Flat-band superconductivity in periodically strained graphene: mean-field and Berezinskii–Kosterlitz–Thouless transition

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Abstract
In the search of high-temperature superconductivity one option is to focus on increasing the density of electronic states. Here we study both the normal and s-wave superconducting state properties of periodically strained graphene, which exhibits approximate flat bands with a high density of states, with the flatness tunable by the strain profile. We generalize earlier results regarding a one-dimensional harmonic strain to arbitrary periodic strain fields, and further extend the results by calculating the superfluid weight and the Berezinskii–Kosterlitz–Thouless (BKT) transition temperature $T_{BKT}$ to determine the true transition point. By numerically solving the self-consistency equation, we find a strongly inhomogeneous superconducting order parameter, similarly to twisted bilayer graphene. In the flat-band regime the order parameter magnitude, critical chemical potential, critical temperature, superfluid weight, and BKT transition temperature are all approximately linear in the interaction strength, which suggests that high-temperature superconductivity might be feasible in this system. We especially show that by using realistic strain strengths $T_{BKT}$ can be made much larger than in twisted bilayer graphene, if using similar interaction strengths. We also calculate properties such as the local density of states that could serve as experimental fingerprints for the presented model.

Keywords: graphene, superconductivity, strain engineering, flat bands, BCS theory

Supplementary material for this article is available online
(Some figures may appear in colour only in the online journal)

1. Introduction
Graphene was long waiting for superconductivity to be added to its long list of miraculous properties. It took over ten years after its discovery before superconductivity was demonstrated in chemically doped graphene [1–4] with a critical temperature $T_c$ of a few kelvin. Recently the experiments on magic-angle twisted bilayer graphene (TBG) [5–7] have drawn much more attention, demonstrating superconductivity in a carbon-only material (although the role of the hexagonal boron nitride substrates is being disputed [8]) similarly with a $T_c$ of a few kelvin.

Lack of superconductivity in pristine graphene can be understood from the small-$\nu$ limit of the standard Bardeen–Cooper–Schrieffer (BCS) result for the critical temperature, $T_c \sim \omega_c e^{-1/|\lambda|\nu}$ [9, 10], with $|\lambda|$ describing the strength of the attractive electron–electron interaction, $\nu$ being the density of states (DOS) at the Fermi level, and $\omega_c$ being the cutoff (Debye) frequency. Since for intrinsic, undoped, graphene the density of states at the Fermi level is $\nu = 0$, according to this result we have also $T_c = 0$. The doping experiments can be understood from the same result. Since
close to the Dirac point $\nu$ increases linearly with chemical potential, doping can be utilized to render $T_c$ finite. But due to the exponential suppression of the critical temperature, to produce $T_c$ of a few kelvin, the chemical potential shift has to be of the order of eV [1, 3], corresponding to a very heavy doping level.

TBG provides an alternative mean to render $T_c$ finite: increase the density of states by flattening the electronic bands through moiré-modulated interlayer coupling. In the limit of a large $\nu$ (the flat-band limit), BCS theory gives a linear relationship $T_c \sim |\lambda|\Omega$ [10], where $\Omega$ is the area of the flat band, instead of the exponential one. The linear relation allows in principle to increase $T_c$ much higher even with a small interaction $|\lambda|$. Here the limiting factor seems to be the area $\Omega$ of the flat band, which in the case of TBG is roughly the superlattice (moire) Brillouin zone (SBZ), fixed by the rotation angle $\theta$. Since $\theta$ fixes also the interlayer coupling modulation, the whole dispersion is fixed by the rotation alone. From experiments [5, 11] and theories [12, 13] we know that in order to yield flat bands $\theta$ has to be close to the magic angle $\theta^* \approx 1^\circ$, for which $\Omega$ is only about 0.04% [14] of the original graphene Brillouin zone (BZ). An increase of a few kelvin in $T_c$ has been successfully demonstrated [6] by applying high pressure to slightly increase $\theta^*$ and thus also $\Omega$. In TBG the flat bands are in fact not exactly at zero energy, but of the order of meV higher and lower. But compared to chemically doped graphene where ~eV doping levels are needed, a thousandfold reduction in the needed chemical potentials allows using much simpler and more easily tunable electrical doping.

In this work we study yet another mechanism to produce flat bands in graphene, which is possibly free of the limitations in TBG: periodic strain [15–21]. Instead of periodically modulating interlayer hopping in TBG, we modulate the intralayer hopping in monolayer graphene by periodic strain. In this system we can, in principle, separately choose the strain period $d$ (and thus the SBZ area $\sim \Omega$) and its strength $\beta$ (and thus the flatness of the bands), potentially allowing us to increase $T_c$ higher than in TBG by engineering strains with high amplitude and small period.

At low energies, near the $K$ and $K' = -K$ points where graphene can be described as a Dirac material, strain is modeled by a pseudo vector potential $A$ [16, 18, 22–24], similarly to an external magnetic field. While the external magnetic field breaks the time-reversal symmetry and usually suppresses superconductivity, the strain-induced $A$ has opposite signs on different valleys, preserving time-reversal symmetry and thus preserving and even promoting spin-singlet superconductivity. Moreover, strain-induced pseudo vector potentials can easily reach an effective magnetic field strength of tens of Tesla or even hundreds [21, 26] of tesla, opening the possibility for extreme tuning of electronic properties.

Possibilities for experimentally producing periodic strain in graphene are numerous. In fact, flat bands have already been observed in an experiment by Jiang et al [21], where both 1D and 2D periodic strains were created by boundary conditions. In this experiment the displacement amplitude was of the order of 1 Å and the period $d$ was tunable between 8 and 25 nm. Even better control of the strain pattern could perhaps be achieved by optical forging [27], which allows drawing arbitrary out-of-plane strain patterns in graphene, even below the diffraction limit [28]. On the other hand the small secondary ripples observed in the simulations [27] could be exploited, similarly to the Jiang et al experiment [21], but with better control.

Another experimentally demonstrated method is to use an AFM tip to evaporate hydrogen from a Ge-H substrate to produce a pressurized H$_2$ gas under specific locations of graphene [29]. One option could be graphene on a corrugated surface [30, 31]. Applying in-plane compression has been predicted to produce periodic wrinkles both in simply-supported [30, 32] and encapsulated [33, 34] graphene, with amplitude and period of the order of 0.2 Å and 2 nm, respectively. In the same spirit the proposed graphene cardboard material could be manufactured [35]. Also an ultracold atom gas in a tunable optical honeycomb lattice [36] could be used.

It has been predicted [37–41] and observed [42] that TBG exhibits moiré-periodic strain due to lattice mismatch and the following structural relaxation. The relative magnitude of the moiré and strain effects can be, however, difficult to disentangle, as superconductivity by both effects has been predicted by BCS theory [14, 19]. But if the moiré effect is enhancing superconductivity, as it seems to be, we get a lower bound for $T_c$ by studying the strain effects. Similarly periodic strain can be expected with other mismatch lattices, such as graphene on hBN [6].

In this work we generalize the model and results of Kaupila et al [19], where both the normal and superconducting spin-singlet, s-wave state in periodically strained graphene (PSG) have been studied in the case of a cosine-like 1D potential $A(x, y) = \frac{\pi}{d}(\cos(2\pi x/d))$, to arbitrary periodic pseudo vector potentials $A$. This generalization is motivated by the experiment of Jiang et al [21], where a variety of periodic strain patterns, both 1D and 2D, were manufactured. On the other hand generalizing the theory to 2D strains bridges the gap between PSG [19] and TBG [14] by showing how similar these two systems are in many aspects.

The main conclusions of Kaupila et al are that (i) approximate flat bands are formed in the normal state, (ii) the superconducting order parameter $\Delta(x)$ becomes inhomogeneous and is peaked near the minima/maxima of $\nabla \times A$, similarly to the local density of states (LDOS), (iii) magnitude of $\Delta$ can be tuned by the amplitude $\beta$, (iv) $T_c$ is linear in $\lambda$ in the flat-band regime (large $\lambda$ or $\beta$), and (v) even though $\Delta$ is strongly inhomogeneous and anisotropic, supercurrent is only slightly anisotropic. We show that these results continue to hold even when we change the shape of $A$ and move to 2D potentials. In addition we show how the shape of $A$ and its dimensionality affect the superconducting order parameter $\Delta$, the critical chemical potential $\mu_c$, and the critical temperature $T_c$. We furthermore extend the calculations by calculating the superfluid weight [43, 44] $D^*$ and the Berezinskii–Kosterlitz–Thouless (BKT) transition temperature $T_{cKT}$ to determine the proper transition temperature in a 2D system.

In addition to Kaupila et al, spin-singlet, s-wave superconductivity in strained graphene was studied also by Uchoa et al [16]. They, however, concentrated on strain fields with
a homogeneous pseudomagnetic field \( \mathbf{B} = \nabla \times \mathbf{A} \) and correspondingly to a superconducting state with a homogeneous order parameter \( \Delta \). This constraint allowed them to derive analytical formulas e.g. for \( \Delta \) and \( T_c \), yielding a similar \( T_c \)-linear-in-\( \lambda \) result as what Kaupilla et al found out. Here we instead focus on periodic and inhomogeneous strain accompanying also a periodic and inhomogeneous \( \mathbf{B} \), which is probably more accessible experimentally, since a constant \( \mathbf{B} \) might be difficult to obtain [26]. Our approach is also complementary to Uchoa et al in the sense that the periodicity of \( \mathbf{B} \) allows us to use the notion of (flat) electronic bands, whereas in the pseudo-Landau-level perspective of Uchoa et al the electronic levels are the pseudo-Landau levels.

This article is organized as follows. In section 2 we derive the Bogoliubov–de Gennes (BdG) theory to describe the superconducting state of PSG at low energies, details of which are shown in the supplementary material [2]. In section 3 we present the results of applying some selected periodic pseudo vector potentials \( \mathbf{A} \) by numerically solving the self-consistency equation. In section 4 we summarize the main results and discuss open questions and future prospects.

2. Model

In the low-energy limit, after adding an in-plane displacement field \( \mathbf{u} \) and an out-of-plane displacement field \( h \), the graphene continuum Hamiltonian for valley \( \rho \in \{+,-\} \) is

\[
\mathcal{H}^\rho(r) = \hbar v_F \mathbf{\sigma}^\rho \cdot (-i \nabla + \rho \mathbf{A}(r)) - \mu,
\]

where the pseudo vector potential is given by [22, 23]

\[
\mathbf{A} = -\frac{\beta_G}{2a_0}(u_{xx} - u_{yy}, -2u_{xy})
\]

and the strain tensor is

\[
u_{ij} = \frac{1}{2}(\partial_i u_j + \partial_j u_i) + \frac{1}{2} \partial_i h \partial_j h.
\]

Here \( v_F \) is the graphene Fermi velocity, \( \mu \) is the chemical potential, \( \beta_G = -d \ln \gamma / d \ln a_0 \approx 2 \) is the graphene Grüneisen parameter [23], \( a_0 \) is the carbon–carbon bond length, \( \mathbf{\sigma}^\rho = (\rho \mathbf{\sigma}_x, \mathbf{\sigma}_y) \) is a vector of sublattice-space Pauli matrices, and the graphene zigzag direction is assumed to be in the \( x \) direction. Note that \( \mathbf{A} \) works exactly like a vector potential related to an external magnetic field, but with the important difference that it changes sign on valley exchange \( \rho \to -\rho \), preserving time-reversal symmetry \( \mathcal{H}^\rho = \mathcal{H}^{-\rho} \). Because of the relation (2) we use the words 'strain' and 'pseudo vector potential' interchangeably. Note that for the linear elasticity theory to be valid we should have \( \beta \)

\[
\|\mathbf{u}(r + \delta_1) - \mathbf{u}(r)\|, \|h(r + \delta_1) - h(r)\| \ll a_0,
\]

where \( \delta_1 \), \( \delta_2 \), and \( \delta_3 \) are the graphene nearest neighbor vectors.

We model the possible superconducting state by a (slightly generalized) BCS theory using BdG formalism. We assume an intervalley, local (also in sublattice) interaction of strength \( \lambda \) (negative for attractive interaction considered here), which has been widely used in the past graphene literature [45–50] to model \( s \)-wave superconductivity. In this case the effective interacting mean-field continuum Hamiltonian can be shown to be \( 2 \)

\[
\mathbf{H}_{\text{BdG}} = \sum_{\sigma \rho} \int d\mathbf{r} \psi^\dagger_{\sigma \rho}(r) \mathcal{H}^\rho(\mathbf{r}) \psi_{\sigma \rho}(r) + \frac{1}{2} \sum_{\sigma \rho} \int d\mathbf{r} \psi^\dagger_{\sigma \rho}(r) \Delta_{\sigma}(r) \psi_{\sigma \rho}^\dagger(r) + \text{h.c.} + \text{const},
\]

where \( \sigma \in \{\uparrow, \downarrow\} \) denotes spin, the real-space integrals are over the Born–von Kármán cell \( \mathbb{R}^2 / L_{\text{Bak}} \), and \( \psi_{\sigma \rho}(r) = (\psi_{\sigma \rho A}(r), \psi_{\sigma \rho B}(r))^T \) is a sublattice-space vector of the electron annihilation operators. Furthermore the superconducting order parameter in the sublattice space is \( \Delta_{\sigma}(r) = \text{diag}(\Delta_{\sigma A}(r), \Delta_{\sigma B}(r)) \), where

\[
\Delta_{\sigma, \alpha} = \lambda \sum_{\rho} \langle \psi_{\sigma \rho A} \psi_{\sigma \rho A} \rangle
\]

with angle brackets denoting the thermal average and \( \alpha \in \{A, B\} \) denoting the sublattice. Note that this kind of a local interaction corresponds to spin-singlet type of superconductivity, since from the fermionic anticommutation relations it directly follows that \( \Delta_{\sigma, \alpha} = -\Delta_{\sigma, \bar{\alpha}} \). Furthermore due to locality \( r \) denotes the center-of-mass coordinate of the Cooper pair, while the relative coordinate is always zero, meaning that this interaction corresponds to \( s \)-wave superconductivity.

Utilizing the fermionic anticommutation relations and by doubling the basis set we can bring \( \mathbf{H}_{\text{BdG}} \) in (5) into the Nambu form

\[
\mathbf{H}_{\text{BdG}} = \frac{1}{2} \sum_{\sigma \rho} \int d\mathbf{r} \psi^\dagger_{\sigma \rho}(r) \mathcal{H}_{\text{BdG}}(\mathbf{r}) \psi_{\sigma \rho}(r) + \text{const},
\]

where the BdG Hamiltonian in Nambu space and the Nambu-vector are

\[
\mathcal{H}_{\text{BdG}}^\rho = \begin{pmatrix} \mathcal{H}^\rho & \Delta \\ \Delta^* & -\mathcal{H}^\rho \end{pmatrix}, \quad \psi_{\sigma \rho} = \begin{pmatrix} \psi_{\sigma \rho A} \\ s(\sigma) \psi_{\sigma \rho B}^\dagger \end{pmatrix},
\]

respectively. Here the spin-independent order parameter is \( \Delta = \Delta_1 = s(\sigma) \Delta_{\sigma A}, s(\uparrow) = 1 \), and \( s(\downarrow) = -1 \).

Using the spectral theorem, a symmetry between the positive- and negative-energy states, and defining the fermionic Bogoliubov operators as

\[
\gamma_{\sigma \rho \delta} = \frac{1}{\sqrt{V}} \int d\mathbf{r} \psi_{\sigma \rho \delta}^\dagger(\mathbf{r}) \psi_{\sigma \rho}(\mathbf{r})
\]

we may bring \( \mathbf{H}_{\text{BdG}} \) into the diagonal form \( 2 \)

\[
H = \frac{1}{2} \sum_{\sigma \rho \delta} E_{\sigma \rho \delta} \gamma_{\sigma \rho \delta}^\dagger \gamma_{\sigma \rho \delta} + \text{const}.
\]
Here \( \mathbf{k} \) together with the band index \( b \) enumerate the positive-energy solutions of the BdG equation

\[
\mathcal{H}_{\text{BdG}}^\rho(\mathbf{r})w_{\rho \mathbf{k}}(\mathbf{r}) = E_{\rho \mathbf{k}}w_{\rho \mathbf{k}}(\mathbf{r})
\]

(11)

and \( V = |\mathbf{r}|^2/L_{\text{Bk}} \) is the area of the Born–von Kármán cell. According to the calculation above, diagonalizing \( \mathcal{H}_{\text{BdG}}^\rho \), i.e., bringing it to the form (10), is equivalent to solving the BdG equation (11).

By inverting the Bogoliubov transformation (9) we write the definition of the order parameter (6) as the self-consistency equation

\[
\Delta_\alpha(\mathbf{r}) = -\frac{\lambda}{V} \sum_{\rho \mathbf{k}} u_{\rho \mathbf{k}, \alpha}(\mathbf{r}) v_{\rho \mathbf{k}, \alpha}^*(\mathbf{r}) \tanh \left( \frac{E_{\rho \mathbf{k}}}{2k_B T} \right),
\]

(12)

at temperature \( T \), where we denoted the Nambu components of \( w \) as \( w = (u, v) \). Note that \( \Delta_\alpha \) might depend on sublattice \( \alpha \), while Kauppila et al [19] defined \( \Delta \) by summing over \( \alpha \).

As we see below, the self-consistent \( \Delta_\alpha \) is, in fact, sublattice dependent, leading to a different \( T \) than in [19].

In real space the self-consistency equation (12) is local in space but the BdG equation (11) is a group of 2D differential eigenvalue equations. The equations can be made easier to solve by utilizing periodicity of \( \mathbf{A} \) and writing them in Fourier space. We assume both the pseudo vector potential \( \mathbf{A} : \mathbb{R}^2 / \text{SL} \rightarrow \mathbb{R}^2 \) (and thus the strain) and the order parameter \( \Delta \) to be periodic in translations of the arbitrary superlattice \( \text{SL} = \text{span}_\mathbb{Z}\{\mathbf{t}_1, \mathbf{t}_2\} \subset \mathbb{R}^2 \), allowing us to use the Fourier series

\[
A(\mathbf{r}) = \sum_G e^{i \mathbf{G} \cdot \mathbf{r}} \tilde{A}(\mathbf{G}), \quad \Delta(\mathbf{r}) = \sum_G e^{i \mathbf{G} \cdot \mathbf{r}} \tilde{\Delta}(\mathbf{G}).
\]

(13)

Here the sums are over \( \text{SL}^*_G \), where \( \text{SL}^*_G = \text{SL}^* = \text{span}_\mathbb{Z}\{\mathbf{G}_1, \mathbf{G}_2\} \) is the reciprocal lattice of \( \text{SL} \). \( \text{SL}^*_G = \text{span}_\mathbb{Z}\{\mathbf{G}_1\} \) is a one-dimensional sublattice of \( \text{SL}^* \), and \( S \in \{\text{RZ, MZ}\} \) denotes either the reduced zone scheme or the mixed zone scheme (the terms are justified below), the latter of which being applicable only if \( \mathbf{A} \) and \( \Delta \) are constant in the \( t_2 \) direction, which we call the 1D potential case. Otherwise we call \( \mathbf{A} \) a 2D potential.

Together with the assumption of the eigenfunctions \( w_{\rho \mathbf{k}} \) being periodic in the Born–von Kármán cell, the Fourier series (13) imply the existence of the Bloch-type Fourier series

\[
w_{\rho \mathbf{k}}(\mathbf{r}) = e^{i \mathbf{G} \cdot \mathbf{r}} \sum_{G} e^{i \mathbf{G} \cdot \mathbf{r}} \tilde{w}_{\rho \mathbf{k}}(\mathbf{G})
\]

(14)

and the Fourier space version of the BdG equation

\[
\sum_{\mathbf{G}'} \tilde{\mathcal{H}}_{\text{BdG,G'}}^\rho(k) \tilde{w}_{\rho \mathbf{k}}(k + \mathbf{G'}) = E_{\rho \mathbf{k}} \tilde{w}_{\rho \mathbf{k}}(k + \mathbf{G}).
\]

(15)

In the matrix form (15) can be written as

\[
\tilde{\mathcal{H}}_{\text{BdG}}^\rho(k) \tilde{w}_{\rho \mathbf{k}} = E_{\rho \mathbf{k}} \tilde{w}_{\rho \mathbf{k}},
\]

(16)

where the underlined variables are matrices or vectors in the \( \mathbf{G} \) space. Here \( k \in L_{\text{Bk}}^*/\text{SL}^*_G \) belongs to the superlattice Brillouin zone (SBZ) in the scheme \( S \), \( b \) enumerates the positive-energy bands for each \( k \), and the Nambu-space BdG Hamiltonian is

\[
\tilde{\mathcal{H}}_{\text{BdG,G'}}^\rho(k) = \begin{pmatrix}
\Delta_\mathbf{G}_2^\rho & \Delta_\mathbf{G}_1^\rho \\
\Delta_\mathbf{G}_1^\rho & -\Delta_\mathbf{G}_2^\rho
\end{pmatrix}
\]

(17)

with the noninteracting (normal state) Hamiltonian

\[
\tilde{\mathcal{H}}_{\text{G'}}^\rho(k) = \hbar v_T \sigma^\rho \cdot (k + \mathbf{G}) \delta_{\mathbf{G}'} + \mu \delta_{\mathbf{G}'} - \mu \delta_{\mathbf{G}'},
\]

(18)

Note the similarity to the Dirac-point low-energy TBG model in [14, 50, 51]: while here \( \mathbf{A} \) couples the sublattices and \( \mathbf{G} \) vectors within the layer, in TBG the Hamiltonian (18) has a two-layer structure, \( \mathbf{A} \) is absent, and the interlayer coupling \( t_{\|} \) couples sublattices and \( \mathbf{G} \) vectors between the layers.

As we show in this paper, the second layer is not necessary for yielding flat bands, but what seems to be enough is coupling in the \( \mathbf{G} \) space. To generalize the theory to study the combined effect of periodic strain and moiré physics, which should yield even more pronounced flat bands, would be thus easy: add the second rotated layer to the noninteracting Hamiltonian (18) and couple the layers by \( t_{\perp}(k + \mathbf{G}) \).

Let us discuss the notion of the reduced and the mixed zone schemes. In the reduced zone scheme \( k = k_{1, \mathbf{G}_1} + k_{2, \mathbf{G}_2} \in L_{\text{Bk}}^*/\text{SL}^*_G \) is periodic both in the \( \mathbf{G}_1 \) and \( \mathbf{G}_2 \) directions, with both \( k_{1, \mathbf{G}_1}, k_{2, \mathbf{G}_2} \in [-\frac{1}{2}, \frac{1}{2}] \) being periodic Bloch momenta. This is also traditionally called the reduced zone (or the repeated zone) scheme. In the case of \( \mathbf{A} \) and \( \Delta \) being constant in the \( t_2 \) direction (the 1D potential case), we are also allowed to use the mixed zone scheme, where \( k = k_{1, \mathbf{G}_1} + k_{2, \mathbf{G}_2} \in L_{\text{Bk}}^*/\text{SL}^*_M \) is periodic only in the \( \mathbf{G}_1 \) direction but not in the \( \mathbf{G}_2 \) direction, with \( k_{1, \mathbf{G}_1} \in [-\frac{1}{2}, \frac{1}{2}] \) being a periodic Bloch momentum and \( k_{2, \mathbf{G}_2} \in [-\infty, \infty] \) being a nonperiodic real momentum. Thus in the traditional notion the \( \mathbf{G}_1 \) direction is in the reduced (or repeated) zone and the \( \mathbf{G}_2 \) direction in the extended zone scheme, justifying the term mixed zone scheme.

The reduced zone scheme is convenient if one wants to compare the effects of the 1D and 2D potentials, since the dispersions look similar and the notion of a band is the same, but the calculations are heavy due to the \( \mathbf{G} \) space being two-dimensional. On the other hand the mixed zone scheme produces cleaner-looking dispersions and is computationally much lighter due to the \( \mathbf{G} \) space being one-dimensional, but with the cost of more difficult comparison between the 1D and 2D potentials. Thus in all the 1D potential calculations we use the mixed zone scheme unless otherwise stated. Also Kauppila et al [19] used the mixed zone scheme in all the calculations and visualizations.

Using the Fourier series (13) and (14) in (12) and approximating the \( k \) sum as an integral (assuming the Born–von Kármán cell to be large), the Fourier-space self-consistency equation becomes

\[
\Delta_\alpha^\rho(\mathbf{G}) = -\frac{\lambda}{(2\pi)^2} \sum_G \int d\mathbf{k} \tanh \left( \frac{E_{\rho \mathbf{k}}}{2k_B T} \right) \times \tilde{u}_{\rho \mathbf{k}, \alpha}(k + \mathbf{G}) \tilde{v}_{\rho \mathbf{k}, \alpha}^*(k + \mathbf{G} - \mathbf{G}_2),
\]

(19)
where the integral is over the continuum superlattice Brillouin zone \( \mathbb{R}^2 / SL^*_5 \) in the scheme \( S \), which in the reduced zone scheme can be interpreted as the parallelogram defined by \( G_1 \) and \( G_2 \), and in the mixed zone scheme as the semi-infinite parallellogram with the finite side being \( G_1 \) and the infinite side being in the direction of \( G_2 \).

In summary, in Fourier space we are solving the BdG equation (15) together with the self-consistency equation (19). Now the BdG equation is a normal matrix eigenvalue equation, but the price to pay is that the corresponding matrix has countably infinite dimension \( 2 \times 2 \times | SL^*_5 | \), and the self-consistency equation becomes nonlocal in the Fourier components. Numerically, however, they are easy to solve, provided we truncate the Fourier-component set \( SL^*_5 \) and the band sum, and in the case of 1D potential add a momentum cutoff in the \( k \) integral in the \( G_2 \) direction. These cutoffs we choose so large that the results (dispersion, \( \Delta \)) start to become saturated, and together they correspond to the energy cutoff \( \hbar \omega_c \) introduced earlier.

In a 2D system, however, we know that the superconducting transition is not properly described by the mean-field critical temperature \( T_c \) determined from the order parameter \( \Delta \), but by the BdG transition temperature determined from the superfluid weight \( D^s \), which describes the linearized supercurrent density response \( \langle j \rangle = \frac{\hbar e}{4 \pi} D^s (\mathbf{A}) \) to an external (real) vector potential \( \mathbf{A} \), where the angle brackets denote average over position. For the present model we may calculate the \( \mu, \nu \in \{ x, y \} \) component of the superfluid weight from \( \frac{\hbar^2 k}{2m_e} D^s_{\mu \nu} = \frac{\hbar^2 k}{2m_e} \sum_{\rho \rho'} \int d\mathbf{k} \frac{f(E_{\rho \rho'}) - f(E_{\rho' \rho})}{E_{\rho \rho'} - E_{\rho' \rho}} \times \left( \sigma_{\rho \rho'}^\mu \sigma_{\rho \rho'}^\nu \tilde{w}_{\rho \rho'}^{\dagger} \tilde{w}_{\rho \rho'}^{\dagger} - \tilde{w}_{\rho \rho'}^{\dagger} \tau^\rho \sigma_{\rho \rho'}^\mu \tilde{w}_{\rho \rho'}^{\dagger} \tilde{w}_{\rho \rho'}^{\dagger} \tau^\rho \sigma_{\rho \rho'}^\nu \right), \)

where the \( b, b' \) band sums are calculated over both the positive- and negative-energy bands, \( \tau^\mu \) is the Pauli-\( z \) matrix in Nambu space, \( f \) is the Fermi–Dirac distribution, the difference quotient is interpreted as the derivative \( f(E_{\rho \rho'}) \big|_{E_{\rho \rho'} = E_{\rho' \rho}} \), and where we suppressed the \( k \) dependence.

From the temperature dependence of \( D^s \) we can then calculate the BdG transition temperature \( T_{\text{BKT}} \) from the generalized KT–Nelson criterion \( 52–54 \)

\[
k_B T_{\text{BKT}} = \frac{\pi}{8} \sqrt{\det D^s (T_{\text{BKT}})}
\]

for an anisotropic superfluid weight, which also needs to be calculated self-consistently, unless \( D^s (T_{\text{BKT}}) \approx D^s (0) \).

3. Results

We solve\(^3\) the order parameter \( \Delta \), the superfluid weight \( D^s \), and the Berezinskii–Kosterlitz–Thouless transition temperature \( T_{\text{BKT}} \) for a selection of periodic pseudo vector potentials

\[^3\]The MATHEMATICA code we have used to compute the normal-state properties, the mean-field order parameter, the superfluid weight, and the BKT transition temperature is available at https://gitlab.jyu.fi/jyucmt/psg-and-tbg.
primitive vectors are $G_1 = (2\pi/d, 0)$ and $G_2 = (0, 2\pi/d)$. But since for the 1D potentials the components are multiplied by $\delta_{m,0}$, we may as well use a one-dimensional Fourier series and define in the mixed zone scheme

$$A_{\cos}^{1D}(m_1 G_1) = \frac{\beta}{2d} (0, \delta_{m_1,-1} + \delta_{m_1,1}), \quad (25)$$

where $m_1 G_1 \in \text{SL}_{AZ}$ belongs to SBZ in the mixed zone scheme.

On the other hand in the 2D case we concentrate on the simplest generalization of the 1D cosine-like potential $A_{\cos}^{1D}$, the potential

$$A_{\cos}^{2D}(x, y) = \frac{\beta}{d} (\cos(2\pi y/d), \cos(2\pi x/d)) \quad (26)$$

with the lattice of periodicity being the square superlattice $\text{SL} = \text{span}_Z \{ t_1, t_2 \}$, with the primitive vectors $t_1 = (d, 0)$ and $t_2 = (0, d)$. Note that we are allowed to choose a potential periodic in any superlattice, whereas in TBG the (moiré) superlattice is fixed by the rotation angle. Thus in principle the periodic strain allows much more freedom in tuning the system. The Fourier components of the 2D cosine potential are

$$A_{\cos}^{2D}(m_1 G_1 + m_2 G_2) = \frac{\beta}{2d} (\delta_{m_2,-1} + \delta_{m_2,1}, \delta_{m_1,-1} + \delta_{m_1,1}) \quad (27)$$

in the reduced zone scheme, where $m_1 G_1 + m_2 G_2 \in \text{SL}_{AZ}$ with $G_1 = (2\pi/d, 0)$ and $G_2 = (0, 2\pi/d)$.

According to (2) the potentials $A_{\cos}$ and $A_{\cos}^{2D}$ can be produced for example in the in-plane displacement fields

$$u_{\cos}^{1D}(x, y) = \frac{\beta_0}{\beta_G \pi} (0, \sin(2\pi x/d)), \quad (28)$$

$$u_{\cos}^{2D}(x, y) = \frac{\beta_0}{\beta_G \pi} (0, \sin(2\pi x/d) + \sin(2\pi y/d)), \quad (29)$$

respectively. The pseudomagnetic fields $B = \nabla \times A = \partial_y A_x - \partial_x A_y$ produced by the 1D and 2D cosine potentials, together with these example displacement fields, are shown in figures 2(a)–(d). The amplitude $B$ of $B$, which is an important factor determining the flatness of the bands and the magnitude of the superconducting order parameter $\Delta_{ls}$, is

$$B_{\cos}^{1D} = \frac{2\pi \beta}{d^2}, \quad B_{\cos}^{2D} = \frac{c \beta}{d^2}, \quad B_{\cos}^{2D} = \frac{4\pi \beta}{d^2} \quad (30)$$

for the potential $A_{\cos}^{1D}, A_{\cos}^{2D}$, or $A_{\cos}^{2D}$, respectively. To give a realistic scale for $\beta$, in the experiment by Jiang et al [21] a pseudomagnetic field of $\frac{1}{2}B \approx 100 \text{T}$ was observed for a strain period of $d = 14 \text{ nm}$, which corresponds to $\beta \approx 5$ for the 1D cosine potential. To be better in the flat-band regime, we mostly use a factor of 4–8 times larger values of $\beta$ in this study.

Corresponding typical profiles of $\Delta_{ls}$ for the cosine potentials are shown in figures 2(e)–(f), from where it is clear that $\Delta_{ls}$ is always peaked at the minima/maxima of the pseudomagnetic field $B$. For comparison in TBG [14] $\Delta$ is localized around the AA stacking regions and is independent of the sublattice and layer. Note that the sublattice dependence was not present in the work by Kauppila et al [19] due to sublattice-summation in the self-consistency equation. As we see below, it is approximately the maximum (over position $r$) of the order parameter that is important in describing the strength of the superconducting state. As for all the studied potentials the maximum of the order parameter is independent of the sublattice, we simply denote $\Delta := \max A_{ls}$.

The typical dispersion relations in the normal state are shown in figure 3 together with the conical unstrained graphene dispersions. For an easier comparison the 1D potential $A_{\cos}^{1D}$ dispersion is shown both in the mixed zone and reduced zone schemes, while the 2D potential $A_{\cos}^{2D}$ dispersion only in the reduced zone scheme (the only possibility in this case). We find similar-looking approximate flat bands as in TBG [14, 51], with the difference that here the number and the flatness of the flat bands can be controlled by $\beta$ and $c$. Also all the successive bands are touching, while in TBG many models predict the flat bands to be isolated [39, 41, 51, 55].

We calculate most of the superconducting state results at optimal doping $\mu = \mu_{\text{opt}}$, which is the energy of the density...
we show the ratio in the flat-band regime the behavior of the order parameter is exponentially suppressed and we also which we call the sive behavior of the lowest energy bands starts playing a role, On the other hand for small enough except for zone scheme (RZ). (c) The 2D cosine potential \( \beta = \mu \) (c) at \( B \) the pseudomagnetic field dispersions in orange.

\[
\rho = \pm \frac{30}{7} \text{opt and } \Omega \text{ for all the potentials in figures 4(b) and (c) in the mixed zone scheme (MZ) and (b) in the reduced zone scheme (RZ). (c) The 2D cosine potential } A_{\text{cos}}^{2D} \text{ (shown for } \beta = 15) \text{ in the reduced zone scheme. The strained dispersions are shown in blue and for comparison the conical unstrained graphene dispersions in orange.}
\]

Figure 3. Typical dispersions in the normal state at the valley \( \rho = \pm \) with \( \mu = 0 \). (a, b) The 1D cosine potential \( A_{\text{cos}}^{1D} \) (shown for \( \beta = 30 \)) (a) in the mixed zone scheme (MZ) and (b) in the reduced zone scheme (RZ). (c) The 2D cosine potential \( A_{\text{cos}}^{2D} \) (shown for \( \beta = 15 \)) in the reduced zone scheme. The strained dispersions are shown in blue and for comparison the conical unstrained graphene dispersions in orange.

\[
\text{max } \Delta = -\zeta B \lambda,
\]

we show the ratio \( \zeta \) for all the potentials in figures 4(b) and (c) at \( \mu = \mu_{\text{opt}} \) and \( T = 0 \). In the flat-band regime \( \zeta \) tends approximately to a constant \( \zeta \approx 0.15 \), which holds as long as \( c \lesssim 20 \). For \( c \approx 20 \) we start seeing deviations from this result, with \( \zeta \approx 0.05 \) for the extreme case of \( c = 100 \). The small variation in \( \zeta \) due to \( c \) even in the flat-band regime is most likely due to the fact that the maximum of \( \Delta \) is not exactly the correct quantity to calculate, but it gives a very good estimate. We may compare this to the exact-flat-band result \([14]\) with a constant \( \Delta_{\text{FB}}^{1D} \), for which \( \Delta_{\text{FB}}^{1D} = -\frac{1}{\pi} n \Omega \lambda \) with \( \Omega = 1/d^2 \) and \( n \) being the number of flat bands. In PSG it is the amplitude \( B \) of the pseudomagnetic field \( B \) that effectively determines \( n \Omega \), the number of approximate flat bands in the system with the SBZ area of \( 1/d^2 \). A similar linear relationship was found by Uchoa et al \([16]\) in the case of a homogeneous pseudomagnetic field \( B \), although with a somewhat larger prefactor \( \zeta \).

3.1. Order parameter profile, dispersion, and density of states

In figure 5 we show a cross section of the self-consistent \( \Delta_{\text{FB}}^{1D} \) [as in figures 2(e) and (f)] along the line \((x, 0) \) [1D potentials] or \( r(1, -1) \) [2D potential] for different potentials \( A \), strain strengths \( \beta \), and slope parameters \( c \). The effect of \( \beta \) is to simply linearly increase the amplitude of \( \Delta_{\text{FB}}^{1D} \). On the other hand increasing \( c \) not only increases the amplitude of \( \Delta_{\text{FB}}^{1D} \),
In the normal state the overall energy dependence shows \(\Delta V\) varying the 2D cosine potential parameter \(c\). But makes it also more localized. We also see that for the 2D potential \(A_{2D}^{\cos}\) with the strain strength \(\beta\) along the diagonal behaves similarly as \(\Delta A_{1D}/\lambda\) in the x direction for the 1D potential \(A_{1D}^{\cos}\) with \(2\beta\).

These effects we can further see in the dispersions and densities of states in figures 6 and 7, respectively, which are plotted at \(\mu = 0\) for clarity. In figure 6 we show the cross section of the dispersions in figure 3 along the line (0, \(k_y\)) [1D potentials] or \(k(1, 1)\) [2D potential], both in the normal and superconducting states, and in the different schemes to allow for easier comparison between the 1D and 2D potentials. In figure 7 we show the corresponding densities of states (DOS). We clearly see in the normal state how increasing \(\beta\) and \(c\) both suppress the group velocity, thus increasing flatness of the bands. The density of states becomes correspondingly more and more peaked at zero energy. The superconducting energy gap also increases both with increasing \(\beta\) and \(c\), and the peculiar double-peak structure in the superconducting DOS is also better revealed for higher \(\beta\) or \(c\). In the 2D case it is notable how increasing \(\beta\) generates multiple peaks in the normal state DOS, and thus also in the superconducting state DOS, in a way that separates it from the 1D potentials.

To determine more properties that could be measured e.g. by STM [21, 56], we show in figure 8 the local densities of states (LDOS) along the line (\(x, 0\)) [1D potentials] or \((1, −1)\) [2D potential], which further illustrate the results discussed so far. In the normal state the overall energy dependence shows the clear peak at zero energy for the 1D potentials, as well as the multiple-peak structure for the 2D potential. In the superconducting state the energy dependence also shows the superconducting gap, as already seen in the total DOS in figure 7. The position dependence gives us more information about the underlying strain field. They clearly show the high density of low-energy states near the points \(x = ±d/4\) (1D potentials) or \(r = ±d/4\) (2D potential), that is, points where \(B\) has extrema. Furthermore the states on the positive (negative) \(x\) or \(r\) side are those coming from the sublattice \(A\) (\(B\)), which, by comparison to figures 2(c) and (d), means that the \(A\) (\(B\)) sublattice states are localized at the minima (maxima) of \(B\). This kind of localization and sublattice polarization was also experimentally observed by Jiang et al [21]. Since the low-energy states are the ones contributing to superconductivity,
Figure 7. Effect of the strain strength $\beta$ and the slope parameter $c$ on the density of states (DOS) at $\mu = 0$, $T = 0$, and $\lambda/(\hbar v_F d) = -0.01$ (normal state: solid, superconducting state: dashed lines) for (a) $A_{1D}^{\cos}$, (b) $A_2^{1D}$, and (c) $A_{2D}^{\cos}$. For clarity the successive curves in the DOS plots are shifted vertically by 15 in (a, c) and by 10 in (b). Each curve is normalized such that the shown area integrates to unity.

their localization explains the similar localization of the order parameter $\Delta_{A/B}$, as seen in figures 2(e) and (f).

In the normal state LDOS we further see the localization pattern splitting at higher energies for the 1D potentials. This is contrasted with the 2D potential, where the higher-energy peaks are separated not only in position but also in energy. Furthermore in the superconducting state LDOS we see the same behavior in the energy gaps as in the total DOS: increasing $\beta$ or $c$ leads to an increasing gap size, with the localization pattern staying the same. Again the 2D potential behaves slightly differently: the gap is largest at $r = \pm d/4$, while for the 1D potentials the gap at $x = \pm d/4$ is smallest.

3.2. Critical doping level and temperature

We can in principle calculate the critical doping level $\mu_c$ and the critical temperature $T_c$ by solving the self-consistency equation (19) for various $\mu$ and $T$ and by solving for the point where $\Delta$ vanishes. But since the fixed-point iteration scheme converges slowly when $\Delta$ is small, we calculate $\mu_{50\%}$ [$T_{50\%}$] instead, corresponding to the chemical potential [temperature] at which $\Delta$ has decreased to $\Delta(\mu = 0)/2$ [$\Delta(T = 0)/2$].

We show in figure 9(a) the $\mu$-dependence of $\Delta$ at $T = 0$ in the case of $A_{1D}^{\cos}$, from where $\mu_{50\%}$ is determined. We see how doping away from the flat band, which in the flat-band regime is located at zero energy, kills superconductivity. In this case $\mu_{50\%}$ approaches $\sim 0.7$ max $\Delta(\mu = 0)/2$ in the

Figure 8. Local density of states (LDOS) at $\mu = 0$ and $T = 0$ along the line $(x, 0)$ [1D potentials] or $(r, 1)$ [2D potential] both in the (a, c, e, g, i) normal and (b, d, f, h, j) superconducting (SC) states. In the superconducting state $\lambda/(\hbar v_F d) = -0.01$. In each plot the states on the positive (negative) $x$ or $r$ side come from the sublattice $A$ ($B$). Each plot is normalized such that the total visible area integrates to unity.
at zero doping. Perservative regimes so that superconductivity would be observed in PSG we can in principle tune PSG if the interaction strength \(a\) non-zero doping level \([5]\), and the same might happen also in DOS because of the small energy scale of \(\epsilon\) ene gies flat-band regime, kills superconductivity. The behavior is the same for \(\mu\). The four curves for each \(\beta\) are those for \(-\lambda/(\hbar v_F d) = 0.005, 0.01, 0.015, 0.02\) (from top to bottom). (b) The ratio \(\mu_{50%}/\max\Delta(\mu = 0)\) as a function of \(\lambda\) for different potentials \(A\). Each 1D potential has three curves corresponding to \(\beta = 20, 30, 40\) (from top to bottom). In the flat-band regime the ratio tends approximately to a constant as in \((32)\).

In flat-band limit. In the flat-band regime the results fit very well the relation \(\max\Delta(\mu) = \sqrt{(\max\Delta(\mu = 0)^2 - (\mu/b)^2)}\) with \(b\) as the fitting parameter, as compared to the result \(57\) \(\Delta_F(\mu) = \sqrt{\Delta_F^2(\mu = 0) - \mu^2}\) for exactly flat bands and homogeneous \(\Delta_F^2\). On the other hand in the dispersive regime \(\Delta\) is not maximized at zero chemical potential, but around \(\mu \approx 0.9\) max \(\Delta(\mu = 0) \approx 0.9\) max \(\Delta(\mu = \mu_{opt}) \approx 0.02\hbar v_F/\epsilon_d\) instead, which corresponds to the DOS peak position shown in figure \(7a\). This is exactly the same behavior as seen in TBG \([14, 50]\): in the flat-band regime the energy scale of \(\Delta\) exceeds the DOS peak separation (the ‘bandwidth’) and the smeared DOS is centered at zero energy, while in the dispersive regime \(\Delta\) can ‘see’ the double-peaked DOS because of the small energy scale of \(\Delta\). In TBG this might explain \([14, 50]\) why superconductivity is observed at a nonzero doping level \([5]\), and the same might happen also in PSG if the interaction strength \(A\) is small enough. But note that in PSG we can in principle tune \(A\) (its functional dependence, \(\beta, c, d\) and \(d\)) to move the interface between the flat-band and dispersive regimes so that superconductivity would be observed at zero doping.

To further verify that \(\mu_{50%}\) is linear in \(\max\Delta(\mu = 0)\) in the flat-band regime,

\[
\mu_{50%} = \eta \max\Delta(\mu = 0), \quad (32)
\]

we show in figure \(9b\) the ratio \(\eta\) at \(T = 0\) for a selection of potentials. In the flat-band regime the ratio tends approximately to a constant \(\eta \approx 0.7\) as long as \(c \lesssim 10\). For \(c \gtrsim 10\) we start seeing slight deviations from this, with \(\eta \approx 0.6\) and 0.5 for \(c = 20\) and 100, respectively. The critical chemical potential \(\mu_c\) is slightly larger, approximately \(\mu_c \approx \max\Delta(\mu = 0)\) for \(A_{cos}\) in the flat-band regime according to figure \(9a\). This coincides with the case of perfectly flat bands and a constant \(\Delta_F^2\) for which \((58)\), supplemental material of reference \([14]\) \(\mu_c = \Delta_F^2(\mu = 0)\).

In experiments the filling \(\nu\) is more easily directly controlled \([5]\) than the chemical potential \(\mu\), which we use as a parameter. While we do not calculate the filling, hints for the \(\nu(\mu)\) dependence in PSG can be found from the work of Uchoa et al \([16]\) in the case of strained graphene with a homogeneous pseudomagnetic field\(^4\) or from references \([14, 51]\) in the case of TBG.

In figure \(10\) we show the corresponding plots for determining \(T_{50%}\) at \(\mu = \mu_{opt}\). Again the ratio \(\xi\) in

\[
k_b T_{50%} = \xi \max\Delta(T = 0), \quad (33)
\]
tends approximately to a constant \(\xi \approx 0.35\) in the flat-band regime as long as \(c \lesssim 10\). For \(c \gtrsim 10\) we start seeing deviations from this, with \(\xi \approx 0.3\) for \(c = 20\) and \(\xi \approx 0.25\) for \(c = 100\). The critical temperature \(T_c\) is slightly larger, approximately \(k_b T_c \approx 0.4\) max \(\Delta(T = 0)\) for \(A_{cos}\) in the flat-band regime according to figure \(10a\). For comparison, in the case of perfectly flat bands and a constant \(\Delta_F^2\) we have the result \((10)\) \(k_b T_F^2 = \frac{1}{2} \Delta_F^2(T = 0)\) and in TBG \([14]\) within the same interaction model \(k_b T_c \approx 0.25\) max \(\Delta(T = 0)\).

3.3. Superfluid weight and Berezinskii–Kosterlitz–Thouless transition temperature

To determine the true superconducting transition temperature we calculate the superfluid weight \(D^s\) and the Berezinskii–Kosterlitz–Thouless transition temperature \(T_{BKT}\) from \((20)\) and \((21)\). In figure \(11a\) we show the total superfluid weight \(\sqrt{\det D^s}\), together with the different components \(D^s_{\mu}\).

\(^4\) When comparing to Uchoa et al \([16]\), in their language we concentrate on the zeroth pseudo-Landau level, which in figure 1 of Uchoa et al means \(-1 \leq \mu/(\hbar v_F) \leq 1\) or \(-2 \leq \nu \leq 2\).
as a function of the interaction strength $\lambda$ for $A_{1D}^{1D}$. The behavior is very similar to that of max $\Delta$ in figure 4(a); it is linear in the flat-band regime and also increases linearly with increasing $\beta$. To further verify that $\sqrt{\det D}$ is linear in max $\Delta$,

$$\sqrt{\det D} = \chi \max \Delta,$$  

(34)

we show the ratio $\chi$ in figures 11(c) and (d) at $\mu = \mu_{\text{opt}}$ and $T = 0$. In the flat-band regime the ratio tends approximately to a constant $\chi \approx 0.15 \ldots 0.4$, which has more variation than $\eta$ and $\xi$ for $\mu_{\text{50%}}$ and $T_{\text{50%}}$ in the flat-band regime. For comparison, in TBG we found [51] within the same interaction model that $\chi \approx 0.35$ in the flat-band regime.

We may again compare (34) to the case of exactly flat bands and a constant $\Delta_{\text{FB}}$. But since the superfluid weight depends heavily on the Hamiltonian itself and not only its eigenvalues, we need to specify which flat-band model to use. We take the ‘graphene flat-band limit’, that is, graphene with $v_F \to 0$. In this case [44, 48] $D_{\text{FB}} = \frac{1}{2} \Delta_{\text{FB}}$ at $\mu = 0 \approx \mu_{\text{opt}}$ and $T = 0$, which in fact holds for any $v_F$.

What is intriguing in figure 11(a) is that for the studied 1D potentials the superfluid weight is almost isotropic although the potentials are highly anisotropic. There is, however, a slight anisotropy, $D_{\text{xy}} \neq D_{\text{yx}}$, for large $\beta$ and $\lambda$. For the 2D potential $A_{2D}^{1D}$ (not shown) the superfluid weight is isotropic, $D_{\text{xx}} = D_{\text{yy}}$. The off-diagonal components $D_{\text{xy}} = D_{\text{yx}} = 0$ are zero for all the studied potentials. The (an)isotropy is consistent with the symmetries of the studied potentials. In (a) both potentials have three curves corresponding to $-\lambda/(\hbar v_F d) = 0.01, 0.015, 0.02$, from bottom to top. Also the dashed line $\sqrt{\det D} = 8k_B T/\pi$ is shown, from intersections of which $T_{\text{BKT}}$ is determined through (21). (c, d) The ratio $\sqrt{\det D}/\max \Delta$ as a function of the interaction strength $\lambda$ for (c) the cosine potentials and (d) $A_{1D}^{1D}$ with varying $c$. In (c) each $A$ has three curves corresponding to (from top to bottom) $\beta = 20, 30, 40$ (1D potentials) or $\beta = 10, 15, 20$ (2D potential), while in (d) each $A$ has two curves corresponding to (from top to bottom) $\beta = 30, 40$. In the flat-band regime the ratio is approximately a constant depending slightly on the potential, as in (34).

Figure 11. Behavior of the superfluid weight $D^*$ at optimal doping $\mu = \mu_{\text{opt}}$ and (a, c, d) $T = 0$. (a) $D^*$ as a function of $\lambda$ for $A_{1D}^{1D}$ showing linearity in the flat-band regime. The superfluid weight for the 1D potentials is slightly anisotropic, $D_{\text{xy}} \neq D_{\text{yx}}$, for large $\beta$ and $\lambda$. For the 2D potential $A_{2D}^{1D}$ (not shown) the superfluid weight is isotropic, $D_{\text{xx}} = D_{\text{yy}}$. The off-diagonal components $D_{\text{xy}} = D_{\text{yx}} = 0$ are zero for all the studied potentials. The (an)isotropy is consistent with the symmetries of the studied potentials. In (a) both potentials have three curves corresponding to $-\lambda/(\hbar v_F d) = 0.01, 0.015, 0.02$, from bottom to top. Also the dashed line $\sqrt{\det D} = 8k_B T/\pi$ is shown, from intersections of which $T_{\text{BKT}}$ is determined through (21). (c, d) The ratio $\sqrt{\det D}/\max \Delta$ as a function of the interaction strength $\lambda$ for (c) the cosine potentials and (d) $A_{1D}^{1D}$ with varying $c$. In (c) each $A$ has three curves corresponding to (from top to bottom) $\beta = 20, 30, 40$ (1D potentials) or $\beta = 10, 15, 20$ (2D potential), while in (d) each $A$ has two curves corresponding to (from top to bottom) $\beta = 30, 40$. In the flat-band regime the ratio is approximately a constant depending slightly on the potential, as in (34).

Figure 12. Behavior of the Berezinskii–Kosterlitz–Thouless transition temperature $T_{\text{BKT}}$ at optimal doping $\mu = \mu_{\text{opt}}$. The ratio $k_B T_{\text{BKT}}/\max \Delta(T = 0)$ for (a) the cosine potentials and (b) $A_{1D}^{1D}$ with varying $c$. In (a) both potentials have three curves corresponding to $\beta = 20, 30, 40$ (1D potential) or $\beta = 10, 15, 20$ (2D potential), from top to bottom in the flat-band regime, while in (b) each $c$ has two curves corresponding to $\beta = 30, 40$, from top to bottom. In the flat-band regime the ratio tends approximately to a constant depending slightly on the potential, as in (35).

Although in this work we do not separate the superfluid weight into the conventional and geometric contributions [44], from general knowledge [44] and calculations in TBG [51, 60] we expect the geometric contribution to dominate in the flat-band regime.

In figure 11(b) we further show $\sqrt{\det D}$ as a function of temperature $T$ for $A_{1D}^{1D}$, from where $T_{\text{BKT}}$ is determined through (21) by solving for the intersection point with the line $\sqrt{\det D} = 8k_B T/\pi$. We immediately see that in the flat-band regime $D^*(T_{\text{BKT}}) \approx D^*(0)$ is a rather good approximation so that the self-consistency in (21) is not essential. This is very different from TBG [51], where the temperature dependence is essential due to $T_{\text{BKT}}$ being closer to $T_c$. We nevertheless need to solve the full self-consistent equation for all the potentials, as the relative magnitude of $T_c$ and $T_{\text{BKT}}$ is not known beforehand.

The resulting ratio $k_B T_{\text{BKT}}/\max \Delta(T = 0)$ is shown in figure 12 for the different potentials at $\mu = \mu_{\text{opt}}$, further confirming that $D^*(T_{\text{BKT}}) \approx D^*(0)$: apart from the different scale, the $T_{\text{BKT}}$ plots in figure 12 are very similar to the $D^*$ plots in figures 11(c) and (d). Furthermore in the linear relation

$$k_B T_{\text{BKT}} = \kappa \max \Delta(T = 0),$$

(35)
the ratio $\kappa$ tends approximately to a constant $\kappa \approx 0.05 \ldots 0.15$ in the flat-band regime. Again in (35) we see similarity to the ‘graphene flat-band limit’ result with a homogeneous $\Delta_{FB}$, for which $k_B T_{FB(\text{BKT})} = \frac{\pi}{4} D_{FB}(T_{FB(\text{BKT})}) \approx \frac{1}{4} \Delta_{FB}(T=0) \approx 0, \mu = 0 \approx \mu_{opt}$ if we furthermore assume $D_{FB}(T_{FB(\text{BKT})}) \approx D_{FB}(0)$.

Combining (34) and (35) we get in the flat-band regime at $\mu = \mu_{opt}$ the ratio $T_{FB(\text{BKT})}/T_{50\%}(1D \text{ FB}) \approx \frac{\kappa}{\xi} 0.2 \ldots 0.4$ depending on the potential. For $A_{1D}$ this yields $T_{FB(\text{BKT})}/T_{50\%}(1D) \approx 0.4$, and within the same accuracy $T_{FB(\text{BKT})}/T_{c} \approx 0.4$. For comparison in TBG we found within the same interaction model in the flat-band regime $k_B T_{FB(\text{BKT})} \approx 0.16 \ldots 0.2 \max \Delta(T=0)$ [51] (depending slightly on $\lambda$), $k_B T_{c} \approx 0.25 \max \Delta(T=0)$ [14], and thus $T_{FB(\text{BKT})}/T_{c} \approx 0.6 \ldots 0.8$.

By combining (31) and (35) we get $k_B T_{FB(\text{BKT})} = -\kappa \mathcal{B} \lambda$ at $\mu = \mu_{opt}$. Let us calculate an estimate of $T_{FB(\text{BKT})}$ by using $\lambda = -1eV a^2 \approx -6 \text{ eV } \text{ Å}^2$, which roughly corresponds [14, 51] to $T_{FB(\text{BKT})} \approx 1 \text{ K}$ measured in TBG [5]. Here $a = \sqrt{3} \text{Å} \approx 2.46 \text{Å}$ is the graphene lattice constant. For $A_{1D}$ we have $\mathcal{B} = 2 \pi \beta/d^2$ and in the flat-band regime $\kappa \approx 0.15$ and $\zeta \approx 0.15$, yielding a similar $T_{FB(\text{BKT})} \approx 1 \text{ K}$ if we apply strain for example such that $\beta = 40$ and $d = 60 \text{ nm}$ [then $\lambda/(hv_{\text{FD}}) \approx -0.002$ if using $v_{\text{FD}} = 1 \times 10^4 \text{ m s}^{-1}$, which is in the flat-band regime according to figures 4(b) and 12(a)]. In the case of the in-plane displacement field $u_{\text{1D}}(\text{cos})$ (28) this corresponds to the displacement amplitude $\beta_{\text{in}}/(\beta_{\text{1D}} G) \approx 1 \text{ nm}$ if $\beta_{\text{G}} = 2$.

Since in this case the elasticity theory assumes $\beta_{\text{G}} \ll d/a_0$ and $d/a_0 \gg 1$, we are very well in the validity regime. On the other hand, if we are able to decrease the strain period to $d = 10 \text{ nm}$ [then $\lambda/(hv_{\text{FD}}) \approx -0.0009$], we get to a high-temperature superconductor value of $T_{FB(\text{BKT})} \approx 40 \text{ K}$, which is still in the validity regime. Note the optimization problem in increasing $T_{FB(\text{BKT})}$: decreasing $d$ directly enhances $T_{FB(\text{BKT})}$ but at the same time it makes the validity limit for $\beta$ too tight, while at the same time we have to increase $\beta$ as possible. But this might only be a limiting factor in our linear elasticity theory, while a more complete microscopic theory could, perhaps, yield a result that increasing $\beta$ or decreasing $d$ always increases $T_{FB(\text{BKT})}$.

The experiments of Jiang et al [21] with $2B \approx 100 \text{ T}$ and $d = 14 \text{ nm}$ can be described by the 1D cosine potential with $\beta \approx 5$. When $\lambda = -6 \text{ eV } \text{ Å}^2$, $\lambda/(hv_{\text{FD}}) \approx -0.007$ is not in the flat-band regime. Hence $T_{FB(\text{BKT})}$ cannot be obtained from the simple estimate used above, and is likely much lower than 1 K. Increasing the strain amplitude by a factor of 4, so that $\beta = 20$, would yield $\zeta \approx 0.05$, $\kappa = 0.17$, and thus $T_{FB(\text{BKT})} \approx 0.007 h v_{\text{FD}}/k_B$ $\approx 4 \text{ K}$. Further decreasing the period to $d = 8 \text{ nm}$, a period which was already observed by Jiang et al, would yield already $\lambda/(hv_{\text{FD}}) \approx -0.01$ and thus $T_{FB(\text{BKT})} \approx 11 \text{ K}$.

4. Conclusions

We have studied both the normal and superconducting s-wave state properties of periodically strained graphene (PSG) in the continuum low-energy model. We have shown that periodic strain might be a mechanism that allows increasing the critical temperature $T_c$ higher than a few kelvin, observed in doped graphene and in twisted bilayer graphene (TBG), or possibly even to tens of kelvins. Especially we have generalized the results of Kauppila et al [19], where the authors studied the same problem in the case of a 1D cosine-like pseudo vector potential $\mathcal{A}$, to potentials with arbitrary shape and dimension. We furthermore calculated the superfluid weight and the Berezinskii–Kosterlitz–Thouless transition temperature $T_{\text{BKT}}$ to determine the true transition temperature observed in experiments. In the normal state we observed flat bands in the spectrum and localization of low-energy states near the extremum points of the effective pseudomagnetic field $B = \nabla \times \mathcal{A}$.

We modeled the superconducting state by the Bogoliubov–de Gennes mean-field theory assuming a local interaction between the Cooper pairs leading to s-wave pairing. Because of the inhomogeneous strain field we observed a highly inhomogeneous order parameter $\Delta_{\mathcal{A}/B}$ that is localized near the extremum points of $B$, similarly to the localization of the low-energy states. We also noticed how the superconducting $T_c$ or $T_{\text{BKT}}$ can be linearly increased by increasing the strain strength $\beta$, decreasing the period $d$, or by increasing the slope (near the extremum points of $B$) of the corresponding pseudo vector potential $\mathcal{A}$. On the other hand increasing the slope makes the order parameter also more localized.

While between the 1D potentials we observed only quantitative differences in the results, for the 2D cosine potential we saw also some qualitative differences when compared to the 1D potentials. The main differences are the localization pattern of $\Delta_{\mathcal{A}/B}$ and the more peaked structure of the (local) density of states both in the normal and superconducting states. In the 2D case we studied only the cosine potential, but on the other hand the qualitative similarity in the results between the different 1D potentials gives us certainty that changing the shape of the potential would not change the qualitative results in the 2D case neither. However, it should be noted that it is the shape of $\mathcal{B}$ that matters and not that of the potential $\mathcal{A}$ itself, and thus even a 2D potential can produce results that are effectively those of a 1D potential.

We chose all our potentials to be periodic in a square (super)lattice, but note that any other lattice could be chosen as well, with different shapes and different periodicities in the two directions. Properties of this lattice are then directly seen in the dispersion, as well as in the localization of $\mathcal{B}$ and $\Delta_{\mathcal{A}/B}$. We also observed the symmetry $\Delta_{\mathcal{B}}(r) = \Delta_{\mathcal{A}}(-r)$ of the order parameter for all the chosen potentials. This is due to the inversion symmetry $A(r) = A(-r)$ present in all of them. The relative magnitude between $\Delta_{\mathcal{A}}$ and $\Delta_{\mathcal{G}}$ can then be tuned by breaking this symmetry, e.g. by using a sawtooth-wave potential.

We also observed some very peculiar structures in the (local) density of states, which could serve as an experimental fingerprint of the physics described by this model. We furthermore found that in the flat-band regime the superconducting order parameter maximum max $\Delta$ at $\mu = \mu_{opt}$ and $T = 0$, the ‘critical’ chemical potential $\mu_{opt}$, at $T = 0$, the ‘critical’ temperature $T_{50\%}$ at $\mu = \mu_{opt}$, the superfluid weight $\sqrt{\text{det} D}$ at $\mu = \mu_{opt}$ and $T = 0$, and the BKT transition temperature $T_{\text{BKT}}$ at $\mu = \mu_{opt}$ are all approximately linear in the interaction.
strength \( \lambda \). The linear relations, instead of exponential ones in usual bulk superconductors, suggest that high-temperature superconductivity might be possible in PSG.

As is known from the closely-related materials twisted bilayer graphene [5, 7, 11, 61–63], twisted trilayer graphene [64], twisted double bilayer graphene [65–67], rhombohedral graphite [68], or other graphite-based composites [69–73], other phases like correlated insulators might be present. These are obviously excluded from the present study, but as we showed in previous studies [14, 51], the superconductivity-only model gives a plausible explanation for the observed superconducting states in TBG. This view of competing phases is supported by recent experiments where superconductivity could be seen without the correlated insulating phases [62, 63]. Thus we expect our similar model to work also in PSG when concentrating only on superconductivity. If the competing phase (if any) is magnetic, we know from a recent study [74] that in a pure flat-band system superconductivity is favored over magnetism whenever (in the weak-coupling regime) the effective attractive electron–electron interaction strength \( \tilde{\lambda}t_Hc = \frac{g^2}{\Omega_{FB}/\Omega_{BG}} \) is stronger than the repulsive one \( u = U\Omega_{FB}/\Omega_{BG} \). Here \( g \) is the electron–phonon coupling constant, \( U \) is the repulsive Hubbard coupling constant, \( \tilde{\lambda}_c \) is the characteristic phonon energy (in this case the Einstein energy \( \hbar\omega_c \)), and \( \Omega_{FB}/\Omega_{BG} \) is the ratio of the flat-band area to the Brillouin zone area. The competition between superconductivity and magnetism was studied also by Löthman & Black-Schaffer [58], who showed how the two phases in the flat-band regime have somewhat different dependence on the doping.

An interesting future prospect would be to study the other phases which, by the analogue of TBG, are highly probable. Secondly the combination of moiré [14, 51] and strain [this work] physics would perhaps advance the understanding of superconductivity in TBG, where intrinsic periodic strain is inevitable. Thirdly, generalizing the Eliashberg formalism [74, 75] to the case of PSG with inhomogeneous superconductivity would make handling the electron–phonon interaction more accurate, especially in the strong-coupling limit. From the experimental point of view the challenge is to manufacture periodically strained graphene samples with large amplitudes and small periods and to perform low-temperature conductivity measurements in this (electrically doped) system to reveal the possible superconducting and/or correlated insulator states. The periodic strain and flat bands observed by Jiang et al [21] are already an intriguing starting point, but according to our calculations a \( T_{BKT} \) of the order of 4 K would need a strain amplitude 4 times larger than in the experiment. On the other hand, further decreasing the period to 8 nm would yield already \( T_{BKT} \approx 11 \) K.

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