A Tight-Binding Approach to Creating van der Waals Metamaterials

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Abstract

Van der Waals heterostructures are a fertile frontier for discovering emergent phenomena in condensed matter systems. They are constructed by stacking elements of a large library of two-dimensional materials, which couple together through van der Waals interactions. However, the number of possible combinations within this library is staggering, and fully exploring their potential is a daunting task. Van der Waals metamaterials are designed to reshape the flow of acoustic waves to mimic electron motion. It has been previously shown that the van der Waals interaction can be mimicked in metamaterials through the use of an interlayer coupling membrane. I now present a method for designing new van der Waals metamaterials following tight-binding principles. I first present the connected-cavity model for designing two-dimensional metamaterials. I then demonstrate the effectiveness of the model by creating metamaterial graphene, AA- and AB-stacked metamaterial bilayer graphene, and metamaterial twisted bilayer graphene.
List of Figures

1.1 (a) A one-dimensional Bravais lattice has periodicity $a$. (b) A two-dimensional rectangular Bravais lattice has primitive lattice vectors $\vec{a}_1$ and $\vec{a}_2$.

1.2 There is no unique choice of primitive vectors for a Bravais lattice. Here we see just four possible sets of primitive vectors for the triangular lattice, but there are in fact an infinite number of possibilities.

1.3 (a) Each point in the triangular lattice is equivalent to every other point, making it a Bravais lattice. (b) In the honeycomb lattice, points A and B are equivalent but A and C are not, meaning it is not a Bravais lattice.

1.4 The honeycomb lattice has many possible unit cells (red) and primitive cells (green). While generic unit cells may contain any number of lattice points, each primitive cell contains only one lattice point. Widely-used conventional unit cells (blue) contain two lattice points.

1.5 A two-dimensional hexagonal Brillouin zone has high-symmetry $k$ points $\Gamma$ (green, at the center of the Brillouin zone), K (blue, at the vertices of the hexagon, with adjacent K points differentiated as K and $K'$), and M (red, at the midpoint of the hexagon edges).

1.6 The energy needed for an electron to hop to a nearest neighbor if $t$. Each gray bar connects nearest neighbors. One pair of nearest neighbors in shown in red; one pair of next-nearest neighbors is shown in green; one pair of next-next-nearest neighbors in shown in blue.

2.1 Two honeycomb lattices twisted relative to each another form periodic Moiré patterns at certain commensurate angles. The conventional unit cell of the overall Moiré pattern (red) increases in size as the twist angle ($\theta$) decreases. When $\theta = 0^\circ$, the system is bilayer graphene and its unit cell is the same size as graphene’s. When $\theta = 21.8^\circ$, the twisted bilayer supercell contains 7 graphene unit cells; when $\theta = 9.43^\circ$, the supercell contains 37 graphene unit cells; when $\theta = 5.09^\circ$, the supercell contains 127 graphene unit cells.
3.1 Here we visualize a 2SB honeycomb lattice of connected cavities as the basic design for phononic metamaterial graphene. Channels connect nearest neighbor hopping sites. Nearest neighbor hopping \( t \) occurs either from an \( A_1 \) site to an \( A_2 \) site or vice versa.  

3.2 (a) This unit cell for 2SB metamaterial graphene shows air cavities in purple, steel in opaque gray, and HDPE bounding layers in semi-transparent gray. Here \( a = 10 \) mm, \( R = 3.5 \) mm, \( w = R/4 = 0.875 \) mm, and \( D = 1 \) mm. (b) The metamaterial’s resulting band structure displays an isolated Dirac-like crossing (blue) mimicking graphene.

3.3 Unconnected-cavity sites \( (A_3) \) have been added to the original honeycomb lattice of connected cavities. Channels connect nearest neighbor hopping sites (with hopping energy \( t \)); there is no hopping from \( A_1 \) or \( A_2 \) sites to \( A_3 \) sites.

3.4 (a) The unit cell for 3SB connected-cavity metamaterial graphene has the same properties as the unit cell in Figure 3.2(a). The additional unconnected cavities have radius \( r = 0.75R = 2.625 \) mm. (b) The metamaterial’s resulting band structure displays an isolated Dirac-like crossing (blue) identical to the one shown in Figure 3.2(b). The extra band (gray) represents an isolated air mode within the extra cavities.

3.5 Here we visualize the coupling between 2SB honeycomb cavity networks in either layer of AB-stacked metamaterial bilayer graphene. Connective channels within each layer indicate intralayer hopping \( t \); Overlapping sites have interlayer hopping \( \Delta \). \( A_2 \) and \( B_1 \) sites overlap; \( A_1 \) and \( B_2 \) do not.

3.6 (a) This unit cell for 2SB AB-stacked metamaterial bilayer graphene shows air cavities in purple and steel in opaque gray. The metamaterial layers and outer HDPE bounding layers have the same properties \( (a, R, w, \text{and } D) \) as in Figure 3.2(a). The center coupling layer is HDPE with thickness \( D \), identical to the outer bounding layers. (b) The metamaterial’s resulting band structure displays abnormal behavior at the K point not matching the expected parabolic “kissing” structure present in AB-stacked bilayer graphene.

3.7 Here we visualize the coupling between 3SB honeycomb cavity networks in either layer of AB-stacked metamaterial bilayer graphene. Connective channels within each layer indicate intralayer hopping \( t \); Overlapping sites have interlayer hopping \( \Delta \). \( A_2 \) and \( B_1 \) sites overlap as usual. \( A_1 \) sites overlap with \( B_3 \) sites and \( B_2 \) with \( A_3 \).
3.8 (a) This unit cell for 3SB AB-stacked metamaterial bilayer graphene has the same properties as in Figure 3.6(a), with additional unconnected air cavities \( r = 2.626 \text{ mm} \). (b) The metamaterial’s resulting band structure displays parabolic “kissing” at the K point mimicking AB-stacked bilayer graphene. The extra two bands (gray) represent isolated air modes in \( A_3 \) and \( B_3 \) sites, respectively.

3.9 Here we visualize the coupling between 2SB honeycomb cavity networks in either layer of AA-stacked metamaterial bilayer graphene. Connective channels within each layer indicate intralayer hopping \( (t) \); Overlapping sites have interlayer hopping \( (\Delta) \). Each cavity network overlaps entirely with the other; that is, \( A_1 \) sites overlap with \( B_1 \) sites and \( A_2 \) with \( B_2 \).

3.10 (a) This unit cell for 2SB AA-stacked metamaterial bilayer graphene has the same properties as in Figure 3.6(a), with one metamaterial layer rotated 60° so that both cavity networks entirely overlap. (b) The metamaterial’s resulting band structure displays double-Dirac-like crossings at the K point mimicking AA-stacked bilayer graphene.

3.11 Here we visualize the coupling between the 3SB honeycomb cavity networks in either layer of AA-stacked metamaterial bilayer graphene. Connective channels within each layer indicate intralayer hopping \( (t) \); Overlapping sites have interlayer hopping \( (\Delta) \). There is overlap between \( A_1 \) and \( B_1 \), \( A_2 \) and \( B_2 \), and \( A_3 \) and \( B_3 \) sites.

3.12 (a) This unit cell for 3SB AA-stacked metamaterial bilayer graphene has the same properties as in Figure 3.8(a), with one metamaterial layer rotated 60° so that both cavity networks entirely overlap. (b) The metamaterial’s resulting band structure retains the expected double-Dirac-like crossings at the K point seen in Figure 3.10(b). The extra two bands (gray) represent isolated air modes in \( A_3 \) and \( B_3 \) sites.

3.13 TBG has regions of AB stacking (blue triangles) and AA stacking (red circles) connected by transition regions that have neither AA nor AB stacking.

3.14 The Dirac-like crossings (blue) originally present in 3SB AA-stacked metamaterial bilayer graphene become flatter as \( \theta \) decreases. Each band structure spans a smaller frequency range than the previous. Extra bands (gray) represent various other modes in the TBG heterostructure and are not relevant to the flattening bands.
3.15 (a) The metamaterial TBG supercells grow in size as $\theta$ decreases. Each supercell is composed of two 3SB metamaterial graphene layers (Figure 3.4) with an intervening HDPE coupling membrane, all sandwiched between two outer HDPE bounding membranes. As in all previously shown models, $a = 10$ mm, $R = 3.5$ mm, $r = 2.625$ mm, $w = 0.875$ mm, and $D = 1$ mm. (b) The band structures from Figure 3.14 are shown here plotted on the same frequency scale to visualize the flattening effect. Data is unavailable for the gray regions in the latter three band structures, but would contain additional extra bands (gray) irrelevant to the flattening Dirac-like crossings (blue).
# Contents

Acknowledgements i  
Abstract ii  
List of Figures iii  

1 Introduction  
1.1 Crystal Structure 1  
1.2 Reciprocal Space 2  
1.3 Tight-Binding 5  

2 Motivating Metamaterials 8  
2.1 Two-dimensional Materials and van der Waals Heterostructures 8  
2.2 Metamaterials 10  
2.3 Phononic Metamaterials as Quantum Mimics 10  
2.4 Work Presented 11  

3 Design Approach and Results 13  
3.1 The Connected Cavity Model 13  
3.2 Designing Two-dimensional Metamaterials 14  
3.2.1 Graphene 15  
3.3 Designing Stacked van der Waals Metamaterials 17  
3.3.1 AB-Stacked Bilayer Graphene 17  
3.3.2 AA-Stacked Bilayer Graphene 18  
3.4 Designing Twisted van der Waals Metamaterials 21  
3.4.1 Twisted Bilayer Graphene 22  

4 Conclusion and Future Works 29  
4.1 Expanding the Library of Two-dimensional Metamaterials 29  
4.2 Magic-Angle Metamaterial Twisted Bilayer Graphene 29  
4.3 Flat Band Metamaterials 30
Chapter 1

Introduction

1.1 Crystal Structure

Though the word “crystal” may call to mind angled gemstones, a true crystal is defined as a solid whose atoms are arranged in an ordered, periodic structure. As such, most solids, including metals, are crystalline in nature. The ideal crystal is structured as a \textbf{Bravais lattice}, an infinitely repeated pattern of identical units.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig1.png}
\caption{(a) A one-dimensional Bravais lattice has periodicity $a$. (b) A two-dimensional rectangular Bravais lattice has primitive lattice vectors $\vec{a}_1$ and $\vec{a}_2$.}
\end{figure}

Let us consider the simplest Bravais lattice: a one-dimensional chain of points (Figure 1.1(a)). If we define the chain’s periodicity as $a$, the position of any point $r$ is

$$r_n = na \quad n \in \mathbb{Z}.$$

Extending the lattice into two dimensions, let us consider a rectangular lattice of points (Figure 1.1(b)). We define this lattice’s periodicity through \textbf{primitive lattice vectors} $\vec{a}_1$ and $\vec{a}_2$. The position of any point $\vec{r}$ is then:

$$\vec{r}_{[n_1,n_2]} = n_1\vec{a}_1 + n_2\vec{a}_2 \quad n_1, n_2 \in \mathbb{Z}.$$
According to this definition, the choice of primitive lattice vectors is not unique (Figure 1.2). From these Bravais lattice definitions we see each lattice point has the same environment. In other words, the arrangement and orientation of the surrounding lattice appears exactly the same from the perspective of any point in a Bravais lattice.

In a non-Bravais lattice, however, each point is not necessarily equivalent to every other point. Consider the honeycomb lattice, which is similar to the triangular Bravais lattice but is not itself a Bravais lattice (Figure 1.3). To establish the periodicity of a non-Bravais lattice, we define a unit cell. A lattice unit cell is any region of a lattice that, when tiled out in space, reproduces the full lattice. A lattice’s primitive unit cell is its smallest possible unit cell, containing only one point. However, for convenience, we often use a conventional unit cell, usually chosen to be as small as possible while maintaining the lattice’s overall symmetries. There is no unique choice for any type of unit cell (Figure 1.4).

1.2 Reciprocal Space

While crystal structure is best described in real space, waves propagating through crystals are best described in reciprocal space. The relationship between real space and reciprocal space is entirely analogous to the relationship between the time domain and frequency domain; we can switch between real and reciprocal space via Fourier transform.

If points in a real space lattice represent positions, a lattice in reciprocal space will

---

1I have left out the explicit definitions for three-dimensional lattices because my research involves only two-dimensional materials. However, the mathematical definition for two-dimensional lattices can easily be extended to three dimensions by adding a third primitive lattice vector term. Again, the choice for all three vectors in that case would not be unique.

2I will henceforth specify all Bravais lattices as such; the term “lattice” alone can refer to non-Bravais lattices like the honeycomb lattice.

3Reciprocal space is often called “momentum space” or “$k$-space”.
Figure 1.3: (a) Each point in the triangular lattice is equivalent to every other point, making it a Bravais lattice. (b) In the honeycomb lattice, points A and B are equivalent but A and C are not, meaning it is not a Bravais lattice.

have points representing wavevectors. Such a lattice is called the reciprocal lattice, and we can generate it by taking the Fourier transform of a real space lattice\(^4\).

Let us consider a lattice of points in real space defined \(\vec{R}\). Consider also a generic plane wave \(e^{ikt}\). The periodicity of plane wave will match the periodicity of the lattice only at certain values of \(\vec{k}\). The set of such \(\vec{k}\) values in reciprocal space form the reciprocal lattice, which we will define as \(\vec{K}\). From this definition we can say \(\vec{K}\) is the reciprocal lattice of \(\vec{R}\) if and only if

\[
e^{i\vec{K} \cdot \vec{R}} = 1
\]

for all points in the direct lattice.

The relationship between the periodicity of the direct and reciprocal lattices is outlined by the formula

\[
\vec{b}_i \cdot \vec{a}_j = 2\pi \delta_{ij},
\]

where \(\delta_{ij}\) is the Kronecker delta\(^6\).

For a one-dimensional lattice \(R\) with periodicity \(a\), the reciprocal lattice periodicity is simply \(b = 2\pi/a\). That is, given \(R_n = na\), \(K_m = mb\), where \(m\) is an integer.

Similarly, a two-dimensional lattice \(\vec{R}\) with primitive lattice vectors \(\vec{a}_1\) and \(\vec{a}_2\) has a

\(^4\)When described in respect to a reciprocal lattice, the real space lattice is known as the “direct lattice”.

\(^5\)I am following Ashcroft & Mermin’s notation here [1], but other solid-state textbooks (like Simon’s [2]) may refer to the reciprocal lattice as \(\vec{G}\).

\(^6\)\(\delta_{ij} = 1\) for \(i = j\) and is zero otherwise
Figure 1.4: The honeycomb lattice has many possible unit cells (red) and primitive cells (green). While generic unit cells may contain any number of lattice points, each primitive cell contains only one lattice point. Widely-used conventional unit cells (blue) contain two lattice points.

The reciprocal lattice with reciprocal primitive vectors

\[
\vec{b}_1 = 2\pi \frac{\vec{a}_2 \times \hat{z}}{\hat{z} \cdot (\vec{a}_1 \times \vec{a}_2)} \quad \text{and} \\
\vec{b}_2 = 2\pi \frac{\hat{z} \times \vec{a}_1}{\hat{z} \cdot (\vec{a}_1 \times \vec{a}_2)}.
\]

Or, for \( \vec{R}_n = n_1\vec{a}_1 + n_2\vec{a}_2 \), \( \vec{K}_m = m_1\vec{b}_1 + m_2\vec{b}_2 \).

The primitive unit cell of the reciprocal lattice is known as the Brillouin zone. It contains all unique \( k \) points representing the periodicity of waves allowed in the original structure. Certain high-symmetry points exist in a Brillouin zone depending on the lattice’s geometry\(^8\) (Figure 1.5).

Recalling the plane wave traveling through the direct lattice, the relationship between its frequency and wavevector is known as a dispersion relation. We can therefore plot the full dispersion of the wave by plotting its frequency at all points in the Brillouin zone. We only need to span the Brillouin zone because it is periodic. For example, if the

\(^7\) Again, we can see how this formula may be applied to find the reciprocal primitive vectors in three dimensions, though I will not outline that here.

\(^8\) \( \Gamma \) always refers to the center of the Brillouin zone regardless of its geometry.
reciprocal lattice periodicity is $b = 2\pi/a$, the Brillouin zone centered around wavevector $k = 0$ will span $-\pi/a \leq k \leq \pi/a$, and $k \to k + 2\pi/a$.

The dispersion relation for an electron is analogously between its energy and momentum. The plotted dispersion for an electron is called a band structure.

1.3 Tight-Binding

To describe electron dynamics, we use the Hamiltonian operator $\hat{H}$, which encodes the total energy of the system. When detailing the Hamiltonian for electrons in a crystalline solid, we may be tempted to incorporate each electron’s interactions with every other electron; the resulting Hamiltonian would however be non-linear and almost certainly impossible to solve exactly. Instead, we consider the non-interacting Hamiltonian of a crystalline solid, simplified to the form

$$\hat{H} = \hat{T} + \hat{V},$$

where $\hat{T}$ is the kinetic energy operator and $\hat{V}$ is the potential energy operator. $\hat{V}$ arises from the ionic cores of atoms in the solid lattice and is therefore periodic in nature:

$$\hat{V} = V(\vec{r}) = \sum_n V_0(\vec{r} - \vec{r}_n)$$

There are two conventional techniques for solving this Hamiltonian to describe electron dynamics through a particular solid. First is the nearly-free electron model, which takes $\hat{V}$ to be weak compared to $\hat{T}$. Qualitatively, under the nearly-free model, electrons are free to move throughout a solid with small perturbations due to the atomic lattice’s periodic potential.

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9This is known as the “first Brillouin zone”.

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Figure 1.5: A two-dimensional hexagonal Brillouin zone has high-symmetry $k$ points $\Gamma$ (green, at the center of the Brillouin zone), $K$ (blue, at the vertices of the hexagon, with adjacent $K$ points differentiated as $K$ and $K'$), and $M$ (red, at the midpoint of the hexagon edges).
The second technique, called the tight-binding model, takes the opposite limit of the nearly-free model, considering $\hat{T}$ to be weak compared to $\hat{V}$. We then consider the single-electron bound states for each atom. With the approximation that the width of these bound states is small compared to the distance between lattice sites, the continuum model becomes a discrete one. That is, under the tight-binding model, electrons are bound to their atoms, rather than existing freely through the material.

The degree to which on-site wavefunctions between neighboring atoms in the lattice overlap determines electron coupling between sites. If two lattice sites are coupled, an electron has the opportunity to “hop” between sites. Electrons therefore travel through a material by hopping between neighboring lattice sites. From the perspective of one lattice site, the closest surrounding site the electron can hop to is called the “nearest neighbor”. Further sites are denoted “next-nearest neighbor”, “next-next nearest neighbor” and so on (Figure 1.6). We will assume there is a uniform distance between nearest neighbors\textsuperscript{10}. The energy needed for an electron to hop to a nearest neighbor is $t$. If we let $i$ and $j$ denote nearest neighbor sites, we can write the tight-binding Hamiltonian as

$$\hat{H}_{i,j} = t,$$

where the hopping rate between nearest neighbor sites is $t^2$. It is more common, however, to examine the tight-binding Hamiltonian in its second quantized form:

$$\hat{H} = t \sum_{i,j} (a_i^\dagger a_j + a_j^\dagger a_i)$$

where $a_i^\dagger$ denotes the creation operator which projects an input state onto the site $i$ and $a_i$ is the annihilation operator, which removes an input state from $i$. In English, the tight-binding Hamiltonian shows the energy needed for an electron to be annihilated from a site $j$ and then created at a site $i$, or vice versa.

\textsuperscript{10}Graphically, a line connecting two lattice points denotes a pair of nearest neighbors (Figure 1.6).
Qualitatively, the tight-binding model allows us to intuitively consider electrons as particles hopping from site to site. Although this approximation may seem too simplistic to be accurate, the tight-binding model provides reasonably accurate solutions for the low-energy portions of electron spectra.
Chapter 2

Motivating Metamaterials

2.1 Two-dimensional Materials and van der Waals Heterostructures

Two-dimensional (2D) materials are crystalline structures comprised of very few atomic layers. This reduced dimensionality typically leads to enhanced quantum effects; 2D materials often display exceptional electronic, optical and magnetic properties [3, 4, 5, 6].

The first successfully isolated single-atom thick 2D material was graphene, formally discovered in 2004 [3]. Since then, the library of 2D materials has grown rapidly, providing a promising platform for studying emergent condensed matter phenomena.

2D materials can be stacked together, providing a wealth of novel quantum phenomena to explore. Individual 2D layers are held together by strong in-plane covalent bonds, while multiple layers in a stack are held together by weak out-of-plane van der Waals forces, making it easy to remove and replace 2D layers from a stack without damaging them [6]. Macroscopic stacks of 2D materials are appropriately called van der Waals (vdW) heterostructures.

Layers within a vdW heterostructure can be twisted relative to one another, providing an additional degree of freedom for the exploration of novel quantum phenomena. Introducing a twist angle between two atomic lattices creates a new combined structure with unique properties. At certain commensurate angles, this combined structure displays a periodic Moiré Pattern which has a larger conventional unit cell\(^1\) containing multiple primitive cells of the original lattices 2.1.

While 2D materials and vdW heterostructures are an exciting platform for discovering emergent condensed matter phenomena, the expanding library of 2D materials and

\(^1\)This larger unit cell is often called a “supercell”.
Figure 2.1: Two honeycomb lattices twisted relative to each another form periodic Moiré patterns at certain commensurate angles. The conventional unit cell of the overall Moiré pattern (red) increases in size as the twist angle ($\theta$) decreases. When $\theta = 0^\circ$, the system is bilayer graphene and its unit cell is the same size as graphene’s. When $\theta = 21.8^\circ$, the twisted bilayer supercell contains 7 graphene unit cells; when $\theta = 9.43^\circ$, the supercell contains 37 graphene unit cells; when $\theta = 5.09^\circ$, the supercell contains 127 graphene unit cells.
increasing number of vdW heterostructure possibilities makes fully exploring their capabilities an arduous task. Aside from the daunting breadth of systems, experimental work with vdW heterostructures can be difficult and expensive. Certain systems are very difficult to examine even theoretically; twisted bilayer graphene, for example, displays interesting effects at very low twist angles [7, 8] that produce very large unit cells difficult to model computationally. Prototyping and perfecting devices utilizing vdW heterostructures is a time-consuming and expensive process.

2.2 Metamaterials

Metamaterials are engineered to exhibit unique properties due to their macroscopic structure rather than their individual components. They are typically created from many elements assembled into a periodic pattern, allowing the composite structure to manipulate certain physical phenomena.

Because metamaterials’ capabilities rely only on their structure, there is great flexibility in their production. So long as the basic periodic structure of a metamaterial is sustained, the size parameters and materials used to build it can be tuned to adjust the resulting physical effects as needed. We can build metamaterials at the nanoscale out of photonic crystals or at the macroscale using a 3D printer, and so on.

Metamaterials are further attractive to work with because they can mimic quantum materials. Although metamaterials have length and energy scales vastly different from quantum materials, they can be macroscopically constructed to exhibit physical phenomena similar to those of nanoscale quantum materials. Already metamaterials have been used to mimic topological insulators, quantum Hall systems, and Weyl semimetals [9, 10, 11].

2.3 Phononic Metamaterials as Quantum Mimics

Phononic metamaterials are engineered to control acoustic waves to produce interesting effects. We can therefore engineer a phononic metamaterial to manipulate the flow of ultrasound so that it mimics the way electrons move through solids, effectively creating a phononic metamaterial quantum mimic.

Electron behavior in solids is governed by the Schrödinger wave equation:

\[ i\hbar \frac{\partial \psi}{\partial t} = -\frac{\hbar^2}{2m} \nabla^2 \psi, \]

which atomic orbital eigenstates that can be mapped onto the atomic lattice to produce an energy band structure.
The acoustic wave equation in a periodic solid is:

\[ B_r(\vec{r}) \left( \nabla \cdot \left[ \frac{1}{\rho_r(\vec{r})} \nabla p \right] \right) = -\frac{\omega^2}{c_1^2} p, \]

where \( p \) is the pressure, \( \rho_r(\vec{r}) = \rho(\vec{r})/\rho_1 \) is the relative mass density of the material, \( B_r(\vec{r}) = B(\vec{r})/B_1 \) is the relative bulk modulus of the material, and \( c_1 = \sqrt{B_1/\rho_1} \) is the speed of sound in the material. The acoustic wave equation has pressure mode solutions that can be mapped onto the metamaterial lattice to produce a frequency band structure.

Although the former is quantum mechanical the latter classical, they can be solved through analogous means [12], implying not only that the atomic orbitals of the quantum material are comparable to the pressure modes of the metamaterial, but also that the band structures of the two materials should display matching features\(^2\).

Viewed through a tight-binding lens, the parameters of this quantum material – phononic metamaterial analogy are as follows: the hopping particle of interest is in the quantum material the electron and in the phononic metamaterial the normal mode of an acoustic vibration, or a phonon\(^3\). Lattice hopping sites are in the quantum material atoms and in the phononic metamaterial some sort of periodic structure capable of hosting a phonon.

Phononic metamaterials present a compelling pathway for studying emergent condensed matter phenomena because they are easier to fabricate and characterize than quantum materials. These quantum mimics can be fabricated macroscopically, and their governing equations are classical and well-understood.

Phononic crystals have been engineered to mimic graphene [14] and more recently, macroscopic phononic metamaterials have been used to mimic graphene and bilayer graphene [15], presenting phononic metamaterials as an attractive platform for modeling 2D materials and vdW heterostructures.

### 2.4 Work Presented

This thesis develops an intuitive model for phononic metamaterial analogs of 2D materials and vdW heterostructures following tight-binding principles. This model can be used to expand our library of 2D metamaterials and vdW metamaterials, providing a compelling platform to rapidly explore and prototype interesting condensed matter phenomena.

First, I describe my “connected-cavity” model for designing 2D metamaterials according to tight-binding principles. I then create phononic metamaterial graphene.

\(^2\)This phononic scheme can further be translated to photonics through simple variable-matching [13]

\(^3\)A phonon is the quantum of vibration, similar to how a photon is the quantum of light [2].
Next, I extend my model to accurately mimic stacked vdW heterostructures and create all standard configurations of phononic metamaterial bilayer graphene.

Finally, I introduce a twist angle to the bilayer system to create phononic metamaterial\textsuperscript{4} twisted bilayer graphene, providing a pathway for achieving a magic-angle phononic flat band.

All work was conducted theoretically using the acoustics module of the commercial finite element analysis software COMSOL Multiphysics.

\textsuperscript{4}I will henceforth drop the “phononic” specification; unless otherwise specified, all metamaterials presented in this thesis are phononic.
Chapter 3

Design Approach and Results

3.1 The Connected Cavity Model

Recall the following requirements of the tight-binding approximation:

1. Particles are bound to lattice sites.

2. The hopping parameter \( t \) is equal between all nearest-neighbor lattice sites.

3. A particle’s hopping ability depends on the coupling between neighboring sites.

A phononic metamaterial mimicking a 2D quantum material through tight-binding principles must also meet these three requirements. I have thus developed what I refer to as the “connected-cavity” metamaterial model, which fulfills the above requirements as follows:

1. Lattice sites are represented in the metamaterial as cavities in a solid that can hold standing acoustic waves; in other words, phonons are bound to these cavity lattice sites.

2. The hopping parameter \( t \) is represented by the distance between cavities. The physical distance between nearest-neighbor cavities in the metamaterial is therefore made equal (see Figure 3.2).

3. Coupling in this metamaterial refers to an acoustic wave’s ability to travel from one cavity to another. Thin channels are introduced between nearest-neighbor sites to activate coupling and effectively guide the wave along allowed hopping paths.

To prevent acoustic waves from leaking out, we simply place thin bounding membranes above and below the metamaterial\(^1\).

\(^1\)The thickness and density of these bounding layers is variable, as are the other properties of metamaterials built using this design.
There are several implications to this design. Firstly and perhaps most noticeably, it is highly intuitive. Even without any understanding of the quantum materials these metamaterials mimic or the tight-binding principles they follow, we can reasonably understand the behavior of acoustic waves within them.

This design also implies great flexibility in fabrication. A connected-cavity metamaterial can be produced macroscopically at a range of sizes. The metamaterial’s scale influences the frequency at which acoustic waves display interesting effects; generally, metamaterials built at smaller length scales will produce band structures with features at higher frequencies, and vice versa. There is also flexibility in the material used to fabricate a connected-cavity metamaterial. At the most abstract level, so long as the material within the connected-cavity network has a density much smaller than the surrounding material’s density, the metamaterial should recognizably manipulate acoustic waves according to tight-binding principles. Intuitively, the greater the difference in densities between the two materials, the more effective the model will be. For simplicity, we would imagine the cavities would contain air and the surrounding material would be some solid\textsuperscript{2}.

Finally, there is room to modify the design as needed to mimic a range of 2D materials. So long as nearest-neighbor cavity sites are equally distant to each other, we can conceivably build any type of lattice as a connected-cavity metamaterial. What’s more, because site coupling is controlled by physical channels connecting cavities, additional unconnected features can be inserted into the metamaterial as needed. For example, we can insert extra cavities into the honeycomb connected-cavity metamaterial without significantly altering its band structure (Figures 3.2 and 3.4). The ability to insert these extra features will prove relevant when coupling multiple metamaterial layers using tight-binding principles\textsuperscript{3}.

### 3.2 Designing Two-dimensional Metamaterials

To create a metamaterial analog of a 2D quantum material, we must first identify the electron hopping patterns within the quantum material in question.

\textsuperscript{2}In all following calculations, metamaterials were built from steel sheets containing connected air cavities, with length scales on the order of millimeters. We could, however, imagine fabricating these metamaterials from 3D-printer high-density polyethylene (HDPE), or from laser-cut acrylic.

\textsuperscript{3}This also has implications for conserving resources during metamaterial fabrication. If we were, for example, 3D-printing these metamaterials, inserting extra holes could save material and reduce printing time.
3.2.1 Graphene

In graphene, carbon atoms form a honeycomb lattice with a two-atom basis. Let the two atoms be $A_1$ and $A_2$. If we restrict electrons to nearest neighbor hopping only, an electron can hop from an $A_1$ atom to a surrounding $A_2$ atom, but not to another $A_1$ atom, and vice versa. If the nearest neighbor hopping energy is $t$, the tight-binding Hamiltonian for graphene is:

$$\hat{H} = \begin{bmatrix} 0 & t \\ t & 0 \end{bmatrix}. \quad (3.1)$$

Graphene’s band structure displays an isolated Dirac crossing at the K point due to its $C_6$ symmetry [3].

To create connected-cavity metamaterial graphene, cavities are arranged in a honeycomb pattern with a two-site basis (2SB). $A_1$ and $A_2$ cavities are coupled with connecting channels. (Figure 3.1).

I modeled this metamaterial as air cavities in steel sandwiched between bounding HDPE layers. The metamaterial has cavity spacing $(a)$, cavity radius $(R)$, channel width $(w)$, and layer thickness $(D)$ on the order of millimeters. The band structure for this metamaterial displays an isolated Dirac-like crossing, effectively behaving like graphene for frequencies near 7.42 kHz (Figure 3.2).

![Figure 3.1: Here we visualize a 2SB honeycomb lattice of connected cavities as the basic design for phononic metamaterial graphene. Channels connect nearest neighbor hopping sites. Nearest neighbor hopping $(t)$ occurs either from an $A_1$ site to an $A_2$ site or vice versa.](image)
Figure 3.2: (a) This unit cell for 2SB metamaterial graphene shows air cavities in purple, steel in opaque gray, and HDPE bounding layers in semi-transparent gray. Here $a = 10 \text{ mm}$, $R = 3.5 \text{ mm}$, $w = R/4 = 0.875 \text{ mm}$, and $D = 1 \text{ mm}$. (b) The metamaterial’s resulting band structure displays an isolated Dirac-like crossing (blue) mimicking graphene.

To demonstrate the stability of this model, I introduced extra unconnected cavities throughout the metamaterial. This change is the equivalent of adding a third site ($A_3$) to the lattice basis. $A_3$ sites in this three-site basis (3SB) metamaterial graphene have zero coupling to the $A_1$ and $A_2$ sites (Figure 3.3). Let the radius of the $A_3$ cavities be $r$.

Figure 3.3: Unconnected-cavity sites ($A_3$) have been added to the original honeycomb lattice of connected cavities. Channels connect nearest neighbor hopping sites (with hopping energy $t$); there is no hopping from $A_1$ or $A_2$ sites to $A_3$ sites.

The tight-binding Hamiltonian for such a material is:

$$\hat{\mathcal{H}} = \begin{bmatrix} 0 & t & 0 \\ t & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}.$$ (3.2)
The band structure for 3SB metamaterial graphene is identical to that of 2SB metamaterial graphene, with the addition of a flat band (shown in gray) (Figure 3.4(b)). The extra band represents an isolated air mode within the extra cavities and does not affect the metamaterial’s graphene-like behavior. Altering the size of the additional cavities will change the frequency of the extra band\(^4\). Our ability to insert unconnected cavities without negatively impacting the metamaterial’s behavior will prove useful when extending the connected-cavity model to bilayer systems.

### 3.3 Designing Stacked van der Waals Metamaterials

Layered 2D materials couple together through the vdW interaction\(^[16]\). We can mimic the vdW interaction in metamaterials by placing a coupling membrane between metamaterial layers. Acoustic waves propagating in either metamaterial layer induce oscillations in the coupling membrane producing the effects of interlayer hopping; the strength $\Delta$ of this interlayer hopping can be tuned by adjusting the density and thickness of the coupling membrane\(^[15]\). To complete the stacked metamaterial, we must identify interlayer hopping patterns.

#### 3.3.1 AB-Stacked Bilayer Graphene

Bilayer graphene is a stacked system of two honeycomb lattices, each with a two-atom basis. Let the atoms in one layer be $A_1$ and $A_2$ and the atoms in the second layer be $B_1$ and $B_2$. We already know the hopping parameters for $A_1$ with $A_2$ and $B_1$ with $B_2$ (Equation 3.1); we must now determine how the layers interact with each other.

Bilayer graphene naturally occurs in the AB stacking configuration in which the two atomic lattices are offset so that $A_2$ overlaps with $B_1$. There is allowed hopping $\Delta$ between the two overlapping atoms. $A_1$ and $B_2$ do not overlap, meaning no hopping is allowed between the two. The resulting Hamiltonian for AB-stacked bilayer graphene is:

$$
\hat{H} = \begin{bmatrix}
0 & t & 0 & 0 \\
t & 0 & \Delta & 0 \\
0 & \Delta & 0 & t \\
0 & 0 & t & 0
\end{bmatrix}.
$$

(Equation 3.3)

Intralayer hopping ($t$) contributes one Dirac cone per graphene layer to the overall band structure. The two Dirac cones then couple through interlayer hopping to produce

\(^4\)If the extra cavities become too large there is a risk of the air within them coupling through the solid material to the connected-cavity system, thereby altering the metamaterial’s graphene-like behavior. In this model, extra cavities with radius $r = 0.75R$ work well.
parabolic “kissing” bands at the K point [17].

To create metamaterial bilayer graphene, we might consider simply placing a coupling membrane between two connected-cavity metamaterial graphene layers. Utilizing 2SB metamaterial graphene (Figure 3.5), however, will not yield the correct band structure. Rather than achieving the expected parabolic “kissing” structure, 2SB AB-stacked metamaterial bilayer graphene’s band structure displays a triple degeneracy at the K point (Figure 3.6(b)).

This occurs because the non-overlapping sites ($A_2$ and $B_1$) were coupling partially to the steel in their respective counterpart layers. That is, oscillations in the air in $A_2$ and $B_1$ sites were propagating through the coupling membrane into solid steel, which then disturbed the acoustic waves traveling through the connected cavities enough to alter the band structure.

To remedy this issue, I built the same bilayer structure instead using 3SB metamaterial graphene (Figure 3.7).

From 3SB metamaterial graphene we learned unconnected cavities do not have interlayer coupling to the surrounding connected-cavity network. By using 3SB metamaterial graphene to make AB-stacked metamaterial bilayer graphene, $A_2$ and $B_1$ sites overlap with $B_3$ and $A_3$ sites respectively; interlayer coupling between these sites is contained and prevented from affecting the rest of the material. The tight-binding Hamiltonian for this structure is

\[
\hat{H} = \begin{bmatrix}
0 & t & 0 & 0 & 0 & 0 \\
t & 0 & 0 & \Delta & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 \\
0 & \Delta & 0 & 0 & t & 0 \\
0 & 0 & 0 & t & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0
\end{bmatrix}.
\]  

The resulting band structure shows the desired parabolic “kissing” bands at the K point, mimicking AB bilayer graphene (Figure 3.8(b)).

There are two additional bands (shown in gray) representing isolated air modes within $A_3$ and $B_3$ sites, respectively. Again, altering the size of the additional cavities will change the frequencies of the extra bands\(^5\).

### 3.3.2 AA-Stacked Bilayer Graphene

AA-stacked bilayer graphene has entirely overlapping atomic layers; there is allowed hopping $\Delta$ between $A_1$ and $B_1$ sites, as well as between $A_2$ and $B_2$ sites. AA-stacked bilayer graphene

\(^5\)There is a balance to be reached between making the extra cavities large enough to contain interlayer coupling but small enough to avoid intralayer coupling. Again, $r = 0.75R$ works well here.
Figure 3.4: (a) The unit cell for 3SB connected-cavity metamaterial graphene has the same properties as the unit cell in Figure 3.2(a). The additional unconnected cavities have radius $r = 0.75R = 2.625$ mm. (b) The metamaterial’s resulting band structure displays an isolated Dirac-like crossing (blue) identical to the one shown in Figure 3.2(b). The extra band (gray) represents an isolated air mode within the extra cavities.

Figure 3.5: Here we visualize the coupling between 2SB honeycomb cavity networks in either layer of AB-stacked metamaterial bilayer graphene. Connective channels within each layer indicate intralayer hopping ($t$); Overlapping sites have interlayer hopping ($\Delta$). $A_2$ and $B_1$ sites overlap; $A_1$ and $B_2$ do not.
Figure 3.6: (a) This unit cell for 2SB AB-stacked metamaterial bilayer graphene shows air cavities in purple and steel in opaque gray. The metamaterial layers and outer HDPE bounding layers have the same properties \((a, R, w, \text{ and } D)\) as in Figure 3.2(a). The center coupling layer is HDPE with thickness \(D\), identical to the outer bounding layers. (b) The metamaterial’s resulting band structure displays abnormal behavior at the K point not matching the expected parabolic “kissing” structure present in AB-stacked bilayer graphene.

bilayer graphene has the tight-binding Hamiltonian

\[
\hat{H} = \begin{bmatrix}
0 & t & \Delta & 0 \\
t & 0 & 0 & \Delta \\
\Delta & 0 & 0 & t \\
0 & \Delta & t & 0
\end{bmatrix}.
\] (3.5)

This stacking configuration produces a band structure with two spaced Dirac crossings at the K point [16].

While AA-stacked bilayer graphene has interesting electronic properties, it is difficult to fabricate and examine, as bilayer graphene tends to relax back to the AB stacking configuration\(^6\) [16].

Conversely, the metamaterial’s connected-cavity sites have rigid positions and do not relax like atoms in a lattice. We can therefore easily create AA-stacked metamaterial bilayer graphene by layering two metamaterial graphene layers around a coupling membrane.

I first created 2SB AA-stacked metamaterial bilayer graphene (Figure 3.9).\(^6\)

\(^6\)We mainly focus on AB and AA because they are periodic and have reasonable unit cells, but there are bilayer graphene stacking configurations beyond just AB and AA. Two layers of graphene can conceivably be stacked in any configuration. All configurations aside from AB are difficult to fabricate due to bilayer graphene’s tendency to relax back to AB stacking.
This structure produced the desired double-Dirac-like crossing at the K point, mimicking AA-stacked bilayer graphene (Figure 3.10(b)).

For completeness, I also created 3SB AA-stacked metamaterial bilayer graphene (Figure 3.11), which has the tight-binding Hamiltonian

$$\hat{H} = \begin{bmatrix}
0 & t & 0 & \Delta & 0 & 0 \\
t & 0 & 0 & 0 & \Delta & 0 \\
0 & 0 & 0 & 0 & 0 & 0 \\
\Delta & 0 & 0 & 0 & t & 0 \\
0 & \Delta & 0 & t & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0
\end{bmatrix}.$$  \hspace{1cm} (3.6)

This structure also produced the desired double-Dirac-like crossing at the K point (Figure 3.12(b)). The two extra bands (shown in gray) again represent isolated air modes within $A_3$ and $B_3$ sites, respectively, and do not affect the graphene-like properties of the metamaterial.

This second model worked because the extra unconnected cavities ($A_3$ and $B_3$) had interlayer coupling to each other but no intralayer coupling to their respective layer’s connected-cavity network. Our ability to create an accurate mimic of AA-stacked bilayer graphene with extra cavities will prove useful as we move towards mimicking twisted bilayer graphene.

3.4 Designing Twisted van der Waals Metamaterials

Introducing a twist angle between stacked 2D materials gives rise to novel and sometimes exotic physical phenomena [6, 5]. If we can extend the vdW metamaterials model to also
mimic twisted vdW heterostructures, we can confidently present the model as a means for rapidly exploring phenomena in their quantum counterparts.

For eased computation, we often consider twisted systems at commensurate twist angles that create periodic Moiré patterns in the heterostructure (Figure 2.1).

### 3.4.1 Twisted Bilayer Graphene

Twisted bilayer graphene (TBG) has interesting band structure features, especially at lower twist angles. As the twist angle decreases, the two Dirac crossings of bilayer graphene become flatter [18, 19, 20]. Flattening bands correspond to electrons moving at slower and slower speeds. This is significant because slowed electrons are more likely to interact within a material, resulting in unconventional and exotic behavior like high-temperature superconductivity [7, 8].

Let us consider the geometry of bilayer graphene as a twist angle ($\theta$) is introduced between layers. AB and AA are the two bilayer graphene stacking configurations that produce a periodic heterostructure. While we might consider both AA and AB-stacked bilayer graphene as untwisted models, each configuration is actually a twisted version of the other; twisting one layer in AB-stacked bilayer graphene by 60° produces AA-stacked bilayer graphene, and vice versa. Beginning twist calculations from the AB or AA stacking configuration will produce parabolic touching or linear crossing behavior at the K point, respectively. Aside from K point behavior, TBG band structures remain
Figure 3.9: Here we visualize the coupling between 2SB honeycomb cavity networks in either layer of AA-stacked metamaterial bilayer graphene. Connective channels within each layer indicate intralayer hopping \((t)\); Overlapping sites have interlayer hopping \((\Delta)\). Each cavity network overlaps entirely with the other; that is, \(A_1\) sites overlap with \(B_1\) sites and \(A_2\) with \(B_2\).

largely the same between AB and AA starting configurations [20].

However, the choice of metamaterial bilayer graphene to twist is not arbitrary. As \(\theta\) decreases, the TBG supercell size increases, and zones of AB and AA stacking configurations form within the overall structure (Figure 3.13). I was previously able to mimic AA-stacked bilayer graphene with both the 2SB and 3SB metamaterial graphene, but I could only mimic AB-stacked bilayer graphene with 3SB metamaterial graphene. I therefore began twisting from 3SB AA-stacked metamaterial bilayer graphene (Figures 3.11 and 3.12) to ensure AB-stacked regions in the heterostructure would be accurate.

I built 3SB metamaterial TBG unit cells for commensurate angles 21.8°, 9.43°, 6.01°, and 5.09° (Figure 3.15(a)). Their resulting band structures displayed flattening Dirac-like crossings very closely resembling features found in TBG band structures (Figure 3.15(b)\(^7\)). From \(\theta = 21.8^\circ\) to \(\theta = 5.09^\circ\), the Dirac-like crossings’ bandwidth decreased from 2.13 kHz to 0.06 kHz (Figure 3.14).

The incredible success of this metamaterial TBG model has several implications.

First, the metamaterial bilayer graphene design required no additional changes to be extended to metamaterial TBG. Specifically, the additional unconnected sites in the 3SB metamaterial graphene did not cause unusual behavior once the bilayer system was twisted, meaning regions of the twisted supercell that had neither AA nor AB stacking configurations were not impacted by the presence of extra cavities. Any interlayer coupling to these unconnected cavities was effectively contained and did not cause unwanted intralayer coupling.

\(^7\)Note: the metamaterial TBG band structures span the Brillouin zone differently than previous band structures shown. Here we span the Brillouin zone from K point to K point to more easily visualize how the Dirac-like crossings flatten at different twist angles.
Figure 3.10: (a) This unit cell for 2SB AA-stacked metamaterial bilayer graphene has the same properties as in Figure 3.6(a), with one metamaterial layer rotated 60° so that both cavity networks entirely overlap. (b) The metamaterial’s resulting band structure displays double-Dirac-like crossings at the K point mimicking AA-stacked bilayer graphene.

Second, this metamaterial TBG closely mimics its quantum counterpart without relaxation effects. As mentioned previously, while atoms in bilayer graphene tend to relax back to an AB stacking configuration, sites in these phononic metamaterials are rigid; this fact allowed us to create AA-stacked metamaterial bilayer graphene just as easily as AB-stacked metamaterial bilayer graphene. However, atoms in TBG also tend to relax back to the AB stacking configuration [21]. In reference to Figure 3.13, relaxation effects in TBG would minimize the circular AA stacking regions and maximize the triangular AB stacking regions. Metamaterial TBG clearly lacks relaxation effects, yet mimics the quantum system very well.

Most important to note is that while metamaterial TBG’s band structures display flattening features closely resembling those present in TBG’s band structures, these features appear at much higher twist angles in the metamaterial system than in the quantum system. The twist angle at which flattening features appear is dependent on interlayer coupling in system; in general, the greater the coupling between the two graphene layers, the higher the twist angle at which we see flatter features. In TBG, coupling is increased by applying pressure to the system, which can be difficult to achieve experimentally [22]. In metamaterial TBG, however, interlayer coupling is controlled by the density and thickness of the coupling membrane [15]. The coupling membrane’s properties can be tweaked easily to pull interesting flat features up to higher twist angles. Higher twist angles result in smaller TBG supercells (see Figures 2.1 and 3.14(a)), which are far easier to
Figure 3.11: Here we visualize the coupling between the 3SB honeycomb cavity networks in either layer of AA-stacked metamaterial bilayer graphene. Connective channels within each layer indicate intralayer hopping ($t$); Overlapping sites have interlayer hopping ($\Delta$). There is overlap between $A_1$ and $B_1$, $A_2$ and $B_2$, and $A_3$ and $B_3$ sites.

computationally model than the larger supercells created from very small twist angles$^8$.

$^8$In fact, the metamaterial TBG band structures presented in this paper each took a few hours (or less) to generate. Compared to the days required to run band structure calculations for quantum systems, a few hours is a massive improvement.
Figure 3.12: (a) This unit cell for 3SB AA-stacked metamaterial bilayer graphene has the same properties as in Figure 3.8(a), with one metamaterial layer rotated 60° so that both cavity networks entirely overlap. (b) The metamaterial's resulting band structure retains the expected double-Dirac-like crossings at the K point seen in Figure 3.10(b). The extra two bands (gray) represent isolated air modes in $A_3$ and $B_3$ sites, respectively.
Figure 3.13: TBG has regions of AB stacking (blue triangles) and AA stacking (red circles) connected by transition regions that have neither AA nor AB stacking.

Figure 3.14: The Dirac-like crossings (blue) originally present in 3SB AA-stacked metamaterial bilayer graphene become flatter as $\theta$ decreases. Each band structure spans a smaller frequency range than the previous. Extra bands (gray) represent various other modes in the TBG heterostructure and are not relevant to the flattening bands.
Figure 3.15: (a) The metamaterial TBG supercells grow in size as $\theta$ decreases. Each supercell is composed of two 3SB metamaterial graphene layers (Figure 3.4) with an intervening HDPE coupling membrane, all sandwiched between two outer HDPE bounding membranes. As in all previously shown models, $a = 10$ mm, $R = 3.5$ mm, $r = 2.625$ mm, $w = 0.875$ mm, and $D = 1$ mm. (b) The band structures from Figure 3.14 are shown here plotted on the same frequency scale to visualize the flattening effect. Data is unavailable for the gray regions in the latter three band structures, but would contain additional extra bands (gray) irrelevant to the flattening Dirac-like crossings (blue).
Chapter 4

Conclusion and Future Works

4.1 Expanding the Library of Two-dimensional Meta-
materials

Obviously, to use vdW metamaterials to rapidly prototype and screen their quantum
counterparts, the library of 2D metamaterials must be expanded. Using the connected-
cavity model presented here, we can carry out that expansion by developing new 2D
metamaterials, which can then be stacked and twisted to create new vdW metamaterials.

The first additional 2D metamaterial we may consider designing is metamaterial
hexagonal boron nitride (hBN). It shares the overall honeycomb geometry of graphene,
but is actually composed of two triangular lattices of boron and nitrogen, respectively.
We might also consider it a 2SB honeycomb lattice where \( A_1 \) atoms are boron and \( A_2 \)
atoms are nitrogen. Because hBN does not have perfect \( C_6 \) symmetry, the Dirac crossing
is gapped out at the K point [23].

We might consider creating metamaterial hBN by using the same connected-cavity
lattice as metamaterial graphene but with altered cavity radii corresponding to boron-like
and nitrogen-like sites.

4.2 Magic-Angle Metamaterial Twisted Bilayer Graphene

TBG has been shown to superconduct at certain “magic” angles that flatten the Dirac
crossings completely [19, 20, 7, 8]. As a result, TBG has further promoted vdW hetero-
structures as a compelling platform for studying poorly understood correlated behav-
ior.

Already my results for metamaterial TBG reflect the ability to adjust the twist angle
at which we see flatter features. Although COMSOL metamaterial TBG simulations are
restricted to commensurate angles, we can tweak the coupling membrane to achieve a flat band at a commensurate angle, effectively turning it into a magic angle. We can further alter the coupling membrane to achieve flat bands at relatively high commensurate angles for greater computational ease.

Creating metamaterial magic-angle TBG would not only give us a platform to study its quantum counterpart, but also yield interesting and useful phononics. A phononic flat band is equivalent to a localized vibrational wave or “frozen” sound. Metamaterials with phononic flat bands can be interesting platforms for non-linear acoustics and may aid the field of super-resolution imaging.

4.3 Flat Band Metamaterials

While this thesis explored the potential to translate interesting quantum physics to the phononic realm for the purposes of exploration, we can also use metamaterials to translate interesting phononics to the quantum realm. By taking advantage of the tight-binding-like connected-cavity model, we can design 2D metamaterials that produced flat bands simply based on their cavity lattice geometry.

For example, the kagome lattice displays a flat band in its band structure due to destructive interference in electron hopping that confines electrons to certain sites [24]. We can create a connected-cavity metamaterial build with a kagome lattice of cavities to create a phononic flat band. We can further explore how tweaking the lattice geometry can further flatten this band, resulting in better phonon localization. Such an exploration could provide the blueprints for an eventual flat band quantum material.
References


