Nodal Surfaces in Photoemission from Twisted Bilayer Graphene

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(Dated: May 10, 2014)

Selection rules and interference effects in angle resolved photoemission spectra from twisted graphene bilayers are studied within a long wavelength theory for the electronic structure. Using a generic model for the interlayer coupling, we identify features in the calculated ARPES momentum distributions that are controlled by the singularities and topological character of its long wavelength spectrum. We distinguish spectral features that are controlled by single-layer singularities in the spectrum, their modification by gauge potentials in each layer generated by the interlayer coupling, and new energy-dependent interference effects that directly probe the interlayer coherence. The results demonstrate how the energy- and polarization-dependence of ARPES spectra can be used to characterize the interlayer coupling in twisted bilayer graphenes.

PACS numbers: 73.22.Pr, 78.67.Wj, 79.60.-i

I. INTRODUCTION

Angle resolved photoemission spectroscopy (ARPES) is a well developed tool for mapping out electronic bands in solids. Recent applications to single layer and few-layer graphenes have demonstrated that, in addition to obtaining the energies and momenta of the quasiparticle bands with high resolution, the measured variations of the photoemission intensities also encode information about the phase structure of the wavefunctions. The matrix elements that couple the initial quasiparticle states to the outgoing free electron states are momentum dependent and produce modulations in the intensities of ARPES momentum distributions at a constant initial state energy. These matrix elements typically contain a coherent superposition of amplitudes for emission from the different sites and layers that are represented in the initial state wavefunctions. These in turn depend on the (conserved) value of the crystal momentum parallel to the surface plane as well as the energy and polarization state of the exciting radiation.

Interpreting these intensity modulations presents a complex problem that has been addressed recently for the case of single layer and some forms of bilayer graphenes. Graphene presents a nearly ideal platform for exploring this phenomena for two reasons: (i) Graphene is a two dimensional material in which there is no dispersion of the quasiparticle bands due to a third (perpendicular) component of the crystal momentum. (ii) The quasiparticle band structure contains point singularities around which the internal phases in its wavefunctions undergo a complete twist. Indeed, interfering amplitudes in ARPES from single layer graphene have been observed and have striking consequences. The photoemission intensity on a constant energy momentum space contour vanishes along a particular direction (labelled the “dark corridor”) where the emission amplitudes from the two sublattices turns out to exactly cancel. The orientations of these dark corridors are rotated for photoemission from states near symmetry-related zone corner points when resolved in the extended Brillouin zone. For AB-stacked bilayer graphene the emission patterns are more complex but they have been analyzed similarly to identify the sign of the dominant interlayer Hamiltonian matrix elements between neighboring sites in the adjacent layer. Theory predicts that single layer graphene with a strong Rashba spin-orbit interactions should have a distinct photoemission signature in its spin-resolved ARPES reflecting the entanglement of its spin and orbital (pseudospin) degrees of freedom.

All of these previous theoretical analyses take as their starting points a (presumed known) model for the low energy electronic Hamiltonian and deduce the consequences for the ARPES momentum distributions as a function of the initial state energy. In this paper we take the complementary point of view and consider the ARPES signatures of a generic Hamiltonian for a graphene bilayer. Our approach is motivated by our interest in better understanding the electronic structure of “twisted” multi-layer graphenes where the symmetry axes of the two layers are rotationally misaligned. It is now widely appreciated that this misalignment has a profound effect on the electronic coherence between neighboring layers though developing a microscopic theory for it has proven to be an elusive goal. Current theories indicate that this problem is very rich since kinematical constraints define different regions of energy and momentum where the two layers can act as “strongly coupled” or “nearly decoupled” even for a single structure. These energy-momentum sectors depend on the misalignment angle in a way that is not yet fully understood. For small angles of misalignment theory predicts that new spectral features and narrow bands emerge at low energy. Existing experiments are providing conflicting information about the nature of the interlayer coupling a difficulty that may arise from physical differences between twisted graphenes that are made by different experimental methods.

Since both intra-layer and interlayer electronic coher-
ence in multilayer graphenes can be identified by their interference signatures in ARPES, we suggest that deducting the coupling from experiment on twisted graphenes rather than attempting to calculate it from microscopic theory is a promising direction. In this paper we follow this approach and present calculations of ARPES intensities using a generic model for a twisted graphene bilayer. The inputs to the model are the Dirac models for two decoupled layers, a momentum offset due to the rotational misalignment, an electrostatic asymmetry (bias) between the layers and a general long wavelength matrix that couples the pseudospin amplitudes on neighboring layers. From this starting point we calculate the electron momentum distributions at fixed energy and the ARPES distributions which are weighted by the momentum dependent photoemission probabilities for various polarization states of the exciting radiation. A careful analysis of the ARPES momentum distributions shows that the “dark corridor” known for single layer graphene changes its shape for twisted graphene and allows one to discriminate between various models for the coupling between layers. We present calculations of the ARPES intensities that illustrate this effect and a symmetry analysis of the relevant matrix that allows us to interpret the spectra. The results demonstrate how experimental study of the ARPES spectra as a function of the energy and polarization of the exciting radiation can be used to fully characterize the interlayer coherence in these multilayer structures.

In this paper we first briefly review the long wavelength description of twisted bilayer graphene with a theory that is appropriate to small rotation angles and in Section II we derive an expression for the photoemission transition matrix elements based on this model. In section III we survey the topological transitions in the band structure that occur in this model as the Hamiltonian parameters are varied. Section IV shows the calculated ARPES momentum distributions showing a rich assortment of interference phenomena which are unique to twisted multilayer graphenes. Section V gives an analysis of the interference effects with in terms of the underlying symmetries of the model. We present an analysis of the interference patterns in terms of its “energy-flattened” nodal surfaces in momentum space which provides a useful diagnostic for various forms of interlayer coherence. A further discussion of the experimental signatures of these predictions is given in Section VI.

II. LONG WAVELENGTH MODEL

A. Small Angle Twisted Bilayers

Our work uses a long wavelength low energy theory of the electronic states in graphene bilayers. In single layer graphene the electrons propagate on a honeycomb lattice with two sublattice sites labelled $A$ and $B$. At low energies there are two electronic bands that touch at $E = 0$ at discrete points located at the Brillouin zone corners $\nu \mathbf{K}$ where $\nu = 1\,(-1)$ denote the $\mathbf{K}(\mathbf{K'})$ points. These bands disperse linearly around the contact points and are described by a Hamiltonian at crystal momentum $\mathbf{k}$ that can be linearized in the difference $\mathbf{q} = \mathbf{k} - \nu \mathbf{K}$

$$\mathcal{H}_S = \hbar v_F \mathbf{\sigma} \cdot (\nu q_x \hat{e}_x + q_y \hat{e}_y)$$ (1)

where $\mathbf{\sigma}$ are Pauli matrices acting in the space of sublattice (pseudospin) degree of freedom.

For a graphene bilayer we adopt a four-component basis with a conventional ordering of its four site-layer degrees of freedom $(A_1, B_1, A_2, B_2)$. When the symmetry axes of the two layers are aligned and the interlayer coupling is set to zero the Hamiltonian for the two layers is simply a doubled version of the single layer Hamiltonian and can be written in the form

$$\mathcal{H}_B = \mathcal{H}_S \tau_0$$ (2)

where $\mathbf{\tau}$ are Pauli matrices acting on the layer degree of freedom. In $AB$ stacked graphene the symmetry axes of the two layers are aligned and the two layers are coupled through a hybridization of amplitudes on different sublattice sites in the two layers e.g. $(A_1, B_2)$ as described by the bilayer Hamiltonian

$$\mathcal{H}_{\text{Bernal}} = \mathcal{H}_S \tau_0 + \frac{\gamma_1}{2} (\sigma_x \tau_x - \sigma_y \tau_y)$$ (3)

where $\gamma_1$ is the strength of the interlayer coupling.

In this work we are concerned with the situation where the symmetry axes of the two layers are rotated by a relative angle $\theta$. It is convenient to regard each of the layers as rotated by angles $\pm \theta/2$ with respect to a common reference structure in which the $\mathbf{K}$ point is oriented along the $x$ direction as shown in Figure 1. With this convention the $\mathbf{K}$ points in the two layers shift by momenta $\Delta \mathbf{K}_\pm$ which are, in complex vector notation

$$\Delta \mathbf{K}_\pm = \left( e^{\pm i\theta/2} - 1 \right) \mathbf{K}$$ (4)

The components of $\mathbf{q}$ are resolved in the original unrotated frame so that the effect of the rotation is to induce both a shift of origin and the phases of the complex momenta appearing in $\mathcal{H}_B$ according to the replacement rule

$$q_x + iq_y \mapsto q'_x, \pm + iq'_y, \pm$$

$$q'_x, \pm + iq'_y, \pm = \left( q_x + iq_y - \left( e^{\pm i\theta/2} - 1 \right) \mathbf{K} \right) e^{\mp i\theta/2}$$ (5)

Expressed in a common frame, the offset single-layer Dirac operators of Eqn. (1) are

$$\mathcal{H}_\pm = \hbar v_F \mathbf{\sigma} \cdot (\nu q'_{x,\pm} \hat{e}_x + q'_{y,\pm} \hat{e}_y)$$ (6)
dospins in each layer. The smoothest periodic matrix-
and local $2\times 2$ symmetries. In a periodic supercell with its
be further simplified by exploiting lattice and rotational
where $\hat{A}$
In this situation
When
When the rotation angle $\theta$ is small the interlayer coupling
across a twisted bilayer admits a description analogous to
the coefficients $\hat{A}$
layer off diagonal blocks

$$\mathcal{H}_{\text{lattice}} = \begin{pmatrix} \mathcal{H}_1 & \mathcal{T}(\vec{r}) \\ \mathcal{T}^T(\vec{r}) & \mathcal{H}_2 \end{pmatrix}$$

(7)

When $\theta$ is small the registry between layers is modulated
in a supercell evolving smoothly through local zones with
$(A_1B_2)$ stacking, $(B_1A_2)$ stacking and $(A_1A_2)$ stacking.
In this situation $\mathcal{T}(\vec{r})$ in Eqn. (7) is a smooth, periodic
and local $2 \times 2$ matrix function of $\vec{r}$ acting on the pseu-
dospins in each layer. The smoothest periodic matrix-
valued function satisfying these constraints is

$$\mathcal{T}(\vec{r}) = \hat{t}_0 + \sum_{n=1}^{6} \hat{t}_n e^{i\vec{G}_n \cdot \vec{r}}$$

(8)

where $\hat{t}_n (n = 0, 6)$ are $2 \times 2$ matrix-valued constants and
$\vec{G}_n (n = 1, 6)$ are vectors in the first star of superlattice
reciprocal lattice vectors. The matrix coefficients $\hat{t}_n$ can
be further simplified by exploiting lattice and rotational
symmetries. In a periodic supercell with its $AB$, $BA$ and
$AA$ zones centered at positions $\vec{r}_1$, $\vec{r}_2$ and $\vec{r}_3$, respectively,
the coefficients $\hat{t}_n$ for the three even elements of $\vec{G}_n$ are

$$\hat{t}_{n,\text{even}} = t_0 \begin{pmatrix} e^{-i\vec{G}_n \cdot \vec{r}_1} & e^{-i\vec{G}_n \cdot \vec{r}_2} \\ e^{-i\vec{G}_n \cdot \vec{r}_3} & e^{-i\vec{G}_n \cdot \vec{r}_4} \end{pmatrix} = t_0 \begin{pmatrix} z & 1 \\ z^* & z \end{pmatrix}$$

(9)

where $z = \exp(2\pi i/3)$ and $\hat{t}_{n,\text{odd}} = \hat{t}_{n,\text{even}}$. Sublattice
symmetry requires that the constant matrix $\hat{t}_0$ has the form

$$\hat{t}_0 = \begin{pmatrix} c_{AA} & c_{AB} \\ c_{BA} & c_{BB} \end{pmatrix}$$

(10)

In the small $\theta$ limit, the interlayer coupling is described
by three real parameters $c_{AA} = c_{BB}, c_{AB} = c_{BA}$ and
$t_0$. One can choose these parameters by specifying the
direct ($\gamma_1$) and skew ($\gamma_3$, $\gamma_4$) interlayer hopping parameters
in the $AB$ registered zones of a twisted bilayer. Table I makes this translation by giving the constants in the
modulated interlayer hopping model of a twisted bilayer
in terms of the three interlayer intersite amplitudes. Note
that if the interlayer hopping amplitudes is long range,
i.e. if it varies slowly on the scale of the graphene primitive
cell, then $t_0$ is small compared to the constant term.
When the skew hopping terms are isotropic ($\gamma_3 = \gamma_4$) the
diagonal and off diagonal terms in the constant matrix
have the same amplitude ($c_{AA} = c_{AB}$).

As examples, Table I evaluates these constants for two
models. In Model I a direct interlayer term $\gamma_1$ is re-
tained and the skew hopping terms are arbitrarily set to
zero. Even in this extremely short range model one finds
that the constant terms dominate the spatially modu-
lated part of the interlayer Hamiltonian. In Model II
the constants are fit to the Slonczewski-Weiss-McClure
(SWMc) parameterization of the interlayer amplitudes in
Bernal graphite retaining its skew hopping terms, and
as expected the amplitude of the modulated term is even
weaker. Note also that the SWMc parameterization also
shows a strong asymmetry between $\gamma_3$ and $\gamma_4$. These
two amplitudes describe interlayer hopping processes at
the same range but in different crystallographic direc-
tions with respect to the graphene symmetry axes. This
asymmetry, deduced from fitting experimental data, is
not contained in two-center tight binding theories that
posit an isotropic interlayer hopping model. The asym-
metry arises from a crystal field anisotropy in the Wan-
nier states appropriate to single layer graphene.

The constant matrix $\hat{t}_0$ is the supercell-average of the
interlayer hopping operator $\hat{A}$. One can distinguish three
different kinds of behavior depending on whether its off
diagonal elements are dominant, its diagonal elements are
dominant or they are equal. The SWMc parameterization
realizes the first type of behavior since, as shown in Table I, it is dominated by its site off-diagonal parts: $c_{AB} \gg c_{AA}$. Alternatively, any two center tight bind-
ing model that assumes an isotropic interlayer hopping
model is a member of the last family which describes
a different type of interlayer coherence as we show ex-

\begin{table}[h]
\centering
\begin{tabular}{|c|c|c|}
\hline
Coefficient & Parameterization & I & II \\
\hline
$t_0$ & $(\gamma_1-\gamma_3)/9$ & 43.3 & 8.3 \\
$c_{AA}$ & $\gamma_4 + (\gamma_1-\gamma_3)/3$ & 130.0 & 69.0 \\
$c_{AB}$ & $(\gamma_1+2\gamma_3)/3$ & 130.0 & 340.0 \\
\hline
\end{tabular}
\caption{Parameters (meV units) in interlayer hopping Hamiltonian for a twisted bilayer fitted to the interlayer intersite amplitudes $\gamma_1$, $\gamma_3$ and $\gamma_4$ in its Bernal stacked zones. Model I: $\gamma_1 = 390$ meV, $\gamma_3 = 315$ meV and $\gamma_4 = 44$ meV.
}
\end{table}
Eqn. (11) describing emission from states near a single valley is obtained by substituting the related Hamiltonian for a free particle state whose energy \( \nu \) is conserved, laterally phase matching the outgoing state with projection of the original lattice Hamiltonian for \( \nu = -1 \) is obtained by the substitution \( \sigma \to -\sigma^* \). The Hamiltonian Eqn. (11) is the starting point for the calculations we present in this paper. It is a function of the rotation angle \( \theta \), the electrostatic potential difference \( V \) and coupling coefficients \( \Gamma \). Choices of \( \Gamma \) allow us to explore the space of symmetry allowed interlayer coupling models.

Note that the projection of the original lattice Hamiltonian onto the Dirac basis changes the phases of the interlayer terms because of the oscillation of the pseudospin-layer orbital space for each of the two orthogonal polarizations. These operators are independent of momentum \( q \) but depend on the rotation angles \( \pm \theta/2 \).

The matrix elements for photoexcitation with photon polarization \( \hat{e} \) are

\[
\mathcal{M}(p, q) = \langle \psi_p^\rangle |\hat{e} \cdot \hat{\sigma}|\psi_q \rangle
\]

To evaluate Eqn. (14) we project the outgoing plane wave state \( \psi_p^\rangle \) onto the same basis used to represent the initial state \( \psi_q \). Crucially, this projection introduces a phase difference \( \phi = p \cdot \hat{d}/\hbar \) between plane wave amplitudes on the two layers separated by a vertical distance \( d \). For photoemission near the graphene Brillouin zone corner, by using excitation energies in the range 30-100 eV, \( \phi \) can be varied over a range of approximately \( 4\pi \). Note that the parallel component of the momentum \( p_\parallel \) is conserved, laterally phase matching the outgoing state to the initial Bloch state. Therefore, under typical experimental conditions the analogous lateral interference effects are small and nearly energy independent. (They would be exactly zero for perpendicular emission.) These small effects are not included in our calculations. Thus, written in the orbital-layer basis, the final state appearing in Eqn. (14) in our model is

\[
\psi_p^\rangle = \frac{1}{2} \begin{pmatrix} 1 \\ e^{i\phi} \\ e^{i\phi} \end{pmatrix} e^{ip \cdot r}
\]

The matrix element in Eqn. (14) is the inner product of the operator Eqn. (13) between an occupied eigenstate of Eqn. (11) and the plane wave final state given in Eqn. (15).

Our analysis presents three different momentum distributions for interpretation of the ARPES intensities. The first is a map of the spectral function at \( A(q, E_i) \) as a function of initial state energy \( E_i \) unweighted by the transition matrix elements

\[
A(q, E_i) = \sum_n \delta(E_n(q) - E_i)
\]
summed over occupied bands \( n \) of Eqn. (11). The second is a map of a momentum distribution of the simulated ARPES intensities \( I(q, E_i) \) which includes the state dependent transition matrix elements

\[
I(q, E_i) = \sum_n |M_n(p, q)|^2 \delta(E_n(q) - E_i)
\]  

As noted above the matrix elements \( |M(p, q)|^2 \) also depend on the polarization state of the incident light. The third is a map that “flattens” the energy resolution of Eqn. (17), giving instead for each of the occupied bands the distribution of just its momentum- and polarization-dependent squared matrix elements

\[
P_n(q) = |M_n(p, q)|^2
\]  

The summand in Eqn. (17) is the product of the spectral functions \( \delta(E_0 - E_1) \) that encode the phase information. Maps of the unconstrained distributions \( P_n(q) \) are useful for exposing the quantum geometric origins of the interference patterns that are accessible in the ARPES intensities. Experimentally one can determine the nodal structure of Eqn. (18) by the evolution of a series of spectra that sweep \( E_i \) through the occupied bands.

### III. TOPOLOGICAL TRANSITIONS

The spectrum of the Hamiltonian in Eqn. (11) exhibits topological transitions as a function of its interlayer coupling parameters \( \Gamma_1, \Gamma_2 \) as shown in Figure 2. In the literature on twisted graphenes a frequently used model treats the case \( \Gamma_1 = \Gamma_2 \) where the long wavelength coupling generates a scalar (i.e. sublattice diagonal) interlayer coupling matrix. In that analysis one finds that the interlayer coupling generates a gauge potential in each layer that displaces the two Dirac nodes towards each other as the rotation angle is increased. Decreasing the rotation angle increases the effective coupling strength \( \Gamma_1 \), i.e. the supercell averaged interlayer coupling is dominated by its sublattice off diagonal terms. An extreme version of this situation one can examine the case \( \Gamma_1 \neq 0, \Gamma_2 = 0 \). If the coupling strength \( \Gamma_1 \) is weak this model shares many features of the isotropic model, with a slightly lower symmetry as shown in the middle panel of Figure 2. In our earlier work\(^{12}\) we noted that after a gauge transformation this theory can be cast into a form where it describes two momentum-offset Dirac cones with opposite helicity coupled by a scalar (i.e. sublattice diagonal) interlayer coupling matrix. In that analysis one finds that the interlayer coupling generates a gauge potential in each layer that displaces the two Dirac nodes towards each other as the rotation angle is decreased. Decreasing the rotation angle increases the effective coupling strength \( \Gamma_1 \). Ultimately, at a critical rotation angle the nodes merge and annihilate as shown in Figure 3. For still smaller angles two new point singularities emerge describing two new Dirac modes of a strongly interlayer-coupled limit.

It is important to appreciate that the momentum difference between the Dirac nodes is not determined purely geometrically by the rotation angle, but instead it is generally changed by the strength and symmetry of the coupling between the layers. Also note that in the crossover regime the system has a spectrum reminiscent of \( AB \) stacked graphene, albeit on a reduced energy scale. The presence of the singularities at \( E = 0 \) is symmetry protected and, as is the case in \( AB \)-stacked graphene, cannot be removed unless the layer symmetry is lifted.

In the opposite limit \( \Gamma_1 = 0, \Gamma_2 \neq 0 \) one finds two low energy features that have the same helicity and repel each other in momentum space so that no such \( E = 0 \) merger occur regardless of the coupling strength \( \Gamma_2 \). Instead, at finite energy the two Dirac cones cross to produce new symmetry protected points which are second
FIG. 2: Energy surfaces in the four band model for a twisted graphene bilayer for three different models for the long wavelength interlayer coupling matrix: (left panel) Model I with $\hat{V}_{\text{int}} \propto (\mathbbm{i} + \sigma_x)$, (middle panel) ($\Gamma_1 = \Gamma_2 = 0.05$) Model II with $\hat{V}_{\text{int}} \propto \sigma_x$ in the weak coupling regime ($\Gamma_1 = 0.13, \Gamma_2 = 0$), (right panel) Model III with $\hat{V}_{\text{int}} \propto \mathbbm{1}$ ($\Gamma_1 = 0, \Gamma_2 = 0.15$). The blue dots denote the positions of the Dirac points of a pair of twisted but decoupled sheets. The actual Dirac points at $E = 0$ are displaced from these points in Models II and III due to gauge potentials generated by the interlayer coupling.

FIG. 3: Energy surfaces showing a transition from weak to strong coupling in interaction Model II with $\hbar v_F \Delta K = 0.15$. As the coupling parameter $\Gamma_1$ is increased the renormalized Dirac points evolve from weak coupling (left panel: $\Gamma_1 = 0.13, \Gamma_2 = 0$) and merge at a critical point (central panel: (left panel: $\Gamma_1 = 0.15, \Gamma_2 = 0$)) and produce two new Dirac points in the strong coupling phase of the model (right panel: (left panel: $\Gamma_1 = 0.20, \Gamma_2 = 0$)). The blue dots denote the Dirac points for a pair of twisted by decoupled layers.

generation Dirac singularities near $\pm \Delta$ as shown in the right hand panel of Figure 2. Despite the appearance of a symmetry-allowed band crossing at the midpoints between the $E = 0$ Dirac nodes, the system retains a saddle point structure where the density of states is enhanced in an extended region of momentum space that is laterally displaced away from the position of the crossing point. The low energy features of this type of coupling are quite distinct from those of Model I and II. Notably, a perturbative velocity renormalization of the low energy Dirac nodes is symmetry forbidden for this class of interaction models. This can provide an explanation for the absence of a velocity renormalization inferred from Landau level spectroscopy\textsuperscript{22} and from ARPES\textsuperscript{21}, both of which find a Dirac velocity for twisted BLG that is the same as the velocity for single layer graphene within experimental error. It also provides a interpretation for ARPES experiments that clearly show a crossing of the Dirac cones cross instead of a hybridization induced anticrossing even in the limit of low rotation angles\textsuperscript{21}.

The spectra displayed in Figure 2 and 3 show a distinct low energy topology controlled by the symmetry of the long wavelength interlayer coupling matrix. The singular points in these spectra have a striking signature in the simulated ARPES momentum distributions as discussed below. Additionally there are special lines in momentum space where ARPES selection rules are operative and these can be used to discriminate between these models and even to directly identify the momentum shifts of their Dirac features due to the gauge coupling from the interlayer potential\textsuperscript{18,25}. The details of this analysis are presented in the following sections.
IV. ARPES MOMENTUM DISTRIBUTIONS

In this section we discuss the constant energy momentum distributions calculated for the three limiting models discussed in Section III, presented both as unweighted distributions and as distributions that are weighted by their photoemission transition probabilities.

The top panel of Figure 4 shows the unweighted momentum distribution for the fully symmetric model with \( \Gamma_1 = \Gamma_2 = \Gamma \) where the interlayer matrix has the form \( \hat{V}_{\text{int}} = \Gamma(I + \sigma_1) \) (Model I). At energies well below the interlayer scale \( -\hbar v_F \Delta K \) the constant energy contours are self intersecting loops. The inner loop maps out a constant energy contour on the lowest valence band and as the energy is increased (i.e. as it is made less negative) this inner loop collapses to a point and vanishes at the extremum of the lowest band. The remaining single surface develops a “bean” shape that encircles both Dirac points highlighted by the bold dots. At still higher energy this remaining contour fissions at energy of the saddle point producing two new closed curves that separately encircle the two Dirac points. At positive energy these constant energy contours expand and reconnect at a positive saddle point energy to form a single surface. At still higher energy these surface “folds” contacting at an energy corresponding to the minimum of its highest energy band leading again to a constant energy surface in the form of a self intersecting loop. In the presence of an interlayer bias (shown in the far right panel) the surfaces maintain the same topology but are inflated/deflated around the two Dirac points in response to the layer symmetry breaking potential.

The lower two panels of Figure 4 shows these distributions at the same sampling energies but weighted by the squared modulus of their transition matrix elements for photoemission using light that is \( x \) polarized (top row) and \( y \) polarized (bottom row). For either polarization we observe a modulation of the simulated emission intensities the emergence of a pattern of null points where the probability for photoemission vanishes. A symmetry analysis of these nodal surfaces is presented in Section V. A comparison of these density plots shows that the intensity patterns undergo a complex evolution as a function of energy with the nodal points undergoing pairwise creation and annihilation on constant energy surfaces in each band at critical values of the initial state energy. As shown in the rightmost two panels, these null points are robust features of the simulated ARPES momentum distributions and occur even when the layer symmetry is broken by an interlayer potential.

The top panel of Figure 5 shows the unweighted momentum distributions using the interlayer coupling model \( \hat{V}_{\text{int}} = \Gamma_1 \sigma_x \) (Model II). This describes an interlayer Hamiltonian that is dominated by its hopping terms that connect different sublattices, a situation that is very nearly realized by the SWMc parameterization. This breaks the symmetry of Model I, so that there are two separate (nonintersecting) constant energy surfaces at energies below \( -\hbar v_F \Delta K \). As the energy increases the innermost loop collapses to a point and vanishes (not shown) identifying the band extremum of the lowest energy occupied band. At higher energy the remaining
ring shrinks and fissions at a saddle point energy forming two loops which in turn collapse around the two layer-coupled Dirac points singularities. As noted in Section III these points are generally displaced from the unperturbed Dirac points of the two decoupled layers due to a gauge potential generated by the interlayer coupling. In the weak coupling limit (i.e. for large rotation angle)

The lower two panels of Figure 5 show the probability weighted distributions calculated for Model II for the same representative energies. Again we observe a complex modulation of the emission intensities with null points identified in the constant energy surfaces for both polarizations. The locations of the nodal points in these surfaces exhibit an interesting oscillation as a function of band index and light polarization. Note for example the interchange of the positions of the emission maxima and nodal points in the second and third bands under exchange of the \( x \) and \( y \) polarization.

Figure 6 presents the unweighted momentum distributions obtained from a similar analysis using a coupling model of the form \( V_{\text{int}} = \Gamma_2 \sigma_0 \) (Model III). In this limit the interlayer coupling is dominated by amplitudes connecting sites on the same sublattice. A coupling of this type would occur in a putative \( AA \) stacked geometry which, though apparently excluded by the conventional SWMc parameters of Table I, would provide an explanation for the absence of a Fermi velocity renormalization and the intersection of Dirac bands observed in twisted graphenes grown on SiC (0001). At low energy the constant energy surfaces form two disconnected loops. The signature of coupling in the form of Model III is the collapse of the inner ring at critical energy identifying second generation Dirac point of Figure 2 and its re-emergence as a closed loop immediately above this energy (not shown). As the energy is increased still further this ring bifurcates at a pair of saddle points positioned displaced along the horizontal \( (q_x) \) axes. These two surfaces collapse around the two \( E = 0 \) Dirac nodes in the
model. Here, because of the gauge potential generated by the interlayer coupling the $E = 0$ nodes are repelled from the geometrically defined Dirac points. As one crosses to positive energies this evolution is reversed leading to the reconstruction of two ringlike surfaces for large positive energies.

The lower two panels of Figure 6 show the distributions of Model III weighted by their transition probabilities. Again we observe the complementary positions of the emission maxima and nodal points from a single band for $x$ and $y$ polarized excitation.

V. Symmetry Analysis

The nodal patterns identified in Section IV can be understood by a consideration of the analytic structure of the complex transition amplitudes $M(p, q)$ in Eqn. (14). The key observation is that in particular circumstances identified below this complex transition amplitude degenerates to a real function of $q$. In this situation $\mathcal{M}$ will generically vanish along lines in reciprocal space (where it changes sign) rather than at points (where it wraps its phase). The evolution of the nodal patterns in ARPES distributions then correspond to the evolution of constant energy surfaces through a network of lines along which $M$ is zero. These lines can terminate at singular points in the band structure. In the simplest picture, these points can be identified with the rotationally offset Dirac points. The actual situation is richer however since, as we have seen, these points are generally displaced from the rotationally defined Dirac points by gauge potentials produced by the interlayer coupling and in addition second generation symmetry protected singularities can appear elsewhere the band structure. The identification of these lines, their continuity and termination points in $q$ space provides a powerful probe of the nature of the interlayer coherence.

A. Intralayer Interference of the Transition Amplitudes

The layer diagonal blocks in Eqn. (11) are momentum-displaced and -rotated versions of the Hamiltonian and have the symmetry

$$\mathcal{H}_S = \sigma_x \mathcal{H}_S^* \sigma_x$$

so that the eigenfunctions are equal weight states

$$\chi(q) = \frac{1}{\sqrt{2}} \begin{pmatrix} e^{-i\alpha(q)} \\ \pm e^{i\alpha(q)} \end{pmatrix}$$

The velocity operators of Eqn. (13) are layer block-diagonal and $\theta$-dependent and can be represented compactly

$$v_\mu = \begin{pmatrix} 1 & 0 \\ 0 & e^{i\theta/2} \end{pmatrix} \sigma_\mu \begin{pmatrix} 1 & 0 \\ 0 & e^{-i\theta/2} \end{pmatrix}$$

in the top layer with positive rotation angle $\theta$ where $\sigma_\mu$ is the $\mu$-th $2 \times 2$ Pauli matrix. Reversing the sign of $\theta$ in this expression gives the related velocity operator projected on the bottom layer. Combining Eqns. (20) and (21) we
identify the transition states

\[ \chi^\pm_x(q) = v_x \cdot \chi(q) = \frac{1}{\sqrt{2}} \begin{pmatrix} \pm e^{i(\alpha(q) - \theta/2)} \\ e^{-i(\alpha(q) - \theta/2)} \end{pmatrix} \quad \text{x-polarized} \]

\[ \chi^\pm_y(q) = v_y \cdot \chi(q) = \frac{i}{\sqrt{2}} \begin{pmatrix} \mp e^{i(\alpha(q) - \theta/2)} \\ e^{-i(\alpha(q) - \theta/2)} \end{pmatrix} \quad \text{y-polarized} \]

The photoemission matrix elements obtained by projecting these transition states onto the plane wave final state \([15]\) for \(x\) polarized excitation are

\[ M^+_{x}(p, q) = \frac{1}{\sqrt{2}} \cos(\alpha(q) - \theta/2) \]

\[ M^-_{x}(p, q) = -\frac{i}{\sqrt{2}} \sin(\alpha(q) - \theta/2) \] (22)

and for \(y\)-polarized excitation

\[ M^+_{y}(p, q) = \frac{1}{\sqrt{2}} \sin(\alpha(q) - \theta/2) \]

\[ M^-_{y}(p, q) = \frac{i}{\sqrt{2}} \cos(\alpha(q) - \theta/2) \] (23)

(24)

The structure of these matrix elements has been studied previously in the context of single layer graphene. Note the reversal of the \(q\)-space modulations of the of \(x\) and \(y\) polarized transition amplitudes for the positive and negative energy eigenstates. Importantly the transition amplitudes vanish along lines where \(\alpha(q) = \theta/2\) or \(\alpha(q) = \theta/2 \pm \pi/2\). These nodal lines terminate at points where the layer projected Hamiltonian \(\propto I\) and \(\chi^+\) and \(\chi^-\) are degenerate. Crossing this contact point reverses the assignment of \(\pm\) indices to the upper and lower energy bands, thereby terminating the nodal line in a single band and transferring a nodal line to its partner. This phenomenon is responsible for the one sided “dark corridor” seen in photoemission from single layer graphene\([19]\). Equations (22) and (23) are rotated versions of that result where the new dark corridors are rotated at angles \(\pm \theta/2\) in the two layers due to the rotational misalignment.

Any real combination of the transition amplitudes in Eqs. (22) and (23) describes linearly polarized excitation at some general angle of polarization. We see that its effect is simply to rotate the nodal line for photoemission while preserving an endpoint anchored at a singular point of degeneracy. By contrast elliptically or circularly polarized light requires a phase lag between two orthogonal polarizations of the radiation. In this situation the transition matrix element is necessarily represented by a two dimensional parameter space (it must be a complex scalar function) and it can vanish only at special points in momentum space where two independent transition amplitudes vanish. Such points would be difficult to measure in ARPES since this would require fine tuning the initial state to a single critical energy.

These conclusions hold quite generally for any system described by the symmetry of Eqn. (19) in its layer projected Hamiltonians. For coupled BLG, if the interlayer coupling matrix \(\hat{V}\) respects sublattice symmetry (as occurs in models in the form of Eqn. (11)), one can obtain an effective layer projected theory that respects sublattice symmetry by integrating out one of the layers. Such a system generically exhibits one-sided line nodes in the photoemission matrix elements. Since the eigenfunctions of this layer projected Hamiltonian are equal weight states of the form given in Eqn. (20) its matrix elements are controlled by the evolution of a single scalar parameter \(\alpha(q)\). This requires that its nodal lines are continuous except at point singularities where \(\nabla \alpha\) is undefined and a nodal line can either terminate or switch between bands that are degenerate at the singularity. As noted in Section IV the endpoints of these nodal lines will generally deviate from the rotationally defined points of degeneracy if a residual gauge potential is induced by integrating out the interlayer coupling. Nonetheless the physics responsible for this behavior is essentially the singular structure of the single layer Hamiltonian perturbatively renormalized by interactions between layers.

B. Interlayer Interference of the Transition Amplitudes

Transition matrix elements derived from the Hamiltonian in Eqn. (11) support additional interference features that arise directly from its interlayer coupling amplitudes. These provide a more sensitive probe of the symmetry of the interlayer Hamiltonian in twisted BLG and are analyzed in this section.

Our approach exploits a symmetry in Eqn. (11) along the line \(q_y = 0\) where because of the reversal of the momentum offsets \(\pm \Delta K/2\) the layer-projected Hamiltonians in the top and bottom layers are related

\[ \mathcal{H}_{\text{bottom}} = \mathcal{H}^*_{\text{top}} \] (25)

This symmetry is broken for a general two dimensional momentum \(q\) but it remains a local symmetry for all momenta \(q_x\) along the midline. Along this line the eigenfunctions are equal-weight combinations of the four layer-orbital degrees of freedom. Additionally, since the Hamiltonian commutes with the operator \(R = \tau_x \sigma_x\), these eigenstates can be indexed by their parities \((\pm)\) under \(R\) and take the form

\[ \Psi^\pm(q) = \frac{1}{2} \begin{pmatrix} e^{-i\alpha(q)} \\ e^{i\alpha(q)} \\ \pm e^{i\alpha(q)} \\ \pm e^{-i\alpha(q)} \end{pmatrix} \] (26)

The four-component velocity operators are
and the transition states are obtained by application of these operators to the initial states $\Psi^\pm$, giving for $x$ polarized light:

$$
\Psi^\pm_x = V_x \cdot \Psi^\pm = \frac{1}{2} \begin{pmatrix}
\sigma_\mu \\
\sigma_\mu
\end{pmatrix}
\begin{pmatrix}
e^{i(\alpha - \theta/2)} & e^{-i(\alpha - \theta/2)} \\
e^{-i(\alpha - \theta/2)} & e^{i(\alpha - \theta/2)}
\end{pmatrix}
\begin{pmatrix}1 \\ e^{\mp i\theta/2}
\end{pmatrix}
\begin{pmatrix}1 \\ e^{-i\theta/2}
\end{pmatrix}
\begin{pmatrix}1 \\ e^{i\theta/2}
\end{pmatrix}
\begin{pmatrix}
e^{-i(\alpha - \theta/2)} & e^{i(\alpha - \theta/2)} \\
e^{i(\alpha - \theta/2)} & e^{-i(\alpha - \theta/2)}
\end{pmatrix}
\begin{pmatrix}1 \\ e^{-i\theta/2}
\end{pmatrix}
\begin{pmatrix}1 \\ e^{i\theta/2}
\end{pmatrix}
$$

for $y$ polarized light

$$
\Psi^\pm_y = V_y \cdot \Psi^\pm = \frac{1}{2} \begin{pmatrix}
\sigma_{\mu} \\
\sigma_{\mu}
\end{pmatrix}
\begin{pmatrix}
e^{-i(\alpha - \theta/2)} & e^{i(\alpha - \theta/2)} \\
-e^{i(\alpha - \theta/2)} & e^{-i(\alpha - \theta/2)}
\end{pmatrix}
\begin{pmatrix}1 \\ e^{\mp i\theta/2}
\end{pmatrix}
\begin{pmatrix}1 \\ e^{-i\theta/2}
\end{pmatrix}
\begin{pmatrix}1 \\ e^{i\theta/2}
\end{pmatrix}
\begin{pmatrix}
e^{i(\alpha - \theta/2)} & e^{-i(\alpha - \theta/2)} \\
-e^{-i(\alpha - \theta/2)} & e^{i(\alpha - \theta/2)}
\end{pmatrix}
\begin{pmatrix}1 \\ e^{-i\theta/2}
\end{pmatrix}
\begin{pmatrix}1 \\ e^{i\theta/2}
\end{pmatrix}
$$

The matrix elements for photoemission in this case depend on the relative phase $\phi$ for the layer amplitudes in the outgoing state in Eqn. 15. For $x$ polarized light the matrix elements are

$$
M^+_x(p, q) = \cos(\alpha - \theta/2) \cos(\phi/2)
$$

and for $y$ polarized light

$$
M^-_x(p, q) = -i \cos(\alpha - \theta/2) \sin(\phi/2)
$$

and

$$
M^+_y(p, q) = \sin(\alpha - \theta/2) \sin(\phi/2)
$$

and

$$
M^-_y(p, q) = i \sin(\alpha - \theta/2) \cos(\phi/2)
$$

In Eqns. 30 and 31 the phases $\alpha$ and $\theta$ are independent variables (the former is determined by coefficients in the Hamiltonian and the latter is the rotation angle) so that generically $\alpha - \theta/2$ is not a multiple of $\pi/2$. Thus the relevant interference physics is fully controlled by the interlayer phase $\phi$. When the outgoing waves from the two layers are in phase ($\phi = 2m\pi$) emission from states with even $R$-parity is allowed for $x$ polarized and forbidden for $y$ polarized light. Conversely, emission from the odd $R$-parity states is allowed for $y$ polarized light and forbidden for $x$ polarization. When the phase lag between layers $\phi = (2m + 1)\pi$ these selection rules are exactly reversed. As is the case for intralayer interference, nodal lines are absent for circular or elliptical polarization since this requires a simultaneous vanishing of the matrix elements in two orthogonal excitation channels.

When $\phi \neq (0, \pi)$ the emission between layers is neither exactly in or out of phase and nodal lines do not occur along the midline. Instead they are replaced by troughs (local minima) in the ARPES momentum distributions.

The $\phi$-derivatives of the momentum distributions measured for two orthogonal linear polarizations allow one to extract the internal phases $\alpha$ that define the wavefunctions $\Psi^\pm$ along this line. Physically these $\phi$ derivatives can be measured by measuring the differential change of the emission intensity at each position in $q$ space as a function of the excitation energy. Using Eqns. 30 and 31 one finds that the intensities for even $R$-parity state in two orthogonal polarizations are

$$
I_x = \cos^2(\alpha - \theta/2) \left(\frac{1 + \cos(\phi)}{2}\right)
$$

and therefore the ratio $r_{yx} = -(dI_y/d\phi)/(dI_x/d\phi)$ is $\phi$ (energy) independent and allows one to identify $\alpha(q)$

$$
\alpha(q) = \theta/2 + \arctan(r_{yx}(q))
$$

Eqn. 33 holds everywhere along the midline when the interlayer bias $V = 0$. When the interlayer bias is nonzero a similar expression can be used to determine the wavefunctions along a hyperbolic locus in $q$-space where the wavefunction $\Psi^\pm$ is an equal weight state shared between the two layers. Illustrative examples are given in Section V.D.

Eqns. 30 and 31 predict that nodal lines at $q_y = 0$ extend over the entire midline when the $R$-parity of the initial state is constant along this line. This is always the case when the bands are nonintersecting and illustrations of this are given in the following sections. Note however that when a symmetry protected band crossing occurs along the midline, the $R$-parity changes its sign at a singular point of degeneracy identifiable by the termination of a nodal line in one band (ordered by energy) and its appearance in another (also ordered by energy). The $q_y = 0$ midline is perpendicular to the offset $\Delta K$ between the bare Dirac nodes of the two layers so that crossings of this type do not occur in the simplest model of the rotated bilayer (Model I). However, in refined models for twisted bilayer graphene symmetry protected crossings do occur, notably along the line $q_y = 0$ near $E = 0$ in the strong coupling limit of Model II and at the energy of the second generation Dirac point (near the interlayer scale $\hbar v_F \Delta K$) for any coupling strength in Model III. This provides a unique spectroscopic diagnostic of singularities in their band structures.
The interlayer selection rules can be understood from the transformation of the current operators $V_\alpha$ under the symmetry operation $R$, $V_{x(y)}$ have even(odd) $R$-parities, and for $\phi = 0(\pi)$ the outgoing plane wave state in Eqn. (15) is even(odd). Thus the even parity initial state $\Psi^+$ can be excited into the outgoing final state with $\phi = 0(\pi)$ only using $x(y)$ polarization. Similar considerations apply for general interlayer phase difference $\phi$ with a rotation of the principal axes as noted above.

C. Branching Nodal Surfaces

The nodal patterns discussed in Section IV A arise from cancellation of photoemission amplitudes from the degrees of freedom in the individual layers, and in Section IV B they arise from cancellation of amplitudes from different layers. There are special points in $q$ space where both destructive interference conditions are satisfied. These can be identified as branching points where a nodal line running along the midline $q_y = 0$ develops branches that continue to the (renormalized) Dirac points of the two layers. The locations of these branching points can be determined from consideration of the bilayer coherent wavefunctions along the high symmetry $q_y = 0$ midline and are analyzed in this section.

Using the Hamiltonian in Eqn. (11) we partition the Hamiltonian at $q_y \neq 0$ into a Hamiltonian along the midline at $q_y = 0$ and a $q_y$ dependent piece

$$\mathcal{H}(q_y) = \mathcal{H}(q_y = 0) + \mathcal{H}'(q_y) = \mathcal{H}(q_y = 0) + q_y \tau_0 \sigma_2$$ (34)

Therefore the eigenstates near the midline can be expanded in terms of the eigenstates along the midline

$$\Psi_m(q_y) = \Psi_m(q_y = 0) + q_y \sum_n \langle \Psi_n(0) | \sigma_2 \tau_0 | \Psi_m(0) \rangle \frac{E_m - E_n}{E_m - E_n}$$ (35)

The midline eigenfunctions are also eigenfunctions of $R = \sigma_x \tau_z$, and since $\mathcal{H}'$ is odd under $R$ the sum connects only states of opposite $R$-parity. In a similar way the photoemission matrix element for the $m$-th band in the $\mu$-th polarization $\mathcal{M}_{\mu}$ can be expanded

$$\mathcal{M}_{m,\mu}(q_y) = \mathcal{M}_{m,\mu}(0) + q_y \sum_n \langle \Psi_n(0) | j_{\mu} | \Psi_m(0) \rangle \frac{\langle \Psi_n(0) | \sigma_2 \tau_0 | \Psi_m(0) \rangle}{E_m - E_n}$$ (36)

The sum in Eqn. (36) runs over the two states that reverse the $R$ parity of the initial state $\Psi_m(0)$ and using Eqns. (30) and (31) when $\mathcal{M}_{m,\mu}(0) = 0$ one sees that $\mathcal{M}_{n,\mu}(0)$ is nonzero for each of these admixed states. This means that the photoemission matrix element turns on linearly as a function of the transverse momentum $q_y$ and the nodal surface is an unbranched line for a general momentum $q_x$.

However a singular point can occur along the midline $(q_x,0)$ where the two contributions in the sum cancel. To locate such a point we label the two energy denominators between the state $m$ and its two intermediate states $n_1$ and $n_2$ by $\Delta_1 = E_m - E_{n_1}$ and $\Delta_2 = E_m - E_{n_2}$. Then the $q_y$-linear changes to the matrix element can be written as a real function of the phase angle $\alpha$

$$\delta \mathcal{M} = q_y \left( \frac{\cos 2\alpha \cos(\alpha - \theta/2)}{\Delta_1} + \frac{\sin 2\alpha \sin(\alpha - \theta/2)}{\Delta_2} \right)$$

where $\beta = \arctan((\Delta_1/\Delta_2) \tan 2\alpha)$ and $B$ is a constant. A critical branching point occurs when the right hand side of Eqn. (37) is zero. A nodal line forms two branches above and below the $q_x$ axis at such a critical point, once formed these arms continue to the Dirac points along the $q_y$ axis. This can be understood as a consequence of the continuity of the intralayer nodal lines that must connect the two Dirac points on opposite sides of the $q_x$ axis.

D. Examples

1. Model I

The nodal lines for Model I with $V_{\text{int}} \propto (1 + \sigma_x)$ are shown in Figure 7. We observe that the nodal lines that occur along the midline $q_y = 0$ propagate without termination. This occurs because the presence or absence of a nodal line is determined by the polarization state of the light and by the $R$-parity of the state, which does not change as a function of $q_x$ along the midline. The interlayer nodal lines also occur as complementary pairs, i.e. they are present for a particular band in $x$ polarization only when absent in $y$ polarization and vice versa. This occurs whenever the phase lag $\phi$ for emission between layers is a multiple of $\pi$. The phase difference $\phi$ can be tuned continuously by varying the excitation energy and for $\phi \neq m\pi$ the nodal lines are absent in both $x$ and $y$ polarizations. However for intermediate $\phi$ these lines are recovered when the principal polarization axes are also rotated according to the formulas given in Eqns. (30) and (31). No such energy dependence occurs for the nodal surfaces that terminate on the Dirac points which arise essentially from intralayer interference in the transition amplitudes and are therefore $\phi$-independent. The energy and polarization dependence of the nodal lines can therefore be used as a smoking gun to discriminate between intralayer and interlayer interference phenomena in the momentum distributions.
FIG. 7: Nodal lines in the photoemission amplitudes calculated for interlayer coupling Model I with $\hat{V}_{\text{int}} \propto (1 + \sigma_x)$. The top row is for $x$-polarized light and the bottom row is for $y$ polarized light. The four columns are for the four bands of the model ordered by energy. The $x$ polarized amplitudes in bands 2 and 3 have one-sided nodal lines at constant values of $q_y$ while the $y$ polarized amplitudes have nodal curves due to a gauge potential generated by the interlayer coupling. In bands 2 and 3 the nodal lines terminate at points (red dots) that can be identified with the Dirac points of the decoupled layers (green dots).

There are additional nodal lines in the second and third bands that terminate at the Dirac points. The termination and the appearance of these features correspond to a transferring a line node between the bands that touch at the Dirac points. The signature of the nodal patterns produced by the coupling in Model I is the appearance, in $x$ polarized excitation, of one-sided horizontal nodal lines that terminate at the Dirac points. This occurs because the interlayer operator in Model I has one zero eigenvalue, implying that there is one state with a particular pseudospin polarization in each layer that cannot be transported to its neighboring layer. This state is the antisymmetric eigenfunction of $\sigma_x$ and it cannot be coupled to the outgoing $\sigma_x$-symmetric final state by $x$ polarized radiation. Since these states are localized to the individual layers, they are eigenfunctions of the single layer $2 \times 2$ Dirac operators for which the pseudospin polarization remains constant along radially directed lines in $q$ space. Thus there are two such states with nodal lines that terminate at the unperturbed Dirac nodes of the two layers. In $y$ polarization the related lines nodes share the same termination points but the pseudospin polarization is a function of both $q_x$ and $q_y$ and is constant along a curve in momentum space, as shown in the lower panel. The curvature of this line is a manifestation of the momentum dependence of the gauge potential produced by the interaction with the neighboring layer. The termination points for these lines are the same for $x$ and $y$ polarized excitation, and thus the point singularities in this model can be identified with the geometrically determined Dirac points of two decoupled layers. This is a special feature of this class of interaction models.

2. Model II

In Model II $\hat{V}_{\text{int}} \propto \sigma_x$ so that the interlayer coupling is controlled by terms that connect opposite sublattice sites on neighboring layers. The signature of this model is the existence of a weak and strong coupling limit separated by a critical state where the renormalized Dirac points of the two layers are merged. The nodal structure in these two regimes is illustrated in Figures 8 and 9. The dimensionless coupling strength $\tilde{\Gamma}_1 = c_{AB}/\hbar v_F \Delta K$ so that the strong coupling limit can be realized for sufficiently small rotation angles. Using the data of Table I, one expects the strong coupling regime to describe the physics for rotation angles $\theta < 4^\circ$.

The nodal structure for Model II in the weak coupling regime is shown in Figure 8. It shares some features in common with Model I and the nodal pattern in $y$ polarization is quite similar. The main difference is in $x$ polarization where “flat” one-sided nodal lines of Model I evolve into dispersive nodal curves in Model II. As noted above the curvature of these lines is a momentum-space manifestation of the interlayer gauge potential for the twisted bilayer which is absent by symmetry for a single polarization in Model I.
FIG. 8: Nodal lines in the photoemission amplitudes calculated for interlayer coupling Model II with $\hat{V}_{\text{int}} \propto \sigma_x$. The top row is for $x$-polarized light and the bottom row is for $y$ polarized light. The four columns are for the four bands of the model ordered by energy. The data are shown for Model II in its weak coupling regime before the merger and reconstruction of the $E = 0$ Dirac points. The renormalized Dirac points (red dots) are displaced from the geometrically defined Dirac points of two decoupled layers (open green circles). The nodal lines in bands 2 and 3 terminate on the renormalized Dirac points. An additional nodal line along the midline is produced by interlayer interference of the transition amplitudes.

FIG. 9: Nodal lines in the photoemission amplitudes calculated for interlayer coupling Model II with $\hat{V}_{\text{int}} \propto \sigma_x$ in the strong coupling regime. The top row is for $x$-polarized light and the bottom row is for $y$ polarized light. The four columns are for the four band of the model ordered by energy. The data show the effect of reconstruction of the $E = 0$ Dirac points (red dots) in the strong coupling regime which are positioned along the $\pm q_x$ axis with the geometrically defined Dirac points (open green circles) along the $\pm q_y$ axis. The nodal lines in bands 2 and 3 terminate on these renormalized Dirac points. An additional nodal line along the midline occurs due to interlayer interference of the transition amplitudes.

In the strong coupling regime the nodal patterns in Model II shown in Figure 9 are more interesting. Here the reconstructed strong coupling Dirac points occur along the high symmetry $q_x$ axis. Thus the interlayer nodal lines can and do terminate at singular points along the $q_y$ axis. Interestingly these spectra also support branching points as shown. The nodal lines that branch away from these branch points propagate without termination to large momenta as shown.

3. Model III

In Model III the interlayer coupling matrix $\hat{V}_{\text{int}} \propto I$ so that the coupling is dominated by amplitudes that connect the same sublattice in the two layers. The nodal patterns calculated for this model are shown in Figure 10. The signature of this family of interaction models is the appearance of second generation symmetry-protected Dirac points at finite energy.
FIG. 10: Nodal lines in the photoemission amplitudes calculated for interlayer coupling Model III with $\hat{V}_{\text{int}} \propto \mathbb{I}$. The top row is for $x$-polarized light and the bottom row is for $y$ polarized light. The four columns are for the four bands of the model ordered by energy. Red dots denote the renormalized $E = 0$ Dirac points in the model that are displaced away from each other along the $\pm q_y$ axis by gauge potentials generated by the interlayer coupling. The open green circles denote the undisplaced Dirac nodes of two decoupled layers. The nodal lines in bands 2 and 3 terminate on these renormalized Dirac points. An additional one-sided nodal line terminating at a second generation Dirac point occurs along the midline occurs in all four bands due to interlayer interference of the transition amplitudes.

Interlayer nodal lines propagating along the $q_x$ axis collide with these second generation points where they switch between bands. The fundamental signature of this class of models is the appearance of new one-sided nodal lines along the midline $q_y = 0$. This behavior is clearly evident in both polarizations in Figure 10. The nodal lines that terminate on the renormalized $E = 0$ Dirac points are again curves rather than straight rays, manifesting the $\mathbf{q}$-dependence of the gauge potential generated by the interlayer coupling.

4. Phases along the midline

Figure 11 shows the internal phases $\alpha$ in the wavefunctions $\Psi_\pm$ calculated along the midline $q_y = 0$ using Eqn. 33 for the three interaction models discussed in the previous sections. We have verified that the internal phase $\alpha$ determined by differentiation of the emission intensities is independent of the value of the phase lag $\phi$ for emission from the two layers. A careful inspection of these plots allows one to further discriminate between the various interaction models.

All the models are characterized by an evolution from $\alpha = 0$ to $\alpha = \pi/2$ as a function of increasing $q_x$. The phase angle $\alpha$ is half the phase difference between sublattice amplitudes in the same layer. Thus this evolution describes the transition from states composed of the asymptotic (large $|q_x|$) layer eigenstates $(1, 1)$ to $(1, -1)$ which is a common feature in the negative energy solutions in all three interaction models.

The intermediate behavior is quite different in these models and can be used to analyze the symmetry of the interlayer coupling in the intermediate and small $q_x$ region. For example, in Model II (middle panel) the interlayer coupling matrix $\hat{V}_{\text{int}} \propto \sigma_x$ and using Eqn. 25 the phase $\alpha$ is then identified as the geometrical Dirac angle $\arctan(\Delta K/q_x)$. Note that this is the same for both negative energy bands at all values of $q_x$ in this model. By contrast Model III (right panel) shows a different phase angle $\alpha$ for the two negative energy states as a consequence of their interlayer coupling. The signature of this family of interaction models is both this phase splitting and a jump discontinuity that occurs at $q_x = 0$ where the two negative energy bands touch and the $R$-parity of a given band changes. Model I combines features of both: the phase angle $\alpha$ deviates from the Dirac angle defined by the rotational misorientation and the angles $\alpha$ evolve smoothly from 0 to $\pi/2$ in both manifolds.

VI. DISCUSSION

In this paper we have seen how the energy and polarization dependence of ARPES momentum distributions from bilayer graphene can be used to measure the phase structure of its layer-coupled wavefunctions. The nodal surfaces discussed in this paper can be measured in two
FIG. 11: Variation of the internal phase parameter $\alpha$ as a function of $q_x$ along the midline $q_y = 0$. The three panels are for three limiting coupling models: left panel: $\hat{V}_{\text{int}} \propto 1 + \sigma_x$, middle panel: $\hat{V}_{\text{int}} \propto \sigma_x$, right panel: $\hat{V}_{\text{int}} \propto 1$. The solid (dashed) curves give the variation of the phase angle $\alpha$ in the first (second) occupied band.

ways. One may sweep the excitation energy and keep the outgoing electron energy fixed, in which case one measures the momentum distribution as a function of a varying initial state energy. This protocol effectively sweeps the constant initial state energy contours through a pattern of nodal lines in their transition probabilities and allows a determination of momentum space nodal structures shown in Figures 7-10. The evolution of the momentum distributions during this type of sweep are illustrated in Figure 12 for interaction Models II and III. Alternatively, one can sweep the excitation energy and detection energy simultaneously, thereby measuring the momentum distribution for a single initial state energy. This protocol probes the symmetry of single initial state wavefunctions. Both methods are useful. The fingerprint of intralayer interference are nodal patterns which do not change as a function of the excitation energy, and this seems best suited to the first method. By contrast, interlayer interference effects are energy dependent and our analysis exploits the differential energy dependence of the emission intensities in orthogonal polarizations. This requires following the emission from the same initial state as the excitation energy is varied and can be analyzed most easily by the second method.

It will be useful to first confirm these predicted energy dependences experimentally. We expect there will be a clean separation between intra- and inter-layer interference effects seen in the experimental spectra with linearly polarized light and a suppression of these effects for circularly polarized light. Then focusing on the linearly polarized spectra, observation of the curvature of the intra-layer nodal surfaces that terminate on the primary Dirac points will provide a definitive experimental signature of the momentum dependence of the gauge potentials produced by the interlayer coupling. Conversely, observation of $\phi$-periodic variations of the emission intensities along particular lines in momentum space identify those features that are controlled by the interlayer coherence of the electronic states and differential dependence of their intensities on the excitation energy can be used to determined the sublattice symmetry of the matrix valued interlayer coupling potential.

Although we have focused on the situation with no vertical electrostatic potential difference between the layers the results can be generalized to graphene bilayers with scalar layer asymmetry. In this case the relevant momentum space contours deform to a family of mutually orthogonal confocal hyperbolas and ellipses. The extension of our methods to that situation is relatively straightforward and will likely be needed for a quantitative analysis of experimental data. The methods developed in this paper are also quite general and can be implemented with more sophisticated models for the electronic structure of BLG. It is hoped that measurement and analysis along these lines will provide a clean experimental resolution of the nature of the electronic states in twisted bilayer graphenes.

VII. ACKNOWLEDGEMENTS

This work was supported by the Department of Energy, Office of Basic Energy Sciences under Grant No. DE-

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FIG. 12: Illustration of the evolution of ARPES momentum distributions for three initial energies crossing through the nodal lines for two different interaction models. (Left panel) Model III with $\hat{V}_{\text{int}} = 0.15$. (Right panel) Model II in its strong coupling regime with $\hat{V}_{\text{int}} = 0.2\sigma$. The floor(ceiling) in these plots show the nodal surfaces superposed from the negative(positive) energy bands respectively.

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