, $n_1 = [x_0] \in \mathbb{N}$, $r_1 = x - [x] \in \mathbb{I}$, and $x_1 = 1/r_1 \in \mathbb{I}$, $r_n \neq 0$. For $k = 1$ this generates on the left the next CFA parameter $n_2 = [x_1]$ and $x_2 = 1/r_2 = 1/(x_0 - [x_0])$ from $n_0$ and $x_0$. Find $[n, x]_{k+1}^T$ for $k = 2, 3, 4, 5$.

---

\(^{25}\)This notation is highly nonstandard due to the nonlinear operations. The matrix elements are derived from the vector rather than multiplying them. These calculation may be done with the help of Matlab/Octave.
Chapter 3

Algebraic Equations: Stream 2

3.1 The physics behind nonlinear Algebra (Euclidean geometry)

Stream 2 is geometry, which led to the merging of Euclid’s geometrical methods and the development of algebra by al-Khwarizmi 830 CE (Fig. 1.1, p. 13). This migration of ideas led Descartes and Fermat to develop analytic geometry (Fig. 1.2, p. 15).

The mathematics up to the time of the Greeks, documented and formalized by Euclid, served students of mathematics for more than two thousand years. Algebra and geometry were, at first, independent lines of thought. When merged, the focus returned to the Pythagorean theorem. Algebra generalized the analytic conic section into the complex plane, greatly extending the geometrical approach as taught in Euclid’s *Elements*. With the introduction of algebra, numbers, rather than lines, could be used to represent geometrical lengths in the complex plane. Thus the appreciation for geometry grew after the addition of rigorous analysis using numbers.

<table>
<thead>
<tr>
<th>History of Mathematics after the 15th Century</th>
</tr>
</thead>
<tbody>
<tr>
<td>16th Bombelli 1526–1572; Galileo 1564–1642; Kepler 1571–1630; Mersenne 1588–1648;</td>
</tr>
<tr>
<td>17th Huygens 1629–1695; Newton 1642–1727, <em>Principia</em> 1687; Bernoulli, Jakob 1655–1705; Bernoulli, Johann 1667–1748; Fermat, Pierre de 1607–1665; Pascal, Blaise 1623–1662; Descartes 1596–1648</td>
</tr>
<tr>
<td>18th Bernoulli, Daniel 1700–1782; Euler 1707–1783; d’Alembert 1717–1783; Lagrange 1736–1833; Laplace 1749–1827; Fourier 1768–1830; Gauss 1777–1855; Cauchy 1789–1857</td>
</tr>
<tr>
<td>20th Bode, H. 1905–1982</td>
</tr>
</tbody>
</table>

Physics inspires algebraic mathematics: The Chinese used music, art, and navigation to drive mathematics. Much of their knowledge has been handed down as either artifacts, such as musical bells and tools, or mathematical relationships documented, but not created, by scholars such as Euclid, Archimedes, Diophantus, and perhaps Brahmagupta. With the invention of algebra in 830 CE by al-Khwarizmi, mathematics became more powerful and blossomed. During the 16th and 17th centuries, it became clear that differential equations (DEs), such as the wave equation, can characterize a law of nature at a single point in space and time. This principle was not obvious. A desire to understand the motions of objects and planets precipitated many new discoveries. This period, centered around Galileo, Newton, and Euler, is shown on the timeline in Fig. 1.2 (p. 15).

As we have described, the law of gravity was first formulated by Galileo using the concept of conservation of energy, which determines how masses are accelerated when friction is not considered and the mass is constant. Tycho Brahe investigated the motion of the planets. Starting in early 1600, Kepler began an extended visit with Brahe. In 1604, working with Brahe’s data Kepler described the inverse square law of light, and studied the workings of the human eye. Kepler was also the first to predict that the orbits of planets are described by ellipses. It seems he underappreciated the significance of his finding, as he continued working on his incorrect epicycle planetary model. Building on Galileo (1638) (see the discussion on pages 7-9), Newton demonstrated that there must be a gravitational potential between two masses \(m_1, m_2\) of the form

\[
\phi_{\text{New}}(r(t)) \propto \frac{m_1 m_2}{r(t)}, \quad (3.1.0.1)
\]
where \( r = |x_1 - x_2| \) is the Euclidean distance between the two point masses at locations \( x_1 \) and \( x_2 \). Using algebra and his calculus, Newton formalized the equations of gravity, forces, and motion (Newton’s three laws), the most important being

\[
f(t) = \frac{d}{dt} M(t) v(t),
\]

and showed that Kepler’s discovery of planetary elliptical motion naturally follows from these laws (see p. 9). With the discovery of Uranus in 1781, “Kepler’s theory was ruined” (i.e., proven wrong) (Stillwell, 2010, p. 23).

possibly the first measurement of the speed of sound, 1380 Paris feet per second was made by Marin Mersenne in 1630.\textsuperscript{a}

\textsuperscript{a}1 English foot is 1.06575 Paris feet.

**Newton and the speed of sound:** After Newton proposed the basic laws of gravity and explained the elliptical motion of the planets, he proposed the first model of the speed of sound.

In 1630 Mersenne showed that the speed of sound was approximately 1000 [ft/s]. This may have been done by finding the difference between the time of the flash of an explosion and the time it is heard. For example, if the explosion is 1 [mi] away, the delay is about 5 [s]. Thus with a simple clock, such as a pendulum, and an explosive, the speed may be accurately measured. If we say the speed of sound is \( c_0 \), then the equation for the wavefront is \( f(x,t) = u(x - c_0 t) \), where the function \( u(t) = 0 \) for \( t < 0 \) and \( u(t) = 1 \) for \( t > 0 \). If the wave is traveling in the opposite direction, then the formula is \( f(x,t) = u(x + c_0 t) \). If one also assumes that sounds add in an independent manner (superposition holds) (see Postulate P2 on p. 121), then the general solution for the acoustic wave is

\[
f(x,t) = Au(x - c_0 t) + Bu(x + c_0 t),
\]

where \( A \) and \( B \) are the amplitudes of the two waves. This is the solution proposed by d’Alembert in 1747 for the acoustic wave equation

\[
\frac{\partial^2}{\partial x^2} g(x,t) = \frac{1}{c_0^2} \frac{\partial^2}{\partial t^2} g(x,t),
\]

one of the most important equations of mathematical physics (see Eq. 4.4.0.1, p. 142), 20 years after Newton’s death.

It was well established, at least by the time of Galileo, that the wavelength \( \lambda \) and frequency \( f \) of a pure tone sound wave obey the relation

\[
f \lambda = c_0.
\]
3.1. ALGEBRA AND GEOMETRY AS PHYSICS

Given what we know today, the general solution to the wave equation may be written in terms of a sum over
the complex exponentials, famously credited to Euler, as
\[ q(x, t) = Ae^{2\pi i (ft-x/\lambda)} + Be^{2\pi i (ft+x/\lambda)}, \]
where \( t \) is time, \( x \) is position, and \( ft \) and \( x/\lambda \) are dimensionless. This equation describes only the steady-state
solution, with no onsets or dispersion. Thus this solution must be generalized to include these important effects.

The basics of sound propagation were within Newton’s grasp and were finally published in
Principia in 1687. The general solution to Newton’s wave equation [i.e., \( p(x, t) = G(x \pm x/c) \)], where \( G \) is any function, was first
published 60 years later by d’Alembert in 1747.

Newton’s value for the speed of sound in air \( c_o \) was incorrect by the thermodynamic constant \( \sqrt{\gamma_o} = \sqrt{\frac{T}{\rho}} \), a problem
that would take between 129 and 163 years to rectify, by Laplace in 1816, and experimentally by Rankine
in 1850 (Rayleigh, 1896, p. 19-23, Vol. II). What was needed was the adiabatic process (the concept of constant-
heat energy). For audio frequencies (0.02-20 [kHz]), the temperature gradients cannot diffuse the distance of a
wavelength in one cycle, “trapping” the heat energy in the wave (Tisza, 1966; Pierce, 1981; Boyer and Merzbach,
2011).\(^1\)

To repair Newton’s formula for the sound speed it was necessary to define the dynamic stiffness of air \( \eta_o P_o \),
where \( P_o \) is 1 [atm], or \( 10^5 \) [Pa] (1 [Pa] is 1 [N/m\(^2\)].

This required replacing Boyle’s Law (\( PV/T = \) constant) with the adiabatic expansion law (\( PV^{\gamma_o} = \) constant).
But this fix still ignores the important viscous and thermal losses, as discussed in Appendix D (Kirchhoff, 1868;
Rayleigh, 1896; Mason, 1927; Pierce, 1981).

Today we know that when ignoring viscous and thermal losses, the speed of sound is given by
\[ c_o = \sqrt{\frac{\eta_o P_o}{\rho_o}} = 343 \text{ [m/s]}, \]
which is a function of the density \( \rho_o = 1.12 \text{ [kg/m}^3] \), \( P_o = 10^5 \) [Pa], and the dynamic stiffness \( \eta_o P_o \) of air.\(^2\) The
speed of sound stated in other units is 434 [m/s], 1234.8 [km/h], 1.125 [ft/ms], 1125.3 [ft/s], 4.692 [s/mi], 12.78
[mi/min], 0.213 [mi/s], and 767 [mi/h].

Newton’s success was important because it quantified the physics behind the speed of sound and demonstrated
that momentum (\( mv \)), not mass \( m \), is transported by the wave. His concept was correct, and his formulation
using algebra and calculus represented a milestone in science, assuming no visco-elastic losses. When losses are
included, the wave number becomes a complex function of frequency, leading to Eq. D.1.1.4 (p. 250).

In periodic structures, again the wave number becomes complex due to diffraction, as commonly observed
in optics (e.g., diffraction gratings) and acoustics (creeping surface waves). Thus Eq. 3.1.0.4 holds for only the
simplest cases. In general, Eq. 3.1.0.6; the complex analytic (thus causal and dispersive) function propagation vector \( \kappa(x, s) \) must be considered (see Eq. 3.1.0.6).

The corresponding discovery of the formula for the speed of light was made 174 years after Principia by
Maxwell (ca.1861). Maxwell’s formulation also required great ingenuity, as it was necessary to hypothesize an
experimentally unmeasured term in his equations to get the mathematics to correctly predict the speed of light
(and gravity waves). This parallel with the speed of sound is notable.

It is somewhat amazing that to this day we have failed to fully understand gravity significantly better than
Newton’s theory, although this may too harsh given Einstein’s famous work on general relativity in about 1920.\(^3\)

Case of dispersive wave propagation: This classic relationship \( \lambda f = c \) is deceptively simple, yet confusing,
because the wave number\(^4\) \( k = 2\pi / \lambda \) becomes a complex function of frequency (has both real and imaginary
parts) in dispersive media when losses are considered, as discussed in Appendix D. (Kirchhoff, 1868; Mason,
1928).

A second important example is the case of electron waves in silicon crystals, where the wave number \( k(f) =
2\pi f / c \) is replaced by the complex analytic function of \( s \), the propagation vector \( \kappa(s) \). In this case the wave
becomes the eigenfunction of the vector (3D) wave equation
\[ p(\mathbf{x}, t) = P_o(s)e^{ist}e^{\pm \kappa(x, s) \cdot x}, \]
\(^1\)There were other physical enigmas, such as the observation that sound disappears in a vacuum, or Pascal’s observation that a vacuum
cannot draw water up a column by more than 34 feet.
\(^2\)\( \eta_o = c_p/c_v = 1.4 \) is the ratio of two thermodynamic constants, and \( P_o = 10^5 \) [Pa] is the barometric pressure of air.
\(^3\)Gravity waves were first observed experimentally while I was writing this chapter.
\(^4\)This term is a misnomer, since the wave number is a complex function of the complex Laplace frequency \( s = \sigma + \omega \), thus not a number
in the common sense. Much worse, \( \kappa(s) = s/c \) must be complex analytic in \( s \), which an even stranger condition. The term wave number is
so well established, that there is little hope for recovery at this point.

\[ \frac{\gamma_o}{c^2} \]
where \(|\kappa(x, s)|\) is the vector eigenvalue (Brillouin, 1953). In these more general cases, \(\kappa(x, s)\) must be a vector complex analytic function of the Laplace frequency \(s = \sigma + \omega j\), and inverted with the Laplace transform (Brillouin, 1960, with help from Sommerfeld). This is because electron “waves” in the dispersive semiconductor (e.g., silicon) are “causally filtered” in three dimensions – in magnitude, phase, and directions \(x\). These 3D dispersion relationships are known as Brillouin zones.

Silicon is a highly dispersive “wavefilter,” forcing the wavelength to be a function of both \(s\) and direction. This view is elegantly explained by Brillouin (1953, Chap. 1) in his historic text. Although the most famous examples come from quantum mechanics (Condon and Morse, 1929), modern acoustics contains a rich source of related examples (Morse, 1948; Beranek, 1954; Ramo et al., 1965; Fletcher and Rossing, 2008).

### 3.1.1 The first algebra

Prior to the invention of algebra, people worked out mathematical problems as sentences, using an obtuse description of the problem (Stillwell, 2010, p. 93). Algebra changed this approach and led to a compact language of mathematics, where numbers are represented as symbols (e.g., \(x\) and \(a\)). The problem to be solved could be formulated in terms of sums of powers of smaller terms, the most common being powers of some independent variable (i.e., time or frequency). If we set \(a_n = 1\), then

\[
P_N(z) \equiv z^n + a_{n-1}z^{n-1} + \cdots + a_0z^0 = z^n + \sum_{k=0}^{n-1} a_kz^k = \prod_{k=0}^{n} (z - z_k)
\]

(3.1.1.7)

is called a monic polynomial. The coefficient \(a_n\) cannot be zero, or the polynomial would not be of degree \(n\). The resolution is to force \(a_n = 1\), since this simplifies the expression and does not change the roots.

The key question is What values of \(z = z_k\) result in \(P_N(z_k) = 0\)? In other words, what are the roots \(z_k\) of the polynomial? Answering this question consumed thousands of years, with intense efforts by many aspiring mathematicians. In the earliest attempts, it was a competition to demonstrate mathematical acumen. Results were held as a secret to the death bed. It would be fair to view this effort as an obsession. Today the roots of any polynomial may be found, to high accuracy, by numerical methods. Finding roots is limited by the numerical limits of the representation—namely, by IEEE 754 (see p. 29). There are also a number of important theorems.

Of particular interest is the problem of drawing a circle and a line, and finding the intersection (root). There was no solution to this venerable problem using geometry. The resolution is addressed in the solution of Euclid’s formula (Problem # 2 of Assignment 2.2).

### 3.1.2 Finding roots of polynomials

The problem of factoring polynomials has a history more than a millennium in the making. While the quadratic (degree \(N = 2\)) was solved by the time of the Babylonians (i.e., the earliest recorded history of mathematics), the cubic solution was finally published by Cardano in 1545. The same year, Cardano’s student solved the quartic (\(N = 4\)). In 1826 (281 years later) it was proved that the quintic (\(N = 5\)) could not be factored by analytic methods.

As a concrete example we begin with the important but trivial case of the quadratic

\[
P_2(s) = as^2 + bs + c.
\]

(3.1.2.8)

First note that if \(a = 0\), the quadratic reduces to the monomial \(P_1(s) = bs + c\). Thus we have the necessary condition that \(a \neq 0\). The best way to proceed is to divide \(a\) out and work directly with the monic \(\hat{P}_2(s) = \frac{1}{a}P_2(s)\). In this way we do not need to worry about the \(a = 0\) exception.

The roots are the values of \(s = s_k\) such that \(\hat{P}_2(s_k) = 0\). One of the earliest mathematical results, recorded by the Babylonians in about 2000 BCE, was the factoring of the quadratic by completing the square. We can isolate \(s\) by rewriting Eq. 3.1.2.8 as

\[
\hat{P}_2(s) \equiv \frac{1}{a}P_2(s) = (s + b/2a)^2 - (b/2a)^2 + c/a.
\]

(3.1.2.9)

The factorization may be verified by expanding the squared term and canceling \((b/2a)^2\):

\[
\hat{P}_2(s) = (s^2 + (b/a)s + (b^2/2a)^2) - (b^2/2a)^2 + c/a.
\]

Setting Eq. 3.1.2.9 equal to zero and solving for the two roots \(s_{\pm}\) give the quadratic formula:

\[
s_{\pm} = \frac{-b \pm \sqrt{b^2 - 4ac}}{2a}
\]

(3.1.2.10)
3.1. ALGEBRA AND GEOMETRY AS PHYSICS

Role of the discriminant: Equation 3.1.2.10 can be further simplified. The term \((b/2)^2 - c > 0\) under the square root is called the discriminant. Nominally in physics and engineering problems, the discriminant is negative and \(b/2 \ll \sqrt{c}\) may be ignored (the damping is small compared to the resonant frequency), leaving only \(-c\) under the radical. Thus the most natural way (i.e., corresponding to the most common physical cases) of writing the roots (Eq. 3.1.2.10) is

\[
s_\pm \approx -b/2 \pm j\sqrt{|c|} = -\sigma_0 \pm s_0. \tag{3.1.2.11}
\]

This form separates the real and imaginary parts of the solution in a natural way. The term \(\sigma_0 = b/2\) is called the damping, which accounts for losses in a resonant circuit; the term \(\omega_0 = \sqrt{|c|}\), for mechanical, acoustical, and electrical networks, is called the resonant frequency. The last approximation ignores the (typically) minor correction \(b/2\) to the resonant frequency, which in engineering applications is almost always ignored. Knowing that there is a correction is highlighted by this formula, which makes us aware that the small approximation exists (thus can be ignored).

It is not required that \(a, b, c \in \mathbb{R} > 0\), but for physical problems of interest, this is almost always true (>99.99% of the time).

Summary: The quadratic equation and its solution are ubiquitous in physics and engineering. It seems obvious that instead of memorizing the meaningless quadratic formula (Eq. 3.1.2.10), one should learn the physically meaningful solution (Eq. 3.1.2.11), obtained via Eq. 3.1.2.9 with \(a = 1\). Arguably, the factored and normalized form (Eq. 3.1.2.9) is easier to remember as a method (completing the square) rather than as a formula to be memorized.

Additionally, the real \((b/2)\) and imaginary (\(\pm j\sqrt{c}\)) parts of the two roots have physical significance as the damping and resonant frequency. Equation 3.1.2.10 has none (it is useless).

No insight is gained by memorizing the quadratic formula. To the contrary, an important concept is gained by learning how to complete the square, which is typically easier than identifying \(a, b, c\) and blindly substituting them into Eq. 3.1.2.10. Thus it’s worth learning the alternative solution (Eq. 3.1.2.11), since it is more common in practice and requires less algebra to interpret the final answer.

Exercise #1

By direct substitution, demonstrate that Eq. 3.1.2.10 is the solution of Eq. 3.1.2.8. Hint: Work with \(\tilde{P}_2(x)\).

**Solution:** Setting \(a = 1\), we can write the quadratic formula as

\[
s_\pm = \frac{-b \pm j\sqrt{4c - b^2}}{2}.
\]

Substituting this into \(\tilde{P}_2(s)\) gives

\[
\tilde{P}_\pm(s) = s_\pm^2 + bs_\pm + c = \frac{-b \pm \sqrt{b^2 - 4c}}{2} \frac{-b \pm \sqrt{b^2 - 4c}}{2} + c
\]

\[
= \left(\frac{-b + \sqrt{b^2 - 4c}}{2}\right)^2 + b \left(\frac{-b + \sqrt{b^2 - 4c}}{2}\right) + c
\]

\[
= \frac{1}{4} \left(b^2 - 2b\sqrt{b^2 - 4c} + (b^2 - 4c)\right) + \frac{1}{4} \left(b^2 + 2b\sqrt{b^2 - 4c} + (b^2 - 4c)\right)
\]

\[
= 0.
\]

In third grade I learned the times-table trick for 9:

\[
9 \cdot n = (n - 1) \cdot 10 + (10 - n).
\]

With this simple rule I did not need to depend on my memory for the 9 times table. For example: \(9 \cdot 7 = (7 - 1) \cdot 10 + (10 - 7) = 60 + 3 = 63\) and \(9 \cdot 3 = (3 - 1) \cdot 10 + (10 - 3) = 20 + 7 = 27\). By expanding, one can see why it works: \(9n = n10 + (-10) + (10 - n) = n(10 - 1). \) Note that the two terms \((n - 1)\) and \((10 - n)\) add to 9.

Learning an algorithm is much more powerful than memorizing the 9 times tables. How you think about a problem can have a great impact on your perception.

---

5This is the case for mechanical and electrical circuits that have small damping. Physically \(b > 0\) is the damping coefficient and \(\sqrt{c} > 0\) is the resonant frequency. One may then simplify and factor the form as \(s^2 + 2bs + c^2 = (s + b + j\sqrt{c})(s + b - j\sqrt{c}).\)
Newton’s method for finding the roots of $P_N(s)$: Newton is well known for an approximate but efficient method to find the roots of a polynomial. Consider the polynomial $s$, $P_N(s) \in \mathbb{C}$:

$$P_N(s) = c_N(s - s_0)^N + c_{N-1}(s - s_0)^{N-1} + \cdots + c_1(s - s_0) + c_0,$$  \hspace{1cm} (3.1.2.12)

where we use Taylor’s formula (see p. 76) to determine the coefficients

$$c_k = \frac{1}{k!} \frac{d^k}{ds^k} P_N(s) \bigg|_{s=s_0}. \hspace{1cm} (3.1.2.13)$$

If our initial guess for the root $s_1$ is close to a root $s_0$ (i.e., $s_1 - s_0$ is within the radius of convergence), then $|(s_1 - s_0)^k| \ll |(s_1 - s_0)|$ for $k \geq 2 \in \mathbb{N}$. Thus we may truncate $P_N(s_1)$ to its linear term $c_1$:

$$P_N(s_1) \approx (s_1 - s_0) \frac{d}{ds} P_N(s) \bigg|_{s=s_0} + P_N(s_0)$$

$$= (s_1 - s_0)P'_N(s_0) + P_N(s_0),$$

where $P'_N(s_0)$ is shorthand for $dP_N(s_0)/ds$.

Newton’s approach (approximation) was to define a recursion such that the next guess $s_{n+1}$ is closer to the root $s_o$ than the previous guess $s_n$. Replacing $s_1$ by $s_{n+1}$ and $s_0$ by $s_n$ gives

$$P_N(s_{n+1}) = (s_{n+1} - s_n)P'_N(s_n) + P_N(s_n) \to 0.$$  

Here we assume $P_N(s_{n+1}) \to 0$ because $s_{n+1} \to s_0$ as $n \to \infty$.

Solving for $s_{n+1}$, we get

$$s_{n+1} = s_n - \frac{P_N(s_n)}{P'_N(s_n)}.$$  \hspace{1cm} (3.1.2.14)

Everything on the right is known, thus $s_{n+1}$ should converge to the root $s_0$ as $n \to \infty$.

In practice, it takes only a few steps to approach the root. In experimental trials (see Fig. 3.2) fewer than 10 steps give double-precision floating-point machine accuracy. If any value $s_n$ is close to a root of $P'_N$, the recursion fails, giving a large value for $s_{n+1}$ and forcing the method to restart at $s_{n+1}$, far from the root. In such cases the solution typically converges to a different root. It should not be difficult to detect these large nonconvergent steps by monitoring $|s_{n+1} - s_n|$, which should be monotonically decreasing.

However, if one assumes that the initial guess $s_1 \in \mathbb{R}$ and then evaluates the polynomial using real arithmetic, the estimate $s_{n+1} \in \mathbb{R}$. Thus the iteration will not converge if $s_0 \in \mathbb{C}$.

Root $s_0 \in \mathbb{C}$ may be found by a recursion that defines a sequence $s_n \to s_0$, $n \in \mathbb{N}$, such that $P_N(s_n) \to 0$ as $n \to \infty$. As shown in Fig. 3.2, solving for $s_{n+1}$ using Eq. 3.1.2.14 always gives one of the roots, due to the analytic behavior of the complex logarithmic derivative $P'_N/P_N$.

With every step, $s_{n+1}$ is closer to the root, finally converging to the root in the limit. As it comes closer, the linearity assumption becomes more accurate, resulting in a better approximation and thus a faster convergence.

Equation 3.1.2.14 depends on the log-derivative $d\log P(x)/dx = P'(x)/P(x)$. It follows that even for cases where fractional derivatives of roots are involved (see p. 150), Newton’s method should converge, since the logarithmic derivative linearizes the equation.\(^5\)

Newton’s view: Newton believed that imaginary roots and numbers have no meaning (see p. 134), thus he sought only real roots (Stillwell, 2010, p. 119). In this case Newton’s relationship may be explored as a graph, which puts Newton’s method in the realm of analytic geometry.

Example: Given a polynomial $P_2 = 1 - x^2$ that has roots $\pm 1$, we can use Newton’s method to find the roots. For $P'_2(x) = -2x$, Newton’s iteration becomes

$$x_{n+1} = x_n + \frac{1 - x^2_n}{2x_n}.$$  

\(^5\)This seems like a way to understand fractional, even irrational, roots.
3.1. ALGEBRA AND GEOMETRY AS PHYSICS

Figure 3.2: Newton’s method applied to the polynomial that has real roots \(1, 2, 3, 4\) (left) and 5 complex roots (right). A random starting point was chosen, and each curve shows the values of \(s_n\) as Newton’s method converges to the root. Different random starting points converge to different roots. The method always results in convergence to a root. Claims to the contrary (Stewart, 2012, p. 347) are a result of forcing the roots to be real. For convergence, one must assume \(s_n \in \mathbb{C}\). For a related discussion see Stillwell (2010, Sec. 14.7).

From the Gauss-Lucas theorem, for the case of \(N = 2\), the root of \(P'_2(x)\) is always the average of the roots of \(P_2(x)\). To start the iteration \((n = 0)\) we need an initial guess for \(x_0\), which is an initial random guess of where a root might be. The only place we may not start is at the roots of \(P'_N\). For \(P_2(x) = 1 - x^2\),

\[
x_1 = x_0 + \frac{1 - x_0^2}{2x_0} = x_0 + \frac{1}{2} (-x_0 + 1/x_0).
\]

Exercise #2
Let \(P_2(x) = 1 - x^2\). Choose the expansion point as \(x_0 = 1/2\). Draw a graph describing the first step of the iteration.

Solution: We start with an \((x, y)\) coordinate system and put points at \(x_0 = (1/2, 0)\) and the vertex of \(P_2(x) = (0, 1)\) \((P_2(0) = 1)\). Then we draw \(1 - x^2\), along with a line from \(x_0\) to \(x_1\).

Exercise #3
From Exercise #3 calculate \(x_1\) and \(x_2\). What number is the algorithm approaching? Is it a root of \(P_2\)?

Solution: First we must find \(P'_2(x) = -2x\). Thus the equation we will iterate is

\[
x_{n+1} = x_n + \frac{1 - x_n^2}{2x_n} = \frac{x_n^2 + 1}{2x_n}.
\]

By hand,

\[
x_0 = 1/2
x_1 = \frac{(1/2)^2 + 1}{2(1/2)} = \frac{1}{4} + 1 = 5/4 = 1.25
x_2 = \frac{(5/4)^2 + 1}{2(5/4)} = \frac{25/16 + 1}{10/4} = \frac{41}{40} = 1.025.
\]

These estimates rapidly approach the positive real root \(x = 1\). Note that if one starts at the root of \(P'(x) = 0\) (i.e., \(x_0 = 0\)), the first step is indeterminate.

Exercise #4
Write an Octave/Matlab script to check your answer for part (a).

Solution:
```octave
x=1/2;
for n = 1:3
    x = x+(1-x*x)/(2*x);
end
```

Exercise #5
For \( n = 4 \), what is the absolute difference between the root and the estimate, \(|x_r - x_4|\)?

Solution: \(4.6 \times 10^{-8}\) (very small!)

Exercise #6
What happens if \(x_0 = -1/2\)?

Solution: The solution converges to the negative root, \(x = -1\).

Exercise #7
Does Newton’s method (Kelley, 2003) work for \(P_2(x) = 1 + x^2\)? Hint: What are the roots in this case?

Solution: In this case \(P_2'(x) = 2x\), thus the iteration gives

\[x_{n+1} = x_n - \frac{1 + x_n^2}{2x_n}.
\]

The roots are purely imaginary, \(x_{\pm} = \pm 1i\). Obviously Newton’s method fails because there is no way for the answer to become complex. If, like Newton, you didn’t believe in complex numbers, your method would fail to converge to the complex roots (i.e., Real in \(\rightarrow\) Real out). This is because Octave/Matlab assumes \(x \in \mathbb{R}\) if it is initialized as \(\mathbb{R}\). □

Exercise #8
What if you let \(x_0 = (1 + j)/2\) for the case of \(P_2(x) = 1 + x^2\)?

Solution: By starting with a complex initial value, we fix the Real in \(\Rightarrow\) Real out problem. □

Basic properties of polynomials
In some sense polynomials such as \(P_N(z)\) are the simplest constructions used in algebra, and a summary of their most basic properties is helpful.

1. The degree of a polynomial is \(n\).
2. Polynomials are single-valued; that is, for every \(z_o\), there is precisely one value for \(P_N(z_o)\).
3. In mathematical physics and engineering it is common to have real coefficients \(a_n\), but complex coefficients are possible.
4. The coefficients of every polynomial are determined by its Taylor series—namely, Eq. 3.2.2 (see p. 76).
5. If the coefficients are real and positive, then the \(P_N(x)\) is positive and real if \(x \geq 0\)
6. The fundamental theorem of algebra states that \(P_N(z)\) has exactly \(n\) roots.
7. The roots of polynomials with positive and real coefficients typically have complex roots – that is, if \(P_N(z_k) = 0\), then \(z_k \in \mathbb{C}\).
8. The region of convergence (RoC) of every polynomial about the expansion point is infinite.
9. The roots of the derivative of a polynomial lie within the convex hull defined by the roots of \(P_N(z)\), as described by the Gauss-Lucas theorem (see p. 69).

Exercise #9
Find the logarithmic derivative of \(f(x)g(x)\).

Solution: From the definition of the logarithmic derivative and the chain rule for the differentiation of a product, we have

\[
\frac{d}{dx} \ln f(x)g(x) = \frac{d}{dx} \ln f + \frac{d}{dx} \ln g = \frac{1}{f} \cdot \frac{d}{dx} f + \frac{1}{g} \cdot \frac{d}{dx} g.
\]
Example: If we assume that function \( P_3(s) = (s - a)^2/(s - b)^2 \), then
\[
\ln P_3(s) = 2 \ln(s - a) - \pi \ln(s - b)
\]
and
\[
\frac{d}{ds} \ln P_3(s) = \frac{2}{s - a} - \frac{\pi}{s - b}.
\]

Reduction by the logarithmic derivative to simple poles: As shown in the Example \( P_3(s) \) above, a function that has poles of arbitrary degree (i.e., \( \pi \) in the example) may be reduced to the sum of two functions having simple poles by taking the logarithmic derivative since
\[
L_N(s) = \frac{N(s)}{D(s)} = \frac{d}{ds} \ln P_N(s) = \frac{P'_N(s)}{P_N(s)}.
\]  
(3.1.2.15)

Here the polynomial is the denominator \( D(s) = P_N(s) \), while the numerator \( N(s) = P'_N(s) \) is the derivative of \( D(s) \). Thus the logarithmic derivative can play a key role in the analysis of complex analytic functions, as it reduces higher-order poles, even those of irrational degree, to simple poles (those of degree 1).

The logarithmic derivative \( L_N(s) \) has the following special properties:

1. \( L_N(s) \) has simple poles \( s_p \) and zeros \( s_z \).
2. The poles of \( L_N(s) \) are the zeros of \( P_N(s) \).
3. The zeros of \( L_N(s) \) (i.e., \( P'_N(s_z) = 0 \)) are the zeros of \( P'_N(s) \).
4. \( L_N(s) \) is analytic everywhere other than at its poles.
5. Since the zeros of \( P_N(s) \) are simple (no second-order poles), the zeros of \( L_N(s) \) always lie close to the line connecting the two poles. One may easily demonstrate the truth of the statement numerically, and it has been quantified by the Gauss-Lucas theorem, which specifies the relationship between the roots of a polynomial and those of its derivative. Specifically, the roots of \( P'_{N-1} \) lie inside the convex hull of the roots of \( P_N \).

To understand the meaning of convex hull, consider the following construction: If stakes are placed at each of the \( N \) roots of \( P_N(x) \), and a string is then wrapped around the stakes, with all the stakes inside the string, the convex hull is then the closed set inside the string. One can begin to imagine how the \( N - 1 \) roots of the derivative must evolve with each set inside the convex hull of the previous set. This concept may be recursed to smaller values of \( N \).

6. Newton’s method may be expressed in terms of the reciprocal of the logarithmic derivative, since
\[
s_{k+1} = s_k + \epsilon_o/L_N(s),
\]
where \( \epsilon_o \) is called the step size, which is used to control the rate of convergence of the algorithm. If the step size is too large, the root-finding path may jump to a different domain of convergence and thus a different root of \( P_N(s) \).

7. Not surprisingly, given all the special properties, \( L_N(x) \) plays an key role in mathematical physics.

Euler’s product formula: Counting may be written as a linear recursion simply by adding 1 to the previous value, starting from 0. The even numbers may be generated by adding 2, starting from 0. Multiples of 3 may be similarly generated by adding 3 to the previous value, starting from 0. Such recursions are fundamentally related to prime numbers \( \pi_k \in P \), as first investigated by Euler. This logic is the basis of the sieve (see Sec. Sec. 2.5, p. 42). The basic idea is both simple and important, taking almost everyone by surprise, likely even Euler. It is related to the old idea that the integers may be generated by the geometric series when viewed as a recursion.

Example: Let’s look at counting modulo prime numbers. For example, if \( k \in \mathbb{N} \), then
\[
k \cdot \text{mod}(k, 2), \quad k \cdot \text{mod}(k, 3), \quad k \cdot \text{mod}(k, 5)
\]
are all multiples of the primes \( \pi_1 = 2, \pi_2 = 3, \) and \( \pi_3 = 5 \).
Exercise #10
Show that \( N_n = n \) follows from the above recursion. **Solution:** If \( n = -1 \), we have \( N_n = 0 \) and \( u_n = 0 \). For \( n = 0 \) the recursion gives \( N_1 = N_0 + u_0 \), thus \( N_1 = 0 + 1 \). When \( n = 1 \), we have \( N_2 = N_1 + 1 = 1 + 1 = 2 \). For \( n = 2 \), the recursion gives \( N_3 = N_2 + 1 = 3 \). Continuing the recursion, we find that \( N_n = n \). Today we denote recursions of this form as a digital filter. The state diagram for \( N_n \) is shown in Fig. 3.3. ■

To start the recursion, we define \( u_n = 0 \) for \( n < 0 \). Thus \( u_0 = u_{-1} + 1 \). But since \( u_{-1} = 0 \), \( u_0 = 1 \). The counting numbers follow from this recursion. A more understandable notation is convolution of the step function with itself—namely,

\[
m_n = u_n \ast u_n = \sum_{m=0}^{\infty} u_m u_{m-n} \leftrightarrow \frac{1}{(1-z)^2},
\]

which says that the counting numbers \( \hat{n} \in \mathbb{N} \) are easily generated by convolution, which corresponds to a second-order pole at \( z = 1 \) in the \( z \)-transform frequency domain (see Sec. 3.4.1).

Exercise #11
Write an Octave/Matlab program that generates the odd numbers \( N_n = \{1, 0, 3, 0, 5, 0, 7, 0, 9, \ldots\} \) by removing the even numbers.

**Solution:**
```matlab
M=50; N=(0:M-1); u=ones(1,M); u(1)=0; Dem=[1 1]; Num=[1]; n=filter(Num,Dem,u); y2=n.*N; F1=N-y2
```
which generates: \( F1 = [0, 1, 0, 3, 0, 5, 0, 7, 0, 9, 0, \ldots] \). ■

An alternative is to use the \( \text{mod}(n,N) \) function:
```matlab
M=20; n=0:M; k=mod(n,2); m=(k==0).*n;
```
which generates \( m = [0, 1, 0, 3, 0, 5, \ldots] \).
Exercise #12
Write a program to recursively down-sample \( N \) by 2:1.

**Solution:**
\[
N = [1 0 3 0 5 0 7 0 9 0 11 0 13 0 15]
\]
\[M = N(2:2:end);\]
which gives: \( M = [1, 3, 5, 7, 9, 11, 13, 15, \ldots] \]

For the next step toward a full sieve (Fig. 2.3, p. 43), we generate all the multiples of 3 (the second prime) and subtract these from the list. This will either zero out these numbers from the list or create negative items, which may then be removed. Numbers are negative when the number has already been removed because it has a second factor of that number. For example, 6 is already removed because it is a multiple of 2 and thus was removed with the multiples of prime number 2.

### 3.1.3 Matrix formulation of the polynomial

There is a simple relationship between every constant coefficient differential equation, its characteristic polynomial, and the equivalent matrix form of that differential equation, defined by the *companion matrix*. The roots of the monic polynomial are the eigenvalues of the companion matrix \( C_N \) (Horn and Johnson, 1988, p. 147).

The **companion matrix**: The \( N \times N \) companion matrix is defined as

\[
C_N = \begin{bmatrix}
0 & -c_0 \\
1 & 0 & 0 \\
0 & 1 & 0 \\
\vdots & 0 & 1 & 0 & \ddots & \vdots \\
0 & \cdots & 0 & 1 \\
& & & & & & -c_N-2 \\
& & & & & -c_N-1
\end{bmatrix}_{N \times N}
\]

(3.1.3.17)

The constants \( c_{N-n} \) are from the monic polynomial of degree \( N \),

\[
P_N(s) = s^N + c_{N-1}s^{N-1} + \cdots + c_2s^2 + c_1s + c_0
\]

\[
= s^N + \sum_{n=0}^{N-1} c_ns^n,
\]

that has coefficient vector

\[
c_N = [1, c_{N-1}, c_{N-2}, \ldots, c_0]^T.
\]

Any transformation of a matrix that leaves the eigenvalues invariant (e.g., the transpose) results in an equivalent definition of \( C_N \). Note that the Octave and Matlab companion matrix function \( \text{C} = \text{compan}(\text{A}) \) returns the coefficient vector along the top row.

**Exercise #13**
Show that the eigenvalues of the \( 3 \times 3 \) companion matrix are the same as the roots of \( P_3(s) \).

**Solution:** Expanding the determinant of \( C_3 - sI_3 \) along the rightmost column, we get

\[
P_3(s) = -s & 0 & -c_0 \\
1 & -s & -c_1 \\
0 & 1 & -(c_2 + s)
\]

\[
= c_0 + c_1s + (c_2 + s)s^2 = s^3 + c_2s^2 + c_1s + c_0.
\]

Setting this to zero gives the requested result.

**Exercise #14**
Find the companion matrix for the Fibonacci sequence defined by the recursion (i.e., difference equation)

\[
f_{n+1} = f_n + f_{n-1}
\]

and initialized with \( f_n = 0 \) for \( n < 0 \) and \( f_0 = 1 \).
**Solution:** Taking the z-transform gives the polynomial \((z^1 - z^0 - z^{-1})F(z) = 0\), which has the coefficient vector \(c = [1, -1, -1]^T\), resulting in the Fibonacci companion matrix

\[
C = \begin{bmatrix}
0 & 1 \\
1 & 1
\end{bmatrix}.
\]

The Matlab/Octave companion matrix routine `compan(C)` uses an alternative definition that has the same eigenvalues (see p. 53).

**Example:** A polynomial is represented in Matlab/Octave in terms of its coefficient vector. When the polynomial vector for the poles of a differential equation is

\[c_N = [1, c_{N-1}, c_{N-2}, \ldots, c_0]^T,\]

the coefficient \(c_N = 1\). This normalization guarantees that the leading term is not zero and the number of roots \((N)\) is equal to the degree of the monic polynomial.

### 3.1.4 Working with polynomials in Matlab/Octave

In Matlab/Octave there are eight related functions you must become familiar with:

1. \(R = \text{roots}(A)\): Vector \(A = [a_N, a_{N-1}, \ldots, a_0] \in \mathbb{C}\) are the complex coefficients of polynomial \(P_N(z) = \sum_{n=0}^{N} a_n z^n \in \mathbb{C}\), where \(N \in \mathbb{N}\) is the degree of the polynomial. It is convenient to force \(a_N = 1\), corresponding to dividing the polynomial by this value, when it is not 1, thus guaranteeing it cannot be zero. Further, \(R\) is the vector of roots \([z_1, z_2, \ldots, z_n] \in \mathbb{C}\) such that \(\text{polyval}(A, z_k) = 0\).

   **Example:** \(\text{roots}([1, -1]) = 1\)

2. \(y = \text{polyval}(A, x)\): This evaluates the polynomial defined by vector \(A \in \mathbb{C}^N\) evaluated at \(x \in \mathbb{C}\), returning vector \(y(x) \in \mathbb{C}\).

   **Example:** \(\text{polyval}([1 -1], 1) = 0, \text{polyval}([1, 1], 3) = 4\)

3. \(P = \text{poly}(R)\): This is the inverse of \(\text{root}\), returning a vector of polynomial coefficients \(P \in \mathbb{C}^N\) of the corresponding characteristic polynomial, starting from either a vector of roots \(R\) or a matrix \(A\), for example, defined with the roots on the diagonal. The characteristic polynomial is defined as the determinant of \(|A - \lambda I| = 0\) that has roots \(R\).

   **Example:** \(\text{poly([1])} = [1, -1], \text{poly([1,2])} = [1, -3, 2]\)

   Due to IEEE 754 scaling issues, this can give strange results that are numerically correct, but only within the limits of IEEE 754 accuracy.

4. \(R = \text{polyder}(C)\): This routine takes the \(N\) coefficients of polynomial \(C\) and returns the \(N - 1\) coefficients of the derivative of the polynomial. This is useful when working with Newton’s method, where each step is proportional to \(P_N(x)/P'_{N-1}(x)\).

   **Example:** \(\text{polyder([1,1])} = [1]\)

5. \([K, R] = \text{residue}(N, D)\): Given the ratio of two polynomials \(N, D, \text{residue}(N, D)\) returns vectors \(K, R\) such that

\[
\frac{N(s)}{D(s)} = \sum_k \frac{K_k}{s - s_k}, \quad (3.1.48)
\]

   where \(s_k \in \mathbb{C}\) are the roots of the denominator \(D\) polynomial and \(K \in \mathbb{C}\) is a vector of residues, which characterize the roots of the numerator polynomial \(N(s)\). The use of \(\text{residue}(N, D)\) is discussed on page 152. This is one of the most valuable time-saving routines I know.

   **Example:** \(\text{residue}(2, [1 0 -1]) = [1 -1]\)
6. $C = \text{conv}(A, B)$: Vector $C \in \mathbb{C}^{N + M - 1}$ contains the polynomial coefficients of the convolution of the two vectors of coefficients of polynomials $A, B \in \mathbb{C}^N$ and $B \in \mathbb{C}^M$.

Example: $[1, 2, 1] = \text{conv}([1, 1], [1, 1])$

7. $[C, R] = \text{deconv}(N, D)$: Vectors $C, N, D \in \mathbb{C}$. This operation uses long division of polynomials to find $C(s) = N(s)/D(s)$ with remainder $R(s)$, where $N = \text{conv}(D, C) + R$—namely,

$$C = \frac{N}{D} \text{ remainder } R.$$  \hfill (3.1.4.19)

Example: By defining the coefficients of two polynomials as $A = [1, a_1, a_2, a_3]$ and $B = [1, b_1, b + 2]$, we can find the coefficients of the product from $C = \text{conv}(A, B)$ and recover $B$ from $C$ with $B = \text{deconv}(C, A)$.

8. $A = \text{compan}(D)$: Vector $D = [1, d_{N-1}, d_{N-2}, \ldots, d_0]^T \in \mathbb{C}$ contains the coefficients of the monic polynomial

$$D(s) = s^N + \sum_{k=1}^{N} d_{N-k} s^k,$$

and $A$ is the companion matrix of vector $D$ (Eq. 3.1.3.17, p. 71). The eigenvalues of $A$ are the roots of the monic polynomial $D(s)$.

Example: $\text{compan}([1 -1 -1]) = [1 1; 1 0]$

Exercise #15
Practice the use of Matlab’s/Octave’s related functions that manipulate roots, polynomials, and residues: $\text{root}()$, $\text{conv}()$, $\text{deconv}()$, $\text{poly}()$, $\text{polyval}()$, $\text{polyder}()$, $\text{residue}()$, $\text{compan}()$.

Solution: We try Newton’s method for various polynomials. We use $N = \text{poly}(R)$ to provide the coefficients of a polynomial given the roots $R$. Then we use $\text{root}()$ to factor the resulting polynomial. Finally, we use Newton’s method and show that the iteration converges to the nearest root. \hfill $\blacksquare$

3.2 Eigenanalysis

At this point we turn a corner in the discussion toward the important topic of eigenanalysis, which starts with the computation of the eigenvalues and their eigenvectors of a matrix. As briefly discussed on page 34, eigenvectors are mathematical generalizations of resonances, or modes, naturally found in physical systems.

When you pluck the string of a violin or guitar, or hammer a bell or tuning fork, there are natural resonances that occur. These are the eigenmodes of the instrument. The frequency of each mode is related to the eigenvalue, which in physical terms is the frequency of the mode. But this idea goes way beyond simple acoustical instruments. Wave-guides and atoms are resonant systems. The resonances of the hydrogen atom are called the Lyman series, a special case of the Rydberg series and Rydberg atom (Bohr, 1954; Gallagher, 2005).

Thus this topic runs deep in both physics and eventually mathematics. In some real sense, eigenanalysis was what the Pythagoreans were seeking to understand. This relationship is rarely spoken about in the literature, but once you see it, it can never be forgotten, as it colors your entire view of all aspects of modern physics.

3.2.1 Eigenvalues of a matrix

The method for finding eigenvalues is best described with an example. Starting from the matrix Eq. 2.5.7.15 (p. 54), the eigenvalues are defined by the eigenmatrix equation

$$\frac{1}{2} \begin{bmatrix} 1 & 1 \\ 2 & 0 \end{bmatrix} \begin{bmatrix} \nu_1 \\ \nu_2 \end{bmatrix} = \lambda \begin{bmatrix} \nu_1 \\ \nu_2 \end{bmatrix}.$$
CHAPTER 3. ALGEBRAIC EQUATIONS: STREAM 2

The unknowns here are the eigenvalue \( \lambda \) and the eigenvector \( \mathbf{e} = [e_1, e_2]^T \). First we find \( \lambda \) by subtracting the right from the left:

\[
\frac{1}{2} \begin{bmatrix} 1 & 1 \\ 2 & 0 \end{bmatrix} \begin{bmatrix} e_1 \\ e_2 \end{bmatrix} - \lambda \begin{bmatrix} e_1 \\ e_2 \end{bmatrix} = \frac{1}{2} \begin{bmatrix} 1 - 2\lambda & 1 \\ 2 & -2\lambda \end{bmatrix} \begin{bmatrix} e_1 \\ e_2 \end{bmatrix} = 0. \tag{3.2.1.1}
\]

The only way this equation for \( \mathbf{e} \) can have a solution is if the matrix is singular. If it is singular, the determinant of the matrix is zero.

**Example:** The determinant in the above equation is the product of the diagonal elements minus the product of the off-diagonal elements, which results in the quadratic equation

\[-2\lambda(1 - 2\lambda) - 2 = 4\lambda^2 - 2\lambda - 2 = 0.
\]

Completing the square gives

\[(\lambda - 1/4)^2 - (1/4)^2 - 1/2 = 0, \tag{3.2.1.2}
\]

thus the roots (i.e., eigenvalues) are \( \lambda_{\pm} = \frac{1\pm\sqrt{5}}{4} = \{1, -1/2\} \).

**Exercise #16**
Expand Eq. 3.2.1.2 and recover the quadratic equation.

**Solution:**

\[(\lambda - 1/4)^2 - (1/4)^2 - 1/2 = \lambda^2 - \lambda/2 + (1/4)^2 - (1/4)^2 - 1/2 = 0.
\]

Thus completing the square is the same as the original equation. ■

**Exercise #17**
Find the eigenvalues of the matrix in Eq. 2.5.6.7 (see p. 52).

**Solution:** This is a minor variation on the previous example. Briefly, we have

\[\det \begin{bmatrix} 1 - \lambda & N \\ 1 & 1 - \lambda \end{bmatrix} = (1 - \lambda)^2 - N = 0.
\]

Thus \( \lambda_{\pm} = 1 \pm \sqrt{N} \). ■

**Exercise #18**
Starting from \([x_n, y_n]^T = [1, 0]^T\) compute the first 5 values of \([x_n, y_n]^T\).

**Solution:** Here is a Matlab/Octave code for computing \(x_n\):

\[
x(1:2,1)=[1;0];
A=[1 1;2 0]/2;
for k=1:10; x(k+1)=A*x(:,k); end
\]

which gives the rational \((x_n \in \mathbb{Q})\) sequence: 1, 1/2, 3/4, 5/8, 11/2^4, 21/2^5, 43/2^6, 85/2^7, 171/2^8, 341/2^9, 683/2^10, ... ■

**Exercise #19**
Show that the solution to Eq. 2.5.7.14 (see p. 54) is bounded, unlike that of the divergent Fibonacci sequence. Explain what is going on.

**Solution:** Because the next value is the mean of the last two, the sequence is bounded. To see this one needs to compute the eigenvalues of the matrix in Eq. 2.5.7.15 (p. 54). ■

The key to the analysis of such equations is called the eigenanalysis, or modal-analysis method. These are also known as resonant modes in the physics literature. Eigenmodes describe the naturally occurring “ringing” found in physical wave-dominated boundary value problems. Each mode’s eigenvalue quantifies the mode’s natural frequency. Complex eigenvalues result in damped modes, which decay in time due to energy losses. Common examples include tuning forks, pendulums, bells, and the strings of musical instruments, all of which have a characteristic frequency.
Two modes with the same frequency are said to be degenerate. This is a very special condition, with a high degree of symmetry.

Cauchy’s residue theorem (see p. 151) is used to find the time-domain response of each frequency-domain complex eigenmode. Thus eigenanalysis and eigenmodes of physics are the same thing (see p. 142) but are described using different notional methods.\(^9\) The “eigen method” is summarized in Appendix B.3, (see p. 237).

Taking a simple example of a 2 \(\times\) 2 matrix \(T \in \mathbb{C}\), we start from the definition of the two eigenvalues
\[
\lambda_{\pm} \in \mathbb{C}.
\]
and two 2x1 eigenvectors \(e_{\pm} \in \mathbb{C}\).

**Example:** Assume that \(T\) is the Fibonacci matrix in Eq. 2.5.7.12 (see p. 53.)

The eigenvalues \(\lambda_{\pm}\) may be merged into a 2 \(\times\) 2 diagonal eigenvalue matrix
\[
\Lambda = \begin{bmatrix}
\lambda_+ & 0 \\
0 & \lambda_-
\end{bmatrix},
\]
while the two eigenvectors \(e_+\) and \(e_-\) are merged into a 2 \(\times\) 2 eigenvector matrix
\[
E = [e_+, e_-] = \begin{bmatrix}
e_1^+ & e_1^- \\
e_2^+ & e_2^-
\end{bmatrix},
\]
corresponding to the two eigenvalues. Using matrix notation, this may be compactly written as
\[
T E = E \Lambda.
\]

Note that while \(\lambda_{\pm}\) and \(E_{\pm}\) commute, \(E \Lambda \neq \Lambda E\).

From Eq. 3.2.1.5 we may obtain two very important forms:

1. the diagonalization of \(T\),
\[
\Lambda = E^{-1} T E,
\]
and

2. the eigenexpansion of \(T\),
\[
T = E \Lambda E^{-1},
\]

which is used for computing powers of \(T\) (i.e., \(T^{100} = E^{-1} \Lambda^{100} E\)).

**Example:** If we take
\[
T = \begin{bmatrix}
1 & 1 \\
1 & -1
\end{bmatrix},
\]
then the eigenvalues are given by \((1 - \lambda_+)(1 + \lambda_-) = -1\); thus \(\lambda_{\pm} = \pm \sqrt{2}\). This method of eigenanalysis is discussed on page 50 and in appendix B.2 (see p. 235).

**Exercise #20**
Show that the geometric series formula holds for 2 \(\times\) 2 matrices. Starting with the 2 \(\times\) 2 identity matrix \(I_2\) and \(a \in \mathbb{C}\), with \(|a| < 1\), show that
\[
I_2(a I_2 - a I_2)^{-1} = I_2 + a I_2 + a^2 I_2^2 + a^3 I_2^3 + \cdots.
\]

**Solution:** Since \(a^k I_2^k = a^k I_2\), we may multiply both sides by \(I_2 - a I_2^2\) to obtain
\[
I_2 = I_2 + a I_2 + a^2 I_2^2 + a^3 I_2^3 + \cdots - a I_2(a I_2 + a^2 I_2^2 + a^3 I_2^3 + \cdots) = [1 + (a + a^2 + a^3 + \cdots) - (a + a^2 + a^3 + a^4 + \cdots)] I_2 = I_2.
\]

This equality requires that the two series converge, which requires that \(|a| < 1\).

\(^9\)During the discovery or creation of quantum mechanics, two alternatives were developed: Schrödinger’s differential equation method and Heisenberg’s matrix method. Eventually it was realized the two were equivalent.
Exercise #21
When the matrix $T$ is not a square matrix, Eq. 3.2.1.3 may be generalized as

$$ T_{m,n} = U_{m,m} \Lambda_{m,n} V_{n,n}^\dagger. $$

**Solution:** This useful generalization of eigenanalysis is called a singular value decomposition (SVD). To see this use the Matlab/Octave command $[U,L,V]=\text{svd}(A)$ where $A$ is a rectangular (nonsquare) matrix.

**Summary:** The GCD (Euclidean algorithm), Pell’s equation, and the Fibonacci sequence may all be written as compositions of $2 \times 2$ matrices. Thus Pell’s equation and the Fibonacci sequence are special cases of the $2 \times 2$ matrix composition

$$ \begin{pmatrix} x \\ y \end{pmatrix}_{n+1} = \begin{pmatrix} a & b \\ c & d \end{pmatrix} \begin{pmatrix} x \\ y \end{pmatrix}_n. $$

This is an important and common thread of these early mathematical findings. This $2 \times 2$ linearized matrix recursion plays a special role in physics, mathematics, and engineering because one-dimensional system equations are solved using the $2 \times 2$ eigenanalysis method. More than several thousand years of mathematical trial and error set the stage for this breakthrough. But it took even longer to be fully appreciated.

The key idea of the $2 \times 2$ matrix solution, widely used in modern engineering, can be traced back to Brahmagupta’s solution of Pell’s equation for arbitrary $N$. Brahmagupta’s recursion, identical to that of the Pythagoreans’ $N=2$ case (see Eq. 2.5.6.9, p. 52), eventually led to the concept of linear algebra, defined by the simultaneous solutions of many linear equations. The recursion by the Pythagoreans (6th century BCE) predated the creation of algebra by al-Khwarizmi (ninth century CE), as seen in Fig. 1.1 (see p. 13).

### 3.2.2 Taylor series

An analytic function is one that meets these criteria:

1. It may be expanded in a Taylor series:

$$ P(x) = \sum_{n=0}^{\infty} c_n (x - x_o)^n. \tag{3.2.2.8} $$

2. It converges for $|x - x_o| < 1$, called the region of convergence (RoC), with coefficients $c_n$.

3. The Taylor series coefficients $c_n$ are defined by taking derivatives of $P(x)$ and evaluating them at the expansion point $x_o$—namely,

$$ c_n = \frac{1}{n!} \frac{d^n}{dx^n} P(x) \bigg|_{x=x_o}. \tag{3.2.2.9} $$

4. Although $P(x)$ may be multivalued, the Taylor series is always single-valued.

**Properties:** The Taylor formula is a prescription for how to uniquely define the coefficients $c_n$. Without the Taylor series formula, we would have no way of determining $c_n$. The proof of the Taylor formula is transparent—simply taking successive derivatives of Eq. 3.2.2.8 and then evaluating the result at the expansion point. If $P(x)$ is analytic, then this procedure always works. If $P(x)$ fails to have a derivative of any order, then the function is not analytic and Eq. 3.2.2.8 is not valid for $P(x)$. For example, if $P(x)$ has a pole at $x_o$, then it is not analytic at that point.

The Taylor series representation of $P(x)$ has special applications for solving differential equations for these reasons:

1. It is single-valued.
2. All its derivatives and integrals are uniquely defined.
3. It may be continued into the complex plane by extending $x \in \mathbb{C}$. Typically this involves expanding the series about a different expansion point.
Analytic continuation: A limitation of the Taylor series expansion is that it is not valid outside of its RoC. One method for avoiding this limitation is to move the expansion point. This is called analytic continuation. However, analytic continuation is a nontrivial operation because it (1) requires manipulating an infinite number of derivatives of \( P(x) \), (2) at the new expansion point \( x_o \), where (3) \( P(x-x_o) \) may not have derivatives, due to possible singularities. (4) Thus one needs to know where the singularities of \( P(s) \) are in the complex \( s \) plane. Due to these many problems, analytic continuation is rarely used, other than as an important theoretical concept.

Example: The trivial case is the geometric series \( P(x) = 1/(1-x) \) about the expansion point \( x = 1 \). The function \( P(x) \) is defined everywhere, except at the singular point \( x = 1 \), whereas the geometric series is valid for only \( |x| < 1 \).

Exercise #22
Verify that \( c_0 \) and \( c_1 \) of Eq. 3.2.2.8 follow from Eq. 3.2.2.9.

Solution: To obtain \( c_0 \), for \( n = 0 \), there is no derivative \((d^n/dx^n) \) indicates no derivative is taken), so we must simply evaluate \( P(x-x_o) = c_0 + c_1(x-x_o) + \cdots \) at \( x = x_o \), leaving \( c_0 \). To find \( c_1 \), we take one derivative, which results in \( P'(x) = c_1 + 2c_2(x-x_o) + \cdots \). Evaluating this at \( x = x_o \) leaves \( c_1 \). Each time we take a derivative we reduce the degree of the series by 1, leaving the next constant term.

Exercise #23
Suppose we truncate the Taylor series expansion to \( N \) terms. What is the name of such functions?

Solution: When an infinite series is truncated, the resulting function is called an \( N \)th-degree polynomial:

\[ P_N(x) = \sum_{n=0}^{N} c_n(x-x_o) + c_2(x-x_o)^2 \cdots c_N(x-x_o)^N. \]

We can find \( c_0 \) by evaluating \( P_N(x) \) at the expansion point \( x_o \), since from the above formula \( P_N(x_o) = c_0 \). From the Taylor formula, \( c_1 = P_N'(x)|_{x_o} \).

Exercise #24
How many roots do \( P_N(x) \) and \( P_N'(x) \) have?

Solution: According to the fundamental theorem of algebra, \( P_N(x) \) has \( N \) roots and \( P_N'(x) \) has \( N - 1 \) roots. The Gauss-Lucas theorem states that the \( N - 1 \) roots of \( P_N'(x) \) lie inside the convex hull of the \( N \) roots of \( P_N(x) \) (see p. 69).

Exercise #25
Would it be possible to find an inverse Gauss-Lucas theorem that states where the roots of the integral of a polynomial might be?

Solution: To the best of my knowledge this problem has not been addressed. However it seems a question worthy of some thought.

With each integral there is a new degree of freedom that must be accommodated. Thus this problem is difficult. But since there is only one extra degree of freedom, it does not seem impossible. To solve this problem a constraint is needed.

Role of the Taylor series: The Taylor series plays a key role in the mathematics of differential equations and their solution, as the coefficients of the series uniquely determine the analytic series representation via its derivatives. The implications and limitations of the power series representation are very specific: If the series fails to converge (i.e., outside the RoC), it is essentially meaningless.

A very important fact about the RoC: It is relevant to only the series, not the function being expanded. Typically the function has a pole at the radius of the RoC, where the series fails to converge. However, the function being expanded is valid everywhere other than at the pole. It seems that this point has been poorly explained in many texts. In addition, the RoC is the region of divergence (RoD), which is the RoC’s complement.

The Taylor series does not need to be infinite to converge to the function it represents, since it obviously works for any polynomial \( P_N(x) \) of degree \( N \). But in the finite case \((N < \infty)\), the RoC is infinite and the series is the function \( P_N(x) \) exactly, everywhere. Of course, \( P_N(x) \) is called a polynomial of degree \( N \). When \( N \to \infty \), the Taylor series is valid only within the RoC, and it is (typically) the representation of the reciprocal of a polynomial.
These properties are both the curse and the blessing of the analytic function. On the positive side, analytic functions are the ideal starting point for solving differential equations, which is exactly how they were used by Newton and others. Analytic functions are “smooth,” since they are infinitely differentiable, with coefficients given by Eq. 3.2.2.9. They are single-valued, so there can be no ambiguity in their interpretation. On the negative side, they only represent the function within the RoC, which depends on the expansion point.

Two well-known analytic functions are the geometric series ($|x| < 1$)

$$\frac{1}{1 - x} = 1 + x + x^2 + x^3 + \cdots = \sum_{n=0}^{\infty} x^n \quad \text{(3.2.2.10)}$$

and the exponential series ($|x| < \infty$)

$$e^x = 1 + x + \frac{1}{2}x^2 + \frac{1}{3 \cdot 2}x^3 + \frac{1}{4 \cdot 3 \cdot 2}x^4 + \cdots = \sum_{n=0}^{\infty} \frac{1}{n!}x^n. \quad \text{(3.2.2.11)}$$

**Exercise #26**

Provide the Taylor series expression for the following functions:

$$F_1(x) = \int \frac{1}{1 - x} \, dx \quad \text{(3.2.2.12)}$$

**Solution:**

$$F_1(x) = x + \frac{1}{2}x^2 + \frac{1}{3}x^3 + \cdots \quad \text{■}$$

$$F_2(x) = \frac{d}{dx} \frac{1}{1 - x} \quad \text{(3.2.2.13)}$$

**Solution:**

$$F_2(x) = 1 + 2x + 3x^2 + \cdots \quad \text{■}$$

$$F_3(x) = \ln \frac{1}{1 - x} \quad \text{(3.2.2.14)}$$

**Solution:**

$$F_3(x) = 1 + \frac{1}{2}x + \frac{1}{3}x^2 + \cdots \quad \text{■}$$

$$F_4(x) = \frac{d}{dx} \ln \frac{1}{1 - x} \quad \text{(3.2.2.15)}$$

**Solution:**

$$F_4(x) = 1 + x + x^2 + x^3 + \cdots \quad \text{■}$$

**Exercise #27**

Using symbolic manipulation (Matlab, Octave, Mathematica), expand the function $F(s)$ in a Taylor series and find the recurrence relationships among the Taylor coefficients $c_n, c_{n-1}, c_{n-2}$. Assume $a \in \mathbb{C}$ and $T \in \mathbb{R}$.

$$F(s) = e^{as}$$

**Solution:**

A Google search on "octave syms taylor" is useful. The Matlab/Octave code to expand this in a Taylor series is

```
syms s
Taylor = taylor(exp(s), s, 0, 'order', 10)
```

**Exercise #28**

Find the coefficients of the following functions by the method of Eq. 3.2.2.9 and give the RoC.

1. $w(x) = \frac{1}{1-x^2}$.

**Solution:** From a straightforward expansion we know the coefficients are

$$\frac{1}{1-x^2} = 1 + x + (x^2)^2 + (x^2)^3 + \cdots = 1 + x - x^2 - x^3 + \cdots.$$
Determining the RoC for a given analytic function is quite important and may not always be obvious. In general the RoC is a circle whose radius extends from the expansion point out to the nearest pole. Thus when the expansion point is moved, the RoC changes, since the location of the pole is fixed.

Example: For the geometric series (Eq. 3.2.2.10), the expansion point is \( x_o = 0 \) and the RoC is \(|x| < 1\), since \( 1/(1 - x) \) has a pole at \( x = 1 \). We may move the expansion point by a linear transformation – for example, by
replacing $x$ with $z + 3$. Then the series becomes $1/((z + 3) - 1) = 1/(z + 2)$, so the RoC becomes 3 because in the $z$ plane the pole has moved to $-2$.

**Example:** A second important example is the function $1/(x^2 + 1)$, which has the same RoC as the geometric series, since it may be expressed in terms of its residue expansion (also called its partial fraction expansion)

$$
\frac{1}{x^2 + 1} = \frac{1}{(x + 1)(x - 1)} = \frac{1}{2j} \left( \frac{1}{x - 1} - \frac{1}{x + 1} \right).
$$

Each term has an RoC of $|x| < 1$, $|j| = 1$. The amplitude of each pole is called the residue, defined in Eq. 4.5.1.4 (see p. 152). The residue for the pole at $1j$ is $1/2j$.

The roots must be found by factoring the polynomial (e.g., Newton’s method). Once the roots are known, the residues are best found with linear algebra.

In summary, the function $1/(x^2 + 1)$ is the sum of two geometric series, with poles at $\pm 1j$, which is not initially obvious because the roots are complex and conjugate. Only when the function is factored does it become clear what is going on.

**Exercise #30**

Verify that the above expression is correct and show that the residues are $\pm 1/2j$.

**Solution:** We cross-multiply and cancel, leaving 1, as required. The RoC is the coefficient on the pole. Thus the residue of the pole at $xj$ is $j/2$. ■

**Exercise #31**

Find the residue of $\frac{d}{dz} z^\pi$.

**Solution:** Taking the derivative gives $\pi z^{\pi-1}$, which has a pole at $z = 0$. Applying the formula for the residue (Eq. 4.5.1.4, p. 152), we find

$$
c^{-1} = \pi \lim_{z \to 0} zz^{\pi-1} = \pi \lim_{z \to 0} z^\pi = 0.
$$

Thus the residue is zero. ■

### 3.2.3 Analytic functions

Any function that has a Taylor series expansion is called an analytic function. Within the RoC, the series expansion defines a single-valued function. Polynomials $1/(1 - x)$ and $e^x$ are examples of analytic functions that are real functions of their real argument $x$.

Every analytic function has a corresponding differential equation, which is determined by the coefficients $a_k$ of the analytic power series. An example is the exponential, which has the property that it is the eigenfunction of the derivative operation

$$
\frac{d}{dx} e^{ax} = ae^{ax},
$$

which may be verified using Eq. 3.2.2.11. This relationship is a common definition of the exponential function, which is special because it is the eigenfunction of the derivative.

The complex analytic power series (i.e., complex analytic functions) may also be integrated term by term, since

$$
\int^x f(x)dx = \sum \frac{a_k}{k + 1} x^{k+1}. \quad (3.2.3.19)
$$

Newton took full advantage of this property of the analytic function and used the analytic series (Taylor series) to solve analytic problems, especially for working out integrals. This allowed him to solve differential equations. To fully understand the theory of differential equations, one must master single-valued analytic functions and their analytic power series.

**Single- vs. multivalued functions** Polynomials and their $\infty$-degree extensions (analytic functions) are single-valued: For each $x$ there is a single value for $P_N(x)$. The roles of the domain and codomain may be swapped to obtain an inverse function with properties that can be very different from those of the function. For example, $y(x) = x^2 + 1$ has the inverse $x = \pm \sqrt{y - 1}$, which is double-valued and complex when $y < 1$. Periodic functions such as $y(x) = \sin(x)$ are even more “exotic,” since $x(y) = \arcsin(x) = \sin^{-1}(x)$ has an infinite number of $x(y)$ values for each $y$. This problem was first addressed in Bernhard Riemann’s 1851 PhD thesis, written while he was working with Gauss.
Exercise #32
Let \( y(x) = \sin(x) \). Then \( dy/dx = \cos(x) \). Show that \( dx/dy = \pm 1/\sqrt{1 - y^2} \).

Solution: Since \( \sin^2 x + \cos^2 x = 1 \), it follows that \( y^2(x) + (dy/dx)^2 = 1 \). Thus \( dy/dx = \pm \sqrt{1 - y^2} \). Taking the reciprocal gives the result result.

To fully understand, Google “implicit function theorem” (D’Angelo, 2017, p. 104).

Exercise #33
Evaluate the integral
\[
I(y) = \int_x^y \frac{dy}{\sqrt{1 - y^2}}.
\]

Solution: From Exercise #32 we know that
\[
x(y) = \int^x dy = \int^y dy = \int_x^y \frac{dy}{\sqrt{1 - y^2}}.
\]

But since \( y(x) = \sin(x) \), it follows that \( x(y) = \sin^{-1} y = \arcsin(y) \).

Exercise #34
Find the Taylor series coefficients of \( y = \sin(x) \) and \( x = \sin^{-1}(y) \). Hint: Use symbolic Octave. Note \( \sin^{-1}(y) = \arcsin(y) \).

Solution: syms s; taylor(sin(s), 'order', 10);
\[
\sin(s) = s - s^3/3! + s^5/5! - s^7/7! + \cdots
\]

and syms s; taylor(asin(s), 'order', 15);
\[
\arcsin(s) = s + \frac{1}{6} s^2 + \frac{3}{40} s^3 + \frac{5}{112} s^4 + \frac{35}{288} s^5 + \frac{63}{1344} s^6 + \frac{231}{139024} s^7 + \cdots
\]

Note that every complex analytic function may be expanded in a Taylor series, within its RoC. It follows that the inverse is also complex analytic, as demonstrated in this case using symbolic algebra.

Exercise #35
What is the necessary condition so that if \( dy/dx = F(x) \), then \( dx/dy = 1/F(x) \).

Solution: This will be true when \( df(x)/dx = F(x) \) is complex analytic because the Fundamental Theorem of Complex Calculus (FTCC) (see p. 135) defines the antiderivative. In this case \( dy/dx = (dx/dy)^{-1} \) (except at singular points, where it is not analytic).

3.2.4 Complex analytic functions
We are given that the argument of an analytic function \( F(x) \) is complex, that is; \( x \in \mathbb{R} \) is replaced by \( s = \sigma + \omega t \in \mathbb{C} \). Recall that \( \mathbb{R} \subset \mathbb{C} \). Thus
\[
F(s) = \sum_{n=0}^{\infty} c_n (s - s_0)^n
\]

(3.2.4.20)

with \( c_n \in \mathbb{C} \). In this case, that function is said to be a complex analytic.

An important example is when the exponential becomes complex, since
\[
e^{\sigma t} = e^{(\sigma + \omega t)} = e^{\sigma t}e^{\omega t} = e^{\sigma t} [\cos(\omega t) + j \sin(\omega t)]
\]

Taking the real part gives
\[
\Re \{e^{\sigma t}\} = e^{\sigma t} e^{\omega t} + e^{-\omega t} = e^{\sigma t} \cos(\omega t)
\]

and \( \Im \{e^{\sigma t}\} = e^{\sigma t} \sin(\omega t) \). Once the argument is allowed to be complex, it becomes obvious that the exponential and circular functions are fundamentally related. This exposes the family of entire circular functions \( \{e^s, \sin(s), \cos(s), \tan(s), \cosh(s), \sinh(s)\} \) and their inverses \( \{\ln(s), \arcsin(s), \arccos(s), \arctan(s), \cosh^{-1}(s), \sinh^{-1}(s)\} \), first fully elucidated by Euler in about 1750 (Stillwell, 2010, p. 315).

Note that because \( \sin(\omega t) \) is periodic, its inverse must be multivalued. What was needed is some systematic way to account for this multivalued property. This extension to multivalued functions is called a branch cut, invented by Riemann in his 1851 PhD thesis, supervised by Gauss in the final years of Gauss’s life.
The Taylor series of a complex analytic function: However, there is a fundamental problem: We cannot formally define the Taylor series for the coefficients $c_k$ until we have defined the derivative with respect to the complex variable $dF(s)/ds$, with $s \in \mathbb{C}$. Thus simply substituting $s$ for $x$ in an analytic function leaves a major hole in one’s understanding of the complex analytic function.

It was Cauchy in 1814 (Fig. 3.1, p. 62) who uncovered the much deeper relationships within complex analytic functions (see p. 125) by defining differentiation and integration in the complex plane, leading to several fundamental theorems of complex calculus, including the fundamental theorem of complex calculus and Cauchy’s formula. We shall explore these fundamental theorems on pages 135-135.

There seems to be some disagreement as to the status of multivalued functions: Are they functions, or is a function strictly single-valued? If so, then we are missing out on a host of interesting possibilities, including all the inverses of nearly every complex analytic function. For example, the inverse of a complex analytic function is a complex analytic function (e.g., $e^s$ and $\log(s)$).

Impact of complex analytic mathematics on physics: It seems likely, if not obvious, that the success of Newton was his ability to describe physics by the use of mathematics. He was inventing new mathematics at the same time he was explaining new physics. The same might be said for Galileo. It seems likely that Newton was extending the successful techniques and results of Galileo’s work on gravity (Galileo, 1638). Galileo died on January 8, 1642, and Newton was born January 4, 1643, just short of one year later. Obviously Newton was well aware of Galileo’s great success and naturally would have been influenced by him (see p. 16).

The application of complex analytic functions to physics was dramatic, as may be seen in the six volumes on physics written by Arnold Sommerfeld (1868–1951), and from the productivity of his many (36) students (e.g., Debye, Lenz, Ewald, Pauli, Guillemin, Bethe, Heisenberg, Morse, and Seebach, to name a few), notable coworkers (Leon Brillouin), and others (John Bardeen) upon whom Sommerfeld had a strong influence. Sommerfeld is famous for training many students who were awarded the Nobel Prize in Physics, yet he never won a Nobel Prize (the prize is not awarded in mathematics). Sommerfeld brought mathematical physics (the merging of physical and experimental principles via mathematics) to a new level with the use of complex integration of analytic functions to solve otherwise difficult problems, thus following the lead of Newton, who used real integration of Taylor series to solve differential equations (Brillouin, 1960, Ch. 3 by Sommerfeld).
3.3 Problems AE-1

Topics of this homework: Fundamental theorem of algebra, polynomials, analytic functions and their inverse, convolution, Newton’s root finding method, Riemann zeta function. Deliverables: Answers to problems

Note: The term analytic is used in two different ways. (1) An analytic function is a function that may be expressed as a locally convergent power series; (2) analytic geometry refers to geometry using a coordinate system.

Polynomials and the fundamental theorem of algebra (FTA)

Problem # 1: A polynomial of degree $N$ is defined as

$$P_N(x) = a_0 + a_1 x + a_2 x^2 + \cdots + a_N x^N.$$  

– 1.1: How many coefficients $a_n$ does a polynomial of degree $N$ have?

– 1.2: How many roots does $P_N(x)$ have?

Problem # 2: The fundamental theorem of algebra (FTA)

– 2.1: State and then explain the FTA.

– 2.2: Using the FTA, prove your answer to the preceding problem. Hint: Apply the FTA to prove how many roots a polynomial $P_N(x)$ of order $N$ has.

Problem # 3: Consider the polynomial function $P_2(x) = 1 + x^2$ of degree $N = 2$ and the related function $F(x) = 1/P_2(x)$. What are the roots (e.g., zeros) $x_{\pm}$ of $P_2(x)$? Hint: Complete the square on the polynomial $P_2(x) = 1 + x^2$ of degree 2, and find the roots.

Problem # 4: $F(x)$ may be expressed as $(A, B, x_{\pm} \in \mathbb{C})$

$$F(x) = \frac{A}{x - x_+} + \frac{B}{x - x_-},$$  

(AE-1.1)

where $x_{\pm}$ are the roots (zeros) of $P_2(x)$, which become the poles of $F(x)$; $A$ and $B$ are the residues. The expression for $F(x)$ is sometimes called a partial fraction expansion or residue expansion, and it appears in many engineering applications.

– 4.1: Find $A, B \in \mathbb{C}$ in terms of the roots $x_{\pm}$ of $P_2(x)$.

– 4.2: Verify your answers for $A$ and $B$ by showing that this expression for $F(x)$ is indeed equal to $1/P_2(x)$.

– 4.3: Give the values of the poles and zeros of $P_2(x)$.

– 4.4: Give the values of the poles and zeros of $F(x) = 1/P_2(x)$. 


Analytic functions

Overview: Analytic functions are defined by infinite (power) series. The function \( f(x) \) is analytic at any value of \( x = x_0 \) where there exists a convergent power series

\[
P(x) = \sum_{n=0}^{\infty} a_n x^n
\]

such that \( P(x_0) = f(x_0) \). The local power series for \( f(x) \) near \( x = x_0 \) is often obtained by finding the Taylor series:

\[
f(x) \approx f(x_0) + \left. \frac{df}{dx} \right|_{x=x_0} (x-x_0) + \frac{1}{2!} \left. \frac{d^2f}{dx^2} \right|_{x=x_0} (x-x_0)^2 + \cdots
\]

\[
= \sum_{n=0}^{\infty} \frac{1}{n!} \left. \frac{d^n f}{dx^n} \right|_{x=x_0} (x-x_0)^n.
\]

The point \( x = x_0 \) is called the expansion point. When the expansion point is at \( x_0 = 0 \), the series is called a MacLaurin series.

Two classic examples are the geometric series where \( a_n = 1 \),

\[
\frac{1}{1-x} = 1 + x + x^2 + x^3 + \cdots = \sum_{n=0}^{\infty} x^n, \quad (AE-1.2)
\]

and the exponential function where \( a_n = 1/n! \), Eq. 3.2.2.11 (p. 78). The coefficients for both series may be derived from the Taylor formula (or the MacLaurin formula when the expansion point is zero).

Problem #5: The geometric series

- 5.1: What is the region of convergence (RoC) for the power series Eq. AE-1.2 of \( 1/(1-x) \) given above—for example, where does the power series \( P(x) \) converge to the function value \( f(x) \)? State your answer as a condition on \( x \). Hint: What happens to the power series when \( x > 1 \)?

- 5.2: In terms of the pole, what is the RoC for the geometric series in Eq. AE-1.2?

- 5.3: How does the RoC relate to the location of the pole of \( 1/(1-x) \)?

- 5.4: Where are the zeros, if any, in Eq. AE-1.2?

- 5.5: Assuming \( x \) is in the RoC, prove that the geometric series correctly represents \( 1/(1-x) \) by multiplying both sides of Eq. AE-1.2 by \( (1-x) \).

Problem #6: Use the geometric series to study the degree \( N \) polynomial. It is very important to note that all the coefficients \( c_n \) of this polynomial are 1.

\[
P_N(x) = 1 + x + x^2 + \cdots + x^N = \sum_{n=0}^{N} x^n. \quad (AE-1.3)
\]

- 6.1: Prove that

\[
P_N(x) = \frac{1 - x^{N+1}}{1-x}. \quad (AE-1.4)
\]

The geometric series is not defined as the function \( 1/(1-x) \), it is defined as the series \( 1 + x + x^2 + x^3 + \cdots \), such that the ratio of consecutive terms is \( x \).
3.3. PROBLEMS AE-1

– 6.2: What is the RoC for Eq. AE-1.3?

– 6.3: What is the RoC for Eq. AE-1.4?

– 6.4: How many poles does \( P_N(x) \) (Eq. AE-1.3) have? Where are they?

– 6.5: How many zeros does \( P_N(x) \) (Eq. AE-1.4) have? State where are they in the complex plane.

– 6.6: Explain why Eqs. AE-1.3 and AE-1.4 have different numbers of poles and zeros.

– 6.7: Is the function \( 1/(1 - x) \) analytic outside of the RoC stated in the first question in Problem 5? Hint: Can it be represented by a different power series outside this RoC?

– 6.8: Extra credit. Evaluate \( P_N(x) \) at \( x = 0 \) and \( x = 0.9 \) for the case of \( N = 100 \), and compare the result to that from Matlab.

```matlab
%sum the geometric series and P_100(0.9)
clear all; close all; format long
N=100; x=0.9; S=0;
for n=0:N
    S=S+xˆn
end
P100=(1-xˆ(N+1))/(1-x);
disp(sprintf('S= %g, P100= %g, error= %g',S,P100, S-P100))
```

Problem # 7 The exponential series

– 7.1: What is the RoC for the exponential series Eq. 3.2.2.11?

– 7.2: Let \( x = j \) in Eq. 3.2.2.11, and write out the series expansion of \( e^x \) in terms of its real and imaginary parts.

– 7.3: Let \( x = j\theta \) in Eq. 3.2.2.11, and write out the series expansion of \( e^x \) in terms of its real and imaginary parts. How does your result relate to Euler’s identity (\( e^{j\theta} = \cos(\theta) + j\sin(\theta) \))? 

Inverse analytic functions and composition

Overview: It may be surprising, but every analytic function has an inverse function. Starting from the function \((x, y \in \mathbb{C})\)

\[ y(x) = \frac{1}{1-x} \]

the inverse is

\[ x = \frac{y - 1}{y} = 1 - \frac{1}{y} \]

Problem # 8: Considering the inverse function described above

– 8.1: Where are the poles and zeros of \( x(y) \)?

– 8.2: Where (for what condition on \( y \)) is \( x(y) \) analytic?
Problem #9 Considering the exponential function $z(x) = e^x$ ($x, z \in \mathbb{C}$).

- 9.1: Find the inverse $x(z)$.
- 9.2: Where are the poles and zeros of $x(z)$?
- 9.3: If $y(s) = 1/(1 - s)$ and $z(s) = e^{s}$. compose these two functions to obtain $(y \circ z)(s)$. Give the expression for $(y \circ z)(s) = y(z(s))$.
- 9.4: Where are the poles and zeros of $(y \circ z)(s)$?
- 9.5: Where (for what condition on $x$) is $(y \circ z)(x)$ analytic?

Convolution

Multiplying two short or simple polynomials is not demanding. However, if the polynomials have many terms, it can become tedious. For example, multiplying two 10th-degree polynomials is not something one would want to do every day.

An alternative is a method called convolution, as described in Sec. 3.4 (p. 89).

Problem #10: Convolution of sequences. Practice convolution (by hand!!) using a few simple examples. Show your work!!! Check your solution using Matlab.

- 10.1: Convolve the sequence \{0 1 1 1\} with itself.
- 10.2: Calculate \{1, 1\} $\star$ \{1, 1\} $\star$ \{1, 1\}

Problem #11: Multiplying two polynomials is the same as convolving their coefficients.

$$
\begin{align*}
f(x) &= x^3 + 3x^2 + 3x + 1 \\
g(x) &= x^3 + 2x^2 + x + 2.
\end{align*}
$$

- 11.1: In Octave/Matlab, compute $h(x) = f(x) \cdot g(x)$ in two ways: (1) use the commands roots and poly, and (2) use the convolution command conv. Confirm that both methods give the same result.
- 11.2: What is $h(x)$?

Newton’s root-finding method

Problem #12: Use Newton’s iteration to find the roots of the polynomial

$$
P_3(x) = 1 - x^3.
$$

- 12.1: Draw a graph describing the first step of the iteration starting with $x_0 = (1/2, 0)$.
- 12.2: Calculate $x_1$ and $x_2$. What number is the algorithm approaching?
- 12.3: Here is a Octave/Matlab script for the $P_2(x)$ case. Modify it to find $P_3(x)$:
3.3. PROBLEMS AE-1

\[ x(1) = 1/2; \quad \% x(1) = 0.9; \quad \% x(1) = -10 \]

\[ y(1) = x(1); \]

\[
\text{for } n=2:10 \\
x(n) = x(n-1) + (1-x(n-1)^2)/(2*x(n-1)); \\
y(n) = (1+y(n-1)^2)/(2*x(n-1)); 
\]

\text{end}

\text{semilogy(abs(x)-1); hold on}

\text{semilogy(abs(7)-1,’or’); hold off}

\section*{5 The Riemann zeta function \( \zeta(s) \)}

\textbf{Verify the sol not removed. Ck edits carefully.}

\textbf{Definitions and preliminary analysis:}

The zeta function \( \zeta(s) \) is defined by the complex analytic power series

\[ \zeta(s) \equiv \sum_{n=1}^{\infty} \frac{1}{n^s} = \frac{1}{1^s} + \frac{1}{2^s} + \frac{1}{3^s} + \frac{1}{4^s} + \cdots. \]

This series converges, and thus is valid, only in the RoC given by \( \Re s = \sigma > 1 \) since there \( |n^{-\sigma}| \leq 1 \). To determine its formula in other regions of the \( s \) plane, one must extend the series via analytic continuation.

\textbf{Euler product formula:} As Euler first published in 1737, one may recursively factor out the leading prime term, which results in Euler’s product formula.\(^{11}\) Multiplying \( \zeta(s) \) by the factor \( 1/2^s \) and subtracting from \( \zeta(s) \) remove all the terms \( 1/(2n)^s \) (e.g., \( 1/2^s + 1/4^s + 1/6^s + 1/8^s + \cdots \))

\[ \left(1 - \frac{1}{2^s}\right) \zeta(s) = 1 + \frac{1}{3^s} + \frac{1}{5^s} + \frac{1}{7^s} + \cdots - \left( \frac{1}{2^s} + \frac{1}{4^s} + \frac{1}{6^s} + \frac{1}{8^s} + \cdots \right), \quad (AE-1.5) \]

which results in

\[ \left(1 - \frac{1}{2^s}\right) \zeta(s) = 1 + \frac{1}{3^s} + \frac{1}{5^s} + \frac{1}{7^s} + \frac{1}{9^s} + \frac{1}{11^s} + \frac{1}{13^s} + \cdots. \quad (AE-1.6) \]

\textbf{Problem # 13: Questions about the Riemann zeta function.}

\begin{itemize}
  \item \textbf{13.1:} What is the RoC for Eq. AE-1.6
  
  \item \textbf{13.2:} Repeat the algebra of Eq. AE-1.5 using the lead factor of \( 1/3^s \).
  
  \item \textbf{13.3:} What is the RoC for Eq. AE-1.7
\end{itemize}

\(^{11}\)This is known as Euler’s sieve, as distinguished from the Eratosthenes sieve.
13.4: Repeat the algebra of Eq. AE-1.5 for all prime scale factors (i.e., \(1/5^s, 1/7^s, \ldots, 1/\pi_k^s, \ldots\)) to show that

\[
\zeta(s) = \prod_{\pi_k \in \mathbb{P}} \frac{1}{1 - \pi_k^{-s}} = \prod_{\pi_k \in \mathbb{P}} \zeta_k(s),
\]

(AE-1.7)

where \(\pi_p\) represents the \(p\)th prime.

13.5: Given the product formula, identify the poles of \(\zeta_p(s) (p \in \mathbb{Z})\), which is important for defining the RoC of each factor. For example, the \(p\)th factor of Eq. AE-1.8, expressed as an exponential, is

\[
\zeta_p(s) \equiv \frac{1}{1 - \pi_p^{-s}} = \frac{1}{1 - e^{-sT_p}},
\]

(AE-1.8)

where \(T_p \equiv \ln \pi_p\).

13.6: Plot Eq. AE-1.9 using \(zviz\) for \(p = 1\). Describe what you see.
3.4 Root classification by convolution

Following the exploration of algebraic relationships by Fermat and Descartes, the first theorem was being formulated by d’Alembert. The idea behind this theorem is that every polynomial of degree \( N \) (Eq. 3.1.1.7) has at least one root. This may be written as the product of the root and a second polynomial of degree of \( N-1 \). By the recursive application of this concept, it is clear that every polynomial of degree \( N \) has \( N \) roots. Today this result is known as the fundamental theorem of algebra:

\[
\text{Every polynomial equation } P(z) = 0 \text{ has a solution in the complex numbers. As Descartes observed,} \\
a \text{ solution } z = a \text{ implies that } P(z) \text{ has a factor } z - a. \text{ The quotient} \\
Q(z) = \frac{P(z)}{z - a} = \frac{P(z)}{a} \left[ 1 + \frac{z}{a} + \left( \frac{z}{a} \right)^2 + \left( \frac{z}{a} \right)^3 + \cdots \right] \tag{3.4.0.1}
\]

is then a polynomial of one lower degree. . . . We can go on to factorize \( P(z) \) into \( n \) linear factors.


The ultimate expression of this theorem is given by Eq. 3.1.1.7 (p. 64), which indirectly states that an \( n^{th} \) degree polynomial has \( n \) roots. We shall use the term degree when speaking of polynomials and the term order when speaking of differential equations. A general rule is that order applies to the time domain and degree to the frequency domain, since the Laplace transform of a differential equation, having constant coefficients, of order \( N \), is a polynomial of degree \( N \) in Laplace frequency \( s \).

Today this theorem is so widely accepted we fail to appreciate it. Certainly about the time you learned the quadratic formula, you were prepared to understand the concept of polynomials having roots. The simple quadratic case may be extended to a higher degree polynomial. The Octave/Matlab command \texttt{roots ([1, a_2, a_1, a_0])} provides the roots \([s_1, s_2, s_3]\) of the cubic equation, defined by the coefficient vector \([1, a_2, a_1, a_0]\). The command \texttt{poly ([s_1, s_2, s_3])} returns the coefficient vector. I don’t know the largest degree that can be accurately factored by Matlab/Octave, but I’m sure its well over \( N = 10^3 \). Today, finding the roots numerically is a solved problem.

The best way to gain insight into the polynomial factorization problem is through the inverse operation, multiplication of monomials. Given the roots \( x_k \), there is a simple algorithm for computing the coefficients \( a_k \) of \( P_N(x) \) for any \( n \), no matter how large. This method is called convolution. Convolution is said to be a trapdoor function, since it is easy, while the inverse, factoring (deconvolution), is hard, and analytically intractable for degree \( N \geq 5 \) (Stillwell, 2010, p. 102).

3.4.1 Convolution of monomials

As outlined by Eq. 3.1.1.7, a polynomial has two equivalent descriptions, first as a series with coefficients \( a_n \) and second in terms of its roots \( x_r \). The question is What is the relationship between the coefficients and the roots? The simple answer is that they are related by convolution.

Let us start with the quadratic

\[
(x + a)(x + b) = x^2 + (a + b)x + ab, \tag{3.4.1.2}
\]

where in vector notation \([-a, -b]\) are the roots and \([1, a + b, ab]\) are the coefficients.

To see how the result generalizes, we may work out the coefficients for the cubic (\( N = 3 \)). Multiplying the following three factors gives

\[
(x - 1)(x - 2)(x - 3) = (x^2 - 3x + 2)(x - 3) = x(x^2 - 3x + 2) - 3(x^2 - 3x + 2) = x^3 - 6x^2 + 11x - 6. \tag{3.4.1.3}
\]

When the roots are \([1, 2, 3]\), the coefficients of the polynomial are \([1, -6, 11, -6]\). To verify, we can substitute the roots into the polynomial and show that they give zero. For example, \( r_1 = 1 \) is a root, since \( P_3(1) = 1 - 6 + 11 - 6 = 0 \).

As the degree increases, the algebra becomes more difficult. Imagine trying to work out the coefficients for \( N = 100 \). What is needed is a simple way of finding the coefficients from the roots. Fortunately, convolution keeps track of the bookkeeping, formalizing the procedure, along with Newton’s deconvolution method for finding the roots of polynomials (see p. 66).
**Convolution of two vectors:** To get the coefficients by convolution, we write the roots as two vectors \([1, a]\) and \([1, b]\). To find the coefficients, we must convolve the vectors, indicated by \([1, a] \ast [1, b]\), where \(\ast\) denotes convolution. Convolution is a recursive operation. The convolution of \([1, a] \ast [1, b]\) is done as follows: Reverse one of the two monomials, padding unused elements with zeros. Next slide one monomial against the other, forming the local scalar product (element-wise multiply and add):

\[
\begin{array}{ccccccccc}
  a & 1 & 0 & 0 & a & 1 & 0 & 0 & a & 1 \\
 0 & 0 & 1 & b & 0 & 1 & b & 0 & 1 & b \\
 0 & = & x^2 & = & (a + b)x & = & abx^0 & = & 0
\end{array}
\]

resulting in coefficients \([\ldots, 0, 0, 1, a + b, ab, 0, 0, \ldots]\).

If we reverse one of the polynomials and then take successive scalar products, all the terms in the sum of the scalar product correspond to the same power of \(x\). This explains why the convolution of the coefficients gives the same answer as the product of the polynomials.

As seen from the above example, the positions of the first monomial coefficients are reversed and then slide across the second set of coefficients, the scalar product is computed, and the result is placed in the output vector. Outside the range shown, all the elements are zero. In summary,

\[[1, -1] \ast [1, -2] = [1, -1 - 2, 2] = [1, -3, 2],\]

In general,

\[[a, b] \ast [c, d] = [ac, bc + ad, bd].\]

Convolving a third term \([1, -3]\) with \([1, -3, 2]\) gives (Eq. 3.4.1.3)

\[[1, -3] \ast [1, -3, 2] = [1, -3 - 3, 9 + 2, -6] = [1, -6, 11, -6],\]

which is identical to the cubic example found by the algebraic method.

When we convolve one monomial factor at a time, the overlap is always two elements; thus it is never necessary to compute more than two multiplications and one addition for each output coefficient. This greatly simplifies the operations (i.e., they are easily done in your head). Thus the final result is more likely to be correct. Comparing this to the algebraic method, we see that convolution has the clear advantage.

**Exercise #36**

What three nonlinear equations would we need to solve to find the roots of a cubic?

**Solution:** From our formula for the convolution of three monomials, we may find the nonlinear deconvolution relationships between the roots \([-a, -b, -c]\) and the cubic’s coefficients \([1, \alpha, \beta, \gamma]\):\(^{12}\)

\[
(x + a) \ast (x + b) \ast (x + c) = (x + c) \ast (x^2 + (a + b)x + ab) = x \cdot (x^2 + (a + b)x + ab) + c \cdot (x^2 + (a + b)x + ab) = x^3 + (a + b + c)x^2 + (ab + ac + cb)x + abc = [1, a + b + c, ab + ac + cb, abc].
\]

It follows that the nonlinear equations must be

\[
\alpha = a + b + c \\
\beta = ab + ac + bc \\
\gamma = abc.
\]

These equations may be solved by the classic cubic solution, which therefore is a deconvolution problem, also known as long division of polynomials. Therefore the following long division of polynomials must be true:

\[
\frac{x^3 + (a + b + c)x^2 + (ab + ac + cb)x + abc}{x + a} = x^2 + (b + c)x + bc.
\]

---

\(^{12}\)By working with the negative roots, we may avoid an unnecessary and messy alternating sign problem.
The product of a monomial $P_i(x)$ and a polynomial $P_N(x)$ gives $P_{N+1}(x)$: This is another way of stating the fundamental theorem of algebra. Each time we convolve a monomial with a polynomial of degree $N$, we obtain a polynomial of degree $N + 1$. The convolution of two monomials results in a quadratic (degree 2 polynomial). The convolution of three monomials gives a cubic (degree 3). In general, the degree $k$ of the product of two polynomials of degree $n, m$ is the sum of the degrees $(k = n + m)$. For example, if the degrees are each 5 ($n = m = 5$), then the resulting degree is 10.

While we all know this theorem from high school algebra class, it is important to explicitly identify the fundamental theorem of algebra.

Note that the degree of a polynomial is one less than the length of the vector of coefficients. Since the leading term of the polynomial cannot be zero, or else the polynomial would not have degree $N$, when we look for roots, the coefficient can (and should always) be normalized to 1.

In summary, the product of two polynomials of degree $m, n$ having $m$ and $n$ roots is a polynomial of degree $m + n$. This is an analysis process of merging polynomials by coefficient convolution. Multiplying polynomials is a merging process into a single polynomial.

**Composition of polynomials:** Convolution is not the only important operation between two polynomials. Another is composition, which may be defined for two functions $f(z)$ and $g(z)$. The composition $c(z) = f(z) ◦ g(z) = f(g(z))$. As a specific example, suppose $f(z) = 1 + z + z^2$ and $g(z) = e^{z^2}$. With these definitions,

\[ f(z) ◦ g(z) = 1 + e^{z^2} + (e^{z^2})^2 = 1 + e^{z^2} + e^{4z}. \]

Note that $f(z) ◦ g(z) \neq g(z) ◦ f(z)$.

**Exercise #37**

Find $g(z) ◦ f(z)$.

**Solution:**

\[ e^{2f(z)} = e^{2(1+z+z^2)} = e^2e^{1+z+z^2} = e^3e^{z^2}. \]

### 3.4.2 Residue expansions of rational functions

As we discussed on page 72, there are eight important Matlab/Octave routines that are closely related: `conv()`, `deconv()`, `poly()`, `polyder()`, `polyval()`, `residue()`, `root()`, `deconv()`. Several of these are complements of each other or do a similar operation in a slightly different way. The routines `conv()` and `poly()` build polynomials from the roots, while `root()` solves for the roots given the polynomial coefficients. The operation `residue()` converts the ratio of two polynomials and expands it in a partial fraction expansion, with poles and residues.

When lines and planes are defined, the equations are said to be linear in the independent variables. In keeping with this definition of linear, we say that the equations are nonlinear when the equations have degree greater than 1 in the independent variables. The term bilinear has a special meaning: Both the domain and codomain are linearly related by lines (or planes). As an example, impedance is defined in frequency as the ratio of the voltage over the current, but it often has a representation as the ratio of two polynomials, $N(s)$ and $D(s)$:

\[ Z(s) = \frac{N(s)}{D(s)} = sL_0 + R_0 + \sum_{k=0}^{K} \frac{K_k}{s - s_k}. \]  (3.4.2.4)

Here $Z(s)$ is the impedance, and $V$ and $I$ are the voltage and current at radial frequency $\omega$.\(^\text{13}\)

Such an impedance is typically specified as a rational or bilinear function—namely, the ratio of two polynomials, $P_N(s) = N(s) = [a_N, a_{n-1}, \ldots, a_0]$ and $P_K(s) = D(s) = [b_K, b_{K-1}, \ldots, b_0]$ of degrees $N, K \in \mathbb{N}$, as functions of complex Laplace frequency $s = \sigma + j\omega$ with simple roots. Most impedances are rational functions, since they may be written as $D(s)V = N(s)I$. Since $D(s)$ and $N(s)$ are both polynomials in $s$, a rational function is also called a bilinear transformation, or in the mathematical literature a Möbius transformation, which comes from a corresponding scalar differential equation of the form

\[ \sum_{k=0}^{K} b_k \frac{d^k}{dt^k} v(t) = \sum_{n=0}^{N} a_n \frac{d^n}{dt^n} v(t) \leftrightarrow I(\omega) \sum_{k=0}^{K} b_k s^k = V(\omega) \sum_{n=0}^{N} a_n s^n. \]  (3.4.2.5)

\(^{13}\)Note that the relationship between the impedance and the residues $K_k$ is a linear one, best solved by setting up a linear system of equations in the unknown residues.
This construction is also known as the ABCD method in the engineering literature (Eq. 3.7.0.1, p. 110). This equation, as well as Eq. 3.4.2.4, follows from the Laplace transform (see p. 119) of the differential equation (on left) by forming the impedance \( Z(s) = V/I = A(s)/B(s) \). This form of the differential equation follows from Kirchhoff’s voltage and current laws (KCL, KVL) or from Newton’s laws (for the case of mechanics).

The physical properties of an impedance: Based on d’Alembert’s observation that the solution to the wave equation is the sum of forward and backward traveling waves, the impedance may be rewritten in terms of forward and backward traveling waves (see p. 142)

\[
Z(s) = \frac{V}{I} = \frac{V^+ + V^-}{I^+ - I^-} = r_o \frac{1 + \Gamma(s)}{1 - \Gamma(s)},
\]

(3.4.2.6)

where \( r_o = P^+/I^+ \) is called the characteristic impedance of the transmission line (e.g., wire) connected to the load impedance \( Z(s) \), and \( \Gamma(s) = V^-/V^+ = I^-/I^+ \) is the reflection coefficient corresponding to \( Z(s) \). Any impedance of this type is called a Brune impedance due to its special properties (Brune, 1931a; Van Valkenburg, 1964a). Like \( Z(s) \), \( \Gamma(s) \) is causal and complex analytic. The impedance and the reflectance function \( \Gamma(s) \) must both be complex analytic, since they are related to the bilinear transformation, which assures the mutual complex analytic properties.

Due to the bilinear transformation, the physical properties of \( Z(s) \) and \( \Gamma(s) \) are very different. Specifically, the real part of the load impedance is nonnegative (\( \mathbb{R}\{Z(\omega)\} \geq 0 \)) if and only if \( |\Gamma(s)| \leq 1 \). In the time domain, the impedance \( z(t) \leftrightarrow Z(s) \) must have a value of \( r_o \) at \( t = 0 \). Correspondingly, the time-domain reflectance \( \gamma(t) \leftrightarrow \Gamma(s) \) must be zero at \( t = 0 \).

This is the basis of conservation of energy, which may be traced back to the properties of the reflectance \( \Gamma(s) \).

Exercise #38
Show that if \( \mathbb{R}\{Z(s)\} \geq 0 \), then \( |\Gamma(s)| \leq 1 \).

Solution: Taking the real part of Eq. 3.4.2.6, which must be \( \geq 0 \), we find

\[
\mathbb{R}\{Z(s)\} = \frac{r_o}{2} \left[ \frac{1 + \Gamma(s)}{1 - \Gamma(s)} + \frac{1 + \Gamma^*(s)}{1 - \Gamma^*(s)} \right] = \frac{r_o}{|1 + \Gamma(s)|^2} \geq 0.
\]

Thus \(|\Gamma| \leq 1 \). ■

3.5 Introduction to Analytic Geometry

Analytic geometry came about as Euclid’s geometry merged with algebra. The combination of Euclid’s (323 BCE) geometry and al-Khwarizmi’s (830 CE) algebra resulted in a totally new and powerful tool, analytic geometry, independently worked out by Descartes and Fermat (Stillwell, 2010). The addition of matrix algebra during the 18th century enabled analysis in more than three dimensions, which today is one of the most powerful tools used in artificial intelligence, data science, and machine learning. The utility and importance of these new tools cannot be overstated. The timeline for this period of development in mathematics is shown in Fig. 1.2 (see p. 15).

There are many important relationships between Euclidean geometry and 16th-century algebra. Table 3.1 is an attempt at a detailed comparison. Important similarities include vectors, their Pythagorean lengths \([a, b, c]\),

\[
c = \sqrt{(x_2 - x_1)^2 + (y_2 - y_1)^2},
\]

(3.5.0.1)

\( a = x_2 - x_1 \), and \( b = y_2 - y_1 \), and the angles. Euclid’s geometry had length and angles but no concept of coordinates or thus of vectors. One of the main innovations of analytic geometry is that we could compute with real, and soon after, complex numbers.

Several new concepts came with the development of analytic geometry:

1. Composition of functions: If \( y = f(x) \) and \( z = g(y) \), then the composition of functions \( f \) and \( g \) is denoted \( z(x) = g \circ f(x) = g(f(x)) \).
2. Elimination: Given two functions \( f(x, y) \) and \( g(x, y) \), elimination removes either \( x \) or \( y \). This procedure, known to the Chinese, is called Gaussian elimination.
3. Intersection: One may speak of the intersection of two lines to define a point or two planes to define a line. This is a special case of elimination when the functions \( f(x, y) \) and \( g(x, y) \) are linear in their arguments. The term intersection is also an important but very different concept in set theory.
4. Vectors: Analytic geometry provides the concept of a vector (see Appendix A, p. 221) as a line with length and orientation (i.e., direction). Analytic geometry defines vectors in any number of dimensions as ordered sets of points.

5. Scalar and vector products: Analytic geometry extends the ideas of Euclidean geometry with the introduction of the scalar (dot) product of two vectors $f \cdot g$ and the vector (cross) product $f \times g$ (see Fig. 3.4).

Table 3.1: An ad hoc comparison of Euclidean geometry and analytic geometry. I am uncertain of the classification of the items in the third column.

<table>
<thead>
<tr>
<th>Euclidean geometry: $\mathbb{R}^3$</th>
<th>Analytic geometry: $\mathbb{R}^n$</th>
<th>Uncertain</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Proof</td>
<td>Numbers</td>
<td>Recursion</td>
</tr>
<tr>
<td>• Line length</td>
<td>Algebra</td>
<td>Iteration $\in \mathbb{C}^2$, (Newton’s method)</td>
</tr>
<tr>
<td>Line intersection</td>
<td>Power series</td>
<td></td>
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<tr>
<td>Point</td>
<td>Analytic functions</td>
<td></td>
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<tr>
<td>Projection (scalar product)</td>
<td>Complex analytic functions:</td>
<td></td>
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<tr>
<td>Line direction</td>
<td>$\sin \theta$, $\cos \theta$, $e^{\theta i}$, $\log z$</td>
<td></td>
</tr>
<tr>
<td>Vector (sort of)</td>
<td>Scalar product</td>
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<tr>
<td>Conic section</td>
<td>Cross product</td>
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<tr>
<td>Square roots (spiral of Theodorus)</td>
<td>Generalized vector product</td>
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<td>Normed vector spaces</td>
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<td>Composition</td>
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<tr>
<td></td>
<td>Fundamental theorem of algebra</td>
<td></td>
</tr>
</tbody>
</table>

What algebra also added to geometry was the ability to compute with complex numbers. For example, in geometry the length of a line (Eq. 3.5.0.1) was measured with a compass; numbers played no role. Once algebra was available, the line’s Euclidean length could be computed numerically, directly from the coordinates of the two ends, defined by the 3-vector

$$e = x\mathbf{\hat{x}} + y\mathbf{\hat{y}} + z\mathbf{\hat{z}} = [x, y, z]^T,$$

which represents a point at $(x, y, z) \in \mathbb{R}^3 \subset \mathbb{C}^3$ in three dimensions, having direction from the origin $(0, 0, 0)$ to $(x, y, z)$. An alternative matrix notation is $e = [x, y, z]^T$, a column vector of three numbers. These two notations are different ways of representing the vector $e$.

By defining the vector, analytic geometry allows Euclidean geometry to become quantitative, beyond the physical drawing of an object (e.g., a sphere, triangle, or line). With analytic geometry we have the Euclidean concept of a vector, a line that has a magnitude (length) and direction, but analytic, defined in terms of physical coordinates (i.e., numbers). The difference between two vectors defines a third vector, a concept already present in Euclidean geometry. For the first time, complex numbers were allowed into geometry (but rarely used before Cauchy and Riemann).

Scalar product of two vectors: When we use algebra, many concepts that are obvious with Euclid’s geometry may be made precise. There are many examples of how algebra extends Euclidean geometry, the most basic being the scalar product (also known as the dot product) between vectors $x \in \mathbb{R}^3$ and $\kappa \in \mathbb{C}^3$:

$$x \cdot \kappa = (x\mathbf{\hat{x}} + y\mathbf{\hat{y}} + z\mathbf{\hat{z}}) \cdot (\alpha\mathbf{\hat{x}} + \beta\mathbf{\hat{y}} + \gamma\mathbf{\hat{z}}) \in \mathbb{C}$$

$$= \alpha x + \beta y + \gamma z.$$
Represented by a convergent series that has Taylor coefficients and are integrable term by term. Functions are analytic, there is no obvious reason that this should be a problem, since analytic functions may be vector space. This intuitive and somewhat obvious idea is powerful. In this case the scalar product can be defined (i.e., \( \langle s \rangle \in \mathbb{R} \)).

The length is a concept of Euclidean geometry, and it must always be positive and real. A complex (or negative) generalization of the length, in any number of dimensions, that forces the sign of the square root to be nonnegative. The norm of a vector \( \| \mathbf{e} \| \equiv +\sqrt{\mathbf{e} \cdot \mathbf{e}} \geq 0 \)

is defined as the positive square root of the scalar product of the vector with itself (see Appendix A.3). This is a generalization of the length, in any number of dimensions, that forces the sign of the square root to be nonnegative. The length is a concept of Euclidean geometry, and it must always be positive and real. A complex (or negative) length is not physically meaningful. More generally, the Euclidean length of a line is given as the norm of the difference between two real vectors \( \mathbf{e}_1, \mathbf{e}_2 \in \mathbb{R} \):

\[
||\mathbf{e}_1 - \mathbf{e}_2||^2 = (\mathbf{e}_1 - \mathbf{e}_2) \cdot (\mathbf{e}_1 - \mathbf{e}_2)
= (x_1 - x_2)^2 + (y_1 - y_2)^2 + (z_1 - z_2)^2 \geq 0.
\]

From this formula we see that the norm of the difference of two vectors is a compact expression for the Euclidean length. A zero-length vector, such as a point, is the result of the fact that

\[
||x - x||^2 = (x - x) \cdot (x - x) = 0.
\]

**Integral definition of a scalar product:** Up to this point, following Euclid, we have only considered a vector to be a set of elements \( \{x_n\} \in \mathbb{R} \), index over \( n \in \mathbb{N} \), as defining a linear vector space with scalar product \( x \cdot y \), with the scalar product defining the norm or length of the vector \( ||x|| = \sqrt{x \cdot x} \). Given the scalar product, the norm naturally follows.

Now an obvious question presents itself: Can we extend our definition of vectors to differentiable functions (i.e., \( f(t) \) and \( g(t) \)) indexed over \( t \in \mathbb{R} \) with coefficients labeled by \( t \in \mathbb{R} \) rather than by \( n \in \mathbb{N} \)? Clearly, if the functions are analytic, there is no obvious reason that this should be a problem, since analytic functions may be represented by a convergent series that has Taylor coefficients and thus are integrable term by term.

Specifically, under certain conditions, the function \( f(t) \) may be thought of as a vector, defining a normed vector space. This intuitive and somewhat obvious idea is powerful. In this case the scalar product can be defined

![Figure 3.4: Vectors \( \mathbf{A}, \mathbf{B}, \mathbf{C} \) are used to define the scalar product \( \mathbf{A} \cdot \mathbf{B} \in \mathbb{R} \), vector wedge-product \( \mathbf{A} \wedge \mathbf{B} \in \mathbb{R} \), and triple wedge-product \( \mathbf{C} \cdot (\mathbf{A} \wedge \mathbf{B}) \). The vector wedge-product is the same as the vector cross-product except the output is a scalar rather than a vector. As shown, the scalar dot and vector wedge-products complement each other, since one is proportional to the \( \sin \theta \) of the angle \( \theta \) between them, and the other to the \( \cos \theta \). The scalar product computes the projection of one vector on the other (the length of the base of the triangle formed by the two vectors), while the vector wedge-product \( \mathbf{A} \wedge \mathbf{B} \) computes the area of the parallelogram (area = base \cdot \text{height} = 2\mathbf{A} \cdot \mathbf{B} ||\mathbf{L}|| \) formed by the two vectors. Thus \( ||\mathbf{A} \cdot \mathbf{B}||^2 + ||\mathbf{A} \wedge \mathbf{B}||^2 = ||\mathbf{A}||^2 ||\mathbf{B}||^2 \). The scalar triple product \( \mathbf{C} \cdot (\mathbf{A} \times \mathbf{B}) \) is the volume of the parallelepiped (i.e., prism) defined by the three vectors \( \mathbf{A}, \mathbf{B}, \) and \( \mathbf{C} \). When all the angles are \( 90^\circ \), the volume becomes a cuboid (see p. 94). 3.4 p 94

Scalar products play an important role in vector algebra and calculus (see Appendix A.3, p. 225).

In vector notation the scalar product is written as

\[
\mathbf{x} \cdot \mathbf{y} = \begin{bmatrix} x^T \\ y \\ z \end{bmatrix} \cdot \begin{bmatrix} \alpha \\ \beta \\ \gamma \end{bmatrix} = \alpha x + \beta y + \gamma z. \tag{3.5.0.2}
\]

If \( \mathbf{x}(s) \in \mathbb{C}^3 \) is a complex function of frequency \( s \), then the scalar product is a complex function of \( s \).
3.5. INTRODUCTION TO ANALYTIC GEOMETRY

in terms of the integral

\[ f(t) \cdot g(t) = \int_t f(t)g(t) dt = \|f(t)\| \|g(t)\| \cos \theta \]

summed over \( t \in \mathbb{R} \), rather than a sum over \( n \in \mathbb{N} \).

This definition of the vector scalar product allows for a significant but straightforward generalization of our vector space, which will turn out to be both useful and an important extension of the concept of a normed vector space. In this space we can define the derivative of a norm with respect to \( t \), which is not possible for the discrete case, indexed over \( n \). The distinction introduces the concept of analytic continuity in the index \( t \), which does not exist for the discrete index \( n \in \mathbb{N} \).

**Pythagorean theorem and the Schwarz inequality:** Regard Fig. 3.4, suppose we compute the difference between vector \( A \in \mathbb{R} \) and \( \alpha B \in \mathbb{R} \) as \( L = \|A - \alpha B\| \in \mathbb{R} \), where \( \alpha \in \mathbb{R} \) is a scalar that modifies the length of \( B \). We seek the value of \( \alpha \), which we denote as \( \alpha^* \), that minimizes the length of \( L \). From simple geometrical considerations, \( L(\alpha) \) will be minimum when the difference vector is perpendicular to \( B \), as shown in the figure by the dashed line from the tip of \( A \perp B \).

To show this algebraically, we write the expression for \( L(\alpha) \), take the derivative with respect to \( \alpha \), and set it to zero, which gives the formula for \( \alpha^* \). The argument does not change, but the algebra greatly simplifies if we normalize \( A \) and \( B \) to be unit vectors \( \hat{A} \) and \( \hat{B} \), that minimizes the length of \( \alpha \). We seek the value of \( \alpha \) in terms of the integral \( \alpha \in \mathbb{R} \), this example violates the common assumption that \( \alpha \in \mathbb{C} \).

Thus the length is shortest \( (L = L_\alpha) \), as shown in Fig. 3.4) when

\[
\frac{d}{d\alpha} L_\alpha^2 = -2a \cdot b + 2\alpha^* = 0.
\]

Solving for \( \alpha^* \in \mathbb{R} \), we find \( \alpha^* = a \cdot b \). Since \( L_\alpha > 0 \) \( (a \neq b) \), Eq. 3.5.0.4 becomes

\[
1 - 2|a \cdot b|^2 + |a \cdot b|^2 = 1 - |a \cdot b|^2 > 0.
\]

In conclusion, \( \cos \theta \equiv |a \cdot b| < 1 \). In terms of \( \hat{A} \) and \( \hat{B} \) this is \( |A \cdot B| < ||A|| ||B|| \cos \theta \), as shown adjacent to \( B \) in Fig. 3.4. Thus the scalar product between two vectors is their direction cosine. Furthermore, since this forms a right triangle, the Pythagorean theorem must hold. The triangle inequality says that the sum of the lengths of the two sides must be greater than the length of the hypotenuse. Note that \( \Theta \in \mathbb{R} \notin \mathbb{C} \). This derivation is an abbreviated version of a related discussion on page 99. Equality cannot be obtained because the Fourier space forms an open set, which gives rise to Gibbs ringing.

**Vector cross (\( \times \)) and wedge (\( \wedge \)) products of two vectors:** The vector product (cross-product) \( A \times B \) and the exterior product (wedge product) \( A \wedge B \) are the second and third types of vector products. As shown in Fig. 3.4,

\[
C = A \times B = (a_1 \hat{x} + a_2 \hat{y} + a_3 \hat{z}) \times (b_1 \hat{x} + b_2 \hat{y} + b_3 \hat{z}) = \begin{vmatrix} \hat{x} & \hat{y} & \hat{z} \\ a_1 & a_2 & a_3 \\ b_1 & b_2 & b_3 \end{vmatrix}
\]

is \( \perp \) to the plane defined by \( A \) and \( B \). The cross-product is strictly limited to two input vectors \( A \) and \( B \in \mathbb{R}^2 \) taken from three real dimensions (i.e., \( \mathbb{R}^3 \)).

The exterior (wedge) product generalizes the cross-product, since it may be defined in terms of any two vectors \( A, B \in \mathbb{C}^2 \) taken from \( n \) dimensions \( (\mathbb{C}^n) \) with output in \( \mathbb{C}^1 \). Thus the cross-product is composed of three wedge-products.

**Example:** If we define \( A = 3\hat{x} - 2\hat{y} + 0\hat{z} \) and \( B = 1\hat{x} + 1\hat{y} + 0\hat{z} \), then the cross-product is

\[
A \times B = \begin{vmatrix} \hat{x} & \hat{y} & \hat{z} \\ 3 & -2 & 0 \\ 1 & 1 & 0 \end{vmatrix} = (3\hat{j} + 2\hat{z}).
\]

Since \( a_1 \in \mathbb{C} \), this example violates the common assumption that \( A \in \mathbb{R}^3 \).
2. The wedge-product $A \wedge B$ takes two vectors and returns a scalar, which is the magnitude of a vector $\perp$ to the plane defined by the two input vectors (see Fig. 3.4). It is defined as

$$A \wedge B = \begin{vmatrix} a_1 & b_1 \\ a_2 & b_2 \end{vmatrix} = \begin{vmatrix} 3j & 1 \\ -2 & 1 \end{vmatrix} = (3j\mathbf{x} - 2\mathbf{y}) \wedge (\mathbf{x} + \mathbf{y}) = 3 \cdot 0 \mathbf{x} \wedge \mathbf{y} - 3j\mathbf{y} \wedge \mathbf{y} - 2 \mathbf{x} \wedge \mathbf{y} = (3j + 2)\mathbf{y} + (3j + 2)\mathbf{z} = 3j + 2.$$

This defines a compact and useful algebra (Hestenes, 2003).

From the example above we see that the absolute value of the wedge-product $|a \wedge b|$ is defined as $|a \times b|$, namely,

$$|(a_2\mathbf{y} + a_3\mathbf{z}) \wedge (b_2\mathbf{y} + b_3\mathbf{z})| = |a \times b|.$$

The wedge-product is especially useful because it is zero when the two vectors are colinear, – that is, $\mathbf{x} \wedge \mathbf{x} = 0$ and $\mathbf{x} \wedge \mathbf{y} = 1$, where $\mathbf{x}$ and $\mathbf{y}$ are unit vectors. Since

$$\vec{a} \cdot \vec{b} = ||\vec{a}|| ||\vec{b}|| \cos \theta \quad \text{and} \quad \vec{a} \wedge \vec{b} = ||\vec{a}|| ||\vec{b}|| \sin \theta,$$

it follows that

$$\vec{a} \cdot \vec{b} + j\vec{a} \wedge \vec{b} = ||\vec{a}|| ||\vec{b}|| e^{j\theta}$$

which may be viewed as the complex scalar product, with the right-hand side the polar form.

The main advantage of the wedge product is that it is valid in $n \geq 3$ dimensions since it is defined for any two vectors in any number of dimensions. Like the cross-product, the magnitude of the wedge-product is equal to the area of the trapezoid formed by the two vectors.

**Scalar triple product:** The triple of a third vector $C$ with the vector product $A \times B \in \mathbb{R}$ is

$$C \cdot (A \times B) = \begin{vmatrix} c_1 & c_2 & c_3 \\ a_1 & a_2 & a_3 \\ b_1 & b_2 & b_3 \end{vmatrix} \in \mathbb{R}^3,$$

which equals the volume of a parallelepiped.

### 3.5.1 Generalized vector product

As shown in Fig. 3.4, any two vectors $A, B \in \{\mathbf{x}, \mathbf{y}\}$ define a plane. There are two types of vector products:14

1. the scalar product

$$A \cdot B = ||A|| ||B|| \cos \theta \in \mathbb{R},$$

and

2. the vector wedge-product

$$A \wedge B = ||A|| ||B|| \sin \theta \in \mathbb{R},$$

each a real scalar.15

As shown in the figure, these two products form a right triangle and thus may be naturally merged, defining the complex analytic vector (wedge) product

$$A \wedge B = A \cdot B + jA \wedge B = ||A|| ||B|| e^{j\theta} \in \mathbb{C}.$$

Important examples, based on the Poynting theorem, come from Maxwell’s equations

$$\mathcal{P} = E \wedge H = E \cdot H + jE \wedge H \quad [\text{W/m}^2],$$

(Sommerfeld, 1952, p. 26), and the corresponding momentum equation (Johnson et al., 1994)

$$\mathcal{M} = D \wedge B = D \cdot B + jD \wedge B = \frac{1}{c_0^2} \mathcal{P} \quad [\text{J s/m}^4].$$

While the solar constant $\mathcal{P}$ is $1.3$ [kW/m²] is large, solar gravity, which follows from Maxwell’s equations, is $1/c_0^2 = 10 \times 10^{-12}$ smaller.

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14http://en.wikipedia.org/wiki/Bivector

15In some texts the wedge-product is called the vector exterior-product.
Impact of Analytic Geometry: The most obvious impact of analytic geometry was its detailed analysis of the conic sections using algebra rather than drawings with a compass and ruler. An important example is the composition of the line and circle, a venerable construction presumably going back to long before Euclid.

Once algebra was invented, the composition could be done using formulas. With this analysis came complex numbers. The first two mathematicians to appreciate this mixture of Euclid’s geometry and the new algebra were Fermat and Descartes. Soon Newton contributed to this effort by adding physics (e.g., calculations in acoustics, orbits of the planets, and the theory of gravity and light, significant concepts for 1687). (Stillwell, 2010, p. 115-117).

Given these new methods, many new solutions to problems emerged. The complex roots of polynomials continued to appear, without any obvious physical meaning. Newton called them imaginary. Complex numbers seem to have been viewed as more of an inconvenience than a problem. Newton’s solution to this dilemma was to simply ignore the “imaginary” cases (Stillwell, 2010, p. 115-19).

3.5.2 Development of Analytic Geometry

The first “algebra” (al-jabr) is credited to al-Khwarizmi (830 CE). Its invention advanced the theory of polynomial equations in one variable, Taylor series, and composition versus intersections of curves. The solution of the quadratic equation had been worked out thousands of years earlier, but with algebra a general solution could be defined. The Chinese had found the way to solve several equations in several unknowns—for example, finding the values of the intersections of two circles. With the invention of algebra by al-Khwarizmi, a powerful tool became available to solve more difficult problems.

In algebra there are two contrasting operations on functions: composition and elimination (e.g., intersection). Intersection sits between.

Composition:

Composition is the merging of functions by feeding one into the other. If the two functions are \( f \) and \( g \), then their composition is indicated by \( f \circ g \), meaning the function \( y = f(x) \) is substituted into the function \( z = g(y) \), giving \( z = g(f(x)) \).

Composition is not limited to linear equations, even though that is where it is most frequently applied. To compose two functions, we must substitute one equation into the other. That requires solving for that substitution variable, which is not always possible in the case of nonlinear equations. However, many tricks are available that may work around this restriction. For example, if one equation is in \( x^2 \) and the other in \( x^3 \) or \( \sqrt{x} \), it may be possible to multiply the first by \( x \) or square the second. The point is that one of the variables must be isolated so that when it is substituted into the other equation, the variable is removed from the mix.

Example: Let \( y = f(x) = x^2 - 2 \) and \( z = g(y) = y + 1 \). Then

\[
g \circ f = g(f(x)) = (x^2 - 2) + 1 = x^2 - 1.
\]  

(3.5.2.6)

In general, composition does not commute (i.e., \( f \circ g \neq g \circ f \)), as is easily demonstrated. Swapping the order of composition for our example gives

\[
f \circ g = f(g(y)) = z^2 - 2 = (y + 1)^2 - 2 = y^2 + 2y - 1.
\]  

(3.5.2.7)

Intersection:

Complementary to composition is intersection (i.e., decomposition). For example, the intersection of two lines is defined as the point where they meet. This is not to be confused with finding roots. A polynomial of degree \( N \) has \( N \) roots, but the points where two polynomials intersect has nothing to do with the roots of the polynomials. The intersection is a function (equation) of lower degree, implemented by Gaussian elimination.

A system of linear equations \( Ax = y \) has many interpretations, and one should not be biased by the notation. As engineers we are trained to view \( x \) as the input and \( y \) as the output, in which case then \( y = Ax \) seems natural, much like the functional relation \( y = f(x) \). But what does the linear relation \( x = Ay \) mean, when \( x \) is the input? The obvious answer is that \( y = A^{-1}x \).

But when working with systems of equations, there are many uses of equations, and we need to become more flexible in our interpretation. For example \( y = A^2x \) has a useful meaning, and in fact we saw this type of
relationship when working with Pell’s equation (p. 52) and the Fibonacci sequence (p. 53). As another example consider
\[
\begin{pmatrix} z_1 \\ z_2 \end{pmatrix} = \begin{pmatrix} a_{1x} & a_{1y} \\ a_{2x} & a_{2y} \end{pmatrix} \begin{pmatrix} x \\ y \end{pmatrix},
\]
which is reminiscent of a three-dimensional surface \( z = f(x, y) \). We shall find that such generalizations are much more than a curiosity.

### Intersection of two lines:
Unless they are parallel, two lines meet at a point. In terms of linear algebra, this may be written as two linear equations (on the left) along with the intersection point \([x_1, x_2]^T\) given by the inverse of the \(2 \times 2\) set of equations (on the right):
\[
\begin{pmatrix} a & b \\ c & d \end{pmatrix} \begin{pmatrix} x_1 \\ x_2 \end{pmatrix} = \begin{pmatrix} y_1 \\ y_2 \end{pmatrix} \quad \begin{pmatrix} x_1 \\ x_2 \end{pmatrix} = \frac{1}{\Delta} \begin{pmatrix} d & -b \\ -c & a \end{pmatrix} \begin{pmatrix} y_1 \\ y_2 \end{pmatrix}.
\]
(3.5.2.8)

By substituting the expression for the intersection point \([x_1, x_2]^T\) into the original equation, we see that it satisfies the equations. Thus the equation on the right is the solution to the equation on the left.

### Elimination:
Note the structure of the inverse: (1) The diagonal values \((a, d)\) are swapped, (2) the off-diagonal values \((b, c)\) are negated, and (3) the \(2 \times 2\) matrix is divided by the determinant \(\Delta = ad - bc\). If \(\Delta = 0\), there is no solution. When the determinant is zero \((\Delta = 0)\), the slopes of the two lines
\[
slope = \frac{b}{a} = \frac{d}{c}
\]
are equal; thus the lines are parallel. Only if the slopes differ can there be a unique solution.

### Exercise #39
Show that the equation on the right is the solution of the equation on the left.

**Solution:** By direct substitution (composition) of the right equation into the left equation, we have
\[
\begin{pmatrix} a & b \\ c & d \end{pmatrix} \begin{pmatrix} d & -b \\ -c & a \end{pmatrix} \begin{pmatrix} y_1 \\ y_2 \end{pmatrix} = \frac{1}{\Delta} \begin{pmatrix} ad - bc & -ab + ab \\ cd - cd & -eb + ad \end{pmatrix} \begin{pmatrix} y_1 \\ y_2 \end{pmatrix} = \frac{1}{\Delta} \begin{pmatrix} \Delta & 0 \\ 0 & \Delta \end{pmatrix} \cdot \begin{pmatrix} y_1 \\ y_2 \end{pmatrix}.
\]
which gives the identity matrix. \(\blacksquare\)

Algebra will give the solution when geometry cannot. When the two curves fail to intersect on the real plane, the solution still exists, but it is complex-valued. In such cases, geometry, which considers only the real solutions, fails. For example, when the coefficients \([a, b, c, d]\) are complex, the solution exists but the determinant can be complex. Thus algebra is much more general than geometry. Geometry fails when the solution has a complex intersection.

### 3.5.3 Applications of scalar products
Another important example of algebraic expressions in mathematics is Hilbert’s generalization of the Pythagorean theorem (Eq. 1.1.1.1), known as the Schwarz inequality and shown in Fig. 3.5. What is special about this generalization is that it proves that when the vertex is \(90^\circ\), the Euclidean length of the leg is minimum.

Vectors may be generalized to have \(\infty\) dimensions: \(\vec{U}, \vec{V} = [v_1, v_2, \ldots, v_\infty]\). The Euclidean inner product (i.e., scalar product) between two such vectors generalizes the finite-dimensional case
\[
\vec{U} \cdot \vec{V} = \sum_{k=1}^{\infty} u_k v_k = \|\vec{U}\| \|\vec{V}\| \cos \theta
\]
where \(\theta\) is the multivalued angle between the two normalized (unit) vectors
\[
\theta = \cos^{-1} \left( \frac{\vec{U}}{\|\vec{U}\|} \cdot \frac{\vec{V}}{\|\vec{V}\|} \right).
\]

---

16When we write the equation \(Ax = y\) in matrix format, the two equations are \(ax_1 + bx_2 = y_1\) and \(dx_1 + ex_2 = y_2\) with unknowns \((x_1, x_2)\), whereas in the original equations \(ay + bx = c\) and \(dy + ex = f\), the unknowns are \(y\) and \(x\). Thus in matrix format, the names are changed. The first time you see this scrambling of variables, it can be confusing.
As with the finite case norm $||\vec{U}|| = \sqrt{\vec{U} \cdot \vec{U}} = \sqrt{\sum u_k^2}$ the scalar product of the vector with itself, defines the length of the infinite component vector. There is an issue of convergence when the norm of the vectors is zero.

It is a somewhat arbitrary requirement that $a, b, c \in \mathbb{R}$ for (Eq. 1.1.1.1). This seems natural enough, since the sides are lengths. But, what if these lengths are taken from high dimensional complex vectors, as for the lossy vector wave equation or the lengths of vectors in the $\mathcal{F}^T \in \mathbb{C}^n$? Then the equation generalizes to $K \to \infty$ dimensions

$$c \cdot c = ||c||^2 = \sum_{k=1}^{\infty} |c_k|^2.$$ 

As before, $||c|| = \sqrt{||c||^2}$ is the norm of vector $c$, akin to a length, which must be finite (converge). This is simply the important case of complex analytic functions, which also must converge, and is the “intersection” between the $\mathcal{F}^T$ and $L^T$.

**Schwarz inequality:** The Schwarz inequality says that the magnitude of the inner product of two vectors is less than or equal to the product of their lengths: \(^{17}\)

$$|\vec{U} \cdot \vec{V}| \leq ||\vec{U}|| ||\vec{V}||.$$

This may be simplified by normalizing the vectors to have unit length ($\vec{U} = \vec{U}/||\vec{U}||$, $\vec{V} = \vec{V}/||\vec{V}||$), in which case $-1 < \vec{U} \cdot \vec{V} \leq 1$. Another simplification is to define the scalar product in terms of the direction cosine

$$\cos \theta = |\vec{U} \cdot \vec{V}| \leq 1.$$

A proof of the Schwarz inequality is as follows: From these definitions we may define the minimum difference between the two vectors as the perpendicular from the end of the first to the intersection with the second. As shown in Fig. 3.5, $\vec{U} \perp \vec{V}$ may be found by minimizing the length of the vector difference:

$$\min_{\alpha} ||\vec{V} - \alpha \vec{U}||^2 = ||\vec{V}||^2 + 2\alpha \vec{V} \cdot \vec{U} + \alpha^2 ||\vec{U}||^2 > 0$$

$$0 = \partial_{\alpha} (\vec{V} - \alpha \vec{U}) \cdot (\vec{V} - \alpha \vec{U})$$

$$= \vec{V} \cdot \vec{U} - \alpha^* ||\vec{U}||^2$$

$$\therefore \quad \alpha^* = \vec{V} \cdot \vec{U}/||\vec{U}||^2.$$ 

The Schwarz inequality follows:

$$I_{\min} = ||\vec{V} - \alpha^* \vec{U}||^2 = ||\vec{V}||^2 - \frac{|\vec{U} \cdot \vec{V}|^2}{||\vec{U}||^2} > 0$$

$$0 \leq |\vec{U} \cdot \vec{V}| \leq ||\vec{U}|| ||\vec{V}||.$$

An important example of such a vector space includes the definition of the Fourier transform, where we may set

$$U(\omega) = e^{-j\omega_0 t} \quad V(\omega) = e^{j\omega t} \quad U \cdot V = \int e^{j\omega t} e^{-j\omega_0 t} \frac{d\omega}{2\pi} = \delta(\omega - \omega_0).$$

\(^{17}\)We provide a simplified derivation on page 95.
It seems that the Fourier transform is a result that follows from a minimization, unlike the Laplace transform that follows from a causal system. This explains the important differences between the two in terms of their properties (unlike the LT, the FT is not complex analytic). Recall that

$$U \cdot V + j U \wedge V = ||U|| ||V|| e^{j\theta}.$$ 

We further explore this topic on page 115.

### 3.5.4 Gaussian Elimination

The method for finding the intersection of equations is based on the recursive elimination of all the variables but one. This method, known as Gaussian elimination (Appendix 5, p. 228), works across a broad range of cases but may be defined as a systematic algorithm when the equations are linear in the variables (Strang et al., 1993). Rarely do we even attempt to solve problems in several variables of degree greater than 1. But Gaussian elimination may still work in such cases (Stillwell, 2010, p. 90).

In Appendix A.3.5 (p. 230) we derive the inverse of a $2 \times 2$ linear system of equations. Even for a $2 \times 2$ case, the general solution requires a great deal of algebra. Working out a numeric example of Gaussian elimination is more instructive. For example, suppose we wish to find the intersection of the two equations

$$x - y = 3$$
$$2x + y = 2.$$ 

This $2 \times 2$ system of equations is so simple that you may immediately visualize the solution: By adding the two equations, $y$ is eliminated, leaving $3x = 5$. But doing it this way takes advantage of the specific example, and we need a method for larger systems of equations. We need a generalized (algorithmic) approach. This general approach is called Gaussian elimination.

We start by writing the equations in matrix form (note this is not in the form $Ax = y$):

$$\begin{bmatrix} 1 & -1 \\ 2 & 1 \end{bmatrix} \begin{bmatrix} x \\ y \end{bmatrix} = \begin{bmatrix} 3 \\ 2 \end{bmatrix}. \quad (3.5.4.9)$$

Next, we eliminate the lower left term ($2x$) using a scaled version of the upper left term ($x$). Specifically, we multiply the first equation by $-2$ and add it to the second equation, replacing the second equation with the result. This gives

$$\begin{bmatrix} 1 & -1 \\ 0 & 3 \end{bmatrix} \begin{bmatrix} x \\ y \end{bmatrix} = \begin{bmatrix} 3 \\ 2 - 3 \cdot 2 \end{bmatrix} = \begin{bmatrix} 3 \\ -4 \end{bmatrix}. \quad (3.5.4.10)$$

Note that the top equation did not change. Once the matrix is “upper triangular” (zero below the diagonal), we have the solution. If we start from the bottom equation, $y = -4/3$. Then the upper equation gives $x - (-4/3) = 3$ or $x = 3 - 4/3 = 5/3$.

In principle, Gaussian elimination is easy, but if you make a calculation mistake along the way, it is very difficult to find your error. The method requires a lot of mental labor and you have a high probability of making a mistake. Thus you do not want to apply this method every time. For example, suppose the elements are complex numbers or polynomials in some other variable such as frequency. Once the coefficients become more complicated, the seemingly trivial problem becomes corrosive. There is a much better way that is easily verified; it puts all the numerics at the end in a single step.

The above operations may be automated by finding a carefully chosen upper-diagonalized matrix $G$. For example, we can define the Gaussian matrix that zeros the element 2 in the matrix in Eq. 3.5.4.9. More generally, we let

$$G = \begin{bmatrix} 1 & 0 \\ a & 1 \end{bmatrix} \quad (3.5.4.11)$$

Multiplying Eq. 3.5.4.9 by $G$, we find

$$\begin{bmatrix} 1 & 0 \\ a & 1 \end{bmatrix} \begin{bmatrix} 1 & -1 \\ 2 & 1 \end{bmatrix} \begin{bmatrix} x \\ y \end{bmatrix} = \begin{bmatrix} 1 & -1 \\ a + 2 & 1 - a \end{bmatrix} \begin{bmatrix} x \\ y \end{bmatrix} = \begin{bmatrix} 3 \\ 3a + 2 \end{bmatrix}. \quad (3.5.4.12)$$

Thus we obtain Eq. 3.5.4.10 if we let $a = -2$ (we choose $a$ to force the lower left to be zero). At this point we can either back-substitute and obtain the solution, as we did above, or find a matrix $L$ that finishes the job by removing elements above the diagonal. The key is that the determinant of this matrix is 1.
Exercise #40
Using $G$ and $A$ from the discussion above, show that $\det(G) = \det(GA) = 3$.

**Solution:** A common convention is to denote $\det(A) = |A|$. The two sides of the identity are

$$|G| = \det \begin{bmatrix} 1 & -1 \\ 2 & 1 \end{bmatrix} = 1 + 2 = 3, \quad |GA| = \det \begin{bmatrix} 1 & -1 \\ 0 & 3 \end{bmatrix} = 3,$$

and $|G| = 1$. Thus $|GA| = |G||A| = 3$. ■

Matrix inverse: In Appendix A.3.5 (p. 230), finding the inverse of a general $2 \times 2$ matrix takes three steps: (1) swap the diagonal elements, (2) reverse the signs of the off-diagonal elements, and (3) divide by the determinant $\Delta = ab - cd$. Specifically,

$$\begin{bmatrix} a & b \\ c & d \end{bmatrix}^{-1} = \frac{1}{\Delta} \begin{bmatrix} d & -b \\ -c & a \end{bmatrix}. \quad (3.5.4.13)$$

There are very few things that you must memorize, but the inverse of a $2 \times 2$ matrix is one of them. It needs to be in your mental toolkit, like completing the squares (see p. 64).

While it is difficult to compute the inverse matrix from scratch (see Appendix 5), it takes only a few seconds (four dot products) to verify it (steps 1 and 2):

$$\begin{bmatrix} a & b \\ c & d \end{bmatrix} \begin{bmatrix} d & -b \\ -c & a \end{bmatrix} = \begin{bmatrix} ad - bc & -ab + ad \\ cd - cd & -bc + ad \end{bmatrix} = \begin{bmatrix} \Delta & 0 \\ 0 & \Delta \end{bmatrix}. \quad (3.5.4.14)$$

Thus dividing by the determinant gives the $2 \times 2$ identity matrix. A good strategy (don’t trust your memory) is to write down the inverse as best you recall and then verify.

Using the $2 \times 2$ matrix inverse on our example (Eq. 3.5.4.9), we find

$$\begin{bmatrix} x \\ y \end{bmatrix} = \frac{1}{1 + 2} \begin{bmatrix} 1 & -2 & 1 \\ 1 & -1 & 4 \end{bmatrix} \begin{bmatrix} 3 \\ 2 \\ 9 \end{bmatrix} = \frac{1}{3} \begin{bmatrix} 5 \\ -6 + 2 \end{bmatrix} = \begin{bmatrix} 5/3 \\ -4/3 \end{bmatrix}. \quad (3.5.4.15)$$

If you use this method, you will rarely (never) make a mistake and the solution is easily verified. Either you can check the numbers in the inverse, as was done in Eq. 3.5.4.14, or you can substitute the solution back into the original equation.

Exercise #41
Move here from Appendix Linear Algebra. Make sure it blended

Find the solution to the following $3 \times 3$ matrix equation $Ax = b$ by Gaussian elimination. Show your intermediate steps. You can check your work at each step using Matlab.

$$\begin{bmatrix} 1 & 1 & -1 \\ 3 & 1 & 1 \\ -1 & -4 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \\ x_3 \end{bmatrix} = \begin{bmatrix} 1 \\ 9 \\ 8 \end{bmatrix}.$$

1. Show (i.e., verify) that the first GE matrix $G_1$, which zeros out all entries in the first column, is

$$G_1 = \begin{bmatrix} 1 & 0 & 0 \\ -3 & 1 & 0 \\ -1 & 0 & 1 \end{bmatrix}.$$  

Identify the elementary row operations that this matrix performs.

**Solution:** We operate with the GE matrix on $A$:

$$G_1[A|b] = \begin{bmatrix} 1 & 0 & 0 \\ -3 & 1 & 0 \\ -1 & 0 & 1 \end{bmatrix} \begin{bmatrix} 1 & 1 & -1 & 1 \\ 1 & 1 & 1 & 9 \\ 3 & 1 & 1 & 8 \end{bmatrix} = \begin{bmatrix} 1 & 1 & -1 & 1 \\ 0 & -2 & 4 & 6 \\ 0 & -2 & 5 & 7 \end{bmatrix}.$$  

The second row of $G_1$ scales the first row by -3 and adds it to the second row

$$(2) \leftarrow -3(1) + (2).$$

The third row of $G1$ scales the first row by -1 and adds it to the third row: $(3) \leftarrow -(1) + (3)$. ■
2. Find a second GE matrix, \( G_2 \), to put \( G_1 A \) in upper triangular form. Identify the elementary row operations that this matrix performs.

Solution:

\[
G_2 = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & -1 & 1 \end{bmatrix},
\]

or \((2) \leftarrow -(2) + (3)\). Thus we have

\[
G_2G_1[A|b] = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & -1 & 1 \end{bmatrix} \begin{bmatrix} 1 & 0 & 0 \\ -3 & 1 & 0 \\ -1 & 0 & 1 \end{bmatrix} \begin{bmatrix} 1 & 1 & -1 \\ 3 & 1 & 1 \\ 1 & -1 & 4 \end{bmatrix} \begin{bmatrix} 1 & 1 & -1 \\ 0 & 2 & 4 \\ 0 & 0 & 1 \end{bmatrix} = \begin{bmatrix} 1 & 1 & -1 \\ 0 & 2 & 4 \\ 0 & 0 & 1 \end{bmatrix}.
\]

3. Find a third GE matrix, \( G_3 \), that scales each row so that its leading term is 1. Identify the elementary row operations that this matrix performs.

Solution:

\[
G_3 = \begin{bmatrix} 1 & 0 & 0 \\ 0 & -1/2 & 0 \\ 0 & 0 & 1 \end{bmatrix},
\]

which scales the second row by \(-1/2\). Thus we have

\[
G_3G_2G_1[A|b] = \begin{bmatrix} 1 & 0 & 0 \\ 0 & -1/2 & 0 \\ 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} 1 & 1 & -1 \\ 0 & 2 & 4 \\ 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} 1 & 1 & -1 \\ 0 & 2 & 4 \\ 0 & 0 & 1 \end{bmatrix} = \begin{bmatrix} 1 & 1 & -1 \\ 0 & 1 & -2 \\ 0 & 0 & 1 \end{bmatrix}.
\]

4. Finally, find the last GE matrix, \( G_4 \), that subtracts a scaled version of row 3 from row 2, and scaled versions of rows 2 and 3 from row 1, such that you are left with the identity matrix \((G_4G_3G_2G_1A = I)\).

Solution:

\[
G_4 = \begin{bmatrix} 1 & -1 & -1 \\ 0 & 1 & 2 \\ 0 & 0 & 1 \end{bmatrix},
\]

Thus we have

\[
G_4G_3G_2G_1[A|b] = \begin{bmatrix} 1 & -1 & -1 \\ 0 & 1 & 2 \\ 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} 1 & 1 & -1 \\ 0 & 2 & 4 \\ 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} 1 & 1 & -1 \\ 0 & 2 & 4 \\ 0 & 0 & 1 \end{bmatrix} = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix}.
\]

5. Solve for \([x_1, x_2, x_3]^T\) using the augmented matrix format \(G_4G_3G_2G_1[A|b]\), where \([A|b]\) is the augmented matrix. Note that if you’ve performed the preceding steps correctly, \(x = G_4G_3G_2G_1b\).

Solution: From the preceding problems, we see that \([x_1, x_2, x_3]^T = [3, -1, 1]^T\).

**Augmented matrix:** There is one minor notational improvement. Rather than writing the matrix equation as Eq. 3.5.4.9 \((Ax = y)\), we place the \(y\) vector next to the elements of \(A\) to remove the equal sign, which is cumbersome. In this case we write \(GA_{aug}\):

\[
GA_{aug} = \begin{bmatrix} 1 & 0 & 3 \\ -2 & 1 & 3 \\ -1 & 2 & 3 \\ 1 & 0 & 3 \\ 0 & 3 & -4 \end{bmatrix}.
\]
3.6 Problems AE-2

Topics of this homework:
Linear systems of equations, Gaussian elimination, matrix permutations, overspecified systems of equations, analytic geometry, Ohm’s law, two-port networks
Deliverables: Answers to problems

Nonlinear (quadratic) to linear equations
In the following problems we deal with algebraic equations in more than one variable that are not linear equations. For example, the circle $x^2 + y^2 = 1$ may be solved for $y(x) = \pm \sqrt{1 - x^2}$. If we let $z = x + yj = x + j\sqrt{1 - x^2} = e^{j\theta}$, we obtain the equation for half a circle ($y > 0$). The entire circle is described by the magnitude of $z$ as $|z|^2 = (x + yj)(x - yj) = 1$.

Problem # 1: Give the curve defined by the equation:

$$x^2 + xy + y^2 = 1$$

- 1.1: Find the function $y(x)$.
- 1.2: Using Matlab/Octave, plot $y(x)$ and describe the graph.
- 1.3: What is the name of this curve?
- 1.4: Find the solution (in $x$, $p$, and $q$) to these equations:

$$x + y = p$$

$$xy = q.$$ 

- 1.5: Find an equation that is linear in $y$ starting from equations that are quadratic (second-degree) in the two unknowns $x$ and $y$:

$$x^2 + xy + y^2 = 1 \quad \text{(AE-2.1)}$$

$$4x^2 + 3xy + 2y^2 = 3. \quad \text{(AE-2.2)}$$

- 1.6: Compose the following two quadratic equations and describe the results.

$$x^2 + xy + y^2 = 1$$

$$2x^2 + xy = 1$$

Intersection and analytic geometry
To derive Euclid’s formula, it was necessary to intersect a circle and a secant line. Consider the unit circle of radius 1, centered at $(x, y) = (0, 0)$

$$x^2 + y^2 = 1$$

and the secant line through $(-1, 0)$

$$y = t(x + 1)$$
CHAPTER 3. ALGEBRAIC EQUATIONS: STREAM 2

Euclidean Proof:
1) $2\phi + \eta = \pi$
2) $\eta + \Theta = \pi$
3) $\therefore \phi = \Theta/2$

Diophantus’s Proof:
1) $c^2 = a^2 + b^2$
2) $b(a) = t(a + c)$
3) $\zeta(t) \equiv a + jb = \frac{1 - t^2 + it}{1 + t^2}$
4) $\zeta = |c|e^{i\theta} = |c|\frac{1 + it}{1 + t^2} = |c|(\cos(\theta) + i\sin(\theta))$

Pythagorean triplets:
1) $t = p/q \in \mathbb{Q}$
2) $a = p^2 - q^2$
3) $b = 2pq$
4) $c = p^2 + q^2$

Figure 3.6: Derivation of Euclid's formula for the Pythagorean triplets $[a, b, c]$, based on a composition of a line, having a rational slope $t = p/q \in \mathbb{F}$, and a circle $c^2 = a^2 + b^2$, $[a, b, c] \in \mathbb{N}$. This analysis is attributed to Diophantus (Di·o·phant′-us) (250 CE), and today such equations are called Diophantine (Di·o·phant′-ine) equations. PTs have applications in architecture and scheduling, and many other practical problems. Most interesting is their relation to Rydberg’s formula for the eigenstates of the Hydrogen atom.

having slope $t$ and intercept $x = -1$. If the slope $0 < t < 1$, the line intersects the circle at a second point $(a, b)$ in the positive $x, y$ quadrant. The goal is to find $a, b \in \mathbb{N}$ and then show that $c^2 = a^2 + b^2$. Since the construction gives a right triangle with short sides $a, b \in \mathbb{N}$, then it follows that $c \in \mathbb{N}$.

**Problem # 2: Derive Euclid’s formula**

- 2.1: Draw the circle and the line, given a positive slope $0 < t < 1$.

**Problem # 3:** Substitute $y = t(x + 1)$ (the line equation) into the equation for the circle, and solve for $x(t)$.

*Hint: Because the line intersects the circle at two points, you will get two solutions for $x$. One of these solutions is the trivial solution $x = -1$.*

- 3.1: Substitute the $x(t)$ you found back into the line equation, and solve for $y(t)$.

- 3.2: Let $t = q/p$ be a rational number, where $p$ and $q$ are integers. Find $x(p, q)$ and $y(p, q)$.

- 3.3: Substitute $x(p, q)$ and $y(p, q)$ into the equation for the circle, and show how Euclid’s formula for the Pythagorean triples is generated.

For full points you must show that you understand the argument. Explain the mean of the comment “magic happens” when $t^4$ cancels.

**Gaussian elimination**

**Problem # 4: Gaussian elimination**

- 4.1: Find the inverse of

$$A = \begin{bmatrix} 1 & 2 \\ 4 & 3 \end{bmatrix}.$$
3.6. PROBLEMS AE-2

– 4.2: Verify that \( A^{-1}A = AA^{-1} = \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix} \).

**Problem #5**: Find the solution to the following \(3 \times 3\) matrix equation \(Ax = b\) by Gaussian elimination. Show your intermediate steps. You can check your work at each step using Octave/Matlab.

\[
\begin{bmatrix} 1 & 1 & -1 \\ 3 & 1 & 1 \\ 1 & -1 & 4 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \\ x_3 \end{bmatrix} = \begin{bmatrix} 1 \\ 9 \\ 8 \end{bmatrix}.
\]

– 5.1 Show (i.e., verify) that the first Gaussian elimination (GE) matrix \(G_1\) that zeros out all entries in the first column is given by

\[
G_1 = \begin{bmatrix} 1 & 0 & 0 \\ -3 & 1 & 0 \\ -1 & 0 & 1 \end{bmatrix}.
\]

Identify the elementary row operations that this matrix performs.

– 5.2 Find a second GE matrix, \(G_2\), to put \(G_1A\) in upper triangular form. Identify the elementary row operations that this matrix performs.

– 5.3 Find a third GE matrix \(G_3\) that scales each row so that its leading term is 1. Identify the elementary row operations that this matrix performs.

– 5.4: Find the last GE matrix, \(G_4\), that subtracts a scaled version of row 3 from row 2, and scaled versions of rows 2 and 3 from row 1, such that you are left with the identity matrix \((G_4G_3G_2G_1A = I)\).

– 5.5: Solve for \(\{x_1, x_2, x_3\}^T\) using the augmented matrix format \(G_4G_3G_2G_1\{A|b\}\) (where \(\{A|b\}\) is the augmented matrix). Note that if you’ve performed the preceding steps correctly, \(x = G_4G_3G_2G_1b\).

– 5.6: Find the pivot matrix \(G\) that rescales the second row of the augmented matrix \(A|b\) by \(1/3\).

**Two linear equations**

**Problem #6** In this problem we transition from a general pair of equations

\[
f(x, y) = 0 \quad \quad \quad \quad g(x, y) = 0
\]

to the important case of two linear equations

\[
y = ax + b 
\quad \quad \quad \quad y = \alpha x + \beta.
\]

Note that to help keep track of the variables, roman coefficients \((a, b)\) are used for the first equation and Greek \((\alpha, \beta)\) for the second.

– 6.1: What does it mean, graphically, if these two linear equations have a unique solution, a nonunique solution, or no solution?
– 6.2: Assuming the two equations have a unique solution, find the solution for $x$ and $y$.

– 6.3: When will this solution fail to exist (for what conditions on $a$, $b$, $\alpha$, and $\beta$)?

– 6.4: Write the equations as a $2 \times 2$ matrix equation of the form $A\vec{x} = \vec{b}$, where $\vec{x} = \{x, y\}^T$.

– 6.5: Finding the inverse of the $2 \times 2$ matrix, and solve the matrix equation for $x$ and $y$.

– 6.6: Discuss the properties of the determinant of the matrix ($\Delta$) in terms of the slopes of the two equations ($a$ and $\alpha$).

**Problem # 7: The application of linear functional relationships between two variables**

We use $2 \times 2$ matrices to describe two-port networks, as discussed in Sec. 3.7 (p. 110). Transmission lines are a great example. Both voltage and current must be tracked as they travel along the line. Figure 3.10 (p. 114) shows an example segment of a transmission line.

Suppose you are given the following pair of linear relationships between the input (source) variables $V_1$ and $I_1$ and the output (load) variables $V_2$ and $I_2$ of the transmission line:

\[
\begin{aligned}
V_1 & = f(1) \ \text{and} \ I_1 = 1 \\
V_2 & = I_2 = 2 \ (i.e., V_2/I_2 = 1/2 \ \Omega) .
\end{aligned}
\]

– 7.1: Let the output (the load) be $V_2 = 1$ and $I_2 = 2$ (i.e., $V_2/I_2 = 1/2 \ \Omega$). Find the input voltage and current $V_1$ and $I_1$.

– 7.2: Let the input (source) be $V_1 = 1$ and $I_1 = 2$. Find the output voltage and current $V_2$ and $I_2$.

**Integer equations: applications and solutions**

Any equation for which we seek only integer solutions is called a *Diophantine* equation.

**Problem # 8: A practical example of using a Diophantine equation:**

“A merchant had a 40-pound weight that broke into 4 pieces. When the pieces were weighed, it was found that each piece was a whole number of pounds and that the four pieces could be used to weigh every integral weight between 1 and 40 pounds. What were the weights of the pieces?” *-Bachet de Bèziriac (1623)*

Here, weighing is performed using a balance scale that has two pans, with weights on either pan. Thus, given weights of 1 and 3 pounds, one can weigh a 2-pound weight by putting the 1-pound weight in the same pan with the 2-pound weight, and the 3-pound weight in the other pan. Then the scale will be balanced. A solution to the four weights for Bachet’s problem is $1 + 3 + 9 + 27 = 40$ pounds.

\[
\begin{array}{c}
\begin{array}{c}
1 \\
\text{Left side ‘-’}
\end{array} \\
\begin{array}{c}
?+2 \\
\text{Right side ‘+’}
\end{array}
\end{array}
\]

**Solution:** $2 = -1 + 3$

---

18Taken from: Joseph Rotman, (Rotman, 1996, p. 50)
3.6 PROBLEMS AE-2

Show how the combination of 1-, 3-, 9-, and 27-pound weights can be used to weigh 1, 2, 3, \ldots, 8, 28, and 40 pounds of milk (or something else, such as flour). Assuming that the milk is in the left pan, provide the position of the weights using a negative sign $-$ to indicate the left pan and a positive sign $+$ to indicate the right pan. For example, if the left pan has 1 pound of milk, then 1 pound of milk in the right pan, $+1$, will balance the scales.

Hint: It is helpful to write the answer in matrix form. Set the vector of values to be weighed equal to a matrix indicating the pan assignments, multiplied by a vector of the weights $[1, 3, 9, 27]^T$. The pan assignments matrix should contain only the values $-1$ (left pan), $+1$ (right pan), and 0 (leave out). You can indicate these using $-$, $+$, and blanks.

### Ohm’s Law

In general, impedance is defined as the ratio of a force to a flow. For electrical circuits, the voltage is the force and the current is the flow. Ohm’s law states that the voltage across and the current through a circuit element are related by the impedance of that element (which may be a function of frequency). For resistors, the voltage over the current is called the resistance and is a constant (e.g., the simplest case is $V/I = R$). For inductors and capacitors, the voltage over the current is a frequency-dependent impedance (e.g., $V/I = Z(s)$, where $s$ is the complex frequency $s \in \mathbb{C}$).

As shown in Table 3.2 (p. 114), the impedance concept also holds in mechanics and acoustics. In mechanics, the force is equal to the mechanical force on an element (e.g., a mass, dashpot, or spring) and the flow is the velocity. In acoustics, the force is pressure and the flow is the volume velocity or particle velocity of air molecules.

**Problem # 9:** The resistance of an incandescent (filament) lightbulb, measured cold, is about 100 ohms. As the bulb lights up, the resistance of the metal filament increases.

Ohm’s law says that the current

$$\frac{V}{I} = R(T),$$

where $T$ is the temperature. In the United States, the voltage is 120 volts (RMS) at 60 [Hz]. Find the current when the light is first switched on.

**Problem # 10:** The power in watts is the product of the force and the flow. What is the power of the lightbulb of Problem 9?

**Problem # 11:** State the impedance $Z(s)$ of each of the following circuit elements: a resistor with resistance $R$, an inductor with inductance $L$, and a capacitor with capacitance $C$.

For the capacitor,

**Problem # 12:** Consider what happens at the triple point of water. As water freezes or thaws, the temperature remains constant at 0 (C°). Once all the water is frozen and more heat is removed, the temperature drops below 0°. As heat is added, water thaws but the temperature remains at 0°. Once all the ice has melted, what is the temperature as more heat is added?

Model the triple point using a Zener diode, a resistor, and a capacitor. A Zener diode holds the voltage constant independent of current. For the case of water’s triple point, the voltage represents the temperature of water at the triple point, clamped at 0 [C°]. The current represents the heat flux. The latent heat of water at the triple point is 32 Cal/gm. Thus as the temperature rises from below freezing, the water is clamped at 0° once the triple point is reached. At that point, adding more heat flux has no effect on the temperature until all the ice melts. Once the ice has melted, the temperature again begins to rise until it hits the boiling point, where it again stays at 100° until all the water has evaporated.

### Vector algebra in $\mathbb{R}^3$.

Definitions of the scalar (also called a dot product) $A \cdot B$, cross $A \times B$ and triple product $A \cdot (B \times C)$, may be found in Appendix A (p. 221), where $A, B, C$ in $\mathbb{R}^3 \subset \mathbb{C}^3$, as shown in Fig. 3.4 page $\text{fig:VecDev}$. A fourth “double-cross” ($\mathcal{G}$) vector product is:$^{10}$

$$A \times (B \times C) = \alpha_aB - \beta_aC.$$

$^{10}$Greenberg p. 694, Eq. 8.
where \( \alpha_0 = \mathbf{A} \cdot \mathbf{C} \) and \( \beta_0 = \mathbf{A} \cdot \mathbf{B} \) (Note: \( \mathbf{A} \times (\mathbf{B} \times \mathbf{C}) \neq (\mathbf{A} \times \mathbf{B}) \times \mathbf{C} \)).

**Problem #13: Scalar product \( \mathbf{A} \cdot \mathbf{B} \)**

- 13.1: If \( \mathbf{A} = a_x \mathbf{\hat{x}} + a_y \mathbf{\hat{y}} + a_z \mathbf{\hat{z}} \) and \( \mathbf{B} = b_x \mathbf{\hat{x}} + b_y \mathbf{\hat{y}} + b_z \mathbf{\hat{z}} \), write out the definition of \( \mathbf{A} \cdot \mathbf{B} \).

- 13.2: The dot product is often defined as \( ||\mathbf{A}|| ||\mathbf{B}|| \cos(\theta) \), where \( ||\mathbf{A}|| = \sqrt{\mathbf{A} \cdot \mathbf{A}} \) and \( \theta \) is the angle between \( \mathbf{A} \) and \( \mathbf{B} \). If \( ||\mathbf{A}|| = 1 \), describe how the dot product relates to the vector \( \mathbf{B} \).

**Problem #14: Vector (cross) product \( \mathbf{A} \times \mathbf{B} \)**

- 14.1: If \( \mathbf{A} = a_x \mathbf{\hat{x}} + a_y \mathbf{\hat{y}} + a_z \mathbf{\hat{z}} \) and \( \mathbf{B} = b_x \mathbf{\hat{x}} + b_y \mathbf{\hat{y}} + b_z \mathbf{\hat{z}} \), write out the definition of \( \mathbf{A} \times \mathbf{B} \).

- 14.2: Show that the cross product is equal to the area of the parallelogram formed by \( \mathbf{A} \) and \( \mathbf{B} \), namely \( ||\mathbf{A}|| ||\mathbf{B}|| \sin(\theta) \), where \( ||\mathbf{A}|| = \sqrt{\mathbf{A} \cdot \mathbf{A}} \) and \( \theta \) is the angle between \( \mathbf{A} \) and \( \mathbf{B} \).

**Problem #15: Triple product \( \mathbf{A} \cdot (\mathbf{B} \times \mathbf{C}) \)**

Let \( \mathbf{A} = [a_1, a_2, a_3]^T \), \( \mathbf{B} = [b_1, b_2, b_3]^T \), \( \mathbf{C} = [c_1, c_2, c_3]^T \) be three vectors in \( \mathbb{R}^3 \).

- 15.1: Starting from the definition of the dot and cross product, explain using a diagram and/or words, how one shows that: \( \mathbf{A} \cdot (\mathbf{B} \times \mathbf{C}) = \begin{vmatrix} a_1 & a_2 & a_3 \\ b_1 & b_2 & b_3 \\ c_1 & c_2 & c_3 \end{vmatrix} \).

- 15.2: Describe why \( |\mathbf{A} \cdot (\mathbf{B} \times \mathbf{C})| \) is the volume of parallelepiped generated by \( \mathbf{A}, \mathbf{B} \) and \( \mathbf{C} \).

- 15.3: Explain why three vectors \( \mathbf{A}, \mathbf{B}, \mathbf{C} \) are in one plane if and only if the triple product \( \mathbf{A} \cdot (\mathbf{B} \times \mathbf{C}) = 0 \).

**Problem #16: Given two vectors \( \mathbf{A}, \mathbf{B} \) in the \( \mathbf{\hat{x}}, \mathbf{\hat{y}} \) plane shown in Fig. 2.3 (same as 3.4 on page 94), with \( \mathbf{B} = \mathbf{\hat{y}} \) (i.e., \( ||\mathbf{B}|| = 1 \)).**

- 16.1: Show that \( \mathbf{A} \) may be split into two orthogonal parts, one in the direction of \( \mathbf{B} \) and the other perpendicular (\( \perp \)) to \( \mathbf{B} \). Hint: Express the vector products of \( \mathbf{A} \) and \( \mathbf{B} \) (dot and
cross) in polar coordinates (Greenberg, 1988).

\[ A = (A \cdot B)B + B \times (A \times B) \]
\[ = A_\parallel + A_\perp. \]
3.7 Transmission (ABCD) matrix composition method

Matrix composition: Matrix multiplication represents a composition of $2 \times 2$ matrices because the input to the second matrix is the output of the first [this follows from the definition of composition: $f(x) \circ g(x) = f(g(x))]$. Thus the ABCD matrix is also known as the transmission matrix, or occasionally the chain matrix. The general expression for the transmission matrix $T(s)$ is

$$
\begin{bmatrix}
V_1 \\
I_1
\end{bmatrix} =
\begin{bmatrix}
\mathcal{A}(s) & \mathcal{B}(s) \\
\mathcal{C}(s) & \mathcal{D}(s)
\end{bmatrix}
\begin{bmatrix}
V_2 \\
I_2
\end{bmatrix}.
$$

(3.7.0.1)

The four coefficients $\mathcal{A}(s), \mathcal{B}(s), \mathcal{C}(s), \mathcal{D}(s)$ are all complex functions of the Laplace frequency $s = \sigma + j\omega$ (see p. 120). Typically they are polynomials in $s$—$\mathcal{C}(s) = s^2 + 1$, for example. A sum and parallel combination of inductors, capacitors, and resistors always results in an impedance given by the ratio of two polynomials (Brune impedance). Thus such methods are called lumped-element networks. A symbolic eigenanalysis of $2 \times 2$ matrices may be found in Appendix B.3 (see p. 237).

It is a standard convention to always define the circuit into the node. Since the input current on the left of Eq. 3.7.0.1 is the same as the output current on the right ($I_2$), we need the negative sign on $I_2$ to match the sign convention of current into every node. When we use this construction, all the currents will all agree.

We have already used $2 \times 2$ matrix composition for: (1) representing complex numbers (see p. 26), (2) computing the $\gcd(m, n)$ of $m, n \in \mathbb{N}$ (see p. 44), (3) computing Pell’s equation (see p. 52), and (4) computing the Fibonacci sequence (see p. 53). It appears that $2 \times 2$ matrices have high utility.

Definitions of $\mathcal{A}, \mathcal{B}, \mathcal{C}, \mathcal{D}$: By writing the equations that correspond to Eq. 3.7.0.1, we can show that

$$
\begin{align*}
\mathcal{A}(s) &= \left. \frac{V_1}{V_2} \right|_{I_2=0}, & \mathcal{B}(s) &= -\left. \frac{V_1}{I_2} \right|_{V_2=0}, & \mathcal{C}(s) &= \left. \frac{I_1}{V_2} \right|_{I_2=0}, & \mathcal{D}(s) &= -\left. \frac{I_1}{I_2} \right|_{V_2=0}.
\end{align*}
$$

(3.7.0.2)

Each equation has a physical interpretation and a corresponding name. Functions $\mathcal{A}$ and $\mathcal{C}$ are said to be blocked because the output current $I_2$ is zero. Functions $\mathcal{B}$ and $\mathcal{D}$ are said to be short-circuited because the output voltage $V_2$ is zero. These two terms (blocked vs. short-circuited) are electrical engineering-centric, arbitrary, and fail to generalize to other cases; thus we should avoid these terms.

For example, in a mechanical system blocked would correspond to an output isometric (no length change) velocity of zero. In mechanics the isometric force is defined as the maximum applied force conditioned on zero velocity (the blocked force). Thus the short-circuited force ($\mathcal{B}$) would correspond to zero force, which is nonsense. These engineering-centric terms do not gracefully generalize, so better terminology is needed. Much of this was sorted out by Thévenin in about 1883 (Van Valkenburg, 1964a; Johnson, 2003; Kennelly, 1893).

$\mathcal{A}$ and $\mathcal{D}$ are called voltage (force) and current (velocity) transfer functions, since they are ratios of voltages and currents, whereas $\mathcal{B}$ and $\mathcal{C}$ are known as the transfer impedance and transfer admittance. For example, the unloaded (blocked) ($I_2 = 0$) output voltage $V_2 = I_1 / C$ corresponds to the isometric force in mechanics. In this way each term expresses an output (port 2) in terms of an input (port 1) for a given load condition.

Example: Figure 3.8 Two examples of networks that may be analyzed using the ABCD transmission matrix method.

Exercise #42
Derive the formula for $\mathcal{C}$ in terms of the input and output currents and voltages. Hint: See Eq. 3.7.0.2.
Solution: Writing out the lower equation gives $I_1 = C V_2 - \mathcal{D} I_2$ and setting $I_2 = 0$, we may obtain the equation for $C = I_1 / V_2 |_{I_2=0}$. 

Exercise #43
Can $C = 0$?
Solution: Yes, if $I_2 = 0$ and $I_1 = I_2$, then $C = 0$. For $C \neq 0$, there needs to be a finite shunt impedance across $V_1$, so that $I_1 \neq I_2 = 0$. 

3.7.1 Thévenin parameters of a source
An important concept in circuit theory is that of the Thévenin parameters: the open-circuit voltage and the short-circuit current. Their ratio defines the Thévenin impedance (Johnson, 2003). The open-circuit voltage is defined as the voltage $V_2$ when the load current $I_2 = 0$, which was shown in Eq. 3.7.0.2 to be $V_2 / I_1 |_{I_2=0} = 1 / C$. 

Thévenin Voltage: From Eq. 3.7.0.1 there are two definitions for the Thévenin voltage $V_{\text{Théven}}$, depending on the source on the left:

\[
\frac{V_{\text{Théven}}}{I_1} |_{I_2=0} = \frac{1}{C} \quad \text{and} \quad \frac{V_{\text{Théven}}}{V_1} |_{I_2=0} = \frac{1}{A}. \tag{3.7.1.3}
\]

A more general expression is needed when the source impedance is finite (neither voltage nor current).

Thévenin impedance The Thévenin impedance is the impedance looking into port 2 with $V_1 = 0$; thus

\[
Z_{\text{Théven}} = \left. \frac{V_2}{I_2} \right|_{V_1=0}. \tag{3.7.1.4}
\]

From the upper equation of Eq. 3.7.0.1, with $V_1 = 0$, we obtain $\mathcal{A} V_2 = \mathcal{B} I_2$; thus

\[
Z_{\text{Théven}} = \frac{\mathcal{B}}{\mathcal{A}}. \tag{3.7.1.5}
\]

3.7.2 The impedance matrix
With a bit of algebra, we can find the impedance matrix in terms of $\mathcal{A}, \mathcal{B}, \mathcal{C}, \mathcal{D}$ (Van Valkenburg, 1964a, p. 310):

\[
\begin{bmatrix} V_1 \\ V_2 \end{bmatrix} = \begin{bmatrix} z_{11} & z_{12} \\ z_{21} & z_{22} \end{bmatrix} \begin{bmatrix} I_1 \\ I_2 \end{bmatrix} = \frac{1}{C} \begin{bmatrix} \mathcal{A} & \Delta_T \\ \mathcal{D} & -\mathcal{D} \end{bmatrix} \begin{bmatrix} I_1 \\ -I_2 \end{bmatrix}. \tag{3.7.2.6}
\]

The determinate of the transmission matrix is $\Delta_T = \pm 1$, and if $C = 0$, the impedance matrix does not exist (See (P6) on p. 122 for a discussion).

Definitions of $z_{11}(s), z_{12}(s), z_{21}(s), z_{22}(s)$: The definitions of the matrix elements are easily read off of the equation as

\[
z_{11} \equiv \left. \frac{V_1}{I_1} \right|_{I_2=0}, \quad z_{12} \equiv \left. \frac{V_1}{I_2} \right|_{I_1=0}, \quad z_{21} \equiv \left. \frac{V_2}{I_1} \right|_{I_2=0}, \quad z_{22} \equiv \left. \frac{V_2}{I_2} \right|_{I_1=0}. \tag{3.7.2.7}
\]

These definitions follow trivially from Eq. 3.7.2.6 and each element has a physical interpretation. For example, the unloaded ($I_2 = 0$, also called blocked or isometric) input impedance is $z_{11}(s) = \mathcal{A}(s)/\mathcal{C}(s)$, while the unloaded transfer impedance is $z_{21}(s) = 1/\mathcal{C}(s)$. For reciprocal systems (Postulate 6, p. 122), $z_{12} = z_{21}$, since $\Delta_T = 1$. For antireciprocal systems, such as dynamic (also called magnetic) loudspeakers and microphones (Kim and Allen, 2013), $\Delta_T = -1$; thus $z_{21} = -z_{12} = 1/C$. Finally, $z_{22}$ is the impedance looking into port 2 with port 1 open/blocked ($I_1 = 0$). The problem with these basic definitions is that their physical interpretation is unclear. We can solve this problem by referring to Fig. 3.9, which is easier to understand than Eq. 3.7.2.6, as it allows us to quickly visualize the many relationships. Specifically, the circuit of Fig. 3.9 is given by the $\mathcal{T}(s)$ matrix

\[
\begin{bmatrix} V_1 \\ I_1 \end{bmatrix} = \begin{bmatrix} 1 + z_a y_b \\ y_b \end{bmatrix} \begin{bmatrix} z_c (1 + z_a y_b) + z_a \\ 1 + y_b z_c \end{bmatrix} \begin{bmatrix} V_2 \\ -I_2 \end{bmatrix}. \tag{3.7.2.8}
\]
Note that it is trivial to invert the $T(s)$ matrix because $\Delta_T = \pm 1$.

From the circuit elements defined in Fig. 3.9 (i.e., $z_a, z_c, y_b$) we can easily compute the impedance matrix elements of Eq. 3.7.2.6 (i.e., $z_{11}, z_{12}, z_{21}, z_{22}$). For example, the impedance matrix element $z_{11}$, in terms of $z_a$ and $y_b$, is easily read off of Fig. 3.9 as the sum of the series and shunt impedances:

$$z_{11}(s)|_{t_2=0} = z_a + \frac{1}{y_b} = \frac{A}{C}.$$

Given the impedance matrix, we can then compute transmission matrix $T(s)$—namely, from Eq. 3.7.2.6,

$$\frac{1}{C(s)} = z_{21}, \quad \frac{A(s)}{C(s)} = z_{11}.$$

Figure 3.9: Equivalent circuit for a transmission matrix. This allows us to better visualize the matrix elements in terms of complex impedances $z_a(s), z_c(s), y_b(s)$, as defined in this figure.

The theory is best modeled using the transmission matrix (Eq. 3.7.0.1), while experimental data are best modeled using the impedance matrix (Eq. 3.7.2.6).

**Rayleigh reciprocity:** Figure 3.9 is particularly helpful in understanding the Rayleigh reciprocity Postulate P6 ($\mathcal{P}(s) = \pm \mathcal{C}(s)$, pages 122, 260):

$$\left.\frac{V_2}{I_2}\right|_{I_2=0} = \left.\frac{V_1}{I_1}\right|_{I_1=0}.$$

This says that the output voltage over the input current is symmetric, which is obvious from Fig. 3.9.

The Thévenin voltage ($V_{\text{Th}}$), impedance ($Z_{\text{Th}}$) and reciprocity are naturally explained in terms of Fig. 3.9. This is the normal case of magnetic circuits, such as loudspeakers, where $y_b$ is represented by a gyrator (Kim and Allen, 2013) or electron spin in quantum mechanics.

### 3.7.3 Network power relationships

Impedance is a general concept, closely tied to the definition of power $\mathcal{P}(t)$ (and energy). Power is defined as the product of the effort (force) and the flow (current). As described in Table 3.2, these concepts are very general, applying to mechanics, electrical circuits, acoustics, thermal circuits, and any other case where conservation of energy applies. Two basic variables are defined, generalized force and generalized flow, also called conjugate variables. The product of the conjugate variables is the power, and the ratio is the impedance. For example, for the case of voltage and current,

$$\mathcal{P}(t) \equiv v(t)i(t), \quad v(t) = z(t) \ast i(t), \quad i(t) = y(t) \ast v(t)$$

where $\ast$ defines convolution (Sec. 4.7.4 page 164)

$$v(t) = z(t) \ast i(t) \equiv \int_{t=0}^{\infty} z(\tau) i(t-\tau) d\tau \leftrightarrow Z(\omega)I(\omega).$$

**Power vs. power series, linear vs. nonlinear** Another place where second-degree equations appear in physical applications is in energy and power calculations. The electrical power is given by the product of the voltage $v(t)$ and current $i(t)$ (or in mechanics as the force times the velocity). For example, if we define $\mathcal{P} = v(t)i(t)$ to be the power $\mathcal{P}$ [watts], then the total energy [joules] at time $t$ is (Van Valkenburg, 1964a, Sec. 14)

$$E(t) = \int_0^t v(t)i(t) dt.$$
3.7. TRANSMISSION (ABCD) MATRIX COMPOSITION METHOD

From this observe that the power is the rate of change of the total energy

\[ P(t) = \frac{d}{dt} E(t), \]

reminiscent of the fundamental theorem of calculus (Eq. 4.2.0.2, p. 135).

3.7.4 Ohm’s law and impedance

The ratio of voltage to current is called the impedance and it has units of [ohms]. For example, given a resistor of \( R = 10 \) [ohms],

\[ v(t) = R \, i(t); \]

namely, 1 [amp] flowing through the resistor would give 10 [volts] across it. Merging the linear relationship due to Ohm’s law with the definition of power shows that the instantaneous power in a resistor is quadratic in voltage and current:

\[ P(t) = v(t)^2 / R = i(t)^2 R. \]  \hspace{1cm} (3.7.4.9)

Note that Ohm’s law is linear in its relationship between voltage and current, whereas power and energy are nonlinear.

Ohm’s law generalizes in a very important way that allows the impedance (e.g., resistance) to be a linear complex analytic function of complex frequency \( s = \sigma + \omega \) (Kennelly, 1893; Brune, 1931a). Impedance is a fundamental concept in many fields of engineering. For example: Newton’s second law \( F = ma \) obeys Ohm’s law, with mechanical impedance \( Z(s) = sm \). Hooke’s law \( F = kx \) for a spring is described by a mechanical impedance \( Z(s) = k/s \). In mechanics a resistor is called a dashpot and its impedance is a positive-real constant.

Kirchhoff’s laws: KCL and KVL: The laws of electricity and mechanics may be written using Kirchhoff’s current and voltage laws (KCL and KVL), which lead to linear systems of equations in the currents and voltages (velocities and forces) of the system under study, with complex coefficients having positive-real parts.

Points of major confusion are a number of terms that are misused, and overused, in the fields of mathematics, physics, and engineering. Some of the most obviously abused terms are linear/nonlinear, energy, power, and power series. These have multiple meanings that can be, and are, fundamentally in conflict.

Transfer functions (transfer matrix): The only method that seems to work to sort out these terms is to cite the relevant physical application in specific contexts. The most common standard reference is a physical system that has an input \( x(t) \) and an output \( y(t) \). If the system is linear, then it may be represented by its impulse response \( h(t) \). In such cases, the system equation is

\[ y(t) = h(t) * x(t) \leftrightarrow Y(\omega) = H(s)|_{s=0} X(\omega); \]

namely, the convolution of the input with the impulse response gives the output. From Fourier analysis this relationship may be written in the real frequency domain as a product of the Laplace transform of the impulse response evaluated on the \( \omega \)-axis and the Fourier transform of the input \( X(\omega) = x(t) \) and output \( Y(\omega) = y(t) \).

If the system is nonlinear, then the output is not given by a convolution, and the Fourier and Laplace transforms have no obvious meaning.

The question that must be addressed is why the power is nonlinear, whereas a power series of \( H(s) \) is linear: Both have powers of the underlying variables. This is confusing and rarely, if ever, addressed. The quick answer is that powers of the Laplace frequency \( s \) correspond to derivatives, which are linear operations, whereas the product of the voltage \( v(t) \) and current \( i(t) \) is nonlinear. The important and interesting question will be addressed on page 121 in terms of the system postulates of physical systems.

Ohm’s law: In general, impedance is defined as the ratio of a force to a flow. For electrical circuits (Table 3.2), the voltage is the force and the current is the flow. Ohm’s law states that the voltage across and the current through a circuit element are linearly related by the impedance of that element (which is typically a complex function of the complex Laplace frequency \( s = \sigma + \omega \)). For resistors, the voltage over the current is called the resistance and is a constant (e.g., the simplest case is \( V/I = R \in \mathbb{R} \)). For inductors and capacitors, the impedance depends on the Laplace frequency \( s \) (e.g., \( V/I = Z(s) \in \mathbb{C} \)).

\(^{20}\) In acoustics the pressure is a potential, like voltage. The force per unit area is given by \( f = -\nabla p \); thus \( F = -\int \nabla p \, dS \). Velocity is analogous to a current. In terms of the velocity potential, the velocity per unit area is \( v = -\nabla \phi \).
Table 3.2: The generalized impedance is defined as the ratio of a force to a flow, a concept that also holds in mechanics and acoustics. In mechanics, the force is the mechanical force on an element (e.g., a mass, dashpot, or spring) and the flow is the velocity. In acoustics, the force is the pressure and the flow is the volume velocity or particle velocity of air molecules.

<table>
<thead>
<tr>
<th>Case</th>
<th>Force</th>
<th>Flow</th>
<th>Impedance units ohms [Ω]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrical</td>
<td>voltage (V)</td>
<td>current (I)</td>
<td>$Z = V/I$</td>
</tr>
<tr>
<td>Mechanics</td>
<td>force (F)</td>
<td>velocity (U)</td>
<td>$Z = F/U$ mechanical [Ω]</td>
</tr>
<tr>
<td>Acoustics</td>
<td>pressure (P)</td>
<td>particle velocity (V)</td>
<td>$Z = P/V$ specific [Ω]</td>
</tr>
<tr>
<td>Acoustics</td>
<td>mean pressure (p)</td>
<td>volume velocity (V')</td>
<td>$Z = p/V'$ acoustic [Ω]</td>
</tr>
<tr>
<td>Thermal</td>
<td>temperature (T)</td>
<td>entropy (S)</td>
<td>$Z = T/S$ thermal [Ω]</td>
</tr>
</tbody>
</table>

As shown in Table 3.2, the impedance concept also holds for mechanics and acoustics. In mechanics, the force is equal to the mechanical force on an element (e.g., a mass, dashpot, or spring) and the flow is the velocity. In acoustics, the force density is the pressure and the flow is the volume velocity or particle velocity of air molecules.

In this section we shall derive the method of the linear composition of systems, also known as the ABCD transmission matrix method, or in the mathematical literature, the M"{o}bius (bilinear) transformation. With the method of matrix composition, we can use a linear system of 2 × 2 matrices to represent a significant family of networks. By the application of Ohm’s law to the circuit shown in Fig. 3.10, we can model a cascade of such cells, which characterize transmission lines (Campbell, 1903).

\[ L = \frac{V_1}{I_1}, \]

\[ C = \frac{V_2}{I_2}. \]

Figure 3.10: A single LC cell of the LC transmission line. Every cell of any transmission line may be modeled by the ABCD method as the product of two matrices. For the example shown here, the inductance \( L \) of the coil and the capacitance \( C \) of the capacitor are in units of [henry/m] and [farad/m]; thus they depend on length \( \Delta x \) [m] that the cell represents. Note the flows are always defined as into the + node.

Example of the use of the ABCD matrix composition: Figure 3.10 shows a network composed of a series inductor (mass) that has an impedance \( Z_l = sL \) and a shunt capacitor (compliance) that has an admittance \( Y_c = sC \in C \). As determined by Ohm’s law, each equation describes a linear relationship between the current and the voltage. For the inductive impedance, applying Ohm’s law gives

\[ Z_l(s) = (V_1 - V_2)/I_1, \]

where \( Z_l(s) = Ls \in C \) is the complex impedance of the inductor. For the capacitive impedance, applying Ohm’s law gives

\[ Y_c(s) = (I_1 + I_2)/V_2, \]

where \( Y_c = sC \in C \) is the complex admittance of the capacitor.

Each of these linear impedance relationships may be written in a 2 × 2 matrix format. The series inductor (\( C = 0 \)) equation gives \( I_1 = -I_2 \)

\[ \begin{bmatrix} V_1 \\ I_1 \end{bmatrix} = \begin{bmatrix} 1 & Z_l \\ 0 & 1 \end{bmatrix} \begin{bmatrix} V_2 \\ -I_2 \end{bmatrix}, \]

(3.7.4.10)

while the shunt capacitor (\( L = 0 \)) equation yields \( V_1 = V_2 \)

\[ \begin{bmatrix} V_1 \\ I_1 \end{bmatrix} = \begin{bmatrix} 1 & 0 \\ Y_c & 1 \end{bmatrix} \begin{bmatrix} V_2 \\ -I_2 \end{bmatrix}. \]

(3.7.4.11)

When the second matrix equation for the shunt admittance (Eq. 3.7.4.11) is substituted into the series impedance equation (Eq. 3.7.4.10), we find that the ABCD matrix composition \( (T_{12} = T_1 \circ T_2) \) for the cell is the product of two matrices:

\[ \begin{bmatrix} V_1 \\ I_1 \end{bmatrix} = \begin{bmatrix} 1 & Z_l \\ 0 & 1 \end{bmatrix} \begin{bmatrix} 1 & 0 \\ Y_c & 1 \end{bmatrix} \begin{bmatrix} V_2 \\ -I_2 \end{bmatrix} = \begin{bmatrix} 1 + Z_lY_c & Z_l \\ Y_c & 1 \end{bmatrix} \begin{bmatrix} V_2 \\ -I_2 \end{bmatrix}. \]

(3.7.4.12)
Note that the determinant of the matrix $\Delta = AD - BC = 1$. This is not an accident, since the determinants of the two matrices are each 1; thus the determinant of their product is 1. Every cascade of series and shunt elements will always have $\Delta = 1$.

For the case of Fig. 3.10, Eq. 3.7.4.12 has $A(s) = 1 + s^2LC$, $B(s) = sL$, $C(s) = sC$, and $D = 1$. These equations characterize the four possible relationships of the cell’s input and output voltage and current. For example, the ratio of the output to input voltage, with the output unloaded, is

$$\frac{V_2}{V_1}\bigg|_{I_2=0} = \frac{1}{A(s)} = \frac{1}{1 + Z_t Y_c} = \frac{1}{1 + s^2LC}.$$  

This is known as the voltage divider relationship. To derive the current divider relationship, we use the lower equation with $V_2 = 0$:

$$\frac{-I_2}{I_1}\bigg|_{V_2=0} = 1.$$

**Exercise #44**

What happens if the roles of $Z$ and $Y$ are reversed?

**Solution:**

$$\begin{bmatrix} V_1 \\ I_1 \end{bmatrix} = \begin{bmatrix} 1 & 0 \\ Y_c & 1 \end{bmatrix} \begin{bmatrix} 1 \\ Z_t \end{bmatrix} \begin{bmatrix} V_2 \\ -I_2 \end{bmatrix} = \begin{bmatrix} 1 & Z_t \\ Y_c & 1 + Z_t Y_c \end{bmatrix} \begin{bmatrix} V_2 \\ -I_2 \end{bmatrix}$$  \hspace{1cm} (3.7.4.13)

This is the same network reversed in direction. ■

**Exercise #45**

What happens if the series element is a capacitor and the shunt an inductor?

**Solution:**

$$\begin{bmatrix} V_1 \\ I_1 \end{bmatrix} = \begin{bmatrix} 1 & 1/Y_c \\ 0 & 1 \end{bmatrix} \begin{bmatrix} 1 \\ Z_t \end{bmatrix} \begin{bmatrix} V_2 \\ 1 + 1/Z_t Y_c \end{bmatrix} \begin{bmatrix} 1 \\ 1/Z_t \end{bmatrix} \begin{bmatrix} V_2 \\ -I_2 \end{bmatrix}$$  \hspace{1cm} (3.7.4.14)

This circuit is a high-pass filter rather than a low-pass. ■

**Properties of the transmission matrix:** The transmission matrix is always constructed from the product of elemental matrices of the form

$$\begin{bmatrix} 1 & Z(s) \\ 0 & 1 \end{bmatrix} \quad \text{or} \quad \begin{bmatrix} 1 & 0 \\ Y(s) & 1 \end{bmatrix}.$$  

Thus for the case of reciprocal systems (Postulate P6, p. 122),

$$\Delta_T = \det \begin{bmatrix} A(s) & B(s) \\ C(s) & D(s) \end{bmatrix} = 1,$$

since the determinant of the product of each elemental matrix is 1 and the determinant of their product is 1. An antireciprocal system may be synthesized by the use of a gyrator, and for such cases $\Delta_T = -1$.

The eigenvalue and vector equations for a T matrix are summarized in Appendix B (p. 233) and discussed in Appendix B.3 (p. 237). The basic postulates of network theory also apply to the matrix elements $A(s), B(s), C(s), D(s)$, which place restrictions on their functional relationships. For example, Postulate P1 (p. 121) places limits on the poles and/or zeros of each function, since the time response must be causal.

### 3.8 Signals: Fourier transforms

Two fundamental tools in engineering mathematics are the Fourier and the Laplace transforms, which deal with time-frequency analysis (Papoulis, 1962).

The Fourier transform ($FT$) takes a time domain signal $f(t) \in \mathbb{R}$ and transforms it to the frequency domain by taking the scalar product (also called dot product) of $f(t)$ with the complex time vector $e^{-j\omega t}$:

$$f(t) \leftrightarrow F(\omega) = f(t) \cdot e^{-j\omega t},$$

where $F(\omega)$ and $e^{-j\omega t} \in \mathbb{C}$ and $\omega, t \in \mathbb{R}$. The scalar product between two vectors results in a scalar (number), as discussed in Appendix A.3 (p. 225).
Definition of the Fourier transform: The forward transform takes \( f(t) \) to \( F(\omega) \), while the inverse transform takes \( F(\omega) \) to \( \tilde{f}(t) \). The tilde indicates that, in general, the recovered inverse transform signal can be slightly different from \( f(t) \). Examples are presented in Table 3.3.

\[
F(\omega) = \int_{-\infty}^{\infty} f(t) e^{-j\omega t} dt \quad \tilde{f}(t) = \frac{1}{2\pi} \int_{-\infty}^{\infty} F(\omega) e^{j\omega t} d\omega \quad (3.8.0.1)
\]

It is accepted in the engineering and physics literature to use the case of the variable to indicate the type of argument. A time-domain function is \( f(t) \), where \( t \) has units of seconds [s] and is lowercase. Its Fourier transform is uppercase \( F(\omega) \) and is a function of frequency, having units of either hertz [Hz] or radians [2\(\pi\) Hz]. This case convention helps the reader parse the variable under consideration. This notation is a helpful but not agree with the notation used in mathematics, where units are rarely cited.

Table 3.3: Basic (Level I) Fourier transforms. Note that \( a > 0 \in \mathbb{R} \) has units [rad/s]. To flag this necessary condition, we use \([a]\) to assure this condition will be met. The other constant \( T_o \in \mathbb{R} [s] \) has no restrictions, other than being real. Complex constants may not appear as the argument to a delta function, since complex numbers do not have the order property.

<table>
<thead>
<tr>
<th>( f(t) \leftrightarrow F(\omega) )</th>
<th>Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \delta(t) \leftrightarrow 1(\omega) \equiv 1 \forall \omega )</td>
<td>Dirac</td>
</tr>
<tr>
<td>( 1(t) \equiv 1 \forall t \leftrightarrow 2\pi \delta(\omega) )</td>
<td>Dirac</td>
</tr>
<tr>
<td>( \text{sgn}(t) = \frac{t}{</td>
<td>t</td>
</tr>
<tr>
<td>( \tilde{u}(t) = \frac{1(t) + \text{sgn}(t)}{2} \leftrightarrow \pi \delta(\omega) + \frac{1}{j\omega} \equiv \tilde{U}(\omega) )</td>
<td>step</td>
</tr>
<tr>
<td>( \delta(t - T_o) \leftrightarrow e^{-j\omega T_o} )</td>
<td>delay</td>
</tr>
<tr>
<td>( \tilde{\delta}(t - T_o) * f(t) \leftrightarrow F(\omega)e^{-j\omega T_o} )</td>
<td>delay</td>
</tr>
<tr>
<td>( \tilde{u}(t)e^{-</td>
<td>a</td>
</tr>
<tr>
<td>( \text{rec}(t) = \frac{1}{T_o} [\tilde{u}(t) - \tilde{u}(t - T_o)] \leftrightarrow \frac{1}{T_o} (1 - e^{-j\omega T_o}) )</td>
<td>pulse</td>
</tr>
<tr>
<td>( \tilde{u}(t) * \tilde{u}(t) \leftrightarrow \delta^2(\omega) )</td>
<td>Not defined</td>
</tr>
</tbody>
</table>

Table 3.4: Abbreviations: \( \mathcal{F} \mathcal{T} \): Fourier Transform; FS: Fourier Series; DTFT: Discrete time Fourier transform; DFT: Discrete Fourier transform (the FFT is a “fast” DFT);

<table>
<thead>
<tr>
<th>FREQUENCY \ TIME</th>
<th>continuous ( \omega )</th>
<th>discrete ( \omega_k )</th>
<th>periodic ( ((\omega))_\Omega_n )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \mathcal{F} \mathcal{T} )</td>
<td>continuous ( \omega )</td>
<td>discrete ( \omega_k )</td>
<td>periodic ( ((\omega))_\Omega_n )</td>
</tr>
<tr>
<td>-</td>
<td>DFT (FFT)</td>
<td>FS</td>
<td></td>
</tr>
<tr>
<td>-</td>
<td>DTFT</td>
<td>DFT (FFT)</td>
<td></td>
</tr>
</tbody>
</table>

Types of Fourier Transforms: As summarized in Table 3.4, each \( \mathcal{F} \mathcal{T} \) type is determined by its time and frequency symmetries. A time function \( f(t) \) may be continuous in time, with \(-\infty < t < \infty\), discrete in time, \( f_n = f(t_n) \) with \( t_k = kT_s \), where \( T_o \) is the called the Nyquist sample period, or periodic in time, \( f((t))_{T_p} = f(t + kT_p) \), where \( T_p \) is called the period. Here \( k, n \in \mathbb{Z} \) and \( T_o, T_p \in \mathbb{R} \).

A general rule is that if a function is discrete in one domain (time or frequency), it is periodic in the other domain (frequency or time). For example, the discrete time function \( f_n \) must have a periodic frequency response—namely, \( f_n \leftrightarrow F((\omega))_{T_p} \). This is the case of discrete-time Fourier transform (DTFT). Alternatively, when the time
function is periodic, the frequencies must be discrete—namely, \( f((t)) \leftrightarrow F(\omega) \). This is the case of the Fourier series (FS). When both the time and frequencies are discrete, both the time and frequencies must be periodic. This is the case of the discrete Fourier transform (DFT). These three cases are summarized in Table 3.5.

1. Both time \( t \) and frequency \( \omega \) are real.
2. For the forward transform (time to frequency), the sign of the exponential is negative.
3. The limits on the integrals in both the forward and reverse FTs are \([-\infty, \infty]\).
4. When we take the inverse Fourier transform, the scale factor of \( \frac{1}{2\pi} \) is required to cancel the \( 2\pi \) in the frequency differential \( d\omega = 2\pi df \).
5. The Fourier step function is defined by the use of superposition of 1 and \( \text{sgn}(t) = t/|t| \) as

\[
\tilde{u}(t) \equiv \frac{1 + \text{sgn}(t)}{2} = \begin{cases} 
1 & t > 0 \\
1/2 & t = 0 \\
0 & t < 0 
\end{cases},
\]

Taking the FT of a delayed step function, we get

\[
\tilde{u}(t - T_o) \leftrightarrow \frac{1}{2} \int_{-\infty}^{\infty} [1 - \text{sgn}(t - T_o)] e^{-j\omega t} dt = \pi \tilde{\delta}(\omega) + \frac{e^{-j\omega T_o}}{j\omega}.
\]

Thus the FT of the step function has the term \( \pi \tilde{\delta}(\omega) \) due to the 1 in the definition of the Fourier step. This term introduces a serious flaw with the FT of the step function: While it appears to be causal, it is not. Compare this to the convolution \( u(t) * u(t) \) in Table C.3 on page 242.

6. The convolution \( \tilde{u}(t) * \tilde{u}(t) \) is not defined because both \( 1 * 1 \) and \( \tilde{\delta}^2(\omega) \) are not defined.

7. The inverse FT has convergence issues whenever there is a discontinuity in the time response. We indicate this with a hat over the reconstructed time response. The error between the target time function and the reconstructed is zero in the root-mean sense, but not point-wise.

Specifically, at the discontinuity point for the Fourier step function \( t = 0 \) \( \tilde{u}(t) \neq u(t) \), yet \( \int |\tilde{u}(t) - u(t)|^2 dt = 0 \). At the point of the discontinuity, the reconstructed function displays Gibbs ringing (it oscillates around the step and hence does not converge at the jump). The LT does not exhibit Gibbs ringing and thus is exact.

8. The FT is not always analytic in \( \omega \), as in this example of the step function. The step function cannot be expanded in a Taylor series about \( \omega = 0 \) because \( \tilde{\delta}(\omega) \) is not analytic in \( \omega \).

9. The Fourier \( \delta \) function is denoted \( \tilde{\delta}(t) \) to differentiate it from the Laplace delta function \( \delta(t) \). They differ because the step functions differ due to the convergence problem.

10. One may define

\[
\tilde{u}(t) = \int_{-\infty}^{t} \tilde{\delta}(t) dt
\]

and the somewhat questionable notation

\[
\tilde{\delta}(t) = \frac{d}{dt} \tilde{u}(t),
\]

since the Fourier step function is not analytic.

11. The \( \text{rec}(t) \) function is defined as

\[
\text{rec}(t) = \frac{\tilde{u}(t) - \tilde{u}(t - T_o)}{T_o} = \begin{cases} 
0 & t < 0 \\
1/T_o & 0 < t < T_o \\
0 & t > T_o
\end{cases}.
\]

It follows that \( \tilde{\delta}(t) = \lim_{T_o \to 0} \). Like \( \tilde{\delta}(t) \), the \( \text{rec}(t) \) has unit area.

12. When a function is periodic in one domain \((t, f)\), it must be discrete in the other (Table 3.5).
CHAPTER 3. ALGEBRAIC EQUATIONS: STREAM 2

Table 3.5: The general rule is that if a function is discrete in one domain (time or frequency) it is periodic in the other. Abbreviations: FT: Fourier Transform; FS: Fourier Series; DTFT: Discrete time Fourier transform; DFT: Discrete Fourier transform (the FFT is a “fast” DFT);

<table>
<thead>
<tr>
<th>FREQUENCY \ TIME</th>
<th>continuous t</th>
<th>discrete $t_k$</th>
<th>periodic ($((t))_{T_o}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>continuous $\omega$</td>
<td>$\mathcal{F}T$</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>discrete $\omega_k$</td>
<td>–</td>
<td>DFT (FFT)</td>
<td>FS</td>
</tr>
<tr>
<td>periodic $((\omega))_{\Omega}$</td>
<td>–</td>
<td>DTFT</td>
<td>DFT (FFT)</td>
</tr>
</tbody>
</table>

Table 3.6: Key properties of FTs

\[ \frac{d}{dt} v(t) \leftrightarrow j\omega V(\omega) \quad \text{deriv} \]
\[ f(t) \ast g(t) \leftrightarrow F(\omega)G(\omega) \quad \text{conv} \]
\[ f(t)g(t) \leftrightarrow \frac{1}{2\pi} F(\omega) \ast G(\omega) \quad \text{conv} \]
\[ f(at) \leftrightarrow \frac{1}{a} F \left( \frac{\omega}{a} \right) \quad \text{scaling} \]

Periodic signals: In addition to these two basic types of time-frequency transforms, there are several variants that depend on the symmetry in time and frequency. For example, when the time signal is sampled (discrete in time), the frequency response becomes periodic, leading to the DTFT. When a time response is periodic, the frequency response is sampled (discrete in frequency), leading to the FS. These two symmetries may be simply characterized as periodic in time $\Rightarrow$ discrete in frequency and periodic in frequency $\Rightarrow$ discrete in time. When a function is discrete in both time and frequency, it is necessarily periodic in time and frequency, leading to the DFT. The DFT is typically computed with an algorithm called the FFT, which can dramatically speed up the calculation when the data are a power of 2 in length.

Causal-periodic signals: A special symmetry occurs with functions that are causal and periodic in frequency. The best example is the $z$-transform, which are causal (one-sided in time) discrete-time signals. The harmonic series (Eq. 3.2.2.10, p. 78) is the $z$-transform of the discrete-time step function and is thus, due to symmetry, analytic within the RoC in the complex-frequency ($z$) domain.

The double brackets on $f(())_{T_o}$ indicate that $f(t)$ is periodic in $t$ with period $T_o$—that is, $f(t) = f(t + kT_o)$ for all $k \in \mathbb{N}$. Averaging over one period and dividing by $T_o$ give the average value.

Exercise #46
Consider the Fourier series scalar (dot) product (Eq. 3.5.0.2, p. 94) between “vectors” $f(())_{T_o}$ and $e^{-j\omega k t}$:

\[ F(\omega_k) = f(())_{T_o} \cdot e^{-j\omega_k t} \]

\[ = \frac{1}{T_o} \int_0^{T_o} f(t)e^{-j\omega_k t} dt, \]

where $\omega_0 = 2\pi / T_o$ and $f(t)$ has period $T_o$—that is, $f(t) = f(t + nT_o) = e^{j\omega_n t}$ with $n \in \mathbb{N}$ and $\omega_k = k\omega_0$.

What is the value of the Fourier series scalar product?

Solution: Evaluating the scalar product, we find

\[ e^{j\omega_n t} \cdot e^{-j\omega_k t} = \frac{1}{T_o} \int_0^{T_o} e^{j(\omega_n - \omega_k) t} dt \]

\[ = \frac{1}{T_o} \int_0^{T_o} e^{2\pi j(n-k)t/T_o} dt = \begin{cases} 1 & n = k, \\
0 & n \neq k. \end{cases} \]

The two signals (vectors) are orthogonal. ♦
Table 3.7: As summarized in this table of scalar products (dot products), the various types of Fourier transforms differ in their support in time and frequency. The transform types are Fourier transform, Fourier series, discrete-time Fourier transform, and fast Fourier transform (fast version of the DFT). The support then defines the inner product form. In this way all the various forms of Fourier transforms may be reduced to differences in the scalar product, as dictated by the support of the signals in time and frequency. In the above, 0 = N, 0 = f_k = k/N, represent discrete time and frequency samples, where 0 is one sample period. The signal period for the Fourier series (FS) is 0 [s]. For the discrete Fourier transform (DFT) the signal period is N, where N is the length of the DFT. Typically the transform length is taken to be a power of 2, such as N = 1024 samples. This is done to improve the speed of the transform, known as the fast Fourier transform (FFT). The term form provides the mathematical form of the scalar product which depends on the signal symmetry (finite-duration, periodic, causal/one-sided, discrete-time/frequency, continuous-time/frequency, etc.). ON stands for ortho-normal. This column shows the signals that are used when taking the transform. The signal is projected onto these vectors by the scalar product.

<table>
<thead>
<tr>
<th>Name</th>
<th>Domain</th>
<th>scalar product</th>
<th>form</th>
<th>ON</th>
</tr>
</thead>
<tbody>
<tr>
<td>FT</td>
<td>−∞ &lt; t ∈ ℝ &lt; ∞</td>
<td>x(t) · y(t)</td>
<td>∫ x(t) y(t) dt</td>
<td>e^{−j2πf t}</td>
</tr>
<tr>
<td>FT</td>
<td>−∞ &lt; f ∈ ℝ &lt; ∞</td>
<td>X(f) · Y(f)</td>
<td>∫ X(f) Y(f) df</td>
<td>e^{j2πf t}</td>
</tr>
<tr>
<td>FS</td>
<td>0 ≤ t ∈ ℝ ≤ T</td>
<td>x((t)) · y((t))</td>
<td>1/ T ∫ x(t) y(t) dt</td>
<td>e^{−j2πf t}</td>
</tr>
<tr>
<td>FS</td>
<td>−∞ &lt; f_k = k/T ∈ ℤ &lt; ∞</td>
<td>X_k · Y_k</td>
<td>∑_{k=−∞}^{∞} X_k Y_k</td>
<td>e^{j2πf_k t}</td>
</tr>
<tr>
<td>DTFT</td>
<td>−∞ &lt; t_n &lt; ∞</td>
<td>x_n · y_n</td>
<td>∑_{n=−∞}^{∞} x_n y_n</td>
<td>e^{−j2πt_n Ω}</td>
</tr>
<tr>
<td>DTFT</td>
<td>−π &lt; Ω &lt; π</td>
<td>X((Ω)) · Y((Ω))</td>
<td>∫_{−π}^{π} X(e^{jΩ}) Y(e^{jΩ}) dΩ</td>
<td>e^{j2πt_n Ω}</td>
</tr>
<tr>
<td>DFT/FFT</td>
<td>0 ≤ t_n = nT ≤ (N−1)T</td>
<td>x_n y_n</td>
<td>∑_{n=0}^{N−1} x_n y_n</td>
<td>e^{−j2πt_n f_k}</td>
</tr>
<tr>
<td>DFT/FFT</td>
<td>0 ≤ f_k = k/N ≤ (N−1)/NT</td>
<td>X_k Y_k</td>
<td>1/N ∑_{n=0}^{N−1} X_k Y_k</td>
<td>e^{j2πf_k t}</td>
</tr>
</tbody>
</table>

Exercise #47
Consider the discrete-time FT (DTFT) as a scalar (dot) product (Eq. 3.5.0.2, between “vectors” f_n = f(t) |_t_n, and e^{−jωt_n}, where t_n = nT_s and T_s = 1/2F_max is the sample period.

Solution: The scalar product over n ∈ ℤ is

\[
F((ω))_{2π} = \sum_{n=−∞}^{∞} f_n e^{−jωt_n} = \sum_{n=−∞}^{∞} f_n e^{−jωn},
\]

where \(ω_0 = 2π/T_s\) and \(ω_k = kω_0\) is periodic (i.e., \(F(ω) = F(ω + kω_0)\)).

3.9 Systems: Laplace transforms

The Laplace transform takes real causal signals \(f(t)u(t) ∈ ℝ\), as a function of real time \(t ∈ ℝ\), that are strictly zero for negative time \(f(t) = 0\) for \(t < 0\), and transforms them into complex analytic functions \(F(s) ∈ ℂ\) of complex frequency \(s = σ + jω\). For the case of a Fourier transform, we use the same notation: \(f(t) ↔ F(s)\).

When a signal is zero for negative time \(f(t < 0) = 0\), it is said to be causal, and the resulting transform \(F(s)\) must be complex analytic over significant regions of the s plane. For a function of time to be causal, time must be real \((t ∈ ℝ)\), since if it were complex, it would lose the order property (thus it could not be causal). It is helpful to emphasize the causal nature of \(f(t)u(t)\) to force causality, with the Heaviside step function \(u(t)\).

Any restriction on a function (e.g., real, causal, periodic, positive real part, etc.) is called a symmetry property. There are many forms of symmetry (Sec. 3.9.1). The concept of symmetry is very general and widely used in both mathematics and physics, where it is more generally known as group theory. As shown in Fig. 3.8, the two most common FT symmetries are continuous and discrete-time signal. One-sided periodic transforms also exist, such as the system shown in Fig. 3.3 (p. 70).

Definition of the Laplace transform: The forward and inverse Laplace transforms are shown in Eq. 3.9.0.1. Here \(s = σ + jω ∈ ℂ\) [2πHz] is the complex Laplace frequency in radians and \(t ∈ ℝ\) [s] is the time in seconds.
Tables of the more common transforms are provided in Appendix C Table C.3 (p. 242). Properties of more advanced \(\mathcal{LT}\) s are given in Table C.4 (p. 243).

Forward and inverse Laplace transforms:

\[
F(s) = \int_0^\infty f(t)e^{-st}dt \quad \quad \quad f(t) = \frac{1}{2\pi j} \int_{\sigma - j\infty}^{\sigma + j\infty} F(s)e^{st}ds
\]

(3.9.0.1)

\[F(s) \leftrightarrow f(t)\]

\[f(t) \leftrightarrow F(s)\]

When we deal with engineering problems, it is convenient to separate the signals we use from the systems that process them. We do this by treating signals, such as a music signal, differently from a system, such as a filter. In general, signals may start and end at any time. The concept of causality has no mathematical meaning in signal space. Systems, on the other hand, obey rigid rules (to assure that they remain physical). These physical restrictions are described in terms of the system postulates, which we present on page 121. There is a question as to why postulates are needed and which ones are the best choices. These questions are discussed in lectures by Feynman (1968, 1970a). The original video is also available online in many places—for example, on YouTube.

Table 3.8: Laplace transforms \(\mathcal{LT}\) s are complementary to the class of Fourier transforms \(\mathcal{FT}\) due to the fact that the time function must be a causal function. All \(\mathcal{LT}\) s are complex analytic in the complex frequency \(s = \sigma + j\omega\) domain. As an example, a causal function that is continuous but one-sided in time is the step function \(\mathcal{LT}\), which has the \(\mathcal{LT}\) of \(u(t)\) is \( \frac{1}{1/(1-z^{-n})} \). When a function is discrete in time but one-sided, it has a Zeta-transform. The discrete-time step function is \(u[n] = u[n] \leftrightarrow 1/(1-z^{-n})\).

<table>
<thead>
<tr>
<th>FREQUENCY \ TIME</th>
<th>continuous (\omega)</th>
<th>discrete (t[k])</th>
<th>causal-periodic ((t)) (T_z)</th>
</tr>
</thead>
<tbody>
<tr>
<td>continuous (\omega)</td>
<td>(\mathcal{LT})</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>discrete (\omega[k])</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>periodic (z^n[t])</td>
<td>–</td>
<td>(z)-Transform</td>
<td>–</td>
</tr>
</tbody>
</table>

As we discuss on page 152, to invert the \(\mathcal{LT}\) one must use the Cauchy residue theorem (CT-3), which requires closure of the contour \(C\) at \(\omega j \rightarrow \pm j\infty\),

\[
\oint_C = \int_{\sigma - j\infty}^{\sigma + j\infty} + \int_{C_{\infty}}
\]

where the path represented by \(C_{\infty}\) is a semicircle of infinite radius. For a causal, stable (e.g., doesn’t “blow up” in time) signal, all of the poles of \(F(s)\) must be inside of the Laplace contour, in the left half \(s\)-plane.

Example: Hooke’s law for a spring states that the force \(f(t)\) is proportional to the displacement \(x(t)\)—that is, \(f(t) = Kx(t)\). The formula for a dashpot is \(f(t) = Rv(t)\), and Newton’s famous formula for mass is \(f(t) = d[Mv(t)]/dt\), which for a constant \(M\) is \(f(t) = Mdv/dt\).

The equation of motion for the mechanical oscillator in Fig. 3.11 is given by Newton’s second law; the sum of the forces must balance to zero:

\[
M \frac{d^2x(t)}{dt^2} + R \frac{dx(t)}{dt} + Kx(t) = f(t) \leftrightarrow (M s^2 + Rs + K)X(s) = F(s). \quad \text{(3.9.0.2)}
\]

https://www.youtube.com/watch?v=xnzB_IHGyjg; https://www.youtube.com/watch?v=YaUlqXRPMmY; https://www.youtube.com/watch?v=xnzB_IHGyjg
3.9. SYSTEMS: LAPLACE TRANSFORMS

These three constants—mass $M$, resistance $R$, and stiffness $K$ ($\in \mathbb{R} \geq 0$)—are real and nonnegative. The dynamical variables are the driving force $f(t) \leftrightarrow F(s)$, the position of the mass $x(t) \leftrightarrow X(s)$, and its velocity $v(t) \leftrightarrow V(s)$, with $v(t) = \frac{dx(t)}{dt} \leftrightarrow V(s) = sX(s)$.

Newton’s second law (ca.1650) is the mechanical equivalent of Kirchhoff’s (ca.1850) voltage law (KVL), which states that the sum of the voltages around a loop must be zero. The gradient of the voltage results in a force on a charge (i.e., $F = qE$). The current may be thought of as the flow of charge.

Equation 3.9.0.2 may be re-expressed in the frequency domain in terms of an impedance (i.e., Ohm’s law), defined as the ratio of the force $F(s)$ to velocity $V(s) = sX(s)$, and the sum of three impedances:

$$Z(s) = \frac{F(s)}{V(s)} = \frac{Ms^2 + Rs + K}{s} = Ms + R + \frac{K}{s}. \quad (3.9.0.3)$$

**Example:** The divergent series

$$e^t u(t) = \sum_{n=0}^{\infty} \frac{1}{n!} t^n \leftrightarrow \frac{1}{s-1}$$

is a valid description of $e^t u(t)$, with an unstable pole at $s = 1$. For values of $|x - x_n| < 1$ ($x \in \mathbb{R}$), the analytic function $P(x)$ is said to have a region of convergence (RoC). For cases where the argument is complex ($s \in \mathbb{C}$), this is called the radius of convergence (RoC). We might call the region $|s - s_0| > 1$ the region of divergence (RoD) and $|s - s_0| = 0$ the singular circle. Typically the underlying function $P(s)$, defined by the series, has a pole on the singular circle.

There seems to be a conflict with the time response $f(t) = e^{at} u(t)$, that has a divergent series (unstable pole). I’m not sure how to explain this conflict, other than to point out that $t \in \mathbb{R}$; thus the series expansion of the diverging exponential is real analytic, not complex analytic. First, $f(t)$ has a Laplace transform with a pole at $s = 1$, in agreement with its unstable nature. Second, every analytic function must be single-valued. This follows from the fact that each term in Eq. 3.2.2.8 (p. 76) is single-valued. Third, analytic functions are “smooth,” since they may be differentiated an infinite number of times and the series still converges.

The key idea that every impedance must be complex analytic and $\geq 0$ for $\sigma > 0$ was first proposed by Otto Brune in his PhD thesis at MIT, supervised by Ernst A. Guillemin, an MIT electrical engineering professor who played an important role in the development of circuit theory and was a student of Arnold Sommerfeld.\(^{22}\) Other MIT advisers were Norbert Wiener and Vannevar Bush. Brune’s primary, but non-MIT, advisor was W. Cauer, who was trained in 19th-century German mathematics, perhaps under Sommerfeld (Brune, 1931b).

**Summary:** While the definitions of the FT ($\mathcal{F}$) and LT ($\mathcal{L}$) transforms may appear similar, they are not. The key difference is that the time response of the Laplace transform is causal, leading to a complex analytic frequency response. The frequency response of the Fourier transform is complex but not complex analytic, since the frequency $\omega$ is real. Fourier transforms do not have poles.

The concept of symmetry is helpful in understanding the many different types of time-frequency transforms. Two fundamental types of symmetry are causality and periodicity.

The Fourier transform $\mathcal{F}$ characterizes the steady-state response, while the Laplace transform $\mathcal{L}$ characterizes both the transient and steady-state responses. Given a causal system force response (Eq. 3.9.0.3), $F(s) \leftrightarrow f(t)$ with input velocity $V(s) \leftrightarrow v(t)$, the response is

$$f(t) = z(t) * v(t) \leftrightarrow Z(\omega) = F(s) \big|_{s=j\omega} V(\omega),$$

which says that the force is the convolution of the mechanical impedance $z(t)$ with the input velocity $v(t)$.

### 3.9.1 System postulates

Solutions of differential equations, such as the wave equation, are conveniently described in terms of mathematical properties, which we present here in 11 system postulates (see Appendix F, p. 257, for greater detail):

(P1) **Causality** (noncausal/acausal): Causal systems respond when acted upon. All physical systems obey causality. An example of a causal system is an integrator, which has a response of a step function. Filters are also examples of causal systems. Signals represent acausal responses. They do not have a clear beginning or

\(^{22}\)It must be noted that University of Illinois Professor ‘Mac’ Van VanValkenburg was arguably more influential in circuit theory during the same period. Mac’s books are certainly more accessible, but perhaps less widely cited.
end, such as the sound of the wind or traffic noise. A causal linear system is typically complex analytic and is naturally represented in the complex $s$ plane via Laplace transforms. A nonlinear system may be causal but not complex analytic.

(P2) **Linearity** (nonlinear): Linear systems obey superposition. Let two signals $x(t)$ and $y(t)$ be the inputs to a linear system, producing outputs $x'(t)$ and $y'(t)$. When the inputs are presented together as $ax(t) + by(t)$ with weights $a, b \in \mathbb{C}$, the output is $ax'(t) + by'(t)$. If either $a$ or $b$ is zero, the corresponding signal is removed from the output.

Nonlinear systems mix the two inputs, thereby producing signals that are not present in the input. For example, if the inputs to a nonlinear system are two sine waves, the output contains distortion components that have frequencies not present at the input. One example of a nonlinear system is one that multiplies the two inputs. A second is a diode, which rectifies a signal, letting current flow in only one direction. Most physical systems have some degree of nonlinear response, but this is not always desired. Other systems are designed to be nonlinear, such as the diode example.

(P3) **Passive** (active): An active system has a power source, such as a battery, while a passive system has no power source. Although you may consider a transistor amplifier to be active, it is so only when connected to a power source. Brune impedances satisfy the positive-real condition (Eq. 3.2.2.18, p. 79).

(P4) **Real** (complex) time response: All physical systems are Real in, Real out. They do not naturally have a power source. Although you may consider a transistor amplifier to be active, it is so only when connected to a power source. Brune impedances satisfy the positive-real condition (Eq. 3.2.2.18, p. 79).

(P5) **Time-invariant** (time varying): For a system to be a time-varying system, the output must depend on when the input signal starts or stops. If the output, relative to the input, is independent of the starting time, then the system is said to be time-invariant.

(P6) **Reciprocal** (non- or antireciprocal): In many ways this is the most difficult postulate to understand. It is best characterized by the ABCD matrix (see p. 110). If $\Delta_T = 1$, the system is said to be reciprocal. If $\Delta_T = -1$, it is said to be antireciprocal. The impedance matrix is reciprocal when $z_{12} = z_{21}$ and antireciprocal when $z_{12} = -z_{21}$. Dynamic loudspeakers are antireciprocal and must be modeled by a gyrator, which may be thought of as a transformer that swaps the force and flow variables. For example, the input impedance of a gyrator terminated by an inductor is a capacitor. This property is best explained by Fig. 3.9 (p. 112). For an extended discussion on reciprocity, see page 260.

(P7) **Reversibility** (nonreversible): If swapping the input and output of a system leaves the system invariant, it is said to be reversible. When $A = D$, the system is reversible. Note the distinction between reversible and reciprocal.

(P8) **Space-invariant** (space-variant): If a system operates independently as a function of where it physically is in space, then it is space-invariant. When the parameters that characterize the system depend on position, it is space-variant.

(P9) **Deterministic** (random): Given the wave equation along with the boundary conditions, the system’s solution may be deterministic, or not, depending on its extent. Consider a radar or sonar wave propagating out into uncharted territory. When the wave hits an object, the reflection can return waves that are not predicted due to unknown objects. This is an example where the boundary condition is not known in advance.

(P10) **Quasistatic** ($ka < 1$): Quasistatics follows the Nyquist sampling theorem for systems that have dimensions that are small compared to the local wavelength (Nyquist, 1924). This assumption fails when the frequency is raised (the wavelength becomes short). Thus this is also known as the long-wavelength approximation. Quasistatics is typically stated as $ka < 1$, where $k = 2\pi/\lambda = \omega/c_0$ and $a$ is the smallest dimension of the system. See page 209 for a detailed discussion of the role of quasistatics in acoustic horn wave propagation, and page 193 for a method on how to integrate the transmission matrix and Nyquist sampling.

Postulate P10 is closely related to the Feynman lecture *The "underlying unity" of nature*, where Feynman asks (Feynman, 1970b, Ch. 12-7): “Why do we need to treat the fields as smooth?” His answer is related to the wavelength of the probing signal relative to the dimensions of the object being probed. This raises the fundamental question: Are Maxwell’s equations a band-limited approximation to reality? Today we have no definite answer to this question.
The following quote seems relevant:

The Lorentz force formula and Maxwell’s equations are two distinct physical laws, yet the two methods yield the same results.

Why the two results coincide was not known. In other words, the flux rule consists of two physically different laws in classical theories. Interestingly, this problem was also a motivation behind the development of the theory of relativity by Albert Einstein. In 1905, Einstein wrote in the opening paragraph of his first paper on relativity theory, “It is known that Maxwell’s electrodynamics—as usually understood at the present time—when applied to moving bodies, leads to asymmetries which do not appear to be inherent in the phenomena.” But Einstein’s argument moved away from this problem and formulated special theory of relativity, thus the problem was not solved.

Richard Feynman once described this situation in his famous lecture (The Feynman Lectures on Physics, Vol. II, 1964), “we know of no other place in physics where such a simple and accurate general principle requires for its real understanding an analysis in terms of two different phenomena. Usually such a beautiful generalization is found to stem from a single deep underlying principle. . . . We have to understand the “rule” as the combined effects of two quite separate phenomena.”

(P11) Periodic ↔ discrete: When a function is discrete in one domain (e.g., time or frequency), it is periodic in the other (frequency or time).

Summary of the 11 system postulates: Each postulate has at least two categories. For example, (P1) is causal, noncausal, or acausal, while (P2) is linear or nonlinear. (P6) and (P9) apply to only two-port algebraic networks (those that have an input and an output). The others apply to both two- and one-port networks (e.g., an impedance is a one-port). An important example of a two-port is the antireciprocal transmission matrix of a dynamic (EM) loudspeaker (see p. 257).

Related forms of these postulates may be found in the network theory literature (Van Valkenburg, 1964a,b; Ramo et al., 1965). Postulates P1–P6 were introduced by Carlin and Giordano (1964), and Postulates P7–P9 were added by Kim et al. (2016). While linearity (P2), passivity (P3), realness (P4), and time-invariant (P5) are independent, causality (P1) is a consequence of linearity (P2) and passivity (P3) (Carlin and Giordano, 1964, p. 5).

3.9.2 Probability

Many things in life follow rules we don’t understand, and thus are unpredictable, yet they have structure due to some underlying poorly understood physics (e.g., quantum mechanics). Unlike mathematicians, engineers are taught to deal with uncertainty in terms of random processes using probability theory. For many this starts out as a large set of boring incomprehensible definitions, but once you begin to understand, it becomes interesting mathematics. It needs to be in your skin. If you don’t have an intuition for it, either keep working on it or else find another job. Don’t memorize a bunch of formulas, because that won’t work over the long run.

Some view probability as combinatorics. This is wrong. It is much more than that. From my view, based on the function of the auditory system, probability is about the signal processing of noise and signals (i.e., not combinatorics). The goal of probability is to find correlations in observations, such as the relative frequency of observations in sequential observations of events. Hamming (2004) presents an insightful discussion on probability.

Definitions:

1. An event is an unpredictable outcome (Papoulis and Pillai, 2002). For example, measuring the temperature $T(x, t) \in \mathbb{R}$ with $x \in \mathbb{R}^3$ at time $t$ [s] is an event. Measuring the temperature every hour gives 24 events per day [degrees/h]. Also, the single toss of a coin, resulting in \{H, T\}, is an event.

Exercise #48

What are the units of a temperature event?

Solution: Although we might think the answer is degrees, that unit is not the data that are being observed. Rather, the relative frequency of temperatures is important. For example, how many times was the event between 20° and 21° or between 22° and 27°? Events are dimensionless numbers with no units. ■
2. A trial is \( N \) events.

3. An experiment \( \{M, N\} \) is \( M \) trials of \( N \) events.

4. We must always keep track of the number of events so that we can compute the mean (i.e., average) and the uncertainty of an observable outcome.

5. The mean of many trials is the average.

6. A random variable \( X \) is the outcome from an experiment. A random variable rarely has stated units. For example, flipping a coin \( N = 8 \) times defines the number of trials.

**Exercise #49**
Give the units of coin flips?

\[ X \equiv \{H, H, H, T, H, T, T, T\} \]

**Solution:** The random variable \( \{H, T\} \) does not have units; it has random outcomes.

**Exercise #50**
How do you identify the meaning of a variable that is dimensionless (has no units)?

**Solution:** One must get creative. We can let \( H = 1 \) and \( T = -1 \) so that the mean can be zero.

**Exercise #51**
What are the mean and standard deviation of the coin toss?

**Solution:** To compute the mean (or standard deviation) we assign numbers to \( H \) and \( T \). For example, we let \( H = 1 \) and \( T = 0 \). Then we use the usual formula to compute the numerical values.

7. The expected value is the mean of \( N \) events.

**Exercise #52**
What is the difference between the mean, the expected value, and the average?

**Solution:** These terms all mean the same thing. Having several terms that mean the same thing is one of the many things that make probability theory so arbitrary. It is sloppy to have unclear terminology.

**Exercise #53**
How do you assign a numerical mean to random outcomes \( \{H, T\} \)?

**Solution:** If we let \( H = 1 \) and \( T = 0 \), then the mean is

\[ \mu = (1 + 1 + 1 + 0 + 1 + 0 + 1 + 0)/8 = 5/8. \]

It is critically important to keep track of the number of events \((N = 8\) in the Exercise 52). In some sense \( N \) is more important than the actual measured sequence. It is helpful to think of \( N \) as the independent variable and \( X \) as the dependent variable; that is, think of \( X(N) \), not \( N(X) \).

**Example:** We define a trial by flipping a coin \( N = 10 \) times. We form an experiment by \( M \) repeated trials \((M = 1000)\).
3.10 Complex analytic mappings (domain-coloring)

One of the most difficult aspects of complex functions of a complex variable is visualizing the mappings from the \( z = x + yj \) to \( w(z) = u + vj \) planes. For example, \( w(z) = \sin(x) \) is trivial;

\[
\sin(yj) = \frac{e^{-y} - e^y}{2j} = -j\sinh(y)
\]

is pure imaginary. However, the more general case

\[
w(z) = \sin(z) \in \mathbb{C}
\]

when \( z = x + yj \) is real (i.e., \( y = 0 \)) because \( \sin(x) \) is real. Likewise, the case where \( x = 0 \) is not easily visualized. And when \( u(x,y) \) and \( v(x,y) \) are less well-known functions, \( w(z) \) can be even more difficult to visualize. For example, if \( w(z) = J_0(z) \), then \( u(x,y) = u(x,y) \) and \( v(x,y) \) are the real and imaginary parts of the Bessel function.

Fortunately with computer software today, this problem of representation can be solved by adding color to the chart. An Octave/Matlab script\(^{24}\) \texttt{zviz.m} has been devised to make the charts shown in Fig. 3.12. Such charts are known as \textit{domain-coloring}.

In Fig. 3.12, rather than plotting \( u(x,y) \) and \( v(x,y) \) separately, domain-coloring allows us to display the entire function on one color chart (i.e., colorized plot). For this visualization we see the complex polar form of \( w(s) = |w|e^{j\omega} \) rather than the \( 2 \times 2 \) (four-dimensional) Cartesian graph \( w(x + yj) = u(x,y) + v(x,y)j \). On the left is the reference condition, the identity mapping \( w = s \), and on the right the origin has been shifted to the right and up by \( \sqrt{2} \).

Mathematicians typically use the abstract (i.e., nonphysical) notation \( w(z) \), where \( w = u + vi \) and \( z = x + yi \). Engineers typically work in terms of a physical complex impedance \( Z(s) = R(s) + jX(s) \) that has resistance \( R(s) \) and reactance \( X(s) \) [ohms] as a function of the complex Laplace radian frequency \( s = \sigma + \omega j \) [rad], as used, for example, with the Laplace transform (see p. 119).

In Fig. 3.12 we use both notations, with \( Z(s) = s \) on the left and \( w(s) = s - \sqrt{2}j \) on the right, where we show this color code as a 2 \( \times \) 2 dimensional domain-coloring graph. Intensity (dark to light) represents the magnitude of the function, while hue (color) represents the phase, where red is 0\(^\circ\), sea-green is 90\(^\circ\), blue-green is 135\(^\circ\), blue is 180\(^\circ\), and violet is -90\(^\circ\) (or 270\(^\circ\)).\(^{25}\)

The function \( w = s = |s|e^{j\theta} \) has a dark spot (zero) at \( s = 0 \) and becomes brighter away from the origin. On the right is \( w(s) = s - \sqrt{2}j \), which shifts the zero (dark spot) to \( s = \sqrt{2}j \). Thus domain-coloring gives the full \( 2 \times 2 \) complex analytic function mapping \( w(x,y) = u(x,y) + v(x,y)j \) in colorized polar coordinates.

**Visualizing complex functions:** The mapping from \( s = \sigma + \omega j \) to \( w(s) = u(\sigma,\omega j) + iv(\sigma,\omega j) \) is difficult to visualize because for each point in the domain \( s = \sigma + \omega j \), we would like to represent both the magnitude and phase (or real and imaginary parts) of \( w(s) \). A good way to visualize these mappings is to use color (hue) to represent the phase and intensity (dark to light) to represent the magnitude.

**Example:** Figure 3.13 shows a colorized plot of \( w(z) = \sin(\pi(s - j)/2) \) resulting from the Matlab/Octave command \texttt{zviz sin(pix*(s-j)/2)}. The abscissa (horizontal axis) is the real \( \sigma \)-axis and the ordinate (vertical axis) is the complex \( \xi \omega \)-axis. The graph is offset along the ordinate axis by \( 1j \), since the argument \( s - j \) causes a shift of the sine function by \( 1 \) in the positive imaginary direction. The visible zeros of \( w(s) \) appear as dark regions at \((-2,1), (0,1), (2,1)\). As a function of \( \sigma \), \( w(\sigma + 1j) \) oscillates between red (phase is zero degrees), meaning the function is positive and real, and sea-green (phase is 180\(^\circ\)), meaning the function is negative and real.
To use the program, we use the syntax `zviz <function of s>` (for example, we type `zviz s.^2`). Note the period between `s` and `^2`. This will render a domain-coloring (colorized) version of the function. Examples you can render with `zviz` are given in the comments at the top of the `zviz.m` program. A good example for testing is `zviz z-sqrt(j)`, which will show a dark spot (zero) at `(1 + 1j) / sqrt(2) ≈ 0.707(1 + 1j).

Along the vertical axis, the displayed function is either `cosh(y)` or `sinh(y)`, depending on the value of `x`. The intensity becomes lighter as `|w|` increases.

**What is being plotted?** The axes are either `s = σ` and `ω`, or `z = x` and `y`. Superimposed on the `s`-axis is the function `w(s) = u(σ, ω) + v(σ, ω)j`, represented in polar coordinates by the intensity and color of `w(s)`. The density (dark vs. light) displays the magnitude `|w(s)|`, while the color (hue) displays the angle (angle of `w(s)`) as a function of `s`. Thus the intensity becomes darker as `|w|` decreases and lighter as `|w|` increases. The angle of `w` to color map is defined by Fig. 3.12. For example, `0°` is red, `90°` is green, `-90°` is purple, and `180°` is blue-green.

**Example:** Additional examples are given in Fig. 3.14 using the notation `w(s) = u(σ, ω) + v(σ, ω)j`. We see the two complex mappings `w = e^s` (left) and its inverse `s = ln(w)`. The exponential is relatively easy to understand because `w(s) = |e^σ e^jω| = e^σ`.

![Figure 3.13: Plot of sin(0.5π(z − j)).](image)

![Figure 3.14: This domain-color map allows us to visualize complex mappings by the use of intensity (light/dark) to indicate magnitude and color (hue) to indicate angle (phase). The white and black lines are the iso-real and iso-imaginary contours of the complex mapping. Left: The domain-color map for the complex mapping from the plane to the σ-ω plane, which goes to zero as `σ → −∞`, causing the domain-color map to become dark for `σ < −2`. The white and black lines are always perpendicular because `e^σ` is complex analytic everywhere. Right: The principal value of the inverse function `s(z) = u(x, y) + v(x, y)j = log(z)`, which has a zero (dark) at `x = 1`, since there `log(1) = 0` and the imaginary part is zero. Note the branch cut, where the color is discontinuous, from `x = [0, −∞j]`.](image)

The red region is where `ω ≈ 0`, in which case `w ≈ e^σ`. As `σ` becomes large and negative, `w → 0`; thus the entire field becomes dark on the left. The field is becoming light on the right where `w = e^σ → ∞`. If we let `σ = 0`
and look along the $\omega$-axis, we see that the function is changing phase: sea-green (90°) at the top and violet (-90°) at the bottom.

In the right panel note the zero for $s(z) = \ln(z) = \ln|z| + \omega \phi$ at $z = 1$. The root of the $\log(z)$ function is $\log(z_r) = 0$, $w_r = 1$, $\omega \phi = 0$, since $\log(1) = 0$. More generally, the $\log(z)$ of $z = |z|e^{i\phi}$ is $s(z) = \ln|z| + \omega \phi$. Thus $s(w)$ can be zero only when the angle of $w$ is zero.

The $\ln(z)$ function has a branch cut along the $\phi(z) = \angle z = 180^\circ$ axis. As one crosses over the cut, the phase goes above 180° and the plane changes to the next sheet of the log function. The only sheet with a zero is the principal value, as shown. For all others, the log function is either increasing or decreasing monotonically and there is no zero, as seen for sheet 0 (the one shown in Fig. 3.14).

### 3.10.1 The Riemann sphere

Once algebra was formulated, by about 830 CE, mathematicians were able to expand beyond the limits set by geometry on the real plane and the verbose descriptions of each problem in prose (Stillwell, 2010, p. 93). The geometry of Euclid’s Elements had paved the way, but after 2000 years, the addition of the language of algebra changed everything. The analytic function was a key development, heavily used by both Newton and Euler. Also Cauchy made important headway with his investigation of complex variables. Of special note were integration and differentiation in the complex plane of complex analytic functions, which is the topic of Chapters 4 and 5.

It was Riemann, working with Gauss in the final years of Gauss’s life, who made the breakthrough with the concept of the extended complex plane. This concept was based on the composition of a line with the sphere, similar to the derivation of Euclid’s formula for Pythagorean triplets (see p. 50). While the importance of the extended complex plane was unforeseen, it changed analytic mathematics forever, along with the physics it supported. It unified and thus simplified many important integrals to the extreme. The basic idea is captured by the fundamental theorem of complex integral calculus (see Table 4.1, p. 135).

The idea is outlined in Fig. 3.15. On the left is a circle and a line. The difference between this case and the derivation of the Pythagorean triplets is that the line starts at the north pole and ends on the real $x \in \mathbb{R}$ axis at point $x$. At point $x'$, the line cuts through the circle. Thus the mapping from $x$ to $x'$ takes every point on $\mathbb{R}$ to a point on the circle. For example, the point $x = 0$ maps to the south pole (not indicated). To express $x'$ in terms of $x$ one must compose the line and the circle, similar to the composition used in the derivation of Euclid’s formula (see p. 50). The points on the circle, indicated here by $x'$, require a traditional polar coordinate system, that has a unit radius and an angle defined between the radius and a vertical line passing through the north pole. When $x \to \infty$, the point $x' \to N$, known as the point at infinity. But this idea goes much further, as shown on the right half of Fig. 3.15.

Here the real tangent line is replaced by a tangent complex plane $z \in \mathbb{C}$ and the complex puncture point $z' \in \mathbb{C}$—in this case on the complex sphere, called the extended complex plane. This is a natural extension of the chord/tangent method on the left, but with significant consequences. The main difference between the complex plane $z$ and the extended complex plane, other than the coordinate system, is what happens at the north pole. The point at $|z| = \infty$ is not defined on the plane, whereas on the sphere, the point at the north pole is simply another point, like every other point on the sphere.

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26Gauss did lecture to Riemann but he was only giving elementary courses and there is no evidence that at this time he recognized Riemann’s genius.” Then “In 1849 he [Riemann] returned to Göttingen and his Ph.D. thesis, supervised by Gauss, was submitted in 1851.” See https://www-groups.dcs.st-and.ac.uk/~history/Biographies/Riemann.html.
Open vs. closed sets: Mathematically the plane is said to be an open set, since the limit $z \to \infty$ is not defined, whereas on the sphere, the point $z'$ is a member of a closed set, since the north pole is defined. The distinction between an open and closed sets is important because the closed set allows the function to be complex analytic at the north pole, which it cannot be on the plane (since the point at infinity is not defined).

The $z$ plane may be replaced with another plane—say, the $w = F(z) \in \mathbb{C}$ plane, where $w$ is some function $F$ of $z \in \mathbb{C}$. For the moment we shall limit ourselves to complex analytic functions of $z$—namely, $w = F(z) = u(x, y) + v(x, y)j = \sum_{n=0}^{\infty} a_n z^n$.

In summary, given a point $z = x + yj$ on the open complex plane, we map it to $w = F(z) \in \mathbb{C}$, the complex $w = u + vj$ plane, and from there to the closed extended complex plane $w'(z)$. The point of doing this is that it allows us to allow the function $w'(z)$ to be analytic at the north pole, meaning it can have a convergent Taylor series at the point at infinity $z \to \infty$. Since we have not yet defined $dw(z)/dz$, the concept of a complex Taylor series remains undefined.

3.10.2 Bilinear transformation

In mathematics the bilinear transformation has special importance because it is linear in its action on both the input and output variables. Since we are engineers, we shall stick with the engineering terminology. But if you wish to read about this on the internet, be sure to also search for the mathematical term Möbius transformation.

When a point on the complex plane $z = x + yj$ is composed with the bilinear transformation $(a, b, c, d \in \mathbb{C})$, the result is $w(z) = u(x, y) + v(x, y)j$ (this is related to the Möbius transformation, p. 27):

$$w = \frac{az + b}{cz + d}. \tag{3.10.2.1}$$

The transformation $z \to w$ is a cascade of four independent compositions:

1. Translation ($w = z + b; a = 1, b \in \mathbb{C}, c = 0, d = 1$)
2. Scaling ($w = |a|z; a \in \mathbb{R}, b = 0, c = 0, d = 1$)
3. Rotation ($w = \frac{a}{|a|}z; a \in \mathbb{C}, b = 0, c = 0, d = |a|$)
4. Inversion ($w = \frac{1}{z}; a = 0, b = 1, c = 1, d = 0$)

Each of these transformations is a special case of Eq. 3.10.2.1, with the inversion the most complicated. I highly recommend a video showing the effect of the bilinear (Möbius) transformation on the plane (Arnold, D. and Rogness, J., 2019).\(^{27}\)

The bilinear transformation is the most general way to move the expansion point in a complex analytic expansion. For example, when we start from the harmonic series, the bilinear transformation gives

$$\frac{1}{1 - w} = \frac{1}{1 - \frac{az + b}{cz + d}} = \frac{cz + d}{(c - a)z + (d - b)} = \frac{1}{\frac{1}{c} - \frac{a - b}{c - a}}. $$

The RoC is transformed from $|w| < 1$ to $|(az - b)/(cz - d)| < 1$. An interesting application might be to move the expansion point until it is on top of the nearest pole, so that the RoC goes to zero. This might be a useful way of finding a pole, for example.

When the extended plane (Riemann sphere) is analytic at $z = \infty$, we can take the derivatives there, defining the Taylor series with the expansion point at $\infty$. When the bilinear transformation rotates the Riemann sphere, the point at infinity is translated to a finite point on the complex plane, revealing the analytic nature at infinity. A second way to transform the point at infinity is by the bilinear transformation $\zeta = 1/z$, mapping a zero (or pole) at $z = \infty$ to a pole (or zero) at $\zeta = 0$. Thus this construction of the Riemann sphere and the Möbius (bilinear) transformation allows us to understand the point at infinity and treat it like any other point. If you felt that you never understood the meaning of the point at $\infty$ (likely), this should help.

\(^{27}\)https://www.youtube.com/watch?v=0zlfIsUNh04
3.11 Problems AE-3

Topics of this homework:
Visualizing complex functions, bilinear/Möbius transformation, Riemann sphere. Deliverables: Answers to problems

Two-port network analysis

Problem #1: Perform an analysis of electrical two-port networks, shown in Fig. 3.8 (page 110). This can be a mechanical system if the capacitors are taken to be springs and inductors taken as mass, as in the suspension of the wheels of a car. In an acoustical circuit, the low-pass filter could be a car muffler. While the physical representations will be different, the equations and the analysis are exactly the same.

The definition of the ABCD transmission matrix \( T \) is
\[
\begin{bmatrix}
V_1 \\
I_1
\end{bmatrix} =
\begin{bmatrix}
A & B \\
C & D
\end{bmatrix}
\begin{bmatrix}
V_2 \\
-I_2
\end{bmatrix}.
\]
(AE-3.1)

The impedance matrix, where the determinant \( \Delta_T = AD - BC \), is given by
\[
\begin{bmatrix}
V_1 \\
V_2
\end{bmatrix} = \frac{1}{C}
\begin{bmatrix}
A & \Delta_T \\
1 & D
\end{bmatrix}
\begin{bmatrix}
I_1 \\
I_2
\end{bmatrix}.
\]
(AE-3.2)

– 1.1: Derive the formula for the impedance matrix (Eq. AE-3.2) given the transmission matrix definition (Eq. AE-3.1). Show your work.

Problem #2: Consider a single circuit element with impedance \( Z(s) \).

– 2.1: What is the ABCD matrix for this element if it is in series?

– 2.2: What is the ABCD matrix for this element if it is in shunt?

Problem #3: Find the ABCD matrix for each of the circuits of Fig. 3.8.
For each circuit, (i) show the cascade of transmission matrices in terms of the complex frequency \( s \in \mathbb{C} \), then (ii) substitute \( s = 1j \) and calculate the total transmission matrix at this single frequency.

– 3.1: Left circuit (let \( R_1 = R_2 = 10 \, \text{k} \Omega \) (kilo-ohms) and \( C = 10 \, \text{nF} \) (nano-farads))

– 3.2: Right circuit (use \( L \) and \( C \) values given in the figure), where the pressure \( P \) is analogous to the voltage \( V \), and the velocity \( U \) is analogous to the current \( I \).

– 3.3: Convert both transmission (ABCD) matrices to impedance matrices using Eq. AE-3.2. Do this for the specific frequency \( s = 1j \) as in the previous part (feel free to use Matlab/Octave for your computation).

– 3.4: Right circuit: Using the previous solution and Matlab/Octave:
Algebra

Problem #4: Fundamental theorem of algebra (FTA).

- 4.1: State the fundamental theorem of algebra (FTA).

Problem #5: Order and complex numbers:
One can always say that \( 3 < 4 \)—namely, that real numbers have order. One way to view this is to take the difference and compare it to zero, as in \( 4 - 3 > 0 \). Here we will explore how complex variables may be ordered. Define the complex variable \( z = x + iy \in \mathbb{C} \).

- 5.1: Explain the meaning of \(|z_1| > |z_2|\).

- 5.2: If \( x_1, x_2 \in \mathbb{R} \) (are real numbers), define the meaning of \( x_1 > x_2 \). Hint: Take the difference.

- 5.3: Explain the meaning of \( z_1 > z_2 \).

- 5.4: If time were complex, how might the world be different? (not graded)

Problem #6: It is sometimes necessary to consider a function \( w(z) = u + iv \) in terms of the real functions \( u(x, y) \) and \( v(x, y) \) (e.g. separate the real and imaginary parts). Similarly, we can consider the inverse \( z(w) = x + iy \), where \( x(u, v) \) and \( y(u, v) \) are real functions. Find \( u(x, y) \) and \( v(x, y) \) for \( w(z) = \frac{1}{z} \).

Problem #7: Find \( u(x, y) \) and \( v(x, y) \) for \( w(z) = ce^z \) with complex constant \( c \in \mathbb{C} \) for questions 7.1, 7.2, and 7.3:

- 7.1: \( c = e \)

- 7.2: \( c = 1 \) (recall that \( 1 = e^{jk2\pi k} \) for \( k = 0, 1, 2, \ldots \))

- 7.3: \( c = j \). Hint: \( j = e^{\pi/2 + j2\pi m}, \quad m \in \mathbb{Z} \).

- 7.4: Find \( u(x, y) \) for \( w(z) = \sqrt{z} \). Hint: Begin with the inverse function \( z = w^2 \).

Problem #8: Convolution of an impedance \( z(t) \) and its inverse \( y(t) \):
In the frequency domain a Brune impedance is defined as the ratio of a numerator polynomial \( N(s) \) to a denominator polynomial \( D(s) \).

- 8.1: Consider a Brune impedance defined by the ratio of numerator and denominator polynomials, \( Z(s) = N(s)/D(s) \). Since the admittance \( Y(s) \) is defined as the reciprocal of the impedance, the product must be 1. If \( z(t) \leftrightarrow Z(s) \) and \( y(t) \leftrightarrow Y(s) \), it follows that \( z(t) * y(t) = \delta(t) \). What property must \( n(t) \leftrightarrow N(s) \) and \( d(t) \leftrightarrow D(s) \) obey for this to be true?

- 8.2: The definition of a minimum phase function is that it must have a causal inverse. Show that every impedance is minimum phase.
Schwarz inequality

**Problem # 9:** Figure 2.4 page (same as Fig. 3.5 page 99) shows three vectors for an arbitrary value of $\alpha \in \mathbb{R}$ and a specific value of $\alpha = \alpha^*$. 

- 9.1: Given Fig. 2.4, find the value of $\alpha \in \mathbb{R}$ such that the length (norm) of $E$ (i.e., $||E|| \geq 0$) is minimum. Show your derivation, not the answer ($\alpha = \alpha^*$).

- 9.2: Find the formula for $||E(\alpha^*)||^2 \geq 0$. Hint: Substitute $\alpha^*$ into Eq. AE-3.4 and show that this results in the Schwarz inequality:

$$||\vec{U} \cdot \vec{V}|| \leq ||\vec{U}|| ||\vec{V}||.$$

**Problem # 10:**

- 10.1 What is the geometrical meaning of the dot product of two vectors?

- 10.2: Give the formula for the dot product of two vectors. Explain the meaning based on Fig. 3.4 (page 94).

- 10.3: Write the formula for the dot product of two vectors $\vec{U} \cdot \vec{V}$ in $\mathbb{R}^n$ in polar form (e.g., assume the angle between the vectors is $\theta$).

- 10.4: How is the Schwarz inequality related to the Pythagorean theorem?

- 10.5: Starting from $||\vec{U} + \vec{V}||$, derive the triangle inequality

$$||\vec{U} + \vec{V}|| \leq ||\vec{U}|| + ||\vec{V}||.$$

- 10.6: The triangle inequality $||\vec{U} + \vec{V}|| \leq ||\vec{U}|| + ||\vec{V}||$ is true for two and three dimensions: Does it hold for five-dimensional vectors?

- 10.7: Show that the wedge product $\vec{U} \wedge \vec{V} \perp \vec{U} \cdot \vec{V}$.

**Probability**

**Problem # 11:** Basic terminology of experiments
– 11.1: What is the mean of a trial, and what is the average over all trials?

– 11.2: What is the expected value of a random variable $X$?

– 11.3: What is the standard deviation about the mean?

– 11.4: What is the definition of information of a random variable?

– 11.5: How do you combine events? Hint: If the event is the flip of a biased coin, the events are $H = p$, $T = 1 - p$, so the event is $\{p, 1 - p\}$. To solve the problem, you must find the probabilities of two independent events.

– 11.6: What does the term independent mean in the context of question 11.5? Give an example.

– 11.7: Define odds.
Chapter 4

Stream 3A: Scalar Calculus

Stream 3 is infy, a concept that typically means unbounded (immeasurably large), but in the case of calculus, $\infty$ means infinitesimal (immeasurably small), since taking a limit requires small numbers. Taking a limit means you may never reach the target, a concept that the Greeks called Zeno’s paradox (Stillwell, 2010, p. 76).

When we speak of the class of ordinary (versus vector) differential equations, the term scalar is preferable, since the term ordinary is vague if not meaningless. For scalar calculus, a special subset of fundamental theorems about integration are summarized in Table 4.1 (Sec. 4.2), starting with Leibniz’s theorem.

Following our discussion of the integral theorems on scalar calculus are those on vector calculus, without which there can be no understanding of Maxwell’s equations. Of these, the fundamental theorem of vector calculus (also known as Helmholtz decomposition), Gauss’s law, and Stokes’s theorem form the three cornerstones of modern vector field analysis. These theorems allow us to connect the differential (point) and macroscopic (integral) relationships. For example, we can write Maxwell’s equations either as vector differential equations, as shown by Heaviside (along with Gibbs and Hertz), or in integral form. It is helpful to place these two forms side by side to fully appreciate their significance. To understand the differential (microscopic) view, one must understand the integral (macroscopic) view (see Figs. 5.5 and 5.6 on pp. 197 and 198).

4.1 The beginning of modern mathematics

As shown in Fig. 1.2 (p. 15), mathematics as we know it today began in the 16th to 18th centuries, arguably starting with Galileo, Descartes, Fermat, Newton, the Bernoulli family, and most important Euler. Galileo was formidable because of his fame, fortune, and “successful” stance against the powerful Catholic establishment. His creativity in scientific circles was certainly well known due to his many skills and accomplishments. Descartes and Fermat were at the forefront of merging algebra and geometry. While Fermat kept meticulous notebooks, he did not publish and tended to be secretive. Thus Descartes’s contributions were more widely acknowledged though not necessarily deeper.

Regarding the development of calculus, much was yet to be developed by Newton and Leibniz using term-by-term integration of functions based on Taylor series representation. This was a powerful technique but, as stated earlier, incomplete because the Taylor series can represent only single-valued functions within the RoC. More important, Newton (and others) failed to recognize (i.e., rejected) the powerful generalization to complex analytic functions. The first major breakthrough was Newton’s publication of Principia (1687), and the second was by Riemann (1851), advised by Gauss but possibly more influenced by Cauchy.

Following Newton’s lead, the secretive and introverted behavior of the typical mathematician dramatically changed with the Bernoulli family (Fig. 3.1, p. 62). The oldest brother Jacob taught his much younger brother Johann, who then taught his son Daniel. But Johann’s star pupil was Leonhard Euler. Euler first mastered all the tools and then published with a prolificacy previously unknown.

**Euler and the circular functions:** Euler’s first major task was to understand the family of analytic circular functions—$e^x, \sin(x), \cos(x)$, and $\log(x)$—a task begun by the Bernoulli family. Euler sought relationships among these many functions, some of which may not be thought of as being related, such as the log and sin functions. The connection that may “easily” be made is through their complex Taylor series representation (Eq. 3.2.2.9, p. 76). By the manipulation of the analytic series representations, the relationship between $e^x$ and $\sin(x)$ and $\cos(x)$ was precisely captured with the equation

$$e^{j\omega} = \cos(\omega) + j\sin(\omega) \quad (4.1.0.1)$$
and its analytic inverse (Greenberg, 1988, p. 1135)

\[
\tan^{-1}(z) = \frac{1}{2j} \ln \left( \frac{1j - z}{1j + z} \right) = \frac{j}{2} \ln \left( \frac{1 - zj}{1 + zj} \right).
\] (4.1.0.2)

**Exercise #1**
Starting from Eq. 4.1.0.1, derive Eq. 4.1.0.2.

**Solution:** We let \( z(\omega) = \tan \omega \); then

\[
z(\omega) = \frac{\sin \omega}{\cos \omega} = \tan(\omega) = -\frac{e^{j\omega} - e^{-j\omega}}{e^{j\omega} + e^{-j\omega}} = -j \frac{e^{2j\omega} - 1}{e^{2j\omega} + 1}.
\] (4.1.0.3)

Solving for \( e^{-2j\omega} \), we get

\[
e^{-2j\omega} = \frac{1 - zj}{1 + zj}.
\] (4.1.0.4)

Taking \( \ln() \) of both sides and using the definition of \( z(\omega) \) give Eq. 4.1.0.2:

\[
\omega = \tan^{-1}(z) = \frac{j}{2} \ln \frac{1 - zj}{1 + zj}.
\]

The two sides of this equation are shown in Fig. 4.1. These equations are the basis of transmission lines (TL). Here \( z(\omega) \) of Eq. 4.1.0.3 is the TL’s input impedance and Eq. 4.1.0.4 is the reflectance. ■

**Figure 4.1:** Colorized plots of \( \omega(z) = \tan^{-1}(z) \) and \( \omega(z) = \frac{j}{2} \ln(1 - iz)/(1 + iz) \), verifying that they are the same complex analytic function.

Although many high school students memorize Euler’s relationship, it seems unlikely that they appreciate the utility of complex analytic functions (Eq. 3.2.4.21, p. 81).

**History of complex analytic functions:** Newton (ca.1650) famously ignored imaginary numbers and called them imaginary in a disparaging (pejorative) way. Given Newton’s prominence, his view must have keenly attenuated interest in complex algebra, even though it had been described by Bombelli in 1526, likely based on his serendipitous finding of Diophantus’s book *Arithmetic* in the Vatican library.

Euler derived his relationships using real power series (i.e., real analytic functions). While Euler was fluent with \( j = \sqrt{-1} \), he did not consider functions to be complex analytic. That concept was first explored by Cauchy almost a century later. The missing link to the concept of complex analytic functions is the definition of the derivative with respect to the complex argument

\[
F'(s) = \frac{dF(s)}{ds},
\] (4.1.0.5)

where \( s = \sigma + \omega j \), without which the complex analytic Taylor coefficients may not be defined.

Euler did not appreciate the role of complex analytic functions. They were first fully appreciated well after his death in 1785 by Augustin-Louis Cauchy (1789–1857) and then extended by Riemann in 1851.
4.2 Fundamental theorem of scalar calculus

History of scalar calculus: It some sense, the story of calculus begins with the fundamental theorem of calculus (FTC), also known generically as Leibniz’s formula, as shown in Table 4.1. The simplest integral is the length of a line \( L = \int_{0}^{x} dx \). If we label a point on a line as \( x = 0 \) and wish to measure the distance to any other point \( x \), we form the line integral between the two points. If the line is straight, this integral is the Euclidean length given by the difference between the two ends (Eq. 3.5.0.3, p. 94).

If \( F(\chi) \in \mathbb{R} \) describes a height above the line \( \chi \in \mathbb{R} \), then \( f(x) \),

\[
f(x) - f(0) = \int_{x=0}^{x} F(\chi) d\chi,
\]

may be viewed as the antiderivative of \( F(\chi) \). Here \( \chi \) is a dummy variable of integration. Thus the area under \( F(\chi) \) depends on only the difference in the area evaluated at the end points. It makes intuitive sense to view \( f(x) \) as the antiderivative of \( F(\chi) \).

This property of the area as an integral over an interval, depending on only the end points, has important consequences in physics in terms of conservation of energy, allowing for important generalizations. For example, as long as \( \chi \in \mathbb{R} \), we can let \( F(\chi) \in \mathbb{C} \) with no loss of generality, due to the linear Postulate P1 of the integral (see p. 121).

If \( f(x) \) is analytic (Eq. 3.2.2.8, p. 76), then

\[
F(x) = \frac{d}{dx} f(x)
\]

is an exact real differential. It follows that \( F(x) \) is analytic. This is known as the fundamental theorem of (real) calculus (FTC). Thus Eq. 4.2.0.2 may be viewed as an exact real differential. This is easily shown by evaluating

\[
\frac{d}{dx} f(x) = \lim_{\delta \to 0} \frac{f(x + \delta) - f(x)}{\delta} = F(x)
\]

starting from the antiderivative, Eq. 4.2.0.1. If \( f(x) \) is not analytic then the limit may not exist, so this is a necessary condition.

There are many important variations on this very basic theorem (see Table 4.1). For example, the limits could depend on time. Also when we take Fourier transforms, the integrand depends on both time \( t \in \mathbb{R} \) and frequency \( \omega \in \mathbb{R} \) via a complex exponential “kernel” function \( e^{\pm j \omega t} \in \mathbb{C} \), which is analytic in both \( t \) and \( \omega \).

4.2.1 The fundamental theorem of complex calculus

The fundamental theorem of complex calculus (FTCC) states (Greenberg, 1988, p. 1197) that for any complex analytic function \( F(s) \in \mathbb{C} \) with \( s = \sigma + j\omega \in \mathbb{C} \),

\[
f(s) - f(s_0) = \int_{s_0}^{s} F(\zeta) d\zeta,
\]

Equations 4.2.0.1 and 4.2.1.3 differ because the path of the integral is complex. Thus the line integral is over \( s \in \mathbb{C} \) rather than a real integral over \( \chi \in \mathbb{R} \). The FTCC states that the integral depends on only the end points, since

\[
F(s) = \frac{d}{ds} f(s).
\]

Comparing exact differentials, Eq. 4.1.0.5 (FTCC) and Eq. 4.2.0.2 (FTC), we see that \( f(s) \in \mathbb{C} \) must be complex analytic and have a Taylor series in powers in \( s \in \mathbb{C} \). It follows that \( F(s) \) is also complex analytic.

<table>
<thead>
<tr>
<th>Name</th>
<th>Mapping</th>
<th>p.</th>
<th>Description</th>
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</thead>
<tbody>
<tr>
<td>Leibniz (FTC)</td>
<td>( \mathbb{R}^n \to \mathbb{R}^n )</td>
<td>135</td>
<td>Area under a real curve</td>
</tr>
<tr>
<td>Cauchy (FTCC)</td>
<td>( \mathbb{C}^n \to \mathbb{R}^n )</td>
<td>135</td>
<td>Area under a complex curve</td>
</tr>
<tr>
<td>Cauchy’s theorem</td>
<td>( \mathbb{C}^n \to \mathbb{C}^n )</td>
<td>151</td>
<td>Close integral over analytic region is zero</td>
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<tr>
<td>Cauchy’s integral formula</td>
<td>( \mathbb{C}^n \to \mathbb{C}^n )</td>
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<tr>
<td>residue theorem</td>
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<td>( \mathbb{C}^n \to \mathbb{C}^n )</td>
<td></td>
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</tr>
</tbody>
</table>
Complex analytic functions: The definition of a complex analytic function \( F(s) \) of \( s \in \mathbb{C} \) is that the function may be expanded in a Taylor series (Eq. 3.2.4.20, p. 81) about an expansion point \( s_0 \in \mathbb{C} \). This definition follows the same logic as the FTC. Thus we need a definition for the coefficients \( c_n \in \mathbb{C} \), which most naturally follow from Taylor’s formula

\[
c_n = \frac{1}{n!} \frac{d^n}{ds^n} F(s) \bigg|_{s=s_0}. \tag{4.2.1.5}
\]

The requirement that \( F(s) \) have a Taylor series naturally follows by taking derivatives with respect to \( s \) at \( s_0 \). The problem is that both integration and differentiation of functions of complex Laplace frequency \( s = \sigma + \omega j \) have not yet been defined.

Thus the question is What does it mean to take the derivative of a function \( F(s) \in \mathbb{C} \), \( s = \sigma + \omega j \in \mathbb{C} \), with respect to \( s \), where \( s \) defines a plane rather than a real line? We learned how to form the derivative on the real line. Can the same derivative concept be extended to the complex plane?

The answer is affirmative. The question may be resolved by applying the rules of the real derivative when defining the derivative in the complex plane. However, for the complex case, there is an issue regarding direction. Given any analytic function \( F(s) \), the partial derivative with respect to \( \sigma \) different from the partial derivative with respect to \( \omega j \)? For complex analytic functions, the FTCC states that the integral is independent of the path in the \( s \) plane. Based on the chain rule, the derivative must also be independent of the direction at \( s_0 \). This directly follows from the FTCC. If the integral of a function of a complex variable is to be independent of the path, then the derivative of a function with respect to a complex variable must be independent of the direction. This follows from Taylor’s formula for the coefficients of the complex analytic formula (Eq. 4.2.1.5).

The Cauchy-Riemann conditions: The FTC defines the area as an integral over a real differential (\( ds \in \mathbb{R} \)), while the FTCC relates an integral over a complex function \( F(s) \in \mathbb{C} \) along a complex interval (i.e., path) \( (ds \in \mathbb{C}) \). For the FTC the area under the curve depends on only the end points of the antiderivative \( f(x) \). But what is the meaning of an “area” along a complex path? The Cauchy-Riemann conditions provide the answer.

4.2.2 Cauchy-Riemann conditions

For the integral of \( Z(s) = R(\sigma, \omega) + X(\sigma, \omega)j \) to be independent of the path, the derivative of \( Z(s) \) must also be independent of the path. As we show next, this requirement leads to a pair of equations known as the Cauchy-Riemann conditions.

To define

\[
\frac{d}{ds} Z(s) = \frac{d}{ds} [R(\sigma, \omega) + jX(\sigma, \omega)],
\]

we take partial derivatives of \( Z(s) \) with respect to \( \sigma \) and \( \omega j \), and equate them:

\[
\frac{\partial Z}{\partial \sigma} = \frac{\partial R}{\partial \sigma} + j \frac{\partial X}{\partial \sigma} \quad \text{and} \quad \frac{\partial Z}{\partial \omega} = \frac{\partial R}{\partial \omega} + j \frac{\partial X}{\partial \omega}.
\]

This says that a horizontal derivative, with respect to \( \sigma \), is equivalent to a vertical derivative, with respect to \( \omega j \). Taking the real and imaginary parts gives the two equations

\[
\text{CR-1: } \frac{\partial R(\sigma, \omega)}{\partial \sigma} = \frac{j}{r} \frac{\partial X(\sigma, \omega)}{\partial \omega}, \quad \text{and} \quad \text{CR-2: } \frac{\partial R(\sigma, \omega)}{\partial \omega} = -\frac{j}{r} \frac{\partial X(\sigma, \omega)}{\partial \sigma}, \tag{4.2.2.6}
\]

known as the Cauchy-Riemann (CR) conditions. The \( j \) cancels in CR-1 but introduces a \( j^2 = -1 \) in CR-2. They may also be written in polar coordinates \( (s = re^{\theta j}) \) as

\[
\frac{\partial R}{\partial r} = \frac{1}{r} \frac{\partial X}{\partial \theta} \quad \text{and} \quad \frac{\partial X}{\partial r} = -\frac{1}{r} \frac{\partial R}{\partial \theta}.
\]

The FTCC (Eq. 4.2.1.3) follows from CR-1 and CR-2 (Eq. 4.2.2.6).

If you are wondering what would happen if we took a derivative at 45°, then we only need to multiply the function by \( e^{\pi j/4} \). But doing so will not change the derivative. Thus we may take the derivative in any direction by multiplying by \( e^{\theta j} \), and the CR conditions will not change.

The CR conditions are necessary so that the integral of \( Z(s) \), and thus its derivative, is independent of the path, expressed in terms of conditions on the real and imaginary parts of \( Z \). This is a very strong condition on \( Z(s) \), which follows assuming that \( Z(s) \) may be written as a Taylor series in \( s \):

\[
Z(s) = Z_0 + Z_1 s + \frac{1}{2} Z_2 s^2 + \cdots, \tag{4.2.2.7}
\]
where \( Z_n \in \mathbb{C} \) are complex constants given by the Taylor series formula (Eq. 4.2.1.5). As with the real Taylor series, there is the convergence condition that \(|s| < 1\), called the radius of convergence (RoC). This is an important generalization of the region of convergence (RoC) for real \( s = x \).

Every function that may be expressed as a Taylor series in \( s - s_o \) about point \( s_o \in \mathbb{C} \) is said to be complex analytic at \( s_o \). This series, which must be single-valued, is said to converge within a radius of convergence (RoC). This highly restrictive condition has significant physical consequences. For example, every impedance function \( Z(s) \) obeys the CR conditions over large regions of the \( s \) plane, including the entire right half \( s \) plane (RHP) \((\sigma > 0)\). This condition is summarized by the Brune condition \( \Re\{Z(\sigma > 0)\} \geq 0 \), or alternatively \( \angle Z(s) < \angle s \).

When the CR conditions are generalized to volume integrals, this is called either Gauss’s Law or Green’s theorem, which is used in the solution of boundary value problems in engineering and physics (Kusse and Westwig, 2010).

We may merge these equations into a pair of second-order equations by taking a second round of partials. Specifically, eliminating the real part \( R(\sigma, \omega) \) of Eq. 4.2.2.6 gives

\[
\frac{\partial^2 R(\sigma, \omega)}{\partial \sigma \partial \omega} = \frac{\partial^2 X(\sigma, \omega)}{\partial^2 \omega} = -\frac{\partial^2 X(\sigma, \omega)}{\partial^2 \sigma},
\]

which may be written compactly as \( \nabla^2 X(\sigma, \omega) = 0 \). Eliminating the imaginary part gives

\[
\frac{\partial^2 X(\sigma, \omega)}{\partial \omega \partial \sigma} = \frac{\partial^2 R(\sigma, \omega)}{\partial^2 \sigma} = -\frac{\partial^2 R(\sigma, \omega)}{\partial^2 \omega},
\]

which may be written as \( \nabla^2 R(\sigma, \omega) = 0 \).

In summary, for a function \( Z(s) \) to be complex analytic, the derivative \( dZ/ds \) must be independent of direction (path), which requires that the real and imaginary parts of the function obey Laplace’s equation; that is,

\[
\text{CR-3: } \nabla^2 R(\sigma, \omega) = 0 \quad \text{and} \quad \text{CR-4: } \nabla^2 X(\sigma, \omega) = 0.
\]

The CR equations are easy to work with because they are first-order, but the intuition is best understood by noting two facts: (1) the derivative of a complex analytic function is independent of its direction, and (2) the real and imaginary parts of the function both obey Laplace’s equation. Such relationships are known as harmonic functions.

As we shall see in the next few sections, complex analytic functions must be smooth, since every analytic function may be differentiated an infinite number of times within the RoC. The magnitude must attain its maximum and minimum on the boundary. For example, when you stretch a rubber sheet over a jagged frame, the height of the rubber sheet obeys Laplace’s equation. Nowhere can the height of the sheet rise above or below its value at the boundary.

Harmonic functions define conservative fields, which means that energy (like a volume or area) is conserved. The work done in moving a mass from \( a \) to \( b \) in such a field is conserved. If you return the mass from \( b \) back to \( a \), the energy is retrieved and zero net work has been done.

---

1When the function is the ratio of two polynomials, as in the case of the Brune impedance, they are also related to Möbius transformations, also known as bi-harmonic operators.
4.3 Problems DE-1

Topics of this homework:
Complex numbers and functions (ordering and algebra), complex power series, fundamental theorem of calculus (real and complex); Cauchy-Riemann conditions, multivalued functions (branch cuts and Riemann sheets)

Complex Power Series

Problem # 1: In each case derive (e.g., using Taylor’s formula) the power series of \( w(s) \) about \( s = 0 \) and give the RoC of your series. If the power series doesn’t exist, state why! Hint: In some cases, you can derive the series by relating the function to another function for which you already know the power series at \( s = 0 \).

- 1.1: \( 1/(1 - s) \)
- 1.2: \( 1/(1 - s^2) \)
- 1.3: \( 1/(1 - s)^2 \)
- 1.4: \( 1/(1 + s^2) \). Hint: This series will be very ugly to derive if you try to take the derivatives \( \frac{d^n}{ds^n}[1/(1+s^2)] \). Use the results of questions 1.1 through 1.3 to represent this function as \( w(s) = -0.5i/(s - i) + 0.5i/(s + i) = \frac{i ((s-i)-(s+i))}{2i} \)...
- 1.5: \( 1/s \)
- 1.6: \( 1/(1 - |s|^2) \)

Problem # 2: Consider the function \( w(s) = 1/s \)

- 2.1: Expand this function as a power series about \( s = 1 \). Hint: Let \( 1/s = 1/(1 - 1 + s) = 1/(1 - (1 - s)) \).
- 2.2: What is the RoC?
- 2.3: Expand \( w(s) = 1/s \) as a power series in \( s^{-1} = 1/s \) about \( s^{-1} = 1 \).
- 2.4: What is the RoC?
- 2.5: What is the residue of the pole?

Problem # 3: Consider the function \( w(s) = 1/(2 - s) \)
4.3. PROBLEMS DE-1

– 3.1: Expand $w(s)$ as a power series in $s^{-1} = 1/s$. State the RoC as a condition on $|s^{-1}|$. Hint: Multiply top and bottom by $s^{-1}$.

– 3.2: Find the inverse function $s(w)$. Where are the poles and zeros of $s(w)$, and where is it analytic?

**Problem # 4: Summing the series**
The Taylor series of functions have more than one region of convergence.

– 4.1: If $a = 0.1$, what is the value of

$$x = 1 + a + a^2 + a^3 + \cdots?$$

Show your work.

– 4.2: If $a = 10$, what is the value of

$$x = 1 + a + a^2 + a^3 + \cdots?$$

**Quadratic forms**
A matrix that has positive eigenvalues is said to be positive-definite. The eigenvalues are real if the matrix is symmetric, so this is a necessary condition for the matrix to be positive-definite. This condition is related to conservation of energy, since the power is the voltage times the current. Given an impedance matrix

$$V = ZI,$$

the power $\mathcal{P}$ is

$$\mathcal{P} = I \cdot V = I \cdot ZI,$$

which must be positive-definite for the system to obey conservation of energy.

**Problem # 5: In this problem, consider the $2 \times 2$ impedance matrix**

$$Z = \begin{bmatrix} 2 & 1 \\ 1 & 4 \end{bmatrix}.$$

– 5.1: Solve for the power $\mathcal{P}(i_1, i_2)$ by multiplying out this matrix equation (which is in quadratic form) ($I \equiv \begin{bmatrix} i_1 & i_2 \end{bmatrix}^T$):

$$\mathcal{P}(i_1, i_2) = I^T \begin{bmatrix} 2 & 1 \\ 1 & 4 \end{bmatrix} I.$$

– 5.2: Is the impedance matrix positive-definite? Show your work by finding the eigenvalues of the matrix $Z$.

– 5.3: Should an impedance matrix always be positive-definite? Explain.

**Cauchy-Riemann Equations**

**Problem # 6: For this problem $i = \sqrt{-1}$, $s = \sigma + i\omega$, and $F(s) = u(\sigma, \omega) + iv(\sigma, \omega)$. According to the fundamental theorem of complex calculus (FTCC), the integration of a complex analytic function is independent of the path. It follows that the derivative of $F(s)$ is defined as**

$$\frac{dF}{ds} = \frac{d}{ds} [u(\sigma, \omega) + iv(\sigma, \omega)].$$

(DE-1.1)
If the integral is independent of the path, then the derivative must also be independent of the direction:
\[
\frac{dF}{ds} = \frac{\partial F}{\partial \sigma} = \frac{\partial F}{\partial \omega}.
\]  
(DE-1.2)

The Cauchy-Riemann (CR) conditions
\[
\frac{\partial u(\sigma, \omega)}{\partial \sigma} = \frac{\partial v(\sigma, \omega)}{\partial \omega} \quad \text{and} \quad \frac{\partial u(\sigma, \omega)}{\partial \omega} = -\frac{\partial v(\sigma, \omega)}{\partial \sigma}
\]
may be used to show where Equation DE-1.2 holds.

- 6.1: Assuming Equation DE-1.2 is true, use it to derive the CR equations.
- 6.2: Merge the CR equations to show that \( u \) and \( v \) obey Laplace’s equations
\[
\nabla^2 u(\sigma, \omega) = 0 \quad \text{and} \quad \nabla^2 v(\sigma, \omega) = 0.
\]

What can you conclude?

Problem # 7: Apply the CR equations to the following functions. State for which values of \( s = \sigma + i\omega \) the CR conditions do or do not hold (e.g., where the function \( F(s) \) is or is not analytic). Hint: Review where CR-1 and CR-2 hold.

- 7.1: \( F(s) = e^s \)
- 7.2: \( F(s) = 1/s \)

Branch cuts and Riemann sheets

Problem # 8: Consider the function \( w^2(z) = z \). This function can also be written as \( w(z) = \sqrt{z} \). Assume \( z = re^{\phi} \) and \( w(z) = \rho e^{\theta} \).

- 8.1: How many Riemann sheets do you need in the domain (\( z \)) and the range (\( w \)) to fully represent this function as single-valued?
- 8.2: Indicate (e.g., using a sketch) how the sheet(s) in the domain map to the sheet(s) in the range.
- 8.3: Use \( \text{zviz.m} \) to plot the positive and negative square roots \( \pm \sqrt{z} \). Describe what you see.
- 8.4: Where does \( \text{zviz.m} \) place the branch cut for this function?
- 8.5: Must the branch cut necessarily be in this location?

Problem # 9: Consider the function \( w(z) = \log(z) \). As in Problem 8, let \( z = re^{\phi} \) and \( w(z) = \rho e^{\theta} \).

- 9.1: Describe with a sketch and then discuss the branch cut for \( f(z) \).
- 9.2: What is the inverse of the function \( z(f) \)? Does this function have a branch cut? If so, where is it?
4.3. PROBLEMS DE-1

– 9.3: Using \( z \) \( \text{viz.} \), show that

\[
\tan^{-1}(z) = -\frac{j}{2} \log \frac{j - z}{j + z}. \tag{DE-1.3}
\]

In Fig. 4.1 (p. 134) these two functions are shown to be identical.

– 9.4: Algebraically justify Eq. DE-1.3. Hint: Let \( w(z) = \tan^{-1}(z) \) and \( z(w) = \tan w = \sin w / \cos w \); then solve for \( e^{\pi j} \).

A Cauer synthesis of any Brune impedance

**Problem # 10:** One may synthesize a transmission line (ladder network) from a positive real impedance \( Z(s) \) by using the continued fraction method. To obtain the series and shunt impedance values, we can use residue expansion. Here we shall explore this method.

– 10.1: Starting from the Brune impedance \( Z(s) = \frac{1}{s+1} \), find the impedance network as a ladder network.

– 10.2: Use a residue expansion to mimic the CFA floor function (Sec. 2.5.4, p. 47) for polynomial expansions. Find the residue expansion of \( H(s) = \frac{s^2}{s + 1} \) and express it as a ladder network.

– 10.3: Discuss how the series impedance \( Z(s, x) \) and shunt admittance \( Y(s, x) \) determine the wave velocity \( \kappa(s, x) \) and the characteristic impedance \( z_0(s, x) \) when (1) \( Z(s) \) and \( Y(s) \) are both independent of \( x \); (2) \( Y(s) \) is independent of \( x \), \( Z(s, x) \) depends on \( x \); (3) \( Z(s) \) is independent of \( x \), \( Y(s, x) \) depends on \( x \); and (4) both \( Y(s, x), Z(s, x) \) depend on \( x \).

This shows that a Cauer synthesis may be implemented with the residue expansion replacing the floor function in the CFA. This seems to solve Brune’s network synthesis problem.
4.4 Complex analytic Brune admittance

It is rarely stated that the variable that we are integrating over, either \(x\) (space) or \(t\) (time), is real \((x, t \in \mathbb{R})\), since that fact is implicit, due to the physical nature of the formulation of the integral. But this intuition must be refined once complex numbers are included with \(s \in \mathbb{C}\), where \(s = \sigma + \omega j\).

That time and space are real variables is more than an assumption: it is a requirement that follows from the order property. Real numbers have order. For example, if \(t = 0\) is now (the present), then \(t < 0\) is the past and \(t > 0\) is the future. Since time and space are real \((x, t \in \mathbb{R})\), they obey this order property. To have time travel, time and space would need to be complex (they are not), since if the space axis were complex the order property would be invalid.

Interestingly, it was shown by d’Alembert (1747) that time and space are related by the pure delay, due to the wave speed \(c_o\). To obtain a solution to the governing wave equation, which d’Alembert first proposed for sound waves, \(x, t \in \mathbb{R}\) may be functionally combined as

\[
\zeta_\pm = t \pm x/c_o,
\]

where \(c_o \in \mathbb{R} [\text{m/s}]\) is the wave phase velocity. The d’Alembert solution to the wave equation, describing waves on a string under tension is

\[
u(x, t) = f(t - x/c_o) + g(t + x/c_o),
\]

which describes the transverse velocity (or displacement) of two independent waves \(f(\zeta_-), g(\zeta_+) \in \mathbb{R}\) on the string, which represent forward and backward traveling waves.\(^2\) For example, starting with a string at rest, if one displaces the left end, at \(x = 0\), by a step function \(u(t)\), then that step displacement will propagate to the right as \(u(t - x/c_o)\), arriving at location \(x_o [\text{m}]\), at time \(x_o/c_o [\text{s}]\). Before this time, the string will not move to the right of the wave-front, at \(x_o [\text{m}]\), and after \(t_o [\text{s}]\) it will have a non-zero displacement. Since the wave equation obeys superposition (postulate P2, p. 122), it follows that the “plane-wave” eigenfunction of the wave equation for \(x, k \in \mathbb{R}^3\) are given by

\[
\psi_\pm (x, t) = \delta(t \mp k \cdot x) \leftrightarrow e^{st \mp \frac{k}{c_o}x},
\]

where \(|k| = 2\pi/|\lambda| = \omega/c_o\) is the wave number, \(|\lambda|\) is the wavelength, and \(s = \sigma + \omega j\), the Laplace frequency.

When propagation dispersion and losses are considered, we must replace the wave number \(k\) with a complex analytic vector wave number \(\kappa(s) = k_r(s) + jk_i(s)\), which is denoted as either the complex propagation function or the dispersion relation. The vector propagation function is a subtle and significant generalization of the scalar wave number \(k = 2\pi/\lambda\).

Forms of energy loss, which include viscosity and radiation, require \(\kappa(s) \in \mathbb{C}\). Physical examples include acoustic plane waves, electromagnetic wave propagation, antenna theory, and one of the most difficult cases, that of 3D electron wave propagating in crystals (e.g., silicon), where electrons and electro-magnetic (EM) waves are in a state of quantum mechanical equilibrium.

Even if we cannot solve these more difficult problems, we can still appreciate their qualitative solutions. One of the principles that allows us to do that is the causal nature of \(\kappa(s)\). Namely the \(L^T^{-1}\) of \(\kappa(s)\) must be causal, thus Eq. 4.4.0.2 must be causal. The group delay then describes the nature of the frequency dependent causal delay. For example, if the group delay is large at some frequency, then the solutions will have the largest causal delay at that frequency (Brillouin, 1953; Papoulis, 1962). Qualitatively this gives a deep insight into the solutions, even when we cannot compute the exact solution.

Electrons and photons are simply different EM states, where \(\kappa(x, s)\) describes the crystal’s dispersion relations as functions of both frequency and direction, famously known as Brillouin zones. Dispersion is a property of the medium such that the wave velocity is a function of frequency and direction, as in silicon.\(^3\) Highly readable discussions on the history of this topic may be found in Brillouin (1953).

4.4.1 Generalized admittance/impedance

The most elementary examples of Brune admittance and impedance are those made up of resistors, capacitors, and inductors. Such discrete element circuits arise not only in electrical networks but in mechanical, acoustical, and thermal networks as well (Table 3.2, p. 114). These lumped-element networks can always be represented by ratios of polynomials. This gives them a similar structure, with easily classified properties. Such circuits are called Brune admittances (or impedances).\(^4\) An example of a special structure is when the degrees of the numerator and

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\(^2\) d’Alembert’s solution is valid for functions that are not differentiable, such as \(\delta(t - c_o x)\).

\(^3\) In case you missed it, what I’m suggesting is that photons (propagating waves) and electrons (evanescent waves) are different EM wave “states” (Jaynes, 1991). This difference depends on the medium, which determines the dispersion relation (Papapamikis et al., 2018).

\(^4\) Some texts prefer the term immittance to include both admittance and impedance.
denominator polynomials cannot differ by more than one. This restriction on the degrees comes about because the real part of the admittance/impedance must be positive.

But there is a much broader class of admittances that come from transmission lines and other physical structures, which we refer to as generalized admittances. An interesting example is an admittance of the form \( 1/\sqrt{s} \), called a semicapacitor, or \( \sqrt{s} \), called a seminductor. Generalized admittance/impedance is not the ratio of two polynomials. As a result, they are more difficult to characterize.

When a generalized admittance \( Y(s) \) or its impedance \( Z(s) = 1/Y(s) \) is transformed into the time domain, it must have a real and positive surge admittance \( \gamma_r \in \mathbb{R} \) or surge impedance \( z_r \in \mathbb{R} \), followed by the residual response \( v(t), \zeta(t) \).

We define the following notation for the admittance,

\[
Y(s) = \gamma_r + Y(s) \leftrightarrow y(t) = \gamma_r \delta(t) + v(t),
\]

and the impedance,

\[
Z(s) = z_r + z_s \leftrightarrow z(t) = z_r \delta(t) + \zeta(t).
\]

The complexity of the notation is necessary and follows from the fact that \( z(t) \leftrightarrow Z(s) \) and \( y(t) \leftrightarrow Y(s) \) are positive-real and thus minimum phase.

When we are dealing with a transmission line (i.e., wave guides), the generalized admittance is defined as the ratio of a reflected wave to the incident wave. For an electrical system (voltage \( \Phi \), current \( I \)), the input admittance looking to the right from location \( x \) is

\[
Y_{in}^+(x > 0, s) = \frac{I^+(x, \omega)}{\Phi^+(x, \omega)} \quad (4.4.1.4)
\]

and looking to the left is

\[
Y_{in}^-(x < 0, s) = \frac{I^-(x, \omega)}{\Phi^-(x, \omega)} \quad (4.4.1.5)
\]

These two admittances are typically different.

**Generalized reflectance:** A function related to the generalized impedance is the reflectance \( \Gamma(s) \), defined as the ratio of a reflected wave to the incident wave. For the case of acoustics (pressure \( P \), volume velocity \( V' \)),

\[
Y_{in}(x, s) = \frac{P'(s)}{P(s)} = \frac{\gamma_r^+ - \gamma_r^-}{\gamma_r^+ - \gamma_r^-}
\]

\[
= \frac{\gamma_r^+ - \gamma_r^-}{1 + \frac{\gamma_r^-}{\gamma_r^+}}
\]

\[
= \frac{\gamma_r^+ - \Gamma(x, s)}{1 + \Gamma(x, s)} \quad (4.4.1.6)
\]

When the physical system is continuous at the measurement point \( x \), \( \gamma_r^+ (x) = \gamma_r^- (x) \in \mathbb{R} \). The reflectance \( \Gamma(x, s) \) depends on the area function, the boundary conditions, or both.

There is a direct relationship among a transmission line’s area function \( A(x) \in \mathbb{R} \), its characteristic impedance \( \gamma_r(x) \in \mathbb{R} \), and its eigenfunctions. We shall provide specific examples as they arise in our analysis of transmission lines (e.g., Fig. 5.3, p. 184).

A few papers that deal with the relationship between \( Y_{in}(s) \) and the area function \( A(x) \) are Youla (1964); Sondhi and Gopinath (1971); Rasetshwane et al. (2012). However, the general theory of this important and interesting problem is beyond the scope of this text. (See Homework 3.3, Problem # 2.)

**Complex analytic \( \Gamma(s) \) and \( Y_{in}(s) = Z_{in}^{-1}(s) \)**

When we define the complex reflectance \( \Gamma(s) \), we make a key assumption: Even though \( \Gamma(s) \) is defined by the ratio of two functions of real (radian) frequency \( \omega \), like the impedance, the reflectance must be causal (Postulate P1, p. 121). That \( \gamma(t) \to \Gamma(s) \) and \( \zeta(t) \leftrightarrow Z_{in}(s) = 1/Y_{in}(s) \) are causal is required by the physics.

### 4.4.2 Complex analytic impedance

Conservation of energy (or power) is a cornerstone of modern physics. It may first have been under consideration by Galileo Galilei (1564–1642) and Marin Mersenne (1588–1648). Today the question is not whether it is true, but why. Specifically, what is the physics behind conservation of energy? Surprisingly, the answer is straightforward,
based on its definition and the properties of impedance. Recall that the power is the product of the force and the flow, and impedance is their ratio.

The power is given by the product of two variables, sometimes called conjugate variables, the force and the flow. In electrical terms, these are voltage (force) \( v(t) \leftrightarrow V(\omega) \) and current (flow) \( i(t) \leftrightarrow I(\omega) \); thus, the electrical power at any instant of time is

\[
\mathcal{P}(t) = v(t)i(t).
\]  

(4.4.2.8)

The total energy \( E(t) \) is the integral of the power, since \( \mathcal{P}(t) = dE/dt \). Thus if we start with all the elements at rest (no currents or voltages), then the energy as a function of time is always positive

\[
E(t) = \int_{0}^{t} \mathcal{P}(t) dt \geq 0
\]

(4.4.2.9)

and is simply the total energy applied to the network (Van Valkenburg, 1964a, p. 376). Since the voltage and current are related by either an impedance or an admittance, conservation of energy depends on the property of impedance. From Ohm’s law and Postulate P1 (every impedance is causal), and we have

\[
v(t) = z(t) \ast i(t) = \int_{\tau=0}^{t} z(\tau)i(t - \tau) d\tau \leftrightarrow V(s) = Z(s)I(s).
\]

Example: Let \( i(t) = \delta(t) \). Then \(|w|^2(\tau) = i(t) \ast i(t) = \delta(\tau)\). Thus

\[
I_{xx}(t) = \int_{\tau=0}^{t} z(\tau)|w|^2(\tau) d\tau = \int_{\tau=0}^{t} z(\tau)\delta(\tau) d\tau = \int_{0}^{t} z(\tau) d\tau.
\]

The Brune impedance always has the form \( z(t) = r_{o}\delta(t) + \zeta(t) \). The characteristic impedance (also called surge impedance) may be defined as (Lundberg et al., 2007)

\[
r_{o} = \int_{0}^{\infty} z(t) dt.
\]

This definition requires that the integral of \( \zeta(t) \) is zero, a suspect conclusion that needs further investigation.

Note that convolution by an admittance or impedance linearizes the expression for the power.

These ideas are perhaps easier to visualize when we work in the Laplace frequency domain, where the total energy, equal to the integral of the real part of the power, is

\[
\frac{1}{s} \Re VI = \frac{1}{2s} (V' I + VI') = \frac{1}{2s} (Z' I' I + Z I I') = \frac{1}{2s} \Re Z(s)|I|^2 \geq 0.
\]

Mathematically this is called a positive definite operator, since the positive and real resistance is sandwiched between the current, the definiteness.

In conclusion, conservation of energy is totally dependent on the properties of the impedance. Thus one of the most important and obvious applications of complex functions of a complex variable is the impedance function. This seems to be the ultimate example of the FTCC applied to \( z(t) \).

Every impedance must obey conservation of energy (Postulate P3): The impedance function \( Z(s) \) has resistance \( R \) and reactance \( X \) as a function of complex frequency \( s = \sigma + j\omega \). From the causality postulate (P1) (p. 121), \( z(t < 0) = 0 \). Every impedance is defined by a Laplace transform pair

\[
z(t) \leftrightarrow Z(s) = R(\sigma, \omega) + jX(\sigma, \omega),
\]

with \( R, X \in \mathbb{R} \).

According to Postulate P3 (p. 121), a system is passive if it does not contain a power source. Drawing power from an impedance violates conservation of energy. This property is also called positive-real, which was defined by Brune (1931b,a)

\[
\Re\{Z(s \geq 0)\} \geq 0.
\]

(4.4.2.10)

Positive-real systems cannot draw more power than is stored in the impedance.\(^5\) The region \( \sigma \leq 0 \) is called the left half \( s \) plane (LHP), and the complementary region \( \sigma > 0 \) is called the right half \( s \) plane (RHP). According to the Brune condition, the real part of every impedance must be nonnegative (the RHP).

\(^5\) Does this condition hold for the LHP \( \sigma < 0 ? \) It does for Eq. 4.4.2.13.
4.4. COMPLEX ANALYTIC BRUNE ADMITTANCE

It is easy to construct examples of second-order poles or zeros in the RHP such that Postulate P3 is violated. Thus Postulate P3 implies that the impedance may not have more than simple (first-order) poles and zeros, strictly in the LHP. But there is yet more: These poles and zeros in the LHP must have order to meet the minimum phase condition. This minimum phase condition is easily stated:

\[ \angle Z(s) < \angle s, \]

but difficult to prove.\(^6\) There seems to be no proof that second-order poles and zeros (e.g., second-order roots) are not allowed. However, such roots must violate a requirement that the poles and zeros must alternate on the \( \sigma = 0 \) axis, which follows from Postulate P3. In the complex plane the concept of “alternate” is not defined (complex numbers cannot be ordered). What has been proved (i.e., Foster’s reactance theorem) is that if the poles are on the real or imaginary axis, they must alternate, leading to simple poles and zeros (Van Valkenburg, 1964a). The restriction on poles is sufficient but not necessary, as \( Z(s) = 1/\sqrt{s} \) is a positive-real impedance but is less than a first-degree pole (Kim and Allen, 2013). The corresponding condition in the LHP, and its proof, remains elusive (Van Valkenburg, 1964a).

For example, a series resistor \( R_o \) and capacitor \( C_o \) have an impedance given by (Table C.3, p. 242)

\[ Z(s) = R_o + 1/sC_o \leftrightarrow R_o \delta(t) + \frac{1}{C_o} u(t) = z(t), \]  

(4.4.2.12)

with constants \( R_o, C_o \in \mathbb{R} > 0 \). In mechanics, an impedance composed of a dashpot (damper) and a spring has the same form. A resonant system has an inductor, resistor, and capacitor, with an impedance given by

\[ Z(s) = \frac{sC_o}{1 + s^2 R_o + s^2 C_o M_o} \leftrightarrow C_o \frac{d}{dt} (c_+ e^{s_+ t} + c_- e^{s_- t}) = z(t), \]

(4.4.2.13)

which is a second-degree polynomial with two complex resonant frequencies \( s_{\pm} \). When \( R_o > 0 \), these roots are in the LHP, with \( z(t) \leftrightarrow Z(s) \).

Systems (networks) that contain many elements and transmission lines can be much more complicated, yet still have a simple frequency-domain representation. This is the key to understanding how these physical systems work, as we describe next.

Poles and zeros of positive-real functions must be first-degree: The definition of positive-real (PR) functions requires that the poles and zeros of the impedance function be simple (only first-degree). Second-degree poles would have a reactive “secular” response of the form \( h(t) = t \sin(\omega t + \phi) u(t) \), and these terms would not average to zero, depending on the phase, as is required of an impedance. As a result, only single-degree poles are possible.\(^7\) Furthermore, when the impedance is the ratio of two polynomials, where the lower-degree polynomial is the derivative of the higher-degree one, the poles and zeros must alternate. This is a well-known property of the Brune impedance that has never been adequately explained except for very special cases, denoted as Foster’s theorem (Van Valkenburg, 1964b, pp. 104–107). I believe that no one has ever reported an impedance that has second-degree poles and zeros. That would be a rare impedance. Network analysis books never report second-degree poles and zeros in their impedance functions. Nor has there ever been any guidance about where the poles and zeros might lie in the LHP. Understanding the exact relationships between pairs of poles and zeros, to assure that the real part of the impedance is real, would resolve this longstanding unsolved problem (Van Valkenburg, 1964a).

Calculus on Complex analytic functions: To solve a differential equation or integrate a function, Newton used the Taylor series to integrate one term at a time. However, he used only real functions of a real variable due to the fundamental lack of appreciation of the complex analytic function. This same method is how one finds solutions to scalar differential equations today, but using an approach that makes the solution method less obvious. Rather than working directly with the Taylor series, today we use the complex exponential, since the complex exponential is an eigenfunction of the derivative

\[ \frac{d}{dt} e^{st} = se^{st}. \]

Since \( e^{st} \) may be expressed as a Taylor series that has coefficients \( c_n = 1/n! \), in some real sense the modern approach is a compact way of doing what Newton did. Thus every linear constant coefficient differential equation in time may be simply transformed into a polynomial in complex Laplace frequency \( s \), by looking for solutions

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\(^6\)As best I know, this is an open problem in network theory (Brune, 1931a; Van Valkenburg, 1964a).

\(^7\)Secular terms result from second-degree poles, since \( u(t) \ast u(t) = tu(t) \leftrightarrow 1/s^2 \).
of the form \( A(s)e^{st} \) and transforming the differential equation into a polynomial \( A(s) \) in complex frequency. For example,

\[
\frac{d}{dt} f(t) + \alpha f(t) \leftrightarrow (s + \alpha)F(s).
\]

The root of \( A(s) = s + \alpha = 0 \) is the eigenvalue of the differential equation. A powerful tool for understanding the solutions of differential equations, both scalar and vector, is to work in the Laplace frequency domain. The Taylor series has been replaced by \( e^{st} \), which transforms Newton’s real Taylor series into the complex exponential eigenfunction. In some sense, these are the same methods, since

\[
e^{st} = \sum_{n=0}^{\infty} \frac{(st)^n}{n!}.
\]

(4.4.2.14)

Taking the derivative with respect to time gives

\[
\frac{d}{dt} e^{st} = se^{st} = s \sum_{n=0}^{\infty} \frac{(st)^n}{n!},
\]

(4.4.2.15)

which is also complex analytic. Thus if the series for \( F(s) \) is valid (i.e., it converges), then its derivative is also valid. This was a very powerful concept, exploited by Newton for real functions of a real variable, and later by Cauchy and Riemann for complex functions of a complex variable. The key question here is Where does the series fail to converge? This is the main message behind the FTCC (Eq. 4.2.1.3).

The FTCC (Eq. 4.2.1.3) is formally the same as the FTC (Eq. 4.2.0.2) (Leibniz’s formula), the key (and significant) difference being that the argument of the integrand \( s \in \mathbb{C} \). Thus this integration is a line integral in the complex plane. One would naturally assume that the value of the integral depends on the path of integration. And it does, but in a subtle way, as quantified by Cauchy’s various theorems. If the path stays in the same RoC region, then the integral is independent of that path. If a path includes a different pole, then the integral depends on the path, as quantified by the Cauchy residue theorem. The test is to deform the path from the first to the second. If in that deformation the path crosses a pole, then the integral will depend on the path. All of this is dependent on the causal nature of the integral.

The FTC and FTCC are clearly distinguishable yet the reasoning is the same. If \( F(s) = df(s)/ds \) is complex analytic [i.e., has a power series \( f(s) = \sum_k c_k s^k \), with \( f(s), c_k, s \in \mathbb{C} \)], then it may be integrated, and yet the integral does not depend on the path. At first blush, this is sort of amazing. The key is that \( F(s) \) and \( f(s) \) must be complex analytic, which means they are differentiable. This all follows from the Taylor series formula Eq. 4.2.1.5 (p. 136) for the coefficients of the complex analytic series. For Eq. 4.2.1.3 to hold, the derivatives must be independent of the direction (independent of the path), as discussed on page 136. The concept of a complex analytic function therefore has eminent consequences in the form of several key theorems on complex integration discovered by Cauchy in about 1820.

The complex Taylor series generalizes the functions it describes, with unpredictable consequences, as nicely shown by the domain-coloring diagrams in Fig. 3.12 (p. 125). Cauchy’s tools of complex integration were first exploited in physics by Sommerfeld (1952) to explain the onset (e.g., causal) transients in waves, as explained in detail in Brillouin (1960, Chap. 3).

Up to 1910, when Sommerfeld first published his results using complex analytic signals and saddle point integration in the complex plane, the implications of the causal wave-front were poorly understood. It would be reasonable to say that his insights changed our understanding of wave propagation for both light and sound. Unfortunately this insight has never been fully appreciated, even to this day. If you question this review, please read Brillouin (1960, Chap. 1).

The full power of the complex analytic function was first appreciated by Bernard Riemann (1826–1866) in his University of Göttingen PhD thesis of 1851, under the tutelage of Gauss (1777–1855), which drew heavily on the work of Cauchy.
The key definition of a complex analytic function is that it has a Taylor series representation over a region of the complex frequency plane $s = \sigma + j\omega$ that converges in a region of convergence (RoC) about the expansion point, with a radius determined by the nearest pole of the function. A further surprising feature of all analytic functions is that within the RoC, the inverse of that function also has a complex analytic expansion. Thus given $w(s)$, we can determine $s(w)$ to any desired accuracy, critically depending on the RoC. Given the right software (e.g., zviz.m), this relationship may be made precise.

### 4.4.3 Multivalued functions

In the field of mathematics there seems to have been a tug-of-war regarding the basic definition of a function. The accepted definition today is a single-valued (i.e., complex analytic) mapping from the domain to the codomain (or range). This makes the discussion of multivalued functions somewhat awkward. In 1851 Riemann (working with Gauss) resolved this problem for the complex analytic set of multivalued functions by introducing the geometric concept of single-valued sheets, delineated by branch cuts.

Two simple yet important examples of multivalued functions are the circle $z^2 = x^2 + y^2$ and $w = \log(z)$. For example, if we assume $z$ is the radius of the circle, then solving for $y(x)$ gives the double-valued function

$$y(x) = \pm \sqrt{x^2 - x^2},$$

The vertical axis, as a function of the angle and radius of $x \in \mathbb{C}$.

If we accept the modern definition of a function as the mapping from one set to a second set, then $y(x)$ is not a function, or even two functions. For example, what if $x > z$? Or worse, what if $z = 2i$ with $|x| < 1$? Riemann’s construction, using branch cuts for multivalued function, resolves all these difficulties (as best I know).

To proceed, we need definitions and classifications of the various types of complex singularities:

1. Poles of degree 1 are called simple poles. Their amplitude is called the residue (e.g., $\alpha/s$ has residue $\alpha$). Simple poles are special (see Eq. 4.5.1.3, p. 152); they play a key role in mathematical physics, since their inverse Laplace transform defines a causal eigenfunction.

2. When the numerator and denominator of a rational function (i.e., ratio of two polynomials) have a common root (i.e., factor), that root is said to be removable.

3. A singularity that is not removable, a pole, or a branch point is called essential.

4. A complex analytic function (except for isolated poles) is called meromorphic (Boas, 1987). Meromorphic functions can have any number of poles, even an infinite number. The poles need not be simple.

5. When the first derivative of a function $Z(s)$ has a simple pole at $a$, then $a$ is said to be a branch point of $Z(s)$. An important example is the logarithmic derivative:

$$\frac{d \ln(s - a)\alpha}{ds} = \frac{\alpha}{s - a}, \quad \alpha \in \mathbb{I}.$$ 

However, the converse does not necessarily hold.

6. I am not clear about the interesting case of an irrational pole ($\alpha \in \mathbb{I}$). In some cases (e.g., $\alpha \in \mathbb{F}$) this may be simplified with the logarithmic derivative operation, as mentioned on page 69.

More complex topologies are being researched today, and progress is expected to accelerate due to modern computing technology. It is helpful to identify the physical meaning of these more complex surfaces, to guide us in their interpretation and possible applications.

**Branch cuts:** Up to this point we have considered only poles of degree $\alpha \in \mathbb{N}$ of the form $1/s^n$. The concept of a branch cut allows us to manipulate (and visualize) multivalued functions for which $\alpha \in \mathbb{F}$. This is done by breaking each region into single-valued sheets, as shown in Fig. 4.3 (right). The branch cut is a curve $c \in \mathbb{C}$ that separates the various single-valued sheets of a multivalued function. The concepts of branch cuts, sheets, and the extended plane were first devised by Riemann, as described in his thesis of 1851. It was these three mathematical and geometrical constructions that provided deep insight into complex analytic functions, greatly extending the

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8https://www.maths.ox.ac.uk/about-us/departmental-art/theory

important earlier work of Cauchy (1789–1857) on the calculus of complex analytic functions. For an alternative helpful discussion of Riemann sheets and branch cuts, see Boas (1987, pp. 221–25) and Kusse and Westwig (2010).

To study the properties of multivalued functions and branch cuts, we look at \( w(s) = \sqrt{s} \) and \( w(s) = \log(s) \) along with their inverse functions \( w(s) = s^2 \) and \( w(s) = e^s \). For uniformity we refer to the complex abscissa \( s = \sigma + jo \) and the complex ordinate \( w(s) = u + vjo \). When the complex domain and range are swapped, by taking the inverse of a function, multivalued functions are a common consequence. For example, \( f(t) = \sin(t) \) is single-valued and analytic in \( t \) and thus has a Taylor series. The inverse function \( t(f) \) is multivalued.

The best way to explore the complex mapping from the complex planes \( s \rightarrow w(s) \) is to master the single-valued function \( s = w^2(s) \) and its double-valued inverse \( w(s) = \sqrt{s} \).

Figure 4.3 shows the single-valued function \( w(s) = s^2 \) (left), and its inverse (right), the double-valued mapping of \( s(w) = \pm \sqrt{w} \). Single-valued functions such as \( W(s) = s^2 \) are relatively straightforward. Multivalued functions require the concept of a branch cut, defined in the image \((w(s))\) plane, which is a technique to render the multiple values as single-valued on each of several sheets, defined by the sheet index in the domain \((s)\) plane and delineated by a branch cut (in the \((w)\) plane).

The sheets are labeled in the domain \((s)\) plane by a sheet index \( k \in \mathbb{Z} \), while branch points and cuts are defined in the image \((w)\) plane (also called the range). It is important to understand that the path of every branch cut is not unique and may be moved. However, branch points are unique and thus not movable.

The multivalued nature of \( w(s) = \sqrt{s} \) is best understood by working with the function in polar coordinates. We let
\[
s_k = r e^{\theta_j} e^{2\pi k j},
\]
where \( r = |s|, \theta = \angle s, \in \mathbb{R} \), and \( k \in \mathbb{Z} \) is the sheet index.

This concept of analytic inverses becomes important only when the function is multivalued. For example, since \( w(s) = s^2 \) has a period of 2, \( s(w) = \pm \sqrt{w} \) is multivalued. Riemann dealt with such extensions using the concept of a branch cut with multiple sheets labeled by sheet numbers. Each sheet describes an analytic function (Taylor series) that converges within some RoC that has a radius out to the nearest pole. Thus Riemann’s branch cuts and sheets explicitly deal with the need to define unique single-valued inverses of multivalued functions. Since the square root function has two overlapping regions corresponding to the \( \pm \) due to the radical, there must be two connected region—sort of like mathematical Siamese twins: distinct, yet the same.

Hue: By studying the output of \( \text{viz.m} \) (Fig. 3.12, p. 125), we can appreciate domain-coloring. The domain angles \( \angle s \) go from \(-90^\circ < \theta < 90^\circ \), with \( \theta = 0 \) being red and \( \pm 90^\circ \) being green (between yellow and purple). The angle to the hue map is shown in the left panel of Fig. 4.4. For \( w(s) = s^2 \), the \( \angle s \) is expanded by 2, since \( \psi = 2\theta \). For \( w(s) = \sqrt{s} \), the \( \angle s \) is compressed by a factor of 2, since \( \psi = \theta / 2 \). Thus the principal angle \( k = 0 \) \((-180^\circ < \theta < 180^\circ \) maps to half the \( w \) plane \((-90^\circ < \psi < 90^\circ \)) from purple to yellow, while the \( k = 1 \) branch maps to \( 90^\circ < \psi < 270^\circ \). Note how the panel on the right of Fig. 4.3 matches the right half of \( s \) (purple = \( 90^\circ \), yellow/green = \( 90^\circ \)) while the middle panel above comes from the left side of \( s \) (green to purple). The center panel is green at \(-180^\circ \) and purple at \( +180^\circ \), which matches the left panel at \( \pm 180^\circ \), respectively (i.e., \( e^{\pi j} \sqrt{s} \)).

Furthermore we can let
\[
w = \rho e^{\psi j} = \sqrt{r} e^{\theta / 2} e^{\pi \tau k},
\]
where \( \rho = |w|, \psi \angle w, \in \mathbb{R} \). The generic Cartesian coordinates are \( s = \sigma + jw \) and \( w(s) = u(\sigma, \omega) + v(\sigma, \omega)j \). For single-valued functions such as \( w(s) = s^2 \) on the left in Fig. 4.3 there is no branch cut, since \( \psi = 2\theta \). Note how the red color (\( \theta = 0^\circ \)) appears twice in this mapping. JBA Read this carefully for content! For multivalued functions, a branch cut is required, typically along the negative \( v(\sigma, \omega) \) axis (i.e., \( \psi = \pi \)), but it may be freely distorted, as seen by comparing the right panel of Fig. 4.3 with the right panel of Fig. 4.4.
The principal (i.e., first) Riemann sheet of $\sqrt{s}$, corresponding to $-\pi < \theta < \pi$ (i.e., $k = 0$), is shown in the center of Fig. 4.4. This differs from the neighboring sheet ($k = 1$), shown on the right. Thus $w(s) = \sqrt{s}$ is a multivalued function of $s$ that has two single-valued sheets.

**Moving the branch cut**: It is important to understand that the function is analytic on the branch cut but not at the branch point. One is free to move the branch cut (at will). It does not need to be on a line: it could be cut in almost any connected manner, such as a spiral. The only rule is that it must start and stop at the matching branch points, or at $\infty$, which must have the same degree.

Figure 4.3 (left) shows the single-values function $w(s) = s^2$, and Fig. 4.4 (right) shows its inverse, the double-valued mapping of $s(w) = \pm \sqrt{w}$.

The location of the branch cut may be moved by rotating the $s$ coordinate system of Fig. 4.2. For example, $w(s) = \pm \sqrt{s}$ and $w(s) = \pm \sqrt{-s}$ have different branch cuts, as may be easily verified using the Matlab/Octave commands $j*z*\text{zv}(s)$ and $z*\text{zv}(s)$, as shown in Fig. 4.4. Since the cut may be moved, every function is analytic on the branch cut. If a Taylor series is formed on the branch cut, it will describe the function on the two different sheets. Thus the complex analytic series (i.e., the Taylor formula, Eq. 4.2.1.5) does not depend on the location of a branch cut, as it only describes the function uniquely (as a single-valued function), valid in its local region of convergence.

The second sheet ($k = 1$) in Fig. 4.4 picks up at $\theta = \pi$ [rads] and continues on to $\pi + 2\pi = 3\pi$. The first sheet maps the angle of $w$ (i.e., $\phi = \angle w = \theta/2$) from $-\pi/2 < \phi < \pi/2$ ($w = e^{i\theta/2}$). This corresponds to $u = \Re\{w\} > 0$. The second sheet maps $\pi/2 < \psi < 3\pi/2$ (i.e., $90^\circ$ to $270^\circ$), which is $\Re\{w\} = u < 0$. In summary, twice around the $s$ plane is once around the $w(s)$ plane because the angle is half due to the $\sqrt{s}$.

Branch cuts emanate and terminate at *branch points*, defined as singularities (poles) that can even have fractional degree, as for example $1/\sqrt{s}$, and terminate at one of the matching roots, which includes the possibility of $\infty$. For example, suppose that in the neighborhood of the pole, at $s_o$, the function is

$$f(s) = \frac{w(s)}{(s - s_o)^k}, \quad (4.4.3.18)$$

where $w, s, s_o \in \mathbb{C}$ and $k \in \mathbb{Q}$. When $k = 1$, $s_o = \sigma_o + \omega_o j$ is a first-degree “simple pole,” having degree 1 in the $s$ plane, with residue $w(s_o)$.

Typically the order and degree are positive integers, but fractional degrees and orders are common in modern engineering applications (Kirchhoff, 1868; Lighthill, 1978). Here we allow both the degree and the order to be fractional ($\in \mathbb{F}$). When $k \in \mathbb{F} \subset \mathbb{R}$, $k = n/m$ is a real reduced fraction—namely, when $\gcd(n, m) = 1, n \perp m$. This defines the *degree* of a fractional pole. In such cases there must be two sets

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10This presumes that poles appear in pairs, one of which may be at $\infty$.

11We refer to the *order* of a derivative, or differential equation, and the *degree* of a polynomial, as commonly used in engineering applications.
of branch cuts of degrees \( n \) and \( m \). For example, if \( k = 1/2 \), the singularity (branch cut) is of degree 1/2 and there are two Riemann sheets, as shown in Fig. 4.3.

Figure 4.5: Colorized plots of two \( \mathcal{L}/\mathcal{T} \) pairs: Left: \( \sqrt{\pi/s} \leftrightarrow u(t)/\sqrt{t} \). Right: \( \sqrt{s^2 + 1} \leftrightarrow \delta(t) + \frac{1}{t}J_1(t)u(t) \).

**Fractional-order Bessel function:** An important example is the Bessel function and its Laplace transform (\( \mathcal{L}/\mathcal{T} \))

\[
\delta(t) + \frac{1}{t}J_1(t)u(t) \leftrightarrow \sqrt{s^2 + 1},
\]

as shown in Fig. 4.5, which is related to the solution to the wave equation in two-dimensional cylindrical coordinates (see Table C.4, p. 243). Bessel functions are the solutions (i.e., eigenfunctions) of guided acoustic waves in round pipes, or surface waves on the earth (seismic waves), or waves on the surface of a pond (Table 5.2, p. 190).

There are a limited number of possibilities for the degree, \( k \in \mathbb{Z} \) or \( \in \mathbb{F} \), of Eq. 4.4.3.18. If the degree is drawn from \( \mathbb{R} \not\in \mathbb{F} \), the pole can not have a residue. According to the definition of the residue, \( k \in \mathbb{F} \) has no residue. But there remains open the possibility of generalizing the concept of the Riemann integral theorem to include \( k \in \mathbb{F} \).

One way to do this is to use the logarithmic derivative, which transforms fractional poles to simple poles with fractional residues.

If the singularity has an irrational degree \( k \in \mathbb{I} \), the branch cut has the same irrational degree. Accordingly there are an infinite number of Riemann sheets, as in the case of the log function. An example is \( k = \pi \), for which

\[
F(s) = \frac{1}{s\pi} = e^{-\log(s^\pi)} = e^{-\pi \log(s)} = e^{-\pi \log(\rho)} e^{-\pi \theta},
\]

where the domain is expressed in polar coordinates \( s = \rho e^{i\theta} \). When \( k \in \mathbb{F} \), it may be close (e.g., \( k = \pi_{152}/\pi_{153} = \pi_{152}/(\pi_{152} + 2) = 881/883 \approx 0.99883 \), or its reciprocal \( \approx 1.0023 \)). The branch cut could be subtle (it could even go unnoticed), but it would have a significant impact on the function and on its inverse Laplace transform.

**Exercise #2**

Find the poles, zeros, and residues of \( F(s) \).

1.

\[
F(s) = d \left[ \ln \frac{s + e}{s + \pi} \right]
\]

Solution:

\[
F(s) = d \left[ \ln(s + e) - \ln(s + \pi) \right] = \left( \frac{1}{s + e} - \frac{1}{s + \pi} \right)
\]

The poles are at \( s_1 = -e \) and \( s_2 = -\pi \) with respective residues of \( \pm 1 \). 

2.

\[
F(s) = d \left[ \ln \frac{(s + 3)^\nu}{(s + j)^{-\pi}} \right]
\]
Solution: 
\[ F(s) = e^{\pi \ln s} \]

There is a very important take-home message here regarding the utility of the logarithmic derivative, which “linearizes” the fractional pole. ■

3.

\[ F(s) = e^{\pi \ln s} \]

Solution: We take the derivative 
\[ \frac{d}{ds} F(s) = \frac{d}{ds} \ln s = \pi \frac{d}{ds} \ln s = \pi s. \]

Thus we see that \( F'(s) \) has a pole at \( s = 0 \) with residue \( \pi \). It follows that \( F(s) = \int^s F'(s) ds \) has a second-order pole. Thus the residue must be zero. ■

4.

\[ F(s) = \pi^{-s} \]

Solution: We take the logarithmic derivative, 
\[ d \ln F(s)/ds = F'(s)/F(s) = -\ln \pi. \]

Thus \( F'(s) = -\ln \pi F(s) = -\ln \pi \pi^{-s} \). ■

Log function: Next we discuss the multivalued nature of the log function. In this case there are an infinite number of Riemann sheets, not well captured by Fig. 3.14 (p. 126), which displays only the principal sheet. However, if we look at the formula for the log function, the nature is easily discerned. The abscissa \( s \) may be defined as multivalued, since 
\[ s_k = re^{2\pi kj}e^{\theta j}. \]

Here we have extended the angle of \( s \) by \( 2\pi k \), where \( k \) is the sheet index \( \in \mathbb{Z} \). Now we take the log:
\[ \log(s) = \log(r) + (\theta + 2\pi k)j. \]

When \( k = 0 \), we have the principal value sheet, which is zero when \( s = 1 \). For any other value of \( k \), \( w(s) \neq 0 \), even when \( r = 1 \), since the angle is not zero, except for the \( k = 0 \) sheet.

4.5 Three Cauchy integral theorems

4.5.1 Cauchy’s theorems for integration in the complex plane

There are three basic definitions related to Cauchy’s integral formula. They are closely related and can greatly simplify integration in the complex plane. The choice of names is unfortunate, if not totally confusing.

1. Cauchy’s (integral) theorem (CT-1):
\[ \oint_C F(s) ds = 0 \quad (4.5.1.1) \]

if and only if \( F(s) \) is complex analytic inside of a simple closed curve \( C \) (Stillwell, 2010; Boas, 1987, p. 45). The FTCC (Eq. 4.2.1.3) says that the integral depends on only the end points if \( F(s) \) is complex analytic. With the path (contour \( C \)) closed, the end points are the same and thus the integral must be zero as long as \( F(s) \) is complex analytic.

2. Cauchy’s integral formula (CT-2):
\[ \frac{1}{2\pi j} \oint_{B} \frac{F(s)}{s - s_o} ds = \begin{cases} F(s_o), & s_o \in \mathbb{C} < B \text{ (inside)} \\ 0, & s_o \in \mathbb{C} > B \text{ (outside).} \end{cases} \quad (4.5.1.2) \]

Here \( F(s) \) is required to be analytic everywhere within (and on) the boundary \( B \) of integration (Greenberg, 1988, p. 1200); (Boas, 1987, p. 51); (Stillwell, 2010, p. 220). When the point \( s_o \in \mathbb{C} \) is within the boundary, the value \( F(s_o) \in \mathbb{C} \) is the residue of the pole \( s_o \) of \( F(s)/(s - s_o) \). When the point \( s_o \) lies outside the boundary, the integral is zero.
3. **The (Cauchy) residue theorem (CT-3):** (Greenberg, 1988, p. 1241); (Boas, 1987, p. 73)

\[
\int_{C} f(s) ds = 2\pi j \sum_{k=1}^{K} c_k = \sum_{k=1}^{K} \int_{C} \frac{F(s)}{s - s_k} ds,
\]

(4.5.1.3)

where the residues \( c_k \in \mathbb{C} \) correspond to the \( k \)th pole of \( f(s) \) enclosed by the contour \( C \) ((Greenberg, 1988, p. 1241); (Boas, 1987, p. 73)). Cauchy’s integral formula (CT-1) is equivalent to the rightmost form of the residue theorem CT-3.\(^\text{12}\)

**How to calculate the residue:** The case of first-degree poles has special significance because the Brune impedance allows only simple poles and zeros, thus increasing its utility. The residues for simple poles are

\[ c_k = \lim_{s \to s_k} [(s - s_k)F(s)] \]

(Greenberg, 1988; Boas, 1987, p. 72).

When the pole is an \( N \)th degree, the procedure is much more complicated and requires taking \( N - 1 \) order derivatives of \( f(s) \) followed by the limit process (Greenberg, 1988, p. 1242). Higher-degree poles are rarely encountered; thus it is good to know that this formula exists, but perhaps it is not worth the effort to learn (i.e., memorize) it.

### 4.5.2 Cauchy Integral Formula and Residue Theorem

CT-2 (Eq. 4.5.1.2) is an important extension of CT-1 (Eq. 4.5.1.1) in that a pole has been explicitly injected into the integrand at \( s = s_k \). If the pole location is outside the curve \( C \), the result of the integral is zero, in keeping with CT-1. When the pole is inside \( C \), the integrand is no longer complex analytic at the enclosed pole. When this pole is simple, the residue theorem applies. By a manipulation of the take the limit, the radius may be taken to zero in the limit, isolating the pole.

For the related CT-3 (Eq. 4.5.1.3) the same result holds, except it is assumed that there are \( K \) simple poles in the function \( F(s) \). This requires the repeated application of CT-2, \( K \) times, so it represents a minor extension of CT-2. The function \( F(s) \) may be written as \( f(s)/P_K(s) \), where \( f(s) \) is analytic in \( C \) and \( P_K(s) \) is a polynomial of degree \( K \), with all of its roots \( s_k \in \mathbb{C} \).

**Nonintegral degree singularities:** The key point is that this theorem applies when \( n \in \mathbb{R} \), including fractionals \( n \in \mathbb{F} \), but for these cases the residue is always zero, since by definition the residue is the amplitude of the \( 1/s \) term (Boas, 1987, p. 73). Here are some examples:

1. When \( n = 2/3 \) (i.e., \( n = 2/3 \)), the residue of \( s^n \) is zero, by definition.
2. The function \( 1/\sqrt{s} \) has a zero residue (we apply the definition of the residue, Eq. 4.5.1.4).
3. When \( n = 1 \), the residue is, by definition, zero.
4. When \( n \neq 1 \), the residue is given by Eq. 4.5.1.4.
5. CT-1, CT-2, and CT-3 are essential when when we compute the inverse Laplace transform.

**Summary and examples:** These three CT theorems, all attributed to Cauchy, collectively are related to the fundamental theorems of calculus. The general principles are:

1. In general it makes no sense (nor is there any need) to integrate through a pole; thus the poles (or other singularities) must not lie on \( C \).
2. CT-1 (Eq. 4.5.1.1) follows trivially from the fundamental theorem of complex calculus (Eq. 4.2.1.3, p. 135), since if the integral is independent of the path and the path returns to the starting point, the closed integral must be zero. Thus Eq. 4.5.1.1 holds when \( F(s) \) is complex analytic within \( C \).

\(^{12}\)This theorem is the same as a 2D version of Stokes’s theorem (Boas, 1987).
3. Since the real and imaginary parts of every complex analytic function obey Laplace’s equation (Eq. 4.2.2.10, p. 137), it follows that every closed integral over a Laplace field—that is, one defined by Laplace’s equation—must be zero. In fact, this is the property of a conservative system, corresponding to many physical systems. If a closed box has fixed potentials on the walls, with any distribution whatsoever, and a point charge (i.e., an electron) is placed in the box, then a force equal to \( \vec{F} = q E \) is required to move that charge, and thus work is done. However, if the point is returned to its starting location, the net work done is zero.

4. Work is done in charging a capacitor, and energy is stored. However, when the capacitor is discharged, all of the energy is returned to the load.

5. Soap bubbles and rubber sheets on a wire frame obey Laplace’s equation.

6. These are all cases where the fields are Laplacian, thus closed line integrals must be zero. Laplacian fields are commonly seen because they are so basic.

7. We have presented the impedance as the primary example of a complex analytic function. Physically, every impedance has an associated stored energy, and every system having stored energy has an associated impedance. This impedance is usually defined in the frequency \( s \) domain, as a force over a flow (i.e., voltage over current). The power \( P(t) \) is defined as the force times the flow and the energy \( E(t) \) as the time integral of the power

\[
E(t) = \int_{-\infty}^{t} P(t) dt,
\]

which is similar to Eq. 4.2.0.1 (p. 135) [see Sec. 3.7.3, Eq. 3.7.4.9 (p. 113)]. In summary, impedance and power and energy are all fundamentally related.
4.6 Problems DE-2

Topics of this homework:
Integration of complex functions, Cauchy’s theorem, integral formula, residue theorem, power series, Riemann sheets and branch cuts, inverse Laplace transforms

Two fundamental theorems of calculus

**Fundamental Theorem of Calculus (Leibniz):**
According to the fundamental theorem of (real) calculus (FTC),

\[ f(x) = f(a) + \int_a^x F(\xi) d\xi, \quad (DE-2.1) \]

where \( x, a, \xi, F, f \in \mathbb{R} \). This is an indefinite integral (since the upper limit is unspecified). It follows that

\[ \frac{df(x)}{dx} = \frac{d}{dx} \int_a^x F(x) dx = F(x). \]

This justifies also calling the indefinite integral the *antiderivative*.

For a closed interval \([a, b]\), the FTC is

\[ \int_a^b F(x) dx = f(b) - f(a), \quad (DE-2.2) \]

thus the integral is independent of the path from \( x = a \) to \( x = b \).

**Fundamental Theorem of Complex Calculus:**
According to the fundamental theorem of complex calculus (FTCC),

\[ f(z) = f(z_0) + \int_{z_0}^z F(\zeta) d\zeta, \quad (DE-2.3) \]

where \( z_0, z, \zeta, F \in \mathbb{C} \). It follows that

\[ \frac{df(z)}{dz} = \frac{d}{dz} \int_{z_0}^z F(\zeta) d\zeta = F(z). \quad (DE-2.4) \]

For a closed interval \([s, s_0]\), the FTCC is

\[ \int_{s_0}^s F(\zeta) d\zeta = f(s) - f(s_0), \quad (DE-2.5) \]

thus the integral is independent of the path from \( x = a \) to \( x = b \).

**Problem #1**

1.1: Consider Equation DE-2.1. What is the condition on \( F(x) \) for which this formula is true?
1.2: Consider Equation DE-2.3. What is the condition on $F(z)$ for which this formula is true?

**Problem # 2**: In the following problems, solve the integral

$$I = \int_C F(z) \, dz$$

for a given path $C$.

2.1: Perform the following integrals ($z = x + iy \in \mathbb{C}$):

1. $I = \int_0^1 z \, dz$
2. $I = \int_0^1 \frac{1}{z} \, dz$, but this time make the path explicit: from 0 to 1, with y=0, and then to y=1, with x=1.
3. Discuss whether your results agree with Eq. DE-2.4?

2.2: Perform the following integrals on the closed path $C$, which we define to be the unit circle. You should substitute $z = e^{i\theta}$ and $dz = ie^{i\theta} \, d\theta$, and integrate from $\{\pi, -\pi\}$ to go once around the unit circle.

1. $\int_C z \, dz$
2. $\int_C \frac{1}{z} \, dz$
3. Discuss whether your results agree with Eq. DE-2.4?

**Problem # 3**: FTCC and integration in the complex plane

Let the function $F(z) = c^z$, where $c \in \mathbb{C}$ is given for each question. *Hint: Can you apply the FTCC?*

3.1: For the function $f(z) = c^z$, where $c \in \mathbb{C}$ is an arbitrary complex constant, use the Cauchy-Riemann (CR) equations to show that $f(z)$ is analytic for all $z \in \mathbb{C}$.

3.2: Find the antiderivative of $F(z)$.

3.3: $c = \frac{1}{e} = 1/2.7183$, ... where $C$ is $\zeta = 0 \to i \to z$

3.4: $c = 2$, where $C$ is $\zeta = 0 \to (1 + i) \to z$

3.5: $c = i$, where the path $C$ is an inward spiral described by $z(t) = 0.99^t e^{12\pi t}$ for $t = 0 \to t_0 \to \infty$

3.6: $c = e^{t - \tau_0}$, where $\tau_0 > 0$ is a real number and $C$ is $z = (1-i\infty) \to (1 + i\infty)$. *Hint: Do you recognize this integral? If you do not, please do not spend a lot of time trying to solve it via the “brute force” method.*

**Problem # 4**: Cauchy’s theorems for integration in the complex plane

There are three basic definitions related to Cauchy’s integral formula. They are all related and can greatly simplify integration in the complex plane. When a function depends on a complex variable, we use uppercase notation, consistent with the engineering literature for the Laplace transform.

1. **Cauchy’s (Integral) Theorem CT-1** (Stillwell, 2010, p. 319; Boas, 1987, p. 45)

$$\oint_C F(z) \, dz = 0$$
if and only if \( F(z) \) is complex analytic inside of \( C \). This is related to the FTCC,
\[
f(z) = f(a) + \int_a^z F(z)\,dz,
\]
where \( f(z) \) is the antiderivative of \( F(z) \)—namely, \( F(z) = df/dz \). The FTCC requires \( F(z) \) to be complex analytic for all \( z \in \mathbb{C} \). By closing the path (contour \( C \)), Cauchy’s theorem (and the following theorems) allows us to integrate functions that may not be complex analytic for all \( z \in \mathbb{C} \).

\[
\frac{1}{2\pi j} \oint_C \frac{F(z)}{z-z_0} \, dz = \begin{cases} 
F(z_0), & z_0 \in C \text{ (inside)} \\
0, & z_0 \notin C \text{ (outside)}
\end{cases}
\]
Here \( F(z) \) is required to be analytic everywhere within (and on) the contour \( C \). \( F(z_0) \) is called the residue of the pole.

3. **(Cauchy’s) Residue Theorem CT-3** (Boas (1987), p. 72)
\[
\oint_C F(z)\,dz = 2\pi j \sum_{k=1}^{K} \text{Res}_k,
\]
where \( \text{Res}_k \) are the residues of all poles of \( F(z) \) enclosed by the contour \( C \).

**How to calculate the residues**: The residues can be rigorously defined as
\[
\text{Res}_k = \lim_{z \to z_k} [(z - z_k)f(z)].
\]
This can be related to Cauchy’s integral formula: Consider the function \( F(z) = w(z)/(z - z_k) \), where we have factored \( F(z) \) to isolate the first-order pole at \( z = z_k \). If the remaining factor \( w(z) \) is analytic at \( z_k \), then the residue of the pole at \( z = z_k \) is \( w(z_k) \).

– **4.1: Describe the relationships between the theorems:**

1. CT-1 and CT-2
2. CT-1 and CT-3
3. CT-2 and CT-3

– **4.2: Consider the function with poles at \( z = \pm j \),
\[
F(z) = \frac{1}{1 + z^2} = \frac{1}{(z - j)(z + j)}.
\]
Find the residue expansion.

– **4.3: Apply Cauchy’s theorems to solve the following integrals. State which theorem(s) you used and show your work.**

1. \( \oint_C F(z)\,dz \), where \( C \) is a circle centered at \( z = 0 \) with a radius of \( \frac{1}{2} \)
2. \( \oint_C F(z)\,dz \), where \( C \) is a circle centered at \( z = j \) with a radius of 1
3. \( \oint_C F(z)\,dz \), where \( C \) is a circle centered at \( z = 0 \) with a radius of 2

**Problem # 5: Integration in the complex plane**
In the following questions, you’ll be asked to integrate \( F(s) = u(\sigma, \omega) + iv(\sigma, \omega) \) around the contour \( C \) for complex \( s = \sigma + i\omega \),
\[
\oint_C F(s)\,ds.
\]
Follow the directions carefully for each question. When asked to state where the function is and is not analytic, you are not required to use the Cauchy-Riemann equations (but you should if you can’t answer the question “by inspection”).
4.6. PROBLEMS DE-2

\[4.6. \text{ PROBLEMS DE-2}\]

\(-5.1: \ F(s) = \sin(s)\)

\(-5.2: \ \text{Given function } F(s) = \frac{1}{s}\)

1. State where the function is and is not analytic.
2. Explicitly evaluate the integral when \(C\) is the unit circle, defined as \(s = e^{i\theta}, 0 \leq \theta \leq 2\pi\).
3. Evaluate the same integral using Cauchy’s theorem and/or the residue theorem.

\(-5.3: \ F(s) = \frac{1}{s^2}\)

1. State where the function is and is not analytic.
2. Explicitly evaluate the integral when \(C\) is the unit circle, defined as \(s = e^{i\theta}, 0 \leq \theta \leq 2\pi\).
3. What does your result imply about the residue of the second-order pole at \(s = 0\)?

\(-5.4: \ F(s) = e^{st}\)

1. State where the function is and is not analytic.
2. Explicitly evaluate the integral when \(C\) is the square \((\sigma, \omega) = (1, 1) \rightarrow (-1, 1) \rightarrow (1, -1) \rightarrow (-1, -1) \rightarrow (1, 1)\).
3. Evaluate the same integral using Cauchy’s theorem and/or the residue theorem.

\(-5.5: \ F(s) = \frac{1}{s+2}\)

1. State where the function is and is not analytic.
2. Let \(C\) be the unit circle, defined as \(s = e^{i\theta}, 0 \leq \theta \leq 2\pi\). Evaluate the integral using Cauchy’s theorem and/or the residue theorem.
3. Let \(C\) be a circle of radius 3, defined as \(s = 3e^{i\theta}, 0 \leq \theta \leq 2\pi\). Evaluate the integral using Cauchy’s theorem and/or the residue theorem.

\(-5.6: \ F(s) = \frac{1}{2\pi i} \frac{e^{st}}{(s+4)}\)

1. State where the function is and is not analytic.
2. Let \(C\) be a circle of radius 3, defined as \(s = 3e^{i\theta}, 0 \leq \theta \leq 2\pi\). Evaluate the integral using Cauchy’s theorem and/or the residue theorem.
3. Let \(C\) contain the entire left half \(s\) plane. Evaluate the integral using Cauchy’s theorem and/or the residue theorem. Do you recognize this integral?

\(-5.7: \ F(s) = \pm \frac{1}{\sqrt{s}} \ (e.g., \ F^2 = \frac{1}{s})\)

1. State where the function is and is not analytic.
2. This function is multivalued. How many Riemann sheets do you need in the domain \((s)\) and the range \((f)\) to fully represent this function? Indicate (e.g., using a sketch) how the sheet(s) in the domain map to the sheet(s) in the range.
3. Explicitly evaluate the integral

\[\int_C \frac{1}{\sqrt{s}} \, dz\]

when \(C\) is the unit circle, defined as \(s = e^{i\theta}, 0 \leq \theta \leq 2\pi\). Is this contour closed? State why or why not.
4. Explicitly evaluate the integral
\[ \int_C \frac{1}{\sqrt{z}} \, dz \]
when \( C \) is twice around the unit circle, defined as \( s = e^{i\theta}, 0 \leq \theta \leq 4\pi \). Is this contour closed? State why or why not. Hint: Note that
\[ \sqrt{e^{i(\theta + 2\pi)}} = \sqrt{e^{i2\pi}e^{i\theta}} = e^{i\theta} = e^{i\pi} \sqrt{e^{i\theta}} = -1 \sqrt{e^{i\theta}}. \]

5. What does your result imply about the residue of the (twice-around \( \frac{1}{2} \) order) pole at \( s = 0 \)?

6. Show that the residue is zero. Hint: Apply the definition of the residue.

**Problem # 6: A two-port network application for the Laplace transform**

![Figure 4.6](image)

This three-element electrical circuit is a system that acts to low-pass filter the signal voltage \( V_1(\omega) \), to produce signal \( V_2(\omega) \). It is convenient to define the dimensionless ratio \( s/s_c = RCs \) in terms of a time constant \( \tau = RC \) and cutoff frequency \( s_c = 1/\tau \).

- 6.1: Find the \( 2 \times 2 \) ABCD matrix representation of Fig. 3.1. Express the results in terms of the dimensionless ratio \( s/s_c \), where \( s_c = 1/\tau \) is the cutoff frequency and \( \tau = RC \) is the time constant.

- 6.2: Find the eigenvalues of the \( 2 \times 2 \) ABCD matrix. Hint: See Appendix B.3, page 237.

- 6.3: Assuming that \( I_2 = 0 \), find the transfer function \( H(s) \equiv V_2/V_1 \). From the results of the ABCD matrix you determined in questions 6.1 and 6.2, show that
\[ H(s) = \frac{s_c}{s + s_c}. \]  

- 6.4: The transfer function \( H(s) \) has one pole. Where is the pole and residue?

- 6.5: Find \( h(t) \), the inverse Laplace transform of \( H(s) \).

- 6.6: Assuming that \( V_2 = 0 \), find \( Y_{12}(s) \equiv I_2/V_1 \).

- 6.7: Find the input impedance to the right-hand side of the system \( Z_{22}(s) \equiv V_2/I_2 \) for two cases: (1) \( I_1 = 0 \) and (2) \( V_1 = 0 \).

- 6.8: Compute the determinant of the ABCD matrix. Hint: It is always \pm 1.

- 6.9: Given the result of the previous problem Eq. DE-2.8, compute the derivative of \( H(s) = \frac{V_2}{V_1} \bigg|_{I_2=0} \).
### Problem #7: With the help of a computer

Now we look at a few important concepts using Matlab/Octave’s `syms` commands or Wolfram Alpha’s symbolic math toolbox.\(^{13}\)

For example, to find the Taylor series expansion about \( s = 0 \) of

\[
F(s) = -\log(1 - s),
\]

we first consider the derivative and its Taylor series (about \( s = 0 \))

\[
F'(s) = \frac{1}{1 - s} = \sum_{n=0}^{\infty} s^n.
\]

Then, we integrate this series term by term:

\[
F(s) = -\log(1 - s) = \int_0^s F'(s) \, ds = \sum_{n=0}^{\infty} \frac{s^n}{n}.
\]

Alternatively we can use Matlab/Octave commands:

```matlab
syms s
taylor(-log(1-s), 'order', 7)
```

- **7.1:** Use Octave’s `taylor(-log(1-s))` to the seventh order, as in the example above.

1. Try the above Matlab/Octave commands. Give the first seven terms of the Taylor series (confirm that Matlab/Octave agrees with the formula derived above).

2. What is the inverse Laplace transform of this series? Consider the series term by term.

- **7.2:** The function \( \frac{1}{\sqrt{z}} \) has a branch point at \( z = 0 \); thus it is singular there.

1. Can you apply Cauchy’s integral theorem when integrating around the unit circle?

2. This Matlab/Octave code computes \( \int_0^{4\pi} \frac{dz}{\sqrt{z}} \) using Matlab’s/Octave’s symbolic analysis package:

```matlab
syms z
I=int(1/sqrt(z))
J = int(1/sqrt(z),exp(-j*pi),exp(j*pi))
eval(J)
```

Run this script. What answers do you get for \( I \) and \( J \)?

3. Modify this code to integrate \( f(z) = 1/z^2 \) once around the unit circle. What answers do you get for \( I \) and \( J \)?

- **7.3:** Bessel functions can describe waves in a cylindrical geometry.

The Bessel function has a Laplace transform with a branch cut \( J_0(t)u(t) \leftrightarrow \frac{1}{\sqrt{1 + s^2}} \).

Draw a hand sketch showing the nature of the branch cut. Hint: Use `zviz`.

- **7.4** Try the following Matlab/Octave commands, and then comment on your findings.

```matlab
%Take the inverse LT of 1/sqrt(1+s^2)
syms s
I=ilaplace(1/(sqrt((1+s^2))));
disp(I)

%Find the Taylor series of the LT
T=taylor(1/sqrt(1+s^2),10); disp(T);
```

\(^{13}\)https://www.wolframalpha.com/
7.5: When did Friedrich Bessel live?

7.6: What did he use Bessel functions for?

7.7: Use \texttt{zviz} for each of the following:

1. Describe the plot generated by \texttt{zviz \$=Z\$}.

2. Are the functions that follow legal Brune impedances? [Do they obey $\Re \{Z(\sigma > 0) \geq 0\}$? Hint: Consider the phase (color). Plot \texttt{zviz \$} for a reminder of the color map.

   1. \texttt{zviz 1./sqrt(1+S.^2)}
   2. \texttt{zviz 1./sqrt(1-S.^2)}
   3. \texttt{zviz 1./(1+sqrt(S))}

**Problem #8:** This problem is for extra credit.

8.1: Find the $\mathcal{L}^{-1}$ of one factor of the Riemann zeta function $\zeta_p(s)$, where $\zeta_p(s) \leftrightarrow z_p(t)$. Describe your results in words. Hint: See Eq. AE-1.8, p. 30. Hint: Consider the geometric series representation

$$
\zeta_p(s) = \frac{1}{1-e^{-sT_p}} = \sum_{k=0}^{\infty} e^{-skT_p}, \quad (DE-2.7)
$$

for which you can look up the $\mathcal{L}^{-1}$ of each term.

**Problem #9:** Inverse transform of products:

The time-domain version of Eq. DE-2.9 may be written as the convolution of all the $z_k(t)$ factors:

$$
z(t) \equiv z_2(t) \ast z_3(t) \ast z_5(t) \ast z_7(t) \ast \ldots \ast z_p(t) \ast \ldots, \quad (DE-2.8)
$$

where $\ast$ represents time convolution.

*Figure 4.7:* This feedback network, described by a time-domain difference equation with delay $T_p$, has an all-pole transfer function $\zeta_p(s) \equiv Q(s)/I(s)$ given by Eq. DE-2.12, which physically corresponds to a stub of a transmission line, with the input at one end and the output at the other. To describe the $\zeta(s)$ function we must take $\alpha = -1$. A transfer function $Y(s) = V(s)/I(s)$ that has the same poles as $\zeta_p(s)$, but with zeros as given by Eq. DE-2.14, is the input admittance $Y(s) = I(s)/V(s)$ of the transmission line, defined as the ratio of the Laplace transform of the current $i(t) \leftrightarrow I(s)$ to the voltage $v(t) \leftrightarrow V(s)$.

Explain what this means in physical terms. Start with two terms (e.g., $z_1(t) \ast z_2$).
Physical interpretation: Such functions may be generated in the time domain, as shown in Fig. 3.2 (p. 66), using a feedback delay of $T_p$ seconds, described by the two equations in the Fig. 3.2 with a unity feedback gain $\alpha = -1$. Taking the Laplace transform of the system equation, we see that the transfer function between the state variable $q(t)$ and the input $x(t)$ is given by $\zeta_p(s)$, which is an all-pole function, since

$$Q(s) = e^{-sT_p} Q(s) + V(s), \quad \text{or} \quad \zeta_p(s) \equiv \frac{Q(s)}{V(s)} = \frac{1}{1 - e^{-sT_p}}.$$ (DE-2.9)

Closing the feed-forward path gives a second transfer function $Y(s) = I(s)/V(s)$—namely,

$$Y(s) \equiv \frac{I(s)}{V(s)} = \frac{1 - e^{-sT_p}}{1 + e^{-sT_p}}.$$ (DE-2.10)

If we take $i(t)$ as the current and $v(t)$ as the voltage at the input to the transmission line, then $y_p(t) \leftrightarrow \zeta_p(s)$ represents the input impedance at the input to the line. The poles and zeros of the impedance interleave along the $j\omega$ axis. By a slight modification, $\zeta_p(s)$ may alternatively be written as

$$Y_p(s) = \frac{e^{sT_p/2} + e^{-sT_p/2}}{e^{sT_p/2} - e^{-sT_p/2}} = j \tan(sT_p/2).$$ (DE-2.11)

Every impedance $Z(s)$ has a corresponding reflectance function given by a Möbius transformation, which may be read off of Eq. DE-2.14 as

$$\Gamma(s) \equiv \frac{1 + Z(s)}{1 - Z(s)} = e^{-sT_p},$$ (DE-2.12)

since impedance is also related to the round-trip delay $T_p$ on the line. The inverse Laplace transform of $\Gamma(s)$ is the round-trip delay $T_p$ on the line

$$\gamma(t) = \delta(t - T_p) \leftrightarrow e^{-sT_p}.$$ (DE-2.13)

Working in the time domain provides a key insight, as it allows us to parse out the best analytic continuation of the infinity of possible continuations that are not obvious in the frequency domain. Transforming to the time domain is a form of analytic continuation of $\zeta(s)$ that depends on the assumption that $z(t)$ is one-sided in time (causal).
4.7 Inverse Laplace transform $LT^{-1}$

The inverse Laplace transform $LT^{-1}$ (Eq. 3.9.0.1, p. 120) transforms a function of complex frequency $F(s)$ and returns a causal function of time $f(t)$,

$$f(t) \leftrightarrow F(s),$$

where $f(t) = 0$ for $t < 0$. Examples are provided in Table C.3 (p. 242). Now we discuss the details of finding the inverse transform by using CT-3, and we see how the causal requirement $f(t < 0) = 0$ comes about.

The integrand of the inverse transform is $F(s)e^{st}$ and the limits of integration are $-\sigma_o \mp \omega_j$. To find the inverse, we must close the curve at infinity and specify that the integral at $\omega \to \infty$. There are two ways to close these limits—to the right $\sigma > 0$ (RHP) and to the left $\sigma < 0$ (LHP)—but there needs to be some logical reason for this choice. That logic is determined by the sign of $t$. For the integral to converge, the term $e^{st}$ must go to zero as $\omega \to \infty$. In terms of the real and imaginary parts of $s = \sigma + \omega j$, the exponential may be rewritten as $e^{\sigma t}e^{\omega j t}$. Note that both $t$ and $\omega$ go to $\infty$. Thus it is the interaction between these two limits that determines how we pick the closure, RHP or LHP.

4.7.1 Case for negative time ($t < 0$) and causality:

Let us first consider negative time, including $t \to -\infty$. If we were to close $C$ in the LHP ($\sigma < 0$), then the product $\sigma t$ is positive ($\sigma < 0, t < 0$, thus $\sigma t > 0$). In this case, as $\omega \to \infty$, the closure integral $|s| \to \infty$ will diverge. Thus we may not close in the LHP for negative time. If we close in the RHP ($\sigma > 0$), then the product $\sigma t < 0$ and $e^{st}$ will go to zero as $\omega \to \infty$. This then justifies closing the contour, allowing for the use of the Cauchy theorems.

If $F(s)$ is analytic in the RHP, the FTCC applies, and the resulting $f(t)$ must be zero and the inverse Laplace transform must be causal. This argument holds for any $F(s)$ that is analytic in the RHP ($\sigma > 0$).

Unstable poles: An important but subtle point arises: If $F(s)$ has a pole in the RHP, then the above argument still applies if we pick $\sigma_o$ to be to the right of the RHP pole. This means that the inverse transform may still be applied to unstable poles (those in the RHP). This explains the need for the $\sigma_o$ in the limits. If $F(s)$ has no RHP poles, then $\sigma_o = 0$ is adequate and this factor may be ignored.

4.7.2 Case for zero time ($t = 0$):

When the time is zero, the integral does not, in general, converge, which leaves $f(t)$ undefined. This is most clear in the case of the step function $u(t) \leftrightarrow 1/s$, where the integral may not be closed because the convergence factor $e^{st} = 1$ is lost for $t = 0$.

The fact that $u(t)$ does not exist at $t = 0$ explains the Gibbs phenomenon in the inverse Fourier transform. At times where a jump occurs, the derivative of the function does not exist, and thus the time response function is not analytic. The Fourier expansion cannot converge at places where the function is not analytic. A low-pass filter may be used to smooth the function, but at the cost of temporal resolution.

4.7.3 Case for positive time ($t > 0$)

Next we investigate the convergence of the integral for positive time $t > 0$. In this case we must close the integral in the LHP ($\sigma < 0$) for convergence, so that $\sigma t < 0$ ($\sigma \leq 0$ and $t > 0$). When there are poles on the $\omega_j = 0$ axis, $\sigma_o > 0$ assures convergence by keeping the on-axis poles inside the contour. At this point, CT-3 is relevant. If we restrict ourselves to simple poles (as required for a Brune impedance), the residue theorem may be directly applied.

The simplest example is the step function, for which $F(s) = 1/s$ and thus

$$u(t) = \int_{\text{LHP}} \frac{e^{st}}{s} \frac{ds}{2\pi j} \leftrightarrow \frac{1}{s},$$

which is a direct application of CT-3 (Eq. 4.5.1.3 p. 152). The forward transform of $u(t)$ is straightforward, as discussed on page 119. This is true of most, if not all, of the elementary forward Laplace transforms. In these cases, causality is built into the integral by its limits. An interesting problem is how to prove that $u(t)$ is not defined at $t = 0$.

The inverse Laplace transform of $F(s) = 1/(s + 1)$ has a residue of 1 at $s = -1$, thus that is the only contribution to the integral. More demanding cases are Laplace transform pairs

$$\frac{1}{\sqrt{t}} u(t) \leftrightarrow \sqrt{\frac{\pi}{s}} \quad \text{and} \quad J_o(t) u(t) \leftrightarrow \frac{1}{\sqrt{s^2 + 1}}.$$
4.7. INVERSE LAPLACE TRANSFORM \( \mathcal{L}^{-1} \)

\[ s = \cos(\pi z) \]

\[ s = \text{besselj}(0, \pi z) \]

Figure 4.8: Left: Colorized plot of \( w(z) = \sin(z) \). Right: Colorized plot of \( w(z) = J_0(\pi z) \). Note the similarity of the two functions. The first Bessel zero is at 2.405 and thus appears at \( 0.7655 = 2.405/\pi \), about 1.53 times larger than the root of \( \cos(\pi z) \) at 1/2. Other than this minor distortion of the first few roots, the two functions are basically identical. It follows that their \( \mathcal{L}^{-1} \)’s must have similar characteristics, as documented in Table C.4 (p. 243).

as shown in Fig 4.8 (right), and more in Table C.4 (p. 243). Many of these are easily proved in the forward direction but are much more difficult in the inverse direction due to the properties at \( t = 0 \), unless CT-3 is invoked. The last \( \mathcal{L}^{-1} \) pair gives insight into the properties of Bessel functions \( J_0(\pi z) \) and \( H^{(1)}_0(\pi z) \), with a branch cut along the negative axis (see Fig. 4.4).

The form \( \sqrt{s} \) is called a semiinductor (Kim and Allen, 2013), also known as the skin effect in EM theory. The form \( 1/\sqrt{s} \) is a semicapacitor (see Fig. 4.3 (right)).

Two more examples are given in Fig. 4.9 to show Bessel functions \( J_0(\pi z) \) and the Hankel function \( H^{(1)}_0(\pi z/2) \) colorized maps. Note how the white and black contour lines are always perpendicular where they cross, just as in the calibration plots for the x- and y-axes, shown in Fig. 3.12 (p. 125).

Along the x-axis, \( \cos(\pi x) \) is periodic with a period of \( \pi \). The dark spots are at the zeros at \( \pm\pi/2, \pm3\pi/2, \ldots \). Along the \( jy \)-axis, the function goes to either zero (black) or \( \infty \) (white). This behavior carries the same \( \pi \) periodicity as on the \( x = 0 \) line. On the right in Fig. 4.9 is the Hankel function \( H^{(1)}_0(\pi z) \), which is a mixed and distorted version of \( \cos(\pi z) \) with the zeros pushed downward, and \( e^{\pi z} \). This colorized plot shows that these two functions become the same for \( x = \Re z > 0 \).

Some open questions: Without the use of CT-3 it is difficult to see how to evaluate the inverse Laplace transform of \( 1/s \) directly. For example, how do we show that the integral (Eq. 4.5.1.2) is zero for negative time (or 1 for positive time)? CT-3 neatly solves this difficult problem by the convergence of the integral for negative and positive times. Clearly the continuity of the integral at \( \omega \to \infty \) plays an important role. Perhaps the Riemann sphere plays a role in this that has not yet been explored.

4.7.4 Properties of the \( \mathcal{L}^{-1} \)

As shown in Table C.1 of Laplace transforms, there are integral (i.e., integration, not integer) relationships, or properties, that are helpful to identify. The first of these is a definition, not a property:

\[ f(t) \leftrightarrow F(s). \]

Causality: When we take the \( \mathcal{L}^{-1} \), the time-domain response is in lowercase (e.g., \( f(t) \)) and the frequency-domain transform is in uppercase (e.g., \( F(s) \)). It is required, but not always explicitly specified, that \( f(t < 0) = 0 \); that is, the time function must be causal, as stated by Postulate P1 (121).

Linearity: The most basic property is the linearity (superposition) property of the \( \mathcal{L}^{-1} \), stated by Postulate P2 (p. 122).
Convolution property: The product of two LTs in frequency results in convolution in time:

\[ F(s)G(s) \leftrightarrow f(t) * g(t) = \int_0^t f(\tau)g(t - \tau)\,d\tau, \]

where we use \(*\) to indicate the convolution of two time functions.

A key application of convolution is filtering, which takes many forms. The most basic filter is the moving average, the moving sum of data samples, normalized by the number of samples. Such a filter has very poor performance. It also introduces a delay of half the length of the average, which may or may not constitute a problem, depending on the application. Other important examples are a low-pass filter that removes high-frequency noise and a notch filter that removes line noise (i.e., 60 [Hz] in the United States, and its second and third harmonics, 120 and 180 [Hz]). Such noise is typically a result of poor grounding and ground loops. It is better to solve the problem at its root than to remove it with a notch filter. Still, filters are very important in engineering.

By taking the LT of the convolution we can derive this relationship:

\[
\int_0^\infty [f(t) * g(t)] e^{-st} \,dt = \int_0^\infty \left( \int_0^t f(\tau)g(t - \tau)\,d\tau \right) e^{-st} \,dt \\
= \int_0^t f(\tau) \left( \int_0^\infty g(t - \tau)e^{-st} \,d\tau \right) \,d\tau \\
= \int_0^t f(\tau) \left( e^{-s\tau} \int_0^\infty g(t')e^{-st'} \,dt' \right) \,d\tau \\
= G(s) \int_0^t f(\tau) e^{-s\tau} \,d\tau \\
= G(s)F(s).
\]

We first encountered this relationship on page 89 in the context of multiplying polynomials, which is the same as convolving their coefficients. The parallel should be obvious. In the case of polynomials, the convolution is discrete in the coefficients, and here it is continuous in time. But the relationships are the same.

Time-shift property: When a function is time-shifted by time \(T_o\), the LT is modified by \(e^{sT_o}\), leading to the property

\[ f(t - T_o) \leftrightarrow e^{-sT_o} F(s). \]

This is easily shown by applying the definition of the LT to a delayed time function.
**Time derivative:** The key to the eigenfunction analysis provided by the \( \mathcal{L} \) is the transformation of a time derivative on a time function—that is, \[ \frac{df(t)}{dt} \leftrightarrow sF(s). \]

Here \( s \) is the eigenvalue corresponding to the time derivative of \( e^{st} \). Given the definition of the derivative of \( e^{st} \) with respect to time, this definition seems trivial. Yet that definition was not obvious to Euler. It needed to be extended to the space of the complex analytic function \( e^{st} \), which did not happen until at least Riemann (1851).

Given a differential equation of order \( K \), the \( \mathcal{L} \) results in a polynomial in \( s \) of degree \( K \). It follows that this \( \mathcal{L} \) property is the cornerstone of why the \( \mathcal{L} \) is so important to scalar differential equations, as it was to the early analysis of Pell’s equation and the Fibonacci sequence, presented in earlier chapters. This property was first uncovered by Euler. It is not clear whether he fully appreciated its significance, but by the time of his death, it certainly would have been clear to him. Who first coined the terms *eigenvalue* and *eigenfunction*? The word *eigen* is a German word meaning of one.

**Initial and final value theorems:** There are much more subtle relationships between \( f(t) \) and \( F(s) \) that characterize \( f(0^+) \) and \( f(t \to \infty) \). While these properties can be very important in certain applications, they are beyond the scope of this text. They relate to so-called initial value theorems. If the system under investigation has potential energy at \( t = 0 \), then the voltage (velocity) need not be zero for negative time. An example is a charged capacitor or a moving mass. These are important situations, but better explored in a more in-depth treatment.

**4.7.5 Solving differential equations:**

Many differential equations may be solved by assuming a power series (i.e., Taylor series) solution of the form

\[ y(x) = x^r \sum_{n=0}^{\infty} c_n x^n, \]  

(4.7.5.1)

with \( r \in \mathbb{Z} \) and coefficients \( c_n \in \mathbb{C} \). The method of Frobenius is quite general (Greenberg, 1988, p. 193).

**Exercise #3**
When a solution of this form is substituted into the differential equation, a recursion relationship in the coefficients results. For example, if the equation is

\[ y''(x) = \lambda^2 y(x), \]

the recursion is \( c_n = c_{n-1}/n \). The resulting equation is

\[ y(x) = e^{\lambda x} = x^0 \sum_{n=0}^{\infty} \frac{1}{n!} x^n, \]

namely, \( c_n = 1/n! \), thus \( n c_n = 1/(n-1)! = c_{n-1} \).

**Exercise #4**

Find the recursion relationship for \( y(x) = J_\nu(x) \) of order \( \nu \) that satisfies Bessel’s equation

\[ x^2 y''(x) + xy'(x) + (x^2 - \nu^2)y(x) = 0. \]

**Solution:** If we assume a complex analytic solution of the form of Eq. 4.7.5.1, we find the Bessel recursion relationship for coefficients \( c_k \) (Greenberg, 1988, p. 231):

\[ c_k = - \frac{1}{k(k + 2\nu)} c_{k-2}. \]
4.8 Problems DE-3

Topics of this homework:
Brune impedance, lattice transmission line analysis

Brune Impedance

Problem #1: Residue form
A Brune impedance is defined as the ratio of the force \( F(s) \) to the flow \( V(s) \) and may be expressed in residue form as

\[
Z(s) = c_0 + \sum_{k=1}^{K} \frac{c_k}{s - s_k} = \frac{N(s)}{D(s)}
\]

(DE-3.1)

with

\[
D(s) = \prod_{k=1}^{K} (s - s_k)
\]

and

\[
c_k = \lim_{s \to s_k} (s - s_k)D(s) = \prod_{n'=1}^{K-1} (s - s_{n'}). \]

The prime on the index \( n' \) means that \( n = k \) is not included in the product.

- 1.1: Find the Laplace transform (LT) of a (1) spring, (2) dashpot, and (3) mass.
Express these in terms of the force \( F(s) \) and the velocity \( V(s) \), along with the electrical equivalent impedance:
(1) Hooke’s law \( f(t) = Kx(t) \), (2) dashpot resistance \( f(t) = Rv(t) \), and (3) Newton’s law for mass \( f(t) = Mdv(t)/dt \).

- 1.2: Take the Laplace transform (LT) of Eq. DE-3.2 and find the total impedance \( Z(s) \) of the mechanical circuit.

\[
M \frac{d^2}{dt^2} x(t) + R \frac{dx(t)}{dt} + Kx(t) = f(t) \leftrightarrow (Ms^2 + Rs + K)X(s) = F(s). 
\]

(DE-3.2)

- 1.3: What are \( N(s) \) and \( D(s) \) (see Eq. DE-3.1)?

- 1.4: Assume that \( M = R = K = 1 \) and find the residue form of the admittance \( Y(s) = 1/Z(s) \) (see Eq. DE-3.1) in terms of the roots \( s_{\pm} \). Hint: Check your answer with Octave’s/Matlab’s residue command.

- 1.5: By applying Eq. 4.5.1.3 (page 152), find the inverse Laplace transform (LT\(^{-1} \)). Use the residue form of the expression that you derived in question 1.4.

Problem #2: Train-mission-line We wish to model the dynamics of a freight train that has \( N \) such cars and study the velocity transfer function under various load conditions.

As shown in Fig. 3.3, the train model consists of masses connected by springs.

Use the ABCD method (see the discussion in Appendix B.3, p. 237) to find the matrix representation of the system of Fig. 3.3. Define the force on the \( n \)th train car \( f_n(t) \leftrightarrow F_n(\omega) \) and the velocity \( v_n(t) \leftrightarrow V_n(\omega) \).

Break the model into cells consisting of three elements: a series inductor representing half the mass \( (M/2) \), a shunt capacitor representing the spring \( (C = 1/K) \), and another series inductor representing half the mass \( (L = M/2) \), transforming the model into a cascade of symmetric \( (A = D) \) identical cell matrices \( T(s) \).
4.8. PROBLEMS DE-3

Figure 4.10: Depiction of a train consisting of cars treated as masses \( M \) and linkages treated as springs of stiffness \( K \) or compliance \( C = 1/K \). Below it is the electrical equivalent circuit for comparison. The masses are modeled as inductors and the springs as capacitors to ground. The velocity is analogous to a current and the force \( f_n(t) \) to the voltage \( \phi_n(t) \). The length of each cell is \( \Delta \) [m]. The train may be accurately modeled as a transmission line (TL), since the equivalent electrical circuit is a lumped model of a TL. This method, called a Cauer synthesis, is based on the ABCD transmission line method of Sec. 3.7 (p. 110).

- 2.1: Find the elements of the ABCD matrix \( T \) for the single cell that relate the input node 1 to output node 2

\[
\begin{bmatrix} F \\ V \end{bmatrix}_1 = T \begin{bmatrix} F(\omega) \\ -V(\omega) \end{bmatrix}_2. \tag{DE-3.3}
\]

- 2.2: Express each element of \( T(s) \) in terms of the complex Nyquist ratio \( s/s_c \) < 1 (\( s = 2\pi j f, \ s_c = 2\pi j f_c \)). The Nyquist sampling cutoff frequency \( f_c \) is defined in terms the minimum number of cells (i.e., 2) of length \( \Delta \) per wavelength.

The Nyquist sampling theorem says that there are at least two cars per wavelength (more than two time samples at the highest frequency). From the figure, the distance between cars \( \Delta = c_o T_o \) [m], where

\[ c_o = \frac{1}{\sqrt{MC}} \] [m/s].

The cutoff frequency obeys \( f_c \lambda_o = c_o \), where the Nyquist wavelength is \( \lambda_o = 2\Delta \). Therefore the Nyquist sampling condition is

\[ \omega < \omega_c = 2\pi f_c \equiv \frac{2\pi c_o}{\lambda_c} = \frac{2\pi c_o}{2\Delta} = \frac{\pi}{\Delta\sqrt{MC}} \] [Hz]. \tag{DE-3.4}

- 2.3: Use the property of the Nyquist sampling frequency \( \omega < \omega_c \) (Eq. DE-3.6) to remove higher order powers of frequency

\[ 1 + \left( \frac{s}{s_c} \right)^0 \approx 1 \] \tag{DE-3.5}

to determine a band-limited approximation of \( T(s) \).

**Problem # 3** Now consider the cascade of \( N \) such \( T(s) \) matrices and perform an eigenanalysis. Find the eigenvalues and eigenvectors of \( T(s) \) as functions of \( s/s_c \).

**Problem # 4** The velocity transfer function

- 4.1: Assuming that \( N = 2 \) and \( F_2 = 0 \) (two half-mass problem), find the transfer function \( H(s) \equiv V_2/V_1 \). From the results of the \( T \) matrix, find

\[ H_{21}(s) = \frac{V_2}{V_1} \bigg|_{F_2=0} \]

Express \( H_{12} \) in terms of a residue expansion.

- 4.2: Find \( h_{21}(t) \leftrightarrow H_{21}(s) \).
– 4.3: What is the input impedance $Z_2 = F_2/V_2$, assuming $F_3 = -r_0V_3$?

– 4.4: Simplify the expression for $Z_2$ as follows:
1. Assuming the characteristic impedance $r_0 = \sqrt{M/C}$, 
2. terminate the system in $r_0$: $F_2 = -r_0V_2$ (i.e., $-V_2$ cancels).
3. Assume higher-order frequency terms are less than 1 ($|s/s_c| < 1$).
4. Let the number of cells $N \to \infty$. Thus $|s/s_c|^N = 0$.

When a transmission line is terminated in its characteristic impedance $r_0$, the input impedance $Z_1(s) = r_0$. Thus, when we simplify the expression for $T(s)$, it should be equal to $r_0$. Show that this is true for this setup.

– 4.5: State the ABCD matrix relationship between the first and $N$th nodes in terms of the cell matrix. Write out the transfer function for one cell, $H_{21}$.

– 4.6: What is the velocity transfer function $H_{N1} = \frac{V_N}{V_1}$?
Chapter 5

Stream 3B: Vector Calculus

5.1 Properties of fields and potentials

Before we can define the vector operations $\nabla()$, $\nabla \cdot ()$, $\nabla \times ()$, and $\nabla^2 ()$, we must define the objects they operate on: scalar and vector fields. The word field has two very different meanings: a mathematical one, which defines an algebraic structure, and a physical one, discussed next.

Ultimately we wish to integrate in $\in \mathbb{R}^3$, $\mathbb{R}^n$ and $\in \mathbb{C}^n$. Integration is quantified by several fundamental theorems of calculus, each about integration (see pp. 135–136).

5.1.1 Scalar and vector fields

Scalar fields: We use the term scalar field interchangeably with analytic in a connected region of the spatial vector $x = [x, y, z]^T \in \mathbb{R}^3$. In mathematics, functions that are piece-wise differentiable are called smooth, which is different from analytic. Every analytic function may be written as a single-valued and infinitely differentiable power series. A smooth function has at least one or more derivatives but need not be analytic.

Exercise #1

The simplest example of a scalar field is the voltage between two very large (think $\infty$) conducting parallel planes, bias to $V_o$ [V]. In this case the voltage varies linearly (the voltage is complex analytic) between the two plates. One scalar field is

$$\Phi(x, y, z) = V_o (1 - x) \quad [V]. \quad (5.1.1.1)$$

At $x = 0$ the voltage is $V_o$, and at $x = 1$ the voltage is zero. Between 0 and 1 the voltage varies linearly. Thus $\Phi(x)$ defines a scalar field. The gradient of $\Phi(x)$ is a square pulse

$$\nabla \Phi(x) = -V_o (u(x) - u(x - 1)).$$

Exercise #2

The function $f(t) = tu(t)$ is smooth and has one smooth derivative

$$\frac{d}{dt} tu(t) = u(t) + t\delta(t),$$

$$\frac{d^2}{dt^2} tu(t) = \frac{d}{dt} u(t) = \delta(t),$$

but it does not have a second derivative at $t = 0$. Thus $tu(t)$ is not analytic at $t = 0$. However, it has a Laplace transform $f(t) \leftrightarrow F(s)$,

$$tu(t) = u(t) \ast u(t) \leftrightarrow \frac{1}{s^2}$$

with a second-order pole at $s = 0$ with amplitude 1. The amplitude of $1/s$ must be zero, since $F(s)$ has no such term. Thus the LT is analytic everywhere except at its second-order pole. The derivative $df(t)/dt \leftrightarrow sF(s) = 1/s$ has a simple pole with residue 1.
Exercise #3

Next consider \((a, b, c, d \in \mathbb{C})\)

\[ G(s) = \frac{a + bF(\zeta)}{c + dF(\zeta)} = \frac{as^2 + b}{cs^2 + d}, \]

which has second order poles and zeros.

Exercise #4

The outbound eigenfunction of the lossy scalar wave equation in spherical coordinates (i.e., the spherical Bessel function) is

\[ g(r, t) = \frac{e^{st - \kappa(s)r}}{r}, \]

where \(g(r, t) \in \mathbb{C}\) is the complex pressure, \(\kappa(s) = (s + \beta_\omega \sqrt{s})/c_\omega \in \mathbb{C}\) is the complex wave number (Eq. D.1.0.1, p. 249), \(c_\omega\) is the speed of sound, and \(r = \sqrt{x^2 + y^2 + z^2}\). If we ignore viscous and thermal losses, \(\beta_\omega = 0\) (Mason, 1928).

The pressure, a potential, is the solution to the acoustic wave equation in spherical coordinates. The \(1/r^2\) term compensates for the increasing area of the spherical wave as it propagates, to maintain constant energy. The area of the wavefront is proportional to \(r^2\); thus the pressure must be proportional to \(1/r\) so that the integral of the energy \((\propto g^2 \propto 1/r^2)\) over the area \(\propto r^2\) remains constant as the wave progresses outward. Note that the gradient of the potential (i.e., pressure) is proportional to the flow (mass flux) of the wave. The power flux is the product of the potential and the flux, and the ratio is the impedance. In acoustics, this ratio is called the acoustic impedance, measured in acoustic ohms (see Table 3.2, p. 114).

Note that \(\ln g(r, s) = st - \kappa(s)r - \ln r\) is analytic everywhere except at \(r = 0\), but it is double-valued due to \(\beta_\omega \sqrt{s}\), which forces a branch cut, as required to fully describe it in the complex \(s\) plane.

To keep the discussion simple, initially we limit the definition to an analytic surface \(S(x)\), as shown in Fig. 5.1, that has height \(z(x, y) \in \mathbb{R}\) as a function of \(x, y \in \mathbb{R}^2\) (a plane):

\[ z(x, y, t) = \phi(x, y, t), \]

where \(z(x, y, t)\) describes a surface that is analytic in \(x\). Optionally, we can allow the field to be a single-valued function of time \(t \in \mathbb{R}\), since that is the nature of the solutions of the equations we wish to solve.

For example, in Fig. 5.1 we show a smooth single-valued potential at height \(z\) (e.g., constant temperature at height \(z\)) that has isolines (lines on a surface with constant slope).

Vector fields: A vector field is composed of three scalar fields. For example, the electric field used in Maxwell’s equations, \(E(x, t) = [E_x, E_y, E_z]^T [\text{V/m}]\), has three components, each a scalar field. When the magnetic flux vector \(B(x)\) is static (Postulate P5, p. 122), the potential \(\phi(x) [\text{V}]\) uniquely defines \(E(x, t)\) via the gradient,

\[ E(x, t) = -\nabla \phi(x, t) \quad [\text{V/m}]. \quad (5.1.1.2) \]

The electric force on a charge \(q\) is \(F = qE\); thus \(E\) is proportional to the force, and when the medium is conductive, the current density (a flow) is \(J_m = \sigma_s E [\text{A/m}^2]\). The ratio of the potential to the flow is an impedance, so \(\sigma_s\) is a conductance.

Exercise #5

Suppose we are given the vector field in \(\mathbb{R}^3\)

\[ A(x) = [\phi(x), \psi(x), \theta(x)]^T \quad [\text{Wb/m}], \]

where each of the three functions is a scalar field. Then \(A(x) = [x, xy, xyz]^T\) is a legal vector field that has components analytic in \(x\).
Exercise #6
From Maxwell’s equations, the magnetic flux vector is given by
\[ B(x, t) = \nabla \times A(x, t) \quad \text{[Wb/m}^2\text{]} \] (5.1.1.3)

We shall see that this is always true because the magnetic charge \( \nabla \cdot B(x, t) \) must be 0, which is always true in in-vacuo conditions.

To verify that a field is a potential, we check the units [V, A, °C]. However, a proper mathematical definition is that the potential must be an analytic function of \( x \) and \( t \), so that we can operate on it with \( \nabla () \) and \( \nabla \times () \). Note that the divergence of a scalar field is not a legal vector operation.

Feynman (1970b, pp. 14-1 to 14-3) provides an extended tutorial on the vector potential, with many examples.

**Scalar potentials:** Our discussion described the utility of potentials for defining vector fields (e.g., Eqs. 5.1.1.2 and 5.1.1.3). The key distinction between a potential and a scalar field is that potentials have units and thus have a physical meaning. Scalar potentials (i.e., voltage \( \phi(x, t) \) [V], temperature \( T(x, t) \) [°C], and pressure \( \varphi(x, t) \) [pascals]) are physical scalar fields. All potentials are composed of scalar fields, but not all scalar fields are potentials.

For example, the \( \dot{y} \) component of \( E \), \( E_y(x, t) = \dot{y} \cdot E(x, t) \) [V/m], is not a potential. While \( \nabla E_y \) is mathematically defined as the gradient of one component of a vector field, it has no physical meaning (as best I know).

**Vector potentials:** Vector potentials, like scalar potentials, are vector fields with physically meaningful units. They are more complicated than scalar potentials because they are composed of three scalar fields. Vector fields are composed of laminar and rotational flow, which are mathematically described by the fundamental theorem of vector calculus (also called Helmholtz’s decomposition theorem). One superficial but helpful comparison is the momentum of a mass, which may be decomposed into its forward (linear) and rotational momentum.

Since we find it useful to analyze problems using potentials (e.g., voltage) and then take the gradient (e.g., voltage difference) to find the flow (electric field \( E(x, t) \)), the same logic and utility apply when we use the vector potential to describe the magnetic flux (flow) \( B(x, t) \) (Feynman, 1970c). When operating on a scalar potential, we use a gradient, whereas for the vector potential, we operate with the curl.

In Eq. 5.1.1.2 we assumed that the magnetic flux vector \( B(x) \) was static, and thus \( E(x, t) \) is the gradient of the time-dependent voltage \( \phi(x, t) \). However, when the magnetic field is dynamic (not static), Eq. 5.1.1.2 is not valid due to magnetic induction: A voltage induced into a loop of wire is proportional to the time-varying flux cutting across that loop of wire. This is known as the Ampere-Maxwell law. In the static case the induced voltage is zero.

Thus the electric field strength includes both scalar potential \( \phi(x, t) \) and magnetic flux vector potential \( A(x, t) \) components, while the magnetic field strength depends only on the magnetic potential.

### 5.1.2 Gradient \( \nabla \), divergence \( \nabla \cdot \), curl \( \nabla \times \), and Laplacian \( \nabla^2 \)

Three key vector differential operators are used in linear partial differential equations, such as the wave and diffusion equations. All of these begin with the \( \nabla \) operator:
\[
\nabla = \hat{x} \frac{\partial}{\partial x} + \hat{y} \frac{\partial}{\partial y} + \hat{z} \frac{\partial}{\partial z}.
\]

As outlined in Table 5.1, the official name of this operator is nabla. It has three basic uses: (1) the gradient of a scalar field, (2) the divergence of a vector field, and (3) the curl of a vector field. The shorthand notation \( \nabla \phi(x, t) = (\hat{x} \partial_x + \hat{y} \partial_y + \hat{z} \partial_z) \phi(x, t) \) is convenient.\(^1\)

#### Gradient:
As shown in Fig. 5.1, the gradient transforms a complex scalar field \( \Phi(x, s) \in \mathbb{C} \) into a vector field (\( \mathbb{C}^3 \))
\[
\nabla \Phi(x, s) = \left( \hat{x} \frac{\partial \Phi}{\partial x} + \hat{y} \frac{\partial \Phi}{\partial y} + \hat{z} \frac{\partial \Phi}{\partial z} \right) \Phi(x, s)
\]
\[
= \hat{x} \frac{\partial \Phi}{\partial x} + \hat{y} \frac{\partial \Phi}{\partial y} + \hat{z} \frac{\partial \Phi}{\partial z}.
\]

\(^1\)https://en.wikipedia.org/wiki/Del_in_cylindrical_and_spherical_coordinates
CHAPTER 5. STREAM 3B: VECTOR CALCULUS

Table 5.1: The three vector operators manipulate scalar and vector fields. The gradient converts scalar fields into vector fields. The divergence maps vector fields to scalar fields. The curl maps vector fields to vector fields. Four second-order operators (for example GoD and DoG) are defined in Sec. 5.6.6 (p. 200-200).

<table>
<thead>
<tr>
<th>Name</th>
<th>Input</th>
<th>Output</th>
<th>Operator</th>
<th>Mnemonic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gradient</td>
<td>Scalar</td>
<td>Vector</td>
<td>( \nabla () )</td>
<td>grad</td>
</tr>
<tr>
<td>Divergence</td>
<td>Vector</td>
<td>Scalar</td>
<td>( \nabla \cdot () )</td>
<td>div</td>
</tr>
<tr>
<td>Laplacian</td>
<td>Scalar</td>
<td>Scalar</td>
<td>( \nabla \cdot \nabla = \nabla^2 () )</td>
<td>DoG</td>
</tr>
<tr>
<td>Wedge</td>
<td>Vector</td>
<td>Scalar</td>
<td>( \nabla \wedge () )</td>
<td>wedge</td>
</tr>
<tr>
<td>Curl</td>
<td>Vector</td>
<td>Vector</td>
<td>( \nabla \times () )</td>
<td>curl</td>
</tr>
<tr>
<td>Little God</td>
<td>Vector</td>
<td>Vector</td>
<td>( \nabla^2 () = \nabla (\nabla \cdot ()) )</td>
<td>GoD</td>
</tr>
<tr>
<td>Bull-DoG</td>
<td>Vector</td>
<td>Vector</td>
<td>( \nabla^2 () = \nabla \cdot \nabla () )</td>
<td>Dog</td>
</tr>
<tr>
<td>Curl of Curl</td>
<td>Vector</td>
<td>Vector</td>
<td>( \nabla \times \nabla \times () = \nabla^2 () - \nabla^2 () )</td>
<td>CoC</td>
</tr>
<tr>
<td>Div of Curl</td>
<td>Vector</td>
<td>0</td>
<td>( \nabla \cdot \nabla \times () )</td>
<td>DoC</td>
</tr>
<tr>
<td>Curl of Grad</td>
<td>Scalar</td>
<td>0</td>
<td>( \nabla \times \nabla () )</td>
<td>CoG</td>
</tr>
</tbody>
</table>

The gradient may also be factored into a unit vector \( \hat{n} \), as defined in Fig. 5.1, that gives the direction of the gradient, and the gradient’s length \( ||\nabla|| \), defined in terms of the norm of the gradient. Thus the gradient of \( \Phi(x) \) may be written in “polar coordinates” as \( \nabla \Phi(x) = ||\nabla \Phi|| \hat{n} \), which leads to the unit vector

\[
\hat{n} = \frac{\nabla (\Phi(x))}{||\nabla \Phi||}.
\]

Consider the paraboloid \( z = 1 - (x^2 + y^2) \) as the potential, with isopotential circles of constant \( z \) that have a radius of zero at \( z = 1 \) and unit radius at \( z = 0 \). The negative gradient

\[
E(x) = -\nabla z(x, y) = 2(x\hat{x} + y\hat{y} + 0\hat{z})
\]

is \( \perp \) to the circles of constant radius (constant \( z \)) and thus points in the direction of the radius.

A skier in free fall on this surface would be the first one down the hill. Normally skiers try to stay close to the isoclines (not in the direction of the gradient) so they can stay in control. If you ski on an isocline, you must walk, since there is no pull due to gravity.

Divergence:

The divergence of a vector field results in a scalar field. For example, the divergence of the electric field flux vector \( \mathbf{D}(x) \) [C/m²] equals the scalar field charge density \( \rho(x) \) [C/m³]:

\[
\nabla \cdot \mathbf{D}(x) \equiv (\frac{\partial}{\partial x} + \frac{\partial}{\partial y} + \frac{\partial}{\partial z}) \cdot \mathbf{D}(x) = \frac{\partial D_x}{\partial x} + \frac{\partial D_y}{\partial y} + \frac{\partial D_z}{\partial z} = \rho(x).
\]  

(5.1.2.4)

Thus the divergence is analogous to the scalar (dot) product (e.g., \( A \cdot B \)) between two vectors.

Recall that the voltage is the line integral of the electric field,

\[
V(a) - V(b) = \int_a^b \mathbf{E}(x) \cdot dx = -\int_a^b \nabla V(x) \cdot dx = -\int_a^b \frac{dV}{dx} dx,
\]  

(5.1.2.5)

which is simply the fundamental theorem of calculus (see p. 135). In a charge-free region, this integral is independent of the path from \( a \) to \( b \), which is a property of a conservative system.

When we work with guided waves (narrow tubes of flux) having rigid walls that block flow, such that the diameter is small compared with the wavelength (Postulate P10, p. 122), the divergence simplifies to

\[
\nabla \cdot \mathbf{D}(x) = \nabla_r D_r = \frac{1}{A(r)} \frac{\partial}{\partial r} A(r) D_r(r),
\]  

(5.1.2.6)

where \( r \) is the distance down the horn (range variable), \( A(r) \) is the area of the isoresponse surface as a function of the range \( r \), and \( D_r(r) \) is the radial component of vector \( \mathbf{D} \) as a function of the range \( r \). In spherical, cylindrical, and rectangular coordinates, Eq. 5.1.2.6 provides the correct expression (Table 5.2, p. 190).
Properties of the divergence: The divergence is a direct measure of the flux density of the vector field. A vector field is said to be incompressible if the divergence of that field is zero. It is therefore compressible when the divergence is nonzero [e.g., \( \nabla \cdot \mathbf{D}(x, s) = \rho(x, s) \)].

For example, compared to air, water is considered to be incompressible. The stiffness of a fluid (i.e., the bulk modulus) is a measure of its compressibility. At very low frequencies, air may be treated as incompressible (like water), since as \( s \to 0 \),

\[
-\nabla \cdot \mathbf{u}(x, s) = \frac{s}{\eta_o f_o} \mathbf{F}(x, s) \to 0.
\]

The definition of compressible depends on the wavelength in the medium, so the term must be used with some awareness of the frequencies being in the analysis. As a rule of thumb, if the wavelength \( \lambda = c_o/f \) is much larger than the size of the system, the medium may be modeled as an incompressible fluid.

Curl:
The curl (\( \nabla \times \)) takes a vector in \( \mathbb{R}^3 \) into a second vector in \( \mathbb{R}^3 \). For example, in the case of fluids, the Vorticity is defined as \( \mathbf{\Omega} = \nabla \times \mathbf{v} \), and rotation as \( \mathbf{\Omega} = \omega/2 \). The curl is a measure of the rotation of a vector field in a plane about the axis perpendicular to that plane. In the case of water, it corresponds to the angular momentum, such as in a whirlpool, or in air, a tornado. A spinning solid is called a top which falls over if not spinning. But once spinning, it can stably stand on its pointed tip. These systems are stable due to conservation of angular momentum.

The curl and the divergence are both key when we write out Maxwell’s four equations there is little hope of starting from a scalar field, the gradient produces a vector, which is then operated on by the divergence to take the output of the gradient back to a scalar field. Thus the Laplacian transforms a scalar field back to a scalar field. Since the Laplacian does so much work we nickname it DoG for Div of Grad.

For example, \( \nabla \times \mathbf{H}(x, s) \in \mathbb{C}^3 [\text{A/m}] \) into a complex vector field into a current density \( \mathbf{C}(x, s) \in \mathbb{C}^2 [\text{A/m}^2] \):

\[
\nabla \times \mathbf{H}(x, s) \equiv \begin{vmatrix}
\hat{x} & \hat{y} & \hat{z} \\
\partial_x & \partial_y & \partial_z \\
H_x & H_y & H_z
\end{vmatrix} = \mathbf{C}(x, s). \quad [\text{A/m}^2] \quad (5.1.2.7)
\]

The notation \( |\cdot| \) indicates the determinant (Appendix A.3.1, p. 227), \( \partial_x \) is shorthand for \( \partial/\partial x \), and \( \mathbf{H} = [H_x, H_y, H_z]^{T} \).

Exercise #7
If we let \( \mathbf{H} = -yx \hat{x} + x \hat{y} + z \hat{z} \), \( \nabla \times \mathbf{H} = 2 \hat{z} \), thus \( \mathbf{H} \) has a constant rotation; when \( \mathbf{H} = 0 \hat{x} + 0 \hat{y} + z^2 \hat{z} \), \( \nabla \times \mathbf{H} = 0 \) has a curl of zero and thus is irrotational.

There are simple rules that precisely govern when a vector field is rotational versus irrotational, and compressible versus incompressible. These classifications are dictated by Helmholtz’s theorem, the fundamental theorem of vector calculus (Eq. 5.8.0.5, p. 207).

A special case of the curl is the two-dimensional differential wedge products\(^2\)

\[
\nabla_z \wedge \mathbf{H}(x, t) = \begin{vmatrix}
\partial_y & \partial_z \\
H_y & H_z
\end{vmatrix} = C_z(x, s). \quad [\text{A/m}^2].
\]

The curl is made up of three such differential wedge products.\(^3\)

Laplacian \( \nabla^2() \):
The Laplacian operator \( \nabla^2 \equiv \nabla \cdot \nabla \) (Table 5.1, page 172) is defined as the divergence of the gradient

\[
\nabla^2 \equiv \frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} + \frac{\partial^2}{\partial z^2} . \quad (5.1.2.8)
\]

Since the Laplacian does so much work we nickname it DoG for Div of Grad.

Starting from a scalar field, the gradient produces a vector, which is then operated on by the divergence to take the output of the gradient back to a scalar field. Thus the Laplacian transforms a scalar field back to a scalar field. We have seen the Laplacian before when we defined complex analytic functions (Eq. 4.2.2.10, p. 137).

\(^2\)http://en.wikipedia.org/wiki/Triple_product#As_an_exterior_product

\(^3\)This notation suggests that \( |\nabla \cdot \mathbf{E} + j \nabla \times \mathbf{E}|^2 = |\nabla \cdot \mathbf{E}|^2 + |\nabla \times \mathbf{E}|^2 \), which seems to be related to Helmholtz’s theorem. For example since \( \mathbf{E} = -\nabla \Phi + \nabla \times \mathbf{A} \). Then \( \nabla \cdot \mathbf{E} = -\nabla^2 \Phi \) and \( \nabla \times \mathbf{E} = -s \mathbf{B} = -s \nabla \times \mathbf{A} \).
For example, taking the divergence of the simple example Eq. 5.1.1.1 results in the Laplacian of the voltage
\[ \nabla^2 \Phi(x) = -V_o(\delta(x) - \delta(x-1)) = 0 \]
for \(0 < x < 1\). Thus this example obeys Laplace’s equation.

One classic example of the Laplacian is a voltage scalar field \(\Phi(x)\) [V], which results in the electric field vector
\[ E(x) = [E_x(x), E_y(x), E_z(x)]^T = -\nabla \Phi(x) \quad [\text{V/m}] \]
When this is scaled by the permittivity, we obtain the electric flux \(D = \epsilon_o E\) [C/m²], the charge density per unit area. Here \(\epsilon_o\) [F/m] is the vacuum permittivity, which is \(\approx 8.85 \times 10^{-12}\) [F/m].

Taking the divergence of \(D\) results in the charge density \(\rho(x)\) [C/m³] at \(x\)
\[ \nabla \cdot D = \nabla^2 \Phi(x) = \rho(x). \]
Thus the Laplacian of the voltage, scaled by \(\epsilon_o\), results in the local charge density.

Another classic example of the Laplacian is an acoustic pressure field \(p(x, t)\) [Pa], which defines a vector force density \(f(x, t) = -\nabla p(x, t)\) [N/m²] (Eq. 5.2.1.5, p. 182). When this force density [N/m²] is integrated over an area, the net radial force [N] is
\[ F_r = -\int_S \nabla p(x) d\mathbf{x} \quad [\text{N}]. \quad (5.1.2.9) \]
An inflated balloon with a static internal pressure of 3 [atm], in an ambient pressure of 1 [atm] (sea level), forms a sphere due to the elastic nature of the rubber, which acts as a stretched spring under tension. The net force on the surface of the balloon is its area times the pressure drop of 2 atm across the surface. Thus the static pressure is
\[ p(x) = 3u(r_o - r) + 1 \quad [\text{Pa}], \]
where \(u(r)\) is a step function of the radius \(r = ||x|| > 0\), centered at the center of the balloon, having radius \(r_o\).

Taking the gradient gives the negative⁴ of the radial force density (i.e., perpendicular to the surface of the balloon):
\[ -f_r(r) = \nabla p(x) = \frac{\partial}{\partial r} 3u(r_o - r) + 1 = -2(\delta(r_o) - \delta(r_o - r)) \quad [\text{Pa}]. \]
This equation describes a static pressure that is 1 [atm] \((10^5\text{ [Pa]})\) outside the balloon and 3 [atm] inside. The net positive force density is the negative of the gradient of the static pressure.

Finally, taking the divergence of the force produces a double delta function at \(r = r_o\)—namely, \(\nabla^2 p(x) = -2\delta(2)(r_o - r)\), where 2 is the pressure drop across the balloon. If we take the thickness of the rubber \((l\) [m]) into account, then \(\nabla^2 p = -2(\delta(r_o) - \delta(r_o - l))\).

**Vector Laplacian \(\nabla^2()\):**

A second form of the Laplacian is the vector Laplacian \(\nabla^2()\), defined as the divergence of the gradient \(\nabla^2() \equiv \nabla \cdot \nabla()\), thus nicknamed **Bull-Dog**, operates on a vector to produce a vector (Table 5.1, page 172). We shall first see this when working with Maxwell’s equations.

### 5.1.3 Scalar Laplacian operator in \(N\) dimensions

In general, it may be shown that in \(N = 1,2,3\) dimensions (Sommerfeld, 1949, p. 227),
\[ \nabla^2 \Phi = \frac{1}{r^{N-1}} \frac{\partial}{\partial r} \left( r^{N-1} \frac{\partial \Phi}{\partial r} \right). \quad (5.1.3.10) \]
For each value of \(N\), the area \(A(r) = A_o r^{N-1}\). This result will turn out to be useful when we work with the Laplacian in one, two, and three dimensions. This naturally follows from the Webster horn equation (WHEN) (Eq. 5.2.2.10, p. 183).

---

⁴The force is pointing out, stretching the balloon.
Exercise #8
When \( N = 3 \) (i.e., spherical geometry),
\[
\nabla^2 r P \equiv \frac{1}{r^2} \partial_r r^2 \partial_r r P = \frac{1}{r} \frac{\partial^2}{\partial r^2} r^2 r P,
\]
resulting in the general d’Alembert solutions (Eq. 4.4.0.1 p. 142) for the spherical wave equation,
\[
P^{\pm}(r, s) = \frac{1}{r} e^{\mp \kappa(s)r}.
\]

Exercise #9
Prove the result of Example # 9 by expanding Eqs. 5.1.3.11 and 5.1.3.12 using the chain rule.

Solution: Expanding Eq. 5.1.3.11:
\[
\frac{1}{r^2} \partial_r r^2 \partial_r r P = \frac{1}{r^2} \left( 2r + r \partial_r r P \right) \partial_r r P
= \frac{2}{r} P_r + P_{rr}.
\]
Expanding Eq. 5.1.3.12, we obtain
\[
\frac{1}{r} \partial_r r P = \frac{1}{r} \partial_r \left( P + r P_r \right)
= \frac{1}{r} \left( P_r + P_r + r P_{rr} \right)
= \frac{2}{r} P_r + P_{rr}.
\]
Thus the two are equivalent.

Summary: The radial component of the Laplacian in spherical coordinates (Eq. 5.1.3.11) simplifies to
\[
\nabla^2 \varphi(x) = \frac{1}{r^2} \frac{\partial}{\partial r} r^2 \frac{\partial}{\partial r} \varphi(x) = \frac{1}{r} \frac{\partial^2}{\partial r^2} r \varphi(x).
\]
Since \( \nabla^2 = \nabla \cdot \nabla \), it follows that the net force \( f(x) = [F_r, 0, 0]^T \), Eq. 5.1.2.9 in spherical coordinates has a radial component \( F_r \) and angular components of zero. Thus the force across a balloon may be approximated by a delta function across the thin sheet of stretched rubber.
We can extended the preceding example in an interesting way to the case of a rigid hose, a rigid tube, that terminates on the right in an elastic medium (the above example of a balloon), such as an automobile tire. On the far left let’s assume there is a pump injecting the fluid into the rigid hose. Consider two different fluids: air and water. Air is treated as a compressible fluid, whereas water is incompressible. However, such a classification is relative, determined by the relative compliance of the balloon (i.e., tire) at the relatively rigid pump and hose.
This is a special case of a more general situation: When a fluid is treated as incompressible (rigid), the speed of sound becomes infinite, the wave equation is an invalid description. In this case the motion is best approximated by Laplace’s equation. This represents the transition from short to long wavelengths, from wave propagation having delay, to quasi-statics, having no delay.
This example may be modeled as either an electrical or mechanical system. If we take the electrical analog, the pump is a current source, injecting charge \( Q_o \) into the hose, which being rigid cannot expand (has a fixed volume). The hose may be modeled as a resistor and the tire as a capacitor \( C_o \), which fills with charge as it is delivered via the resistor, from the pump. The capacitor obeys the same equation as Hooke’s law for a spring, \( F = K_o \Delta \), where \( K_o \) is the stiffness of the spring, \( C_o = 1/K_o \) is the spring’s compliance, and \( \Delta \) is the displacement. In electrical terms, \( Q_o = C_o \Phi \) where \( \Phi \) is the voltage, which acts like a force \( F; Q_o \) is the charge, which plays the role of the mass of the fluid. The charge \( Q \) is conserved, just as the mass of the fluid is conserved (they cannot be created or destroyed).
The flow of the fluid is called the flux, which is the general term for the mass flow, heat or electrical current. The two equations may be rewritten directly in terms of the force, either \( F, \Phi \), and flow, either \( I = dQ/dt \), the electrical current, or \( J = dM/dt \), the mass flux. In terms of impedance,

\[
I = C_o \frac{d \Phi}{dt} \quad [A]\] (5.1.3.13)

for the electrical analog, and

\[
J = C_o \frac{d F}{dt} \quad [kgm-s/m].\] (5.1.3.14)

It is common to treat the stiffness of the balloon, which acts as a spring, as a compliance \( C_o = 1/K_o \), in which case the impedance reduces to a single form. The impedance \( Z \) is defined in the frequency domain as the ratio of the generalized force over the generalized flow

\[
Z(s) = \frac{1}{sC_o} \quad [ohms].
\]

In the case of the mechanical system \( Z_m(s) \equiv F/J \), while for the electrical system, \( Z_e(s) \equiv \Phi/I \). It is conventional to use the unit [ohms] when working with any impedance, allowing for uniform terminology for different physical situations and forms of impedance. This greatly simplifies the notation.

While the two systems are very different in their physical realization, they are mathematically equivalent, forming a perfect analog. The formula for the impedance is typically expressed in \( s \), the Laplace frequency, which of course is the \( LT \) of the time variables. In the frequency domain Ohm’s law becomes Eq. 5.1.3.14 for the case of a mechanical compliance \( C_o = 1/K_o \) and Eq. 5.1.3.13 for the electrical capacitor \( C \).

The final solution of this system is solved in the frequency domain. The impedance seen by the source is the sum of the resistance \( R \) and the impedance of the load, giving

\[
Z = R + \frac{1}{sC}.
\]

This results in a simple relationship between the force and the flow, as determined by the action of the source on the load \( Z(s) \). The results are given in terms of the voltage across the compliance in terms of the voltage \( \Phi_e \) (or current \( I_e \)) due to the source. Given some algebra, the voltage across the compliance \( \Phi_e \), divided by the voltage of the source, is

\[
\frac{\Phi_e}{\Phi_{\text{source}}} = \frac{R}{R + 1/sC}.
\]

Thus the problem reduces to some algebra in the frequency domain. The time domain response is found by taking the inverse \( LT \), which in this case has a simple pole at \( s_p = 1/RC \). Cauchy’s residue theorem (p. 152) gives the final answer, which describes how the voltage across the compliance builds exponentially with time, from zero to the final value. Given the voltage, the current may also be computed as a function of time. This then represents the entire process of either blowing up a balloon, or charging a capacitor, the difference being only the physical notation, as the math is identical.

Note that the differential equation is first-order in time, which in frequency means the impedance has a single pole. Thus the equation for charging a capacitor or pumping up a balloon describes a diffusion process. If we had taken the impedance of the mass of the fluid in the hose into account, we would have a lumped-parameter model of the wave equation with a second-order system. This is mathematically the same as the homework assignment about train cars (masses) connected by springs (5.4, Homework DE-3, problem #2).

**Exercise #10**

The voltage

\[
\phi(x,t) = e^{-\kappa \cdot x} u(t - x/c) \leftrightarrow \frac{1}{s} e^{-\kappa \cdot x} \quad [V]\] (5.1.3.15)

represents one of d’Alembert’s solution (Eq. 4.4.0.1, p. 142) of the wave equation (Eq. 3.1.0.5, p. 63) as well as an eigenfunction of the gradient operator \( \nabla \). From the definition of the scalar (dot) product of two vectors (Fig. 3.4, p. 94),

\[
\kappa \cdot x = \kappa_x x + \kappa_y y + \kappa_z z = ||\kappa|| \ · ||x|| \ · \cos \theta_{kx},
\]

where \( ||\kappa|| = \sqrt{\kappa_x^2 + \kappa_y^2 + \kappa_z^2} \) and \( ||x|| = \sqrt{x^2 + y^2 + z^2} \) are the lengths of vectors \( \kappa \) and \( x \), and \( \theta_{kx} \) is the angle between them. As before, \( s = \sigma + \omega j \) is the Laplace frequency.
To keep things simple, we let \( \kappa = [\kappa_x, 0, 0]^T \) so that \( \kappa \cdot x = \kappa_x x \hat{x} \). We shall soon see that \( ||\kappa|| = 2\pi/\lambda \) follows from the basic relationship between a wave’s radian frequency \( \omega = 2\pi f \) and its wavelength \( \lambda \):

\[
\omega \lambda = c_0. \tag{5.1.3.16}
\]

As the frequency increases, the wavelength becomes shorter. This key relationship may have been first researched by Galileo in about 1564, followed by Mersenne in about 1627\(^5\) (Fig. 1.5, p. 19).

Exercise #11
Show that Eq. 5.1.3.15 is an eigenfunction of the gradient operator \( \nabla \).

Solution: Taking the gradient of \( \phi(x,t) \) gives

\[
\nabla e^{-\kappa \cdot x} u(t) = -\nabla \kappa \cdot x e^{-\kappa \cdot x} u(t) = -\kappa e^{-\kappa \cdot x} u(t),
\]

or in terms of \( \phi(x,t) \),

\[
\nabla \phi(x,t) = -\kappa \phi(x,t) \leftrightarrow -\frac{s}{c} e^{-\kappa \cdot x}.
\]

Thus \( \phi(x,t) \) is an eigenfunction of \( \nabla \), having the vector eigenvalue \( \kappa \). As before, \( \nabla \phi \) is proportional to the current, since \( \phi \) is a voltage, and the ratio (i.e., the eigenvalue), may be thought of as a mass, analogous to the impedance of a mass (or inductor). In general, the units provide the physical interpretation of the eigenvalues and their spectra.

Exercise #12
Compute \( \hat{n} \) for \( \phi(x,s) \) as given by Eq. 5.1.3.15.

Solution: \( \hat{n} = \kappa / ||\kappa|| \) represents a unit vector in the \( \kappa \) direction.

Exercise #13
If the sign of \( \kappa \) is negative, what are the eigenvectors and eigenvalues of \( \nabla \phi(x,t) \)?

Solution:

\[
\nabla e^{-\kappa \cdot x} u(t) = -\kappa \cdot \nabla (x)e^{-\kappa \cdot x} u(t) = -\kappa e^{-\kappa \cdot x} u(t).
\]

Nothing changes other than the sign of \( \kappa \). Physically this means the wave is traveling in the opposite direction, corresponding to the forward and retrograde d’Alembert waves.

Exercise #14
Find the velocity \( v(t) \) of an electron in a field \( E \).

Solution: From Newton’s 2nd law, \(-qE = m_e \dot{v}(t) \) [Nt], where \( m_e \) is the mass of the electron. Thus we must solve this first-order differential equation to find \( v(t) \). This is easily done in the frequency domain \( v(t) \leftrightarrow V(\omega) \).

Role of Potentials: Note that the scalar fields (e.g., temperature, pressure, voltage) are all scalar potentials, summarized in Table 3.2 (p. 114). In each case the gradient of the potential results in a vector field, just as in the electric case above (Eq. 5.1.1.2).

It is important to understand the physical meaning of the gradient of a potential, which is typically a generalized force (electric field, acoustic force density, temperature flux), that in turn generates a flow (current, velocity, heat flux). The ratio of the potential over flow determines the impedance. Four examples follow:

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\(^5\)https://www-history.mcs.st-and.ac.uk/Biographies/Mersenne.html

“In the early 1620s, Mersenne listed Galileo among the innovators in natural philosophy whose views should be rejected. However, by the early 1630s, less than a decade later, Mersenne had become one of Galileo’s most ardent supporters.” (Garber, 2004)
1. The voltage drop across a resistor causes a current to flow, as described by Ohm’s law. The difference in voltage between two points is a crude form of gradient when the frequency \( f \) [Hz] is low, such that the wavelength is much larger than the distance between the two points. This is the essence of the quasistatic approximation (Postulate P10, p. 122).

2. The gradient of the pressure gives rise to a force density in the fluid medium (air, water, oil, etc.), that causes a flow (velocity vector) in the medium.

3. The gradient of the temperature also causes a flow of heat, that is proportional to the thermal resistance, given Ohm’s law for heat (Feynman, 1970b, p. 3-7).

4. When a solution contains charged ions, it defines an electrochemical Nernst potential \( N(x, t) \) (Fermi, 1936; Scott, 2002). This electrochemical potential is similar to a voltage or temperature field, the gradient of which defines a virtual force on the charged ions, resulting in a current.

Thus in the above examples there is a potential, the gradient of which is a force, that when applied to the medium (an impedance) causes a flow (flux or current) proportional to that impedance due to the medium. These general concepts are worthy of some thought. The product of the force and flow is a power.

Exercise #15
Show that the integral of Eq. 5.1.1.2 is an antiderivative.

**Solution:** We use the definition of the antiderivative given by the FTC (Eq. 4.2.0.2, p. 135):

\[
\phi(x, t) - \phi(x_o, t) = \int_{x_o}^{x} E(x, t) \cdot dx
\]

\[
= - \int_{x_o}^{x} \nabla \phi(x, t) \cdot dx
\]

\[
= - \int_{x_o}^{x} \left( \hat{x} \frac{\partial \phi}{\partial x} + \hat{y} \frac{\partial \phi}{\partial y} + \hat{z} \frac{\partial \phi}{\partial z} \right) \phi(x, t) \cdot dx
\]

\[
= - \int_{x_o}^{x} \left( \hat{x} \frac{\partial \phi}{\partial x} + \hat{y} \frac{\partial \phi}{\partial y} + \hat{z} \frac{\partial \phi}{\partial z} \right) \cdot (\hat{x} dx + \hat{y} dy + \hat{z} dz)
\]

\[
= - \int_{x_o}^{x} \frac{\partial \phi}{\partial x} dx - \int_{y_o}^{y} \frac{\partial \phi}{\partial y} dy - \int_{z_o}^{z} \frac{\partial \phi}{\partial z} dz
\]

\[
= - \int_{x_o}^{x} d\phi(x, t)
\]

\[
= - \left( \phi(x, t) - \phi(x_o, t) \right).
\]

This may be verified by taking the gradient of both sides

\[
\nabla \phi(x, t) - \nabla \phi(x_o, t) = -\nabla \int_{x_o}^{x} E(x, t) \cdot dx = E(x, t).
\]

If we apply the FTC, the antiderivative must be \( \phi(x, t) = E_x x \hat{x} + 0 \hat{y} + 0 \hat{z} \). This same point is made by Feynman (1970b, p. 4-1, Eq. 4.28).

Given that the force on a charge is proportional to the gradient of the potential, this exercise shows that the integral of the gradient depends on only the end points, the work done in moving a charge depends on only the limits of the integral, which is the definition of a conservative field but which holds only in the ideal case where \( E \) is determined by Eq. 5.1.1.2—that is the medium has no friction (there are no other forces on the charge).

The conservative field: An important question is When is a field conservative? A field is conservative when the work done by the motion is independent of the path of the motion. Thus the conservative field is related to the FTC, which states that the integral of the work depends on only the end points.

A more complete answer must await the introduction of the fundamental theorem of vector calculus (Eq. 5.8.0.5, p. 207). A few examples provide insight:
Exercise #16
The gradient of a scalar potential, such as the voltage (Eq. 5.1.1.2), defines the electric field, which drives a current (flow) across a resistor (impedance). When the impedance is infinite, the flow is zero, leading to zero power dissipation. When the impedance is lossless, the system is conservative.

Exercise #17
At audible frequencies the viscosity of air is quite small and thus, for simplicity, it may be taken as zero. However, when the wavelength is small (e.g., at 100 [kHz] $\lambda = c_0/f = 345/10^5 = 3.45$ [mm]) the lossless assumption breaks down, resulting in a significant propagation loss. When the viscosity is taken into account, the field is lossy and thus the field is no longer conservative.

Exercise #18
If a temperature field is a time-varying constant (i.e., $T(x,t) = T_o(t)$), there is no “heat flux,” since $\nabla T_o(t) = 0$. When there is no heat flux [i.e., flux, or flow], there is no heat power, since the power is the product of the force and the flow.

Exercise #19
The force of gravity is given by the gradient of Newton’s gravitational potential (Eq. 3.1.0.1, p. 61)

$$F = -\nabla_r \phi_G(r) = -\frac{1}{r^2}.$$  

Historically speaking, $\phi_N(r)$ was the first conservative field, used to explain the elliptic orbits of the planets around the sun. Galileo’s law says that bodies fall with constant acceleration, giving rise to a parabolic path and a time of fall proportional to $t^2$. This behavior of falling objects directly follows from the Galilean potential:

$$\phi_G(r) = \frac{1}{(r-r_o)} = \frac{-r_o}{1-r/r_o} = -r_o(1 - r/r_o + (r/r_o)^2 + \cdots) \approx -r_o, \quad r \ll r_o,$$

which, given the large radius $r_o$ of the earth and the small distance of the object from the surface of the earth $r - r_o$, is equal to the distance above the ground. Thus Galileo’s law says that the force a falling body sees is constant:

$$F_G = -\nabla_r \phi_G(r) = 1.$$  

This can be scaled by a constant to account for the magnitude of the gravitational force.

Exercise #20
Galileo discovered that the height of a falling object is proportional to the square of the time it falls. Based on Newton’s follow-up analysis, today we would say this height $h(t)$ is

$$h(t) = \frac{1}{2} m G_o (t-t_o)^2 \text{ [m]},$$

where $m$ is the object’s mass and $G_o$ is the gravitational constant for the earth at its surface $r_o$. Show that $h(t)$ directly follows from the potential $\phi_G = r_o - r$. This formula applies if you toss a ball into the air, or if you drop it from a high place.

**Solution:** Given Galileo’s potential $\phi_G(r) \approx m G_o (r_o - r)$, show that the force is constant, thus that $\ddot{h}(t) = m G_o$. Given Galileo’s formula for the height $h(t)$, the velocity is $v(t) = \dot{r}(t) = m G_o t$, and the acceleration is $\ddot{r}(t) = m G_o$.

Exercise #21
Find the time that it takes to fall from a distance $r = L$. Namely, solve $h(t) = L$ for the time the object takes to fall the distance $L$.

**Solution:** Setting $t_o = 0$ gives $t^2 = 2L/m G_o$. Thus the time to fall is $T(L) = \sqrt{2L/m G_o}$. •
5.2 Partial differential equations and field evolution

The three main classes of partial differential equations (PDEs) are: elliptic, parabolic, and hyperbolic, distinguished by the order of the time derivative. These categories seem to have little mathematical utility (the categories appear as labels).

The Laplacian $\nabla^2$: In the most important case the space operator is the Laplacian $\nabla^2$, the definition of which depends on the dimensionality of the waves— that is, the coordinate system being used. We first discussed the Laplacian as a 2D operator on page 136, when we studied complex analytic functions, and again on page 171. An expression for $\nabla^2$ for one, two, and three dimensions was provided as Eq. 5.1.3.10 (p. 174). In 3D rectangular coordinates, it is defined as (see p. 173)

$$\nabla^2 T(x) = \left( \frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} + \frac{\partial^2}{\partial z^2} \right) T(x). \quad (5.2.0.1)$$

The Laplacian operator is ubiquitous in mathematical physics, starting with simple complex analytic functions (Laplace’s equation) and progressing to Poisson’s equation, the diffusion equation, and finally the wave equation. Only the wave equation expresses delay. The diffusion equation “wave” has an instantaneous spread (the effective “wavefront” velocity is infinite, yet the wavelength is long; it’s not a traveling wave).

Examples of elliptic, parabolic, and hyperbolic equations follow:

1. Laplace’s equation: The equation

$$\nabla^2 \Phi(x) = 0 \quad (5.2.0.2)$$

that describes, for example, the voltage inside a closed chamber that has various voltages on the walls, or the steady-state temperature within a closed container given a specified temperature distribution on the walls. There are no dynamics to the potential, even when it is changing, since the potential instantaneously follows the potential on the walls.

2. Poisson’s equation: In the steady state, the diffusion equation degenerates to either Poisson’s or Laplace’s equation; both are classified as elliptic equations (second-order in space, zero-order in time). As in the diffusion equation, the evolution has a wave velocity that is functionally infinite. For example,

$$\nabla^2 \Phi(x, t) = \rho(x, t)$$

holds for gravitational fields or the voltage around a charge.

3. Fourier diffusion equation: Equation 5.2.0.3 describes the evolution of the scalar temperature $T(x, t)$ (a scalar potential), gradients of solution concentrations (i.e., ink in water), and Brownian motion. Diffusion is first-order in time, which is categorized as parabolic (first-order in time, second-order in space). When these equations are Laplace transformed, diffusion has a single real root, resulting in a real solution (e.g., $\rho \in \mathbb{R}$). There is no wavefront in the case of the diffusion equation. As soon as the source is turned on, the field is nonzero at every point in the bounded container. As an example

$$\nabla^2 T(x, t) = \kappa \frac{\partial T(x, t)}{\partial t} \leftrightarrow s \kappa T(x, s) \quad (5.2.0.3)$$

describes the temperature $T(x, t) \leftrightarrow T(x, \omega)$, as proposed by Fourier in 1822, or the diffusion of two miscible liquids (Fick, 1855) or Brownian motion (Einstein, 1905). The diffusion equation is not a wave equation, since the temperature wavefront propagates instantaneously. The diffusion equation does a poor job of representing the velocity of molecules banging into each other, since such collisions have a mean free path, and thus the velocity cannot be infinite.

4. Two types of wave equations

(a) Scalar wave equations: Equation 3.1.0.3 (p. 62) describes the evolution of a scalar potential field, such as pressure $\rho(x, t)$ (sound) or the displacement of a string or membrane under tension. The wave equation is second-order in time. When transformed into the frequency domain, the solution has pairs of complex conjugate roots, leading to two real solutions (i.e, $\rho(x, t \in \mathbb{R})$). The wave equation is classified as hyperbolic (second-order in time and space).

(b) Vector wave equations: Maxwell’s equations describe the propagation of the EM electric $\mathbf{E}(x, t)$ and magnetic $\mathbf{H}(x, t)$ field strength vectors, as well as the electric $\mathbf{D}(x, t) = \epsilon_0 \mathbf{E}(x, t)$ and magnetic $\mathbf{B}(x, t) = \mu_0 \mathbf{H}(x, t)$ flux vectors.
Solution evolution: The partial differential equation defines the evolution of the scalar field \( \phi(x, t) \) and temperature \( T(x, t) \), or vector field \( \mathbf{E}, \mathbf{D}, \mathbf{B}, \mathbf{H} \), as functions of space \( x \) and time \( t \). There are two basic categories of field evolution: diffusion and propagation.

1. **Diffusion:** The simplest and easiest PDE example, easily visualized, is a static 6 (time-invariant) scalar temperature field \( T(x) \) [°C]. Just like an impedance or admittance, a field has regions where it is analytic, and for the same reasons, \( T(x, t) \) satisfies Laplace’s equation

\[
\nabla^2 T(x, t) = 0.
\]

Since there is no current when the field is static, such systems are lossless and thus are conservative. When \( T(x, t) \) depends on time (is not static), it is described by the diffusion equation (Eq. 5.2.0.3), a rule for how \( T(x, t) \) evolves with time from its initial state \( T(x, 0) \). The constant \( \kappa_o \) is called the thermal conductivity, which depends on the properties of the fluid in the container, with \( s \kappa_o \) being the thermal admittance per unit area. The conductivity is a measure of how the heat gradients induce heat currents \( \mathbf{J} = -\kappa_o \nabla T \), analogous to Ohm’s law for electricity.

Note that when \( T(x, t \to \infty) = 0 \) the temperature reaches a steady state, \( \mathbf{J} = 0 \) and \( \nabla^2 T = 0 \). This all depends on what is happening at the boundaries. When the wall temperature of a container is a function of time, the internal temperature \( T(x, t) \) will continue to change, but with a delay that depends on the thermal conductivity \( \kappa_o \).

Such a system is analogous to an electrical resistor-capacitor series circuit connected to a battery. For example, the wall temperature (voltage across the battery) represents the potential driving the system. The thermal conductivity \( \kappa_o \) (the electrical resistor) is likewise analogous. The fluid (the electrical capacitor) is being heated (charged) by the heat (charge) flux. In all cases Ohm’s law defines the ratio of the potential (voltage) to the flux (current). How this happens can only be understood once the solution to the equations has been established. The fluid has a heat capacity analogous to that of an electrical capacitor (Kirchhoff, 1868, 1974).

2. **Propagation:** Pressure and electromagnetic waves are described by a scalar potential (pressure) (Eq. 3.1.0.3, p. 62) and a vector potential (electromagnets) (Eq. 5.7.2.4, p. 204), resulting in scalar and vector wave equations.

All these partial differential equations, scalar and vector wave equations, and the diffusion equation depend on the Laplacian \( \nabla^2 \), which we first saw with the Cauchy–Riemann conditions (Eqs. 4.2.2.10, p. 137).

The Taylor series of \( f(x) \): Next we expand the concept of the Taylor series of one variable to vector \( x \in \mathbb{R}^3 \). Just as we generalized the derivative with respect to a real frequency variable \( \omega \in \mathbb{R} \) to complex frequency \( s = \sigma + \omega j \in \mathbb{C} \), here we generalize the derivative with respect to \( x \in \mathbb{R} \) to the vector \( x \in \mathbb{R}^3 \).

Since the scalar field is analytic in \( x \), it is a perfect place to start. Assuming we have carefully defined the Taylor series (Eq. 3.2.2.9, p. 76) in one and two (Eq. 4.2.2.7, p. 136) variables, the Taylor series of \( f(x) \) in \( x \in \mathbb{R}^3 \) about \( x = 0 \) may be defined as

\[
f(x + \delta x) = f(x) + \nabla f(x) \cdot \delta x + \frac{1}{2!} \sum_{k=1}^{3} \sum_{l=1}^{3} \frac{\partial^2 f(x)}{\partial x_k \partial x_l} \delta x_k \delta x_l + \text{HOT}
\]

where HOT stands for Higher Order Terms (Greenberg, 1988, p. 639). From this definition it is clear that the gradient is the generalization of the second term in the 1D Taylor series expansion.

Summary: For every potential \( \phi(x, t) \) there exists a force density \( f(x, t) = -\nabla \phi(x, t) \), proportional to the potentials, that drives a generalized flow \( \mathbf{u}(x, t) \). If the normal components of the force and flow are averaged over a surface, the mean force and volume flow (i.e., volume velocity for the acoustic case) are defined. In such cases the impedance \( Z(s) = F(x, s)/|\nabla F(x, s)| \) is the net force through the surface force over the net flow, and Gauss’s law and quasi-statics (Postulate P10, p. 122) come into play (Feynman, 1970a). We call this the generalized impedance.

Assuming linearity (Postulate P2, p. 122), the product of the force and flow is the power, and the ratio (force/flow) is an impedance (Table 3.2, p. 114). This impedance statement is called Ohm’s law, Kirchhoff’s

---

6Postulate P3, p. 122.
laws, Laplace’s law, or Newton’s laws. In the simplest cases, they are all linearized (proportional) complex relationships between a force and a flow. Very few impedance relationships are inherently linear over a large range of force or current, but for physically useful levels, they can be treated as linear. Nonlinear interactions require a more sophisticated approach, typically involving numerical methods.

In electrical circuits it is common to define a zero potential ground point that all voltages use as the reference potential. The ground is a useful convention as a simplifying rule, but it obscures the physics and obscures the fact that the voltage is not the force. Rather, the force is the voltage difference, referenced to the ground, which is defined as zero volts. This results in abstracting away (i.e., hiding) the difference in voltage. It seems misleading (more precisely, it is wrong) to state Ohm’s law as the voltage over the current, since Ohm’s law actually says that the voltage difference (i.e., voltage gradient) over the current defines an impedance (Kennelly, 1893).

When we measure the voltage between two points, it is a crude approximation to the gradient based on the quasistatic approximation (Postulate P10). The pressure is also a potential, the gradient of which is a force density, which drives the volume velocity (flow).

In Sec. 5.6.6 we first introduce the fundamental theorem of vector calculus (otherwise known as Helmholtz’ decomposition theorem), which generalizes Ohm’s law to include circulation (e.g., angular momentum, vorticity, and the related magnetic effects). To understand these generalizations in flow, we need to understand compressible and rotational fields (Table 5.3, p. 199), complex analytic functions, and more mathematical physics history.

In summary, it is the difference in the potential (i.e., voltage, temperature, pressure) that is proportional to the flux. This can be viewed as a major simplification of the gradient relationship, justified by the quasistatic assumption (Postulate P10, p. 122).

The roots of the impedance are related to the eigenmodes of the system equations.

### 5.2.1 Scalar wave equation (Acoustics)

In this section we discuss the general solution to the wave equation. The wave equation has two forms: scalar waves (acoustics) and vector waves (electromagnetics). These have an important mathematical distinction but a similar solution space, one scalar and the other vector. We start with the scalar wave equation.

**The scalar wave equation:** A good starting point for understanding PDEs is to explore the scalar wave equation (Eq. 3.1.0.3, p. 62). Thus we limit our analysis to acoustics, the classic case of scalar waves. Acoustic wave propagation was first analyzed mathematically by Isaac Newton (electricity had yet to be discovered) in his famous book *Principia* (1687), in which he first calculated the speed of sound based on the conservation of mass and momentum.

**Early history:** The study of wave propagation begins at least as early as Huygens (ca. 1678), followed soon after (ca. 1687) by Sir Isaac Newton’s calculation of the speed of sound (Pierce, 1981, p. 15). The acoustic variables are the pressure

\[ p(x, t) \leftrightarrow P(x, \omega), \]

and the particle velocity,

\[ u(x, t) \leftrightarrow U(x, \omega). \]

To obtain a wave, we must include two basic components: the stiffness of air and its mass. The two equations are called (1) Newton’s second law \((F = ma)\) and (2) Hooke’s law \((F = kx)\), respectively. In vector form these equations are (1) Euler’s equation (i.e., conservation of momentum density),

\[-\nabla p(x, t) = \rho_o \frac{\partial}{\partial t} u(x, t) \leftrightarrow \rho_o s U(x, s), \quad (5.2.1.5)\]

which assumes the time-average density \(\rho_o\) is independent of time and position \(x\), and (2) the continuity equation (i.e., conservation of mass density),

\[-\nabla \cdot u(x, t) = \frac{1}{\eta_o P_o} \frac{\partial}{\partial t} p(x, t) \leftrightarrow \frac{s}{\eta_o P_o} P(x, s) \quad (5.2.1.6)\]

(Pierce, 1981; Morse, 1948, p. 295). Here \(P_o = 10^5\) Pa is the barometric pressure and \(\eta_o P_o\) is the dynamic (adiabatic) stiffness, with \(\eta_o = 1.4\). Combining Eqs. 5.2.1.5 and 5.2.1.6 (removing \(u(x, t)\)) results in the 3D scalar pressure wave equation

\[ \nabla^2 p(x, t) = \frac{1}{c_o^2} \frac{\partial^2}{\partial t^2} p(x, t) \leftrightarrow \frac{s^2}{c_o^2} P(x, s) \quad (5.2.1.7)\]
with \( c_o = \sqrt{\eta_o P_o/\rho_o} \) being the sound velocity. Because the merged equations describe the pressure, which is a scalar field, this is an example of the scalar wave equation.

**Exercise #22**

Show that Eqs. 5.2.1.5 and 5.2.1.6 can be reduced to Eq. 5.2.1.7.

**Solution:** Taking the divergence of Eq. 5.2.1.5 gives

\[
-\nabla \cdot \nabla p(x, t) = \rho_o \frac{\partial}{\partial t} \nabla \cdot u(x, t).
\]

(Note that \( \nabla \cdot \nabla = \nabla^2 \) (Table 5.1). Next, substituting Eq. 5.2.1.6 into the above equation results in the scalar wave equation, Eq. 5.2.1.7, since \( c_o = \sqrt{\eta_o P_o/\rho_o} \).

**5.2.2 The Webster horn equation (WHEN)**

An important generalization of the problem of lossless plane-wave propagation in 1D uniform tubes is known as transmission line theory. As depicted in Fig. 5.2, by allowing the area \( A(r) \) [e.g., for the conical horn \( A(r) = A_o (r/L)^2 \)] with \( L = 1 \) m and \( A_o \leq 4\pi \) of an acoustical waveguide (horn) to vary along the range axis \( r \) (the direction of wave propagation), we can explore general solutions to the wave equation. Classic applications of horns include vocal tract acoustics, loudspeaker design, cochlear mechanics, quantum mechanics (e.g., the hydrogen atom), and wave propagation in periodic media (Brillouin, 1953).

We must be precise when defining the area \( A(x) \): The area is not the cross-sectional area of the horn; rather it is the wavefront (isopressure) area which is related to Gauss’ law, since the gradient of the pressure defines the force that drives the mass flow (also called volume velocity).

For the scalar wave equation (Eq. 5.1.3.10, p. 174), the Webster Laplacian is

\[
\nabla_r^2 \Phi(r, t) = \frac{1}{A(r)} \frac{\partial}{\partial r} \left[ A(r) \frac{\partial}{\partial r} \right] \Phi(r, t).
\]

The Webster Laplacian is based on the quasistatic approximation (Postulate P10, p. 122), which requires that the frequency lie below the critical value \( f_c = c_o/2d \) — namely, that a half wavelength be greater than the horn diameter \( d \) (i.e., \( d < \lambda/2 \)).\(^7\) For the adult human ear canal, \( d = 7.5 \) mm and \( f_c = (343/2 \cdot 7.5) \times 10^{-3} \approx 22.87 \) kHz.

The term on the right of Eq. 5.2.2.9, which is identical to Eq. 5.1.3.10 (p. 174), is also the Laplacian for thin tubes (e.g., rectangular, spherical, and cylindrical coordinates). Thus the Webster horn “wave” equation is

\[
\frac{1}{A(r)} \frac{\partial}{\partial r} \left[ A(r) \frac{\partial}{\partial r} \right] \Phi(r, t) = \frac{1}{c_o^2} \frac{\partial^2}{\partial t^2} \Phi(r, t) \Leftrightarrow \frac{s^2}{c_o^2} \Phi(r, s),
\]

where \( \Phi(r, t) \leftrightarrow \Phi(r, s) \) is the acoustic pressure in Pascals [Pa] (Hanna and Slepian, 1924; Mawardi, 1949; Eisner, 1967; Morse, 1948); Olson (1947, p. 101); Pierce (1981, p. 360). Extensive experimental analyses for various types of horns (conical, exponential, parabolic) along with a review of horn theory may be found in Goldsmith and Minton (1924). Of special interest is Eisner (1967) due to his history section and long list of relevant articles.

\(^7\)This condition may be written in several ways, the most common being \( k d < 1 \), where \( k = 2\pi/\lambda \) and \( d \) is the horn radius. This may be expressed in terms of the diameter as \( \frac{2\pi}{\lambda} d < 1 \), or \( d < \lambda/2r < \lambda/2 \). Thus \( d < \lambda/2 \) may be a more precise metric by the factor \( \pi/2 \approx 1.6 \). This is called the half-wavelength assumption, a synonym for the quasistatic approximation.
CHAPTER 5. STREAM 3B: VECTOR CALCULUS

Figure 5.3: Throat acoustical resistance \( r_A \) and acoustical reactance \( x_A \), frequency characteristics of infinite eigenfunctions of the parabolic, conical, exponential, hyperbolic, and cylindrical horns, having a throat area of \( 1 \, [\text{cm}^2] \). Note how the “critical” frequency (defined here as the frequency where the reactive and real parts of the radiation impedance are equal) of the horn reduces dramatically with the type of horn. For the uniform horn, the reactive component is zero, so there is no cutoff frequency. For the parabolic horn (1), the cutoff is around 3 kHz. For the conical horn (2), the cutoff is at 0.6 [kHz]. For the exponential horn (3), the critical frequency is around 0.18 [kHz], which is one-16th that of the parabolic horn. For each horn the cross-sectional area is defined as 100 [cm²] at a distance of \( L = 1 \, [\text{m}] \) from the throat (Olson, 1947, p. 101); (Morse, 1948, p. 283).

The limits of the Webster horn equation: It is commonly stated that the Webster horn equation (WHEN) is fundamentally limited and thus is an approximation that applies only to frequencies much lower than \( f_c \) (Morse, 1948; Shaw, 1970; Pierce, 1981). However, in all these discussions it is assumed that the area function \( A(r) \) is the horn’s cross-sectional area, not the area of the isopressure wavefront.

In the next section we show that this “limitation” may be avoided (subject to the \( f < f_c \) quasi-static limit, Postulate P10, p. 122), making the Webster horn theory an “exact” solution for the lowest-order “plane-wave” eigenfunctions of Eq. 5.2.2.10. The limitation of the quasistatic approximation is that it “ignores” higher-order evanescent modes, which are naturally small since, being evanescent modes below their cutoff frequency, the wave number is real and thus they do not propagate (Hahn, 1941; Karal, 1953). This is the same approximation that is required to define an impedance, since every eigenmode has an impedance (Miles, 1948). This method is frequently called a modal analysis or eigenanalysis. These modes define a Hilbert “vector” space (also called an eigenspace).

As derived in Appendix G (p. 263), the acoustic variables (eigenfunctions) are redefined on the isopressure wavefront boundary for the pressure and the corresponding volume velocity (Hanna and Slepian, 1924; Morse, 1948; Pierce, 1981). The resulting acoustic impedance is then the ratio of the pressure to the volume velocity. This approximation is valid up to the frequency where the first cross-mode begins to propagate \( (f > f_c) \), which may be estimated from the roots of the Bessel eigenfunctions (Morse, 1948). Perhaps it should be noted that these ideas, which come from acoustics, apply equally well to electromagnetics or any other wave phenomena described by eigenfunctions.

Visco-thermal losses: When losses are to be included, the wave number \( \kappa(s) = s/c_o \) must be replaced with Eq. D.1.0.1 (p. 249). This introduces dispersion in the wavefront due to the very small term \( \beta_0 \sqrt{s} \), which contains a branch cut. When calculating the losses, we must be careful that they are always on the correct sheet. In cases where precise estimates of the wave properties and input impedance are required, this term is critical.

The best known examples of wave propagation are electrical and acoustic transmission lines. Such systems are loosely referred to as the telegraph or telephone equations, harking back to the early days of their discovery (Heaviside, 1892; Campbell, 1903; Brillouin, 1953; Feynman, 1970a). In acoustics, waveguides are known as horns, such as the horn connected to the first phonographs from around the turn of the century (Webster, 1919). Thus the names reflect the historical development, back to a time when mathematics and its applications were running in close parallel.

5.2.3 Matrix formulation of the WHEN

Newton’s laws of conservation of momentum (Eq. 5.2.1.5) and mass (Eq. 5.2.1.6) are modern versions of Newton’s starting point for calculating the horn lowest-order plane-wave eigenmode wave speed.
The acoustic equations for the average pressure $P(r, \omega)$ and the volume velocity are derived in Appendix G, where the pressure and particle velocity equations (Eqs. G.1.1.4 and G.1.2.6) are transformed into a $2 \times 2$ matrix of acoustical variables, average pressure $P(r, \omega)$ and volume velocity $V(r, \omega)$:

$$
\frac{d}{dr} \begin{bmatrix} P(r, \omega) \\ V(r, \omega) \end{bmatrix} = \begin{bmatrix} 0 & \frac{s \rho_0}{A(r)} \\ \frac{s A(r)}{\eta_0 P_o} & 0 \end{bmatrix} \begin{bmatrix} P(r, \omega) \\ V(r, \omega) \end{bmatrix}.
$$

(5.2.3.11)

The equations

$$
M(r) = \rho_0 / A(r) \quad \text{and} \quad C(r) = A(r) / \eta_0 P_o
$$

(5.2.3.12)

define the per-unit-length mass and compliance of the horn (Ramo et al., 1965, p. 213). The product of $P(r, \omega)$ and $V(r, \omega)$ defines the acoustic power, while their ratio defines the horn's admittance $Y_{\pm}(r, s)$, looking in the two directions (Pierce, 1981, p. 37-41).

To obtain the Webster horn pressure equation Eq. 5.2.2.10 from Eq. 5.2.3.11, we take the partial derivative of the top equation

$$
-\frac{\partial^2 P}{\partial r^2} = s \frac{\partial M(r)}{\partial r} V' + s M(r) \frac{\partial V'}{\partial r}
$$

and then use the lower equation to remove $\frac{\partial V'}{\partial r}$,

$$
-\frac{\partial^2 P}{\partial r^2} = s \frac{\partial M(r)}{\partial r} V = s^2 M(r) C(r) P = \frac{s^2}{c_o^2} P.
$$

(5.2.3.13)

Note that $c_o^2 = MC = \left( \frac{\rho_0}{A(r)} \right) \cdot \left( \frac{\Delta r}{\eta_0 r_0} \right)$.

We use the upper equation a second time to remove $V'$:

$$
-\frac{\partial^2 P}{\partial r^2} + \frac{1}{A(r)} \frac{\partial A(r)}{\partial r} \frac{\partial P}{\partial r} = \frac{s^2}{c_o^2} P(r, s).
$$

(5.2.3.14)

By use of the chain rule, equations of this form may be directly integrated, since

$$
\nabla_r P = \frac{1}{A(r)} \frac{\partial}{\partial r} \left[ A(r) \frac{\partial}{\partial r} P(r, s) \right] = \frac{\partial^2 P}{\partial r^2} P(r, s) + \frac{1}{A(r)} \frac{\partial A(r)}{\partial r} P(r, s).
$$

(5.2.3.15)

This is equivalent to integration by parts, with integration factor $A(r)$. Next we set $\kappa(s) \equiv s / c_o$, which later will be generalized to include visco-thermal losses (Eq. D.1.0.1, p. 249).

Merging Eqs. 5.2.3.13 and 5.2.3.14 results in the Webster horn equation (WHEN) (Eq. 5.2.2.10, p. 183):

$$
\frac{1}{A(r)} \frac{\partial}{\partial r} A(r) \frac{\partial}{\partial r} P(r, s) = \kappa^2(s) P(r, s) \leftrightarrow \frac{1}{c_o^2} \frac{\partial^2}{\partial r^2} \delta(r, t).
$$

(5.2.3.15)

Equations that have this form are known as Sturm-Liouville equations.

This important class of ordinary differential equations follows from the use of separation of variables of the Laplacian in any (i.e., every) separable coordinate system (Morse and Feshbach, 1953, pp. 494-523). The frequency domain eigensolutions are denoted $P^{\pm}(r, s)$; they have corresponding volume velocities denoted $V^{\pm}(r, s)$.

We transform the 3D acoustic wave equation into acoustical variables (Eq. 5.2.1.7) in Appendix G by the application of Gauss's law, resulting in the 1D Webster horn equation (Eq. 5.2.2.10), which is a nonsingular Sturm-Liouville equation. Thus we demonstrate that Eqs. 5.2.1.7 and 5.2.3.11 reduce to to Eq. 5.2.3.15 in a horn.
5.3 Problems VC-1

Topics of this homework:

Vector algebra and fields in $\mathbb{R}^3$, gradient and scalar Laplacian operators, definitions of divergence and curl, Gauss’s (divergence) and Stokes’s (curl) laws, Schwarz inequality, quadratic forms, system postulates

Scalar fields and the $\nabla$ operator

Problem # 1: Let $T(x,y) = x^2 + y$ be an analytic scalar temperature field in two dimensions (single-valued $\in \mathbb{R}^2$).

- 1.1: Find the gradient of $T(x)$ and make a sketch of $T$ and the gradient.

- 1.2: Compute $\nabla^2 T(x)$ to determine whether $T(x)$ satisfies Laplace’s equation.

- 1.3: Sketch the iso-temperature contours at $T = -10, 0, 10$ degrees.

- 1.4: The heat flux is defined as $J(x,y) = -\kappa(x,y)\nabla T$, where $\kappa(x,y)$ is a constant that denotes thermal conductivity at the point $(x, y)$. Given that $\kappa = 1$ everywhere (the medium is homogeneous), plot the vector $J(x,y) = -\nabla T$ at $x = 2, y = 1$. Be clear about the origin, direction, and length of your result.

- 1.5: Find the vector $\perp$ to $\nabla T(x,y)$—that is, tangent to the iso-temperature contours. Hint: Sketch it for one $(x, y)$ point (e.g., 2, 1) and then generalize.

- 1.6: The thermal resistance $R_T$ is defined as the potential drop $\Delta T$ over the magnitude of the heat flux $|J|$. At a single point the thermal resistance is

$$R_T(x,y) = -\nabla T/|J|.$$  

How is $R_T(x,y)$ related to the thermal conductivity $\kappa(x,y)$?

Problem # 2: Acoustic wave equation

Note: In this problem, we will work in the frequency domain.

- 2.1: The basic equations of acoustics in one dimension are

$$-\frac{\partial}{\partial x} P = \rho_o s \hat{v} \quad \text{and} \quad -\frac{\partial}{\partial x} \hat{v} = \frac{s}{\eta_o} P.$$  

Here $P(x, \omega)$ is the pressure (in the frequency domain), $\hat{v}(x, \omega)$ is the volume velocity (the integral of the velocity over the wavefront with area $A$), $s = \sigma + \omega j$, $\rho_o = 1.2$ is the specific density of air, $\eta_o = 1.4$, and $P_o$ is the atmospheric pressure (i.e., $10^5$ Pa). Note that the pressure field $P$ is a scalar (pressure does not have direction), while the volume velocity field $\hat{v}$ is a vector (velocity has direction).

We can generalize these equations to three dimensions using the $\nabla$ operator

$$-\nabla P = \rho_o s \hat{v} \quad \text{and} \quad -\nabla \cdot \hat{v} = \frac{s}{\eta_o P_o} P.$$  

The heat flux is proportional to the change in temperature times the thermal conductivity $\kappa$ of the medium.
5.3. PROBLEMS VC-I

- 2.2: Starting from these two basic equations, derive the scalar wave equation in terms of the pressure \( P \),

\[
\nabla^2 P = \frac{s^2}{c_0^2} P,
\]

where \( c_0 \) is a constant representing the speed of sound.

- 2.3: What is \( c_0 \) in terms of \( \eta_0, \rho_0, \) and \( P_0 \)?

- 2.4: Rewrite the pressure wave equation in the time domain using the time derivative property of the Laplace transform [e.g., \( \frac{dx}{dt} \leftrightarrow sX(s) \)]. For your notation, define the time-domain signal using a lowercase letter, \( p(x, y, z, t) \leftrightarrow \mathcal{P} \).

Vector fields and the \( \nabla \) operator

Vector algebra

Problem # 3: Let \( \mathbf{R}(x, y, z) \equiv x(t)\hat{x} + y(t)\hat{y} + z(t)\hat{z} \).

- 3.1: If \( a, b, \) and \( c \) are constants, what is \( \mathbf{R}(x, y, z) \cdot \mathbf{R}(a, b, c) \)?

- 3.2: If \( a, b, \) and \( c \) are constants, what is \( \frac{d}{dt} \left( \mathbf{R}(x, y, z) \cdot \mathbf{R}(a, b, c) \right) \)?

Problem # 4: Find the divergence and curl of the following vector fields:

- 4.1: \( \mathbf{v} = \hat{x} + \hat{y} + 2\hat{z} \)

- 4.2: \( \mathbf{v}(x, y, z) = x\hat{x} + xy\hat{y} + z^2\hat{z} \)

- 4.3: \( \mathbf{v}(x, y, z) = x\hat{x} + xy\hat{y} + \log(z)\hat{z} \)

- 4.4: \( \mathbf{v}(x, y, z) = \nabla \left( \frac{1}{x} + \frac{1}{y} + \frac{1}{z} \right) \)

Vector and scalar field identities

Problem # 5: Find the divergence and curl of the following vector fields:

- 5.1: \( \mathbf{v} = \nabla \phi, \) where \( \phi(x, y) = xe^y \)

- 5.2: \( \mathbf{v} = \nabla \times \mathbf{A}, \) where \( \mathbf{A} = x\hat{x} + y\hat{y} + z\hat{z} \)

- 5.3: \( \mathbf{v} = \nabla \times \mathbf{A}, \) where \( \mathbf{A} = y\hat{x} + x^2\hat{y} + z\hat{z} \)

- 5.4: For any differentiable vector field \( \mathbf{V} \), write two vector calculus identities that are equal to zero.
5.5: What is the most general form a vector field may be expressed in, in terms of scalar \( \Phi \) and vector \( A \) potentials?

**Problem #6:** Perform the following calculations. If you can state the answer without doing the calculation, explain why.

- 6.1: Let \( \mathbf{v} = \sin(x) \hat{x} + y \hat{y} + z \hat{z} \). Find \( \nabla \cdot (\nabla \times \mathbf{v}) \).

- 6.2: Let \( \mathbf{v} = \sin(x) \hat{x} + y \hat{y} + z \hat{z} \). Find \( \nabla \times (\nabla \sqrt{\mathbf{v} \cdot \mathbf{v}}) \).

- 6.3: Let \( \mathbf{v}(x, y, z) = \nabla (x + y^2 + \sin(\log(z))) \). Find \( \nabla \times \mathbf{v}(x, y, z) \).

Integral theorems

**Problem #7:** For each of the following problems, in a few words, identify either Gauss’s or Stokes’s law, define what it means, and explain the formula that follows the question.

- 7.1: What is the name of this formula?
  \[
  \int_S \mathbf{n} \cdot \mathbf{v} \, dA = \int_V \nabla \cdot \mathbf{v} \, dV.
  \]

- 7.2: What is the name of this formula?
  \[
  \int_S (\nabla \times \mathbf{V}) \cdot d\mathbf{S} = \oint_C \mathbf{V} \cdot d\mathbf{R}
  \]
  Give one important application.

- 7.3: Describe a key application of the vector identity
  \[
  \nabla \times (\nabla \times \mathbf{V}) = \nabla (\nabla \cdot \mathbf{V}) - \nabla^2 \mathbf{V}.
  \]

System Classification

**Problem #8:** Complete this system classification problem about physical systems using the system postulates.

- 8.1: Provide a brief definition of these classifications
  
  \[
  \text{L/NL} : \text{linear (L)/nonlinear (NL)}
  \]
  
  \[
  \text{TI/TV} : \text{time-invariant (TI)/time-varying (TV)}
  \]
  
  \[
  \text{P/A} : \text{passive (P)/active (A)}
  \]
  
  \[
  \text{C/NC} : \text{causal (C)/noncausal (NC)}
  \]
  
  \[
  \text{Re/Clx} : \text{real (Re)/complex (Clx)}
  \]

- 8.2: Along the rows of the table, classify each system using the abbreviations L/NL, TI/TV, P/A, C/NC, and Re/Clx:
### 5.3. PROBLEMS VC-1

<table>
<thead>
<tr>
<th>#</th>
<th>Case</th>
<th>Definition</th>
<th>L/NL</th>
<th>TI/TV</th>
<th>P/A</th>
<th>C/NC</th>
<th>Re/Clx</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Resistor</td>
<td>( v(t) = r_0 i(t) )</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Inductor</td>
<td>( v(t) = L \frac{di}{dt} )</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Switch</td>
<td>( v(t) = \begin{cases} g &amp; t \leq 0; \ v_0 &amp; t &gt; 0. \end{cases} )</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Transistor</td>
<td>( I_{\text{out}} = g_m(V_{\text{in}}) )</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>Resistor</td>
<td>( v(t) = r_0 i(t + 3) )</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>Modulator</td>
<td>( f(t) = e^{i2\pi t} g(t) )</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### 8.3: Classify each equation:

<table>
<thead>
<tr>
<th>#</th>
<th>Case:</th>
<th>L/NL</th>
<th>TI/TV</th>
<th>P/A</th>
<th>C/NC</th>
<th>Re/Clx</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>( A(x) \frac{d^2y(t)}{dx^2} + D(t)y(x, t) = 0 )</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>( \frac{dy(t)}{dt} + \sqrt{t} y(t) = \sin(t) )</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>( y^2(t) + y(t) = \sin(t) )</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>( \frac{d^2y}{dt^2} + xy(t + 1) + x^2y = 0 )</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>( \frac{dy(t)}{dt} + (t - 1) y^2(t) = ie^t )</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 5.2: Horns and their properties for \( N = 1, 2, \) and 3 dimensions, along with the exponential horn (EXP). In this table the horn’s range variable is \( \xi \) [m], having area \( A(\xi) \) [m²] and diameter \( \xi_o = \sqrt{A(\xi_o)/\pi} \) [m], \( F(r) \) is the coefficient on \( \varphi_r \), \( \kappa(s) \equiv s/c_o \), where \( c_o \) is the speed of sound and \( s = \sigma + \omega \) is the Laplace frequency. The range variable \( \xi \) may be rendered dimensionless (see Fig. 5.3) if scaled by \( L \) (i.e., \( r \equiv L \xi \)), with \( \xi \) [m] the linear distance along the horn axis from \( r_o/L \leq \xi \leq 1 \), corresponding to \( r_o \leq r \leq L \), having area \( A_o(r_o/L)^2 \leq A_o \xi_o^2 \leq 4 \pi L^2 \). The horn’s eigenfunctions are \( \varphi^\pm (\xi, \omega) \equiv \varphi^\pm (\xi, t) \). When \( \pm \) is indicated, the outbound solution corresponds to the negative sign. Eigenfunctions \( H^\pm_2(\xi, s) \) are outbound and inbound Hankel functions. The rightmost column is the input radiation admittance normalized by the characteristic admittance \( Y_r^\pm(r) = A(r)/\rho_o c_o \).

<table>
<thead>
<tr>
<th>( N )</th>
<th>Name</th>
<th>radius</th>
<th>Area/( A_o )</th>
<th>( F(r) )</th>
<th>( \varphi^+ (r, s) )</th>
<th>( \varphi^0 (r_o, t) )</th>
<th>( Y_{rad}/\gamma_r )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1D</td>
<td>uniform</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>( e^{+\kappa(s)r} )</td>
<td>( \delta(t) )</td>
<td>1</td>
</tr>
<tr>
<td>2D</td>
<td>parabolic</td>
<td>( \sqrt{r/r_o} )</td>
<td>( r/r_o )</td>
<td>1/r</td>
<td>( H^+ (j \kappa(s)r) )</td>
<td>( -- )</td>
<td>( -\pi r_o H^+_r )</td>
</tr>
<tr>
<td>3D</td>
<td>conical</td>
<td>( r )</td>
<td>( r^2 )</td>
<td>2/r</td>
<td>( e^{+\kappa(s)r}/r )</td>
<td>( \delta(t) )</td>
<td>( \pi r_o )</td>
</tr>
<tr>
<td>EXP</td>
<td>exponential</td>
<td>( e^{mr} )</td>
<td>( e^{2mr} )</td>
<td>2m</td>
<td>( e^{-\kappa(s_1^2 + \kappa^2)r} )</td>
<td>( e^{-mr} E(t) )</td>
<td>Eq. 5.4.3.11</td>
</tr>
</tbody>
</table>

5.4 Three examples of finite-length horns

Figure 5.3 (p. 184) is taken from the classic book by Olson (1947, p. 101), showing the radiation impedance \( Z_{rad}(r, \omega) \) for five horns. Table 5.2 summarizes the properties of four of these horns: uniform (cylindrical) (\( A(r) = A_o \)), parabolic (\( A(r) = A_o r \)), conical (spherical) (\( A(r) = A_o r^2 \)), and exponential (\( A(r) = A_o e^{mr} \)), we discuss three of these next.

5.4.1 Uniform horn

The one-dimensional wave equation \([A(r) = A_o]\) is

\[
\frac{d^2}{dr^2} \varphi = \kappa^2(s) \varphi,
\]

where we set \( \kappa^2(s) \equiv s^2/c_o^2 \), which later will be generalized to include visco-thermal losses (see Eq. D.1.0.1, p. 249).

Solution: The two eigenfunctions of this equation are the two d’Alembert waves (Eq. 4.4.0.1, p. 142):

\[
\varphi(x, t) = \alpha g^+(t - x/c) + \beta g^-(t + (x - L)/c) \leftrightarrow \alpha e^{-\kappa(s)x} + \beta e^{\kappa(s)(x-L)},
\]

where \( \kappa(s) = s/c_o = \omega/c \) is called the propagation function (also known as the wave-evolution function, propagation constant, and wave number) and \( \alpha \) and \( \beta \) are the amplitudes of the two waves.

Note that for the uniform lossless horn, \( \omega/c_o = 2\pi/\lambda \). It is convenient to normalize \( \varphi^+_0 = 1 \) and \( \varphi^-_L = 1 \), as we did for the general case.

The characteristic admittance \( Y_r^+(r) \) (Table 5.2) is independent of direction. The signs must be physically chosen, with the velocity \( v^\pm \) into the port, to assure that \( Y_r^+ > 0 \).

Applying the boundary conditions: The general solution in terms of the eigenvector matrix, evaluated at \( x = L \), is

\[
\begin{bmatrix}
\varphi(x) \\
\varphi'(x)
\end{bmatrix}_L = \begin{bmatrix}
e^{-\kappa L} & e^{\kappa L-x-L} \\
eg \gamma_r e^{-\kappa L} & \gamma_r e^{\kappa L-x-L}
\end{bmatrix}_L \begin{bmatrix}
\alpha \\
\beta
\end{bmatrix}_L = \begin{bmatrix}
e^{-\kappa L} & 1 \\
-\gamma_r e^{-\kappa L} & -\gamma_r
\end{bmatrix}_L \begin{bmatrix}
\alpha \\
\beta
\end{bmatrix}_L,
\]

where \( \alpha \) and \( \beta \) are the relative weights on the two unknown eigenfunctions, to be determined by the boundary conditions at \( x = 0 \), \( L \), \( \kappa \equiv s/c \), and \( \gamma_r = 1/Z_r = A_o/\rho_o c \).

Solving Eq. 5.4.1.1 for \( \alpha \) and \( \beta \) with determinant \( \Delta = -2\gamma_r e^{-\kappa L} \), we get

\[
\begin{bmatrix}
\alpha \\
\beta
\end{bmatrix}_L = \frac{-1}{2\gamma_r e^{-\kappa L}} \begin{bmatrix}
-\gamma_r & -1 \\
\gamma_r e^{-\kappa L} & -\kappa L
\end{bmatrix}_L \begin{bmatrix}
\varphi(x) \\
\varphi'(x)
\end{bmatrix}_L = \frac{1}{2} \begin{bmatrix}
\kappa L & -Z_r e^{\kappa L} \\
1 & -\gamma_r
\end{bmatrix}_L \begin{bmatrix}
\varphi \\
\varphi'
\end{bmatrix}_L.
\]

In the final step we swapped all the signs, including those on \( \varphi' \), and moved \( Z_r = 1/\gamma_r \) inside the matrix.

We can uniquely determine these two weights given the pressure and velocity at the boundary \( x = L \), which is typically determined by the load impedance (\( \varphi_L/\varphi'_L \)).
The weights may now be substituted back into Eq. 5.4.1.1 to determine the pressure and velocity amplitudes at any point \(0 \leq x \leq L\):

\[
\begin{bmatrix}
P \\
V' \\
\end{bmatrix}_x = \frac{1}{2} \begin{bmatrix}
\epsilon^{-\kappa x} & \epsilon^{\kappa(x-L)} \\
\gamma_r e^{-\kappa x} & -\gamma_r e^{\kappa(x-L)} \\
\end{bmatrix}_x \begin{bmatrix}
\epsilon^{\kappa L} & -Z e^{\kappa L} \\
1 & Z \\
\end{bmatrix}_x \begin{bmatrix}
P \\
V' \\
\end{bmatrix}_L.
\] (5.4.1.3)

Setting \(x = 0\) and multiplying these out gives the final transmission matrix:

\[
\begin{bmatrix}
P \\
V' \\
\end{bmatrix}_0 = \frac{1}{2} \begin{bmatrix}
\epsilon^{\kappa L} + \epsilon^{-\kappa L} & \gamma_r (\epsilon^{\kappa L} - \epsilon^{-\kappa L}) \\
\gamma_r (\epsilon^{\kappa L} - \epsilon^{-\kappa L}) & \epsilon^{\kappa L} + \epsilon^{-\kappa L} \\
\end{bmatrix}_x \begin{bmatrix}
P \\
V' \\
\end{bmatrix}_L.
\] (5.4.1.4)

Note that the diagonal terms are \(\cosh \kappa L\) and the off-diagonal terms are \(\sinh \kappa L\).

Applying the last boundary condition, we evaluate Eq. 5.4.1.2 to obtain the ABCD matrix at the input \((x = 0)\) (Pipes, 1958),

\[
\begin{bmatrix}
P \\
V' \\
\end{bmatrix}_0 = \begin{bmatrix}
\cosh \kappa L & Z_r \sinh \kappa L \\
\gamma_r \sinh \kappa L & \cosh \kappa L \\
\end{bmatrix} \begin{bmatrix}
P \\
V' \\
\end{bmatrix}_L.
\] (5.4.1.5)

Note that the determinant is 1 and thus the system is reciprocal.

**Exercise #23**

Evaluate the expression in terms of the load impedance.

**Solution:** Since \(Z_{load} = -\frac{P_L}{V_L}\), we have

\[
\left| \frac{P}{V} \right|_0 = \frac{Z_{load} \cosh \kappa L - Z_r \sinh \kappa L}{Z_{load} \gamma_r \sinh \kappa L - \cosh \kappa L}.
\] (5.4.1.6)

**Impedance matrix:** Expressing Eq. 5.4.1.5 as an impedance matrix gives (algebra required)

\[
\begin{bmatrix}
P_0 \\
P_L \\
\end{bmatrix} = \frac{Z_r}{\sinh \kappa L} \begin{bmatrix}
\cosh \kappa L & 1 \\
1 & \cosh \kappa L \\
\end{bmatrix} \begin{bmatrix}
P_0 \\
V' \\
\end{bmatrix}_L.
\]

**Exercise #24**

Write out the short-circuit \((V_L' = 0)\) input impedance \(Z_{in}(s)\) for the uniform horn.

**Solution:**

\[
Z_{in}(s) = \left. \frac{P}{V} \right| = Z_r \frac{\cosh \kappa L}{\sinh \kappa L} = Z_r \tanh \kappa L|_{V_L'=0}.
\]

**Input admittance** \(Y_{in}\): Given the input admittance of the horn, it is possible to determine whether it is uniform without further analysis. That is, if the horn is uniform and infinite in length, the input admittance at \(x = 0\) is

\[
Y_{in}(x = 0, s) = \frac{\mathcal{V}(0, \omega)}{\mathcal{P}(0, \omega)} = \gamma_r,
\]

since \(\alpha = 1\) and \(\beta = 0\). For an infinite uniform horn, there are no reflections.

When the uniform horn is terminated with a fixed impedance \(Z_r\) at \(x = L\), we can substitute pressure and velocity measurements into Eq. 5.4.1.2 to find \(\alpha\) and \(\beta\), and given these, we can calculate the pressure reflectance at \(x = L\) (Eq. 3.4.2.6, p. 92),

\[
\Gamma_L(s) = \frac{\beta}{\alpha} = \frac{\mathcal{P}(L, \omega) - Z_r \mathcal{V}(L, \omega)}{\mathcal{P}(L, \omega) + Z_r \mathcal{V}(L, \omega)} = \frac{Z_L - Z_r}{Z_L + Z_r},
\]

given sufficiently accurate measurements of the throat pressure \(\mathcal{P}(0, \omega)\), velocity \(\mathcal{V}(0, \omega)\), and the characteristic impedance of the input \(Z_r = \rho_o c/\alpha(0)\).
5.4.2 Conical horn

Using the conical horn area \( A(r) \propto r^2 \) in Eq. 5.2.2.10, on page 183 (or Eq. 5.2.3.11 on page 185) results in the spherical wave equation

\[
\mathcal{P}_{rr}(r, \omega) + \frac{2}{r} \mathcal{P}_r(r, \omega) = \kappa^2 \mathcal{P}(r, \omega),
\]

(5.4.2.7)

where \( \kappa^2(s) \equiv \frac{s^2}{c^2_o} \).

**Radiation admittance for the conical horn:** The conical horn’s acoustic input admittance \( Y_{in}(r, s) \) at any location \( r \) is found by dividing \( V(r, s) \) by \( P(r, s) \):

\[
Y_{in}^\pm(r, s) = \frac{V^\pm(r, s)}{P^\pm(r, s)} = \frac{-A(r)}{s \rho_o} \left( 1 \pm \frac{c_o}{s r} \right) \ln \frac{P^\pm(r, s)}{A(r)}.
\]

(5.4.2.8)

Note how the pressure pulse is delayed by \( r/c_o \) due to \( e^{-\kappa(s)r} \) as it travels down the horn. As the area of the horn increases, the pressure decreases as \( 1/r = 1/\sqrt{A(r)} \). This results in the uniform backflow \( c_o u(t/r) \) due to conservation of mass and the characteristic admittance \( Y_r(r) \) variation with \( r \).

5.4.3 Exponential horn

If we define the area as \( A(r) = e^{2mr} \), the eigenfunctions of the horn are

\[
P^{\pm}(r, \omega) = e^{-mr} e^{\mp j\sqrt{\omega^2-c^2_o} r/c_o},
\]

(5.4.3.10)

which may be shown by the substitution of \( P^{\pm}(r, \omega) \) into Eq. 5.2.2.10 (p. 183), with \( A(r) = e^{2mr} \).

This case is of special interest because the radiation impedance is purely reactive below the horn’s cutoff frequency \( \omega < \omega_c = mc_o \), as may be seen from curves 3 and 4 of Fig. 5.3 (p. 184). As a result, no energy can radiate from an open horn for \( \omega < \omega_c \) because

\[
\kappa(s) = -m \pm \frac{j}{c_o} \sqrt{\omega^2-c^2_o}
\]

is purely real (this is the case of nonpropagating evanescent waves).

If we use Eq. 4.4.1.7 (p. 143), the input admittance is

\[
Y_{in}^\pm(x, s) = -\frac{A(x)}{s \rho_o} \left( m \pm \sqrt{m^2 + \kappa^2} \right) x.
\]

(5.4.3.11)

Kleiner (2013) gives an equivalent expression for \( Y_{in}(x, \omega) \) that has area \( S(x) = e^{mx} \),

\[
Y_{in}(x, \omega) = \frac{S(x)}{\rho c \omega} \left[ \frac{m}{2} + j \frac{\sqrt{4 \omega^2 - (mc)^2}}{2c} \right],
\]

and impedance

\[
Z_{in}(r, \omega) = \frac{\rho c}{S_T} \left[ \frac{\omega_c}{\omega} + \sqrt{1 - \left( \frac{\omega_c}{\omega} \right)^2} \right],
\]

where \( \omega_c(r) \) is the cutoff frequency.

We can use expansions of \( A(r) \) in a Fourier-like exponential series along with superposition to find the general solution for an arbitrary analytic \( A(r) \).

5.5 Solution methods

Two distinct mathematical techniques are described to model the wave equation in physical systems: 1) partial differential equations (PDEs) and 2) lumped-parameter models (i.e., quasistatics). We shall describe both these methods for the case of the scalar wave equation. The first of these is called separation of variables. This method is limited to a small and restrictive number of separable coordinate systems (SCS). Once the SCS is chosen, the
5.5. **SOLUTION METHODS**

Eigenfunctions are known, for any SCS. The differential equations that result from the separation of variables are known as Sturm-Liouville equations, which are always scalar (ordinary) differential equations (ODEs).

A second method of solution is to limit an upper frequency limit, so that quasi-statics can be assumed (the wavelength must be larger than the size of the object being modeled). This method lends itself to the lumped-element transmission method.

1a. **Separable coordinate systems:** Classically PDEs are solved by separation of variables shows that this method is limited to a few coordinate systems, such as rectangular, cylindrical, and spherical coordinates (Morse, 1948, p. 296-7). Even a slight deviation from separable specific coordinate systems represents a major barrier toward further analysis and understanding, blocking insight into more general cases. Separable coordinate systems have a high degree of symmetry. Note that the solution of the wave equation is not tied to a specific coordinate system.

1b. **Sturm-Liouville methods and eigenvectors:** When the coordinate system is separable, the resulting PDEs are always reduced to Sturm-Liouville equations. The important class of Sturm-Liouville equations are solved since their eigenfunctions are tabulated.

Webster horn theory (Webster, 1919; Morse, 1948; Pierce, 1981) is a generalized Sturm-Liouville equation that adds physics in the form of the horn’s area function. The Webster equation sidesteps the seriously limiting problem of separation of variables by using the alternative quasistatic solution which ignores high-frequency higher-order evanescent modes. This is essentially a one-dimensional low-pass approximation to the wave equation.

Mathematics provides rigor. Although physics provides understanding. While both are important, it is the physical applications that make a theory useful.

2. **Lumped-element method:** As described (see page 110) a system may be represented in terms of lumped-elements, as either electrical inductors, capacitors, and resistors or their mechanical counterparts, masses, springs, and dashpots. Such systems are represented by $2 \times 2$ transmission matrices in the $s$ (i.e., Laplace) domain (Ramo et al., 1965, Appendix IV).

When the system of lumped-element networks contains only resistors and capacitors, or resistors and inductors, the system does not support waves and is related to the diffusion equation in its solution. Depending on the elements in the system of equations, there can be an overlap between a diffusion process and scalar waves, represented as transmission lines, both modeled as lumped-element networks of $2 \times 2$ matrices (Eq. 3.7.0.1, p. 110) (Campbell, 1922; Brillouin, 1953; Ramo et al., 1965).

**Nyquist sampling and quasistatics:** Quasistatic methods provide band-limited solutions below a critical frequency $f_c$, where a half wavelength approaches the element spacing $\Delta$, for a much wider class of geometries by avoiding higher-order, high-frequency cross-modes. The model of the train is depicted in Fig. 5.4. The train may be accurately modeled as a transmission line (TL), since the equivalent electrical circuit is a lumped model of a TL.

**Train-mission-line problem:** Below the mechanical mass-sprint system it is the electrical equivalent circuit. The mass is modeled as an inductor and the springs as capacitors to ground. The velocity is analogous to a current and the force $f_n(t)$ to the voltage $\phi_n(t)$. The length of each cell is $\Delta [m]$.

When the wavelength $\lambda = c_0/f_c$ is greater than twice the physical distance $\Delta$ between the elements

$$\lambda > \lambda_c = 2\Delta \ [m],$$

the approximation is mathematically equivalent to a transmission line. As is described in DE-3, problem #2, the velocity is $c_0 = 1/\sqrt{MC}$ [m/s]. As the frequency increases, the wavelength becomes shorter. When the frequency is equal to the critical frequency $f_c$, the critical wavelength $\lambda_c = c_0/f_c = 2\Delta$. Above the critical frequency the quasistatic (lumped-element) model breaks down and switches from a delay line to a lowpass filter, as discussed in DE-3, problem #2.

The frequency is under the control of the modeling process, as more elements may be added, to represent higher frequencies (shorter wavelengths). If the nature of the solution at high frequencies ($f > f_c$) is desired, we must add more sections, thereby increasing $f_c$. For many (perhaps most) problems, lumped elements are easy to use and accurate as long as we don’t violate the upper-frequency limit (Brillouin, 1953; Ramo et al., 1965).
The wave equation (Eq. 5.2.1.7) is second-order in time, there are two causal independent eigenfunction solutions of the homogeneous (i.e., undriven) Webster horn equation: an outbound (right traveling) and an inbound (left-traveling) wave. These causal eigensolutions may be Laplace-transformed into the frequency domain:

\[ \mathcal{P}^\pm(r, s) = \int_0^\infty \varrho^\pm(r, t) e^{-st} \, dt. \]

They may be normalized so that \( \mathcal{P}^\pm(r_o, s) = 1 \), where \( r_o \) is the source excitation reference point.

Every eigenfunction depends on an area function \( A(r) \) (Eq. 5.2.2.10, p. 183). In theory then, given one, it should be possible to find the other. This is known as the inverse problem, which is generally believed to be an unsolved problem. For example, given the eigenvalues \( \lambda_k \), how does one determine the corresponding area function \( A(r) \)?

Because the characteristic impedance \( Z_c(r) \) of the wave in the horn changes with location, there must be local reflections due to these area variations. Thus there are fundamental relationships among the area change \( dA(r)/dr \), the horn’s eigenfunctions \( \mathcal{P}^\pm(r, s) \), the eigenmodes, and the input impedance.

Complex vs. real frequency: We shall continue to maintain the distinction that functions of \( \omega \) are Fourier transforms and causal functions of Laplace frequency \( s \) correspond to Laplace transforms, which are necessarily complex analytic in \( s \) in the right half-plane (RHP) region of convergence (RoC). This distinction is critical, since we typically describe impedance \( Z(s) \) and admittance \( Y(s) \) as complex analytic functions in \( s \) in terms of their poles and zeros. The eigenfunctions \( \mathcal{P}^\pm(r, s) \) of Eq. 5.2.2.10 are also causal complex analytic functions of \( s \).

Plane-wave eigenfunction solutions: In 1690, three years after Newton’s publication of *Principia*, Christiaan Huygens was the first to gain insight into wave propagation, today known as Huygens’s principle. While his concept showed a deep insight, we now know it was seriously flawed, as it ignored the backward-traveling wave (Miller, 1991). In 1747 d’Alembert published the first correct solution for the plane-wave scalar wave equation,

\[ \varrho(x, t) = f(t - x/c_o) + g(t + x/c_o), \quad (5.5.1.1) \]

where \( f(\cdot) \) and \( g(\cdot) \) are general functions of their argument. Why this is the solution may be shown by use of the chain rule, by taking partials with respect to \( x \) and \( t \).

In terms of physics, d’Alembert’s general solution describes two arbitrary waveforms \( f(\cdot) \) and \( g(\cdot) \) traveling at a speed \( c_o \), one forward and one reversed. Thus his solution is quite easily visualized.

**Exercise #25**

By the use of the chain rule, prove that d’Alembert’s formula satisfies the 1D wave equation.

**Solution:** Taking a derivative with respect to \( t \) and \( r \) gives

\[ \partial_t \varrho(r, t) = -c_o f'(r - c_o t) + c_o g'(r + c_o t) \]
5.6. INTEGRAL DEFINITIONS OF $\nabla()$, $\nabla \cdot ()$, $\nabla \times ()$, AND $\nabla \wedge ()$

\[
\partial_r \varrho(r,t) = f'(r-c_0t) + g'(r+c_0t) \]

and a second derivative gives

\[
\partial_{tt} \varrho(r,t) = c_o^2 f''(r-c_0t) + c_o^2 g''(r+c_0t) \]

\[
\partial_{rr} \varrho(r,t) = f''(r-c_0t) + g''(r+c_0t) \]

From these last two equations we have the 1D wave equation

\[
\partial_{rr} \varrho(r,t) = \frac{1}{c_o^2} \partial_{tt} \varrho(r,t),
\]

which has solutions in Eq. 5.5.1.1. ■

Exercise #26
Assuming $f(\cdot)$ and $g(\cdot)$ are $\delta(\cdot)$, find the Laplace transform of the solution corresponding to the uniform horn $A(x) = 1$.

**Solution:** Using Table C.3 (p. 242) of Laplace transforms on Eq. 5.5.1.1 gives

\[
\varrho(x,t) = \delta(t-x/c_o) + \delta(t+x/c_o) \leftrightarrow e^{-sx/c_o} + e^{sx/c_o}.
\] (5.5.1.2)

Note that the delay $T_o = \pm x/c_o$ depends on the range $x$. ■

Three-dimensional d’Alembert spherical eigenfunctions: We can generalize the d’Alembert solution to spherical waves by changing the area function of Eq. 5.2.2.10 to $A(r) = A_o r^2$ (see Eq. 5.1.3.10, p. 174 and Table 5.2, p. 190). The wave equation then becomes

\[
\nabla^2 \varrho(r,t) = \frac{1}{r} \frac{\partial^2}{\partial r^2} r \varrho(r,t) = \frac{1}{c_o^2} \frac{\partial^2}{\partial t^2} \varrho(r,t).
\]

Multiplying by $r$ results in the general spherical (3D) d’Alembert wave equation solution

\[
\varrho(r,t) = \frac{f(t-r/c_o)}{r} + \frac{g(t+r/c_o)}{r}
\]

for arbitrary waveforms $f(\cdot)$ and $g(\cdot)$. These are the eigenfunctions for the spherical scalar wave equation.

5.6 Integral forms of $\nabla()$, $\nabla \cdot ()$ and $\nabla \times ()$

The vector wave equation describes the evolution of a vector field, such as Maxwell’s electric field vector $E(x,t)$. When these fields are restricted to a one-dimensional domain, they are known as guided waves constrained by wave guides.

These equations use three differential vector operators: the gradient, divergence, and curl. There are two forms of definitions for each of these three operators: differential and integral. The integral form provides a more intuitive view of the operator, which in the limit converges to the differential form. Following a discussion of the gradient, divergence, and curl integral operators, we discuss these two forms.

In addition there are three fundamental vector theorems: Gauss’s law (divergence theorem), Stokes’s law (curl theorem), and Helmholtz’s decomposition theorem. Without the use of these fundamental vector calculus theorems, we could not understand Maxwell’s equations.
### 5.6.1 Gradient: \( \mathbf{E} = -\nabla \phi(\mathbf{x}) \)

As shown in Fig. 5.1 (p. 170), the gradient maps \( \mathbb{R}^3 \to \mathbb{R}^3 \). The gradient is defined as the unit-normal \( \hat{n} \) weighted by the potential \( \phi(\mathbf{x}) \) averaged over a closed surface \( S \):

\[
\nabla \phi(\mathbf{x}) \equiv \lim_{S,V' \to 0} \left\{ \int_{S'} \frac{\phi(\mathbf{x}) \hat{n} \, dS}{V'} \right\} \quad [\text{V/m}],
\]

(5.6.1.1)

having area \( S \) and volume \( V' \) and centered at \( \mathbf{x} \) (Greenberg, 1988, p. 773).

Here \( \hat{n} \) is a dimensionless unit vector perpendicular to the surface \( S \):

\[
\hat{n} = \frac{\nabla \phi}{||\nabla \phi||}.
\]

(5.6.1.2)

The dimensions of Eq. 5.6.1.1 are in the units of the potential times the area, divided by the volume, as needed for a gradient (e.g., [V/m]). The units depend on the potential. If \( \phi \) were temperature, the units would be [deg/m].

**Exercise #27**

Justify the units of Eq. 5.6.1.1.

**Solution:**

The units depend on \( \phi \) per unit length. If \( \phi \) is voltage, then the gradient has units of [V/m]. Under the limit, \( d|S|/||S|| \) must have units of \( m^{-1} \cdot m \).

The natural way to define the surface and volume is to place the surface on the isopotential surfaces, forming either a cube or a pill-box-shaped volume. As the volume \( ||S|| \) goes to zero, so must the area \( |S| \). One must avoid irregular volumes where the area is finite as the volume goes to zero (Greenberg, 1988, footnote p. 762).

A well-known example is the potential around a point charge \( Q \) [SI Units of Coulombs]. The constant \( \epsilon_o \) is the permittivity [F/m²]. A second well-known example is the acoustic pressure potential around an oscillating sphere, which has the same form (see Table 5.2, p. 190).

How does this work? To better understand what Eq. 5.6.1.1 means, consider a three-dimensional Taylor series expansion of the potential in \( \mathbf{x} \) about the limit point \( \mathbf{x}_0 \):

\[
\phi(\mathbf{x}) \approx \phi(\mathbf{x}_0) + \nabla \phi(\mathbf{x}) \cdot (\mathbf{x} - \mathbf{x}_0) + \text{HOT}.
\]

We could define the gradient using this relationship as

\[
\nabla \phi(\mathbf{x}_0) = \lim_{\mathbf{x} \to \mathbf{x}_0} \frac{\phi(\mathbf{x}) - \phi(\mathbf{x}_0)}{\mathbf{x} - \mathbf{x}_0}.
\]

For this definition to apply, \( \mathbf{x} \) must approach \( \mathbf{x}_0 \) along \( \hat{n} \). To compute the higher-order terms (HOT), we need the Hessian matrix.\(^{11}\)

The natural way to define a surface \( |S| \) is to find the isopotential contours. The gradient is in the direction of maximum change in the potential, thus perpendicular to the isopotential contours. The secret to the integral definition is taking the limit. As the volume \( ||S|| \) shrinks to zero, the HOT terms are small and the integral reduces to the first-order term in the Taylor expansion, since the constant term integrates to zero. Such a construction was used in the proof of the Webster horn equation (Appendix G, p. 263; Fig. G.1, p. 264).

The problem with Eq. 5.6.1.1 is that it is recursive, since \( \hat{n} \) is based on the gradient and is the kernel of the integral. Thus the integral definition of the gradient is based on the gradient itself. Equation 5.6.1.1 is actually a statement of the mean value theorem for the gradient.

### 5.6.2 Divergence: \( \nabla \cdot \mathbf{D} = \rho \ [\text{C/m}^3] \)

As briefly summarized by Eq. 5.1.2.4 on page. 172, the definition of the divergence at \( \mathbf{x} = [x, y, z]^T \) is

\[
\nabla \cdot \mathbf{D}(\mathbf{x}, t) \equiv [\partial_x, \partial_y, \partial_z] \cdot \mathbf{D}(\mathbf{x}, t) = \left[ \frac{\partial D_x}{\partial x} + \frac{\partial D_y}{\partial y} + \frac{\partial D_z}{\partial z} \right] (\mathbf{x}, t) = \rho(\mathbf{x}, t),
\]

which maps \( \mathbb{R}^3 \to \mathbb{R}^1 \).

\(^{10}\)For further discussions, see Greenberg (1988, pp. 778, 791, 809).

\(^{11}\) \( H_{ij} = \partial^2(\phi)/\partial x_i \partial x_j \), which exists if the potential is analytic in \( \mathbf{x} \) at \( \mathbf{x}_o \).
5.6. INTEGRAL DEFINITIONS OF $\nabla()$, $\nabla \cdot ()$, $\nabla \times ()$, AND $\nabla \wedge ()$

\[ \nabla \cdot \mathbf{D} \equiv \lim_{V',S \to 0} \left\{ \frac{\iint_{S} \hat{n} \cdot \mathbf{D} \, dS}{V'} \right\} \quad [\text{C/m}^3] \]

\[ Q_{\text{enc}} = \iint_{S} \hat{n} \cdot \mathbf{D} \, dS = \iiint_{V'} \rho_{\text{enc}} \, dV' \quad [\text{C}] \]

---

5.6.3 Divergence and Gauss’s law

Like the gradient, the divergence of a vector field may be defined as the surface integral of a compressible vector field as a limit as the volume enclosed by the surface goes to zero. As for the gradient, for this definition to make sense, the surface $S$ must be closed, defining the volume $V'$. The difference is that the surface integral is over the normal component of the vector field being operated on (Greenberg, 1988, p. 762-763),

\[ \nabla \cdot \mathbf{D} = \lim_{V',S \to 0} \iint_{S} \mathbf{D} \cdot \hat{n} \, dS = \rho_{\text{enc}}(x,y,z) \quad [\text{C/m}^3]. \quad (5.6.3.3) \]

As with the gradient, we have defined the surface as $S$, its area as $S$, and the volume within as $V'$. It is a necessary condition that as the area $S$ goes to zero, so does the volume $V'$.

As defined previously (Eq. 5.6.1.2) and shown here in Fig. 5.5, $\hat{n}$ is a unit vector normal to a closed iso-potential surface $S$. The limit, as the volume and surface simultaneously go to zero, defines the total flux across the surface. Thus the surface integral is a measure of the total flux perpendicular to the surface. It is helpful to compare this formula with that for the gradient, Eq. 5.6.1.1.

**Gauss’s law:** The definitions in Fig. 5.5 resulted in Gauss’s law, a major breakthrough in vector calculus. As summarized by Feynman (1970b, p. 13-2):

The current leaving the closed surface $S$ equals the rate of the charge leaving that volume $V'$, defined by that surface.

For the electrical case, this is equivalent to the observation that the total flux across the surface is equal to the net charge enclosed by the surface. Since the volume integral over the charge density $\rho(x,y,z)$ is the total charge enclosed $Q_{\text{enc}}$,

\[ Q_{\text{enc}} = \iiint_{V'} \nabla \cdot \mathbf{D} \, dV' = \iint_{S} \mathbf{D} \cdot \hat{n} \, dS \quad [\text{C}]. \quad (5.6.3.4) \]

When the surface integral over the normal component of $\mathbf{D}(x)$ is zero, the total charge is zero. If there is only positive (or negative) charge inside the surface, $\nabla \cdot \mathbf{D} = \rho(x) = 0$ charge density must also be zero. It is clear that this result only holds in the quasi-static limit, which is always satisfied because $S \to 0$.

Taking the derivative with respect to time gives the total current normal to the surface:

\[ I_{\text{enc}} = \iint_{S} \mathbf{D} \cdot \hat{n} \, dS = \dot{Q}_{\text{enc}} = \iiint_{V'} \dot{\rho}_{\text{enc}} \, dV' \quad [\text{A}]. \quad (5.6.3.5) \]

Of course, to define a volume, the surface must be closed, a necessary condition for Gauss’s law. This reduces to a common-sense summary that can be grasped intuitively.

5.6.4 Integral definition of the curl: $\nabla \times \mathbf{H} = \mathbf{C}$

As briefly summarized on page 173, the differential definition of the curl maps $\mathbb{R}^3 \nabla \times \mathbb{R}^3$. The curl of the magnetic field strength $\mathbf{H}(x)$ is the current density $\mathbf{C} = \sigma \mathbf{E} + \mathbf{D}$:

\[ \nabla \times \mathbf{H} \equiv \begin{vmatrix} \hat{x} & \hat{y} & \hat{z} \\ \frac{\partial}{\partial x} & \frac{\partial}{\partial y} & \frac{\partial}{\partial z} \\ H_x & H_y & H_z \end{vmatrix} = \mathbf{C} \quad [\text{A/m}^2]. \]
**Curl and Stokes’s law:** Like the gradient and divergence, the curl may be written in integral form, allowing for the physical interpretation of its meaning:

The surface integral definition of $\nabla \times \mathbf{H} = C \ [A/m^2]$, where the current density $C$ is perpendicular to the rotation plane of $\mathbf{H}$.

Stokes’s law states that the open surface integral over the normal component of the curl of the magnetic field strength ($\mathbf{\hat{n}} \cdot \nabla \times \mathbf{H} \ [A/m^2]$) is equal to the line integral $\oint_B \mathbf{H} \cdot d\mathbf{l}$ along the boundary $B$. As summarized in Fig. 5.6, Stokes’s law is

$$I_{\text{enc}} = \oint_B (\nabla \times \mathbf{H}) \cdot \mathbf{\hat{n}} \, dS = \oint_B \mathbf{H} \cdot d\mathbf{l} \quad [A].$$

That is,

The line integral of $\mathbf{H}$ along the open surface’s boundary $B$ is equal to the total current enclosed $I_{\text{enc}}$.

In many texts the normalization (denominator under the integral) is a volume $V$ (Greenberg, 1988, p. 778,823-4). However, because the surface is open, this volume does not exist (when we define a volume, the surface must be closed). The definition must hold even in the limit when the curved surface $S$ degenerates to a plane, with the boundary $B$ enclosing $S$. In this limit there is no volume.

To resolve this problem, we take the normalization to be the surface $S$. Note that in the limit $B \rightarrow 0$, the limiting definition is independent of any curvature, since the integral is over the normal component of $\mathbf{H}$ (i.e., $\mathbf{\hat{n}} \perp \mathbf{H}(x, t)$). The net flux is independent of the curvature of $S$ as $B \rightarrow 0$.

**Summary** Since integration is a linear process (sums of smaller elements), we can tile (tessellate) the surface, breaking it up into smaller surfaces and their boundaries, the sum of which is equal to the integral over the original boundary. This is an important concept that leads to the proof of Stokes’s law.

Table 5.1 (p. 172) provides a description of the three basic integration theorems along with their mapping domains. The integral formulations of Gauss’s and Stokes’s laws use $\mathbf{\hat{n}} \cdot \mathbf{D}$ and $\mathbf{H} \times \mathbf{\hat{n}}$ in the integrands. The key distinction between the two laws naturally follows from the properties of the scalar ($\mathbf{A} \cdot \mathbf{B}$) and vector ($\mathbf{A} \times \mathbf{B}$) products, as discussed in Fig. 3.4 (p. 94). To fully appreciate the differences between Gauss’s and Stokes’s laws, we must master these two types of vector products.

Paraphrasing Feynman (1970b, 3-12), we have

1. $\Phi_2 = \Phi_1 + \int_1^2 \nabla \Phi \cdot dS$
2. $\oint \mathbf{D} \cdot \mathbf{\hat{n}} \, dS = \oint \nabla \cdot \mathbf{D} \, dt'$
3. $\oint \mathbf{E} \cdot d\mathbf{l} = \oint (\nabla \times \mathbf{E}) \cdot \mathbf{\hat{n}} \, dS$
5.6. INTEGRAL DEFINITIONS OF $\nabla()$, $\nabla \cdot ()$, $\nabla \times ()$, AND $\nabla \wedge ()$

5.6.5 Helmholtz’s decomposition theorem

We must now rethink everything we have defined in terms of the two types of vector fields that decompose every analytic vector field (Table 5.3). The *irrotational field* is defined as one that is curl free. An *incompressible field* is one that is divergence free. According to Helmholtz’s decomposition theorem, every analytic vector field may be decomposed into independent rotational and compressible components (Helmholtz, 1978). Another name for Helmholtz’s decomposition theorem is the fundamental theorem of vector calculus (FTVC); Gauss’s and Stokes’s theorems along with Helmholtz’s decomposition theorem are the three fundamental theorems of vector calculus. Portraits of Helmholtz and Kirchhoff are provided in Fig. 5.8.

Table 5.3: The four possible classifications of scalar and vector potential fields: rotational/irrotational and compressible/incompressible. Rotational fields are generated by the vector potential (e.g., $A(x, t)$), while compressible fields are generated by the scalar potentials (e.g., voltage $\phi(x, t)$, velocity $\psi$, pressure $\rho(x, t)$, or temperature $T(x, t)$).

<table>
<thead>
<tr>
<th>Field:</th>
<th>Compressible</th>
<th>Incompressible</th>
</tr>
</thead>
<tbody>
<tr>
<td>$v(x, t)$</td>
<td>$\nabla \cdot v \neq 0$</td>
<td>$\nabla \cdot v = 0$</td>
</tr>
<tr>
<td>Rotational</td>
<td>$v = \nabla \phi + \nabla \times \omega$</td>
<td>$v = \nabla \times \omega$</td>
</tr>
<tr>
<td>$\nabla \times v \neq 0$</td>
<td>Vector wave Eq. $\nabla^2 v = \frac{1}{\epsilon} \rho$</td>
<td>Lubrication theory</td>
</tr>
<tr>
<td>Irrotational conservative</td>
<td>$\nabla \cdot v = 0$</td>
<td>$\nabla^2 \phi = 0$</td>
</tr>
<tr>
<td>$\nabla \times v = 0$</td>
<td>Acoustics</td>
<td>$\nabla^2 \rho(x, t) = \frac{1}{\epsilon} \rho(x, t)$</td>
</tr>
<tr>
<td>$\nabla \times = 0$</td>
<td>Statics</td>
<td>Laplace’s Eq. ($\epsilon \rightarrow \infty$)</td>
</tr>
</tbody>
</table>

A magnetic solenoidal field is a uniform-flux field $B_z(x)$ that is generated by a solenoidal coil and, to an excellent approximation, is uniform inside the coil, making it similar to that of a permanent magnet. As a result, the divergence of a solenoidal field is approximately zero, which makes it incompressible ($\nabla \cdot B = 0$) and rotational ($\nabla \times B \neq 0$).

You should know this term, since it is widely used, but the preferred terms are *incompressible* and *rotational*. Strictly speaking, the term *solenoidal field* applies to only a magnetic field produced by a solenoid and is specific to that case.

**Figure 5.7:** A solenoid is a uniform coil of wire. When a current is passed through the wire, a uniform magnetic field intensity $H$ is created. In its properties, this coil is indistinguishable from a permanent bar magnet that has north and south poles. Depending on the direction of the current, one end of a finite solenoidal coil is the north pole of the magnet, and the other end is the south pole. The uniform field inside the coil is called solenoidal, a confusing synonym for irrotational. (Figure from Wikipedia)

**Helmholtz’s decomposition of a differentiable vector field:** This theorem is easily stated (and proved) but less easily appreciated (Heras, 2016). A physical description is helpful: Every vector field may be split into two independent parts: dilation and rotation. We have seen this same idea in vector algebra, where the scalar and wedge products of two vectors are perpendicular (Fig. 3.4, p. 94).

For example, think of linear versus angular momentum, which are independent in that they represent different ways of delivering kinetic energy via different modalities (degrees of freedom, DOF). Linear and rotational motions are a common theme in physics, rooted in geometry. Thus it seems a natural extension to split a vector field into independent dilation and rotation parts. In the case of fluid mechanics, these modalities can couple through friction due to viscosity.

A fluid with mass and momentum can be moving along a path and independently be rotating. These independent modes of motion correspond to different types (modes) of kinetic energy, such as transnational, compressional, and rotational. Each eigenmode of vibration can be viewed as an independent degree of freedom (DoF).

Helmholtz’s decomposition theorem (FTVC) quantifies these degrees of freedom. Second-order vector identities $\nabla \cdot \nabla \times () = 0$ and $\nabla \times \nabla () = 0$ may be used to verify the FTVC. The role of the FTVC is especially powerful when applied to Maxwell’s equations.

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12These theorems are mathematical relationships that follow from physical principles.
The four categories of linear fluid flow: The following is a summary of the four cases for fluid flow, as shown in Table 5.3:

1.1 Compressible, rotational fluid (general case): $\nabla \psi \neq 0$, $\nabla \times \mathbf{w} \neq 0$. This is wave propagation in a medium where viscosity cannot be ignored, as in acoustics close to the boundaries, where viscosity contributes to losses (Batchelor, 1967).

1.2 Incompressible, rotational fluid (lubrication theory): $\mathbf{v} = \nabla \times \mathbf{w} \neq 0$, $\nabla \cdot \mathbf{v} = 0$, $\nabla^2 \psi = 0$. In this case, the flow is dominated by the walls, while the viscosity and heat transfer introduce shear. This is typical of lubrication theory and solenoidal fields.

2.1 Compressible, irrotational fluid (acoustics): $\mathbf{v} = \nabla \psi$, $\nabla \times \mathbf{w} = 0$. Here losses (viscosity and thermal diffusion) are small (assumed to be zero). We can define a velocity potential $\psi$, the gradient of which gives the air particle velocity, thus $\mathbf{v} = -\nabla \psi$. For an irrotational fluid, $\nabla \times \mathbf{v} = 0$ (Greenberg, 1988, p. 826). When a fluid may be treated as having no viscosity, it is typically assumed to be irrotational, since it is the viscosity that introduces the shear (Greenberg, 1988, p. 814). A fluid’s angular velocity is $\Omega = \frac{1}{2} \nabla \times \mathbf{v} = 0$, thus irrotational fluids have zero angular velocity ($\Omega = 0$).

2.2 Incompressible, irrotational fluid (statics): $\nabla \cdot \mathbf{v} = 0$ and $\nabla \times \mathbf{v} = 0$; thus $\mathbf{v} = \nabla \psi$ and $\nabla^2 \psi = 0$. An example is water in a small space at low frequencies, where the wavelength is long compared to the size of the container; the fluid may be treated as incompressible. When $\nabla \times \mathbf{v} = 0$, the effects of viscosity may be ignored, as it is the viscosity that creates the shear that leads to rotation. This is the case in modeling the cochlea, where losses are ignored and the quasistatic limit is justified.

In summary, each of these cases is an approximation that best applies in the low-frequency limit. This is why it is called quasistatic, meaning low but not zero frequency, where the wavelength is more than twice the diameter.

5.6.6 Second-order operators (JBA to review Sec. 5.5.6)

In addition to the first-order vector derivatives are second-order operators, the most important being the scalar Laplacian $\nabla^2() = \nabla \cdot \nabla ()$ and the vector Laplacian $\nabla^2() = \nabla \cdot \nabla ()$, which operates on vectors.\(^{13}\)

Terminology:

There are six second-order combinations of $\nabla$, requiring six mnemonics (Table 5.1, p. 172):

1. **DoG**: Divergence of the gradient (scalar Laplacian) operates on scalar potentials: (Greenberg, 1988, p. 779)

$$\nabla^2 \phi = (\nabla \cdot \nabla) \phi = \frac{\partial^2 \phi}{\partial x^2} + \frac{\partial^2 \phi}{\partial y^2} + \frac{\partial^2 \phi}{\partial z^2}$$


$$\nabla^2 \mathbf{A} = (\nabla \cdot \nabla) \mathbf{A} = \frac{\partial^2 \mathbf{A}}{\partial x^2} + \frac{\partial^2 \mathbf{A}}{\partial y^2} + \frac{\partial^2 \mathbf{A}}{\partial z^2} = BullDog \mathbf{A} - \nabla \times \nabla \times \mathbf{A}$$

3. **GoD**: (little-GoD) Gradient of the Divergence ($\nabla^2 \mathbf{A} = \nabla (\nabla \cdot \mathbf{A})$)

\(^{13}\)https://en.wikipedia.org/wiki/Del_in_cylindrical_and_spherical_coordinates#Non-trivial_calculation_rules
\[ \nabla^2 A = \nabla(\nabla \cdot A) = \nabla \left( \frac{\partial a_x}{\partial x} + \frac{\partial a_y}{\partial y} + \frac{\partial a_z}{\partial z} \right) = \left( \hat{x} \frac{\partial}{\partial x} + \hat{y} \frac{\partial}{\partial y} + \hat{z} \frac{\partial}{\partial z} \right) \left( \frac{\partial a_x}{\partial x} + \frac{\partial a_y}{\partial y} + \frac{\partial a_z}{\partial z} \right) = \hat{x} \frac{\partial}{\partial x} \nabla \cdot A + \hat{y} \frac{\partial}{\partial y} \nabla \cdot A + \hat{z} \frac{\partial}{\partial z} \nabla \cdot A \]

4. CoC: Curl of the curl (GoD\(A - \text{DoG} A\)) (Sommerfeld, 1952, page 33, Eq. 2b).

\[ \nabla \times \nabla \times A = \nabla(\nabla \cdot A) - (\nabla \cdot \nabla)A = \nabla^2 A - \nabla^2 A \]

5. DoC: Divergence of the curl (\(\nabla \cdot \nabla \phi = 0\))

6. CoG: Curl of the gradient (\(\nabla \times \nabla = 0\))

DoC(\()\) and CoG(\()\) are special because they are always zero

\[ \nabla \cdot \nabla \times A = 0, \quad \nabla \times \nabla \phi = 0, \quad (5.6.6.7) \]

making them useful in proving the fundamental theorem of vector calculus, aka, Helmholtz’ decomposition theorem (Eq. 5.8.0.5, p. 207). A third special vector identity CoC is

\[ \nabla \times \nabla \times A = \nabla(\nabla \cdot A) - \nabla^2 A, \quad (5.6.6.8) \]

which operates on vector fields and is useful for defining the vector Laplacian DoG as the difference between little GoD (gOd) and CoC (i.e., GoD = gOd - CoC)

\[ \nabla^2() = \nabla^2 () - \nabla \times \nabla \times () \]

The role of gOd (\(\nabla^2\)) is commonly ignored because it is zero for the magnetic wave equation, due to there being no magnetic charge (\(\nabla \cdot B(x, t) = 0 \text{ thus } \nabla^2 B(x, t) \equiv 0\)). However for the electric vector wave equation it plays a role

\[ \nabla^2 \phi(x, t) = -\nabla E(x, t) = -\frac{1}{\epsilon_0} \nabla^2 D(x, t) = -\frac{1}{\epsilon_0} \nabla \rho(x, t) \]

or since \(\nabla \cdot D = \rho\),

\[ \nabla^2 D(x, t) = \nabla \nabla \cdot D = -\nabla \rho(x, t) \]

When the charge density is inhomogeneous, such as the case of a plasma (e.g., the sun) this term will play an important role as a source term to the electric wave equation. This case needs to be further explored via some physical examples.

**Exercise #28**

Show that GoD and gOd differ.

**Solution:** Use CoC on \(A(x, t)\) to explore this relationship.

**Discussion:** It is helpful to split these six identities into two groups: the utility operators DoG, gOd, GoD, and the identity operators DoC and CoG (Eq. 5.6.6.7). It is helpful to view these two groups as playing fundamentally different roles.

When using second-order differential operators one must be careful with the order of operations, which can be subtle. Most of this is common sense. For example, don’t operate on a scalar field with \(\nabla \times\), and don’t operate on a vector field with \(\nabla\).\(^{14}\) GoD acts on each vector component \(\nabla^2 A = \nabla^2 A_x \hat{x} + \nabla^2 A_y \hat{y} + \nabla^2 A_z \hat{z}\), which is very different from the action of gOd.

\(^{14}\)This operation defines a dyadic tensor, a generalization of the vector.
5.7 The unification of electricity and magnetism

Once we have mastered the three basic vector operations –gradient, divergence, and curl – we are ready to appreciate Maxwell’s equations. Like the vector operations, these equations may be written in integral or differential form. An important difference is that with Maxwell’s equations, we are dealing with well-defined physical quantities. The scalar and vector fields take on meaning and units. Thus, to understand these important equations, we must master both the names and units of the four fields $E$, $H$, $B$, $D$, as described in Table 5.4.

Table 5.4: The variables of Maxwell’s equations have names (e.g., EF, MI) and units (in square brackets [SI Units]). The units are necessary to obtain a full understanding of each of the four variable and their corresponding equation. For example, Eq. EF has units [V/m]. By integrating $E$ from $x = a, b$, one obtains the voltage difference between the two points. The speed of light in-vacuo is $c = 3 \times 10^8 \text{m/s}$, and the characteristic resistance of light $r_o = 377 \text{[ohms]}$ (i.e., ohms).

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Name</th>
<th>Units</th>
<th>Maxwell’s Eq.</th>
</tr>
</thead>
<tbody>
<tr>
<td>$E$</td>
<td>EF: Electric field strength</td>
<td>[V/m]</td>
<td>$\nabla \times E = -\partial_t B$</td>
</tr>
<tr>
<td>$D = \epsilon_o E$</td>
<td>ED: Electric displacement (flux density)</td>
<td>[C/m$^2$]</td>
<td>$\nabla \cdot D = \rho$</td>
</tr>
<tr>
<td>$H$</td>
<td>MF: Magnetic field strength</td>
<td>[A/m]</td>
<td>$\nabla \times H = J_m + \partial_t D$</td>
</tr>
<tr>
<td>$B = \mu_o H$</td>
<td>MI: Magnetic induction (flux density)</td>
<td>[Wb/m$^2$]</td>
<td>$\nabla \cdot B = 0$</td>
</tr>
</tbody>
</table>

Field strength $E, H$: As summarized by Eqs. 5.7.1.1 there are two field strengths; the electric $E$ with units of [V/m] and the magnetic $H$ with units of [A/m]. The ratio $|E|/|H| = \sqrt{\mu_o/\epsilon_o} = 377$ [ohms] for in-vacuo plane-waves ($\mu_o, \epsilon_o$).

To understand the meaning of $E$, if two conducting plates are placed 1 [m] apart, with 1 [V] across them, the electric field is $E = 1$ [V/m]. If a charge (i.e., an electron) is placed in an electric field, it feels a force $f = qE$, where $q$ is the magnitude of the charge [C].

To help us understand the meaning of $H$, consider the solenoid made of wire, as shown in Fig. 5.7, that carries a current of 1 [A]. The magnetic field $H$ inside such a solenoid is uniform and is pointed along the long axis, with a direction that depends on the polarity of the applied voltage (i.e., the direction of the current in the wire).

Flux $D, B$: Flux is a flow, such as the mass flux of water flowing in a pipe [kg/s] driven by a force (pressure drop) across the ends of the pipe, or the heat flux in a thermal conductor, that has a temperature drop across it (i.e., a window or a wall). The flux is the same as the flow, be it charge, mass, or heat (Table 3.2, p. 114). In Maxwell’s equations there are also two fluxes: the electric flux $D$ and the magnetic flux $B$. The flux density units for $D$ are [A/m$^2$] (flux in [A]), and the magnetic flux $B$ is measured in Weber’s [Wb] [A/m$^2$]) or [Tesla] (henry-amps/area) [H-A/m$^2$].
5.7. **THE UNIFICATION OF ELECTRICITY AND MAGNETISM**

5.7.1 **Maxwell’s equations**

Maxwell’s equations (ME) consist of two curl equations (Eqs. 5.7.1.1) operating on the field strengths \( \mathbf{E} \) and \( \mathbf{H} \), and two divergence equations operating on the field fluxes \( \mathbf{D} \) and \( \mathbf{B} \). In matrix format, the ME are

\[
\nabla \times \begin{bmatrix} \mathbf{E}(x, t) \\ \mathbf{H}(x, t) \end{bmatrix} = \partial_t \begin{bmatrix} -\mathbf{B}(x, t) \\ \mathbf{D}(x, t) \end{bmatrix}
\]

\[
\begin{bmatrix} 0 & -\mu_o \\ \varepsilon_o & 0 \end{bmatrix} \partial_t \begin{bmatrix} \mathbf{E}(x, t) \\ \mathbf{H}(x, t) \end{bmatrix}
\]

\[
\leftrightarrow \begin{bmatrix} 0 & -s\mu_o \\ \sigma_o + s\varepsilon_o & 0 \end{bmatrix} \begin{bmatrix} \mathbf{E}(x, \omega) \\ \mathbf{H}(x, \omega) \end{bmatrix}.
\]

(5.7.1.1)

When the medium is conducting, \( \partial_t \mathbf{D} \) must be replaced by \( \mathbf{C} = \sigma_o \mathbf{E} + \partial_t \mathbf{D} \leftrightarrow (\sigma_o + s\varepsilon_o) \mathbf{E}(x, \omega) \) where \( \sigma_o + s\varepsilon_o \) is an admittance density \([\Omega/m^2]\).

There are also two auxiliary equations:

\[
\nabla \cdot \begin{bmatrix} \mathbf{D} \\ \mathbf{B} \end{bmatrix} = -\partial_t \begin{bmatrix} \rho(x) \\ 0 \end{bmatrix}.
\]

(5.7.1.2)

The top equation states conservation of charge, while the lower states that there is no magnetic charge. When expressed in integral format, Stokes’s law follows from the curl equations and Gauss’s law from the divergence equations.

**Exercise #29**

When a static current is flowing in a wire in the \( \hat{z} \) direction, the magnetic flux is determined by Stokes’s theorem (Fig. 5.6). Thus, just outside the wire we have

\[
I_{\text{enc}} = \oint \mathbf{H} \cdot \hat{n} d|S| = \int \mathbf{H} \cdot d\mathbf{l} [\text{A}].
\]

For this simple geometry, the current in a wire is related to \( \mathbf{H}(x, t) \) by

\[
I_{\text{enc}} = \int \mathbf{H} \cdot d\mathbf{l} = H_\phi 2\pi r.
\]

Here \( H_\phi \) is perpendicular to both the radius \( r \) and the direction of the current \( \hat{z} \). Thus

\[
H_\phi = \frac{I_{\text{enc}}}{2\pi r},
\]

where \( H_\phi \) is attenuated by \( 1/r \) (Ramo et al., 1965, Eq. 9, page 244).

**Exercise #30**

Explain how Stokes’s theorem may be applied to \( \nabla \times \mathbf{E} = -\dot{\mathbf{B}} \), and explain what it means. Hint: This is the same argument given above for the current in a wire, but for the electric case.

**Solution:** Integrating the left side of equation Eq. 5.7.1.3 over an open surface results in a voltage (emf) induced in the loop closing the boundary \( \partial S \) of the surface

\[
\phi_{\text{induced}} = \oint \mathbf{E} \cdot \hat{n} d|S| = \int \mathbf{E} \cdot d\mathbf{l} [\text{V}].
\]

The emf (electromagnetic force) is the same as the Thévenin source voltage induced by the rate of change of the flux. Integrating Eq. 5.7.1.3 over the same open surface \( S \) results in the source of the induced voltage \( \phi_{\text{induced}} \), which is proportional to the rate of change of the flux [Webers]:

\[
\phi_{\text{induced}} = -\frac{\partial}{\partial t} \int \mathbf{B} \cdot \hat{n} dA = L \dot{\psi} \quad [\text{Wb/s}] \text{ or } [\text{V}],
\]

where \( L \) is the inductance of the wire. The area integral on the left is in [Wb/m²], resulting in the total flux crossing normal to the surface \( \psi \) [Wb]. Thus the rate of change of the total flux [Wb/s] is a voltage [V].
If we apply Gauss’s theorem to the divergence equations, we find the total flux that crosses the closed surface.

**Exercise #31**

Apply Gauss’s theorem to equation ED and explain what it means in physical terms.

**Solution:** The area of the normal component of $\mathbf{D}$ is equal to the volume integral over the charge density.

Thus Gauss’s theorem says that the total charge within the volume $Q_{\text{enc}}$, found by integrating the charge density $\rho(x)$ over the volume $\mathcal{V}$, is equal to the normal component of the flux $\mathbf{D}$ that crosses the surface $\mathcal{S}$:

$$Q_{\text{enc}} = \iiint_{\mathcal{V}} \nabla \cdot \mathbf{D} \, dV = \iint_{\mathcal{S}} \mathbf{D} \cdot \hat{n} \, dA.$$

When equal amounts of positive and negative charge exist within the volume, the integral is zero. ■

**Summary:** Maxwell’s four equations relate the field strengths to the flux densities. There are two types of variables: field strengths ($\mathbf{E}, \mathbf{H}$) and flux densities ($\mathbf{D}, \mathbf{B}$). There are two classes: electric ($\mathbf{E}, \mathbf{D}$) and magnetic ($\mathbf{H}, \mathbf{B}$). This is a $2 \times 2$ matrix, with column being field strength and flux densities and rows being electric and magnetic variables.

<table>
<thead>
<tr>
<th>Electric</th>
<th>Flux density</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\mathbf{E}$ [V/m]</td>
<td>$\mathbf{D}$ [C/m$^2$]</td>
</tr>
<tr>
<td>$\mathbf{H}$ [A/m]</td>
<td>$\mathbf{B}$ [Wb/m$^2$]</td>
</tr>
</tbody>
</table>

Applying Stokes’s curl theorem to the forces induces a Thévenin voltage (emf) or Norton current source.

Applying Gauss’s divergence theorem to the flows gives the total charge enclosed. The magnetic charge is zero ($\nabla \cdot \mathbf{B} = 0$) because magnetic monopoles do not exist. However, magnetic dipoles do exist, as in the example of the electron that contains a magnetic dipole.

### 5.7.2 Derivation of the vector wave equation

Next we provide the derivation of the vector wave equation starting from Maxwell’s equations (Eq. 5.7.1.1), which is reminiscent of the derivation of the Webster horn equation (Eq. 5.2.1, p. 182). Working in the frequency domain and taking the curl of both sides give

$$\nabla \times \nabla \times \begin{bmatrix} \mathbf{E} \\ \mathbf{H} \end{bmatrix} = \begin{bmatrix} 0 & -s\mu_0 \\ s\epsilon_0 & 0 \end{bmatrix} \nabla \times \begin{bmatrix} \mathbf{E} \\ \mathbf{H} \end{bmatrix} = \begin{bmatrix} 0 & -s\mu_0 \\ s\epsilon_0 & 0 \end{bmatrix} \begin{bmatrix} \mathbf{E} \\ \mathbf{H} \end{bmatrix} = -\frac{s^2}{c_0^2} \begin{bmatrix} \mathbf{E} \\ \mathbf{H} \end{bmatrix}.$$

Using the CoC identity $\nabla \times \nabla \times () = \nabla^2 () - \nabla^2 ()$ (Eq. 5.6.6.8, p. 201) gives

$$\nabla^2 \begin{bmatrix} \mathbf{E} \\ \mathbf{H} \end{bmatrix} = \nabla^2 \begin{bmatrix} \mathbf{E} \\ \mathbf{H} \end{bmatrix} = \frac{s^2}{c_0^2} \begin{bmatrix} \mathbf{E} \\ \mathbf{H} \end{bmatrix}$$

or finally Maxwell’s vector wave equation

$$\nabla^2 \begin{bmatrix} \mathbf{E} \\ \mathbf{H} \end{bmatrix} = \frac{s^2}{c_0^2} \begin{bmatrix} \mathbf{E} \\ \mathbf{H} \end{bmatrix} = \nabla \left[ \frac{1}{\epsilon_0} \nabla \cdot \mathbf{D} \right] - \nabla \left[ \frac{1}{\mu_0} \nabla \cdot \mathbf{B} \right] + \nabla \rho(x,s)$$

(Eq. 5.7.2.4)

with the electric excitation term $\nabla \rho(x,s)$. Note that if $\mu$ and $\epsilon$ depended on $x$, the terms on the right would not be zero. In deep outer space with its black holes and plasma everywhere (e.g., inside the sun) this seems possible, even likely.

Recall the d’Alembert solutions of the scalar wave equation (Eq. 4.4.0.1, p. 142)

$$E(x,t) = f(x-ct) + g(x+ct),$$

where $f$ and $g$ are arbitrary vector fields. This result applies to the vector case, since it represents three identical, yet independent, scalar wave equations in the three dimensions.
Poynting vector: The EM power flux density $\mathcal{P}$ [W/m$^2$] is perpendicular to $E$ and $B$, denoted as

$$\mathcal{P} = \frac{1}{\mu_0} \mathbf{E} \times \mathbf{B} = \mathbf{E} \times \mathbf{H} \quad [\text{W/m}^2].$$

The corresponding EM momentum flux density $\mathcal{M}$ (hence ME are related to mass, thus gravity) is

$$\mathcal{M} = \epsilon_0 \mathbf{E} \times \mathbf{B} = \mathbf{D} \times \mathbf{B} \quad [\text{C/m}^2 \cdot \text{Wb/m}^2].$$

Since the speed of light is $c_0 = 1/\sqrt{\mu_0\epsilon_0}$, dividing by the momentum flux density gives

$$\mathcal{P} = c_0^2 \mathcal{M} \quad [\text{W/m}^2],$$

which is related to the Einstein energy–mass equivalence formula $E = mc_0^2$ (Sommerfeld, 1952).

For example, the power emitted by the sun is about 1360 [W/m$^2$], with a radiation pressure of $4 \times 10^{-6}$ [N/m$^2$] (i.e., 4 μPa) (Fitzpatrick, 2008). By way of comparison, the threshold audible acoustic pressure at the human eardrum at 1 [kHz] is $2 \times 10^{-3}$ [μPa]. Also

The lasers used in Inertial Confinement Fusion (e.g., the NOVA experiment in Lawrence Livermore National Laboratory) typically have energy fluxes of $10^{18}$ [W/m$^2$]. This translates to a radiation pressure of about $10^5$ atmospheres!


Electrical impedance seen by an electron: Up to now we have considered only the Brune impedance, which is a special case with no branch points or branch cuts. We can define impedance for diffusion, as in the diffusion of heat. There is also the diffusion of electrical and magnetic fields at the surface of a conductor, where the resistance of the conductor dominates the dielectric properties. This is called the electrical skin effect, where the conduction currents are dominated by the conductivity of the metal rather than the displacement currents. In such cases, the impedance is proportional to $\sqrt{\omega}$, which implies that it has a branch cut. Still, the real part of the impedance must be positive in the right $s$ half-plane, the required condition of all impedances, such that Postulate P3 (p. 122) is satisfied. The same effect is observed in acoustics (see Appendix D).

When we deal with Maxwell’s equations, the force is defined by the Lorentz force,

$$\mathbf{f} = q\mathbf{E} + q\mathbf{v} \times \mathbf{B} = q\mathbf{E} + \mathbf{C} \times \mathbf{B},$$

which is the force on a charge (e.g., electron) due to the electric $\mathbf{E}$ and magnetic $\mathbf{B}$ fields. The magnetic field plays a role when the charge has a velocity $\mathbf{v}$. When a charge is moving with velocity $\mathbf{v}$, it may be viewed as a current $\mathbf{C} = q\mathbf{v}$ (see the discussion on p. 3.9.1, p. 122).

The complex admittance density is

$$Y(s) = \sigma_0 + s\epsilon_0 \quad [\Omega/\text{m}^2]$$

(Feynman, 1970b, p. 13-1). Here $\sigma_0$ is the electrical conductivity and $\epsilon_0$ is the electrical permittivity. Since $\omega\epsilon_0 \ll \sigma_0$, this reduces to the resistance of the wire per unit length.\(^{15}\)

5.8 Potential solutions of Maxwell’s equations

The primary reason for using potentials is to generate solutions to Maxwell’s equations. For example, if we extend Eq. 5.1.2 (p. 170), we can express Maxwell’s equations in terms of scalar and vector potentials. These relations hips are (Sommerfeld (1952, p. 146), Feynman (1970d, p. 18-10)):

$$\mathbf{E}(x, t) = -\nabla \phi(x, t) - \frac{\partial \mathbf{A}(x, t)}{\partial t} \quad [\text{V/m}]$$

and

$$\mathbf{H}(x, t) = \frac{1}{\mu_0} \left[ \nabla \times \mathbf{A}(x, t) + \frac{\partial \mathbf{D}(x, t)}{\partial t} \right] \quad [\text{A/m}].$$

\(^{15}\)For copper $\omega \ll \omega_c = \sigma_0/\epsilon_0 \approx 6 \times 10^7/9 \times 10^{-12} \approx 6.66 \times 10^{18}$ [rad/s], or $f_c = 10^{18}$ [Hz], This corresponds to a wavelength of $\lambda_c \approx c_0/f_c = 0.30$ [μm]. For comparison, the Bohr radius (hydrogen) is $\approx 0.053$ [μm] (5.66 times smaller) and the Lorentz radius (of the electron) is estimated to be $2.8 \times 10^{-15}$ [μm] (2.8 [femto meters]).
We have extended \( H(x, t) \) to include the electric potential term

\[
D(x, t) = \epsilon(x, t) E(x, t) = -\epsilon(x, t) \nabla \phi(x, t)
\]

normally taken to be zero, because taking the curl of \( H(t) \) naturally removes any electrical potential term due to CoG.\(^{16}\)

When the permittivity \( \epsilon_o(x, t) \) is both inhomogeneous and time dependent,

\[
\nabla \cdot E = -\nabla^2 \Phi - \nabla \cdot \dot{A} = \rho(x, t) / \epsilon_o(x, t)
\]

and

\[
\nabla \times [\epsilon(x, \omega) \nabla \phi(x, \omega)] = \epsilon(x, t) \nabla \times \Phi + \nabla \epsilon(x, t) \times \nabla \phi \neq 0.
\]

The extension makes the potential solutions symmetric so that \( E \) and \( H \) each have electrical and magnetic excitation.

**Exercise #32**

Explain why some dependence on \( \phi(x, t) \) does not appear in Eq. 5.8.0.2, but does in 5.8.0.1.

**Solution:** For \( H(x, t) \) to depend on \( \phi(x, t) \) it must appear through the electric strength, as \( E(x, t) = -\nabla \Phi(x, t) \). But then \( \nabla \times H(x, t) \) would mean applying CoG (i.e., \( \nabla \times \nabla \phi = 0 \)) on the right side of the equation. Since this term would be zero, it is assumed to be zero, thus \( H(x, t) \) is only the dependent on only \( A(x, t) \). To fill out the symmetry, we have added \( \partial_t D(x, t) \) to Eq. 5.8.0.2, to see what might happen in the general case.

**Use of Helmholtz’s theorem on potential solutions:** The generalized solutions to Maxwell’s equations (Eqs. 5.8.0.1 and 5.8.0.2), pp. 205-205 have been expressed in terms of EM potentials \( \phi(x) \) and \( A(x) \) and Helmholtz’s theorem. These are solutions to Maxwell’s equations expressed in terms of the potentials \( \phi(x, s) \) and \( A(x, s) \), as determined at the boundaries (Sommerfeld, 1952, p. 146). These relations hips are invariant to certain functions added to each potential, as shown below. They are equivalent to Maxwell’s equations following the application of \( \nabla \times \nabla \times \).

Next we show that the potential equations (Eqs. 5.8.0.1, 5.8.0.2, p. 205) are consistent with Maxwell’s equations (Eq. 5.7.1.1).

**ME for \( E(x, t) \):** Taking the curl of Eq. 5.8.0.1, applying CoG = 0, and using Eq. 5.8.0.2, we find that

\[
\nabla \times E = -\nabla \times \nabla \phi - 0 \nabla \times \frac{\partial A}{\partial t}
\]

\[
= -\frac{\partial B}{\partial t}
\]

(5.8.0.3)

reverses Maxwell’s equation for \( E(x) \) (Eq. 5.7.1.1).

Taking the divergence of Eq. 5.8.0.2 and applying DoC = 0 gives Eq. 5.7.1.2 for \( B(x) \):

\[
\nabla \cdot B(x) = \nabla \nabla \times \Phi(x) = 0.
\]

**ME for \( H(x, t) \):** To recover Maxwell’s equation for \( H(x) \) (Eq. 5.7.1.1, \( \nabla \times H = C \)) from the potential equation (Eq. 5.8.0.2), we take the curl and use \( B = \epsilon_o H \) (Table 5.4, p. 202):

\[
\nabla \times B(x) = \mu_o \nabla \times H(x)
\]

\[
= \nabla \nabla \times A(x)
\]

\[
= \nabla A(x, t) - \nabla^2 A(x, t)
\]

\[
= \nabla \nabla \cdot A(x, t) - \frac{1}{c_o^2} \frac{\partial^2}{\partial t^2} A(x, t)
\]

\[
= -\frac{1}{c_o^2} \left[ \dot{A} + \nabla \dot{\Phi} \right] + \mu_o J.
\]

\(^{16}\)“invacuo” \( c_o = 8.85 \times 10^{-12} \) [F/m\(^2\)] is the capacitance, and \( \kappa c_o \) is the electric compliance-density of light. The related magnetic mass-density is the permeability \( \mu_o = 4\pi \times 10^{-7} \) [H/m\(^2\)] having an inductive impedance of \( s \mu_o \) [Ω/m]. It is helpful to think of \( \epsilon_o \) as a capacitance per unit area and \( \mu_o \) as an inductance per unit area (consistent with their units). The speed of light is \( c_o = 1/\sqrt{\epsilon_o \mu_o} = 3 \times 10^8 \) [m/s]. It is is slower when traveling in matter (Brillouin, 1960).
This last equation may be split into two independent equations by the use of Helmholtz theorem
\[ \nabla^2 \mathbf{A} - \frac{1}{c_0^2} \ddot{\mathbf{A}} = -\mu_0 \mathbf{J} \quad \text{and} \quad \nabla \cdot \mathbf{A} + \frac{1}{c_0^2} \dot{\Phi} = 0. \]

Taking the divergence of Eq. 5.8.0.2 and applying DoC = 0 gives Eq. 5.7.1.2 (\( \nabla \cdot \mathbf{D} = -\dot{\rho} \)). Alternatively,
\[ \nabla^2 \Phi - \frac{1}{c_0^2} \Phi = -\frac{\rho}{\epsilon_0}, \]

which is the scalar potential wave equation driven by the charge (Sommerfeld, 1952, p. 146).

**Summary:** In conclusion, Eq. 5.8.0.1, along with DoC = 0 and CoG = 0, gives Maxwell’s Eqs. 5.7.1.1 and 5.7.1.2 for \( \mathbf{E} \). Likewise, Eq. 5.8.0.2, along with DoC = 0 and CoG = 0, gives Maxwell’s Eqs. 5.7.1.1 and 5.7.1.2 for \( \mathbf{H} \). The above derivation for \( H(x, t) \) from \( \mathbf{A} \) and \( \Phi \) derives the magnetic component of the field, expressed in terms of its vector potential, in the same way as Eq. 5.7.1.1 describes \( \mathbf{E}(x, t) \) in terms of the potentials.

We may view the potential equations (Eqs 5.8.0.1 and 5.8.0.2) as equivalent to Maxwell’s equations; thus they are the solutions to ME.

**Exercise #33**

Starting with the values of the speed of light \( c_0 = 3 \times 10^8 \) m/s and the characteristic resistance of light waves \( r_0 = 377 \) ohms, use the formulas \( c_0 = 1/\sqrt{\mu_0 \epsilon_0} \) and \( r_0 = \sqrt{\epsilon_0 / \mu_0} \) to find values for \( \epsilon_0 \) and \( \mu_0 \).

**Solution:** Squaring \( c_0^2 = 1/\mu_0 \epsilon_0 \) and \( r_0^2 = \mu_0 / \epsilon_0 \), we may solve for the two unknowns: \( c_0^2 r_0^2 = 1/\mu_0 \epsilon_0 \epsilon_0 = 1/c_0^2 r_0^2 \); thus \( \epsilon_0 = 1/c_0 r_0 = \frac{10^{-8}}{2.998 \times 377} \approx 8.85 \times 10^{-12} \) Fd/m. Likewise, \( \mu_0 = r_0 / c_0 = (377/2.998) \times 10^{-8} \approx 125.75 \times 10^{-8} \) H/m. The value of \( \mu_0 \) is defined in the international SI units as \( 4\pi 10^{-7} \approx 12.56610^{-7} \) H/m.

It is easier to memorize \( c_0 \) and \( r_0 \), from which \( \epsilon_0 \) and \( \mu_0 \) may be quickly derived. \( \square \)

**Exercise #34**

Take the divergence of Maxwell’s equation for the magnetic intensity
\[ \nabla \times \mathbf{H}(x, t) = \mathbf{J}_m(x, t) + \frac{\partial}{\partial t} \mathbf{D}(x, t), \]

and explain what results (\( \mathbf{J}_m = \sigma \mathbf{E} \)).

**Solution:** The divergence of the curl is zero (DoC = 0),
\[ \nabla \cdot \nabla \times \mathbf{H}(x, t) = \nabla \cdot \mathbf{J}_m(x, t) + \frac{\partial}{\partial t} \mathbf{D}(x, t) = 0, \] (5.8.0.4)

which is conservation of charge (i.e., Gauss’s theorem). \( \square \)

**Helmholtz’s decomposition theorem:** Helmholtz’s decomposition theorem is expressed as the linear sum of a scalar potential \( \phi(x, y, z) \) (think voltage) and a vector potential (think magnetic vector potential). Specifically,
\[ \mathbf{E}(x, s) = -\nabla \phi(x, s) + \nabla \times \mathbf{A}(x, s), \] (5.8.0.5)

where \( \phi \) is the scalar and \( \mathbf{A} \) is the vector potential as a function of the Laplace frequency \( s \). Of course, this decomposition is general (not limited to the electromagnetic case). It applies to linear fluid vector fields, which include most liquids and air. When rotational and dilation become coupled, this relationship must break down.

To show how this relationship splits the vector field \( \mathbf{E} \) into its two parts, we need the two key vector identities that are always zero for analytic fields: the curl of the gradient (CoG),
\[ \nabla \times \nabla \phi(x) = 0, \] (5.8.0.6)

and the divergence of the curl\(^{12}\) (DoC)
\[ \nabla \cdot (\nabla \times \mathbf{A}) = 0. \] (5.8.0.7)

These identities are easily verified by working out a few specific examples based on the definitions of the three operators, gradient, divergence, and curl, or in terms of the operator’s integral definitions (see page 195). The

\(^{12}\)Helmholtz was the first person to apply mathematics in modeling the eye and the ear (Helmholtz, 1863a).
identities have a physical meaning: Every vector field may be split into its transnational and rotational parts. If \( E \) is the electric field \([V/m]\), \( \phi \) is the voltage and \( A \) is the induced rotational part, induced by a current.

By applying these two identities to Helmholtz’s decomposition, we can better appreciate the theorem’s significance. It is a form of proof actually, once we have satisfied ourselves that the vector identities are true. In fact, we can work backward using a physical argument that rotational momentum (rotational energy) is independent of transnational momentum. Once these forces are made clear, the vector operations all take on a well-defined meaning, and the mathematical constructions, centered around Helmholtz’s theorem, begin to provide some common-sense meaning. One could conclude that the physics is simply related to the geometry via the scalar and vector products.

Specifically, if we take the divergence of Eq. 5.8.0.5, and use the DoG, then

\[
-\frac{1}{\epsilon_0} \dot{\rho} = \nabla \cdot E = \nabla \cdot \{ -\nabla \phi + \nabla \times A \}^0 = -\nabla \cdot \nabla \phi = -\nabla^2 \phi,
\]

since the DoG zeros the vector potential \( A(x, y, z) \). If instead we use the CoG, then

\[
-\dot{B} = \nabla \times E = \nabla \times \{ -\nabla \phi + \nabla \times A \} = \nabla \times \nabla \times A = \nabla (\nabla \cdot A) - \nabla^2 A,
\]

since the CoG zeros the scalar field \( \phi(x, y, z) \). The last expression requires GoD.

5.9 The quasistatic approximation

A fundamental question that is begging for an answer, and that I have not yet seen asked, is

*What is the mathematical description of quantum mechanics?*

First, this question must have an answer because quantum mechanics (QM) is a highly mathematical subject. Second, we have seen that QM is related to the Webster horn equation and thus may be defined through its Sturm-Liouville equation parameters specifically, the area function \( A(x) \). Third, QM systems are nearly lossless. If there were zero loss, we would not be able to observe them. For all practical purposes, QM systems can be considered to be lossless until they interact with outside forces or particles. It is likely that in their ideal unperturbed state, there is virtually zero loss. Yet we can see the telltale radiation signature, such as the Rydberg series, for the hydrogen atom.

To characterize a lossless system, such as the hydrogen atom, as a Sturm-Liouville system, we need an area function that is exponential. The propagation function \( \kappa(s) \) has no real part, and the electromagnetic energy is trapped inside the area function (i.e., exponential horn). In this case, \( \kappa(s) \) has a cutoff frequency, below which the waves are trapped. The wave velocity in such cases is highly dispersive, giving rise to an accumulation point of the eigenfrequencies. We see this in the analysis of the Rydberg series Appendix H.1.1, which is the quintessential example of the QM system. In such a system \( \kappa(s) \) has two branches, corresponding to d’Alembert out and inbound electromagnetic waves, that are trapped by the exponential area function.

A number of assumptions and approximations result in special cases, many of which are classic. These manipulations are typically done at the differential equation level, making assumptions that change the basic equations that are to be solved. These approximations are distinct from assumptions we make while solving a specific problem. 18

Here are a few important examples of such approximations:

1. In-vacuo waves (free-space scalar wave equation)
2. The vector wave equation expressed in terms of scalar and vector potentials
3. Quasistatics:
   a. scalar wave equation
   b. Kirchhoff’s low-frequency lumped-element approximation networks
   c. Transmission line equations (telephone and telegraph equations)

Huygens made one of the first insights into wave propagation in about 1640.(see Fig. 1.5, p. 19).

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18Watch this magical video by Carl Sagan about Einstein’s views on the speed of light: [https://www.youtube.com/watch?v=_pEiA0-r5A8](https://www.youtube.com/watch?v=_pEiA0-r5A8).
**Quasistatics and its implications:** Quasistatics (Postulate P10, p. 122) is an approximation used to reduce a partial differential equation to a scalar (one-dimensional) equation (Sommerfeld, 1952); that is, quasistatic is a way of reducing a three-dimensional problem to a one-dimensional problem. So that it is not misapplied, it is important to understand the nature of this approximation, which goes to the heart of transmission line theory. The quasistatic approximation states that the wavelength $\lambda$ is greater than the dimensions of the object $\Delta$ (e.g., $\lambda \gg \Delta$). The best known examples—Kirchhoff’s current and voltage laws (KCL and KVL)—almost follow from Maxwell’s equations given the quasistatic approximation (Ramo et al., 1965). These laws, based on Ohm’s law, state that the sum of the currents at a node must be zero (KCL) and the sum of the voltages around a loop must be zero (KCL).

These well-known laws are the analogs of Newton’s laws of mechanics. The sum of the forces at a point is the analog of the sum of the loop voltages. Voltage $\phi$ is the force potential, since the electric field $E = -\nabla \phi$. The sum of the currents is the analog of the vector sum of velocities (mass) at a point, which is zero.

The acoustic wave equation describes how the scalar field pressure $p(x, t)$ and the vector force density potential $f(x, t) = -\nabla p(x, t) \, [N/m^2]$ propagate in three dimensions. The net force is the integral of the pressure gradient over an area. If the wave propagation is restricted to a pipe (e.g., organ pipe) or to a string (e.g., a guitar or lute), the transverse directions may be ignored because of the quasistatic approximation. What needs to be modeled by the equations is the wave propagation along the pipe (string). Thus we can approximate the restricted three-dimensional wave by a one-dimensional wave.

However, if we wish to be more precise about this reduction in geometry ($\mathbb{R}^2 \rightarrow \mathbb{R}$), we need to consider the quasistatic approximation, as it makes assumptions about what is happening in the other directions and quantifies the effect ($\lambda \gg \Delta$). If we consider wave propagation in a tube–say, the ear canal—the main wave direction is down the tube. But there is also wave propagation in the transverse direction, perpendicular to the direction of propagation. As shown in Table F.1 (p. 261), the key statement of the quasistatic approximation is that the wavelength in the transverse direction is much larger than the radius of the pipe. This is equivalent to saying that the radial wave reaches the walls and is reflected back in a time that is small compared to the distance propagated down the pipe. Clearly, the speed of sound down the pipe and in the transverse direction is the same if the medium is homogeneous (i.e., air or water). Thus the sound reaches the walls and is returned (reflected) to the center line in a time that the axial wave traveled about one diameter along the pipe. So, if the distance traveled is several diameters, the radial parts of the wave have time to come to equilibrium. The question one must ask is What are the properties of this equilibrium? The most satisfying answer is provided by looking at the internal forces on the air due to the gradients in the pressure.

The pressure $p(x, t)$ is a potential and thus its gradient is a force density $f(x, t) = -\nabla p(x, t)$. This equation tells us how the pressure wave evolves as it propagates down the horn. Any curvature in the pressure wavefront induces stresses that lead to changes (strains) in the local wave velocity, in the directions of the force density. The main force is driving the wavefront forward (down the horn), but there are radial (transverse) forces as well, which tend to rapidly go to zero.

For example, if the tube has a change in area (or curvature), the local forces create radial flow, which is immediately reflected by the walls due to the small distance to the walls, causing the forces to average out. After traveling a few diameters, these forces come to equilibrium and the wave trends toward a plane wave (or satisfies Laplace’s equation if the distortions of the tube are severe). The internal stress caused by this change in area quickly equilibrates.

There is a very important caveat, however: Only at low frequencies, such that $ka < 1$, can the plane wave mode dominate. At higher frequencies ($ka \geq 1$), where the wavelength is small compared to the diameter, the distance traveled between reflections is much greater than a few diameters. Fortunately the frequencies where this happens are so high that they play no role in frequencies that we care about in the ear canal. This effect results from cross-modes, which are radial and angular standing waves.

Such modes exist in the ear canal above 20 kHz. However, they are much more obvious on the eardrum, where the sound wave speed is much slower than that in air (Parent and Allen, 2010; Allen, 2014). Because of the slower speed, the eardrum has low-frequency cross-modes, and these may be seen in the ear canal pressure and are easily observable in ear canal impedence measurements. Yet they seem to have a negligible effect on our ability to hear sound with high fidelity. The point here is that the cross-modes are present, but we call upon the quasistatic approximation as a justification for ignoring them to get closer to the first-order physics.

### 5.9.1 Quasistatics and Quantum Mechanics

It is important to understand the meaning of Planck’s constant $h$, which appears in the relationships of both photons (light “particles”) and electrons (mass particles). If we could get a handle on what exactly Planck’s constant means, we might have a better understanding of quantum mechanics and physics in general. By cataloging the dispersion
relations (the relationships between the wavelength \( \lambda(\nu) \) and the frequency \( \nu \)) between electrons and photons, this may be attainable.

Basic relationships from quantum mechanics for photons and electrons include the following:

1. Photons (mass = 0, velocity = \( c \))
   
   (a) \( c = \lambda \nu \): The speed of light \( c \) is the product of its wavelength \( \lambda \) and its frequency \( \nu \). This relationship is for only monochromatic (single-frequency) light.

   (b) The speed of light is
   \[
   c_o = \frac{1}{\sqrt{\mu_o \varepsilon_o}} = 3 \times 10^8 \text{ [m/s]}. 
   \]

   (c) The characteristic resistance of light,
   \[ r_o = \sqrt{\mu_o / \varepsilon_o} = |E|/|H| = 377 \text{ [ohms]}. \]
   is defined as the magnitude of the ratio of the electric \( E \) and magnetic \( H \) fields, of a plane wave invacuo.

   (d) \( E = h\nu \): The photon energy is given by Planck’s constant
   \[ h \approx 6.623 \times 10^{-34} \text{ [joule \cdot s]} \]
times the frequency (i.e., bandwidth) of the photon.

2. Electrons (mass = \( m_e \), velocity \( V = 0 \))
   
   (a) \( E_e = m_e c^2 \approx 0.91 \cdot 10^{-30} \cdot 0.3 \cdot 10^{12} = 8.14 \times 10^{-20} \text{ [J]} \) is the electron rest energy (velocity \( V = 0 \)) of every electron of mass \( m_e = 9.1 \times 10^{-31} \text{ [kgm]}, \) where \( c_o \) is the speed of light.

   (b) \( p = h/\lambda \): The momentum \( p \) of an electron is given by Planck’s constant \( h \) divided by the wavelength of an electron \( \lambda \). It follows that the bandwidth of the photon is
   \[ \nu_e = \frac{E_e}{h} \]
   and the wavelength of an electron is
   \[ \lambda_e = \frac{h}{p_e}. \]

   One might reason that QM obeys the quasistatic (long-wavelength) approximation. If we compare the velocity of the electron \( V \) to the speed of light \( c \), then we see that
   \[ c_o = E/p \gg V = E/p = mV^2/mV. \]

Models of the electron: It is helpful to consider the physics of the electron, a negatively charged particle that is frequently treated as a single point in space. If its size were truly zero, there could be no magnetic moment (spin). The accepted size of the electron is known as the Lorentz radius, \( R = 2.8 \times 10^{-15} \text{ m} \). One could summarize the Lorentz radius as: Here lie many unsolved problems in physics. More specifically, at the dimensions of the Lorentz radius, what exactly is the structure of the electron?

Ignoring these difficulties, we can integrate the charge density of the electron over the Lorentz radius and place the total charge at a single point. Then we can make a grossly oversimplified model of the electron. For example, the electric displacement \( \mathbf{D} = \varepsilon_o \mathbf{E} \) (flux density) around a point charge is
\[
\mathbf{D} = -\varepsilon_o \nabla \phi(R) = -Q \nabla \left\{ \frac{1}{R} \right\} = -Q \delta(R) \text{ [C/m}^2].
\]

This formula is taught in many classic texts, but we should remember how crude this model of an electron is. Still, it describes the electric flux in an easily remembered form. However, computationally, it is less nice due to the delta function. The main limitation of this model is that the electron has a magnetic dipole moment (also called spin) that a simple point charge model does not capture. When placed in a magnetic field, due to the magnetic dipole, the electron aligns itself with the field.

We can apply a similar analysis to the gravitational potential. At the surface of the earth we are so far from the center of the earth that the potential appears to be linear because the height is a tiny fraction of the radius of the earth.
5.9. THE QUASISTATIC APPROXIMATION

5.9.2 Conjecture on photon energy

Photons are seen as quantized because they are commonly generated by atoms, which freely radiate photons (light particles) that have the difference in two energy (quantum, or eigenstates) levels. The relationship $E = h\nu$ does not inherently depend on $\nu$ being a fixed frequency. Planck’s constant $h$ is the EM energy density over the frequency, and $E(\nu_0)$ is the integral over the frequency:

$$E(\nu_0) = h\int_{-\nu_0}^{\nu_0} d\nu = 2h\nu_0.$$

When the photon is generated by an atom, $\nu_0$ is quantized by the energy level difference that corresponds to the frequency (energy level difference) of the photon jump.\(^{19}\)

Summary: Mathematics began as a simple way of counting things. But eventually physics and mathematics cleverly and mysteriously evolved to become tools to help us navigate our environment both locally and globally. These tools were used to: (1) deal with daily concerns such as food, water, and waste management, (2) understand the solar system and the stars, and (3) defend ourselves using tools of war, such as the hydrogen bomb. All powerful ideas have both bright and dark sides.

Based on the historical record of the abacus, one can infer that people precisely understood the concepts of counting, addition, subtraction, multiplication (recursive addition), and division (recursive subtraction with a fractional remainder). There is evidence that the abacus, a simple counting tool that formalized the addition of very large numbers, was introduced by the Romans to the Chinese, who used it for trade.

However, this working knowledge of arithmetic did not show up in written number systems. Roman numerals were not useful for calculations done on the abacus. Only the final answer could be expressed in terms of the Roman number system.

According to the known written record, the number zero had no written symbol until the time of Brahmagupta (628 CE). However, we should not assume the concept of zero was not understood simply because there was no symbol for it in the Roman numeral system. Negative numbers and zero would have been obvious when using the abacus. Numbers between the integers would naturally be represented as fractional numbers (\(\mathbb{F}\)), since any irrational number (\(\mathbb{Q}\)) may be approximated with arbitrary accuracy using fractional (\(\mathbb{F}\)) numbers.

Mathematics is the science of formalizing a repetitive method into a set of rules (an algorithm) and then generalizing it as much as possible. Generalizing the multiplication and division algorithms to different types of numbers becomes increasingly more interesting as we move from integers to rational numbers, irrational numbers, real and complex numbers, and ultimately vectors and matrices. How do we multiply two vectors, or multiply and divide one matrix by another? Multiplying and dividing polynomials (by long division) generalizes these operations even further. Linear algebra is another important generalization, fallout from the fundamental theorem of algebra and essential for the generalizations of the number systems.

Many concepts about numbers naturally evolved from music, where the length of a string (along with its tension) determines the pitch (Stillwell, 2010, pp. 11, 16, 153, 261). Decreasing the string’s length by half increases the frequency by a factor of 2. Quartering the length increases the frequency by a factor of 4. One octave is a factor of 2 and two octaves a factor of 4, while a half octave is $\sqrt{2}$. The musical scale was soon factored into rational parts. This scale almost worked, but did not generalize (sometimes known as the Pythagorean comma (Apel, 2003). Today’s well-tempered scale, is based on 12 equal geometric steps along one octave, or 1/12 octave ($\sqrt[12]{2} \approx 1.05946 \approx 18/17 = 1 + 1/17$).

But the concept of a factor was clear. Every number may be written as either a sum or a product (i.e., a repetitive sum). This led early mathematicians to the concept of a prime number, which is based on a unique factoring of every integer. At this same time (ca.5000 BCE), the solution of a second-degree polynomial was understood, which led to a generalization of factoring, since the polynomial, a sum of terms, may be written in factored form. If we think about this a bit, it is an amazing idea that needed to be discovered. This concept led to an important string of theorems on how to factor polynomials and how to numerically describe physical quantities. Newton was one of the first to master these tools with his proof that the orbits of the planets are ellipses, not circles. This led him to expand functions in terms of their derivatives and power series. Could these sums be factored? The solution to this problem led to calculus.

Mathematics, a product of the human mind, is a highly successful attempt to explain the physical world. All aspects of our lives were and are influenced by these tools. Mathematical knowledge is power. It allows us to think about complex problems in increasingly sophisticated ways. Does mathematics have a dark side? Perhaps

\(^{19}\)There is no better example of this than the properties of very large Rydberg atoms, as beautifully articulated by MIT Professor of Physics Daniel Kleppler: https://www.youtube.com/watch?v=e0IWPEhmMho.
no more than language itself. An equation is a mathematical sentence that expresses deep knowledge. Witness 
\[ E = mc^2 \quad \text{and} \quad \nabla^2 \psi = \ddot{\psi}. \]
5.10 Problems VC-2

Topics of this homework:
Maxwell’s equations (ME) and variables \((E, D; B, H)\), compressible and rotational properties of vector fields, fundamental theorem of vector calculus (Helmholtz’s theorem), wave equation

Notation: The following notation is used in this homework:
1. \(s = \sigma + j\omega\) is the Laplace frequency, as used in the Laplace transform.
2. A Laplace transform pair is indicated by the symbol \(\leftrightarrow\): for example, \(f(t) \leftrightarrow F(s)\).
3. \(\pi_k\) is the \(k\)th prime; for example, \(\pi_k \in \mathbb{P}, \pi_k = [2, 3, 5, 7, 11, 13, \ldots]\) for \(k = 1, \ldots, 6\).

Partial differential equations (PDEs): Wave equation

Problem #1: Solve the wave equation in one dimension by defining \(\xi = t \mp x/c\).

– 1.1: Show that d’Alembert’s solution, \(\varphi(x, t) = f(t - x/c) + g(t + x/c)\), is a solution to the acoustic pressure wave equation in one dimension:

\[
\frac{\partial^2 \varphi(x, t)}{\partial x^2} = \frac{1}{c^2} \frac{\partial^2 \varphi(x, t)}{\partial t^2},
\]

where \(f(\xi)\) and \(g(\xi)\) are arbitrary functions.

Problem #2: Solving the wave equation in spherical coordinates (i.e., three dimensions)

– 2.1: Write the wave equation in spherical coordinates \(\varphi(r, \theta, \phi, t)\). Consider only the radial term \(r\) (i.e., dependence on angles \(\theta\) and \(\phi\) is assumed to be zero). Hint: The form of the Laplacian as a function of the number of dimensions is given in Eq. 5.1.3.10 (page 174). Alternatively, look it up on the internet or in a calculus book.

– 2.2: Show that this equation is true:

\[
\nabla^2 \varphi(r) \equiv \frac{1}{r^2} \frac{\partial}{\partial r} r^2 \frac{\partial}{\partial r} \varphi(r) = \frac{1}{r} \frac{\partial^2}{\partial r^2} r \varphi(r).
\]

(VC-2.1)

Hint: Expand both sides of the equation.

– 2.3: Use the results from Eq. VC-2.4 to show that the solution to the spherical wave equation is

\[
\nabla^2 \varphi(r, t) = \frac{1}{c^2} \frac{\partial^2}{\partial t^2} \varphi(r, t) \quad (VC-2.2)
\]

\[
\varphi(r, t) = \frac{f(t - r/c)}{r} + \frac{g(t + r/c)}{r} \quad (VC-2.3)
\]

– 2.4: Using \(f(\xi) = \sin(\xi)u(\xi)\) and \(g(\xi) = e^\xi u(\xi)\), write the solutions to the spherical wave equation, where \(u(\xi)\) is the Heaviside step function.
– 2.5: Sketch this $f(\xi)$ and $g(\xi)$ for several times (e.g., 0, 1, and 2 seconds), and describe the behavior of the pressure $\rho(r, t)$ as a function of time $t$ and radius $r$.

– 2.6: What happens when the inbound wave reaches the center at $r = 0$?

**Helmholtz formula**

Every differentiable vector field may be written as the sum of a scalar potential $\phi$ and a vector potential $w$. This relationship is best known as the fundamental theorem of vector calculus (also called Helmholtz’s formula):

$$\mathbf{v} = -\nabla\phi + \nabla \times \mathbf{w}, \quad (VC-2.4)$$

This formula seems to be a natural extension of the algebraic products $A \mathbf{B} \perp A \times B$, since $A \mathbf{B} \propto ||A|| ||B|| \cos(\theta)$ and $A \times B \propto ||A|| ||B|| \sin(\theta)$, as developed in Appendix A.3.1, page 225. Thus these orthogonal components have magnitude 1 when we take the norm, due to Euler’s identity ($\cos^2(\theta) + \sin^2(\theta) = 1$).

As shown in Table 5.1 (p. 172), Helmholtz’s formula separates a vector field (i.e., $\mathbf{v}(x)$) into compressible and rotational parts:

1. The rotational (e.g., angular) part is defined by the vector potential $\mathbf{w}$, which requires that $\nabla \times \nabla \times \mathbf{w} \neq 0$.
   A field is irrotational (conservative) when $\nabla \times \mathbf{v} = 0$, meaning that the field $\mathbf{v}$ can be generated using only a scalar potential, $\mathbf{v} = -\nabla \phi$ (note that this is how a conservative field is usually defined, by saying there exists some $\phi$ such that $\mathbf{v} = -\nabla \phi$).

2. The compressible (e.g., radial) part of a field is defined by the scalar potential $\phi$, which requires that $\nabla \cdot \nabla \phi = \nabla^2 \phi \neq 0$. A field is incompressible (solenoidal) when $\nabla \cdot \mathbf{v} = 0$, meaning that the field $\mathbf{v}$ can be generated using only a vector potential, $\mathbf{v} = \nabla \times \mathbf{w}$.

The definitions and generating potential functions of irrotational (conservative) and incompressible (solenoidal) fields naturally follow from two key vector identities: (1) $\nabla \cdot (\nabla \times \mathbf{w}) = 0$ and (2) $\nabla \times (\nabla \phi) = 0$.

**Problem #3:** Define the following:

– 3.1: A conservative vector field
– 3.2: An irrotational vector field
– 3.3: An incompressible vector field
– 3.4: A solenoidal vector field
– 3.5: When is a conservative field irrotational?
– 3.6: When is an incompressible field irrotational?

**Problem #4:** For each of the following, (i) compute $\nabla \cdot \mathbf{v}$, (ii) compute $\nabla \times \mathbf{v}$, and (iii) classify the vector field (e.g., conservative, irrotational, incompressible, etc.).

– 4.1: $\mathbf{v}(x, y, z) = -\nabla (3y x^3 + y \log(xy))$

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20 A note about the relationship between the generating function and the test: You might imagine special cases where $\nabla \times \mathbf{w} \neq 0$ but $\nabla \times \nabla \times \mathbf{w} = 0$ (or $\nabla \phi \neq 0$ but $\nabla^2 \phi = 0$). In these cases, the vector (or scalar) potential can be recast as a scalar (or vector) potential. For example, consider a field $\mathbf{v} = \nabla \phi_0 + \mathbf{b}$, where $\mathbf{b} = x \hat{x} + y \hat{y} + z \hat{z}$. Note that $\mathbf{b}$ can actually be generated by either a scalar potential ($\phi_1 = \frac{1}{2}(x^2 + y^2 + z^2)$, such that $\nabla \phi_0 = b$) or a vector potential ($\mathbf{w}_0 = \frac{1}{2}(x \hat{x} + y \hat{y} + z \hat{z})$, such that $\nabla \times \mathbf{w}_0 = \mathbf{b}$). We find that $\nabla \times \mathbf{v} = 0$; therefore $\mathbf{v}$ must be irrotational. We say this irrotational field is generated by $\nabla \phi = \nabla (\phi_0 + \phi_1)$. 
\[-4.2: \mathbf{v}(x,y,z) = xy\mathbf{\hat{x}} - z\mathbf{\hat{y}} + f(z)\mathbf{\hat{z}}\]

\[-4.3: \mathbf{v}(x,y,z) = \nabla \times (x\mathbf{\hat{x}} - z\mathbf{\hat{y}})\]

Maxwell’s Equations

The variables have the following names and defining equations (see Table 5.4, p. 202):

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Equation</th>
<th>Name</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>(E)</td>
<td>(\nabla \times E = -\dot{B})</td>
<td>Electric field strength</td>
<td>[volts/m]</td>
</tr>
<tr>
<td>(D = \varepsilon_0 E)</td>
<td>(\nabla \cdot D = \rho)</td>
<td>Electric displacement (flux density)</td>
<td>[coul/m²]</td>
</tr>
<tr>
<td>(H)</td>
<td>(\nabla \times H = J + \dot{D})</td>
<td>Magnetic field strength</td>
<td>[amps/m]</td>
</tr>
<tr>
<td>(B = \mu_0 H)</td>
<td>(\nabla \cdot B = 0)</td>
<td>Magnetic induction (flux density)</td>
<td>[webers/m²]</td>
</tr>
</tbody>
</table>

Note that \(J = \sigma E\) is the current density (which has units of [amps/m²]). Furthermore, the speed of light in vacuo is \(c_0 = 3 \times 10^8 = 1/\sqrt{\mu_0 \varepsilon_0}\) [m/s], and the characteristic resistance of light \(r_0 = 377 = \sqrt{\mu_0/\varepsilon_0}\) [Ω] (i.e., ohms).

Speed of light

**Problem #5:** The speed of light in vacuo is \(c_0 = 1/\sqrt{\mu_0 \varepsilon_0} \approx 3 \times 10^8\) [m/s]. The characteristic resistance in vacuo is \(r_0 = \sqrt{\mu_0/\varepsilon_0} \approx 377\) [Ω].

- 5.1: Find a formula for the in-vacuo permittivity \(\varepsilon_0\) and permeability in terms of \(c_0\) and \(r_0\). Based on your formula, what are the numeric values of \(\varepsilon_0\) and \(\mu_0\)?

- 5.2: In a few words, identify the law given by this equation, define what it means, and explain the formula:

\[\int_S \mathbf{n} \cdot \mathbf{v} \, dA = \int_V \nabla \cdot \mathbf{v} \, dV.\]

Application of Maxwell’s equations

**Problem #6:** The electric Maxwell equation is \(\nabla \times E = -\dot{B}\), where \(E\) is the electric field strength and \(\dot{B}\) is the time rate of change of the magnetic induction field, or simply the magnetic flux density. Consider this equation integrated over a two-dimensional surface \(S\), where \(\mathbf{n}\) is a unit vector normal to the surface (you may also find it useful to define the closed path \(C\) around the surface):

\[\iint_S [\nabla \times \mathbf{E}] \cdot \mathbf{n} \, dS = -\frac{\partial}{\partial t} \iint_S \mathbf{B} \cdot \mathbf{n} \, dS.\]

- 6.1: Apply Stokes’s theorem to the left-hand side of the equation.

- 6.2: Consider the right-hand side of the equation. How is it related to the magnetic flux \(\Psi\) through the surface \(S\)?

- 6.3: Assume the right-hand side of the equation is zero. Can you relate your answer in question 6.1 to one of Kirchhoff’s laws?

**Problem #7:** The magnetic Maxwell equation is \(\nabla \times H = C \equiv J + \dot{D}\), where \(H\) is the magnetic field strength, \(J = \sigma E\) is the conductive (resistive) current density, and the displacement current \(\dot{D}\) is the time rate of change of the electric flux density \(D\). Here we defined a new variable \(C\) as the total current density.
7.1: First consider the equation over a two-dimensional surface $S$:

$$\int \int_S (\nabla \times H) \cdot \hat{n} dS = \int \int_S (J + \dot{D}) \cdot \hat{n} dS = \int \int_S C \cdot \hat{n} dS.$$ 

Then apply Stokes’s theorem to the left-hand side of this equation. In a sentence or two, explain the meaning of the resulting equation. Hint: What is the right-hand side of the equation?

Problem #8: Consider the next equation in three dimensions. Take the divergence of both sides and integrate over a volume $V$ (closed surface $S$):

$$\int \int \int_V \nabla \cdot (\nabla \times H) dV = \int \int \int_V \nabla \cdot C dV.$$ 

- 8.1: What happens to the left-hand side of this equation? Hint: Can you apply a vector identity? Apply the divergence theorem (sometimes known as Gauss’s theorem) to the right-hand side of the equation, and interpret your result. Hint: Can you relate your result to one of Kirchhoff’s laws?

Second-order differentials

Problem #9: This problem is about second-order vector differentials.

- 9.1: If $\mathbf{v}(x, y, z) = \nabla \phi(x, y, z)$, then what is $\nabla \cdot \mathbf{v}(x, y, z)$?

- 9.2: Evaluate $\nabla^2 \phi$ and $\nabla \times \nabla \phi$ for $\phi(x, y) = xe^y$.

- 9.3: Evaluate $\nabla \cdot (\nabla \times \mathbf{v})$ and $\nabla \times (\nabla \times \mathbf{v})$ for $\mathbf{v} = x\mathbf{i} + y\mathbf{j} + z\mathbf{k}$.

- 9.4: When $\mathbf{V}(x, y, z) = \nabla(1/x + 1/y + 1/z)$, what is $\nabla \times \mathbf{V}(x, y, z)$?

- 9.5: When was Maxwell born and when did he die? How long did he live (within $\pm 10$ years)?

Capacitor analysis

Problem #10: Find the solution to the Laplace equation between two infinite parallel plates separated by a distance $d$. Assume that the left plate at $x = 0$ is at voltage $V(0) = 0$ and the right plate at $x = d$ is at voltage $V_d \equiv V(d)$.

- 10.1: Write Laplace’s equation in one dimension for $V(x)$.

- 10.2: Write the general solution to your differential equation for $V(x)$.

- 10.3: Apply the boundary conditions $V(0) = 0$ and $V(d) = V_d$ to solve for the constants in your equation from question 10.2.

- 10.4: Find the charge density per unit area ($\sigma = Q/A$, where $Q$ is charge and $A$ is area) on the surface of each plate. Hint: $\mathbf{E} = -\nabla V$, and Gauss’s law states that $\int \int_S \mathbf{D} \cdot \hat{n} dS = Q_{enc}$.

- 10.5: Determine the per-unit-area capacitance $C$ of the system.

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21 We study plates that are infinite because this means the electric field lines are perpendicular to the plates, running directly from one plate to the other. However, we solve for per-unit-area characteristics of the capacitor.
Webster horn equation

**Problem # 11:** Horns illustrate an important generalization of the solution of the one-dimensional wave equation in regions where the properties (i.e., area of the tube) vary along the axis of wave propagation. Classic applications of horns are in vocal tract acoustics, loudspeaker design, cochlear mechanics, and any case that has wave propagation. Write the formula for the Webster horn equation, and explain the variables.
5.11 Further readings

The above concepts come straight from mathematical physics, as developed in the 17th–19th centuries. Much of this was first developed in acoustics by Helmholtz, Stokes and Rayleigh, following in Green’s footsteps, as described by Lord Rayleigh (1896). When it comes to fully appreciating Green’s theorem and reciprocity, I have found Rayleigh (1896) to be a key reference. To repeat my reading experience, start with Brillouin (1953, 1960), followed by Sommerfeld (1952) and Pipes (1958). Second-tier reading contains many items: Morse (1948); Sommerfeld (1949); Morse and Feshbach (1953); Ramo et al. (1965); Feynman (1970a); Boas (1987). A third tier might include Helmholtz (1863a); Fry (1928); Lamb (1932); Bode (1945); Montgomery et al. (1948); Beranek (1954); Fagen (1975); Lighthill (1978); Hunt (1952); Olson (1947). Other physics writings include the impressive series of mathematical physics books by stalwart authors, J.C. Slater and Landau and Lifshitz.22

Appendices
Appendix A

Notation

A.1 Number systems

The notation used in this book is defined in this appendix so that it may be quickly accessed. Where the definition is sketchy, page numbers are provided where these concepts are fully explained, along with many other important and useful definitions. For example a discussion of \( \mathbb{N} \) may be found on page 24. Math symbols such as \( \mathbb{N} \) may be found at the top of the index, since they are difficult to alphabetize.

A.1.1 Units

Strangely, or not, classical mathematics, as taught today in schools, does not seem to acknowledge the concept of physical units. Units seem to have been abstracted away. This makes mathematics distinct from physics, where almost everything has units. Presumably this makes mathematics more general (i.e., abstract). But for the engineering mind, this is not ideal, or worse, as it necessarily means that important physical meaning, by design, has been surgically removed. We shall use SI units whenever possible, which means this book is not not a typical book on mathematics. Spatial coordinates are quoted in meters \([\text{m}]\), and time in seconds \([\text{s}]\). Angles in degrees have no units, whereas radians have units of inverse-seconds \([\text{s}^{-1}]\). A complete list of SI units may be found at https://physics.nist.gov/cuu/pdf/sp811.pdf and Graham et al. (1994) for a discussion of basic math notation.

A.1.2 Symbols and functions

We use \( \ln \) as the log function base \( e \), \( \log \) as base 2, and \( \pi k \) to indicate the \( k \)th prime (e.g., \( \pi_1 = 2, \pi_2 = 3 \)).

When working with Fourier \( FT \) and Laplace \( LT \) transforms, lower case symbols are in the time domain while upper case indicates the frequency domain, as \( f(t) \leftrightarrow F(\omega) \). An important exception is Maxwell’s equations because they are so widely used as upper-case bold letters (e.g., \( \mathbf{E}(x, \omega) \)). It would seem logical to change this to \( \mathbf{e}(x, t) \leftrightarrow \mathbf{E}(x, \omega) \), to conform.

A.1.3 Common mathematical symbols

There are many pre-defined symbols in mathematics, too many to summarize here. We shall only use a small subset, defined here.

- A set is a collection of objects that have a common property, defined by braces. For example, if set \( P = \{a, b, c\} \) such that \( a^2 + b^2 = c^2 \), then members of \( P \) obey the Pythagorean theorem. Thus we could say that \( \{1, 1, \sqrt{2}\} \in P \).
- Number sets: \( \mathbb{N}, \mathbb{P}, \mathbb{Z}, \mathbb{Q}, \mathbb{F}, \mathbb{I}, \mathbb{R}, \mathbb{C} \) are briefly discussed below, and in greater detail in Sec. 2.1 (pp. 23–25).
- One can define sets of sets and subsets of sets, and this is prone (in my experience) to error. For example, what is the difference between the number 0 and the null set \( \emptyset = \{0\} \)? Is \( 0 \in \emptyset? \) Ask a mathematician. This seems a lackluster construction in the world of engineering.
- A vector is a column \( n \)-tuple. For example \([3, 5]^T = \begin{bmatrix} 3 \\ 5 \end{bmatrix} \).

\(^1\)https://en.wikipedia.org/wiki/List_of_mathematical_symbols_by_subject#Definition_symbols
APPENDIX A. NOTATION

- The symbol \( \perp \) is used in different ways to indicate two things are perpendicular, orthogonal, or in disjoint sets. In set theory \( A \perp B \) is equivalent to \( A \cap B = \emptyset \). If two vectors \( \mathbf{E}, \mathbf{H} \) are perpendicular \( \mathbf{E} \perp \mathbf{H} \), then their inner product \( \mathbf{E} \cdot \mathbf{H} = 0 \) is zero. One must infer the meaning of \( \perp \) from its context.

### Table A.1: List of all upper and lower case Greek letters used in the text.

<table>
<thead>
<tr>
<th>Greek letters</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \alpha )</td>
<td>( \alpha )</td>
</tr>
<tr>
<td>( \beta )</td>
<td>( \beta )</td>
</tr>
<tr>
<td>( \gamma )</td>
<td>( \gamma )</td>
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<tr>
<td>( \delta )</td>
<td>( \delta )</td>
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<tr>
<td>( \epsilon )</td>
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<td>( \lambda )</td>
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<tr>
<td>( \nu )</td>
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<tr>
<td>( \xi )</td>
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<td>( \omicron )</td>
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<tr>
<td>( \pi )</td>
<td>( \pi )</td>
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<td>( \sigma )</td>
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<td>( \tau )</td>
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<td>( \upsilon )</td>
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<td>( \phi )</td>
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<td>( \chi )</td>
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<tr>
<td>( \psi )</td>
<td>( \psi )</td>
</tr>
<tr>
<td>( \omega )</td>
<td>( \omega )</td>
</tr>
<tr>
<td>( \Omega )</td>
<td>( \Omega )</td>
</tr>
</tbody>
</table>

### Greek letters

Frequently Greek letters, as provided in Fig. A.1, are associated in engineering and physics with a specific physical meaning. For example, \( \omega \) \([\text{rad}]\) is the radian frequency \( 2\pi f \), \( \rho \) \([\text{kgm/m}^3]\) is commonly the density. \( \phi, \psi \) are commonly used to indicate angles of a triangle, and \( \zeta (s) \) is the Riemann zeta function. Many of these are so well established it makes no sense to define new terms, so we will adopt these common terms (and define them).

Likely you do not know all of these Greek letters, commonly used in mathematics. Some of them are pronounced in strange ways. The symbol \( \xi \) is pronounced “see,” \( \zeta \) is “zeta,” \( \beta \) is “beta,” and \( \chi \) is “kie” (rhymes with pie and sky). I will assume you know how to pronounce the others, which are more phonetic in English. One advantage of learning \LaTeX\, the powerful open-source math-oriented word-processing system used to write this book, is that math symbols are included, making then easily learned.

### Table A.2: Double-bold notation for the types of numbers. (#) is a page number. Symbol with an exponent denote the dimensionality. Thus \( \mathbb{R}^2 \) represents the real plane. An exponent of \( 0 \) denotes point, e.g., \( \mathbb{C}^0 \). It is reasonable to consider negative primes to be primes.

<table>
<thead>
<tr>
<th>Symbol (p. #)</th>
<th>Genus</th>
<th>Examples</th>
<th>Counter Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>( N ) (24)</td>
<td>Counting</td>
<td>( 1, 2, 17, 3, 10^{20} )</td>
<td>( -5, 0, \pi, -10.3, 5j )</td>
</tr>
<tr>
<td>( P ) (24)</td>
<td>Prime</td>
<td>( 2, 3, 17, 199, 23993 )</td>
<td>( 0, 1, 4, 3^2, 12 )</td>
</tr>
<tr>
<td>( Z ) (24)</td>
<td>Integer</td>
<td>( -1, 0, 17, 5j, -10^{20} )</td>
<td>( 1/2, \pi, \sqrt{5} )</td>
</tr>
<tr>
<td>( Q ) (24)</td>
<td>Rational</td>
<td>( 2/1, 3/2, 1.5, 1.14 )</td>
<td>( \sqrt{2}, 3^{-1/3}, \pi )</td>
</tr>
<tr>
<td>( F ) (24)</td>
<td>Fractional</td>
<td>( 1/2, 7/22 )</td>
<td>( 1/\sqrt{2} )</td>
</tr>
<tr>
<td>( \mathbb{H} ) (25)</td>
<td>Irrational</td>
<td>( \sqrt{3}^{-1/3}, \pi, e )</td>
<td>Vectors</td>
</tr>
<tr>
<td>( \mathbb{R} ) (25)</td>
<td>Reals</td>
<td>( \sqrt{2}, 3^{-1/3}, \pi )</td>
<td>( 2\pi j )</td>
</tr>
<tr>
<td>( \mathbb{C} ) (25)</td>
<td>Complex</td>
<td>( 1, \sqrt{2}j, 3^{-\sqrt{3}}, \pi j )</td>
<td>Vectors</td>
</tr>
<tr>
<td>( G )</td>
<td>Gaussian integers</td>
<td>( 3 - 2j \in \mathbb{Z} \cup \mathbb{C} )</td>
<td>complex integers</td>
</tr>
</tbody>
</table>

### Double-Bold notation

Table A.2 indicates the symbol followed by a page number indication where it is discussed, and the Genus (class) of the number type. For example, \( N > 0 \) indicates the infinite set of counting numbers \( \{1, 2, 3, \cdots \} \), not including zero. Starting from any counting number, you get the next one by adding 1. Counting numbers are sometimes called the natural or cardinal numbers.

We say that a number is in the set with the notation \( 3 \in \mathbb{N} \subseteq \mathbb{R} \), which is read as “3 is in the set of counting numbers, which in turn is in the set of real numbers,” or in vernacular language “3 is a real counting number.”

Prime numbers \( \mathbb{P} \cap \mathbb{N} \) are taken from the counting numbers, but do not include 1.
The signed integers \( \mathbb{Z} \) include 0 and negative integers. Rational numbers \( \mathbb{Q} \) are historically defined to include \( \mathbb{Z} \), a somewhat inconvenient definition, since the more interesting class are the fractionals \( \mathbb{F} \), a subset of rationals \( \mathbb{F} \in \mathbb{Q} \) that exclude the integers (i.e., \( \mathbb{F} \perp \mathbb{Z} \)). This is a useful definition because the rationals \( \mathbb{Q} = \mathbb{Z} \cup \mathbb{F} \) are formed from the union of integers and fractionals.

The rationals may be defined, using set notation (a very sloppy notation with an incomprehensible syntax), as

\[
\mathbb{Q} = \{ p/q : q \neq 0 \& p, q \in \mathbb{Z} \},
\]

which may be read as “the set ‘\{\ldots\}’ of all \( p/q \) such that ‘\: q \neq 0, \:’ and ‘\: p, q \in \mathbb{Z} \’. The translation of the symbols is in single (‘\ldots’) quotes.

Irrational numbers \( \mathbb{I} \) are very special: They are formed by taking a limit of fractionals, as the numerator and denominator \( \to \infty \), and approach a limit point. It follows that irrational numbers must be approximated by fractionals.

The reals (\( \mathbb{R} \)) include complex numbers (\( \mathbb{C} \)) having a zero imaginary part (i.e., \( \mathbb{R} \subset \mathbb{C} \)).

The size of a set is denoted by taking the absolute value (e.g., \(|\mathbb{N}|\)). Normally in mathematics this symbol indicates the cardinality, so we are defining it differently from the standard notation.

### A.1.4 Classification of numbers:

From the above definitions there exists a natural hierarchical structure of numbers:

\[
\mathbb{P} \subset \mathbb{N}, \quad \mathbb{Z} : \{ \mathbb{N}, 0, -\mathbb{N} \}, \quad \mathbb{F} \perp \mathbb{Z}, \quad \mathbb{Q} : \mathbb{Z} \cup \mathbb{F}, \quad \mathbb{R} : \mathbb{Q} \cup \mathbb{I} \subset \mathbb{C}
\]

1. The primes are a subset of the counting numbers: \( \mathbb{P} \subset \mathbb{N} \).
2. The signed integers \( \mathbb{Z} \) are composed of \( \pm \mathbb{N} \) and 0, thus \( \mathbb{N} \subset \mathbb{Z} \).
3. The fractionals \( \mathbb{F} \) do not include that of the signed integers \( \mathbb{Z} \).
4. The rationals \( \mathbb{Q} = \mathbb{Z} \cup \mathbb{F} \) are the union of the signed integers and fractionals.
5. Irrational numbers \( \mathbb{I} \) have the special properties \( \mathbb{I} \perp \mathbb{Q} \).
6. The reals \( \mathbb{R} : \mathbb{Q}, \mathbb{I} \) are defined as the union of those rationals and irrational \( \mathbb{I} \).
7. Reals \( \mathbb{R} \) may be defined as a subset of those complex numbers \( \mathbb{C} \) having zero imaginary part.

### A.1.5 Rounding schemes

In Matlab/Octave there are five different rounding schemes (i.e., mappings): \( \text{round}(x) \), \( \text{fix}(x) \), \( \text{floor}(x) \), \( \text{ceil}(x) \), \( \text{roundb}(x) \), with input \( x \in \mathbb{R} \) and output \( k \in \mathbb{N} \). For example \( 3 = \lceil \pi \rceil, 3 = \lfloor e \rfloor = 2.7183 \) rounds to the nearest integer, whereas \( 3 = \text{floor}(\pi) \) rounds down while \( 3 = \text{ceil}(e) = 2.7183 \) rounds up. Rounding schemes are used for quantizing a number and generating a remainder. For example: \( y = \text{rem}(x) \) is equivalent to \( y = x - \lfloor x \rfloor \). Note \( \text{round}(\pi) \equiv \lfloor \pi \rceil \) introduces negative remainders in the remainder when ever a number rounds up (\( \pi = \lfloor \pi \rceil - 0.8541 \)).

The continued fraction algorithm (CFA), Sec. 2.5.4 (p. 47) is a recursive rounding scheme, operating on the reciprocal of the remainder. For example:

\[
\exp(1) = 3 + 1/(−4 + 1/(2 + 1/(5 + 1/(−2 + 1/(−7))))) = o(1.75 \times 10^{−6}),
\]

\[
= [3; −4, 2, 5, −2, −7] − o(1.75 \times 10^{−6}).
\]

The expressions in brackets is a notation for the CFA integer coefficients. The Octave/Matlab function having \( x \in \mathbb{R} \), is either \( \text{rat}(x) \) with output \( \in \mathbb{N} \), or \( \text{rats}(x) \), with output \( \in \mathbb{F} \).

### A.1.6 Fourier series

Fourier series tells us that periodic functions are discrete in frequency, with frequencies given by \( n T_s \), where \( T_s \) is the sample period. The discrete FT (DFT) is a good example. When using the DFT, the sample period is \( T_s = 1/2F_{\text{max}} \) and the minimum and maximum frequencies are given by \( F_{\text{min}} = F_{\text{max}}/(N\text{F}) \) where \( \text{F} \) is the size of the DFT.

This concept is captured by the Fourier series, which is a frequency expansion of a periodic function. This concept is quite general. Periodic in frequency implies discrete in time. Periodic and discrete in time requires
periodic and discrete in frequency (the case of the DFT). The modulo function \( x \equiv y \mod \mathbb{R} \) is periodic with period \( y, x \in \mathbb{R} \).

A periodic function may be conveniently indicated using double-parentheses notation. This is sometimes known as modular arithmetic. For example,

\[
f((t))_T = f(t) = f(t + kT)
\]

is periodic on \( t, T \in \mathbb{R} \) with a period of \( T \) and \( k \in \mathbb{Z} \). This notation is useful when dealing with Fourier series of periodic functions such as \( \sin(\theta) \) where \( \sin(\theta) = \sin((\theta))_2\pi = \mod(\sin(\theta), 2\pi) \).

When a discrete valued (e.g., time \( t \in \mathbb{N} \)) sequence is periodic with period \( N \in \mathbb{Z} \), we use square brackets

\[
f[[n]]_N = f[n] = f[n \pm kN],
\]

with \( k \in \mathbb{Z} \). This notation will be used with discrete-time signals that are periodic, such as the case of the DFT.

It is common for fractions to repeat. For example \( 1/7 = 0.((142857)) \) where the double brackets indicates this number repeats. That is \( 1/7 = 0.142857, 142857, 142857, 142857, \ldots \).  

## A.2 Differential equations vs. polynomials

A polynomial has degree \( N \) defined by the largest power. A quadratic equation is degree 2, and a cubic has degree 3. We shall indicate a polynomial by the notation

\[ P_N(z) = z^N + a_{N-1}z^{N-1} \cdots a_0. \]

It is a good practice to normalize the polynomial so that \( a_N = 1 \). This will not change the roots, defined by Eq. 3.1.1.7 (p. 64). The coefficient on \( z^{N-1} \) is always the sum of the roots \( z_n (a_{N-1} = \sum_n z_n) \), and \( a_0 \) is always their product \( (a_0 = \prod_n z_n) \).

Differential equations have order (polynomials have degree). If a second-order differential equation is Laplace transformed (Lec. 3.9, p. 119), one is left with a degree 2 polynomial. For example,

\[
\frac{d^2}{dt^2} y(t) + b \frac{d}{dt} y(t) + cy(t) = \alpha \left( \frac{d}{dt} x(t) + \beta x(t) \right) \leftrightarrow (A.2.0.1)
\]

\[
(s^2 + bs + c)Y(s) = \alpha(s + \beta)X(s). \quad \leftrightarrow \quad \frac{Y(s)}{X(s)} = \alpha \frac{s + \beta}{s^2 + bs + c} \equiv H(s) \leftrightarrow h(t). \quad \leftrightarrow \quad (A.2.0.2)
\]

Using the same argument as for polynomials, the lead coefficient must always be 1. The coefficient \( \alpha \in \mathbb{R} \) is called the gain. The complex variable \( s \) is the Laplace frequency.

The ratio of the output \( Y(s) \) over the input \( X(s) \) is called the system transfer function \( H(s) \). When \( H(s) \) is the ratio of two polynomials in \( s \), the transfer function is said to be bilinear, since it is linear in both the input and output. The roots of the numerator are called the zeros and those of the denominator, the poles. The inverse Laplace transform of the transfer function is called the system impulse response, which describes the system’s output signal \( y(t) \) for any given input signal \( x(t) \), via convolution (i.e., \( y(t) = h(t) \ast x(t) \)).

## A.3 Matrix algebra: Systems

### A.3.1 Vectors

Vectors as columns of ordered sets of scalars \( \in \mathbb{C} \). When we write them out in text, we typically use row notation, with the transpose symbol:

\[
[a, b, c]^T = \begin{bmatrix} a \\ b \\ c \end{bmatrix}.
\]

This is strictly to save space on the page. The notation for conjugate transpose is \( \dagger \), for example

\[
\begin{bmatrix} a \\ b \\ c \end{bmatrix}^\dagger = [a^* \quad b^* \quad c^*].
\]

The above example is said to be a 3-dimensional vector because it has three components.
Row vs. column vectors: With rare exceptions, vectors are columns, denoted column-major. To avoid confusion, it is a good rule to make your mental default column-major, in keeping with most signal processing (vectorized) software. Column vectors are the unstated default of Matlab/Octave, only revealed when matrix operations are performed. The need for the column (or row) major is revealed as a consequence of efficiency when accessing long sequences of numbers from computer memory. For example, when forming the sum of many numbers using the Matlab/Octave command \( \sum(A) \), where \( A \) is a matrix, Matlab/Octave operates on the columns, returning a row vector of column sums:

\[
\sum \begin{bmatrix} 1 & 2 \\ 3 & 4 \end{bmatrix} = [4, 6].
\]

If the data were stored in row-major order, the answer would be the column vector \( \begin{bmatrix} 3 \\ 7 \end{bmatrix} \). Thus Matlab/Octave is column-major by default.

A.3.2 Vector products

A scalar product (aka dot product) is defined to “weight” vector elements before summing them, resulting in a scalar. The transpose of a vector (a row-vector) is typically used as a scale factor (i.e., weights) on the elements of a vector. For example,

\[
\begin{bmatrix} 1 \\ 2 \\ -1 \end{bmatrix} \cdot \begin{bmatrix} 1 \\ 2 \\ -1 \end{bmatrix}^T = \begin{bmatrix} 1 \\ 2 \\ 3 \end{bmatrix} = 1 + 2 \cdot 2 - 3 = 2.
\]

A more interesting example is

\[
\begin{bmatrix} 1 \\ 2 \\ 4 \end{bmatrix} \cdot \begin{bmatrix} 1 \\ s \\ s^2 \end{bmatrix} = \begin{bmatrix} 1 \\ 2 \\ 4 \end{bmatrix}^T = \begin{bmatrix} 1 \\ 2 \\ 4 \\ -1 \end{bmatrix} = 1 + 2s + 4s^2.
\]

Polar scalar product: The vector-scalar product in polar coordinates is (Fig. 3.4, p. 94)

\[ B \cdot C = ||B|| ||C|| \cos \theta \in \mathbb{R}, \]

where \( \cos \theta \in \mathbb{R} \) is called the direction-cosine between \( B \) and \( C \).

Polar wedge product: The vector wedge product in polar coordinates is (Fig. 3.4, p. 94)

\[ B \wedge C = ||B|| ||C|| \sin \theta \in \mathbb{R}, \]

where \( \sin \theta \in \mathbb{R} \) is therefore the direction-sine between \( B \) and \( C \).

Complex polar vector product: From these two polar definitions and \( e^{i\theta} = \cos \theta + j \sin \theta \),

\[ B \cdot C + jB \wedge C = ||B|| ||C|| e^{i\theta}. \]

Hence

\[ |B \cdot C|^2 + |B \wedge C|^2 = ||B||^2 ||C||^2 \cos^2 \theta + ||B||^2 ||C||^2 \sin^2 \theta = ||B||^2 ||C||^2. \]

This relationship holds true in any vector space, of any number of dimensions, containing vectors \( B \) and \( C \). In this case \( s = \sigma + \omega j \in \mathbb{C} \) can be the Laplace frequency. Jaynes (1991) has an relevant discussion about this type of vector product.

A.3.3 Norms of vectors

The norm of a vector is the scalar product of the vector with itself

\[ ||A|| = \sqrt{A \cdot A} \geq 0, \]

forming the Euclidean length of the vector.

\(^2\)https://en.wikipedia.org/wiki/Row-_and_column-major_order

\(^3\)In contrast, reading words in English is ‘row-major.”
Euclidean distance between two points in $\mathbb{R}^3$: The scalar product of the difference between two vectors $(A - B) \cdot (A - B)$ is the Euclidean distance between the points they define

$$\|A - B\| = \sqrt{(a_1 - b_1)^2 + (a_2 - b_2)^2 + (a_3 - b_3)^2}.$$

Triangle inequality

$$\|A + B\| = \sqrt{(a_1 + b_1)^2 + (a_2 + b_2)^2 + (a_3 + b_3)^2} \leq \|A\| + \|B\|.$$

In terms of a right triangle this says the sum of the lengths of the two sides is greater to the length of the hypotenuse, and equal when the triangle degenerates into a line.

Vector cross product: The vector product (aka cross product) $A \times B = \|A\| \|B\| \sin \theta$ is defined between the two vectors $A$ and $B$. In Cartesian coordinates

$$A \times B = \det \begin{vmatrix} \hat{x} & \hat{y} & \hat{z} \\ a_1 & a_2 & a_3 \\ b_1 & b_2 & b_3 \end{vmatrix}.$$

The triple product: This is defined between three vectors as

$$A \cdot (B \times C) = \det \begin{vmatrix} a_1 & a_2 & a_3 \\ b_1 & b_2 & b_3 \\ c_1 & c_2 & c_3 \end{vmatrix}.$$

This may be indicated without the use of parentheses, since there can be no other meaningful interpretation. However for clarity, parentheses should be used. The triple product is the volume of the parallelepiped (3D-crystal shape) outlined by the three vectors, as shown in Fig. 3.4 p. 94.

Dialects of vector notation: Physical fields are, by definition, functions of space $x \ [\text{m}]$, and in the most general case, time $t \ [\text{s}]$. When Laplace transformed, the fields become functions of space and complex frequency (e.g., $E(x, t) \leftrightarrow \tilde{E}(x, s)$). As before, there are several equivalent vector notations. For example, $E(x, t) = [E_x, E_y, E_z]^T = E_x(x, t)\hat{x} + E_y(x, t)\hat{y} + E_z(x, t)\hat{z}$ is “in-line,” to save space. The same equation may written in “displayed” notation as:

$$\begin{bmatrix} E_x(x, t) \\ E_y(x, t) \\ E_z(x, t) \end{bmatrix} = \begin{bmatrix} E_x \\ E_y \\ E_z \end{bmatrix}^T = E_x \hat{x} + E_y \hat{y} + E_z \hat{z}.$$

Note the three notations for vectors, bold font, element-wise columns, element-wise transposed rows and dyadic format. These are all shorthand notations for expressing the vector. Such usage is similar to a dialect in a language.

Complex elements: When the elements are complex ($\in \mathbb{C}$), the transpose is defined as the complex conjugate of the elements. In such complex cases the transpose conjugate may be denoted with a $\dagger$ rather than $T$

$$\begin{bmatrix} -2j \\ 3j \\ 1 \end{bmatrix}^\dagger = \begin{bmatrix} 2j \\ -3j \\ 1 \end{bmatrix} \in \mathbb{C}.$$

For this case when the elements are complex, the dot product is a real number

$$\mathbf{a} \cdot \mathbf{b} = \mathbf{a}^\dagger \mathbf{b} = \begin{bmatrix} a_1^* & a_2^* & a_3^* \end{bmatrix} \begin{bmatrix} b_1 \\ b_2 \\ b_3 \end{bmatrix} = a_1^* b_1 + a_2^* b_2 + a_3^* b_3 \in \mathbb{R}.$$
A.3. MATRIX ALGEBRA OF SYSTEMS

**Norm of a complex vector:** The dot product of a vector with itself is called the norm of \( \mathbf{a} \)

\[
||\mathbf{a}|| = \sqrt{\mathbf{a}^\dagger \mathbf{a}} \geq 0.
\]

which is always non-negative.

Such a construction is useful when \( \mathbf{a} \) and \( \mathbf{b} \) are related by an impedance matrix

\[
\mathbf{V}(s) = \mathbf{Z}(s)\mathbf{I}(s)
\]

and we wish to compute the power. For example, the impedance of a mass is \( m\mathbf{s} \) and a capacitor is \( 1/s\mathbf{C} \). When given a system of equations (a mechanical or electrical circuit) one may define an impedance matrix.

**Complex power:** In this special case, the complex power \( \mathcal{P}(s) \in \mathbb{R}(s) \) is defined, in the complex frequency domain \( (s) \), as

\[
\mathcal{P}(s) = \mathbf{I}^\dagger(s)\mathbf{V}(s) = \mathbf{I}^\dagger(s)\mathbf{Z}(s)\mathbf{I}(s) \leftrightarrow \mathcal{P}(t) \quad [\mathbb{W}].
\]

The real part of the complex power must be positive. The imaginary part corresponds to available stored energy.

The case of three-dimensions is special, allowing definitions that are not easily defined in more than three dimensions. A vector in \( \mathbb{R}^3 \) labels the point having the coordinates of that vector.

### A.3.4 Matrices

When working with matrices, the role of the weights and vectors can change, depending on the context. A useful way to view a matrix is as a set of column vectors, weighted by the elements of the column-vector of weights multiplied from the right. For example,

\[
\begin{bmatrix}
  a_{11} & a_{12} & a_{13} & \cdots & a_{1M} \\
  a_{21} & a_{22} & a_{23} & \cdots & a_{2M} \\
  \vdots & \vdots & \vdots & \ddots & \vdots \\
  a_{N1} & a_{N2} & a_{N3} & \cdots & a_{NM}
\end{bmatrix}
\begin{bmatrix}
  w_1 \\
  w_2 \\
  \vdots \\
  w_M
\end{bmatrix}
= w_1 \begin{bmatrix} a_{11} \\ a_{21} \\ \vdots \\ a_{N1} \end{bmatrix} + w_2 \begin{bmatrix} a_{12} \\ a_{22} \\ \vdots \\ a_{N2} \end{bmatrix} + \cdots + w_M \begin{bmatrix} a_{1M} \\ a_{2M} \\ \vdots \\ a_{NM} \end{bmatrix},
\]

where the weights are \( [w_1, w_2, \ldots, w_M]^T \). Alternatively, the matrix is a set of row vectors of weights, each of which is applied to the column vector on the right \( ([w_1, w_2, \ldots, W_M]^T) \).

The determinant of a matrix is denoted as either \( \det \mathbf{A} \) or simply \( |\mathbf{A}| \) (as in the absolute value). The inverse of a square matrix is \( \mathbf{A}^{-1} \) or \( \text{inv} \mathbf{A} \). If \( |\mathbf{A}| = 0 \), the inverse does not exist. \( \mathbf{AA}^{-1} = \mathbf{A}^{-1}\mathbf{A} \).

Matlab/Octave’s notational convention for a row-vector is \([a, b, c]\) and a column-vector is \([a; b; c]\). A prime on a vector takes the complex conjugate transpose. To suppress the conjugation, place a period before the prime. The \( : \) argument converts the array into a column vector, without conjugation. A tacit notation in Matlab is that vectors are columns and the index to a vector is a row vector. Matlab defines the notation \( 1:4 \) as the “row-vector” \([1, 2, 3, 4]\), which is unfortunate as it leads users to assume that the default vector is a row. This can lead to serious confusion later, as Matlab’s default vector is a column. I have not found the above convention explicitly stated, and it took me years to figure this out for myself.

When writing a complex number we shall adopt \( \langle 1 \rangle \) to indicate \( \sqrt{-1} \). Matlab/Octave allows either \( 1 \) or \( 1 \). Units are SI; angles are in degrees \([\text{deg}]\) unless otherwise noted. The units for \( \pi \) are always radians \([\text{rad}]\). For example \( \sin(\pi) = e^{i\pi/2} \).

### A.3.5 2 \times 2 complex matrices

Here are some definitions to learn:

1. **Scalar:** A number – for example \( \{a, b, c, \alpha, \beta, \ldots\} \in \{\mathbb{Z}, \mathbb{Q}, \mathbb{I}, \mathbb{R}, \mathbb{C}\} \)

2. **Vector:** A quantity having direction as well as magnitude, often denoted by a bold letter \( \mathbf{x} \), or with an arrow over the top \( \vec{x} \). In matrix notation, this is typically represented as a single row \([x_1, x_2, x_3, \ldots]^T \) or single column \([x_1, x_2, x_3, \ldots]^T \) (where \( T \) indicates the transpose). In this class we will typically use column vectors. The vector may also be written out using unit vector notation to indicate direction. For example: \( \mathbf{x}_{3\times1} = x_1\mathbf{\hat{x}} + x_2\mathbf{\hat{y}} + x_3\mathbf{\hat{z}} = [x_1, x_2, x_3]^T \), where \( \mathbf{\hat{x}}, \mathbf{\hat{y}}, \mathbf{\hat{z}} \) are unit vectors in the \( x, y, z \) Cartesian directions (here the vector’s subscript \( 3 \times 1 \) indicates its dimensions). The type of notation used may depend on the engineering problem you are solving.
3. Matrix: \( A = \begin{bmatrix} a_{11}, a_{21}, a_{31}, \ldots, a_{M1} \\ a_{12}, a_{22}, a_{32}, \ldots, a_{M2} \\ \vdots \end{bmatrix}_{N \times M} = \{a_{n,m}\}_{N \times M} \) can be a non-square matrix if the number of elements in each of the vectors \((N)\) is not equal to the number of vectors \((M)\). When \(M = N\), the matrix is square. It may be inverted if its determinant \(|A| = \prod \lambda_k \neq 0\) (where \(\lambda_k\) are the eigenvalues). In this text we work only with \(2 \times 2\) and \(3 \times 3\) square matrices.

4. Linear system of equations: \( Ax = b \) where \(x\) and \(b\) are vectors and matrix \(A\) is a square.

   (a) Inverse: The solution of this system of equations may be found by finding the inverse \(x = A^{-1}b\).

   (b) Equivalence: If two systems of equations \(A_0x = b_0\) and \(A_1x = b_1\) have the same solution (i.e., \(x = A_0^{-1}b_0 = A_1^{-1}b_1\)), they are said to be equivalent.

   (c) Augmented matrix: The first type of augmented matrix is defined by combining the matrix with the right-hand side. For example, given the linear system of equations of the form \(Ax = y\)

   \[
   \begin{bmatrix}
   a & b \\
   c & d
   \end{bmatrix}
   \begin{bmatrix}
   x_1 \\
   x_2
   \end{bmatrix}
   =
   \begin{bmatrix}
   y_1 \\
   y_2
   \end{bmatrix},
   \]

   the augmented matrix is

   \[
   [A|y] = \begin{bmatrix}
   a & b \\
   c & d
   \end{bmatrix}
   \begin{bmatrix}
   y_1 \\
   y_2
   \end{bmatrix}.
   \]

   A second type of augmented matrix may be used for finding the inverse of a matrix (rather than solving a specific instance of linear equations \(Ax = b\)). In this case the augmented matrix is

   \[
   [A|I] = \begin{bmatrix}
   a & b \\
   c & d
   \end{bmatrix}
   \begin{bmatrix}
   1 & 0 \\
   0 & 1
   \end{bmatrix}.
   \]

   Performing Gaussian elimination on this matrix, until the left side becomes the identity matrix, yields \(A^{-1}\). This is because multiplying both sides by \(A^{-1}\) gives \(A^{-1}A|A^{-1}I = I|A^{-1}\).

5. Permutation matrix (P): A matrix that is equivalent to the identity matrix, but with scrambled rows (or columns). Such a matrix has the properties \(\det(P) = \pm 1\) and \(P^2 = I\). For the \(2 \times 2\) case, there is only one permutation matrix:

   \[
   P = \begin{bmatrix}
   0 & 1 \\
   1 & 0
   \end{bmatrix} \quad P^2 = \begin{bmatrix}
   0 & 1 \\
   1 & 0
   \end{bmatrix} \begin{bmatrix}
   0 & 1 \\
   1 & 0
   \end{bmatrix} = \begin{bmatrix}
   1 & 0 \\
   0 & 1
   \end{bmatrix}.
   \]

   A permutation matrix \(P\) swaps rows or columns of the matrix it operates on. For example, in the \(2 \times 2\) case, pre-multiplication swaps the rows,

   \[
   PA = \begin{bmatrix}
   0 & 1 \\
   1 & 0
   \end{bmatrix}
   \begin{bmatrix}
   a & b \\
   \alpha & \beta
   \end{bmatrix} = \begin{bmatrix}
   \alpha & \beta \\
   a & b
   \end{bmatrix},
   \]

   whereas post-multiplication swaps the columns,

   \[
   AP = \begin{bmatrix}
   a & b \\
   \alpha & \beta
   \end{bmatrix}
   \begin{bmatrix}
   0 & 1 \\
   1 & 0
   \end{bmatrix} = \begin{bmatrix}
   b & a \\
   \beta & \alpha
   \end{bmatrix}.
   \]

   For the \(3 \times 2\) case there are \(3 \cdot 2/2 = 3\) such matrices (swap a row with the other 2, then swap the remaining two rows).

6. Gaussian elimination (GE) operations \(G_k\): There are three types of elementary row operations, which may be performed without fundamentally altering a system of equations (e.g. the resulting system of equations is equivalent). These operations are (1) swap rows (e.g. using a permutation matrix), (2) scale rows, or (3) perform addition/subtraction of two scaled rows. All such operations can be performed using matrices.

   For lack of a better term, we’ll describe these as ‘Gaussian elimination’ or ‘GE’ matrices.4 We will categorize any matrix that performs only elementary row operations (but any number of them) as a ‘GE’ matrix. Therefore, a cascade of GE matrices is also a GE matrix.

   Consider the GE matrix

   \[
   G = \begin{bmatrix}
   1 & 0 \\
   1 & -1
   \end{bmatrix}.
   \]

---

4 The term ‘elementary matrix’ may also be used to refer to a matrix that performs an elementary row operation. Typically, each elementary matrix differs from the identity matrix by a single row operation. A cascade of elementary matrices could be used to perform Gaussian elimination.
(a) This pre-multiplication scales and subtracts row (1) from (2) and returns it to row (2).

\[
GA = \begin{bmatrix} 1 & 0 \\ 1 & -1 \end{bmatrix} \begin{bmatrix} a & b \\ \alpha & \beta \end{bmatrix} = \begin{bmatrix} a & b \\ a - \alpha & b - \beta \end{bmatrix}.
\]

The shorthand for this Gaussian elimination operation is \((1) \leftarrow (1)\) and \((2) \leftarrow (1) - (2)\).

(b) Post-multiplication adds and scales columns.

\[
AG = \begin{bmatrix} a & b \\ \alpha & \beta \end{bmatrix} \begin{bmatrix} 1 & 0 \\ -1 & 1 \end{bmatrix} = \begin{bmatrix} a - b & b \\ \alpha - \beta & \beta \end{bmatrix}.
\]

Here the second column is subtracted from the first, and placed in the first. The second column is untouched. This operation is not a Gaussian elimination. Therefore, to put Gaussian elimination operations in matrix form, we form a cascade of pre-multiply matrices.

Here \(\det(G) = 1\), \(G^2 = I\), which won’t always be true if we scale by a number greater than 1. For instance, if \(G = \begin{bmatrix} 1 & 0 \\ m & 1 \end{bmatrix}\) (scale and add), then we have \(\det(G) = 1\), \(G^n = \begin{bmatrix} 1 & 0 \\ n \cdot m & 1 \end{bmatrix}\).

Exercise #1
Find the solution to the following \(3 \times 3\) matrix equation \(Ax = b\) by Gaussian elimination. Show your intermediate steps. You can check your work at each step using Matlab.

\[
\begin{bmatrix} 1 & 1 & -1 \\ 3 & 1 & 1 \\ 1 & -1 & 4 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \\ x_3 \end{bmatrix} = \begin{bmatrix} 1 \\ 9 \\ 8 \end{bmatrix}.
\]

1. Show (i.e., verify) that the first GE matrix \(G_1\), which zeros out all entries in the first column, is given by

\[
G_1 = \begin{bmatrix} 1 & 0 & 0 \\ -3 & 1 & 0 \\ -1 & 0 & 1 \end{bmatrix}.
\]

Identify the elementary row operations that this matrix performs.

**Solution:** Operate with GE matrix on \(A\)

\[
G_1[A|b] = \begin{bmatrix} 1 & 0 & 0 \\ -3 & 1 & 0 \\ -1 & 0 & 1 \end{bmatrix} \begin{bmatrix} 1 & 1 & -1 \\ 1 & 1 & 1 \\ 3 & 1 & 1 \end{bmatrix} = \begin{bmatrix} 1 & 1 & -1 \\ 0 & -2 & 4 \\ 0 & -2 & 5 \end{bmatrix}.
\]

The second row of \(G_1\) scales the first row by -3 and adds it to the second row

\((2) \leftarrow -3(1) + (2)\).

The third row of \(G_1\) scales the first row by -1 and adds it to the third row \([(3) \leftarrow -(1) + (3)\].

2. Find a second GE matrix, \(G_2\), to put \(G_1A\) in upper triangular form. Identify the elementary row operations that this matrix performs. **Solution:**

\[
G_2 = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & -1 & 1 \end{bmatrix},
\]

or \([(2) \leftarrow -(2) + (3)\]. Thus we have

\[
G_2G_1[A|b] = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & -1 & 1 \end{bmatrix} \begin{bmatrix} 1 & 0 & 0 \\ -3 & 1 & 0 \\ -1 & 0 & 1 \end{bmatrix} = \begin{bmatrix} 1 & 1 & -1 \\ 0 & -2 & 4 \\ 0 & 0 & 1 \end{bmatrix}.
\]

\[
\begin{bmatrix} 1 \\ 1 \\ 1 \end{bmatrix}.
\]

\[\square\]
3. Find a third GE matrix, \( G_3 \), which scales each row so that its leading term is 1. Identify the elementary row operations that this matrix performs. **Solution:**

\[
G_3 = \begin{bmatrix} 1 & 0 & 0 \\ 0 & -1/2 & 0 \\ 0 & 0 & 1 \end{bmatrix},
\]

which scales the second row by -1/2. Thus we have

\[
G_3G_2G_1[A|b] = \begin{bmatrix} 1 & 0 & 0 \\ 0 & -1/2 & 0 \\ 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} 1 & 1 & -1 & 1 \\ 0 & 0 & 1 & 1 \end{bmatrix} = \begin{bmatrix} 1 & 1 & -1 & 1 \\ 0 & 1 & -2 & -3 \end{bmatrix}.
\]

\[
\]

4. Finally, find the last GE matrix, \( G_4 \), that subtracts a scaled version of row 3 from row 2, and scaled versions of rows 2 and 3 from row 1, such that you are left with the identity matrix \((G_4G_3G_2G_1A = I)\). **Solution:**

\[
G_4 = \begin{bmatrix} 1 & 1 & -1 \\ 0 & 1 & 2 \\ 0 & 0 & 1 \end{bmatrix}.
\]

Thus we have

\[
G_4G_3G_2G_1[A|b] = \begin{bmatrix} 1 & 1 & -1 \\ 0 & 1 & 2 \\ 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} 1 & 1 & -1 & 1 \\ 0 & 0 & 1 & 1 \end{bmatrix} = \begin{bmatrix} 1 & 0 & 0 & 3 \\ 0 & 1 & 0 & -1 \end{bmatrix}.
\]

\[
\]

5. Solve for \([x_1, x_2, x_3]^T\) using the augmented matrix format \(G_4G_3G_2G_1[A|b]\) where \([A|b]\) is the augmented matrix. Note that if you’ve performed the preceding steps correctly, \(x = G_4G_3G_2G_1b\). **Solution:** From the preceding problems, we see that \([x_1, x_2, x_3]^T = [3, -1, 1]^T\).

---

**Inverse of the 2 × 2 matrix**

We shall now apply Gaussian elimination to find the solution \([x_1, x_2]\) for the 2 × 2 matrix equation \(Ax = y\) (Eq. 3.5.2.8, left). We assume to know \([a, b, c, d]\) and \([y_1, y_2]\). We wish to show that the intersection (solution) is given by the equation on the right.

Here we wish to prove that the left equation (i) has an inverse given by the right equation (ii):

\[
\begin{bmatrix} a & b \\ c & d \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \begin{bmatrix} y_1 \\ y_2 \end{bmatrix} \quad (i);
\begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \frac{1}{\Delta} \begin{bmatrix} d & -b \\ -c & a \end{bmatrix} \begin{bmatrix} y_1 \\ y_2 \end{bmatrix} \quad (ii).
\]

To take the inverse we:

1. swap the diagonal,
2. change the off-diagonal signs, and
3. normalize by the determinant \(\Delta\).

**Derivation of the inverse of a 2 × 2 matrix**

1. Step 1: To derive (ii) starting from (i), normalize the first column to 1.

\[
\begin{bmatrix} 1 & \frac{b}{c} \\ \frac{a}{c} & 1 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \begin{bmatrix} \frac{1}{\delta} & 0 \\ 0 & \frac{1}{\delta} \end{bmatrix} \begin{bmatrix} y_1 \\ y_2 \end{bmatrix}.
\]

2. Step 2: Subtract row (1) from row (2): \((2) \rightarrow (2) - (1)\)

\[
\begin{bmatrix} 1 & \frac{b}{c} \\ 0 & \frac{a}{c} - \frac{b}{c} \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \begin{bmatrix} \frac{1}{\delta} & 0 \\ -\frac{1}{\delta} & \frac{1}{\delta} \end{bmatrix} \begin{bmatrix} y_1 \\ y_2 \end{bmatrix}.
\]

3. Step 3: Multiply row (2) by \(ca\) and express result in terms of the determinant \(\Delta = ad - bc\).

\[
\begin{bmatrix} 1 & \frac{b}{a} \\ 0 & \Delta \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \begin{bmatrix} \frac{1}{\delta} & 0 \\ -\frac{1}{\delta} & \frac{1}{\delta} \end{bmatrix} \begin{bmatrix} y_1 \\ y_2 \end{bmatrix}.
\]
4. Step 4: Solve row (2) for $x_2$: $x_2 = -\frac{c}{\Delta} y_1 + \frac{a}{\Delta} y_2$.

5. Step 5: Solve row (1) for $x_1$:

$$x_1 = \frac{1}{a} y_1 - \frac{b}{a} x_2 = \left[ \frac{1}{a} + \frac{b}{a} \frac{c}{\Delta} \right] y_1 - \frac{b}{a} \frac{a}{\Delta} y_2.$$  

Rewriting in matrix format, in terms of $\Delta = ad - bc$, gives:

$$\begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \left[ \frac{1}{a} + \frac{b}{\Delta} \frac{c}{\Delta} - \frac{b}{\Delta} \right] \begin{bmatrix} y_1 \\ y_2 \end{bmatrix} = \frac{1}{\Delta} \begin{bmatrix} \Delta + bc \\ -c \end{bmatrix} \begin{bmatrix} a \\ -a \end{bmatrix} \begin{bmatrix} x_1 \\ y_1 \end{bmatrix} = \frac{1}{\Delta} \begin{bmatrix} d \\ -b \end{bmatrix} \begin{bmatrix} y_1 \\ y_2 \end{bmatrix},$$  

since $d = (\Delta + bc)/a$.

**Summary:** This is a lot of messy algebra, so it is essential that you memorize the final result: (1) swap the diagonal, (2) change the off-diagonal signs, and (3) normalize by the determinant $\Delta$. 


Appendix B

Eigenanalysis

Eigenanalysis is ubiquitous in engineering applications. It is useful in solving differential and difference equations, data-science applications, numerical approximation and computing, and linear algebra applications. Typically one must take a course in linear algebra to become knowledgeable in the inner workings of this method. In this appendix we intend to provide sufficient basics to allow one to read the text.

B.1 The eigenvalue matrix ($\Lambda$)

Given $2 \times 2$ matrix $A$, the related matrix eigen-equation is

$$AE = E\Lambda. \quad (B.1.0.1)$$

Pre-multiplying by $E^{-1}$ diagonalizes $A$, resulting in the eigenvalue matrix

$$\Lambda = E^{-1}AE = \begin{bmatrix} \lambda_1 & 0 \\ 0 & \lambda_2 \end{bmatrix}. \quad (B.1.0.2)$$

Post-multiplying by $E^{-1}$ recovers $A$

$$A = E\Lambda E^{-1} = \begin{bmatrix} a_{11} & a_{12} \\ a_{21} & a_{22} \end{bmatrix}. \quad (B.1.0.4)$$

Matrix product formula:

This last relation is the entire point of the eigenvector analysis, since it shows that any power of $A$ may be computed from powers of the eigenvalues. Specifically,

$$A^n = E\Lambda^n E^{-1}. \quad (B.1.0.5)$$

For example, $A^2 = AA = E\Lambda (E^{-1}E) \Lambda E^{-1} = E\Lambda^2 E^{-1}$.

Equations B.1.0.1, B.1.0.3 and B.1.0.4 are the key to eigenvector analysis, and you need to memorize them. You will use them repeatedly throughout this course.

Showing that $A - \lambda_\pm I_2$ is singular:

If we restrict Eq. B.1.0.1 to a single eigenvector (one of $e_\pm$), along with the corresponding eigenvalue $\lambda_\pm$, we obtain a matrix equations

$$Ae_\pm = e_\pm \lambda_\pm = \lambda_\pm e_\pm. \quad (B.1.0.6)$$

Note the swap in the order of $E_\pm$ and $\lambda_\pm$. Since $\lambda_\pm$ is a scalar, this is legal (and critically important), since this allows us to factor out $e_\pm$

$$(A - \lambda_\pm I_2)e_\pm = 0. \quad (B.1.0.6)$$

The matrix $A - \lambda_\pm I_2$ must be singular because when it operates on $e_\pm$, having nonzero norm, it must be zero.

It follows that its determinant (i.e., $|A - \lambda_\pm I_2| = 0$) must be zero. This equation uniquely determines the eigenvalues $\lambda_\pm$. 

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B.1.1 Calculating the eigenvalues $\lambda_{\pm}$

The eigenvalues $\lambda_{\pm}$ of $A$ may be determined from $|A - \lambda_{\pm}I_2| = 0$. As an example we let $A$ be Pell’s equation (Eq. 2.5.6.9, p. 52). In this case the eigenvalues may be found from

$$\begin{vmatrix} 1 - \lambda_{\pm} & N \\ 1 & 1 - \lambda_{\pm} \end{vmatrix} = (1 - \lambda_{\pm})^2 - N = 0,$$

thus $\lambda_{\pm} = (1 \mp \sqrt{N})$.  

B.1.2 Calculating the eigenvectors $e_{\pm}$

Once the eigenvalues have been determined, they are substituted into Eq. B.1.0.6, which determines the eigenvectors $E = [e_+, e_-]$, by solving

$$(A - \lambda_{\pm})e_{\pm} = \begin{bmatrix} 1 - \lambda_{\pm} & 2 \\ 1 & 1 - \lambda_{\pm} \end{bmatrix} e_{\pm} = 0,$$

where $1 - \lambda_{\pm} = 1 - (1 \mp \sqrt{N}) = \pm \sqrt{N}$, thus the Pell equation eigenvalues are $\lambda_{\pm} = 1 \mp \sqrt{N}$.  

Recall that Eq. B.1.0.6 is singular because we are using an eigenvalue, and each eigenvector is pointing in a unique direction (this is why it is singular). You might expect that this equation has no solution. In some sense you would be correct. When we solve for $e_{\pm}$, the two equations defined by Eq. B.1.0.6 are co-linear (the two equations describe parallel lines so their wedge product is zero). This follows from the fact that there is only one eigenvector for each eigenvalue.

Since there is only one eigenvalue we are expecting trouble, yet we may proceeding to solve for $e_+ = [e_1^+, e_2^+]^T$ with eigenvalue $+\sqrt{N}$

$$\begin{bmatrix} \sqrt{N} & N \\ 1 & \sqrt{N} \end{bmatrix} \begin{bmatrix} e_1^+ \\ e_2^+ \end{bmatrix} = 0.$$ 

If we divide the top row by $\sqrt{N}$ the two rows are identical, since the matrix must be singular. Thus this matrix equation gives two identical equations. This is the price of an over-specified equation (the singular matrix is degenerate).

We can determine each eigenvectors direction, but not their magnitudes. Following the same procedure for $\lambda_- = -\sqrt{N}$, the equation for $e_-$ is

$$\begin{bmatrix} -\sqrt{N} & N \\ 1 & -\sqrt{N} \end{bmatrix} \begin{bmatrix} e_1^- \\ e_2^- \end{bmatrix} = 0.$$ 

As before, this matrix is singular. Here $e_1^- - \sqrt{Ne_2^-} = 0$, thus the eigenvector is $e^- = c[\sqrt{N}, 1]^T$ where $c$ is a normalization constant.

Thus the unnormalized eigenmatrix is

$$E = \begin{bmatrix} e_1^+ & e_2^- \\ e_2^+ & e_2^- \end{bmatrix} = \begin{bmatrix} \sqrt{N} & -\sqrt{N} \\ 1 & 1 \end{bmatrix}.$$ 

Normalization of the eigenvectors:

The constant $c$ may be determined by normalizing the eigenvectors to have unit length. Since we cannot determine the length, we set it to 1. In some sense the degeneracy is resolved by this normalization,

$$\left(\pm \sqrt{N}\right)^2 + 1^2 = N + 1 = 1/c^2.$$ 

Thus the normalization factor to force each eigen vector to have length 1 is $c = 1/\sqrt{N + 1}$.

---

1It is a convention to order the eigenvalues from largest to smallest.
B.2 Pell equation solution example

Sec. 2.5.6 (p. 52) showed that the solution \([x_n, y_n]^T\) to Pell’s equation is given by powers of the Pell matrix \(A\).

For \(N = 2\), in Sec. 2.5.6 we found the explicit formula for \([x_n, y_n]^T\), based on powers of the Pell matrix

\[
A = 1_j \begin{bmatrix} 1 & 2 \\ 1 & 1 \end{bmatrix}.
\]

(B.2.0.1)

This recursive solution to Pell’s equation (Eq. 2.5.6.7) is Eq. 2.5.6.9 (p. 52). Thus we need powers of \(A\), that is \(A^n\), which gives an explicit expression for \([x_n, y_n]^T\). By the diagonalization of \(A\), its powers are simply the powers of its eigenvalues.

From Matlab/Octave with \(N = 2\) the eigenvalues of Eq. B.2.0.1 are

\[
\lambda_\pm \approx [2.4142j, -0.4142j].
\]

The solution for \(N = 3\) is shown on page 235.

Once the matrix has been diagonalized, one may compute powers of that matrix as powers of the eigenvalues. This results in the general solution given by

\[
\begin{bmatrix} x_n \\ y_n \end{bmatrix} = 1_j^n A^n \begin{bmatrix} 1 \\ 0 \end{bmatrix} = 1_j^n E \Lambda^n E^{-1} \begin{bmatrix} 1 \\ 0 \end{bmatrix}.
\]

The eigenvalue matrix \(D\) is diagonal with the eigenvalues sorted, largest first. The Matlab/Octave command \([E, D] = \text{eig}(A)\) is helpful to find \(D\) and \(E\) given any \(A\). As we saw above,

\[
\Lambda = 1_j \begin{bmatrix} 1 + \sqrt{2} & 0 \\ 0 & 1 - \sqrt{2} \end{bmatrix} \approx \begin{bmatrix} 2.414 & 0 \\ 0 & -0.414 \end{bmatrix}.
\]

B.2.1 Pell equation eigenvalue-eigenvector analysis

Here we show how to compute the eigenvalues and eigenvectors for the \(2 \times 2\) Pell matrix for \(N = 2\)

\[
A = \begin{bmatrix} 1 & 2 \\ 1 & 1 \end{bmatrix}.
\]

The Matlab/Octave command \([E, D] = \text{eig}(A)\) returns the eigenvector matrix \(E\)

\[
E = [e_+, e_-] = \frac{1}{\sqrt{3}} \begin{bmatrix} \sqrt{2} \\ 1 \end{bmatrix} = \begin{bmatrix} 0.8165 & -0.8165 \\ 0.5774 & 0.5774 \end{bmatrix}
\]

and the eigenvalue matrix \(\Lambda\) (Matlab/Octave’s \(D\))

\[
\Lambda \equiv \begin{bmatrix} \lambda_+ & 0 \\ 0 & \lambda_- \end{bmatrix} = \begin{bmatrix} 1 + \sqrt{2} & 0 \\ 0 & 1 - \sqrt{2} \end{bmatrix} = \begin{bmatrix} 2.4142 & 0 \\ 0 & -0.4142 \end{bmatrix}.
\]

The factor \(\sqrt{3}\) on \(E\) normalizes each eigenvector to 1 (i.e., Matlab/Octave’s command \(\text{norm}([\sqrt{2}, 1])\) gives \(\sqrt{3}\)).

In the following discussion we show how to determine \(E\) and \(D\) (i.e., \(\Lambda\)), given \(A\).

Pell’s equation for \(N = 3\)

In Table B.1, Pell’s equation for \(N = 3\) is given, with \(\beta_0 = \sqrt[3]{2}\). Perhaps try other trivial solutions such as \([-1, 0]^T\) and \([\pm j, 0]^T\), to provide clues to the proper value of \(\beta_0\) for cases where \(N > 3\).

Example: I suggest that you verify \(E \Lambda \neq \Lambda E\) and \(AE = EA\) with Matlab/Octave. Here is the Matlab/Octave program which does this:

```matlab
A = [1 2; 1 1]; %define the matrix
[E,D] = eig(A); %compute the eigenvector and eigenvalue matrices
A*E-E*D %This should be \$\approx 0\$, within numerical error.
E*D-D*E %This is not zero
```

My student Kehan found the general formula for \(\beta_0\).
Table B.1: Summary of the solution of Pell’s equation due to the Pythagoreans using matrix recursion, for the case of $N=3$. The integer solutions are shown on the right. Note that $x_n/y_n \to \sqrt{3}$, in agreement with the Euclidean algorithm. The Matlab/Octave program for generating this data is PellSol3.m. It seems likely that the powers of $\beta_0$ could be absorbed in the starting solution, and then be removed from the recursion.

Pell’s Equation for $N = 3$

Case of $N = 3$ & $[x_0, y_0]^T = [1, 0]^T$, $\beta_0 = j/\sqrt{2}$

Note: $x_n^2 - 3y_n^2 = 1$, $x_n/y_n \to \sqrt{3}$

<table>
<thead>
<tr>
<th>$x_1$</th>
<th>$\beta_0$</th>
<th>$[1, 1]$</th>
<th>$[1, 3, 1, 0]$</th>
<th>$(1/\beta_0)^2 - 3(1/\beta_0)^2 = 1$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$x_2$</td>
<td>$\beta_0^2$</td>
<td>$[4, 2]$</td>
<td>$[1, 3, 1, 1]$</td>
<td>$(4/\beta_0^2)^2 - 3(2/\beta_0)^2 = 1$</td>
</tr>
<tr>
<td>$x_3$</td>
<td>$\beta_0^3$</td>
<td>$[10, 6]$</td>
<td>$[1, 3, 4, 2]$</td>
<td>$(10/\beta_0^3)^2 - 3(6/\beta_0)^2 = 1$</td>
</tr>
<tr>
<td>$x_4$</td>
<td>$\beta_0^4$</td>
<td>$[28, 16]$</td>
<td>$[1, 3, 10, 6]$</td>
<td>$(28/\beta_0^4)^2 - 3(16/\beta_0)^2 = 1$</td>
</tr>
<tr>
<td>$x_5$</td>
<td>$\beta_0^5$</td>
<td>$[76, 44]$</td>
<td>$[1, 3, 28, 16]$</td>
<td>$(76/\beta_0^5)^2 - 3(44/\beta_0)^2 = 1$</td>
</tr>
</tbody>
</table>

Summary:

Thus far we have shown that for the case of Pell matrix with $N = 2$, the normalized eigenmatrix and its inverse is

$$E = [e_+, e_-] = \frac{1}{\sqrt{2}} \begin{bmatrix} \sqrt{2} & -\sqrt{2} \\ 1 & 1 \end{bmatrix}$$

and the eigenmatrix is

$$\Lambda = \begin{bmatrix} \lambda_+ & 0 \\ 0 & \lambda_- \end{bmatrix} = \begin{bmatrix} 1 + \sqrt{2} & 0 \\ 0 & 1 - \sqrt{2} \end{bmatrix}.$$

Note that when working with numeric data it is not necessary to normalize $E$. For example, the form of $e_1^\pm = [1 \pm \lambda^\pm, 1]^T$ is very simple, and easy to work with. Once normalize it becomes $(N = 2) [\sqrt{2}/\sqrt{3}, 1/\sqrt{3}]^T = [0.8165, 0.5773]^T$, obscuring its natural simplicity. The normalization buys little in terms of function.

Exercise #1

Verify that $\Lambda = E^{-1} AE$.

Solution: We shall work with the unnormalized eigenmatrix $eE$, where $e = \sqrt{2^2 + 1} = \sqrt{3}$. To compute the inverse of $eE$, 1) swap the diagonal values, 2) change the sign of the off diagonals, and 3) divide by the determinant $\Delta$:

$$(eE)^{-1} = \frac{1}{2e\sqrt{2}} \begin{bmatrix} 1 & \sqrt{2} \\ -\sqrt{2} & 1 \end{bmatrix} = \frac{1}{2e} \begin{bmatrix} 0.707 & 1 \\ -0.707 & 1 \end{bmatrix}.$$

We wish to show that $\Lambda = E^{-1} AE$

$$\frac{1}{2e} \begin{bmatrix} 0.707 & 1 \\ -0.707 & 1 \end{bmatrix} \begin{bmatrix} 1 & \sqrt{2} \\ -\sqrt{2} & 1 \end{bmatrix} \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix} = \begin{bmatrix} 1 + \sqrt{2} & 0 \\ 0 & 1 - \sqrt{2} \end{bmatrix}$$

which is best verified with Matlab. ♦

Exercise #2

Verify that $A = E\Lambda E^{-1}$.

Solution: We wish to show that

$$\begin{bmatrix} 1 & 2 \\ 1 & 1 \end{bmatrix} = \frac{1}{\sqrt{3}} \begin{bmatrix} \sqrt{2} & -\sqrt{2} \\ 1 & 1 \end{bmatrix} \begin{bmatrix} 1 + \sqrt{2} & 0 \\ 0 & 1 - \sqrt{2} \end{bmatrix} \frac{\sqrt{3}}{2\sqrt{2}} \begin{bmatrix} 1 & \sqrt{2} \\ -1 & 1 \end{bmatrix}.$$
All the above solutions have been verified with Octave. Eigenmatrix diagonalization is helpful in generating solutions for finding the solutions of Pell’s and Fibonacci’s equations using transmission matrices.

**Example:** If the matrix corresponds to a transmission line, the eigenvalues have units of seconds [s]

\[ V_+^n = \begin{bmatrix} e^{-sT_o} & 0 \\ 0 & e^{sT_o} \end{bmatrix} V_-^{n+1}. \]  

(B.2.1.2)

In the time domain the forward traveling wave \( v^+_{n+1}(t-(n+1)T_o) = v^+_n(t-nT_o) \) is delayed by \( T_o \). Two applications of the matrix delays the signal by \( 2T_o \).

**B.3 Symbolic analysis of \( \mathbf{T} \mathbf{E} = \mathbf{E} \Lambda \)**

**B.3.1 The \( 2 \times 2 \) transmission matrix**

Here we assume \( \mathbf{T} = \begin{bmatrix} A & B \\ C & D \end{bmatrix} \) with \( \Delta_T = 1 \).

The eigenvectors \( e_\pm \) of \( \mathbf{T} \) are

\[ e_\pm = \frac{1}{2\sqrt{\Delta_T}} \begin{bmatrix} (A - D) \pm \sqrt{(A - D)^2 + 4BC} \\ 1 \end{bmatrix} \]  

(B.3.1.1)

and eigenvalues are

\[ \lambda_\pm = \frac{1}{2} (A + D) \pm \sqrt{(A - D)^2 + 4BC}. \]  

(B.3.1.2)

The term under the radical (i.e., the discriminant) may be rewritten in terms of the determinant of \( \mathbf{T} \) (i.e., \( \Delta_T = AD - BC \)), since

\[ (A - D)^2 - (A + D)^2 = -4AD. \]

The for the ABCD matrix the expression under the radical becomes

\[ (A - D)^2 + 4BC = A^2 + D^2 - 4AD + 4BC = A^2 + D^2 - 4\Delta_T. \]

Rewriting the eigenvectors and eigenvalues in terms of \( \Delta_T = \pm 1 \), we find

\[ e_\pm = \frac{1}{2\sqrt{\Delta_T}} \begin{bmatrix} (A - D) \pm \sqrt{(A + D)^2 - 4\Delta_T} \\ 1 \end{bmatrix} \]  

(B.3.1.3)

and

\[ \lambda_\pm = \frac{1}{2} (A + D) \pm \sqrt{(A + D)^2 - 4\Delta_T}. \]  

(B.3.1.4)

**B.3.2 Matrices with symmetry**

For the case of the ABCD matrix the eigenvalues depend on reciprocity, since \( \Delta_T = 1 \) if \( \mathbf{T}(s) \) is reciprocal, and \( \Delta_T = -1 \) if it is anti-reciprocal. Thus it is helpful to display the eigenfunctions and values in terms of \( \Delta_T \) so this distinction is explicit.

**Reversible systems:**

When \( A = D \)

\[ \mathbf{E} = \begin{bmatrix} -\sqrt{\frac{A}{C}} & \sqrt{\frac{A}{C}} \\ \sqrt{\frac{A}{C}} & \sqrt{\frac{A}{C}} \end{bmatrix} \quad \Lambda = \begin{bmatrix} A - \sqrt{BC} & 0 \\ 0 & A + \sqrt{BC} \end{bmatrix} \]  

(B.3.2.5)

the transmission matrix is said to be reversible, and the properties greatly simplify.
**Reciprocal systems**

When the matrix is symmetric ($B = C$), the corresponding system is said to be *reciprocal*. Most physical systems are reciprocal. The determinant of the transmission matrix of a reciprocal network $\Delta_T = AD - BC = 1$. For example, electrical networks composed of inductors, capacitors and resistors are always reciprocal. It follows that the complex impedance matrix is symmetric (Van Valkenburg, 1964a).

Magnetic systems such as dynamic loudspeakers are anti-reciprocal, and correspondingly $\Delta_T = -1$. The impedance matrix of a loudspeaker is skew symmetric (Kim and Allen, 2013). All impedance matrices are either symmetric or anti-symmetric, depending on whether they are reciprocal (LRC networks) or anti-reciprocal (magnetic networks). These systems have complex eigenvalues with negative real parts, corresponding to lossy systems. In some sense, all of this follows from conservation of energy, but the precise general case is waiting for enlightenment. The impedance matrix is never Hermitian. It is easily proved that Hermitian matrices have real eigenvalues, which correspond to lossless networks. Any physical system of equations that has any type of loss cannot be Hermitian.

In summary, given a reciprocal system, the $T$ matrix has $\Delta_T = 1$, and the corresponding impedance matrix is symmetric (not Hermitian).

**B.3.3 Impedance matrix**

As previously discussed in Sec. 3.7 (p. 110), the $T$ matrix corresponding to an impedance matrix is

\[
\begin{bmatrix}
V_1 \\
V_2
\end{bmatrix} = Z(s) \begin{bmatrix}
I_1 \\
I_2
\end{bmatrix} = \frac{1}{C} \begin{bmatrix}
A & \Delta_T \\
1 & D
\end{bmatrix} \begin{bmatrix}
I_1 \\
I_2
\end{bmatrix}.
\]

Reciprocal systems have skew-symmetric impedance matrices, namely $z_{12} = z_{21}$ (i.e., $\Delta_T = 1$). This condition is best understood using the $T$ form of the impedance matrix, as shown in Fig. 3.9 (p. 112). When the system is both reversible $A = D$ and reciprocal, the impedance matrix simplifies to

\[
Z(s) = \frac{1}{C} \begin{bmatrix}
A & 1 \\
1 & A
\end{bmatrix}.
\]

For such systems there are only two degrees of freedom, $A$ and $C$. As discussed previously in Sec. 3.7 (p. 110), each of these has a physical meaning: $1/A$ is the Thévenin source voltage given a voltage drive and $B/A$ is the Thévenin impedance (Sec. 3.7.1.3, p. 111).

**Impedance is not Hermitian**: By definition, when a system is Hermitian its matrix is conjugate symmetric

\[
Z(s) = Z^\dagger(s),
\]

a stronger condition than reciprocal, but not the symmetry of the Brune impedance matrix. A reciprocal Brune impedance is symmetric (not Hermitian).
Appendix C

Laplace transforms $\mathcal{L}$

Laplace transforms are discussed in Sec. 3.9 (p. 119), with the definition of the $\mathcal{L}$ in Eq. 3.9.0.1 (p. 120). Level-I (basic) $\mathcal{L}$s are listed in Table 3.3 (p. 116).

C.1 Properties of the Laplace transform

The following is a summary description of the $\mathcal{L}$:

1. Time $t \in \mathbb{R}$ [s] and the Laplace frequency [rad] are defined as $s = \sigma + \omega j \in \mathbb{C}$.

2. Given a Laplace transform ($\mathcal{L}$) pair $f(t) \leftrightarrow F(s)$, in the engineering literature, the time domain is always lowercase $[f(t)]$ and causal [i.e., $f(t < 0) = 0$], and the frequency domain is uppercase $[F(s)]$. Maxwell’s venerable equations are the unfortunate exception to this otherwise universal rule.

3. The target time function $f(t < 0) = 0$ (i.e., it must be causal). The time limits are $0^- < t < \infty$. Thus the integral must start from slightly below $t = 0$ to integrate over a delta function at $t = 0$. For example, if $f(t) = \delta(t)$, the integral must include both sides of the impulse. If we want to include noncausal functions such as $\delta(t + 1)$, we must extend the lower time limit. In such cases we simply set the lower limit of the integral to $-\infty$ and let the integrand ($f(t)$) determine the limits.

4. When we take the forward transform ($t \rightarrow s$), the sign of the exponential is negative. This is necessary to ensure that the integral converges when the integrand $f(t) \rightarrow \infty$ as $t \rightarrow \infty$. For example, if $f(t) = e^t u(t)$ (i.e., without the negative $\sigma$ exponent), the integral does not converge.

5. The limits on the integrals of the reverse $\mathcal{L}$s are $[\sigma_o - \infty j, \sigma_o + \infty j] \in \mathbb{C}$. These limits are further discussed in Sec. 4.7.4 (p. 163).

6. When we take the inverse Laplace transform, the normalization factor of $1/2\pi j$ is required to cancel the $2\pi j$ in the differential $ds$ of the integral.

7. The frequencies for the $\mathcal{L}$ must be complex, and in general $F(s)$ is complex analytic for $\sigma > \sigma_o$. It follows that the real and imaginary parts of $F(s)$ are related by the Cauchy-Riemann conditions. Given $\Re\{F(s)\}$, it is possible to find $\Im\{F(s)\}$ (Boas, 1987). Read more on this in Sec. 4.2.2 (p. 136).

8. To take the inverse Laplace transform, we must learn how to integrate in the complex $s$ plane. This is explained on pages 151–163.

9. The Laplace Heaviside step function is defined as

$$u(t) = \int_{-\infty}^{t} \delta(t)dt = \begin{cases} 1 & \text{if } t > 0 \\ \text{NaN} & \text{if } t = 0 \\ 0 & \text{if } t < 0 \end{cases} .$$

Alternatively, we can define $\delta(t) = du(t)/dt$. 

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10. It is easily shown that \( u(t) \leftrightarrow 1/s \) by direct integration,

\[
F(s) = \int_0^\infty u(t) e^{-st} dt = \left. -\frac{e^{-st}}{s} \right|_0^\infty = \frac{1}{s}.
\]

With the \( \mathcal{L}T \) step \((u(t))\), there is no Gibbs ringing effect.

11. The Laplace transform of a Brune impedance takes the form of a ratio of two polynomials. In such cases, the roots of the numerator polynomial are called the zeros while the roots of the denominator polynomial are called the poles. For example, the \( \mathcal{L}T \) of \( u(t) \leftrightarrow 1/s \) has a pole at \( s = 0 \), which represents integration, since

\[
u(t) * f(t) = \int_{-\infty}^{\infty} f(\tau) d\tau \leftrightarrow \frac{F(s)}{s}.
\]

12. The \( \mathcal{L}T \) is quite different from the FT in terms of its analytic properties. For example, the step function \( u(t) \leftrightarrow 1/s \) is complex analytic everywhere except at \( s = 0 \). The FT of \( 1 \leftrightarrow 2\pi\delta(\omega) \) is not analytic anywhere.

13. The dilated step function \((a \in \mathbb{R})\) is

\[
u(at) \leftrightarrow \int_{-\infty}^{\infty} u(at)e^{-st} dt = \frac{1}{a} \int_{-\infty}^{\infty} u(\tau)e^{-s/a}\tau d\tau = \frac{a}{|a|} \frac{1}{s} = \pm \frac{1}{s},
\]

where we have made the change of variables \( \tau = at \). The only effect that \( a \) has on \( u(at) \) is the sign of \( t \), since \( u(t) = u(2t) \). However, \( u(-t) \neq u(t) \), since \( u(t) \cdot u(-t) = 0 \), and \( u(t) + u(-t) = 1 \), except at \( t = 0 \), where it is not defined.

Once complex integration in the complex plane has been defined (see Sec. 4.2.2, p. 136), we can justify the definition of the inverse \( \mathcal{L}^{-1}T \) (Eq. 3.9.0.1).\(^1\)

### C.2 Tables of Laplace transforms

The following tables of \( \mathcal{L}T \) and \( \mathcal{L}^{-1}T \) are a convenient summary of their properties and evaluations for many different functions. Table C.1 gives basic function properties such as convolution and the properties of step functions and frequency scaling. Table C.2 provides the commands for doing symbolic (computer algebra and calculus) transformations, which includes some unusual \( \mathcal{L}T \)s and Taylor series of the \( \Gamma(s) \) function (Graham et al., 1994), a complex analytic extension of the factorial. Table C.3 gives the basic transforms typically used for more common calculations. Table C.4 provides extended less common transforms, such as the half-derivative and integration and Bessel functions.

These tables are available in most books on differential equations and remain a core technology for analytic methods for solving differential equations.

^1https://en.wikipedia.org/wiki/Laplace_transform#Table_of_selected_Laplace_transforms
### Table C.1: Functional relationships among Laplace transforms.

<table>
<thead>
<tr>
<th>LT functional properties</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$f(t) \ast g(t) = \int_{t=0}^{t} f(t - \tau)g(\tau) d\tau \leftrightarrow F(s)G(s)$</td>
<td>convolution</td>
</tr>
<tr>
<td>$u(t) \ast f(t) = \int_{0-}^{t} f(\tau) d\tau \leftrightarrow \frac{F(s)}{s}$</td>
<td>convolution</td>
</tr>
<tr>
<td>$f(at)u(at) \leftrightarrow \frac{1}{a} F\left(\frac{s}{a}\right)$</td>
<td>scaling</td>
</tr>
<tr>
<td>$f(t)e^{-at}u(t) \leftrightarrow F(s + a)$</td>
<td>damped</td>
</tr>
<tr>
<td>$f(t - T)e^{-a(t-T)}u(t - T) \leftrightarrow e^{-sT}F(s + a)$</td>
<td>damped and delayed</td>
</tr>
<tr>
<td>$f(-t)u(-t) \leftrightarrow F(-s)$</td>
<td>reverse time</td>
</tr>
<tr>
<td>$f(-t)e^{-at}u(-t) \leftrightarrow F(a - s)$</td>
<td>time-reversed and damped</td>
</tr>
<tr>
<td>$\frac{d}{dt}f(t) = \delta'(t) \ast f(t) \leftrightarrow sF(s)$</td>
<td>deriv</td>
</tr>
<tr>
<td>$\frac{\sin(t)u(t)}{t} \leftrightarrow \tan^{-1}(1/s)$</td>
<td>half-sync</td>
</tr>
</tbody>
</table>

### Table C.2: Symbolic relationships among Laplace transforms. $K_3$ is a constant.

<table>
<thead>
<tr>
<th>syms</th>
<th>command</th>
<th>result</th>
</tr>
</thead>
<tbody>
<tr>
<td>syms t s p</td>
<td>laplace($t^{p-1}$)</td>
<td>$\frac{\Gamma(p)s^{-p}}{e^{-s}}$</td>
</tr>
<tr>
<td>syms s</td>
<td>ilaplace(gamma(s))</td>
<td>$e^{s/t}$</td>
</tr>
<tr>
<td>syms s t a</td>
<td>ilaplace(exp(-a*s)/(s,s,t))</td>
<td>Heaviside(t - a)</td>
</tr>
<tr>
<td>syms Gamma s t</td>
<td>taylor(Gamma(s,t))</td>
<td>$\frac{1}{s} - \gamma + s\left(\frac{\gamma^2}{2} + \frac{\pi^2}{12}\right) + \frac{1}{6} \text{polygamma}(2,1) - \frac{7\pi^4}{12^2} - \frac{\gamma^3}{6} + s^2 K_3 + \cdots$</td>
</tr>
</tbody>
</table>
Table C.3: Laplace transforms of \( f(t), \delta(t), u(t), \text{rect}(t), t_0, p, e \in \mathbb{R} \) and \( F(s), G(s), s, a \in \mathbb{C} \). Given a Laplace transform (LT) pair \( f(t) \leftrightarrow F(s) \), the frequency domain is always uppercase [e.g., \( F(s) \)] and the time domain lowercase [\( f(t) \)] and causal (i.e., \( f(t < 0) = 0 \)). An extended table of transforms is given in Table C.4.

<table>
<thead>
<tr>
<th>( f(t) \leftrightarrow F(s) ), ( t \in \mathbb{R}; s,F(s) \in \mathbb{C} )</th>
<th>Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \delta(t) \leftrightarrow 1 )</td>
<td>Dirac</td>
</tr>
<tr>
<td>( \delta(</td>
<td>a</td>
</tr>
<tr>
<td>( \delta(t-t_0) \leftrightarrow e^{-st_0} )</td>
<td>delayed Dirac</td>
</tr>
<tr>
<td>( \delta(t-t_0) \ast f(t) \leftrightarrow F(s)e^{-st_0} )</td>
<td>—</td>
</tr>
<tr>
<td>( \sum_{n=0}^{\infty} \delta(t-nt_0) = \frac{1}{1-\delta(t-t_0)} \leftrightarrow \frac{1}{1-e^{-st_0}} = \sum_{n=0}^{\infty} e^{-sn t_0} )</td>
<td>one-sided impulse train</td>
</tr>
<tr>
<td>( u(t) \leftrightarrow \frac{1}{s} )</td>
<td>Heaviside step</td>
</tr>
<tr>
<td>( u(-t) \leftrightarrow -\frac{1}{s} )</td>
<td>anticausal step</td>
</tr>
<tr>
<td>( u(at) \leftrightarrow \frac{a}{s} )</td>
<td>( a \neq 0 \in \mathbb{R} ) dilated or reversed step</td>
</tr>
<tr>
<td>( e^{at}u(-t) \leftrightarrow \frac{1}{-s+a} )</td>
<td>anticausal damped step</td>
</tr>
<tr>
<td>( e^{-at}u(t) \leftrightarrow \frac{1}{s+a} )</td>
<td>( a &gt; 0 \in \mathbb{R} ) damped step</td>
</tr>
<tr>
<td>( \cos(at)u(t) \leftrightarrow \frac{1}{2} \left( \frac{1}{s-a} + \frac{1}{s+a} \right) )</td>
<td>( a \in \mathbb{R} ) cos</td>
</tr>
<tr>
<td>( \sin(at)u(t) \leftrightarrow \frac{1}{2j} \left( \frac{1}{s-a} - \frac{1}{s+a} \right) )</td>
<td>( a \in \mathbb{C} ) damped sin</td>
</tr>
<tr>
<td>( u(t-t_0) \leftrightarrow \frac{1}{s} e^{-st_0} )</td>
<td>( t_0 &gt; 0 \in \mathbb{R} ) time delay</td>
</tr>
<tr>
<td>( \text{rect}(t) = \frac{1}{t_0} [u(t) - u(t-t_0)] \leftrightarrow \frac{1}{t_0} \left( 1 - e^{-st_0} \right) )</td>
<td>rect-pulse</td>
</tr>
<tr>
<td>( u(t) \ast u(t) = tu(t) \leftrightarrow 1/s^2 )</td>
<td>ramp</td>
</tr>
<tr>
<td>( u(t) \ast u(t) \ast u(t) = \frac{1}{2} t^2 u(t) \leftrightarrow 1/s^3 )</td>
<td>double ramp</td>
</tr>
<tr>
<td>( \frac{1}{\sqrt{t}} u(t) \leftrightarrow \sqrt{\frac{\pi}{s}} )</td>
<td>—</td>
</tr>
<tr>
<td>( t^p u(t) \leftrightarrow \frac{\Gamma(p+1)}{s^{p+1}} )</td>
<td>( \Re p &gt; -1, q \in \mathbb{C} )</td>
</tr>
</tbody>
</table>
### Table C.4: An extended table of Laplace transforms. $J_0$, $K_1$ are Bessel functions of the first and second kind.

<table>
<thead>
<tr>
<th>$f(t)$ ↔ $F(s)$</th>
<th>Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>$d^{1/2} \frac{d^{1/2}}{dt} f(t) u(t)$ ↔ $\sqrt{s} F(s)$</td>
<td>half derivative</td>
</tr>
<tr>
<td>$d^{1/2} \frac{d^{1/2}}{dt} u(t)$ ↔ $\sqrt{s}$</td>
<td>half derivative</td>
</tr>
<tr>
<td>$\frac{d}{dt} \frac{1}{\sqrt{\pi t}} u(t)$ ↔ $\frac{s}{\sqrt{s}} = \sqrt{s}$</td>
<td>semi-inductor</td>
</tr>
<tr>
<td>$\frac{1}{\sqrt{\pi t}} u(t)$ ↔ $\frac{1}{\sqrt{s}}$</td>
<td>half integration</td>
</tr>
<tr>
<td>$\text{erfc}(\alpha \sqrt{t})$ ↔ $\frac{1}{\sqrt{s}} e^{-2\alpha \sqrt{s}}$</td>
<td>(Morse and Feshbach (1953), p. 1582) $\alpha &gt; 0$; erfc</td>
</tr>
</tbody>
</table>

\[
J_0(at) u(t) \leftrightarrow \frac{1}{\sqrt{s^2 + a^2}} \quad \text{Bessel}
\]
\[
J_n(\omega_o t) u(t) \leftrightarrow \left( \frac{s^2 + \omega_o^2}{s^2 + a^2} \right)^n
\]
\[
J_1(t) u(t)/t \leftrightarrow \sqrt{s^2 + 1} - s
\]
\[
J_1(t) u(t)/t + 2u(t) \leftrightarrow \sqrt{s^2 + 1} + s = e^{s\text{inh}^{-1}(s)}
\]
\[
\delta(t) + J_1(t) u(t)/t \leftrightarrow \sqrt{s^2 + 1}
\]
\[
I_0(t) u(t) \leftrightarrow 1/\sqrt{s^2 - 1}
\]
\[
u(t)/\sqrt{t + 1} \leftrightarrow e^s \frac{\pi}{s} \text{erfc}(\sqrt{s})
\]
\[
\sqrt{t} u(t) * \sqrt{1 + \sqrt{t}} u(t) \leftrightarrow e^{s/2} K_1(s/2)/2s
\]
C.2.1 $LT^{-1}$ of the Riemann zeta function

The analytic properties of the zeta function have long been a goal of mathematicians, starting with Euler, and then several others, who may have made their reputation working on this important function. For the neophyte, $\zeta(s)$ is important because it is an analytic extension of the sieve, which is a method for identifying prime numbers.

Understood are the locations of the poles of zeta, which depend on the prime numbers. Not so well understood are the remaining analytic properties over the entire plane, such as the zeros of $\zeta(x)$, namely the poles of $1/\zeta(s)$. This section is a brief review of $\zeta(s)$ for beginners, building on the developments of analytic functions from Chapter C, especially Secs. C.2.4 and 2.1.1, p. 29.

Consider $z \equiv e^{sT}$, where $T$ is the sample period at which data are taken (every $T$ seconds). For example, if $T = 22.676 \times 10^{-6} = 1/44,000$ s, then the data is sampled at 44.10 kHz. This is how a CD player works with high-quality music. Thus the unit-time delay operator $z^{-1}$ is

$$\delta(t - T) \leftrightarrow e^{-sT}.$$  

However, when we deal with the Euler and Riemann zeta functions, the only sampling period that makes sense is $T = 1$ [s] or 1 [Hz] (i.e., $n \in \mathbb{Z}$). In this case, the samples of interest are mod$(n, \pi_k)$. Based on the sieve of Eratosthenes, Euler showed that the counting numbers may be uniquely reduced to multiples of the primes. This is the basis for the fundamental theorem of arithmetic, which states that every integer may be uniquely factored into a product of prime numbers. This is a fundamental idea and the source of the concept of the prime number.

The zeta function  The zeta function depends explicitly on the primes, which makes it a special function. In 1737 Euler proposed the real-valued function $\zeta(x) \in \mathbb{R}$ and $x \in \mathbb{R}$ to prove that the number of primes is infinite (Goldstein, 1973). Euler’s definition of $\zeta(x) \in \mathbb{R}$ is given by the power series,

$$\zeta(x) = \sum_{n=1}^{\infty} \frac{1}{n^x} \quad \text{for } x > 1 \in \mathbb{R}. \tag{C.2.1.1}$$

This series converges for $x > 0$, since $R = n^{-x} < 1$, $n > 1 \in \mathbb{N}$.\(^2\)

In 1860 Riemann extended the zeta function into the complex plane, resulting in $\zeta(s)$, defined by the complex analytic power series, identical to the Euler formula except $x \in \mathbb{R}$ has been replaced by $s \in \mathbb{C}$;

$$\zeta(s) \equiv \frac{1}{1^s} + \frac{1}{2^s} + \frac{1}{3^s} + \frac{1}{4^s} + \cdots = \sum_{n=1}^{\infty} \frac{1}{n^s} = \sum_{n=1}^{\infty} n^{-s} \quad \text{for } \Re{s} = \sigma > 1. \tag{C.2.1.2}$$

This formula converges for $\Re{s} > 1$ (Goldstein, 1973). To determine the formula in other regions of the $s$ plane, we need to extend the series via analytic continuation. As it turns out, Euler’s formulation provided detailed information about the structure of primes, going far beyond his original goal.

Euler product formula

As first published by Euler in 1737, we can recursively factor out the leading prime term, which results in Euler’s product formula. Euler’s procedure is an algebraic implementation of the sieve of Eratosthenes (Fig. 2.3, p. 43).

Multiplying $\zeta(s)$ by the factor $1/2^s$ and subtracting from $\zeta(s)$ remove all the powers of $2$: $1/2^0 + 1/2^2 + 1/2^{2s} + 1/2^{3s} + \cdots$

$$\left(1 - \frac{1}{2^s}\right) \zeta(s) = 1 + \frac{1}{2^s} + \frac{1}{3^s} + \frac{1}{4^s} + \frac{1}{5^s} + \cdots - \left(\frac{1}{2^s} + \frac{1}{4^s} + \frac{1}{8^s} + \frac{1}{16^s} + \cdots\right), \tag{C.2.1.3}$$

which results in

$$\zeta_1(s) = (1 - 2^{-s}) \zeta(s) = 1 + \frac{1}{3^s} + \frac{1}{5^s} + \frac{1}{7^s} + \frac{1}{9^s} + \frac{1}{11^s} + \frac{1}{13^s} + \cdots. \tag{C.2.1.4}$$

Repeating this with a lead factor $1/3^s$ applied to Eq. C.2.1.4 gives\(^3\)

$$\frac{1}{3^s} \left(1 - 2^{-s}\right) \zeta(s) = \frac{1}{3^s} + \frac{1}{9^s} + \frac{1}{15^s} + \frac{1}{21^s} + \frac{1}{27^s} + \frac{1}{33^s} + \cdots. \tag{C.2.1.5}$$

---

\(^2\)Sanity check: For example, let $n = 2$ and $x > 0$. Then $R = 2^{-x} < 1$, where $\epsilon \equiv \lim \epsilon x \to 0^+$. Taking the log gives $\log_2 R = -\epsilon \log_2 2 = -\epsilon < 0$. Since $\log_2 R < 0$, $R < 1$.

\(^3\)This is known as Euler’s sieve, as distinguished from the Eratosthenes sieve.
Subtracting Eq. C.2.1.5 from Eq. C.2.1.4 cancels the terms on the right-hand side of Eq. C.2.1.4, giving

\[ \zeta_2(s) = (1 - 3^{-s}) (1 - 2^{-s}) \zeta(s) = 1 + \frac{1}{5^s} + \frac{1}{7^s} + \frac{1}{11^s} + \frac{1}{13^s} + \frac{1}{17^s} + \frac{1}{19^s} + \cdots. \]

If we express this in terms of the primes \( \pi_k \), we can better visualize the structure:

\[ \zeta_2(s) = (1 - \pi_2^{-s}) (1 - \pi_4^{-s}) \zeta(s) = 1 + \frac{1}{\pi_3^s} + \frac{1}{\pi_5^s} + \frac{1}{\pi_6^s} + \frac{1}{\pi_7^s} + \frac{1}{\pi_8^s} + \cdots. \]

Thus \( \zeta_2 \) has removed primes \( \pi_1, \pi_2 \), leaving \( \pi_3 \) as the lead term in the series on the right-hand side.

This leads to a recursion in \( \zeta_k \):

\[ \zeta_k(s) = \zeta(s) \prod_{l=1}^{k} \zeta_l(s) = 1 + \sum_{l=k+1}^{\infty} \pi_l^{-s}. \]

The series on the right-hand side converges rapidly to 1 as each prime is removed, because the RoC is becoming much larger with each recursion. Each recursive step in this construction ensures that the lead term, along with all of its multiplicative factors, is subtracted out, just like the cancellations with the sieve of Eratosthenes. It is instructive to compare each iteration with that of the sieve (see Fig. 2.3, p. 43).

Repeating this process with the remaining primes removes all the terms on the right-hand side but the first (leaving 1), which results in Euler’s analytic product formula \( (s = x \in \mathbb{R}) \), or Riemann’s complex analytic product formula \( (s \in \mathbb{C}) \):

\[ 1 = \zeta(s)(1 - 2^{-s}) \cdot (1 - 3^{-s}) \cdot (1 - 5^{-s}) \cdot (1 - 7^{-s}) \cdot \cdots \cdot (1 - \pi_k^{-s}) \cdot \cdots. \]

\[ = \zeta(s) \prod_{k=1}^{\infty} (1 - \pi_k^{-s}) \]

\[ \zeta(s) = \prod_{k} \mathcal{P}_k(s), \quad \Re\{s\} = \sigma > 0, \text{ verify!} \]

(C.2.1.6) (C.2.1.7)

where \( \mathcal{P}_k(s) = 1 - \pi_k^{-s} \) defines the poles of \( \zeta(s) \) for prime \( \pi_k \).

**Finding the RoC of the product formula:** It would be interesting to find the RoC for \( \mathcal{P}_k(s) \), and for rigor, this question demands further investigation. To find the RoC, we need to evaluate

\[ |\pi_k^{-s}| = |e^{-sT_k}| = |e^{-\sigma T_k}| = \left( \frac{1}{\pi_k} \right)^{\sigma} < 1 \quad \text{for} \quad \sigma > 0, \quad \text{verify} \]

where \( T_k = \ln \pi_k \). For example,

\[ \frac{1}{\mathcal{P}_5(s)} = \frac{1}{1 - \left( \frac{1}{5} \right)^s} = 1 + \frac{1}{5^s} + \frac{1}{5^{2s}} + \frac{1}{5^{3s}} \cdots, \quad \Re\{s\} = \sigma > 0. \]

The RoC for each root is \( \sigma \geq 0 \) since when \( \sigma < 0 \).

Since \( 1/\pi_k < 1 \) for all \( k \in \mathbb{N} \), the Taylor series of \( \zeta_k(x) \) is entire except at its poles. Note that the RoC of a Taylor series in powers of \( \pi_k^{-s} \) increases with \( k \).

**Exercise #1**

Work out the RoC for \( k = 2 \).

**Solution:** The formula for the RoC is given above, which for \( \pi_2 = 3 \) is

| \pi_k^{-s} | = \begin{cases} \left( \frac{1}{3} \right)^{\sigma_r} < 1 & \text{for} \quad \sigma_r > 0, \\ \left( \frac{1}{3} \right)^{-\sigma_r} < 1 & \text{for} \quad \sigma_r < 0, \end{cases}

where \( \sigma_r \) is the boundary of the RoC.
Exercise #2
Show how to construct \( \zeta_2(t) \leftrightarrow \zeta_2(s) \) by working in the time domain.

Solution: The basic procedure for building a sieve is to sum the integers

\[
S_1 = \sum_{n=1}^{\infty} n 2^{n-1} = 1 \cdot 2^0 + 2 \cdot 2^1 + 3 \cdot 2^2 + \ldots,
\]
while the sieve for the \( k \)th prime \( \pi_k \) is

\[
S_k = \sum_{n=1}^{\infty} n \pi_k^{n-1} = 1 \cdot \pi_k^0 + \pi_k \cdot 2^1 + \pi_k \cdot 2^2 + \ldots.
\]

This sum may be written in terms of the convolution with the Heaviside step function \( u_k \), since

\[
u_k \star u_k = n u_k = 0 \cdot u_0 + 1 \cdot u_1 + 2 u_2 + \ldots + k u_k + \ldots.
\]

Poles of \( \zeta_k(s) \)

Riemann proposed that Euler’s zeta function \( \zeta(s) \in \mathbb{C} \) has a complex argument (first explored by Chebyshev in 1850 (Bombieri, 2000)) that extends \( \zeta(s) \) into the complex plane \( (s \in \mathbb{C}) \), thus making it a complex analytic function. Thus we might presume that \( \zeta(s) \) has an inverse Laplace transform. There seems to be very little written on this topic (Hill, 2007). We explore this question further here.

We can now identify the poles of \( \zeta_k(s) \) \((p \in \mathbb{N})\), which are required to determine the RoC. For example, the \( k \)th factor of Eq. C.2.1.7 expressed as an exponential, is

\[
\zeta_k(s) \equiv \frac{1}{1 - \pi_k^{-s}} = \frac{1}{1 - e^{-sT_k}} = \sum_{k=0}^{\infty} e^{-sT_k},
\]
where \( T_k \equiv \ln \pi_k \). Thus \( \zeta_p(s) \) has poles at \(-snT_p = 2\pi n j \) (when \( e^{-sT_p} = 1 \)), and

\[
\omega_n = \frac{2\pi n}{T_k},
\]
with \(-\infty < n \in \mathbb{Z} < \infty\). These poles are the eigenmodes of the zeta function. Fig. C.1 is a domain-colorized plot of this function. It is clear that the RoC of \( \zeta_k \) is \( > 0 \). It would be helpful to determine why \( \zeta(s) \) has a more restrictive RoC than each of its factors.
In the system equation, we see that the transfer function between the state variable $q(t)$ and the input $x(t)$ is given by $\zeta_p(t)$. Taking the Laplace transform, we see that $\zeta(s)$ is an all-pole function,

$$\zeta_p(s) = e^{-sT_p} \zeta_p(s) + \delta(t) \quad \text{(C.2.1.11)}$$

In terms of the physics, these transmission line equations are telling us that $\zeta(s)$ may be decomposed into an infinite cascade of transmission lines (Eq. C.2.1.10), each having a unique delay given by $T_k = \ln \pi_k, \pi_k \in \mathbb{P}$, the log of the primes. The input admittance of this cascade may be interpreted as an analytic continuation of $\zeta(s)$ that defines the eigenmodes of that cascaded impedance function.

Working in the time domain provides a key insight, as it allows us to determine the analytic continuation of the infinity of possible continuations, which are not obvious in the frequency domain. Transforming to the time domain is a form of analytic continuation of $\zeta(s)$ that depends on the assumption that $\zeta(t)$ is one-sided in time (causal).

Additional relationships: We need to know some important relationships provided by both Euler and Riemann (1859) when we study $\zeta(s)$.

With the goal of generalizing his result, Euler extended the definition with the functional equation

$$\zeta(s) = 2^s \pi^{s-1} \sin \left( \frac{\pi s}{2} \right) \Gamma(1-s) \zeta(1-s). \quad \text{(C.2.1.12)}$$

This seems closely related to Riemann’s time reversal symmetry properties (Bombieri, 2000),

$$\pi^{-s/2} \Gamma \left( \frac{s}{2} \right) \zeta(s) = \pi^{-(1-s)/2} \Gamma \left( \frac{1-s}{2} \right) \zeta(1-s).$$

---

4Here we use a shorthand double-parentheses notation $f(t)_T \equiv \sum_{k=0}^{\infty} f(t - kT)$ to define the one-sided infinite sum.
This equation is of the form $F\left(\frac{1}{2}\right)\zeta(s) = F\left(\frac{1-s}{2}\right)\zeta(1-s)$, where $F(s) = \Gamma(s) / \pi^s$.

As shown in Table C.1, the $\mathcal{L}^{-1}$ of $f(-t) \leftrightarrow F(-s)$ represents time-reversal. This leads to causal and anticausal functions that are symmetric about $\Re\{s\} = 1/2$ (Riemann, 1859) leading to an interpretation of Euler’s functional equation.

Riemann (1859, page 2) provides an alternative integral definition of $\zeta(s)$, based on the complex contour integration,

$$2 \sin(\pi s) \Gamma(s-1) \zeta(s) = \int_{x=-\infty}^{\infty} \int_{y=-\infty}^{\infty} \frac{(-x)^{s-1}}{e^x - 1} \frac{(yj)^{-1}}{e^{-yj} - 1} dx dy.$$

Given $\zeta_k(s)$, it seems important to look at the inverse $\mathcal{L}T$ of $\zeta_k(1-s)$ to gain insight into the analytically extended $\zeta(s)$.

**Integral definition of the complex Gamma function $\Gamma(s)$:** The definition of the complex analytic gamma function (see Table C.2) is

$$\Gamma(s+1) = s\Gamma(s) \equiv \int_0^\infty \xi^s e^\xi d\xi,$$

which is a generalization of the real integer factorial function $n!$,

$$\xi(t) = \int_{-\infty}^{\infty} \Gamma(s+1) e^{\xi ds} \frac{ds}{2\pi j}.$$

**What is the ROC of $\zeta(s)$?** It is commonly stated that Euler’s and thus Riemann’s product formulas are valid only for $\Re s > 1$; however, this does not seem to be actually proved (I could be missing this proof). Here I argue that the product formula is entire except at the poles—namely, that the formula is valid everywhere other than at the poles.

The argument goes as follows: Starting from the product formula (Eq. C.2.1.7, p. 245), we form the log-derivative and study the poles and residues:

$$D(s) \equiv \frac{d}{ds} \ln \Pi_k \frac{1}{1 - e^{-sT_k}}$$

$$= - \sum_{k=1}^{\infty} \frac{T_k e^{-sT_k}}{1 - e^{-sT_k}} \quad \leftrightarrow \quad \sum_{n=1}^{\infty} \sum_{k=1}^{\infty} \delta(t - nT_k).$$

Here $T_k = \ln \pi_k$, as previously defined, and $\leftrightarrow$ denotes the inverse Laplace transform, transforming $D(s) \leftrightarrow d(t)$ into the time domain. Note that $d(t)$ is a causal function, composed of an infinite number of delta functions (i.e., time delays), as shown in Fig. C.2 (p. 247).

**Zeros of $\zeta(s)$** We are still left with the most important question: Where are the zeros of $\zeta(s)$? Equation C.2.1.11 has no zeros; it is an all-pole system. The cascade of many such systems is also all-pole. As I see it, the issue is: What is the actual formula for $\zeta(s)$?

To answer this question, we need to study the properties of the reflectance function $\Gamma(s)$ given by Eq. 77. Frequency-domain transfer functions having unity magnitude on the $j\omega$ axis are called *all-pass filters* in the engineering literature. When the reflectance is loss-less, it is therefore all-pass since $|\Gamma(j\omega)| = 1$. An important property of all-pass filters is that they may be accurately approximated by pole-zero pairs straddling the $j\omega$ axis, with the poles to the left (as required by causality) and the zeros to the right. Given this placement, the phases of the poles and zeros add. The group delay gives the net delay of the all-pass filter, which is twice the delay of the poles alone. It would seem that this careful placement of the zeros exactly across from the poles provides the requirement that the zeros all line up parallel to the $j\omega$ axis, as deemed by the Riemann hypothesis. Could this be the resolution of this long-standing mystery? An alternative possibility is that the convergent product formula has zeros that are obscured by the lack of convergence of Eq. C.2.1.2.

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3 We can verify Riemann’s use of $x$, which is taken to be real rather than complex. This could be more natural (i.e., modern Laplace transformation notation) if $-x \to yj \to z$. 

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Appendix D

Visco-thermal losses

D.1 Adiabatic approximation at low frequencies

At very low frequencies, the adiabatic approximation must break down. At higher frequencies, viscous and thermal damping in air can become significant. In acoustics these two effects are typically ignored by assuming that wave propagation is irrotational and thus is described by the scalar wave equation. However, this is an approximation.

As first explained by Kirchhoff (1868), following Helmholtz (1858), these two loss mechanisms are related, but to understand why is somewhat complicated due to both the history and the mathematics. The full theory was first worked out by Kirchhoff (1868, 1974). Both forms of damping are caused by two different but coupled diffusion effects: first viscous effects, due to shear at the container walls, and second thermal effects, due to small deviations from adiabatic expansion (Kirchhoff, 1868, 1974). I believe Einstein was eventually involved as a result of his studies on Brownian motion (Einstein, 1905),

Newton’s early development understandably ignored viscous and thermal losses, which can be significant in a thin region called the boundary layer; at acoustic frequencies when the radius of the container (i.e., the horn) becomes smaller than the viscous boundary layer (e.g., much less than 1 mm), as first described by Helmholtz (1858, 1863b, 1978). Helmholtz’s analysis was soon extended to include thermal losses by Kirchhoff (1868, 1974), as succinctly summarized by Rayleigh (1896) and then experimentally verified by Mason (1927, 1928).

The mathematical nature of damping is that the propagation function $\kappa(s)$ (i.e., complex wave number) is extended to

$$\kappa(s) = \frac{s + \beta_o \sqrt{s}}{c_o} \quad [m^{-1}], \quad (D.1.0.1)$$

where the forwarded $P_-$ and backward $P_+$ pressure waves propagate as

$$P_{\pm}(s,x) = e^{-\kappa(s)x}, e^{\bar{\kappa}(s)x}, \quad (D.1.0.2)$$

with $\bar{\kappa}(s)$ the complex conjugate of $\kappa(s)$ so that $\Re\kappa(s) > 0$. The term $\beta_o \sqrt{s}$ affects both the real and imaginary parts of $\kappa(s)$. The real part is a frequency-dependent loss, and the imaginary part introduces a frequency-dependent speed of sound (Mason, 1928).

D.1.1 Lossy wave-guide propagation

In lossy wave propagation, the losses are due to viscous and thermal damping. The formulation of viscous loss in air transmission was first worked out by Helmholtz (1863a) and then extended by Kirchhoff (1868) to include thermal damping (Rayleigh, 1896, Vol. II, p. 319). These losses are explained by a modified complex propagation function $\kappa(s)$ (Eq. D.1.0.1). Following his review of these theories, Crandall (1926, Appendix A) noted that the “Helmholtz-Kirchhoff” theory had never been experimentally verified. Acting on this suggestion, Mason (1928) set out to verify their theory experimentally.

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1See Ch. 3 of https://www.ks.uiuc.edu/Services/Class/PHYS498/.
APPENDIX D. VISCO-THERMAL LOSSES

Figure D.1: This figure, taken from Mason (1928), compares the Helmholtz-Kirchhoff theory for $|\kappa(f)|$ to Mason’s 1928 experimental measurements of the loss. The ratio of two powers ($P_1$ and $P_2$) is plotted (see Mason’s discussion immediately below his Fig. 4), and as indicated in the label: “$10 \log_{10} P_1 / P_2$ for 1 [cm] of tube length.” This is a plot of the transmission power ratio in [dB/cm].

Mason’s specification of the propagation function

Mason’s results are reproduced in Fig. D.1 as solid lines for tubes of fixed radius between 3.7 and 8.5 [mm] that have a power reflectance given by

$$|\Gamma_L(f)|^2 = \left| e^{-\kappa(f)L} \right|^2 \text{[dB/cm]}.$$  \hspace{1cm} (D.1.1.3)

The complex propagation function cited by Rayleigh (1896) is (Mason, 1928, Eq. 2)

$$\kappa(\omega) = \frac{P\eta'\sqrt{\omega}}{2c_o S\sqrt{2\rho_o}} + \frac{i\omega}{c_o} \left\{1 + \frac{P\eta'}{2S\sqrt{2\omega\rho_o}} \right\}$$ \hspace{1cm} (D.1.1.4)

and the characteristic impedance is

$$z_o(\omega) = \sqrt{\frac{P_o\gamma \rho_o}{2 \left[1 + \frac{P\eta'}{2S\sqrt{2\omega\rho_o}} - J_2 \frac{P\eta'}{S\sqrt{2\omega\rho_o}} \right]},$$ \hspace{1cm} (D.1.1.5)

where $S$ is the tube area and $P$ is its perimeter. Mason specified the physical constants for air to be $\gamma_o = 1.4$ (ratio of specific heats), $\rho_o = 1.2$ kg/m$^3$ (density), $c_o = \sqrt{\gamma_o \rho_o / \rho_o} = 341.56$ m/s (lossless air velocity of sound at 23.5$^\circ$C), $P_o = 10^5$ Pa (atmospheric pressure), and $\mu_o = 18.6 \times 10^{-6}$ Pa-s (viscosity). Based on these values, $\eta'_o$ is defined as the composite thermodynamic constant (Mason, 1928)

$$\eta'_o = \sqrt{\mu_o} \left[1 + \sqrt{\frac{5}{2}} \left(\eta_o^{1/2} - \eta_o^{-1/2}\right)\right] = 18.6 \times 10^{-3} \left[1 + \sqrt{\frac{5}{2}} \left(\sqrt{1.4} - 1 / \sqrt{1.4}\right)\right] = 6.618 \times 10^{-3}.$$  

D.1.2 Impact of viscous and thermal losses

Assuming air at 23.5$^\circ$C, $c_o = \sqrt{\gamma_o P_o / \rho_o} \approx 344$ m/s is the speed of sound, $\eta_o = c_p/c_v = 1.4$ is the ratio of specific heats, $\mu_o = 18.5 \times 10^{-6}$ Pa-s is the viscosity, $\rho_o \approx 1.2$ kg/m$^2$ is the density, and $P_o = 10^5$ Pa (1 atm).

Equation D.1.1.4 and the measured data are compared in Fig. D.1, reproduced from Mason’s Fig. 4, which shows that the wave speed drops from 344 m/s at 2.6 kHz to 339 m/s at 0.4 kHz, a 1.5% reduction. At 1 kHz the loss is 1 dB/m for a 7.5-mm tube. Note that the loss and the speed of sound vary inversely with the radius. As the radius approaches the boundary layer thickness (i.e., the radial distance such that the loss is $e^{-1}$), the effect of the damping dominates the propagation.

With some significant algebra, Eq. D.1.1.4 may be greatly simplified to Eq. D.1.0.1. Numerically,

$$\beta_o = \frac{P}{2S \sqrt{\rho_o}} = \frac{P}{2S} 6.0415 \times 10^{-3}$$ \hspace{1cm} (3)

The real and imaginary parts of this expression, with $s = i\omega$, give Eq. D.1.0.1.
For the case of a cylindrical wave guide, the radius is $R = 2S/P$. Thus

$$\beta_o = \frac{1}{R} \frac{\eta_o'}{\sqrt{\rho_o}} = \frac{1}{R} 6.0415 \times 10^{-3}.$$  

Here $\beta_o = P\eta'/2S\sqrt{\rho_o}$ is a thermodynamic constant, $P$ is the perimeter of the tube, and $S$ the area (Mason, 1928).

For a cylindrical tube that has radius $R = 2S/P$, $\beta_o = \eta_o'/R\sqrt{\rho_o}$. To get a feeling for the magnitude of $\beta_o$, consider a 7.5-mm tube (i.e., the average diameter of the adult ear canal). Then $\eta' = 6.6180 \times 10^{-3}$ and $\beta_o = 1.6110$. Under these conditions, the wave-number cutoff frequency is $\beta_o/\sqrt{|s|} = 1.6011/\sqrt{2\pi}10^3 \approx 0.4131$ Hz. At 1 kHz the ratio of the loss to the propagation is $\beta_o/\sqrt{|s|} = 1.6011/\sqrt{2\pi}10^3 \approx 2\%$. At 100 Hz, this is a 6.4\% effect.$^4$

**Cut-off frequency** $s_o$: The frequency where the lossless part equals the lossy part is defined as $\kappa(s_o) = 0$—namely,

$$s_o + \beta_o\sqrt{s_o} = 0.$$  

Solving for $s_o$ gives the real, and negative, pole frequency of the system:

$$\sqrt{s_o} = -\beta_o = -6.0415 \times 10^{-3}/R.$$  

To get a feeling for the magnitude of $\beta_o$, let $R = 0.75/2$ [cm] (i.e., the average radius of the adult ear canal). Then

$$-\sqrt{s_o} = \beta_o = 6.0145 \times 10^{-3}/3.75 \times 10^{-3} \approx 1.6.$$  

We conclude that the losses are insignificant in the audio range, since for the human ear canal, $f_o = \beta_o^2/2\pi \approx 0.407$ Hz.$^5$

Note how the propagation function has a Helmholtz-Kirchhoff correction for both the real and imaginary parts. This means that both the speed of sound and the damping are dependent on frequency, proportional to $\beta_o\sqrt{s}/c_o$. Note also that the smaller the radius, the greater the damping.

**Summary:** The Helmholtz-Kirchhoff theory of viscous and thermal losses results in a frequency-dependent speed of sound that has a frequency dependence proportional to $1/\sqrt{\omega}$ (Mason, 1928, Eq. 4). This corresponds to a 2\% change in the sound velocity over the decade from 0.2 to 2 kHz (Mason, 1928, Fig. 5), in agreement with experiment.

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$^4$/home/jba/Mimosa/2C-FindLengths.16/doc.2–c_calib.14/m/MasonKappa.m  
$^5$/home/jba/Mimosa/2C-FindLengths.16/doc.2–c_calib.14/m/MasonKappa.m
Appendix E

Number theory applications

E.1 Division with rounding method

We want to show that the GCD for $m, n, k \in \mathbb{N}$ (Eq. 2.5.3.2, p. 45) may be written in matrix form as

$$
\begin{pmatrix}
m \\
n
\end{pmatrix}_{k+1} =
\begin{bmatrix}
0 & 1 \\
1 & -\lfloor \frac{m}{n} \rfloor
\end{bmatrix}
\begin{pmatrix}
m \\
n
\end{pmatrix}_k.
$$

(E.1.0.1)

Eq. E.1.0.1 implements the $gcd(m, n)$ for $m > n$.

This starts with $k = 0$, $m_0 = a$, and $n_0 = b$. With this method there is no need to test whether $n_k < m_n$, as it is built into the procedure. The method uses the floor function $\lfloor x \rfloor$, which finds the integer part of $x$ ($\lfloor x \rfloor$ rounds toward $-\infty$). After each step we will see that the value $n_{k+1} < m_{k+1}$. The method terminates when $n_{k+1} = 0$ with $gcd(a, b) = m_{k+1}$.

The following vectorized code is more efficient than the direct matrix method:

```matlab
function n=gcd2(a,b)
M=[abs(a);abs(b)]; %Save (a,b) in array M(2,1)

% done when M(1) = 0
while M(1) ~= 0
    disp(sprintf('M(1)=%g, M(2)=%g ',M(1),M(2)));
    M=[M(2)-M(1)*floor(M(2)/M(1)); M(1)]; %automatically sorted
end %done

n=M(2); %GCD is M(2)

With a minor extension in the test for “end,” this code can be made to work with irrational inputs (e.g., $(n\pi, m\pi)$).

This method calculates the number of times $n < m$ must subtract from $m$ using the floor function. This operation is the same as the mod function. Specifically,

$$
n_{k+1} = m_k - \lfloor \frac{m}{n} \rfloor n_k
$$

(E.1.0.2)

so that the output is the definition of the remainder of modular arithmetic. This would have been obvious to anyone using an abacus, which explains why it was discovered so early.

Note that the next value of $m = M(1)$ is always less than $n = M(2)$ and must remain greater than or equal to zero. This one-line vector operation is then repeated until the remainder $M(1)$ is 0. The gcd is then $n = M(2)$. When we use irrational numbers, the code still works except the error is never exactly zero due to IEEE 754 rounding. Thus the criterion must be that the error is within some small factor times the smallest number (which in Matlab/Octave is the number $\text{eps} = 2.220446049250313 \times 10^{-16}$, as defined in the IEEE 754 standard).

Thus, without factoring the two numbers, Eq. E.1.0.2 recursively finds the gcd. Perhaps this is best seen with some examples.

The GCD is an important and venerable method, useful in engineering and mathematics but, as best I know, not typically taught in the traditional engineering curriculum.

\footnote{\url{https://en.wikipedia.org/wiki/Modulo_operation}}
**GCD applied to polynomials:** An interesting generalization is to work with polynomials rather than numbers and apply the Euclidean algorithm.

The GCD may be generalized in several significant ways. For example, what is the GCD of two polynomials? To answer this question, we must factor the two polynomials to identify common roots.

### E.2 Derivation of the CFA matrix

We can define the continued fraction algorithm (CFA) starting from the basic definitions of the floor and remainder formulas. Starting with a decimal number \( x \), we split it into the decimal and remainder parts. If we start with \( n = 0 \) and \( x_0 = x \in \mathbb{I} \), the integer part is

\[
m_0 = \lfloor x \rfloor \in \mathbb{N}
\]

and the remainder is

\[
r_0 = x - m_0.
\]

Corresponding to the CFA, the next target \( x_1 \) for \( n = 1 \) is

\[
x_1 = r_0^{-1}
\]

and the integer part is \( m_1 = \lfloor x_1 \rfloor \). As in the case of \( n = 0 \), the integer part is

\[
m_1 = \lfloor x_1 \rfloor
\]

and the remainder is

\[
r_1 = x_1 - m_1.
\]

The recursion for \( n = 2 \) is similar.

For us to better appreciate what is happening, it is helpful to write these recursions in matrix format. Rewriting the case of \( n = 1 \) and using the remainder formula for the ratio of two numbers \( p \geq q \in \mathbb{N} \) with \( q \neq 0 \), we have

\[
\begin{bmatrix}
p \\
q
\end{bmatrix} =
\begin{bmatrix}
u_1 & 1 \\
1 & 0
\end{bmatrix}
\begin{bmatrix}
r_0 \\
r_1
\end{bmatrix}.
\]

From the remainder formula, \( u_1 = [p/q] \). Continuing with \( n = 2 \):

\[
\begin{bmatrix}
r_0 \\
r_1
\end{bmatrix} =
\begin{bmatrix}
u_2 & 1 \\
1 & 0
\end{bmatrix}
\begin{bmatrix}
r_1 \\
r_2
\end{bmatrix},
\]

where \( u_1 = [r_0/r_1] \). Continuing with \( n = 3 \):

\[
\begin{bmatrix}
r_1 \\
r_2
\end{bmatrix} =
\begin{bmatrix}
u_3 & 1 \\
1 & 0
\end{bmatrix}
\begin{bmatrix}
r_2 \\
r_3
\end{bmatrix},
\]

where \( u_2 = [r_1/r_2] \).

For arbitrary \( n \) we find

\[
\begin{bmatrix}
r_{n-2} \\
r_{n-1}
\end{bmatrix} =
\begin{bmatrix}
u_n & 1 \\
1 & 0
\end{bmatrix}
\begin{bmatrix}
r_{n-1} \\
r_n
\end{bmatrix}, \quad (E.2.0.1)
\]

where \( u_n = [r_{n-1}/r_n] \). This terminates when \( r_n = 0 \) in the above \( n \)th step:

\[
\begin{bmatrix}
r_{n-2} \\
r_{n-1}
\end{bmatrix} =
\begin{bmatrix}
u_n & 1 \\
1 & 0
\end{bmatrix}
\begin{bmatrix}
r_{n-1} \\
r_n = 0
\end{bmatrix}.
\]

**Example:** We let \( p = 355 \) and \( q = 113 \), which are coprime, and set \( n = 1 \). Then Eq. E.2.0.1 becomes

\[
\begin{bmatrix}
355 \\
113
\end{bmatrix} =
\begin{bmatrix}
3 & 1 \\
1 & 0
\end{bmatrix}
\begin{bmatrix}
r_0 \\
r_1
\end{bmatrix}.
\]

---

2The method presented here was developed by Yiming Zhang as a student project in 2019.
E.3. TAKING THE INVERSE TO GET THE GCD

since $u_1 = \left\lfloor \frac{355}{113} \right\rfloor = 3$. Solving for the RHS gives $[r_0; r_1] = [113; 16]$ ($355 = 113 \cdot 3 + 16$). To find $[r_0; r_1]$, we take the inverse:

$$\begin{bmatrix} r_0 \\ r_1 \end{bmatrix} = \begin{bmatrix} 0 & 1 \\ 1 & -3 \end{bmatrix} \begin{bmatrix} 355 \\ 113 \end{bmatrix}.$$  

For $n = 2$, with the RHS from the previous step,

$$\begin{bmatrix} 113 \\ 16 \end{bmatrix} = \begin{bmatrix} u_2 & 1 \\ 1 & 0 \end{bmatrix} \begin{bmatrix} r_1 \\ r_2 \end{bmatrix},$$

since $u_2 = \left\lfloor \frac{113}{16} \right\rfloor = 7$. Solving for the RHS gives $[r_1; r_2] = [16; 1]$ ($113 = 16 \cdot 7 + 1$). It seems we are done, but let’s go one step further.

For $n = 3$ we now have

$$\begin{bmatrix} 16 \\ 1 \end{bmatrix} = \begin{bmatrix} u_3 & 1 \\ 1 & 0 \end{bmatrix} \begin{bmatrix} r_1 \\ r_2 \end{bmatrix},$$

since $u_3 = \left\lfloor \frac{16}{1} \right\rfloor = 16$. Solving for the RHS gives $[r_1; r_2] = [1; 0]$. This confirms that we are done, since $r_2 = 0$.

**Derivation of Eq. E.2.0.1:** Equation Eq. E.2.0.1 is derived as follows: Starting from the target $x \in \mathbb{R}$, we define

$$p = \lfloor x \rfloor \quad \text{and} \quad q = \frac{1}{x - p} \in \mathbb{R}.$$  

These two relationships for truncation and remainder allow us to write the general matrix recursion relation for the CFA (Eq. E.2.0.1). Given $\{p, q\}$, we continue with the above cfa method.

One slight problem with the above is that the output is on the right and the input on the left. Thus we need to take the inverse of these relationships to turn this into a composition.

E.3 Taking the inverse to get the gcd

Variables $p$ and $q$ are the remainders $r_{n-1}$ and $r_n$, respectively. Using this notation with $n - 1$ gives Eq. E.2.0.1. Inverting this gives the formula for the GCD:

$$\begin{bmatrix} r_{n-1} \\ r_n \end{bmatrix} = \begin{bmatrix} 0 & \frac{1}{r_{n-2}} \\ 1 & -\frac{r_{n-2}}{r_{n-1}} \end{bmatrix} \begin{bmatrix} r_{n-2} \\ r_{n-1} \end{bmatrix}. $$

This terminates when $r_n = 0$ and the $\gcd(p, q)$ is $r_{n-1}$. Not surprisingly these equations mirror Eq. 2.5.3.3 (p. 46), but with a different indexing scheme and interpretation of the variables.

This then explains why Gauss called the CFA the *Euclidean algorithm*. He was not confused. But since the equations have an inverse relationship, they are not strictly the same.
Appendix F

Eleven postulates of systems of algebraic networks

Physical systems obey basic rules that follow from the physics. It is helpful to summarize these restrictions as postulates presented in terms of a taxonomy, or categorization method, of the fundamental properties of physical systems. Eleven of these are listed in this appendix from an article by Kim and Allen (2013).

F.1 Representative system

A taxonomy of physical systems comes from a systematic summary of the laws of physics, which includes at least the eleven basic network postulates, described in Sec. 3.9.

To describe the network postulates, it is helpful to start from a two-port matrix representation as discussed in Sec. 3.7 (p. 110).

![Figure F.1: The schematic representation of an algebraic network, defined by its two-port ABCD transmission, that has three elements described by the Hunt parameters (Hunt, 1952): $Z_e(s)$, the electrical impedance, $z_m(s)$, the mechanical impedance, and $T(s)$, the transduction coefficient matrix of an electromechanical transducer network. The port variables are $\Phi(f)$ and $I(f)$: the frequency domain voltage and current, and $F(f)$ and $U(f)$: the force and velocity (Hunt, 1952; Kim and Allen, 2013). This matrix factors the two-port model into three $2 \times 2$ matrices, separating the three physical elements as matrix algebra. It is necessary to apply a negative sign on the velocity $-U$ so that it has an outward flow, as required to match the next cell with its inward flow.

As shown in Fig. F.1, the two-port transmission matrix for an acoustic transducer (loudspeaker) is characterized by the equation

$$
\begin{bmatrix}
\Phi_i \\
I_i
\end{bmatrix}
= \begin{bmatrix}
A(s) & B(s) \\
C(s) & D(s)
\end{bmatrix}
\begin{bmatrix}
F_i \\
-U_i
\end{bmatrix}
= \frac{1}{T}
\begin{bmatrix}
z_m(s) & Z_e(s)z_m(s) + T^2 \\
Z_e(s) & Z_e(s)
\end{bmatrix}
\begin{bmatrix}
F_i \\
-U_i
\end{bmatrix},
$$

shown as a product of three $2 \times 2$ matrices in the figure, with each factor representing one of the three Hunt parameters of the loudspeaker.

This figure represents the electromechanical motor of the loudspeaker and consists of three elements: the electrical input impedance $Z_e(s)$, a gyrator, which is similar to a transformer that relates current to force, and the output mechanical impedance $z_m(s)$. This circuit describes what is needed to fully characterize its operation, from electrical input to mechanical (acoustical) output.

The input is electrical (voltage and current) $[\Phi_i, I_i]$ and the output (load) is the mechanical (force and velocity) $[F_i, U_i]$. The first matrix is the general case, expressed in terms of four unspecified functions $A(s)$, $B(s)$, $C(s)$, and $D(s)$, while the second matrix is for the specific example of Fig. F.1. The three entries are the electrical driving point impedance $Z_e(s)$, the mechanical impedance $z_m(s)$, and the transduction $T = B_m l$, where $B_m$ is the magnetic flux strength and $l$ is the length of the wire crossing the flux. Since the transmission matrix is antireciprocal, its determinant $\Delta_T = -1$, as is easily verified.

Other common examples of cross-modality transduction and current–thermal (thermoelectric effect) and force–voltage (piezoelectric effect). These systems are all reciprocal: thus the transduction has the same sign.
F.1.1 Impedance matrix

These eleven network postulates describe the properties of a system that has an input and an output. For an electromagnetic transducer (loudspeaker) the system is described by the two-port transmission matrix, as shown in Fig. F.1. The electrical input impedance of a loudspeaker is $Z_\text{e}(s)$, defined by

$$Z_\text{e}(s) = \frac{V(\omega)}{I(\omega)} |_{U=0}. $$

Note that this driving-point impedance must be causal since it is a function of $s$; thus it has a Laplace transform. The corresponding two-port impedance matrix for Fig. F.1 is

$$\begin{bmatrix} \Phi_1 \\ F_1 \end{bmatrix} = \begin{bmatrix} z_{11}(s) & z_{12}(s) \\ z_{21}(s) & z_{22}(s) \end{bmatrix} \begin{bmatrix} I_1 \\ U_1 \end{bmatrix} = \begin{bmatrix} Z_\text{e}(s) & -T(s) \\ T(s) & z_m(s) \end{bmatrix} \begin{bmatrix} I_1 \\ U_1 \end{bmatrix}. \tag{F.1.1.2}$$

Such a description allows us to define Thévenin parameters, a concept used widely in circuit analysis and network models from other modalities.

The impedance matrix is an alternative description of the system but with generalized forces $[\Phi, F]$ on the left and generalized flows $[I, U]$ on the right. A rearrangement of terms allows us to go from the ABCD to the impedance parameters (Van Valkenburg, 1964b). The electromagnetic transducer is antireciprocal (Postulate P6), $z_{12} = -z_{21} = T = B_0 l$.

F.2 Taxonomy of algebraic networks

The postulates are extended beyond those defined by Carlin and Giordano (Sec. 3.9, p. 119) when there is an interaction of waves and a structured medium, along with other properties not covered by classic network theory. Assuming quasistatics (QS), the wavelength must be large relative to the medium’s lattice constants. Thus the QS property must be extended to three dimensions and possibly to the cases of anisotropic and random media.

Causality: P1 As we stated, due to causality the negative properties (e.g., negative refractive index) must be limited in bandwidth as a result of the Cauchy–Riemann conditions. However, even causality needs to be extended to include the delay, as quantified by the d’Alembert solution to the wave equation, which means that the delay is proportional to the distance. Thus we generalize Postulate P1 to include the space-dependent delay. When we wish to discuss this property, we call it Einstein causality, which says that the delay must be proportional to the distance $x$, with impulse response $\delta(t - x/c)$.

Linearity: P2 The wave properties of a system may be nonlinear. This is not restrictive, as most physical systems are naturally nonlinear. For example, a capacitor is inherently nonlinear: As the charge builds up on the plates of the capacitor, a stress is applied to the intermediate dielectric due to the electrostatic force $F = qE$. In a similar manner, an inductor is nonlinear. Two wires carrying a current are attracted or repelled due to the force created by the flux. The net force is the product of the two fluxes due to each current.

In summary, most physical systems are naturally nonlinear; it’s simply a matter of degree. An important counterexample is an amplifier with negative feedback and a very large open-loop gain. There are, therefore, many types of nonlinearity, both instantaneous types and those with memory (e.g., hysteresis). Given the nature of Postulate P1, even an instantaneous nonlinearity may be ruled out. The linear model is so critical for our analysis, providing fundamental understanding, that we frequently take Postulates P1 and P2 for granted.

Passive/Active impedances: P3 This postulate is about conservation of energy and Otto Brune’s positive-real (PR, also called physically realizable) condition that every passive impedance must obey. Following on the work of his primary Ph.D. thesis advisor Wilhelm Cauer (1900–1945) and Ernst Guillemin, along with Norbert Weiner and Vannevar Bush at MIT, Brune mathematically characterized the properties of every PR one-port driving point impedance (Brune, 1931b).

When the input resistance of the impedance is real, the system is said to be passive, which means the system obeys conservation of energy. The real part of $Z(s)$ is positive if and only if the corresponding reflectance is less than 1 in magnitude. The reflectance of $Z(s)$ is defined as a bilinear transformation of the impedance, normalized by its surge resistance $r_o$ (Campbell, 1903):

$$\Gamma(s) = \frac{Z(s) - r_o}{Z(s) + r_o} = \frac{Z - 1}{Z + 1}.$$
where $\mathcal{Z} = Z/r_o$. The surge resistance is defined in terms of the inverse Laplace transform of $Z(s) \leftrightarrow z(t)$, which must have the form

$$z(t) = r_o \delta(t) + \rho(t),$$

where $\rho(t) = 0$ for $t < 0$. It naturally follows that $\gamma(t) \leftrightarrow \Gamma(s)$ is zero for negative and zero time—namely, $\gamma(0) = 0, t < 0$.

Given any linear PR impedance $Z(s) = R(\sigma, \omega) + jX(\sigma, \omega)$ that has real part $R(\sigma, \omega)$ and imaginary part $X(\sigma, \omega)$, the impedance is defined as being PR (Brune, 1931b) if and only if

$$\Re Z(s) = R(\sigma \geq 0, \omega) \geq 0. \quad (F.2.0.1)$$

That is, the real part of any PR impedance is nonnegative everywhere in the right half-plane ($\sigma \geq 0$). This is a very strong condition on the complex analytic function $Z(s)$ of a complex variable $s$. This condition is equivalent to any of the following statements (Van Valkenburg, 1964a):

1. There are no poles or zeros in the right half-plane ($Z(s)$ may have poles and zeros on the $\sigma = 0$ axis).
2. If $Z(s)$ is PR, then its reciprocal $Y(s) = 1/Z(s)$ is PR (the poles and zeros swap).
3. If the impedance can be written as the ratio of two polynomials (a limited case related to the quasistatics approximation, Postulate P9) that have degrees $N$ and $L$, then $|N - L| \leq 1$.
4. The angle of the impedance $\theta \equiv \angle Z$ lies within $[-\pi \leq \theta \leq \pi]$.
5. The impedance and its reciprocal are complex analytic in the right half-plane; thus each obeys the Cauchy–Riemann conditions there.

**Energy and power:** Since Postulate P3 requires the impedance PR condition, it ensures that every impedance is positive-definite (PD), thus guaranteeing that conservation of energy is obeyed (Schwinger and Saxon, 1968, p.17). This means that the total energy absorbed by any PR impedance must remain positive for all time. Mathematically we can state this as

$$E(t) = \int_{-\infty}^{t} v(t)i(t) dt = \int_{-\infty}^{t} i(t) * z(t) i(t) dt > 0,$$

where $i(t)$ is any current, $v(t) = z(t) * i(t)$ is the corresponding voltage, and $z(t)$ is the real causal impulse response of the impedance [e.g., $z(t) \leftrightarrow Z(s)$ are a Laplace transform pair]. In summary, if $Z(s)$ is PR, then $E(t)$ is PD.

As discussed in detail by Van Valkenburg, any rational PR impedance can be represented as a partial fraction expansion, which can be expanded into first-order poles as

$$Z(s) = K \prod_{n=1}^{L}(s - n_i) \prod_{k=1}^{N}(s - d_k) = \sum_{n} \frac{\rho_n}{s - s_n} e^{j(\theta_n - \theta_n)}, \quad (F.2.0.2)$$

where $\rho_n$ is a complex scale factor (residue). Every pole in a PR function has only simple poles and zeros, which requires that $|L - N| \leq 1$ (Van Valkenburg, 1964b).

Whereas the PD property clearly follows from Postulate P3 (conservation of energy), the physics is not so clear. Specifically, what is the physical meaning of the constraints on $Z(s)$? In many ways, the impedance concept is highly artificial, as expressed by Postulates P1–P7.

When the impedance is not rational, special care must be taken. An example of this is the semi-inductor $M\sqrt{s}$ and the semicapacitor $K/\sqrt{s}$ due, for example, to the skin effect in EM theory and viscous and thermal losses in acoustics, both of which are frequency-dependent boundary-layer diffusion losses (Vanderkooy, 1989). They remain positive-real but have a branch cut and thus are double-valued in frequency.

**Real-time response:** P4 The impulse response of every physical system is real, not complex. This requires that the Laplace transform have conjugate-symmetric symmetry $H(s) = H^*(s^*)$, where the $*$ indicates conjugation [e.g., $R(\sigma, \omega) + X(\sigma, \omega) = R(\sigma, \omega) - X(\sigma, -\omega)$].

**Time invariance:** P5 The meaning of time-invariant requires that the impulse response of a system does not change over time. This requires that the system coefficients of the differential equation describing the system are constant (independent of time).
Rayleigh reciprocity: P6  Reciprocity is defined in terms of the unloaded output voltage that results from an input current. Specifically (same as Eq. 3.7.2.6, p. 111),

\[
\begin{bmatrix}
z_{11}(s) & z_{12}(s) \\
z_{21}(s) & z_{22}(s)
\end{bmatrix} = \frac{1}{C(s)} \begin{bmatrix} A(s) & \Delta_T \\ 1 & D(s) \end{bmatrix},
\]

where \( \Delta_T = A(s)D(s) - B(s)C(s) = \pm 1 \) for the reciprocal and antireciprocal systems, respectively. This is best understood in terms of Eq. F.1.1.2. The off-diagonal coefficients \( z_{12}(s) \) and \( z_{21}(s) \) are defined as

\[
z_{12}(s) = \frac{\Phi_i}{U_i} \bigg|_{l_1=0}, \quad z_{21}(s) = \frac{F_l}{T_l} \bigg|_{U_i=0}.
\]

When these off-diagonal elements are equal \([z_{12}(s) = z_{21}(s)]\), the system is said to obey Rayleigh reciprocity. If they are opposite in sign \([z_{12}(s) = -z_{21}(s)]\), the system is said to be antireciprocal. If a network has neither reciprocal nor antireciprocal characteristics, then we denote it nonreciprocal (McMillan, 1946). The most comprehensive discussion of reciprocity, even to this day, is that of Rayleigh (1896, Vol. I). The reciprocal case may be modeled as an ideal transformer (Van Valkenburg, 1964a), while for the antireciprocal case the generalized force and flow are swapped across the two-port. An electromagnetic transducer (e.g., a moving-coil loudspeaker or electrical motor) is antireciprocal (Kim and Allen, 2013; Beranek and Mellow, 2012); it requires a gyrator rather than a transformer, as shown in Fig. F.1.

Reversibility: P7  A second two-port property is the reversible/nonreversible postulate. A reversible system is invariant to the input and output impedances being swapped. This property is defined by the input and output impedances being equal.

Referring to Eq. F.2.0.3, when the system is reversible, \( z_{11}(s) = z_{22}(s) \) or, in terms of the transmission matrix variables, \( \frac{A(s)}{C(s)} = \frac{D(s)}{C(s)} \) or simply \( A(s) = D(s) \), assuming \( C(s) \neq 0 \).

An example of a nonreversible system is a transformer with a turns ratio that is not 1. Also, an ideal operational amplifier (when the power is turned on) is nonreversible due to the large impedance difference between the input and output. Furthermore, it is active; it has a power gain due to the current gain at constant voltage (Van Valkenburg, 1964b).

Generalizations of this lead to group theory and Noether’s theorem. These generalizations apply to systems that have many modes, whereas quasistatics hold when they operate below a cutoff frequency (Table F.1), meaning that, as in the case of the transmission line, there are no propagating transverse modes. While this assumption is never exact, it leads to highly accurate results because the nonpropagating evanescent transverse modes are attenuated over a short distance and thus, in practice, may be ignored (Montgomery et al., 1948; Schwinger and Saxon, 1968, Chaps. 9–11).

We extend the Carlin and Giordano postulate set to include Postulate P7, reversibility, which was refined by Van Valkenburg (1964a). To satisfy the reversibility condition, the diagonal components in a system’s impedance matrix must be equal. In other words, the input force and flow are proportional to the output force and flow, respectively (i.e., \( Z_e = Z_m \)).

Spatial invariance: P8  The characteristic impedance and wave number \( \kappa(s,x) \) may be strongly frequency- and/or spatially dependent or even be negative over some limited frequency ranges. Due to causality, the concept of a negative group velocity must be restricted to a limited bandwidth (Brillouin, 1960). As Einstein’s theory of relativity makes clear, all materials must be strictly causal (Postulate P1), a view that must therefore apply to acoustics but at a very different time scale. We first discuss generalized postulates, expanding on those of Carlin and Giordano.

Deterministic (randomness): P9  When the media are uniform and time-invariant, the impedance and transfer functions will be deterministic. When the media are turbulent, the response will be random. When light propagates through the universe, it is strongly time-varying. Thus, astrophysics will be seen as random, whereas experiments in the shelter of the relatively uniform environment of the earth will be deterministic. Model calculation will also be deterministic unless one is trying to create a random, time-varying, turbulent medium.

The quasistatic constraint: P10  When a system is described by the wave equation, delay is introduced between two points in space and depends on the wave speed. When the wavelength is large compared to the delay, we can successfully apply the quasistatic (QS) approximation. This method has widespread application and is frequently used without mention of the assumption. This can lead to confusion, since the limitations of the approximation
may not be appreciated. An example is the use of quasistatics in quantum mechanics. The QS approximation has widespread use when the signals may be accurately approximated by a band-limited signal. Examples include KCL, KVL, wave guides, transmission lines, and most important, impedance. The QS property is not mentioned in the six postulates of Carlin and Giordano (1964); thus they need to be extended in some fundamental ways.

When the dimensions of a cellular structure in the material are much smaller than the wavelength, can the QS approximation be valid? This effect can be viewed as a mode filter that suppresses unwanted (or conversely enhances the desired) modes (Ramo et al., 1965). QS may be applied to a three-dimensional specification, as in a semiconductor lattice. But such applications fall outside the scope of this text (Schwinger and Saxon, 1968).

Although I have never seen the idea discussed in the literature, the QS approximation is applied when Green’s theorem is defined. For example, Gauss’s law is not true when the volume of the container violates QS, since changes in the distribution of the charge have not reached the boundary, when doing the integral. Thus such integral relationships assume that the system is in quasi-steady-state (i.e., that QS holds).

Table F.1: Several ways of indicating the quasistatic (QS) approximation. For network theory there is only one lattice constant, a, which must be much less than the wavelength (wavelength constraint). These three constraints are not equivalent when the object may be a larger structured medium, spanning many wavelengths but with a cell structure size much smaller than the wavelength. For example, each cell could be a Helmholtz resonator or an electromagnetic transducer (e.g., an earphone).

<table>
<thead>
<tr>
<th>Measure</th>
<th>Domain</th>
</tr>
</thead>
<tbody>
<tr>
<td>$ka &lt; 1$</td>
<td>Wave number constraint</td>
</tr>
<tr>
<td>$\lambda &gt; 2\pi a$</td>
<td>Wavelength constraint</td>
</tr>
<tr>
<td>$f_c &lt; c/2\pi a$</td>
<td>Bandwidth constraint</td>
</tr>
</tbody>
</table>

Formally, QS is defined as $ka < 1$, where $k = 2\pi/\lambda = \omega/c$ and $a$ is the cellular dimension or the size of the object. Other ways of expressing this include $\lambda/4 > a$, $\lambda/2\pi > a$, $\lambda > 4a$, or $\lambda > 2\pi a$. It is not clear whether it is better to normalize $\lambda$ by $4$ (quarter-wavelength constraint) or $2\pi \approx 6.28 > 4$, which is more conservative by a factor of $\pi/2 \approx 1.6$. Also $k$ and $a$ can be vectors (e.g., Eq. 3.1.0.5, p. 63).

Sergei Schelkunoff may have been the first to formalize this concept (Schelkunoff, 1943), but he was not the first to use it, as exemplified by the Helmholtz resonator. George Ashley Campbell was the first to use the concept in the important application of a wave filter, some 30 years before Schelkunoff (Campbell, 1903). These two men were 40 years apart and both worked for the telephone company (after 1929, called AT&T Bell Labs) (Fagen, 1975).

There are alternative definitions of the QS approximation, depending on the geometric cell structure. The alternatives are listed in Table F.1.

**The quasistatic approximation:** Since the velocity perpendicular to the walls of a horn must be zero, any radial wave propagation is exponentially attenuated ($\kappa(s)$ is real and negative, i.e., the propagation function $\kappa(s)$ (Sec. 4.4, p. 142) will not describe radial wave propagation), with a space constant of about 1 diameter. The assumption that these radial waves can be ignored (i.e., more than 1 diameter from their source) is called the quasistatic approximation. As the frequency is increased and once $f \geq f_c = 2c_0/\lambda$, the radial wave can satisfy the zero normal velocity wall boundary condition and therefore will not be attenuated. Thus above this critical frequency, radial waves (also known as higher-order modes) are supported ($\kappa$ becomes imaginary). Thus for Eq. 5.2.2.10 (p. 183) to describe guided wave propagation, $f < f_c$. But even under this condition, the solution is not precise within a diameter (or so) of any discontinuities (i.e., rapid variations) in the area.

Each horn, as determined by the area function $A(r)$, has a distinct wave equation and thus a distinct solution. Note that the area function determines the upper cutoff frequency via the quasistatic approximation, since $f_c = c_0/\lambda_c$, $\lambda_c/2 > d$, and $A(r) = \pi(d/2)^2$. Thus to satisfy the quasistatic approximation, the frequency $f$ must be less than the cutoff frequency:

$$f < f_c(r) = \frac{c_0}{4} \sqrt{\frac{\pi}{A(r)}}. \quad (F.2.0.4)$$

We have discussed two alternative matrix formulations of these equations: the ABCD transmission matrix, used for computation, and the impedance matrix, used when working with experimental measurements (Pierce, 1981, Chap. 7). For each formulation, reciprocity and reversibility show up as different matrix symmetries, as addressed in Sec. 3.9 (p. 119) (Pierce, 1981, pp. 195–203).

**Periodic ↔ discrete: P11** As has been shown in the discussion on the Fourier transform, when the time (or frequency) domain response is periodic, the frequency (or time) domain is discrete. This is a fundamental symmetry
property that must always be obeyed. This is closely related to the causal $\leftrightarrow$ complex analytic property of the Laplace and $z$ transforms.

**Summary**

A transducer converts between modalities. We propose the general definitions of the eleven system postulates that include all transduction modalities, such as electrical, mechanical, and acoustical. It is necessary to generalize the concept of the QS approximation (Postulate P9) to allow for guided waves.

Given the combination of the important QS approximation and these space-time, linearity, and reciprocity properties, a rigorous definition and characterization of a system can thus be established. It is based on a taxonomy of such materials formulated in terms of material and physical properties and extended network postulates.
Appendix G

Webster horn equation derivation

G.1 Overview

In this appendix we transform the acoustic equations, Eqs. 5.2.1.5 and 5.2.1.6 (p. 182), into their equivalent integral form, Eq. 5.2.2.10 (p. 183). This derivation is similar (but not identical) to that of Hanna and Slepian (1924) and Pierce (1981, p. 360).

G.1.1 Conservation of momentum

The first step is to integrate the normal component of Eq. 5.2.1.5 (p. 182) over the isopressure surface $S$, defined by $\nabla p = 0$

$$-\int_S \nabla p(x,t) \cdot dA = \rho_o \frac{\partial}{\partial t} \int_S u(x,t) \cdot dA.$$

The average pressure $\varrho(x,t)$ is defined by dividing by the total area:

$$\varrho(x,t) \equiv \frac{1}{A(x)} \int_S p(x,t) \hat{n} \cdot dA.$$ (G.1.1.1)

From the definition of the gradient operator, we have

$$\nabla p = \frac{\partial p}{\partial x} \hat{n},$$ (G.1.1.2)

where $\hat{n}$ is a unit vector perpendicular to the isopressure surface $S$. Thus the left side of Eq. 5.2.1.5 reduces to $\partial \varrho(x,t)/\partial x$.

The integral on the right side defines the volume velocity,

$$\nu(x,t) \equiv \int_S u(x,t) \cdot dA.$$ (G.1.1.3)

Thus the integral form of Eq. 5.2.1.5 becomes

$$-\frac{\partial}{\partial x} \varrho(x,t) = \frac{\rho_o}{A(x)} \frac{\partial}{\partial t} \nu(x,t) \leftrightarrow Z(x,s) V',$$ (G.1.1.4)

where

$$Z(s,x) = \frac{s \rho_o}{A(x)} = s M(x)$$ (G.1.1.5)

and $M(x) = \rho_o/A(x)$ [kgm/m$^5$] is the per-unit-length mass density of air.

G.1.2 Conservation of mass

Integrating Eq. 5.2.1.6 over the volume $V$ gives

$$-\int_V \nabla \cdot u \, dV = \frac{1}{\eta_o P_o} \frac{\partial}{\partial t} \int_V p(x,t) \, dV.$$
Figure G.1: Derivation of the horn equation using Gauss’s law: The divergence of the velocity \( \nabla \cdot \mathbf{u} \) within \( \delta x \), shown as the shaded region, is integrated over the enclosed volume. Next the divergence theorem is applied, transforming the integral to a surface integral normal to the surface of propagation. This results in the difference of the two volume velocities \( \delta \nu = \nu(x + \delta x) - \nu(x) = |\mathbf{u}(x + \delta x) \cdot \mathbf{A}(x + \delta x) - \mathbf{u}(x) \cdot \mathbf{A}(x)| \). The flow is always perpendicular to the isopressure contours.

The volume \( V \) is defined by two isopressure surfaces between \( x \) and \( x + \delta x \) (the shaded region of Fig. G.1). On the right-hand side we use the definition of the average pressure (i.e., Eq. G.1.1.1) integrated over the volume \( dV \).

Applying Gauss’s law to the left-hand side\(^1 \) and using the definition of \( \varrho \) (on the right) in the limit \( \delta x \to 0 \) give

\[
- \frac{\partial}{\partial x} \nu(x, t) = \frac{A(x)}{\eta_o P_o} \frac{\partial}{\partial t} \varrho(x, t) \leftrightarrow \mathcal{Y}(x, s) \mathcal{P}(x, s),
\]

where

\[
\mathcal{Y}(x, s) = s A(x)/\eta_o P_o = s C(x).
\]

This assumes the medium is lossless. For a discussion of lossy propagation, see Appendix D (p. 249).

### G.1.3 Horn properties

#### Speed of sound \( c_o \)

In terms of \( M(x) \) and \( C(x) \), the speed of sound and the acoustic admittance are

\[
c_o = \sqrt{\frac{\text{stiffness}}{\text{mass}}} = \sqrt{\frac{1}{C(x)M(x)}} = \sqrt{\frac{\eta_o P_o}{\varrho_o}},
\]

### G.1.4 Characteristic admittance \( \mathcal{Y}_r(x) \):

Since the horn equation (Eq. 5.2.2.10) is second-order, it has two eigenfunction solutions \( \mathcal{P}^\pm \). The ratios of Eq. G.1.2.7 to Eq. G.1.1.5 are determined by the local stiffness \( 1/C(x) \) and mass \( M(x) \). The ratio \( C/M \) determines the area-dependent characteristic admittance \( \mathcal{Y}_r(x) \) (\( \in \mathbb{R} \)):

\[
\mathcal{Y}_r(x) = \frac{1}{\sqrt{\text{stiffness} \cdot \text{mass}}} = \sqrt{\frac{\mathcal{Y}(x, s)}{\mathcal{Z}(x, s)}} = \sqrt{\frac{C(x)}{M(x)}} = \frac{A(x)}{\rho_o \eta_o P_o} = \frac{A(x)}{\rho_o c_o} > 0
\]

\((\text{Campbell, 1903, 1910, 1922})\). The characteristic impedance is \( \mathcal{Z}_r(x) = 1/\mathcal{Y}_r(x) \). Based on a physical argument, \( \mathcal{Y}_r(x) \) must be positive and real; thus only the positive square root is allowed. As long as \( A(x) \) has no jumps (is continuous), \( \mathcal{Y}_r(x) \) must be the same in both directions. It is locally determined by the isopressure surface and its volume velocity.

#### Radiation admittance

The radiation admittance is defined looking into a horn with no termination (infinitely long) from the input at \( x = 0 \):

\[
Y_{rad}^\pm(s) = \frac{\mathcal{Y}_r^\pm}{\mathcal{P}^\pm} \in \mathbb{C}.
\]

The impedance depends on the direction, with \( + \) looking to the right and \( - \) to the left.

---

\(^1\)As shown in Fig. G.1, taking the limit of the difference between the two volume velocities \( \nu(x + \delta x) - \nu(x) \) divided by \( \delta x \) results in \( \partial \nu/\partial x \).
The input admittance $Y^\pm_{in}(x, s)$ is computed using the upper equation of Eq. 5.2.3.11 (p. 185) for $V^\pm(x, s)$ and then dividing by the pressure eigenfunction $P^\pm$. This results in the logarithmic derivative of $P^\pm$:

$$Y^\pm_{in}(x, s) \equiv \frac{Q^\pm}{P^\pm} = \frac{-1}{sM(x)} \frac{\partial}{\partial r} \ln P^\pm(x, s).$$

For example, for the conical horn (last column of Table 5.2, p. 190)

$$Y^\pm_{in} = Y_r (1 \pm \frac{c_0}{sr_0}). \quad (G.1.4.11)$$

Note that $Y^+_{in}(x, s) + Y^-_{in}(x, s) = 2Y_r = 2A_0r^2/\rho_0c_0 \in \mathbb{R}$, which shows that the frequency-dependent parts of the two admittances, being equal and opposite in sign, exactly cancel.

As the wavefront travels down the variable-area horn, there is a mismatch in the characteristic admittance due to the change in area. This mismatch creates a reflected wave, which in the case of the conical horn is $-\frac{c_0}{sr_0}$. Due to conservation of volume, there is a corresponding identical forward component that travels forward, equal to $+\frac{c_0}{sr_0}$. The sum of these two responses to the change in area must be zero in order to conserve volume velocity.

The resulting equation for the velocity eigenfunctions is therefore

$$V^\pm(x, s) = Y^\pm_{in}(x, s)P^\pm(x, s).$$

**Propagation function $\kappa(s)$** The eigenfunctions of the lossless wave equation propagate as

$$P^\pm(x, s) = e^{\mp\kappa(s)x/\sqrt{A(x)}}$$

where $\kappa(s) = \sqrt{Z(x, s)\gamma(x, s)} = \pm s\sqrt{MC}$. The velocity eigenfunctions $V^\pm(x, s)$ may be computed from Eq. G.1.1.4.

From the above definitions,

$$\kappa(s) = \frac{sp_0}{A(x)} \frac{sA(x)}{n_0P_0} = \frac{s}{c_0}.$$  

Thus $\kappa(s)$ and $s$ are the eigenvalues of the differential operations $\partial/\partial x$ and $\partial/\partial t$ on the pressure $P(x, s)$. See Appendix D for the inclusion of viscothermal losses.
Appendix H

Quantum Mechanics and the WHEN

While it is clear that both Schrödinger’s equation and Dirac’s equations are highly accurate, after about 100 years, it is not clear why. Both of these theories seem to violate classical electromagnetics (EM), such as Ohm’s law, since they are built on energy principles rather than electric and magnetic fields. Here we delve into this question, by providing a classical (i.e., EM-based) derivation for the hydrogen atom, one of the most important and obvious successes of quantum mechanics (QM). The problem with QM is not that it fails—rather, it succeeds, without obvious basis. The problem is that we cannot understand the basic principles, and it seems to be in contradiction with any principles of a physical theory.

Based on the Rydberg series, we determine the reflection coefficient, and thus the radiation impedance seen by the electron, in a radial coordinate system centered on the proton. Since the electron and proton both have spin \( \frac{1}{2} \), their magnetic fields must attractively align, accounting for the near-field vector potential, and complementing the far-field attraction due to their opposite signs. As the electron and proton approach each other, due to their far-field potential attraction, the magnetic near field becomes more attractive at close range, due to the magnetic dipoles of the two “particles,” causing them to merge with neutral net magnetic moment and neutral charge, giving a highly stable hydrogen atom. However, given a sufficiently strong distorting field, this highly symmetric state could be disturbed, leading to photon radiation, constrained by the radial eigenstates. It seems more clear than ever that photons and electrons are in a state of equilibrium at the outskirts of very large Rydberg atoms.\(^1\)

\[ \text{Figure H.1: Diagram of the wavelength spectrum of hydrogen for the Lyman, Balmer, and Paschen series, as a function of each line’s wavelength. The notation “Ly-\( \alpha \)” indicates the longest wavelength } \lambda_{11} = 122 \, \text{[nm]} \text{ (i.e., lowest frequency of } 2.46 \, \text{[GHz]} \text{) for the Lyman series. Figure citation: https://en.wikipedia.org/wiki/Hydrogen_spectral_series} \]

\[ \text{H.1 Equation for Rydberg eigenmodes} \]

Like every tuned resonant circuit, atoms have well-defined resonant frequencies, or eigenmodes. Figure H.1 shows the observed radiation spectra for hydrogen. From the very beginning, it has been clear that there is a pattern to these spectral lines. In 1880 Rydberg easily fitted a formula that quantifies the observed eigen spectral lines of hydrogen in terms of the reciprocals of the radiated wavelengths:

\[
\frac{1}{\lambda_{nm}} = R_\infty \left( \frac{1}{n^2} - \frac{1}{m^2} \right), \quad \frac{f_{nm}}{c_0 R_\infty} = \frac{1}{n^2} - \frac{1}{m^2},
\]

all based on these simple observations. Here \( R_\infty = 1.097 \times 10^7 \, \text{[m}^{-1}] \); \( c_0 = 3 \times 10^8 \, \text{[m/s]} \) is the speed of light; \( f_{nm} \) are the dimensionless Rydberg integer frequencies; and \( n, m \in \mathbb{N} \) are positive integers \( \in \mathbb{N} \), where \( n \) labels\(^1\)

\[ \text{https://physics.aps.org/synopsis-for/10.1103/PhysRevLett.121.193401} \]
the series and $m > n$ ($\lambda > 0$) describes the transition from orbit $m$ to orbit $n$, as described in the caption of Fig. H.3.

### H.1.1 The Rydberg atom model

In 1909 Rutherford demonstrated that the atom consisted of a dense core (the proton) surrounded by electrons. This view was supported by the spectrum of the atom, which allows for a radiation spectrum caused by electrons jumping from one energy level to another. It was then noted by Bohr in 1913 (Bohr, 1954) and others that the wavelengths of hydrogen, as described by Eq. H.1.0.1, are consistent with Fig. H.2, where the reciprocal wavelength [m$^{-1}$] is given by Eq. H.1.0.1, having frequencies $f_{nm} = c/\lambda_{nm}$ [Hz]. The challenge of the 1920s was to explain these intuitive and simple models of hydrogen. This gave rise to the birth of quantum mechanics, the history of which is nicely summarized in Condon and Morse (1929).

![Figure H.3](https://en.wikipedia.org/wiki/Hydrogen_spectral_series)
The lines of this spectrum cumulate at the limit given by the Rydberg constant $R$. The adjoining *continuum* lies in the near ultraviolet range. Both the discrete and the continuous spectrum are given by the Schrödinger equation. This equation reduces to a simple mathematical formula the enigma of the spectral lines, with their finite cumulation point, the behavior of which differs so fundamentally from that of all mechanical systems.

### H.1.2 Rydberg wave equation

The objective of this analysis is to demonstrate that one can define a classical Sturm-Liouville model of the *enigmatic* Rydberg atom, by the use of the Webster horn equation

$$\frac{1}{A(r)} \frac{\partial}{\partial r} A(r) \frac{\partial}{\partial r} \psi(r, t) = \frac{1}{c^2} \frac{\partial^2}{\partial t^2} \psi(r, t),$$

which is a one-dimensional wave equation for the electric potential $\psi(r, t)$ propagating in a wave guide having area $A(r)$ as a function of the range, where $r$ is the *range* variable (the axis of wave propagation).

We shall show that given the Rydberg spectrum (Eq. H.1.0.1), we may accurately estimate the electric reflectance $\Gamma(s)$ looking out from the origin (i.e., the proton location, as indicated by the small red dot in Fig. H.3). The radiation impedance $Z_{\text{rad}}(s)$ seen by the proton is related to the reflectance $\Gamma(s)$ by the relation

$$Z_{\text{rad}}(s) = r_o \frac{1 + \Gamma(s)}{1 - \Gamma(s)}.$$  \hspace{1cm} (H.1.2.3)

This formula is the basis of the *Smith chart* used in both physics and engineering studies. It follows that once $\Gamma(s)$ is known (i.e., evaluated given Eq. H.1.0.1), the radiation impedance may be computed. It has been shown that the area function $A(r)$ may be found given the radiation impedance (Sondhi and Gopinath, 1971; Youla, 1964).

---

**Figure H.4:** The top panel is a plot of Eq. H.1.0.1, showing how the eigenmode frequencies $f_l$ depend on the eigen-number index $l$. As the mode number increases, the frequency reaches an asymptote at $c_o R \approx 3.29$ [GHz], with a wavelength limit near $1/R \approx 91.2$ [nm]. The lower panel shows the inverse mapping from frequency to the mode index number $\phi(f_l)$. This figure is for the Lyman series ($n = 1$ and $m = 1, \ldots, 20$). The inverse of this relationship is $l = \phi^{-1}(f_l)$ may be derived from Eq. H.1.0.1, which provides the pole frequencies required to satisfy Eq. H.1.2.3. Note that for frequencies greater than $c_o/R$ the phase switches from purely real to imaginary, accounting for the free electrons that must exist above the upper accumulation frequency (i.e., 3.29 [GHz] for this example).
**H.2 Rydberg solution methods**

The basic idea behind the method is to use Eq. H.1.2.3, by noting that the poles of the impedance are determined by the roots of the denominator of $Z_{\text{rad}}$. Specifically, if $s_p$ is an impedance pole, then it must satisfy $\Gamma(s_p) \approx 1$. Except for losses due to radiation, the atom is lossless; thus $|\Gamma(s)| = 1$. Namely, it must be of the form

$$\Gamma(s) = e^{-\phi(f)}, \quad \text{(H.2.0.1)}$$

where the phase $\phi(f) \in \mathbb{R}$ and $s = \sigma + \omega f$ is the complex Laplace radian frequency, with $\omega = 2\pi f$ [Hz]. Since we know the eigenmode frequencies, which obey $\phi(f_{nm, m}) = 2\pi m$, we may find $\phi(f)$, as follows: For a given series index $n_o$, and given the eigenfrequencies $f_m$, we seek the phase function $\phi_{n_o}(f)$ that maps the eigenfrequencies to their mode index $m$, i.e.,

$$\phi_{n_o}(f_m) = 2\pi m.$$

### H.2.1 Group delay $\tau(s)$

The phase $\phi(\omega)$ is related to the group delay $\tau(\omega)$ by the relation

$$\tau(\omega) = -\frac{\partial}{\partial \omega} \phi(\omega).$$

Here one may assume that the phase is complex analytic, thus allowing a causal damping term into the reflectance phase Eq. H.2.0.1. This follows naturally because the reflectance must be causal (Postulate 3.9.1, p. 121). In the time domain the delay may be written in terms of the inverse $\mathcal{L}$-transform of the group delay,

$$\Gamma(s) = e^{-\int_{s_0}^s \tau(s)ds}.$$

Typically one uses the reflectance phase $2\pi\phi(f)$; thus the group delay is $\tau(f) = -\partial \phi(f)/\partial f$, which is physically interpreted here as the frequency-dependent delay from the proton to the radius of the electron’s orbit. Thus this delay is given by

$$\tau(f) = n \frac{\partial}{\partial f} \left(1 - \frac{n^2}{c_\omega R} f \right)^{-1/2} = \frac{n^2}{2c_\omega R} \left(1 - \frac{n^2}{c_\omega R} f \right)^{-3/2},$$

which is constant for low frequencies and then rises to $\infty$ as frequency approaches the Rydberg frequency ($f \to c_\omega R/n^2$).

One may solve Eq. H.1.0.1 for $m$, for the case of the Lyman series ($n_o = 1$), by the use of the following identity for the Rydberg eigenfrequencies $f_{nm}$, which follow directly from Eq. H.1.0.1, with $m = n_o + l$ (with $n_o, m, l \in \mathbb{N}$)

$$f_{nm} = \frac{c_\omega}{\lambda_{nm}} = c_\omega R \left(\frac{1}{n_o^2} - \frac{1}{(n_o + l)^2}\right) = \frac{c_\omega R}{n_o^2} \left(1 - \frac{1}{(1 + l/n_o)^2}\right). \quad \text{(H.2.1.2)}$$

Note that as $l \to \infty$, $f_{n_o,l} \to c_\omega R/n_o^2$, which is Sommerfeld’s “finite cumulation point” [Hz] $f_{n_o,\infty}$ for the Lyman series ($n_o = 1$).

We can solve Eq. H.2.1.2 for the mode number $l/n < 1$ as a function of mode frequency:

$$n^2 \frac{f_{nm}}{c_\omega R} = 1 - \frac{1}{(1 + l/n)^2}$$

Starting from Eq. H.2.1.2

$$\frac{1}{(1 + l/n)^2} = 1 - n^2 \frac{f_{nm}}{c_\omega R}$$

Solving for $l/n$

$$(1 + l/n)^2 = \frac{1}{1 - n^2 \frac{f_{nm}}{c_\omega R}}$$

$$l/n = \pm \frac{1}{\sqrt{1 - n^2 \frac{f_{nm}}{c_\omega R}}} - 1$$

$$\phi(f_{nm})/2\pi = l = m - n_o \in \mathbb{N}, \quad \text{(H.2.1.3)}$$

as summarized in the lower panel of Fig. H.4.

---

²It follows that these relationships are related by a Hilbert transform.
H.2.2 Finding the area function

Once the phase has been determined, we can compute the impedance using Eq. H.1.2.3. We may then decompose the impedance by using the analytic continued fraction algorithm (or Cauer synthesis), discussed in Sec. 3.7, p. 110.

H.3 Euclid’s formula and the Rydberg atom model

Fundamental to quantum mechanics is the Rydberg series, which describes the quantized energy levels of atoms:

\[ \nu_{n,m} = c_o R_{\infty} Z_n^2 \left( \frac{1}{n^2} - \frac{1}{m^2} \right), \] (H.3.0.1)

where \( \nu_{n,m} \) are the possible eigenfrequencies, \( c_o \) is the speed of light, \( R_{\infty} \approx 1.097 \times 10^7 \) is the Rydberg constant, \( Z_n \) is the atomic number, along with positive integers \( m > n \in \mathbb{N} \), which represent the principal quantum numbers that label all possible allowed atomic eigenstates. Integer \( n \) indicates the lowest (rest) atomic eigenstate while \( m \) labels the higher (excited) state. When \( n = 1 \), the series is the Lyman series corresponding to hydrogen (\( Z_1 = 1 \)). When \( n = 1, m = 2 \), and \( Z_1 = 1 \), the frequency is

\[ \nu_{1,1} = c_o R_{\infty} \left( \frac{1}{1^2} - \frac{1}{2^2} \right)^{\frac{3}{2}} = 2.5 \times 10^{15} \text{ [Hz]}, \] (H.3.0.2)

and the wavelength is \( \lambda = c_o/\nu = 4 \times 10^8 / R_{\infty} = 36.36 \text{ [m]} \).

An open question in this model is: Why are states either empty or filled? The amplitudes of the modes of a string or organ pipe are not quantized. What is it about the atom that forces the energy to be quantized?

Given observed frequencies \( \nu_{n,m} \) it is possible to find the area function that traps the photons into the Rydberg eigenstates. Eq. H.3.0.1 may be rewritten as

\[ \nu_{n,m} = c_o R Z_n^2 \frac{a^2 - n^2}{(2nm)^2} . \]

It is interesting to compare Eq. H.3.0.1 to Euclid’s formula Eq. 2.5.5.6 (p. 50):

\[ a = m^2 - n^2, \quad b = 2mn, \quad c = m^2 + n^2, \] (H.3.0.3)

where \( m > n \in \mathbb{N} \). Euclid’s formula is equivalent to the Pythagorean theorem for integers, since

\[ c^2 = a^2 + b^2, \] (H.3.0.4)

with \( \{a, b, c\} \in \mathbb{N} \). Here \( a < b < c \).

If we interpret the quantum numbers as multiples of a quarter wavelength, then the Rydberg formula is congruent to the Pythagorean theorem of integers. Given the symmetry, this cannot be an accident.

In terms of the lengths of the right triangle \( \{a, b, c\} \), Rydberg’s formula becomes

\[ \nu_{n,m} = c_o R Z_n^2 \frac{a}{b^2} . \]

But since \( b^2 = c^2 - a^2 \),

\[
\nu_{n,m} = c_o \frac{R Z_n^2}{a} \left( \frac{a^2}{b^2} \right) = c_o \frac{R Z_n^2}{a} \frac{a^2}{c^2} \left( \frac{1}{1 - (a/c)^2} \right) .
\]

---

3https://www.youtube.com/watch?v=e0IWPEhmMho

In terms of quantized (discrete) angles, \( \sin(\theta_{n,m}) = \alpha/c \),

\[
\nu_{n,m} = c_0 \frac{RZ_n^2}{a} \frac{1}{4} \left( \sin^2 \theta \right) \frac{1}{1 - \sin^2 \vartheta} \cos^2 \vartheta
\]

\[
= c_0 \frac{RZ_n^2}{a} \frac{1}{4} \sin^2 \theta \cos^2 \theta
\]

\[
= c_0 \frac{RZ_n^2}{a} \frac{1}{4} \tan^2 \theta_{n,m}.
\]

### H.3.1 Eigenmodes of the Rydberg atom

One way to think of eigenmodes is to make an analogy to a piano string or an organ pipe. In these much simpler systems, there is an almost constant delay, say \( \tau \), due to a characteristic length, say \( L = \tau c_0 \), such that the eigenmodes of a string are given by integer multiples of a half wavelength \( \nu_n = n c_0/2L \), while the eigenmodes of the organ pipe are multiples of a quarter wavelength. The distinction is the boundary conditions. For the string the endpoint boundary conditions are pinned displacement (i.e., zero velocity). The organ pipe is closed at one end and open at the other, resulting in multiples of a quarter wavelength \( \nu_n = nc_0/4L \). We suggest looking at the Rydberg series in the same way, but with very different eigenfrequencies (Eq. H.3.0.1). Sommerfeld (1949, p. 201) makes a very interesting comment regarding Eq. H.3.0.1:

This equation reduces to a simple mathematical formula the enigma of the spectral lines, with their finite cumulation point, the behavior of which differs so fundamentally from that of all mechanical systems.

### H.3.2 Discussion

The Rydberg frequencies \( f_{nl} (n = 1, l = 1, \ldots, \infty) \) has poles in the radiation impedance (Eq. H.1.2.3) when \( \phi_l(f_{nl}) \in \mathbb{N} \). Working backwards from the Rydberg formula (Eq. H.2.0.1), we have solved for \( \phi(f_{nl}) \) indicating where this condition is valid (Eq. H.2.1.3). Since the reflectance and the impedance must be causal complex analytic functions of Laplace frequency \( s \), we must replace the discrete frequency \( f_{nl} \) with \( s \):

\[ j2\pi f_{nl} \to s = \sigma + \omega_j, \]

thereby forcing \( l(s) \) to be a complex analytic function of \( s \). Then the poles of the radiation impedance must satisfy

\[ \Gamma(s_{nl}) = e^{2\pi l(f_{nl})} = 1, \]

resulting in eigenfrequencies at \( f_{nl} \).

The next step in this analysis is to determine the area function \( A(r) \) given \( Z_{rad} \) (Eq. H.1.2.3). To do this we must solve an integral equation for \( A(r) \), as discussed by Sondhi and Gopinath (1971) and by Youla (1964).

Perhaps this could be done using an analytic representation for the area function,

\[ A(r) = \sum_k a_k r^k. \]

### H.4 Relations between Sturm-Liouville and quantum mechanics

If we compare the Schrödinger equation (SE) for hydrogen with the corresponding Sturm-Liouville equation we can begin to appreciate their differences. The QM equation for hydrogen is

\[
\frac{\hbar}{2m_o} \frac{\partial}{\partial t} \psi(x,t) = -\frac{\hbar^2}{2m_o} \nabla^2 r \psi(x,t) + V(r) \psi(x,t)
\]

\[
= -\frac{\hbar^2}{2m_o} \frac{\partial}{\partial r} r^2 \psi(x,t) + V(r) \psi(x,t) \]

\[ (H.4.0.1) \]

whereas the horn equation is given by Eq. H.1.2.2.
There are several obvious and disturbing differences between these two equations. First, the SE is, of course, first-order in time. Diffusion equations have no delay and thus cannot have traditional eigenmodes, which result from standing waves in a wave equation, due to boundary conditions. Second, the EM horn equation is of Sturm-Liouville (SL) form, which is a true wave equation (vs. the SE, which is a diffusion equation). The obvious question arises: Is there a potential \( V \) that would allow these two formulations to be equivalent? If so, then this would provide an explanation as to why the SE is successful in explaining the properties of Rydberg atoms.

To explore this possibility we may expand the two differential equations and directly compare them. Expanding Eq. H.1.2.2 gives

\[
\frac{1}{c_0^2} \frac{\partial^2}{\partial t^2} \psi(r, t) = \frac{1}{A(r)} \frac{\partial}{\partial r} A(r) \frac{\partial}{\partial r} \psi(r, t)
\]

(H.4.0.3)

\[
= \frac{\partial^2}{\partial r^2} \psi(r, t) + \frac{1}{A(r)} \frac{\partial A(r)}{\partial r} \psi(r, t).
\]

(H.4.0.4)

Between these two equations we may remove \( \psi \):

\[
\hbar \frac{\partial}{\partial t} \psi(x, t) = -\frac{\hbar^2}{2m_o} \left[ \frac{2}{r} \frac{\partial}{\partial r} \psi(x, t) + \frac{1}{c_0^2} \frac{\partial^2}{\partial t^2} \psi(r, t) - \frac{1}{A(r)} \frac{\partial A(r)}{\partial r} \psi(r, t) \right] + V(r) \psi(x, t).
\]

(H.4.0.5)

It seems that this may isolate the time and spatial parts (as in separation of variables).

### H.4.1 The exponential horn

A relevant and motivational example is the solution of the exponential horn, having area function \( A(r) = A_o e^{2mr} \).

This case is interesting because it has a closed-form solution, which seems relevant and perhaps even related to the hydrogen atom.

Assuming that \( \rho(r, t) \leftrightarrow \mathcal{P}(r, \omega) \) are a Fourier transform pair, with \( A(r) = A_o e^{2mr} \), Eq. H.1.2.2 reduces to

\[
\frac{\partial^2}{\partial r^2} \mathcal{P}(r, \omega) + 2m \frac{\partial}{\partial r} \mathcal{P}(r, \omega) = \kappa^2 \mathcal{P}(r, \omega) \leftrightarrow \kappa^2 \rho(r, \omega) \leftrightarrow \frac{1}{c_0^2} \frac{\partial^2}{\partial t^2},
\]

(H.4.1.6)

with \( \kappa(s) = s/c_0 \).

**Exercise #1**

Show that Eq. H.4.1.6 follows from Eq. H.1.2.2.

**Solution:** Starting from Eq. H.1.2.2 with area \( A(r) = A_o e^{2mr} \)

\[
\frac{1}{A_o e^{2mr}} \frac{\partial}{\partial r} \left( A_o e^{2mr} \frac{\partial \rho}{\partial r} \right) = \frac{1}{c_0^2} \frac{\partial^2 \rho}{\partial t^2}
\]

\[
\rho_{rr}(r, t) + 2m \rho_r(r, t) = \frac{1}{c_0^2} \frac{\partial^2 \rho}{\partial t^2} \leftrightarrow \kappa^2 \mathcal{P}(r, \omega),
\]

which is the time-domain version of Eq. H.4.1.6.

Since this equation is second-order in time with constant coefficients, it has two closed-form solutions:

\[
\mathcal{P}^\pm(r) = \mathcal{P}_0^\pm(\omega) e^{-mr \sqrt{m^2 + \kappa^2}}
\]

\[
= \mathcal{P}_0^\pm(\omega) e^{-mr \sqrt{\omega^2 - \omega_c^2}},
\]

with \( \omega_c = mc_0 \). The two wave amplitudes \( \mathcal{P}_0^\pm(\omega) \) must be determined from the boundary conditions.

**Exercise #2**

Show that \( \mathcal{P}^\pm(r, \omega) \) satisfy Eq. H.4.1.6.

**Solution:** Taking partials with respect to \( r \),

\[
\frac{\partial}{\partial r} \mathcal{P}^\pm(r, \omega) = \left( -m \mp \sqrt{m^2 + \kappa^2} \right) \mathcal{P}^\pm(r, \omega)
\]

\[
\frac{1}{c_0^2} \frac{\partial^2}{\partial t^2} \mathcal{P}^\pm(r, \omega) = \left( -m \mp \sqrt{m^2 + \kappa^2} \right)^2 \mathcal{P}^\pm(r, \omega)
\]

\[
= \left( 2m^2 + \kappa^2 \pm 2m \sqrt{m^2 + \kappa^2} \right) \mathcal{P}^\pm(r, \omega).
\]
Thus Eq. H.4.1.6 reduces to
\[(2m^2 + \kappa^2 \pm 2m\sqrt{m^2 + \kappa^2}) + 2m(-m \mp \sqrt{m^2 + \kappa^2}) = \kappa^2,\]
which is an identity. ■

Next consider the Fourier series (or Fourier transform) of the area function,
\[A(r) = \sum_k a_k e^{2mkr}.\]

It follows from the linearity of the wave equation that the general solution of Eq. H.4.1.6 is
\[P^{\pm}(r, \omega) = \sum_k a^{\pm}_k(\omega) e^{-mkr} e^{\mp \sqrt{m^2 + \kappa^2}r}.\]

Here we have combined \(P^{\pm}(\omega)\) and \(a_k\) as coefficients \(a^{\pm}_k(\omega)\).
Bibliography


Appendix I

Figure I.1:  *Timeline from the early Asians to Bombelli (p. 13).*

Figure I.2:  *Timeline from Descartes to Cauchy (p. 15).*
**Chronological history: 16\textsuperscript{th} to 19\textsuperscript{th} centuries**

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<td>Maxwell</td>
<td>WWI</td>
<td>Hilbert</td>
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<td>Jacob Bernoulli</td>
<td>Mozart</td>
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Figure I.3:  *Timeline from Bombelli to Gauss* (p. 19).

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<td>Mozart</td>
<td>Beethoven</td>
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Figure I.4:  *Timeline from Newton to Brillouin* (p. 62).
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Figure J.2: Fibonacci spiral (p. 54).

Figure J.3: Newtons method, applied to two polynomials: real roots (left); complex roots (right) (p. 67).
Figure J.4: Colorized plots: Left: trivial case $w(s) = s$; Right: $w(s) = s - \sqrt{j}$ (p. 125).

Figure J.5: $w(s) = \sin(0.5\pi(s - j))$ (p. 126).

Figure J.6: Colorized plots of $w(z) = e^z$ (left) and $w(z) = \ln(z)$ (right) (p. 126).
Figure J.7: Plots of $w(z) = \text{atan}(z)$ (left) and $w(z) = \frac{i}{2} \ln \frac{1-i z}{1+i z}$ (right) (p. 134).

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