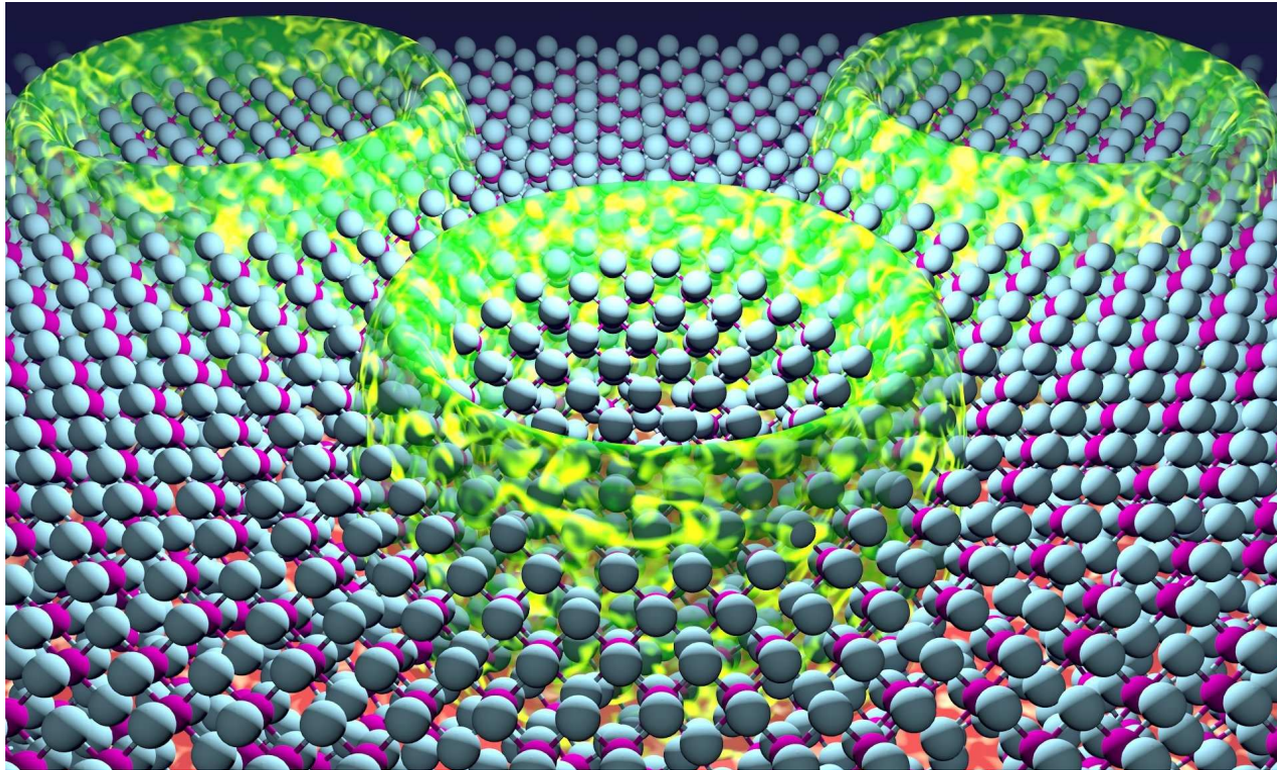




# Moire electronic states and twisted van der Waals heterostructures



# Schedule for the lecture

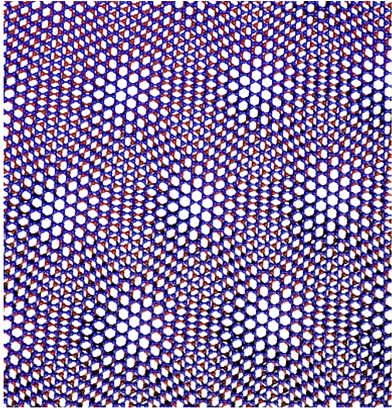
- 40 min lecture
- 15 min break
- 40 min lecture
- 15 min break
- 40 min lecture

# Today's plan

- Moire and quasiperiodicity
- Band structure folding, unfolding and minibands
- Correlations in moire electronic structures
- Topology in moire systems
- Twisted graphene multilayers

# Moire materials

Twisted graphene multilayers

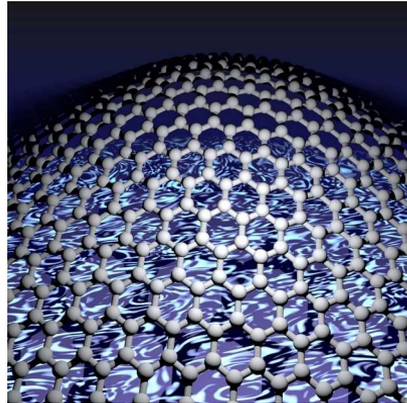


Moire states in several layers

graphene

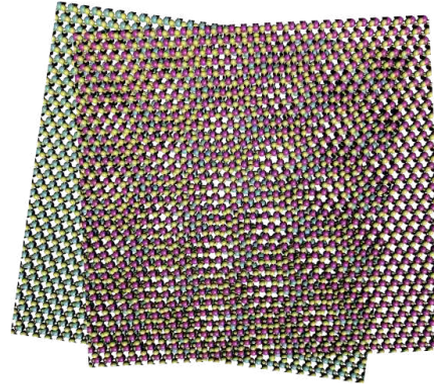
***Moire in electronic properties***

Buckled 2D materials



Moire states in single layer

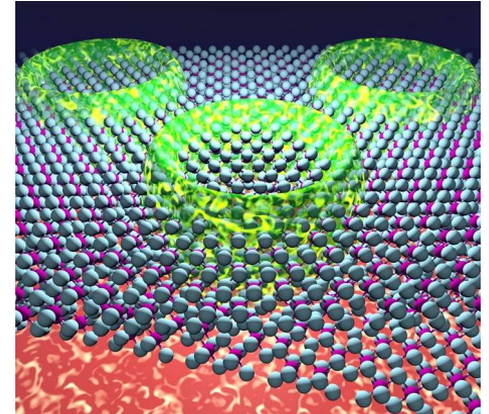
Twisted TMDCs



MoS<sub>2</sub>, WSe<sub>2</sub>  
Moire states in single/several layer

***Moire in magnetic properties***

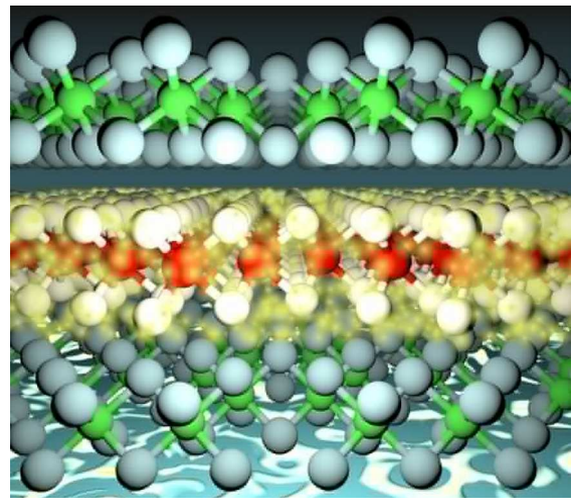
Twisted magnetic 2D materials



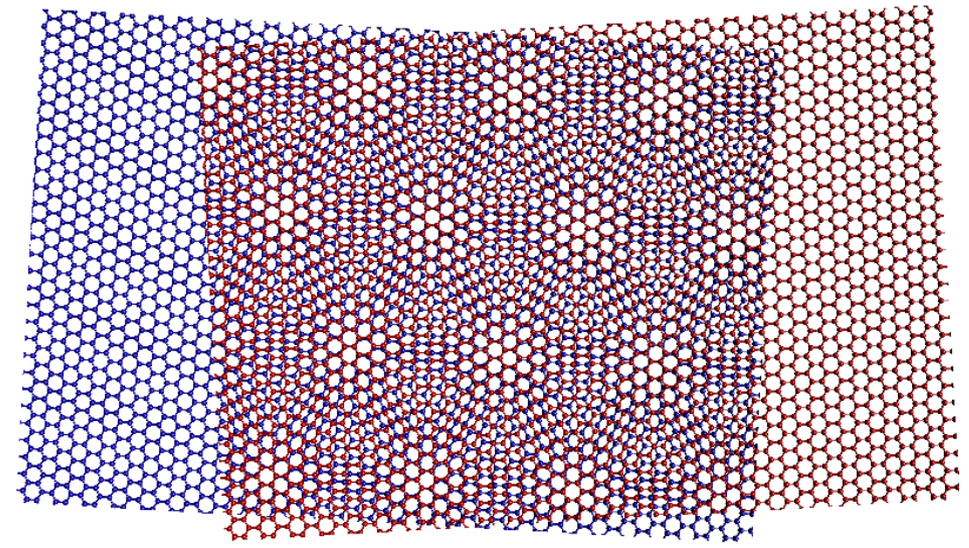
CrCl<sub>3</sub>, CrBr<sub>3</sub>  
Moire magnetism

# How to create moire states with 2D materials

## Stacking



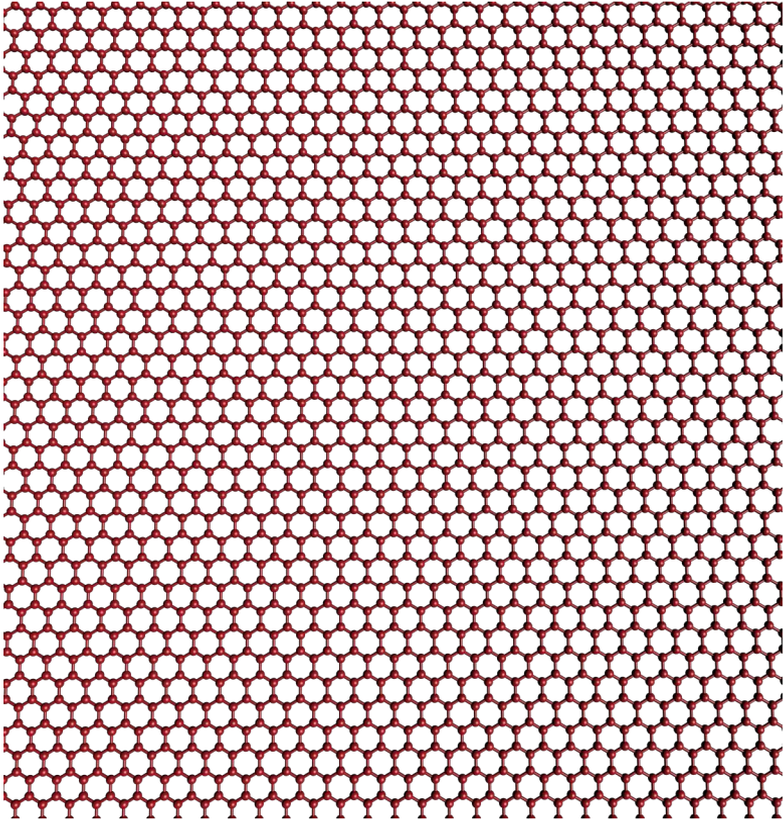
## Rotating



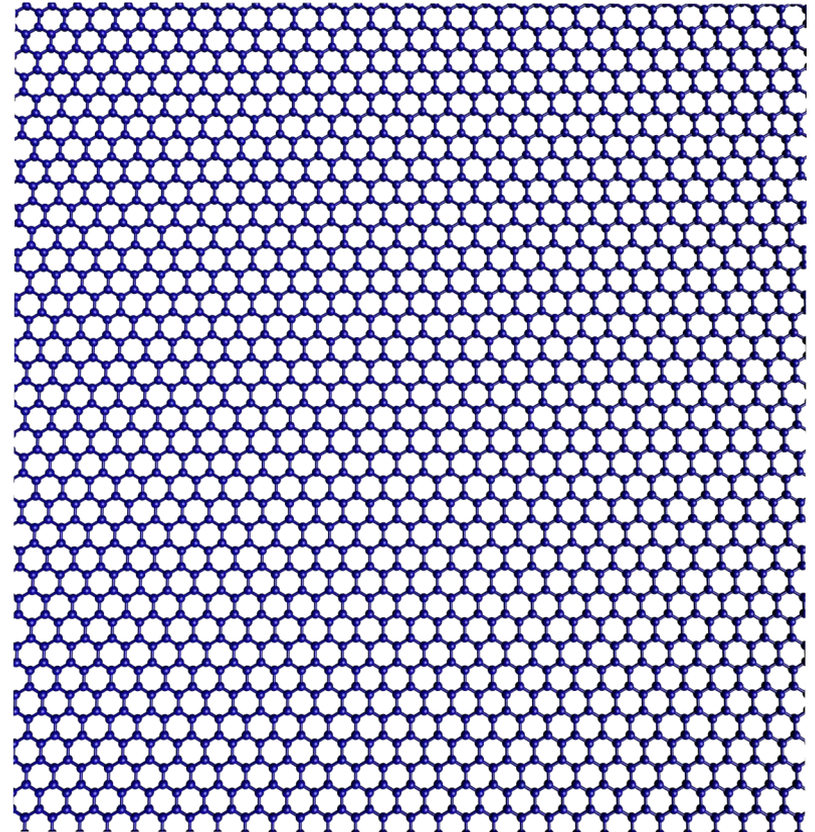
**These are unique features of two-dimensional materials**

# A bilayer van der Waals heterostructure

**Upper graphene layer**

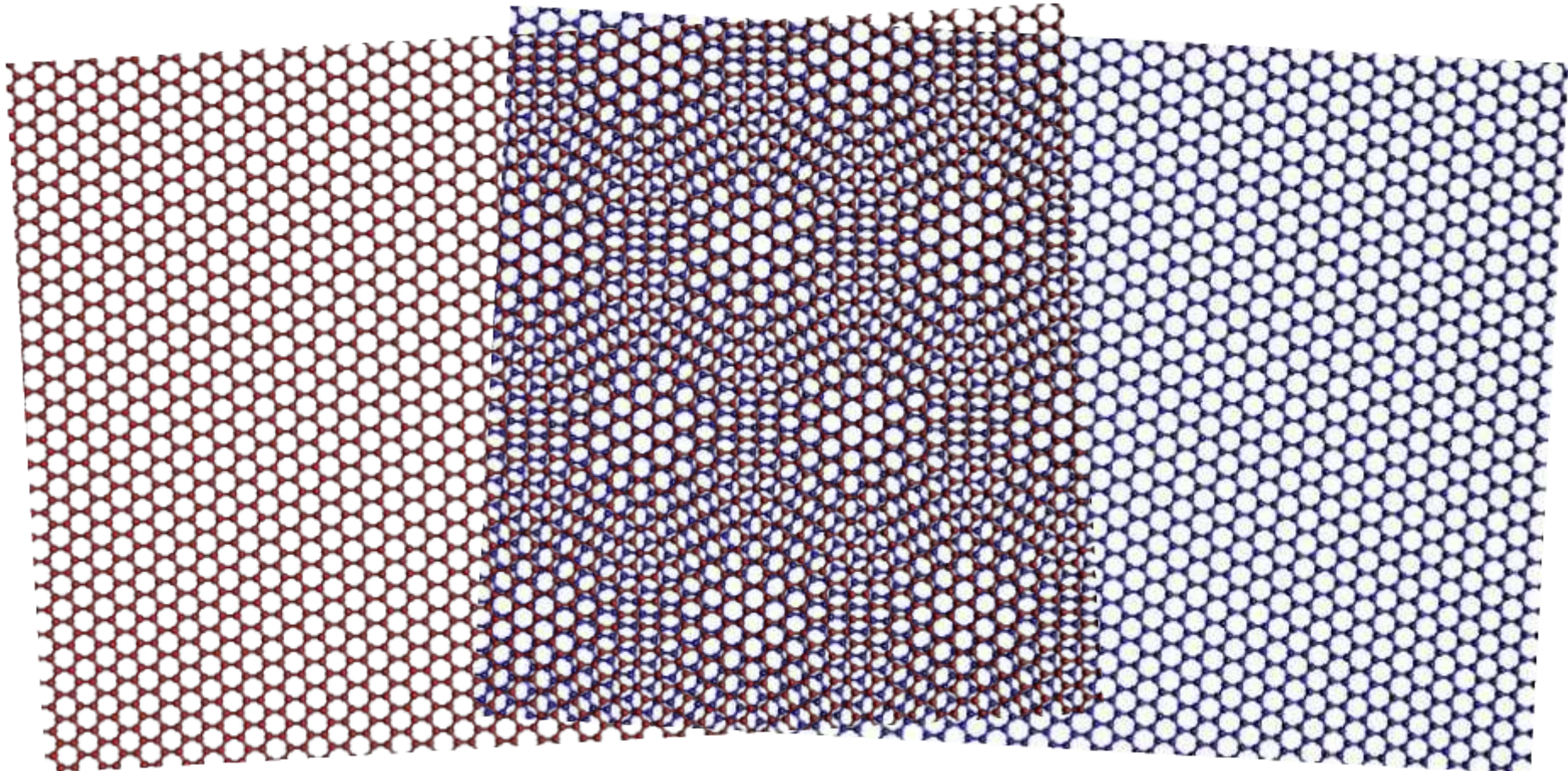


**Lower graphene layer**

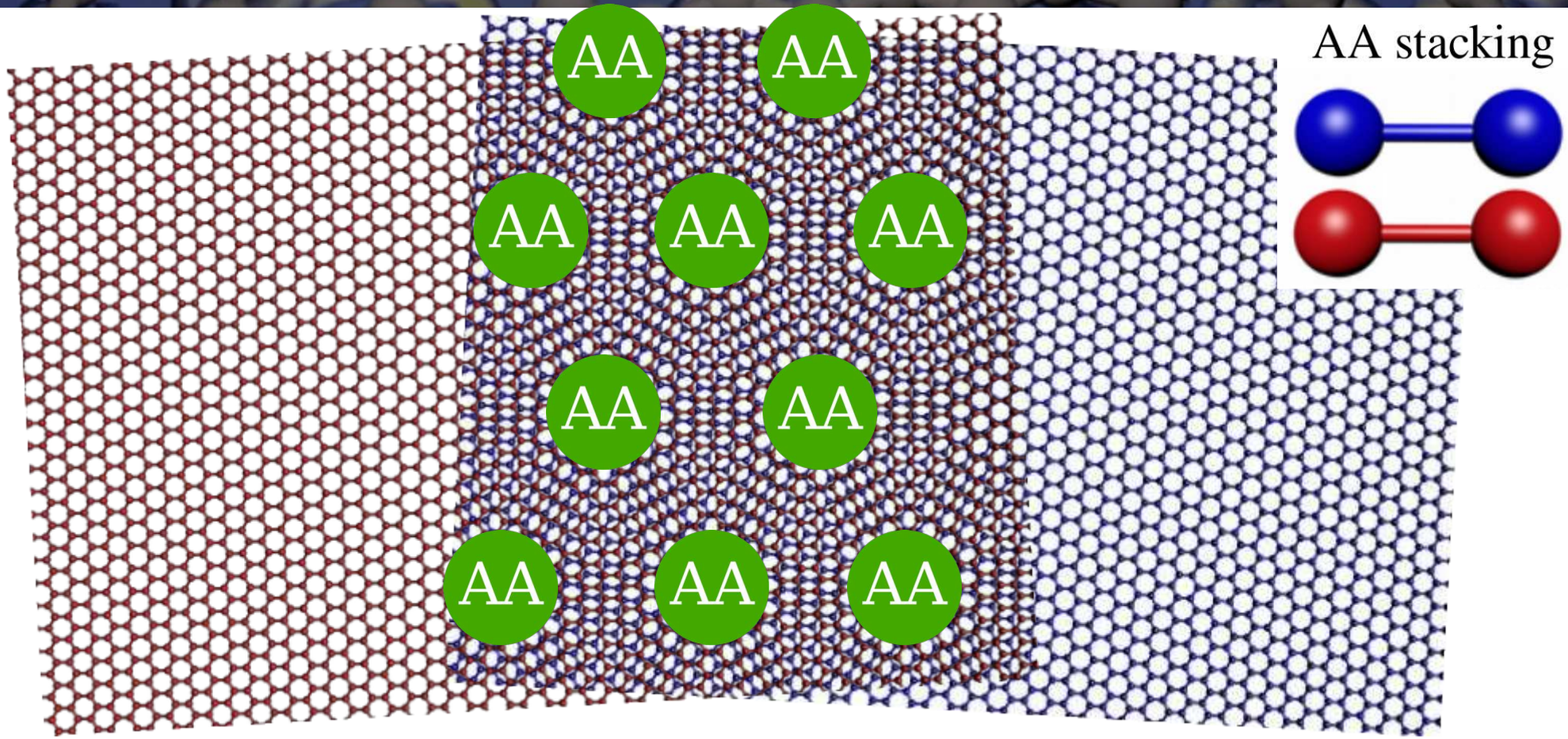




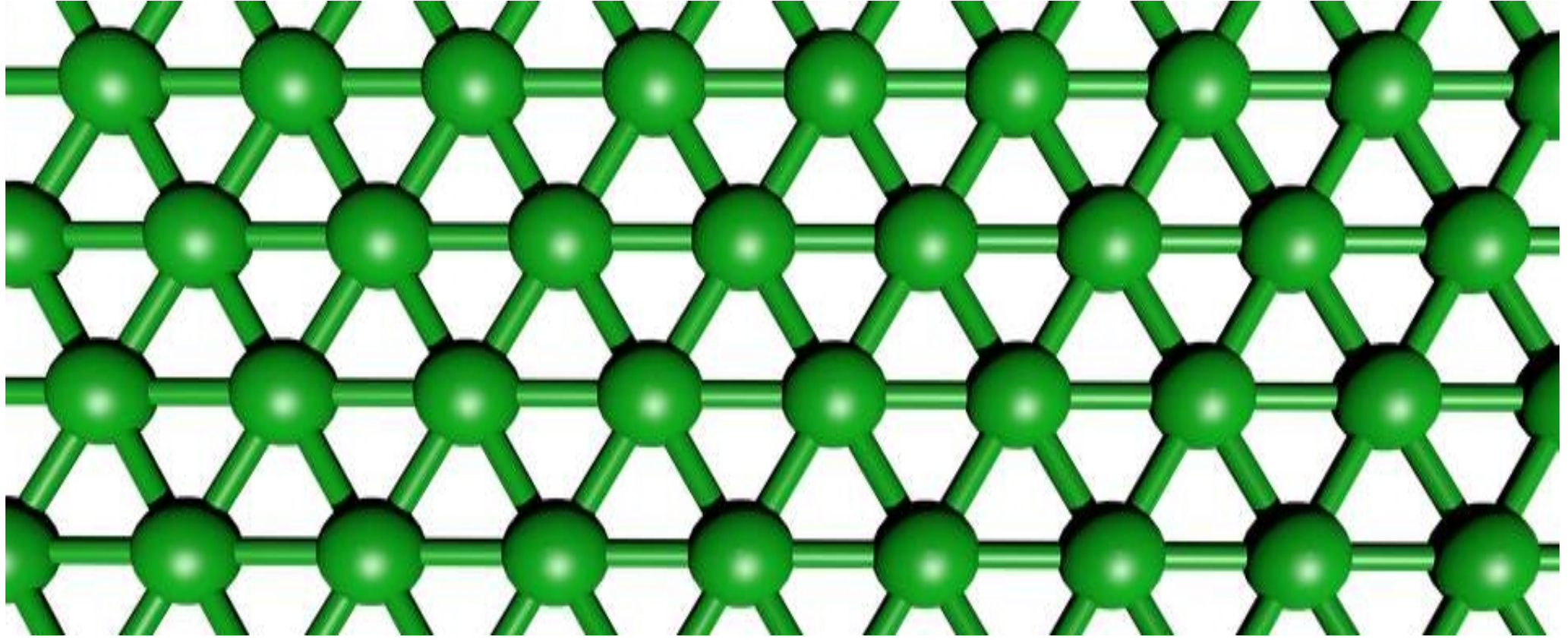
# A bilayer van der Waals heterostructure



# A bilayer van der Waals heterostructure

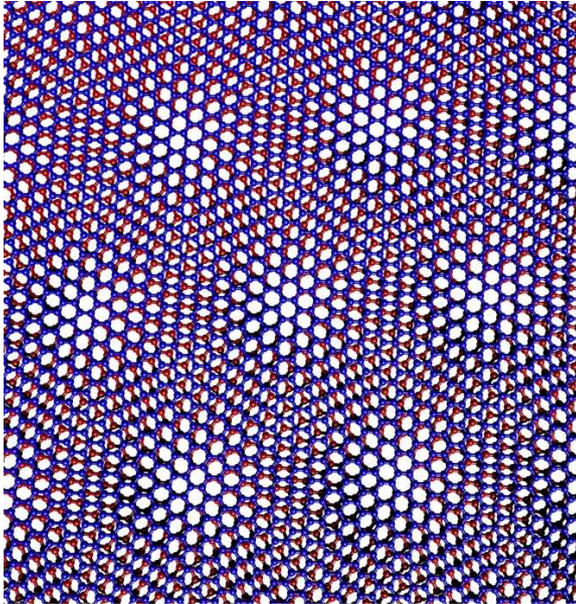


# A bilayer van der Waals heterostructure

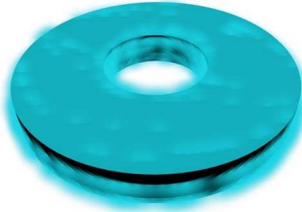


# Electronic states in a single moire superlattice

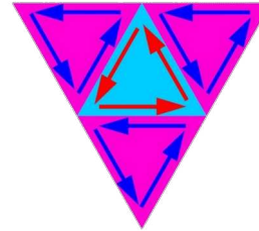
## Twisted bilayer graphene



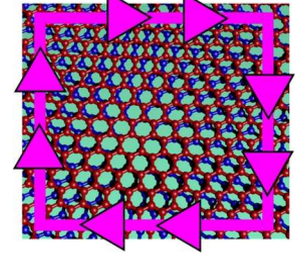
### Superconductivity



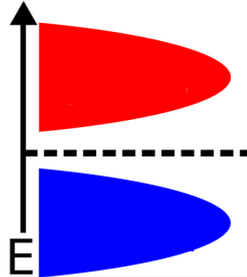
### Topological networks



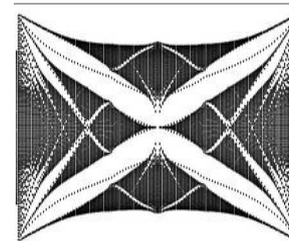
### Chern insulators



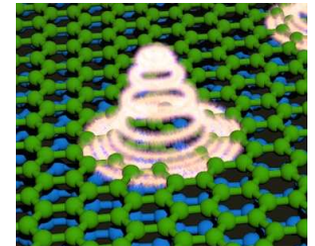
### Correlated insulators



### Quasicrystalline physics



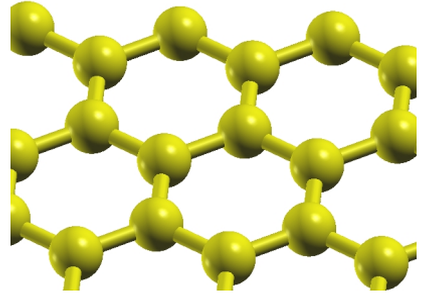
### Fractional Chern insulators



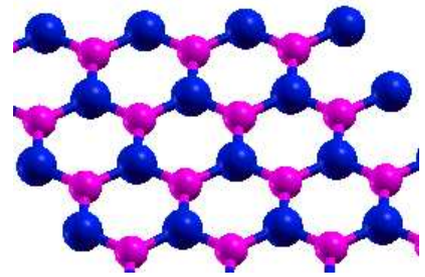
*A single twisted van der Waals material realizes a variety of widely different electronic states*

# The building blocks for twisted van der Waals heterostructure

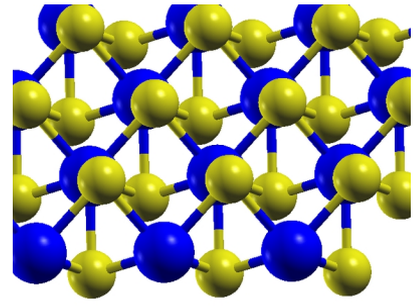
**Semimetal**  
Graphene



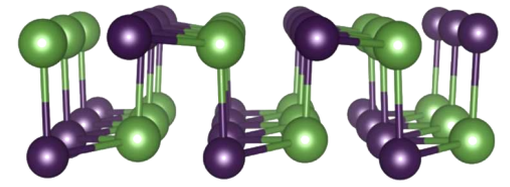
**Insulator**  
BN



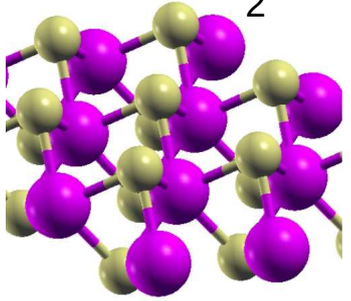
**Superconductor**  
NbSe<sub>2</sub>



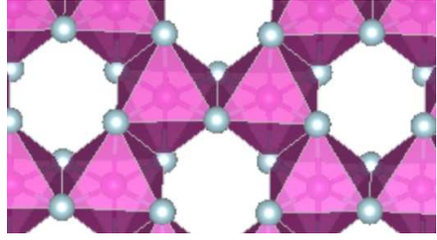
**Ferroelectric**  
SnTe



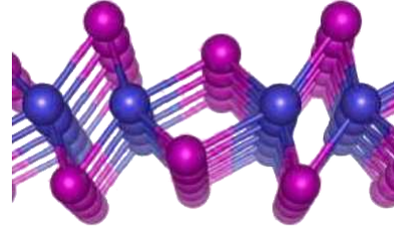
**Semiconductor**  
WSe<sub>2</sub>



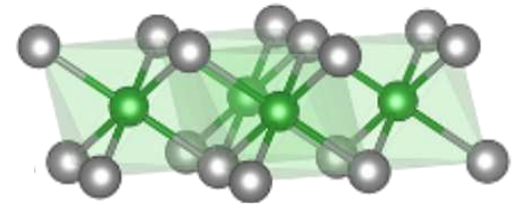
**Ferromagnet**  
CrI<sub>3</sub>



**Quantum spin Hall insulator**  
WTe<sub>2</sub>

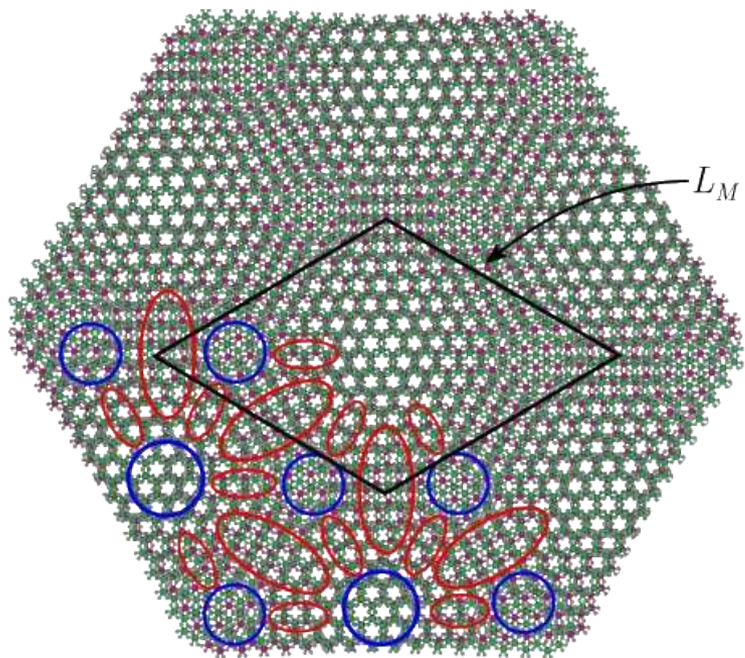


**Multiferroic**  
NiI<sub>2</sub>

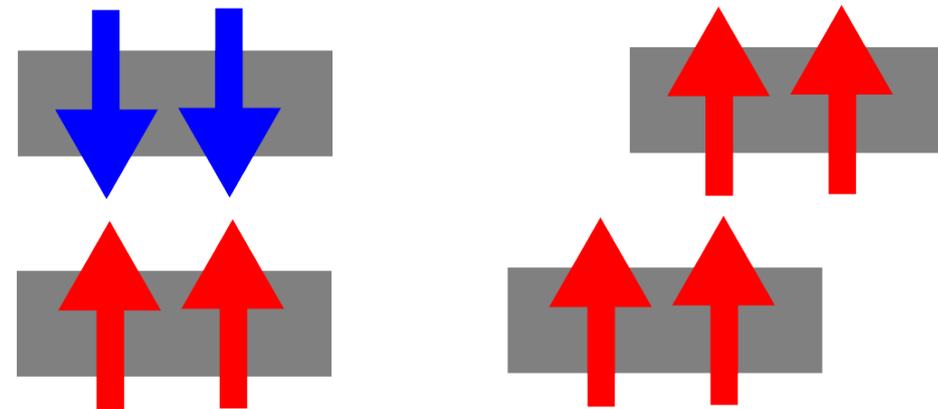


# Twisted 2D magnetic materials

$\text{CrCl}_3$ ,  $\text{CrBr}_3$



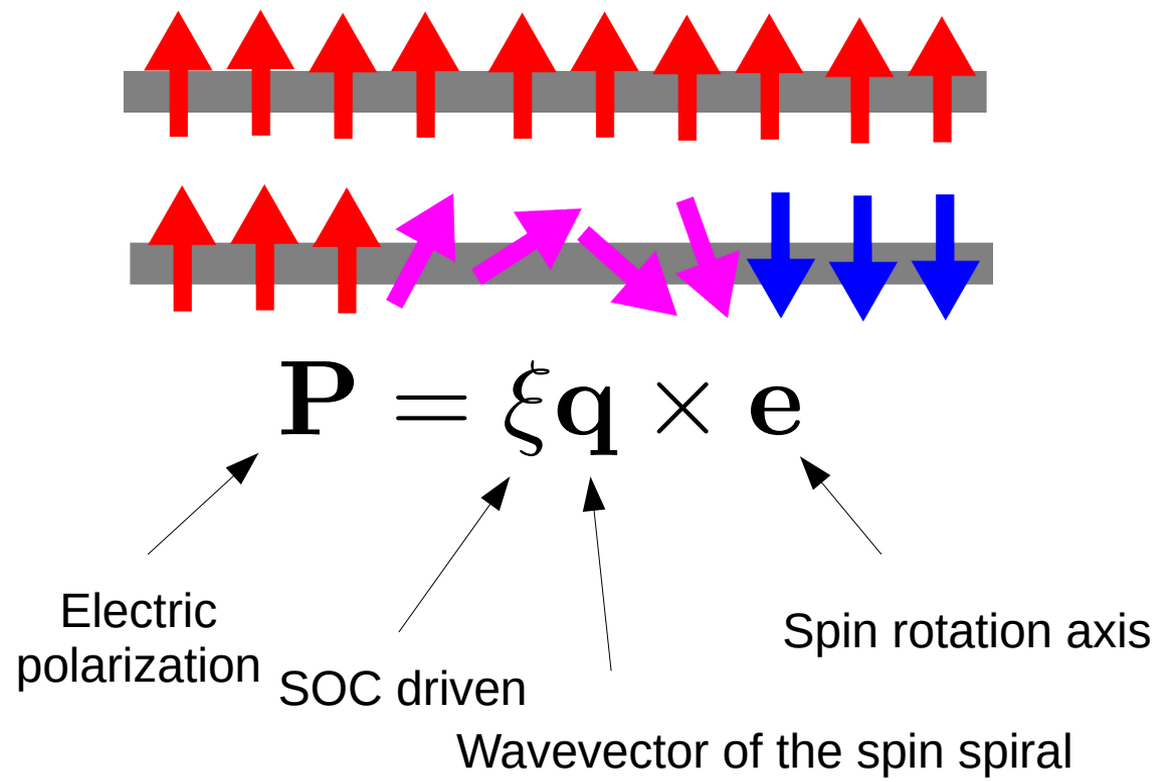
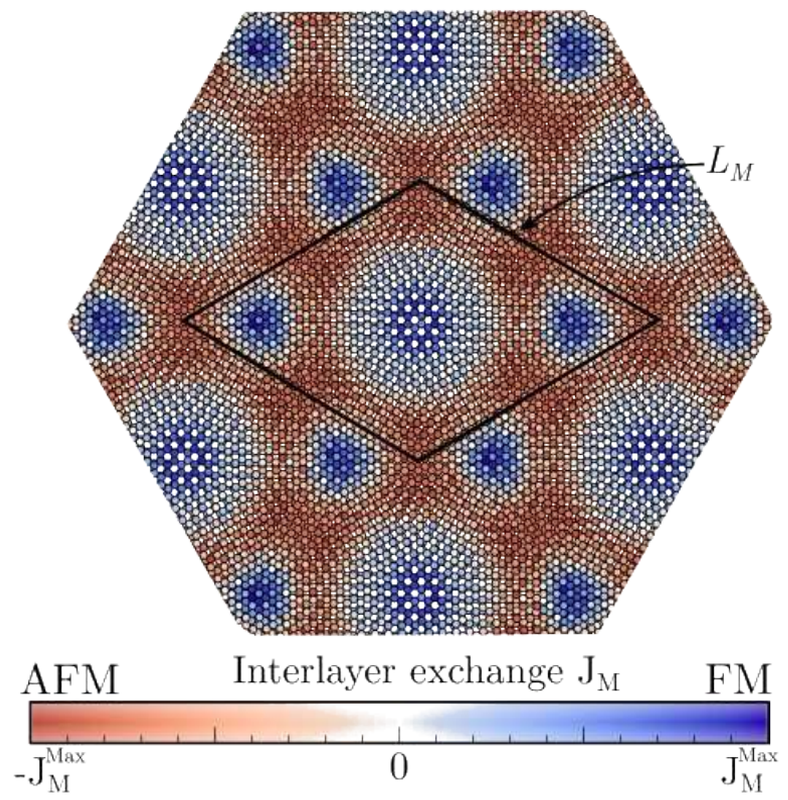
Bilayer Stacking  $\left\{ \begin{array}{l} \text{(M)} \text{ Monoclinic} \\ \text{(R)} \text{ Rhombohedral} \end{array} \right.$



The local stacking determines the coupling between layers

# Twisted 2D magnetic materials

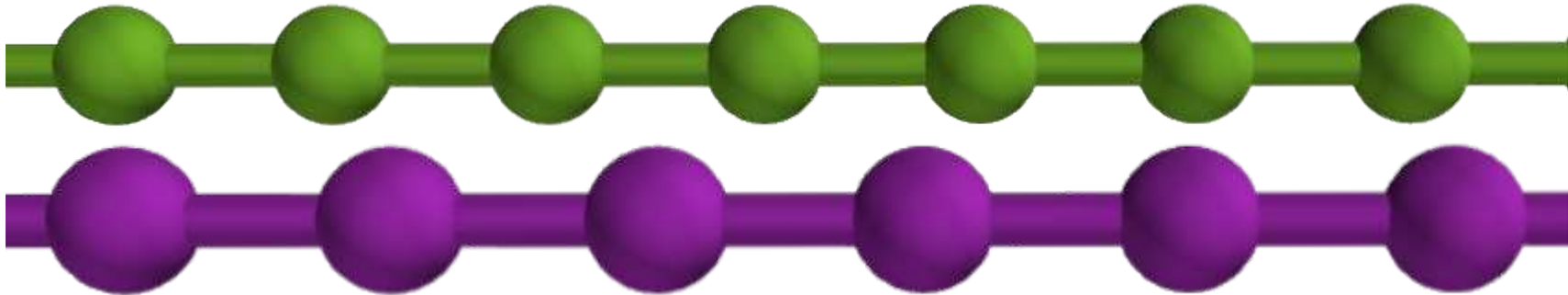
Non-collinear magnetism and multiferroic order appear due to the moire



# Superpotentials and quasiperiodicity

# A minimal moire potential

Let us now take a one dimensional superlattice



We have now two length scales

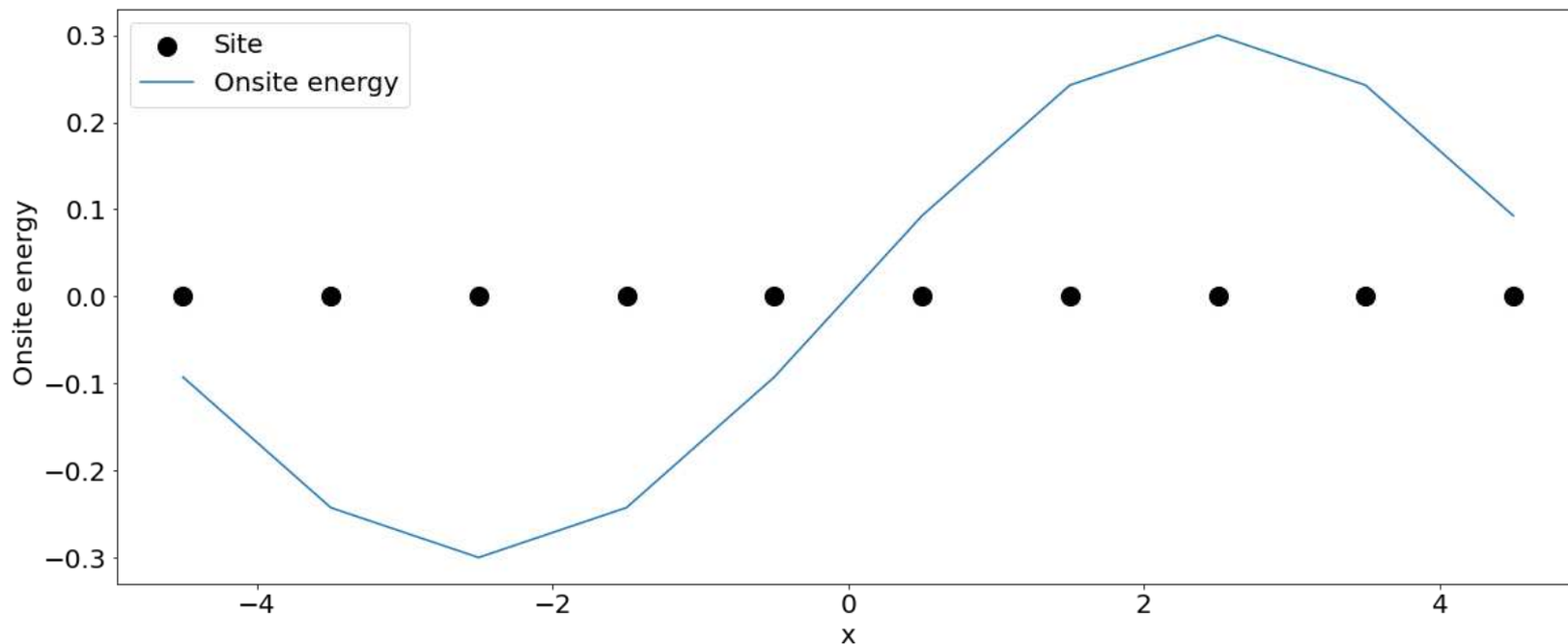
The lattice constant of the top system

The lattice constant of the bottom

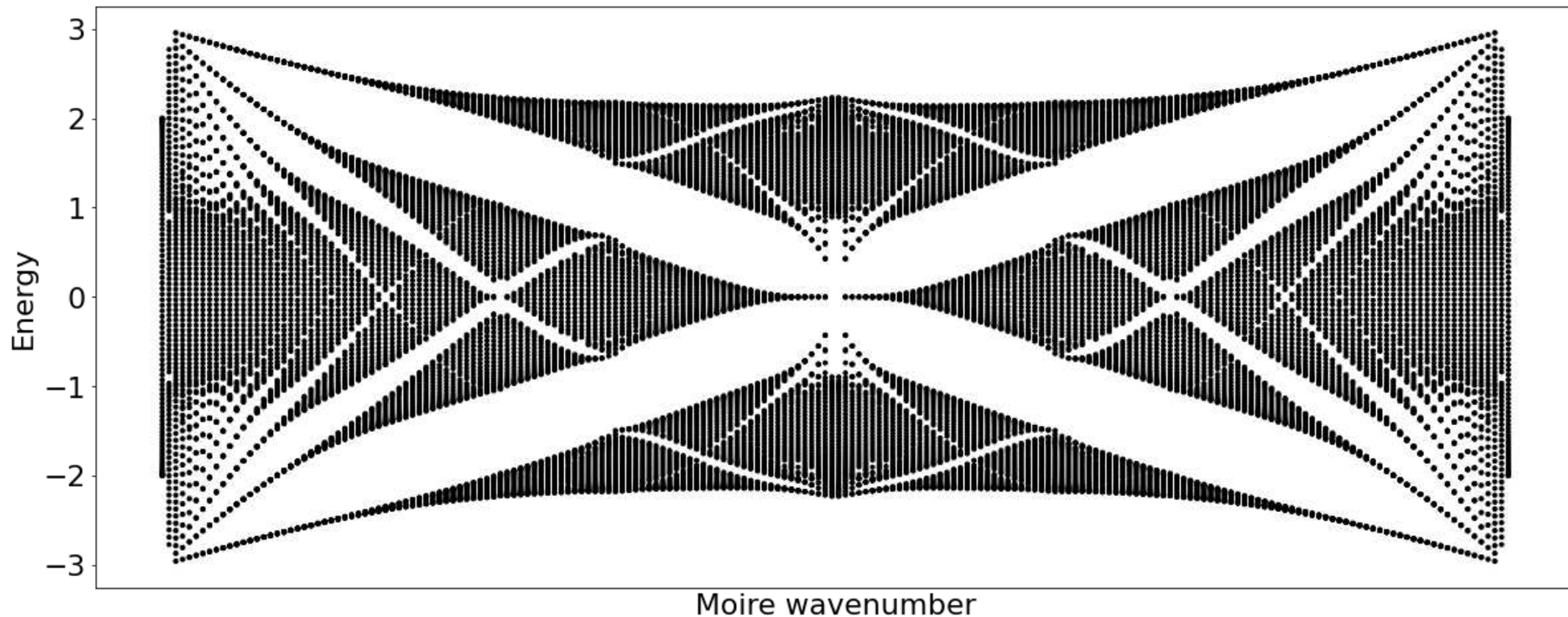
Let us see how the electronic structure gets modified by the superlattice effect

# A minimal moire potential

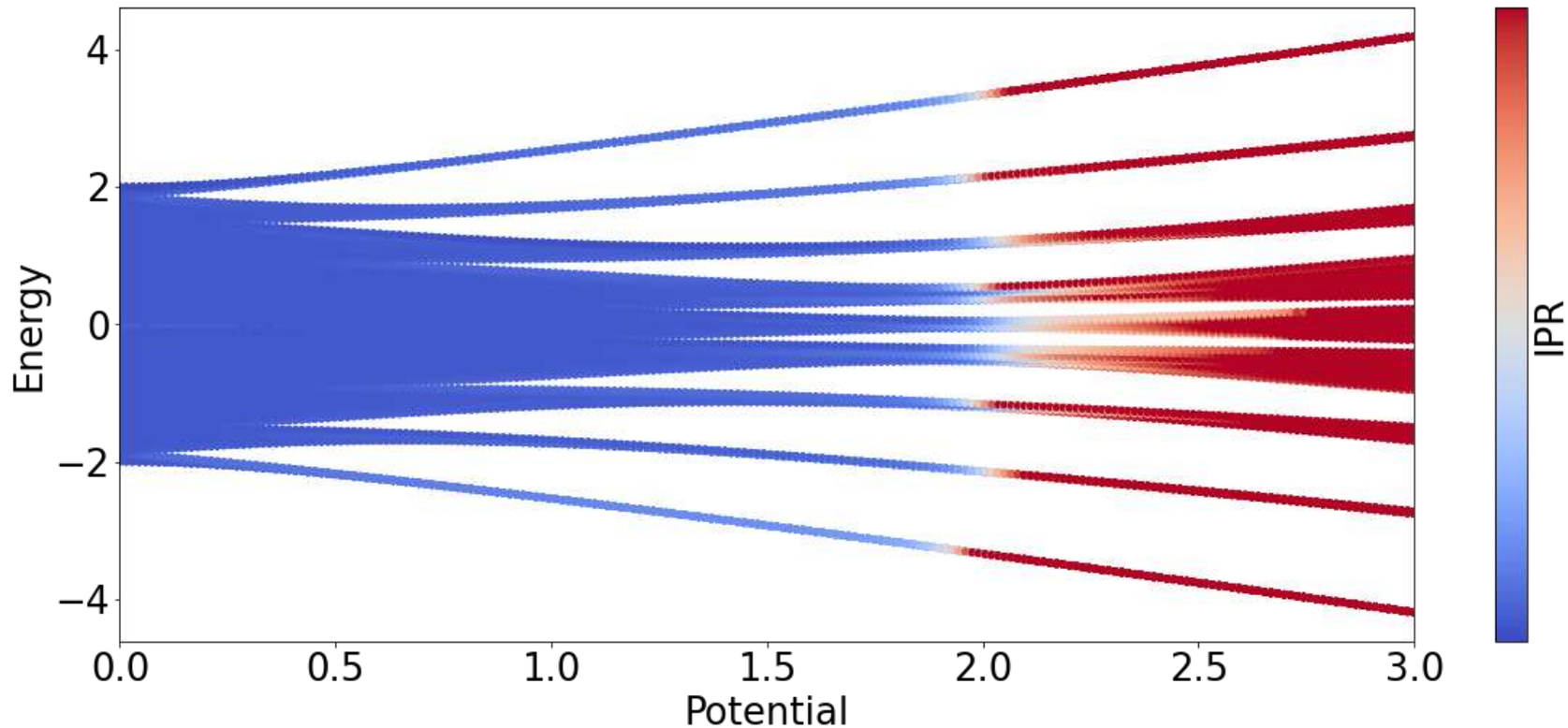
$$H = \sum_n c_n^\dagger c_{n+1} + h.c. + \lambda \sum_n \cos(qn) c_n^\dagger c_n$$



# Spectrum as a function of the moire wavevector



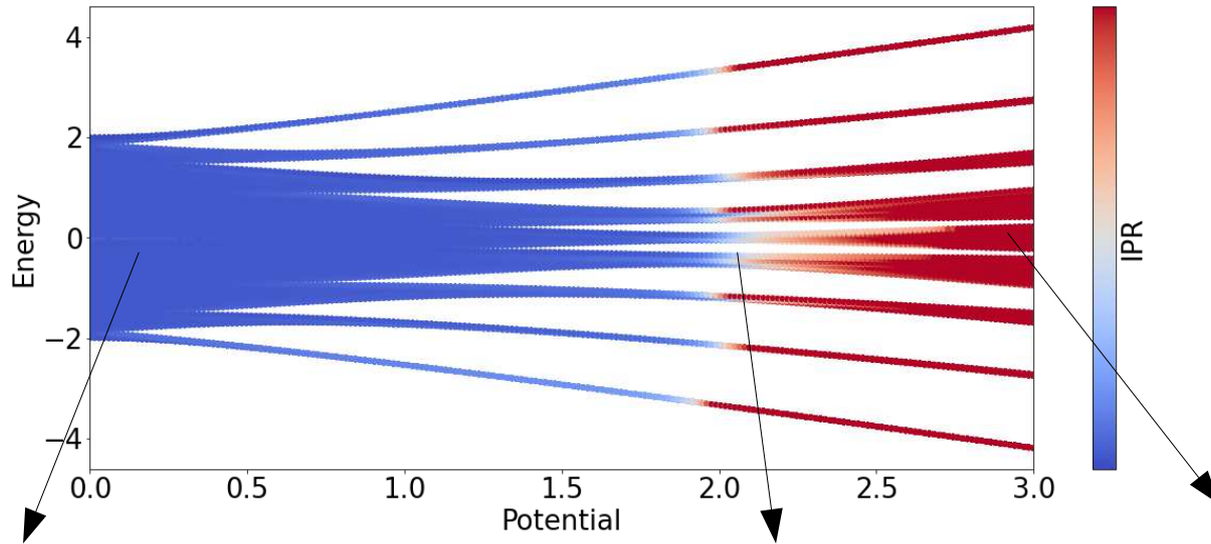
# Superpotentials and criticality



Moire potentials can give rise to critical wavefunctions

$$IPR = \sum_r |\Psi(r)|^4$$

# Superpotentials and criticality



$$IPR = \sum_r |\Psi(r)|^4$$

Extended states

$$|\Psi(\mathbf{r})| = 1/N$$

Critical states

$$|\Psi(\mathbf{r})| = |r|^{-\alpha}$$

Localized states

$$|\Psi(\mathbf{r})| = e^{-\lambda|r|}$$

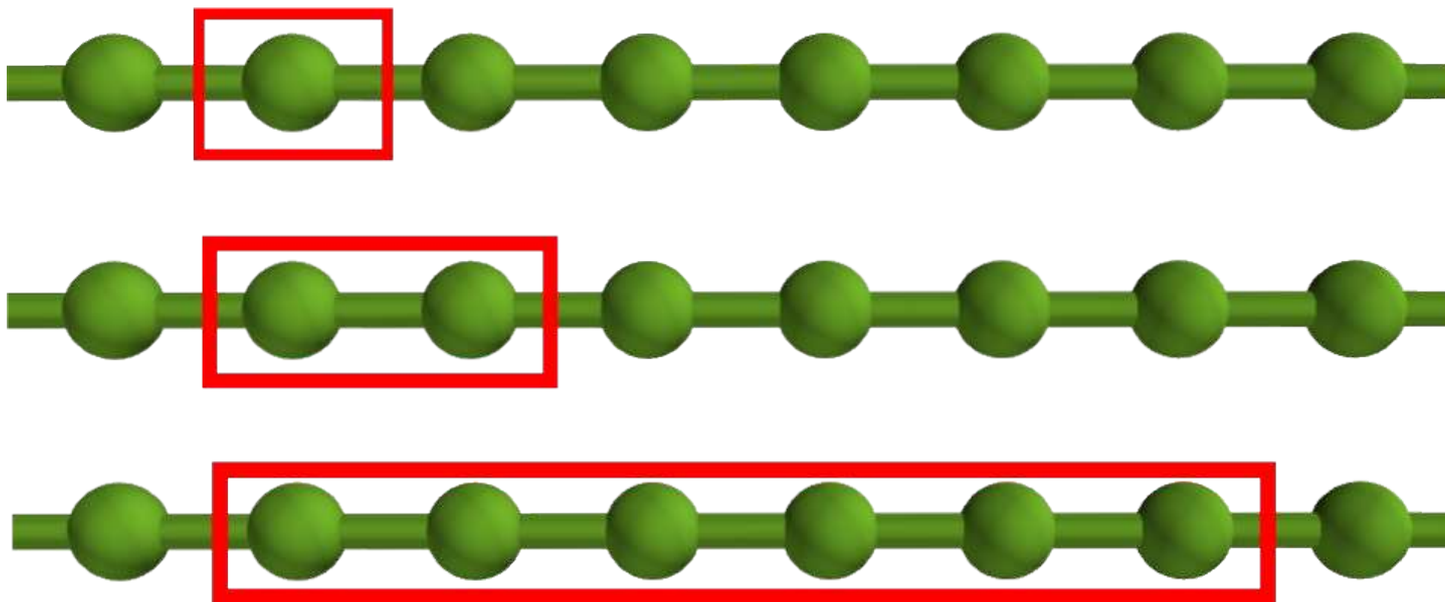
# Minibands and band structure unfolding

# Supercells and band folding

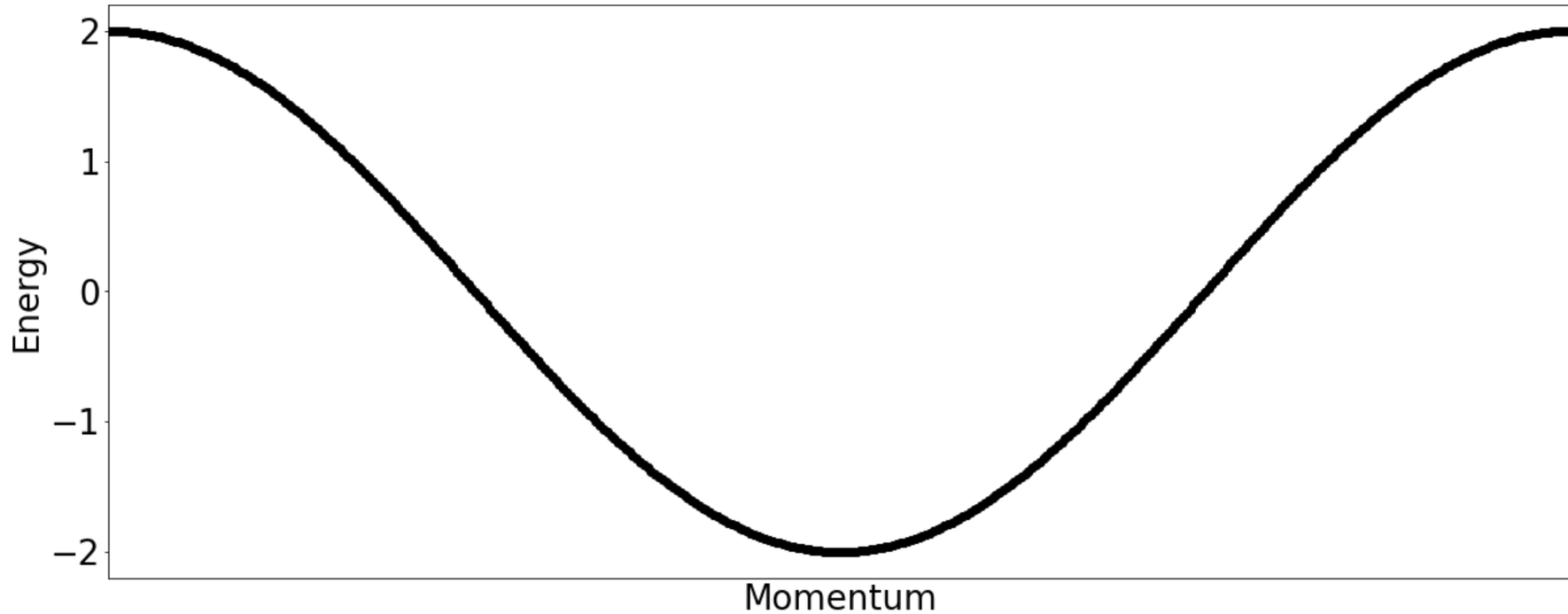
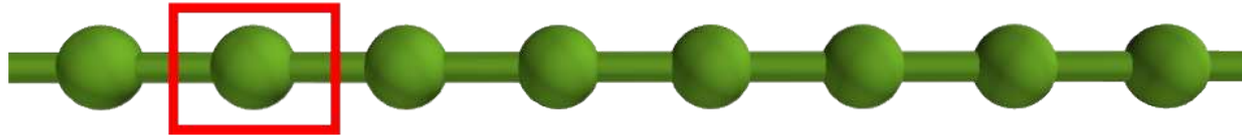
Let us take a 1D chain

$$H = \sum_n c_n^\dagger c_{n+1} + h.c.$$

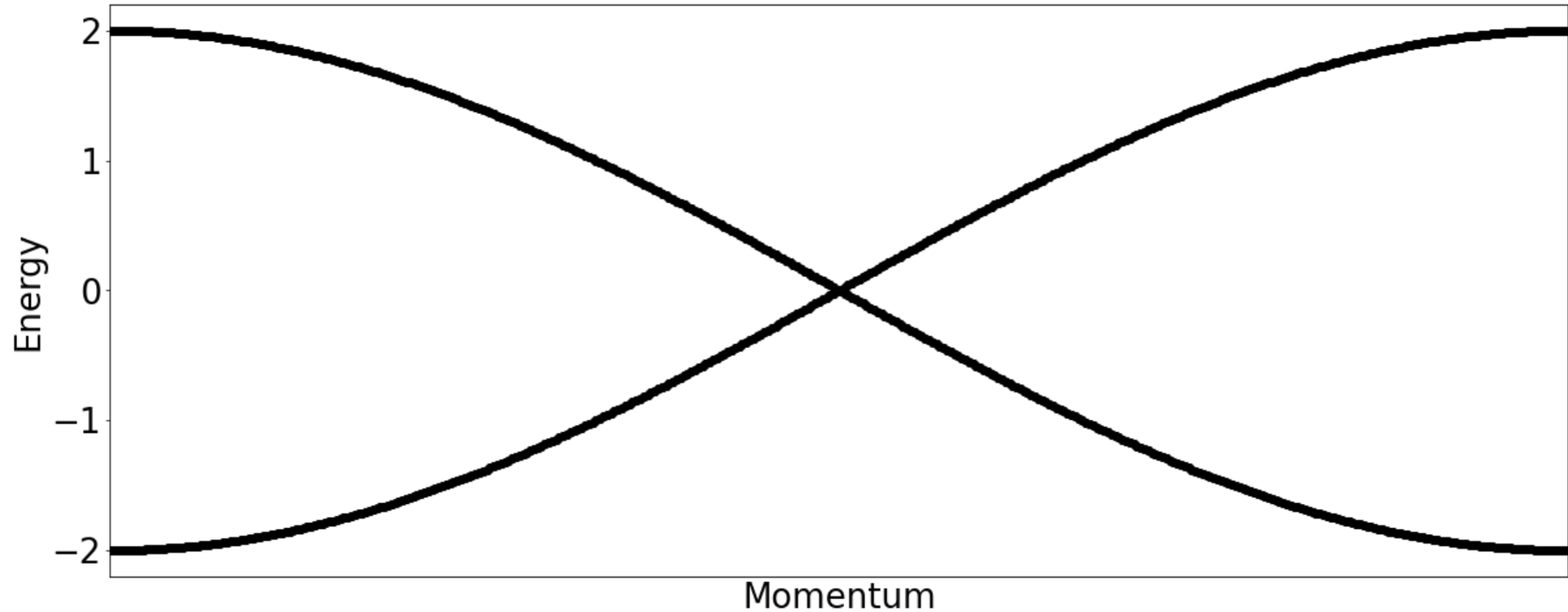
Let us see how the electronic structure changes with the unit cell



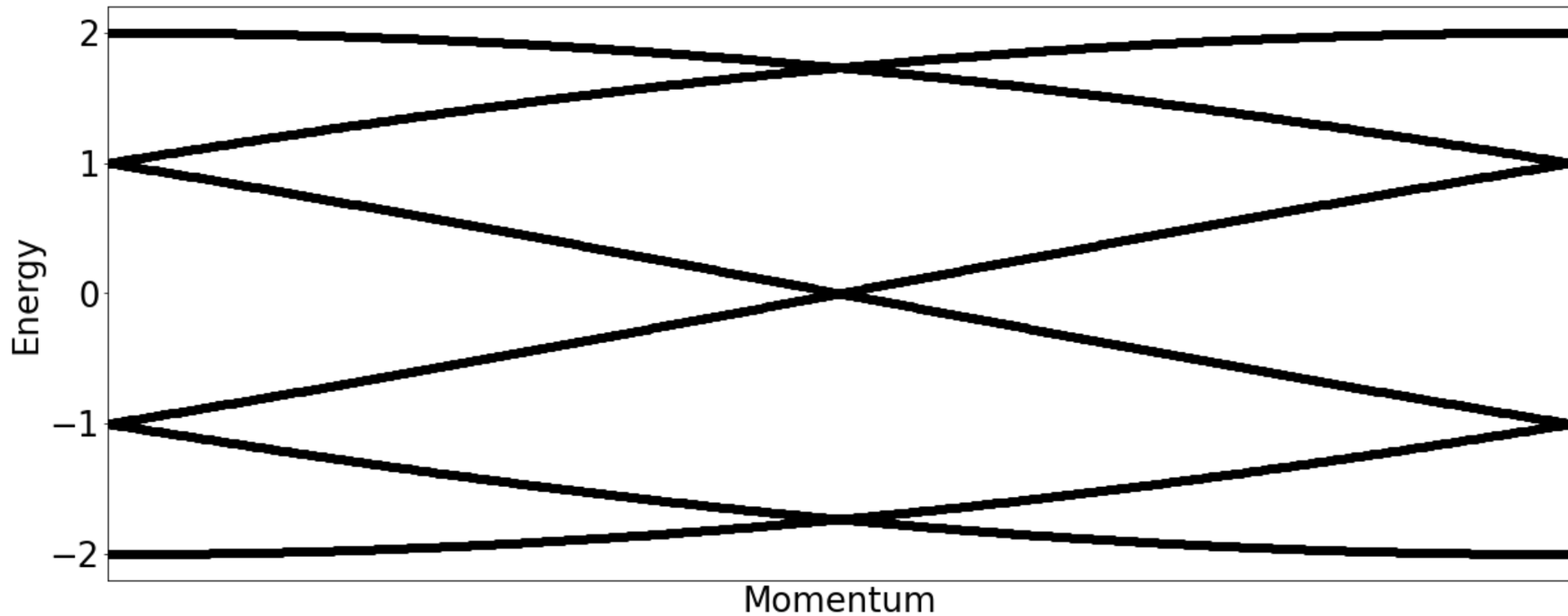
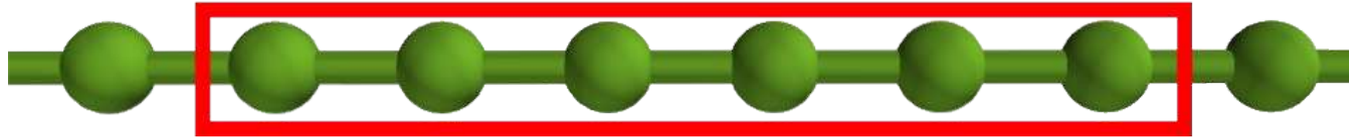
# Supercells and band folding



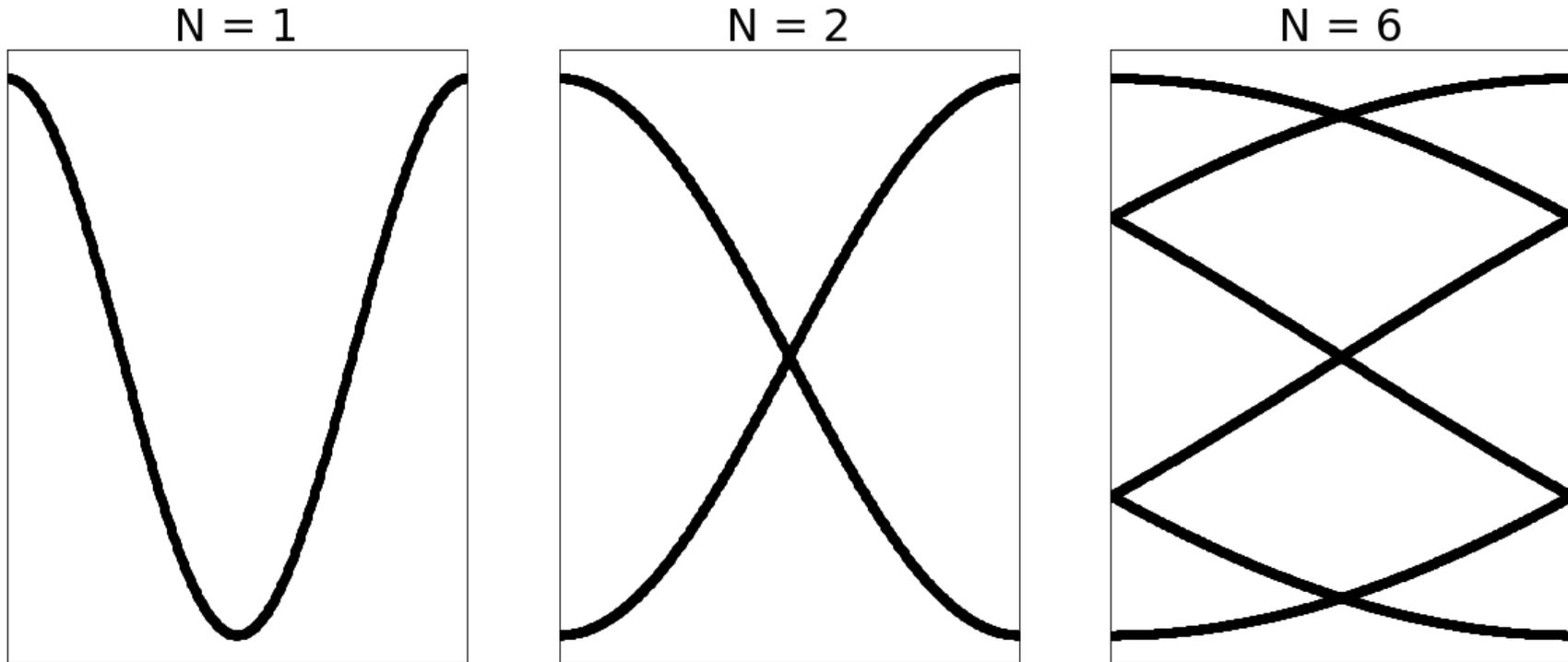
# Supercells and band folding



# Supercells and band folding

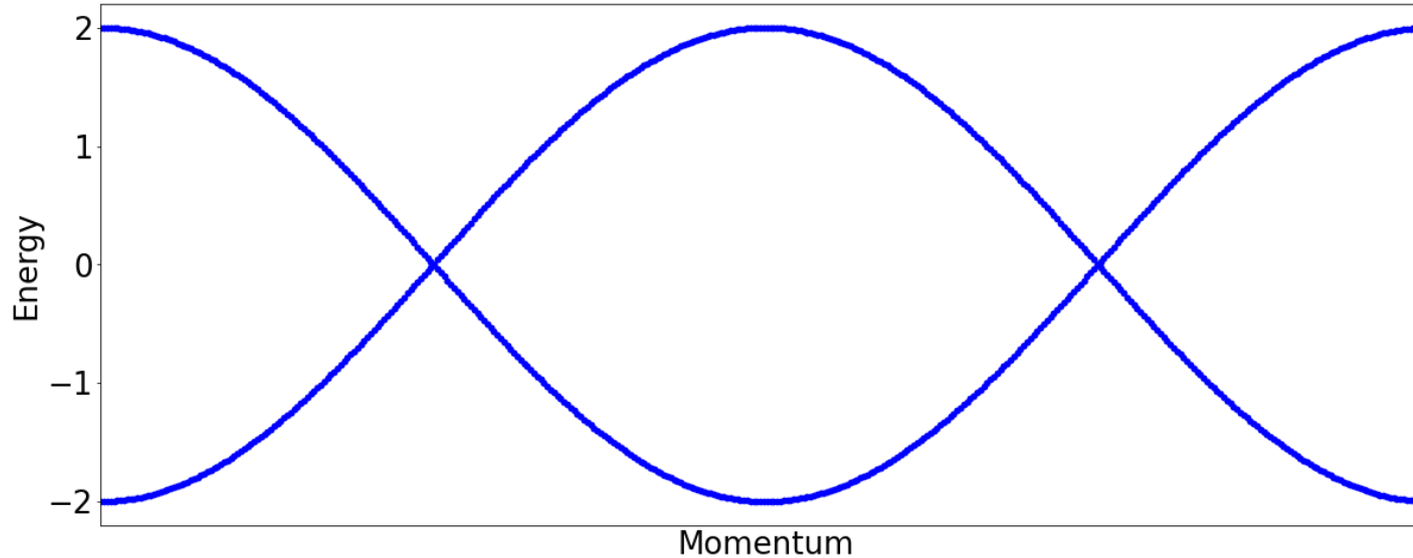
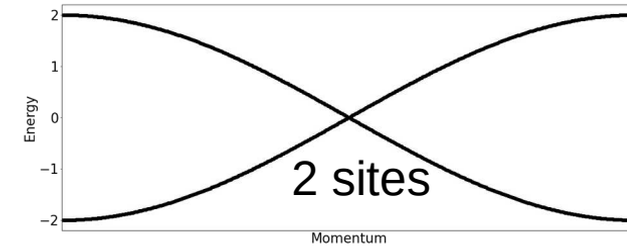
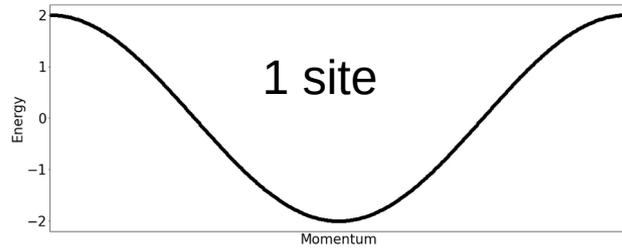


# Supercells and band folding



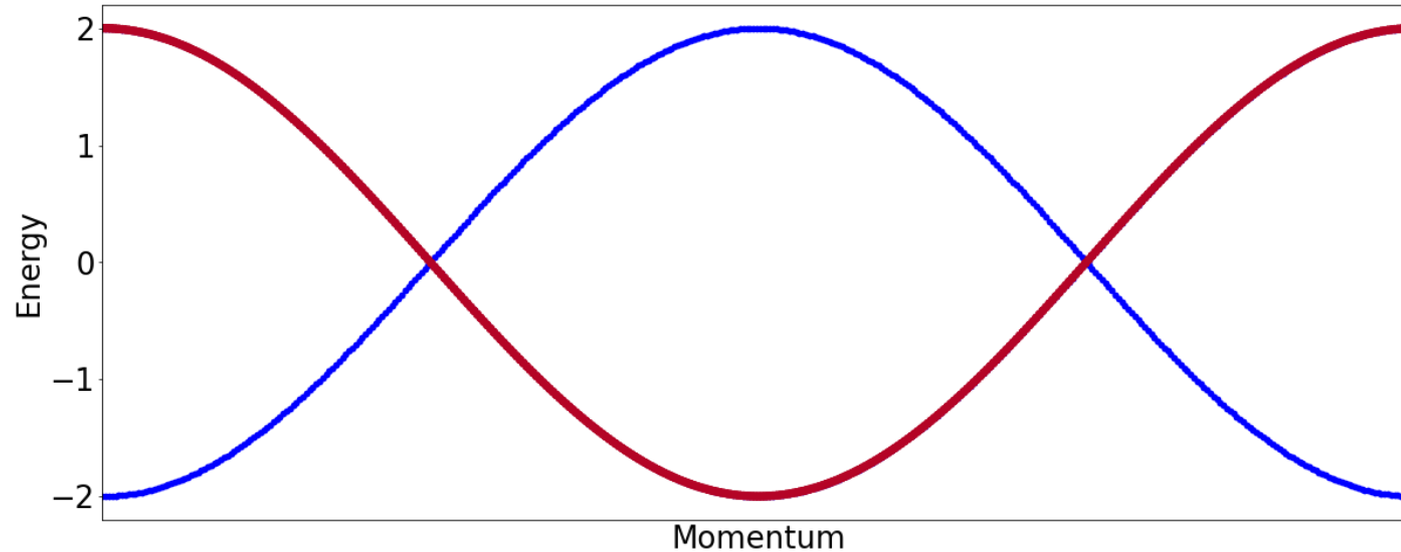
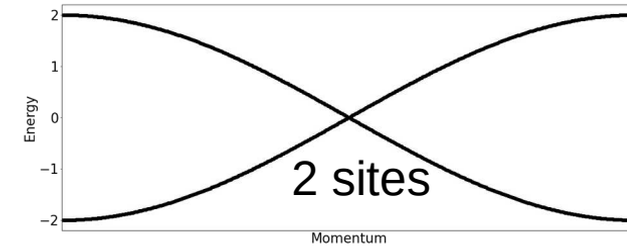
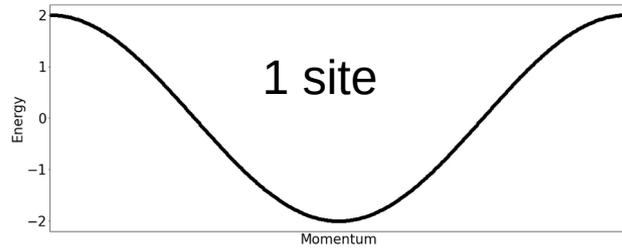
All these electronic structures represent the same physical system, but how do we see that?

# Supercells and band folding



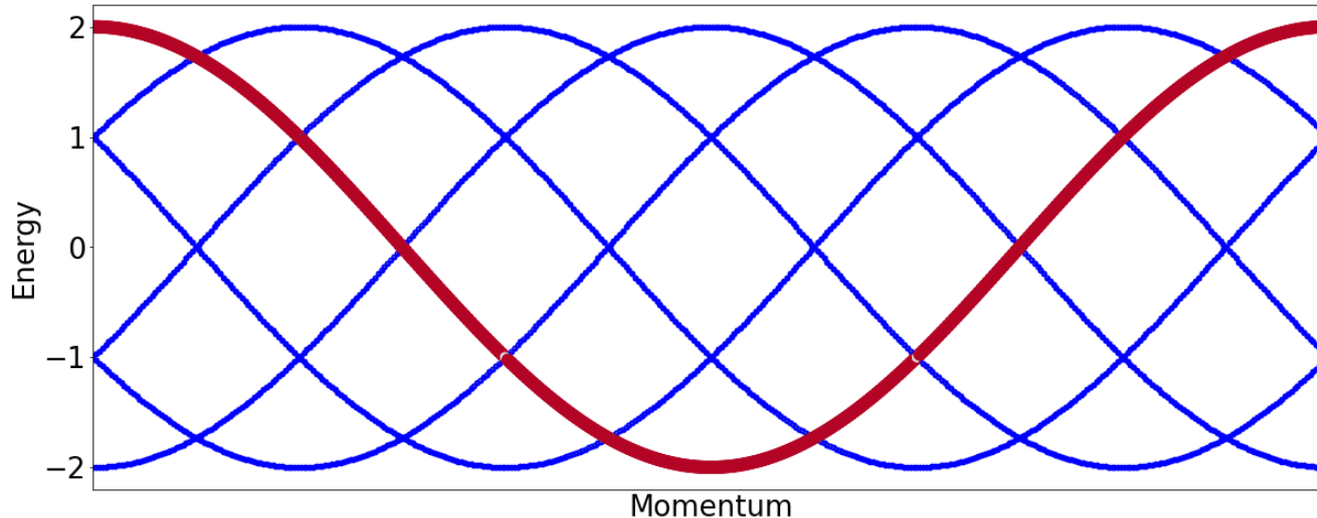
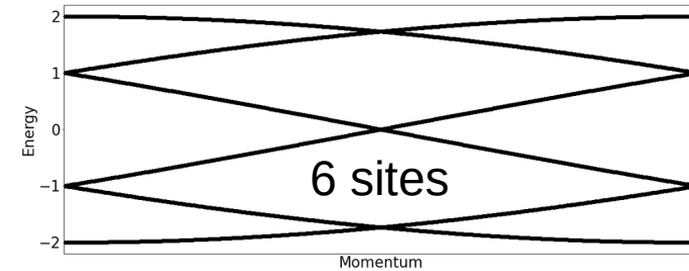
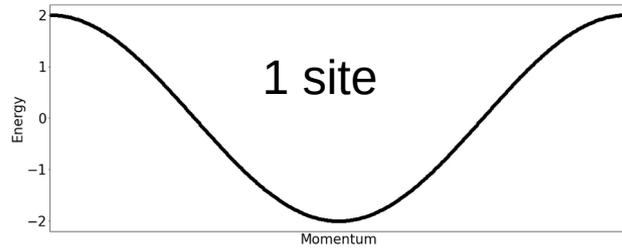
Repeating the electronic structure recovers the original electronic dispersion

# Supercells and band folding



Repeating the electronic structure recovers the original electronic dispersion

# Supercells and band folding



Repeating the electronic structure recovers the original electronic dispersion

# Unfolding and anticrossings in superlattices

Let us now put an impurity every 6 sites (once in a supercell 6)

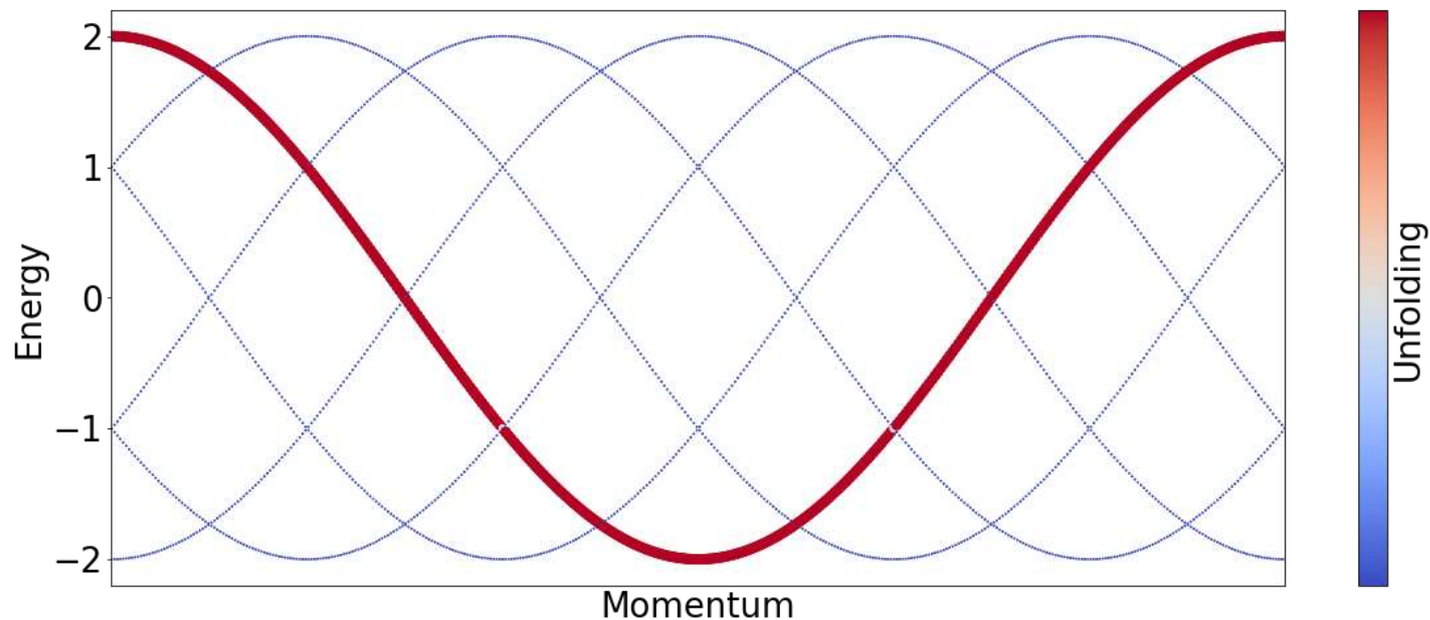


$$H = \sum_n c_n^\dagger c_{n+1} + h.c. + V_0 \sum_{\alpha} c_n^\dagger c_n \quad \alpha \equiv 0 \pmod{6}$$

# Unfolding and anticrossings in superlattices

$$H = \sum_n c_n^\dagger c_{n+1} + h.c. + V_0 \sum_\alpha c_n^\dagger c_n$$

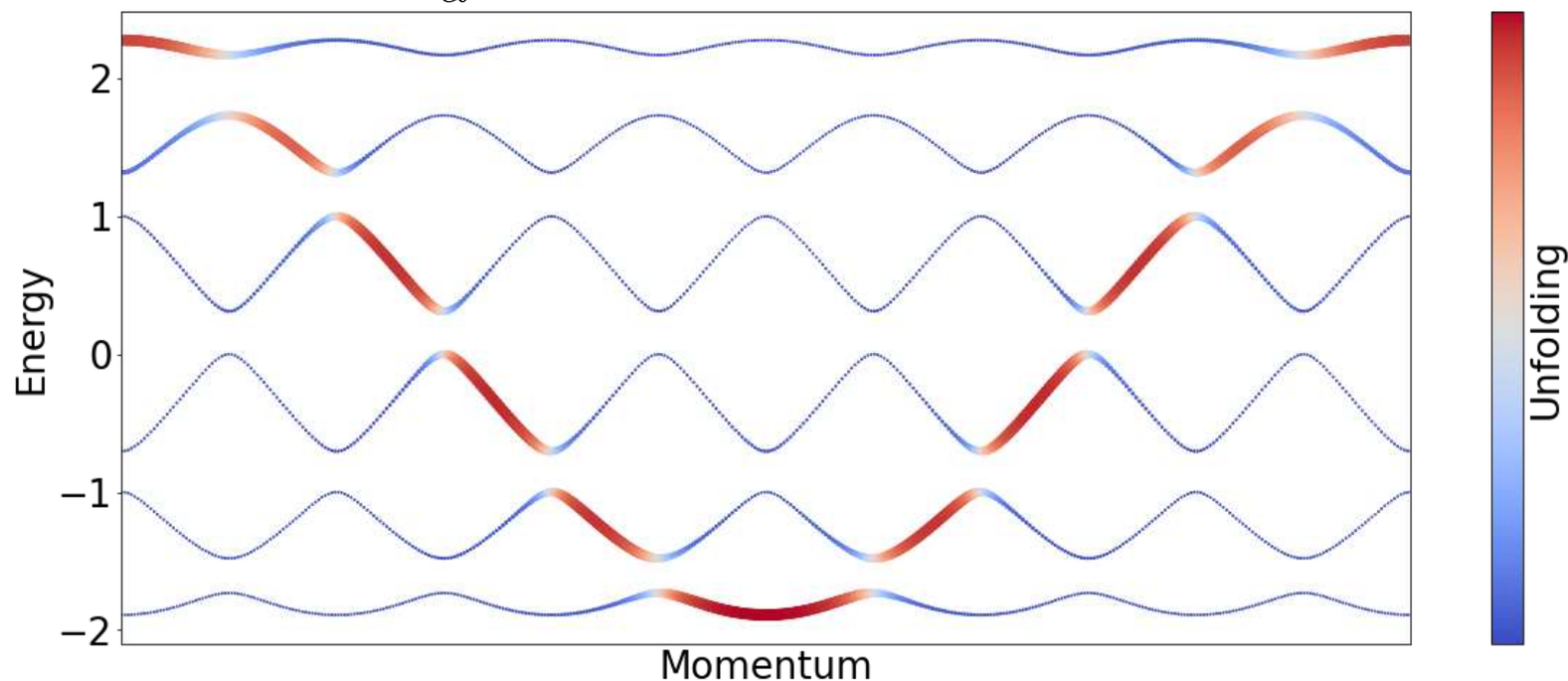
$$V_0 = 0$$



# Electronic structure unfolding

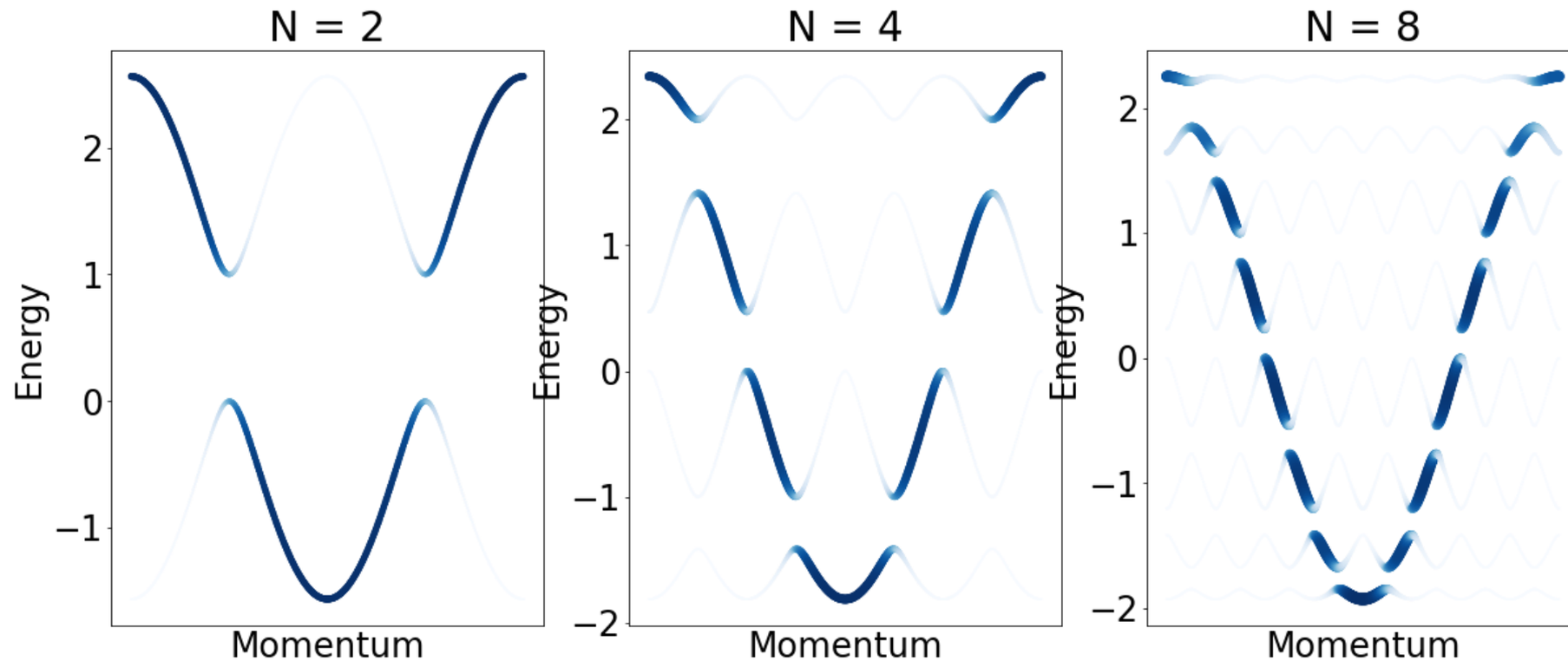
$$H = \sum_n c_n^\dagger c_{n+1} + h.c. + V_0 \sum_\alpha c_n^\dagger c_n$$

$$V_0 \neq 0$$



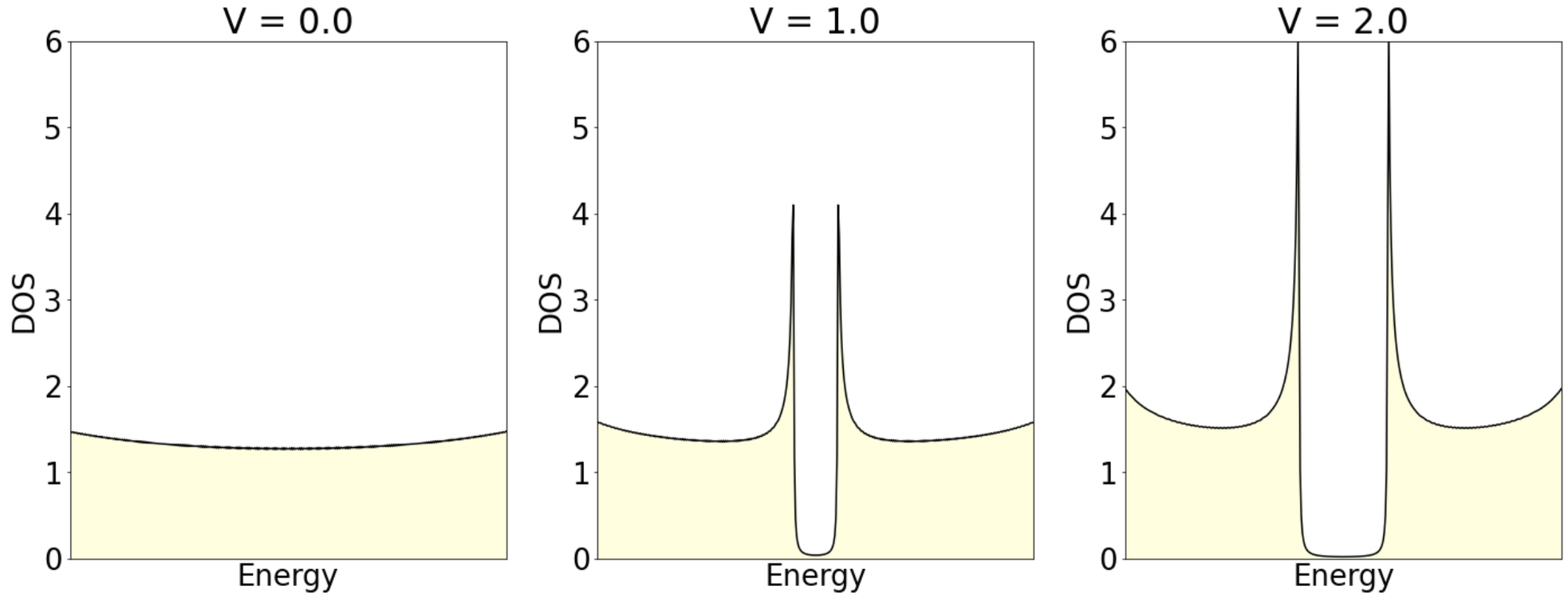
Anticrossing between the bands appear due to the superlattice potential

# Electronic structure unfolding



As the periodicity of the superlattice is increased, more minibands appear

# Moire electronic structure



As the strength of the moire potential increases, the density of states gets enhanced

# Break

10-15 min break

*(optional) to discuss during the break*

$$H = \lambda \sum_n \cos(qn) c_n^\dagger c_n$$

For a lattice with  $L$  sites, what is the value of  $q$  that makes the potential commensurate?

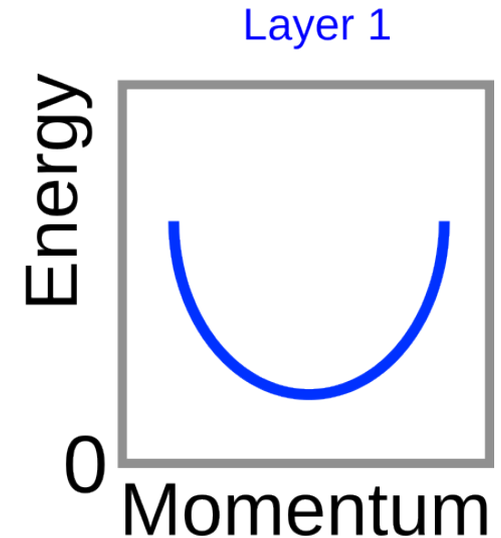
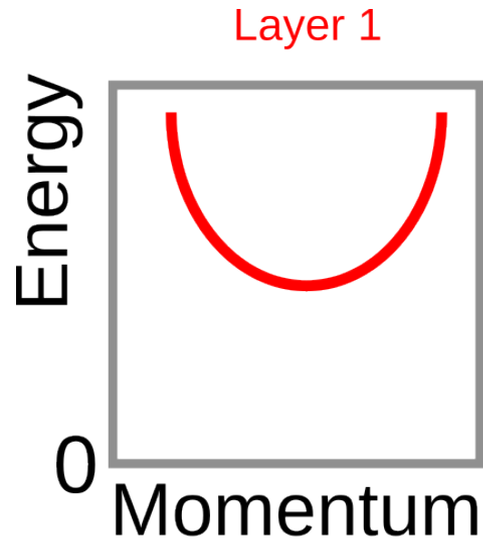
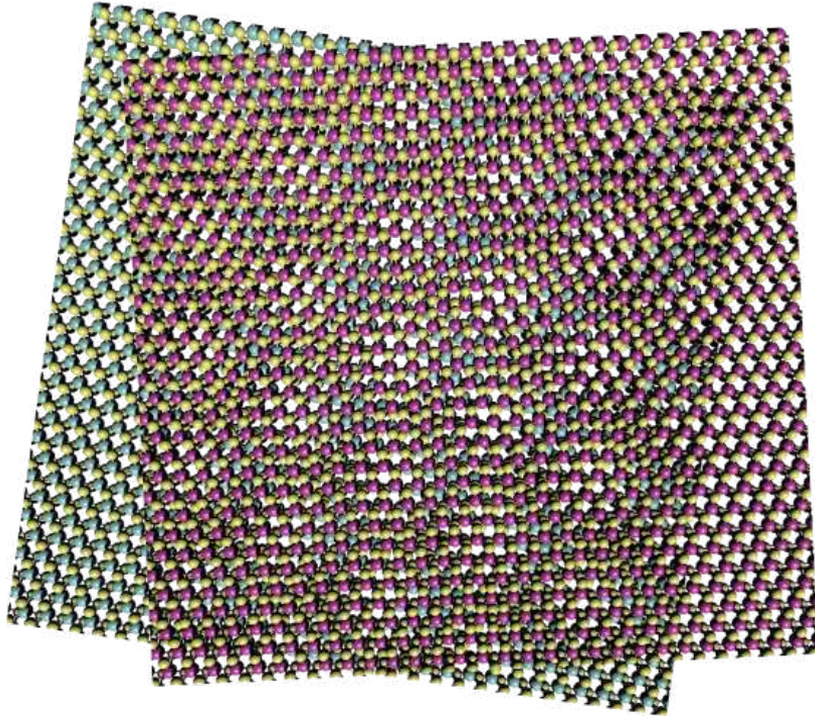
$$q = \frac{\alpha 2\pi}{L}$$

$$q = \frac{\alpha\pi}{L}$$

$$\alpha = 1, 2, 3, 4, \dots$$

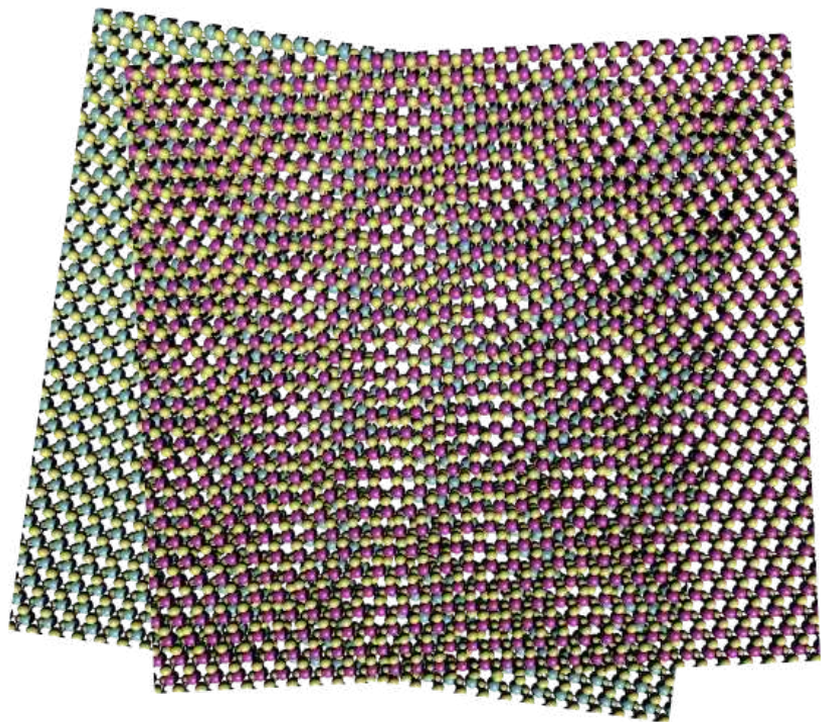
# Moire in twisted TMDC

In twisted TMDC heterobilayers, the moire modulates the band off set of one layer

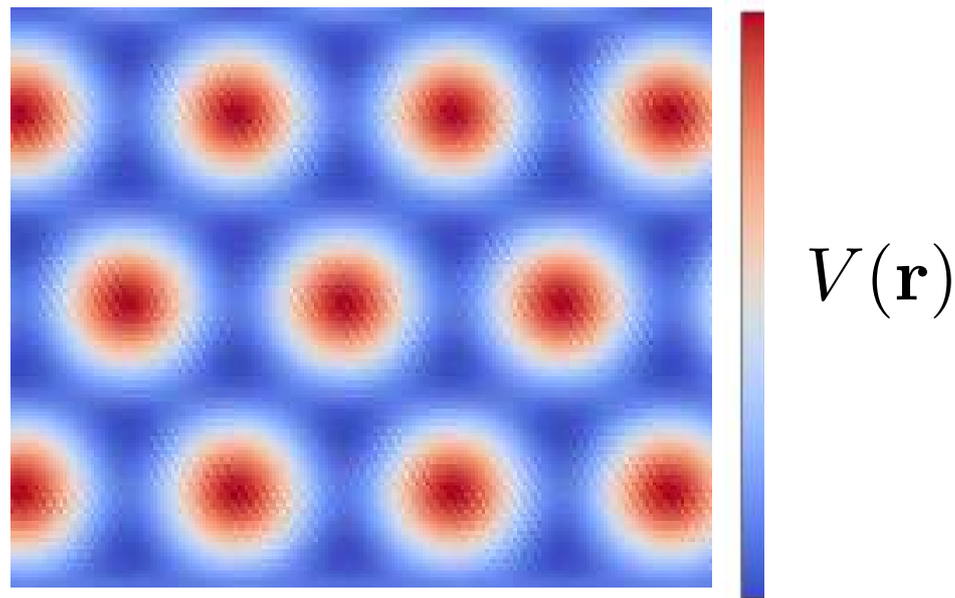


# Moire in twisted TMDC

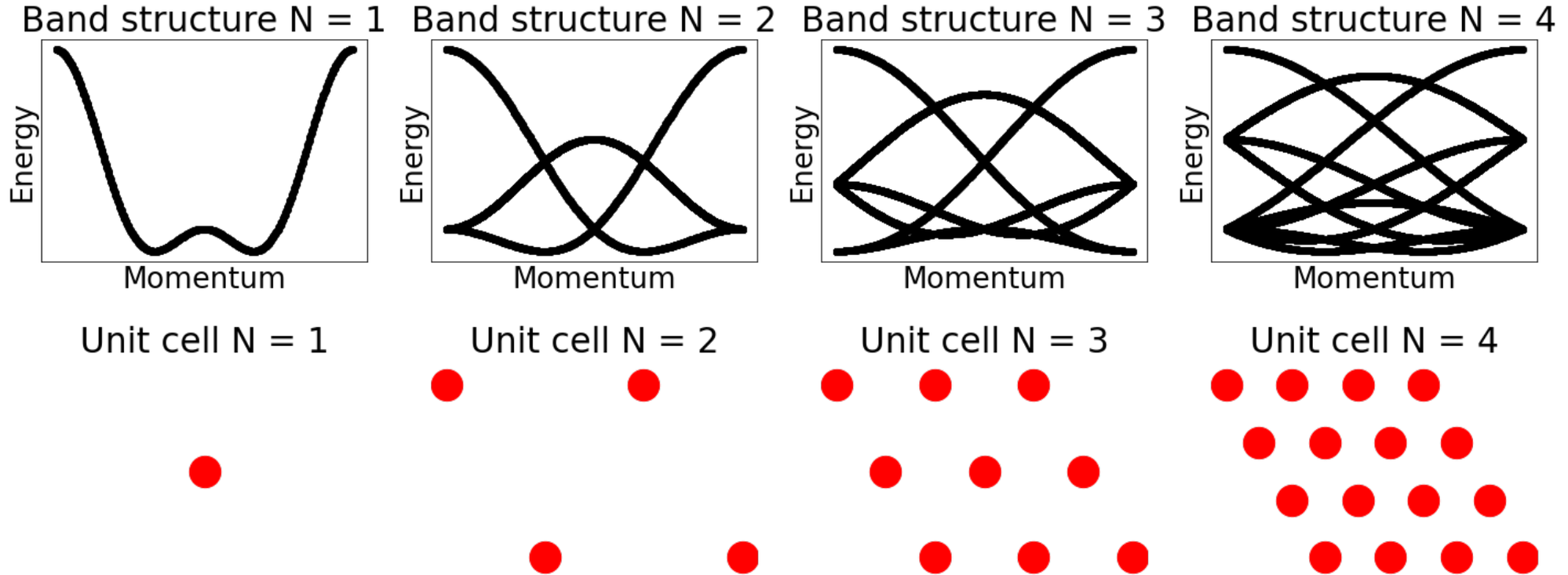
In twisted TMDC heterobilayers, the moire modulates the band offset of one layer



$$H = \sum_{ij} c_i^\dagger c_j + h.c. + \sum_n V(\mathbf{r}_n) c_n^\dagger c_n$$



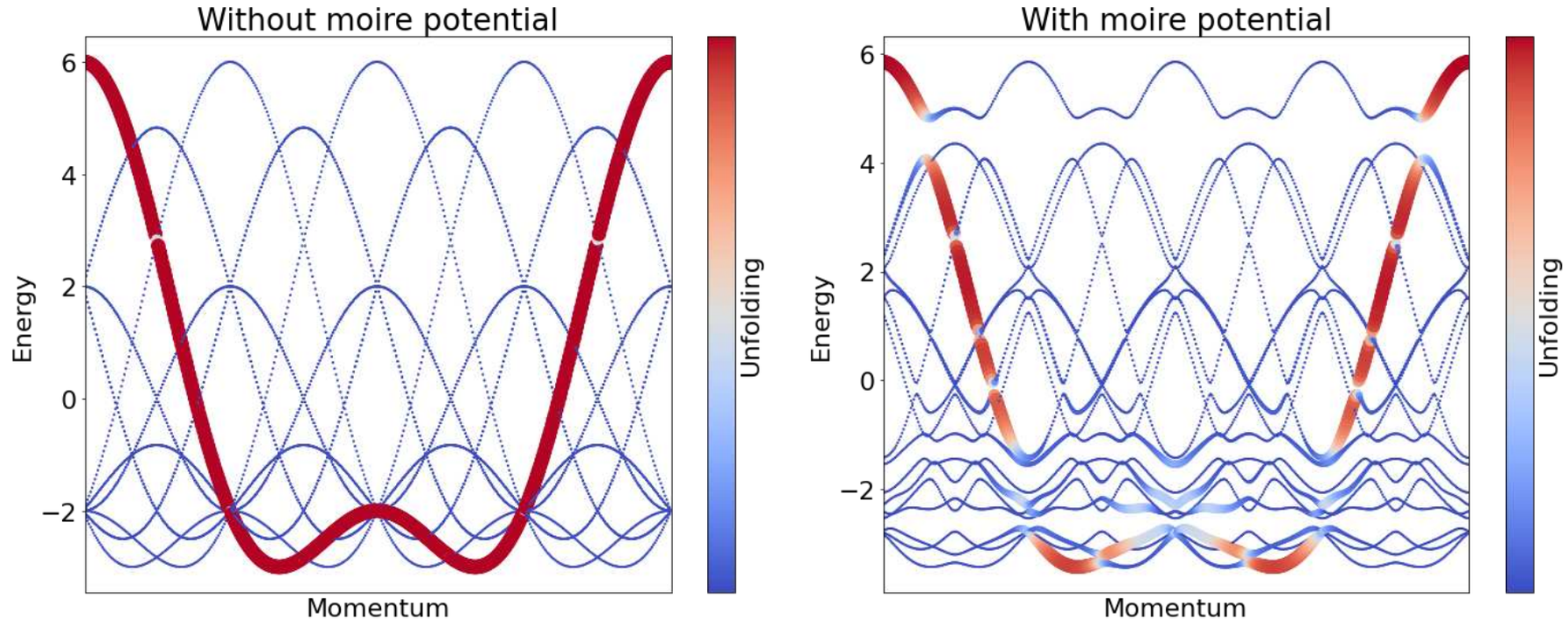
# Band structure folding in 2D superlattices



A 2D superlattice gives rise to a complex folding of the electronic structure

# Band structure folding in 2D superlattices

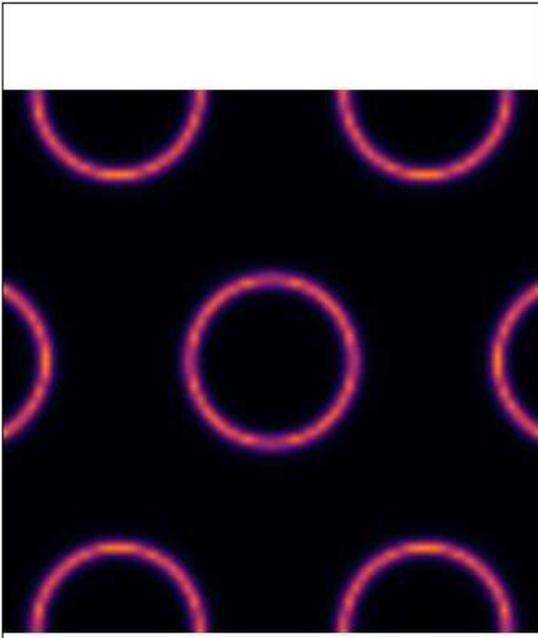
Mini-bands appear due to the moire in one of the layers



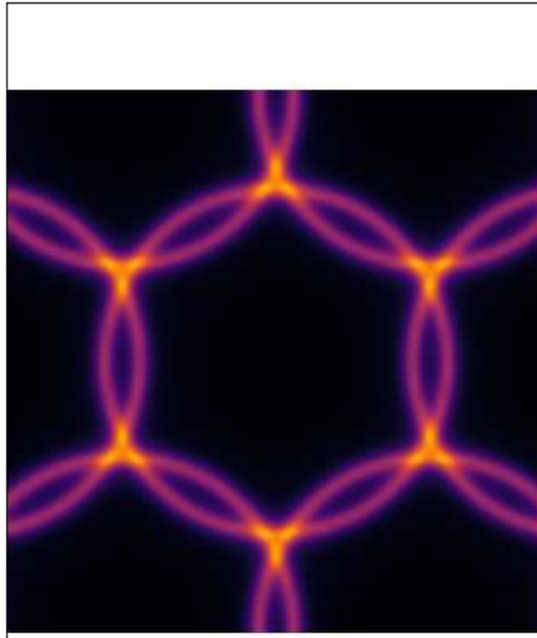
# Fermi surface folding in superlattices

The Fermi surface in 2d supercell gets folded, turning one Fermi surface into many

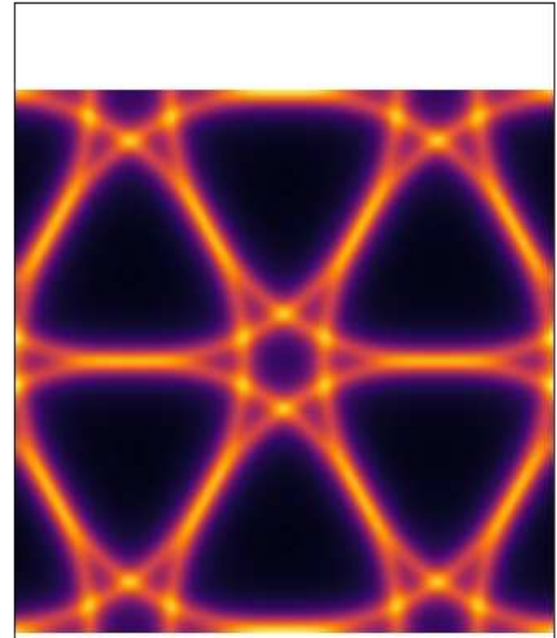
$N = 1$



$N = 2$

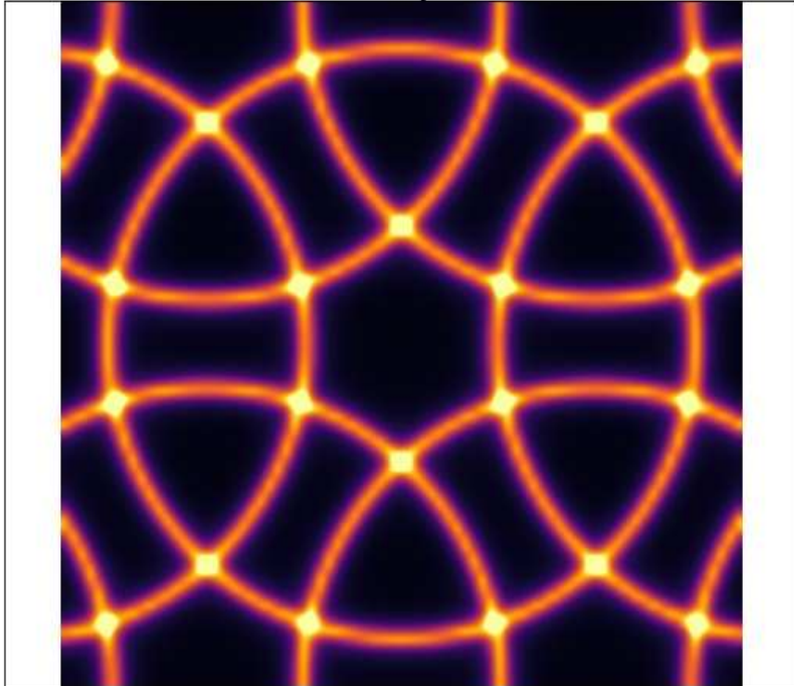


$N = 3$

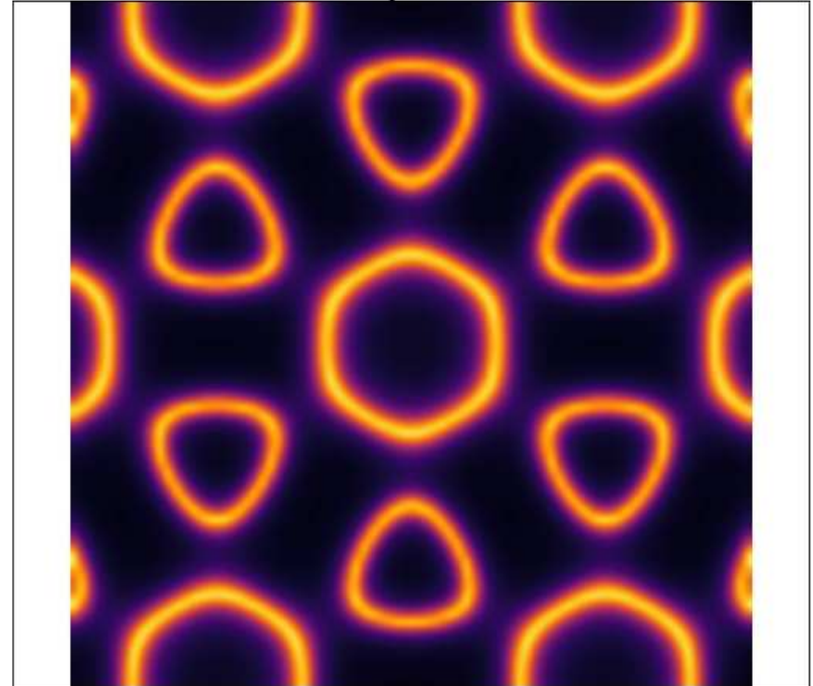


# Fermi surface folding in superlattices

Without superlattice



With superlattice



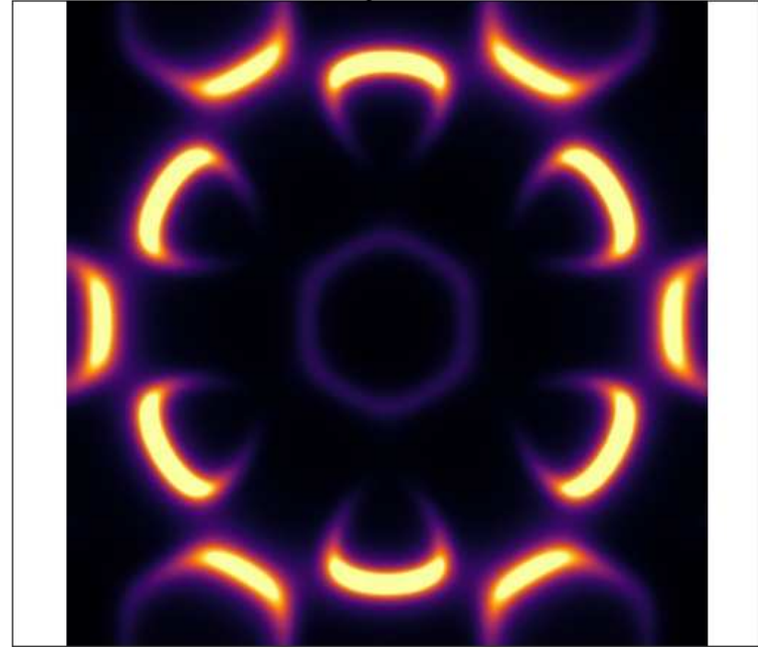
The superlattice changes the Fermi surface topology on the underlying 2D material

# Fermi surface unfolding in superlattices

Without superlattice



With superlattice

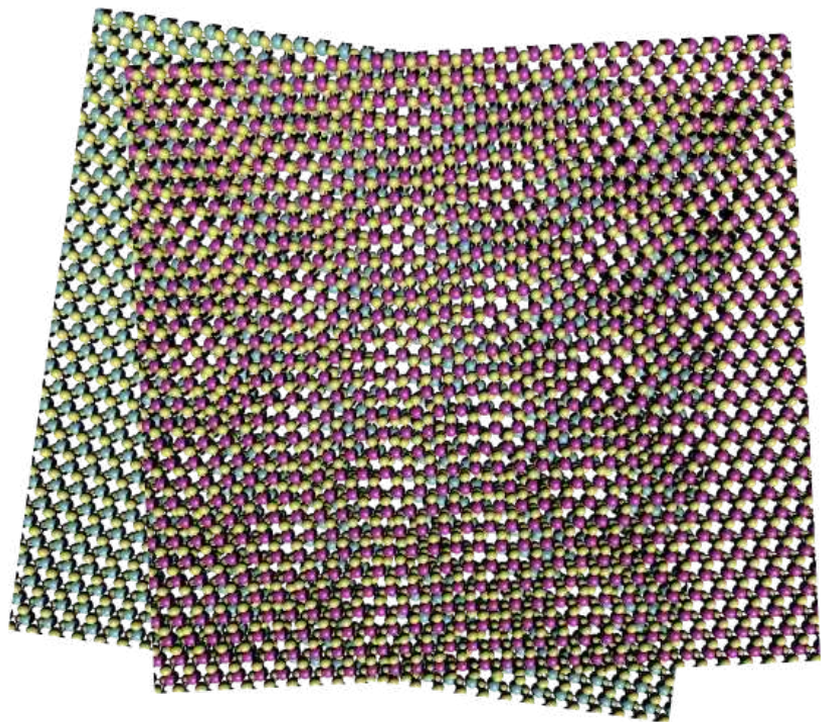


Superlattices fragment the original Fermi surface when unfolded to the original unit cell

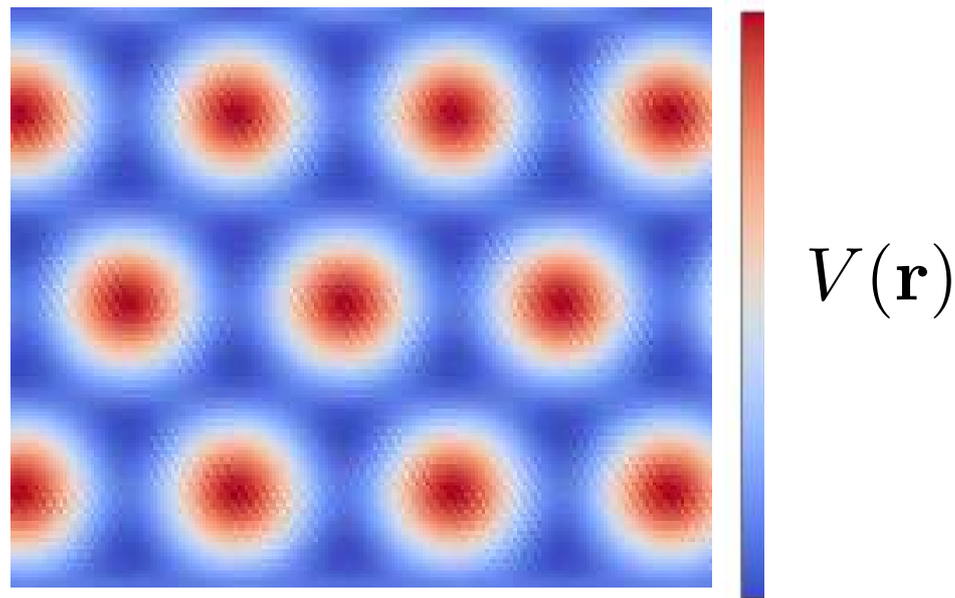
# Moire-driven correlated states

# Moire in twisted TMDC

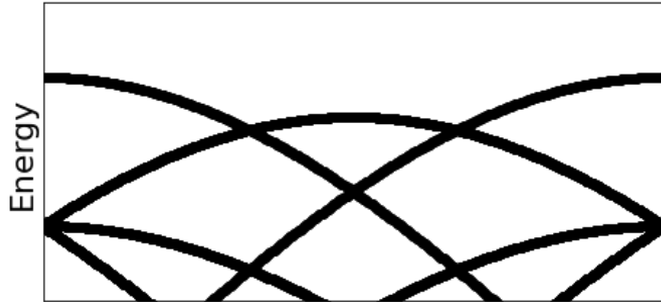
In twisted TMDC heterobilayers, the moire modulates the band off set of one layer



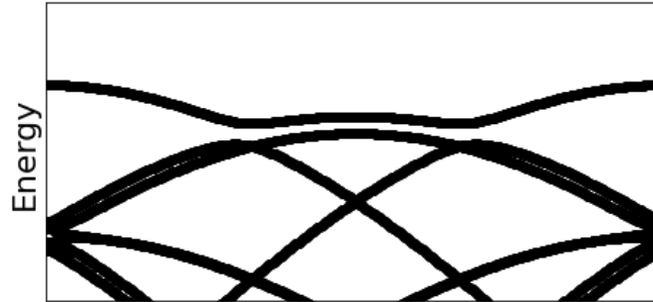
$$H = \sum_{ij} c_i^\dagger c_j + h.c. + \sum_n V(\mathbf{r}_n) c_n^\dagger c_n$$



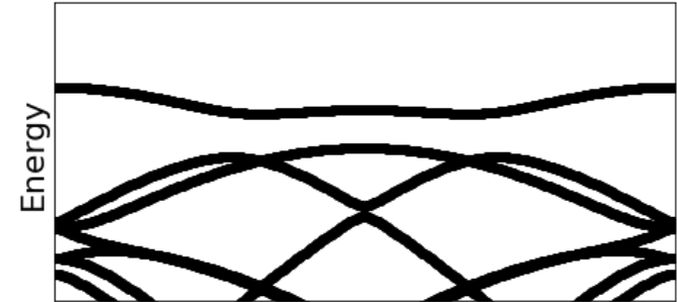
# Band flattening by a moire

 $V = 0.0$ 

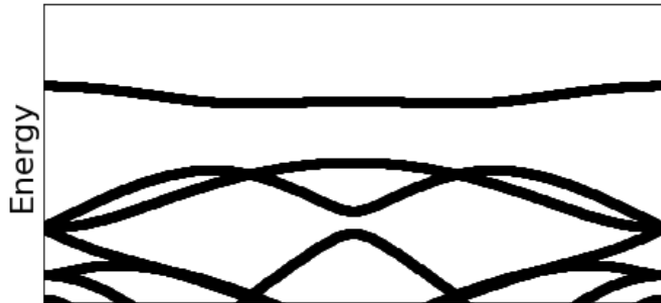
Momentum

 $V = 0.4$ 

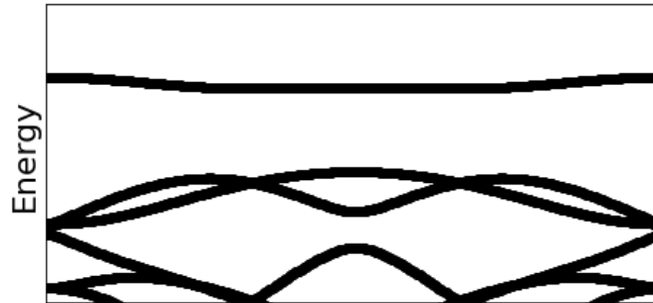
Momentum

 $V = 0.8$ 

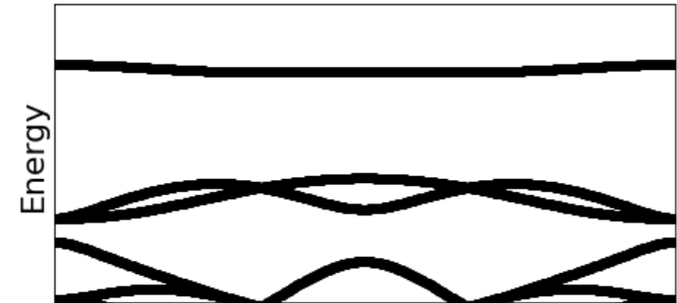
Momentum

 $V = 1.2$ 

Momentum

 $V = 1.6$ 

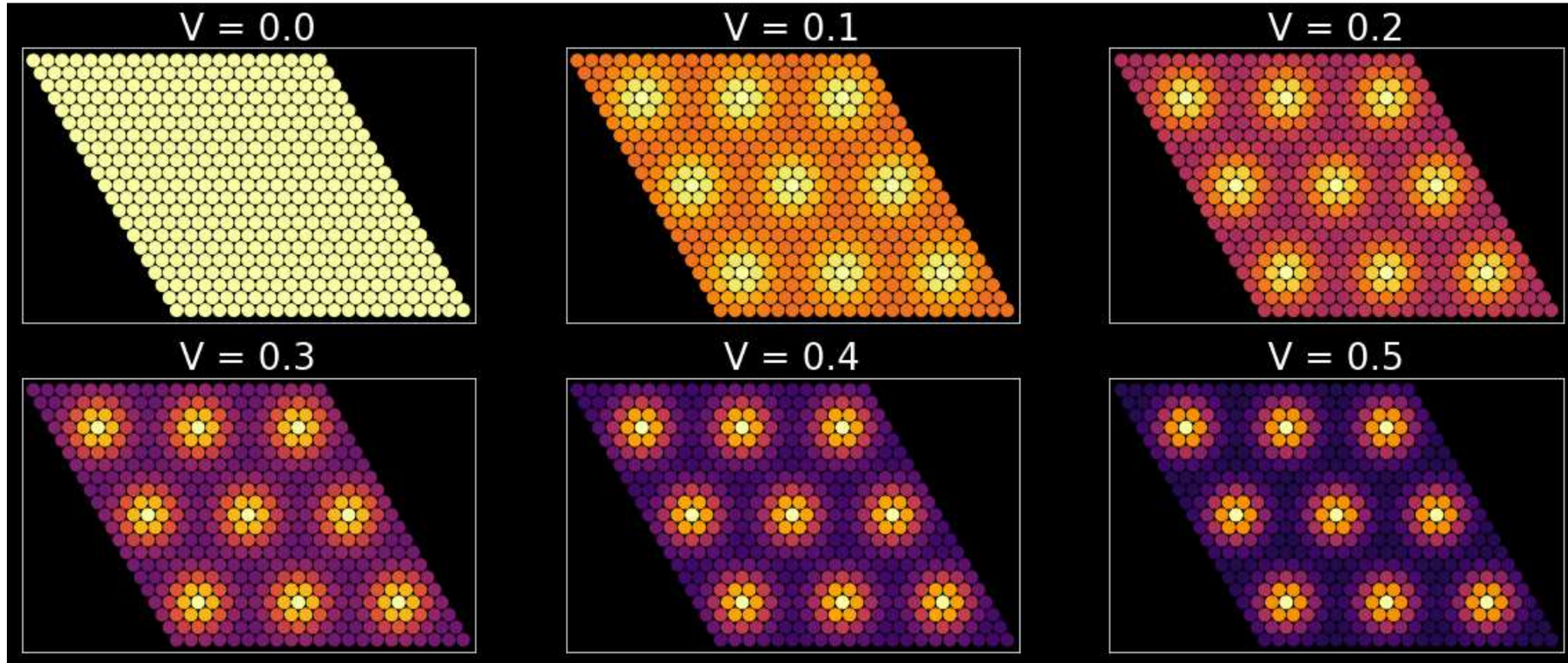
Momentum

 $V = 2.0$ 

Momentum

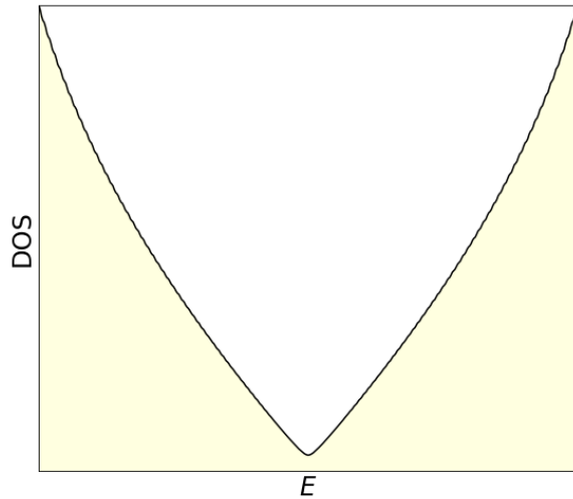
As the strength of a modulation is increased, flat bands appear

# Emergence of moire states

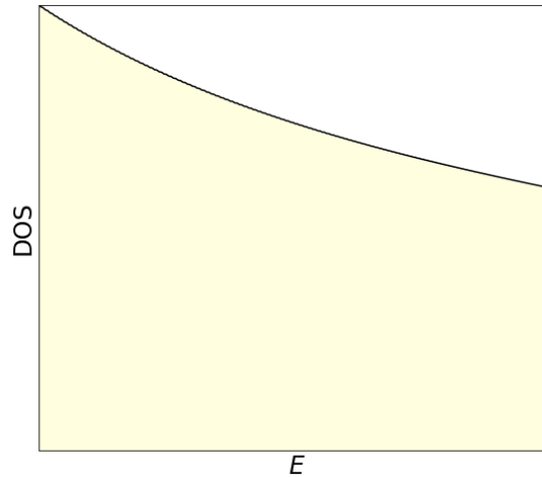


# Controlling the electronic spectra with the moire angle

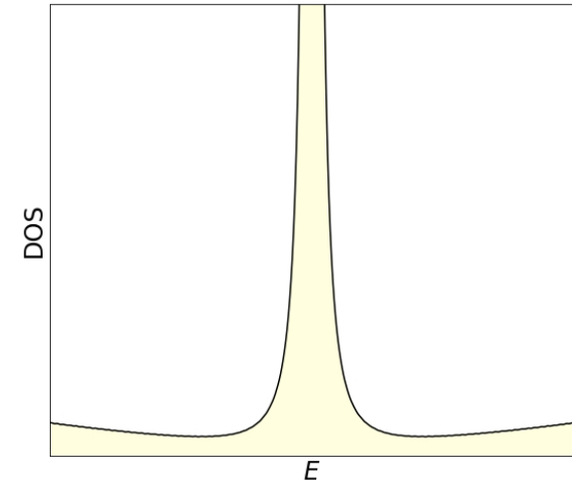
## Semimetals



## Metals



## Flat bands

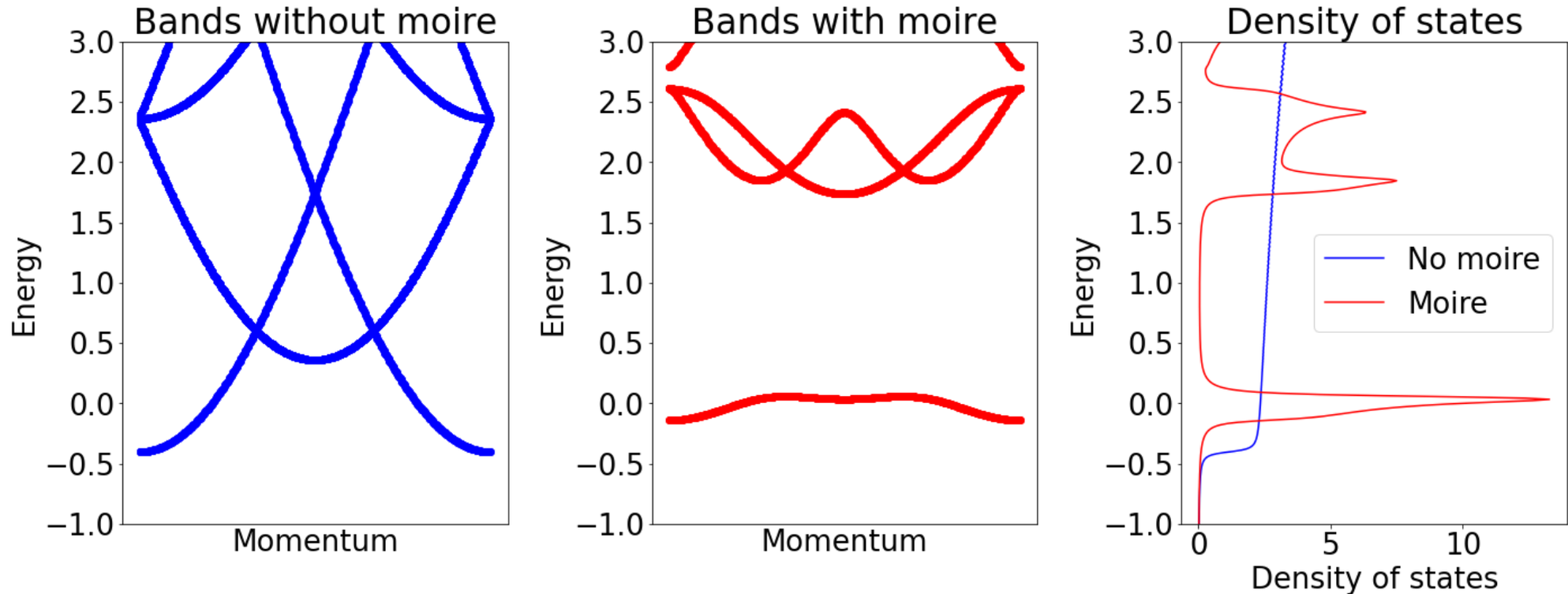


Moire/twist angle

Moire potentials allow driving systems to the correlated limit

$$U_C \sim \frac{1}{D(E_F)}$$

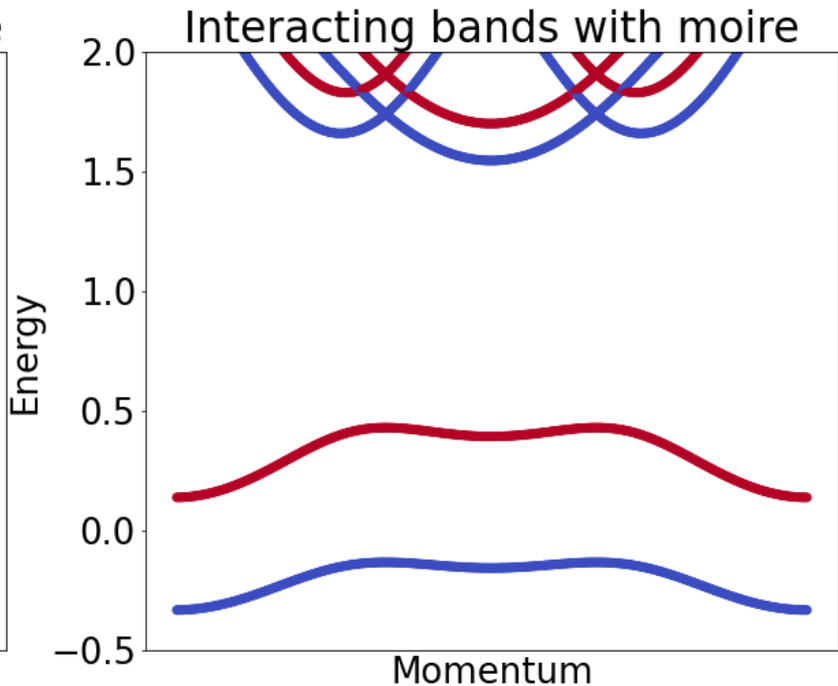
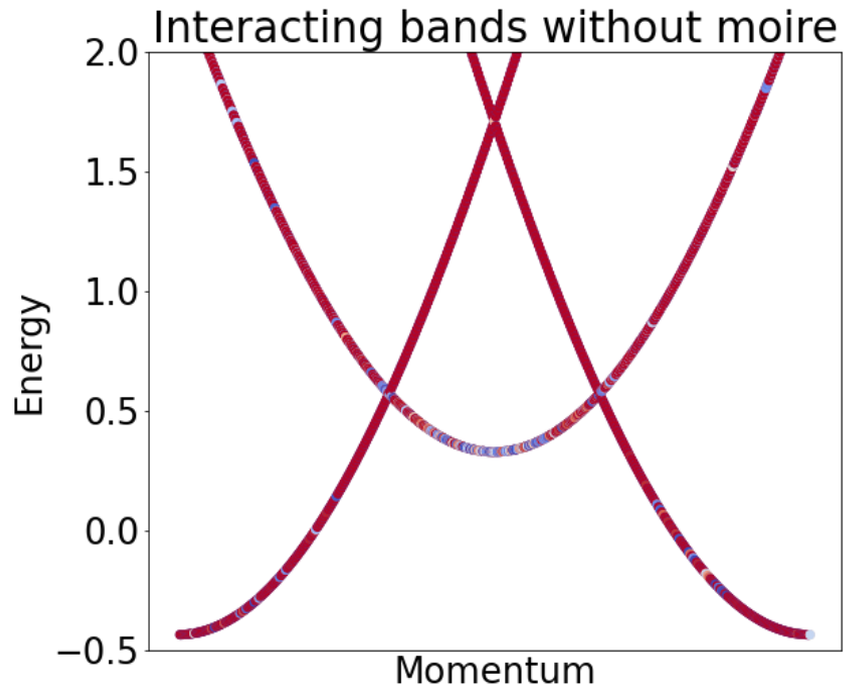
# Moire-enhanced DOS



**The moire potential gives rise to an enhanced DOS**

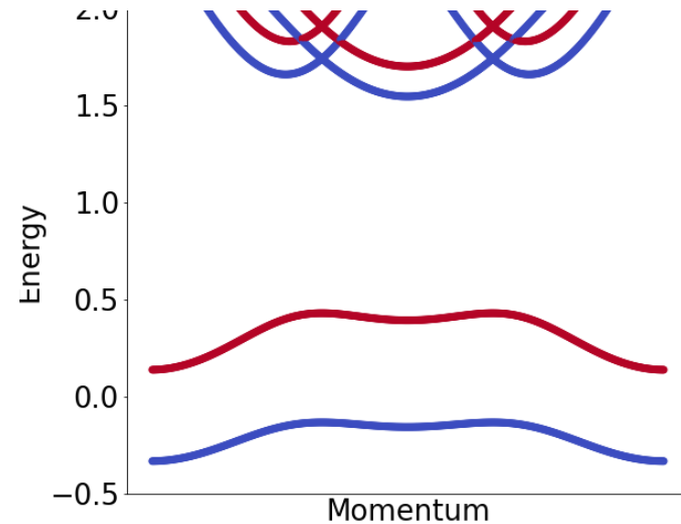
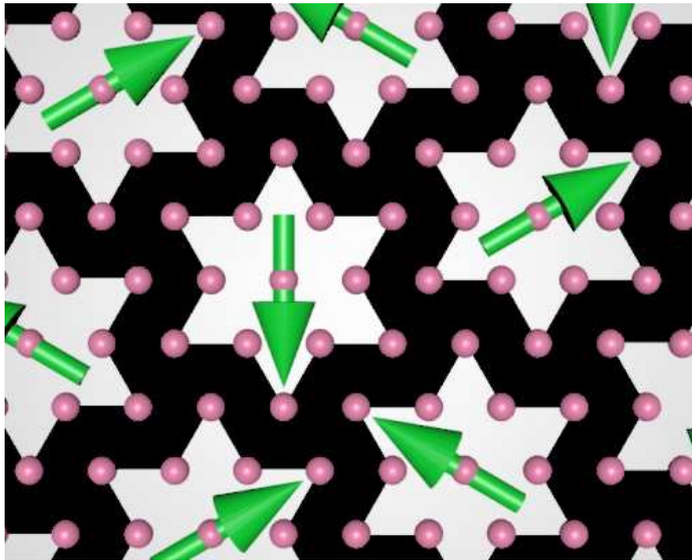
# Interactions in the absence and presence of a moire

$$H = \sum_{ij,s} t_{ij} c_{i,s}^\dagger c_{j,s} + \sum_i U c_{i\uparrow}^\dagger c_{i\uparrow} c_{i\downarrow}^\dagger c_{i\downarrow} + h.c.$$



# Correlated state in 1T-TaS<sub>2</sub> from miniband formation

Charge-density wave reconstruction, leading to a localized orbital in a  $\sqrt{13} \times \sqrt{13}$  unit cell



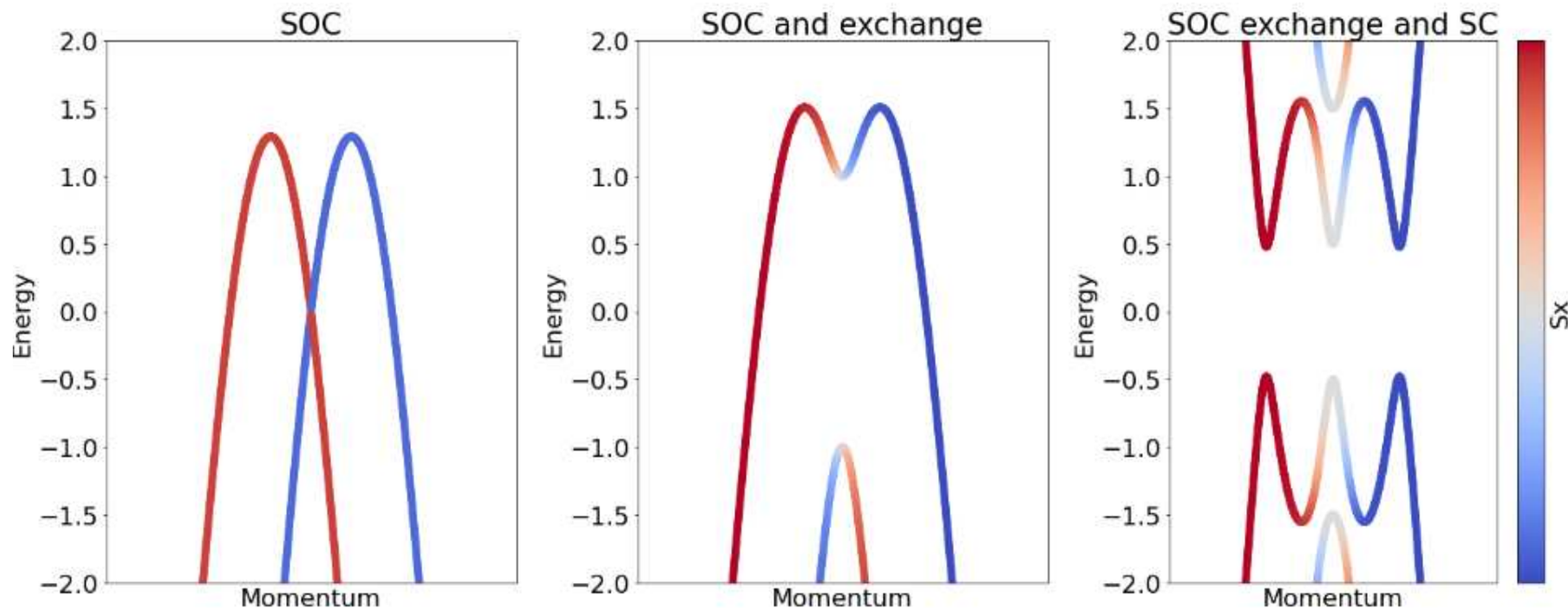
A charge density wave distortion acts as a superlattice modulation

Strong interactions give rise to local moment formation in the mini-bands

# Moire-driven topological states

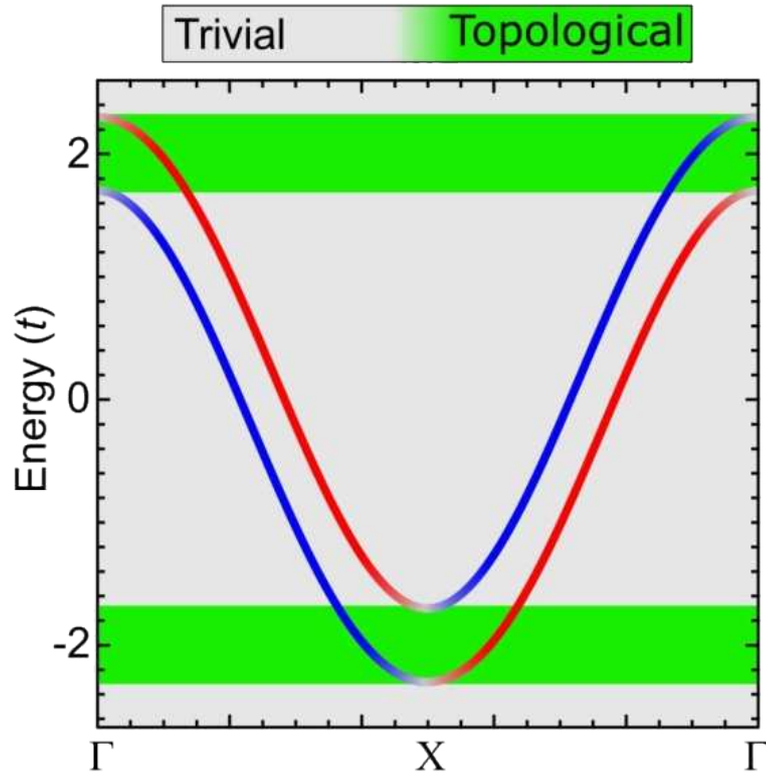
# Artificial topological superconductivity

## Bulk electronic structure



The combination of SOC and exchange creates helical states  
Superconductivity gaps out the helical states in a non-trivial way

# Artificial topological superconductor

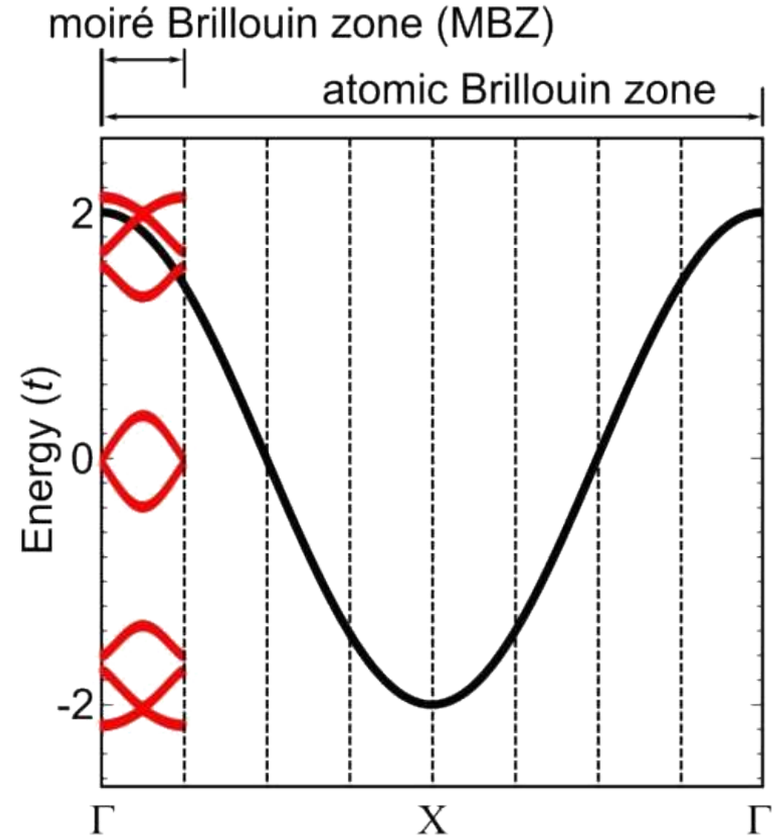
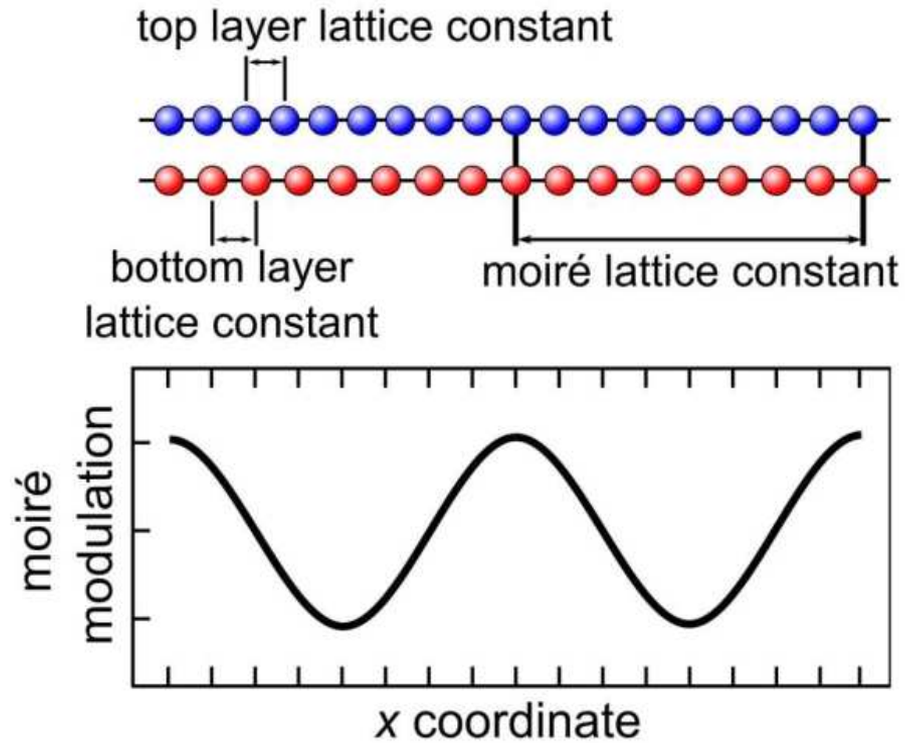


$$H = H_0 + H_J + H_R$$

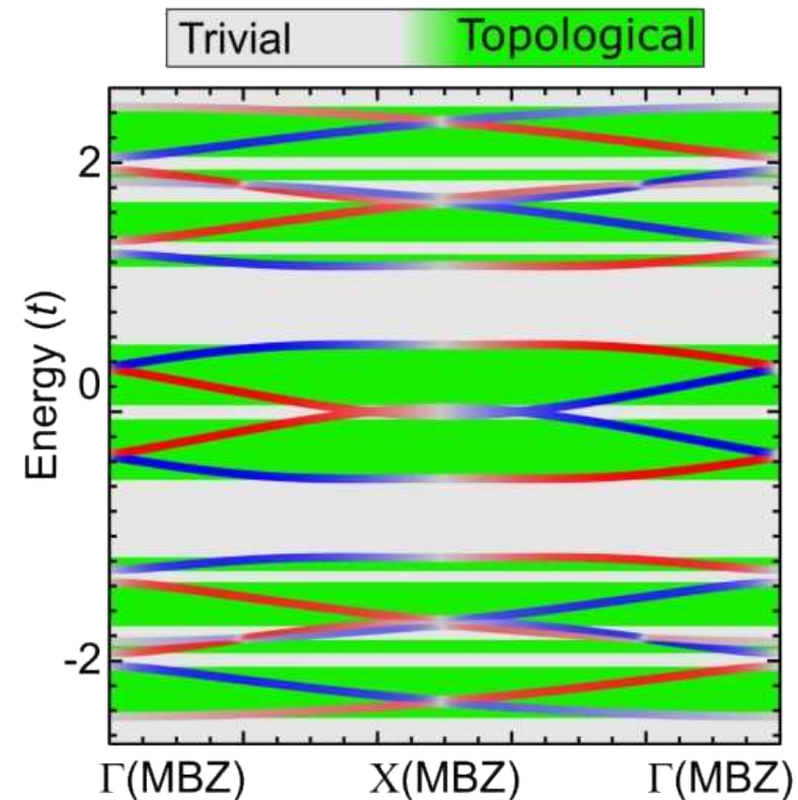
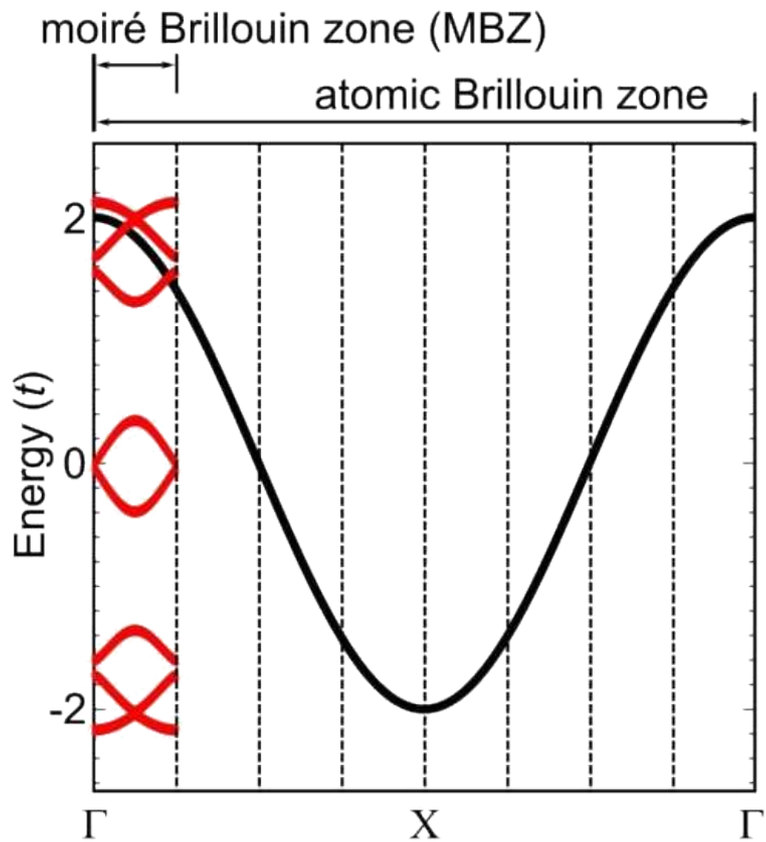
Kinetic energy  $\nearrow$  Exchange field  $\nearrow$  Rashba spin-orbit coupling  $\nearrow$

When switching on an s-wave pairing, green regions lead to topological superconductivity

# Artificial moiré topological superconductor

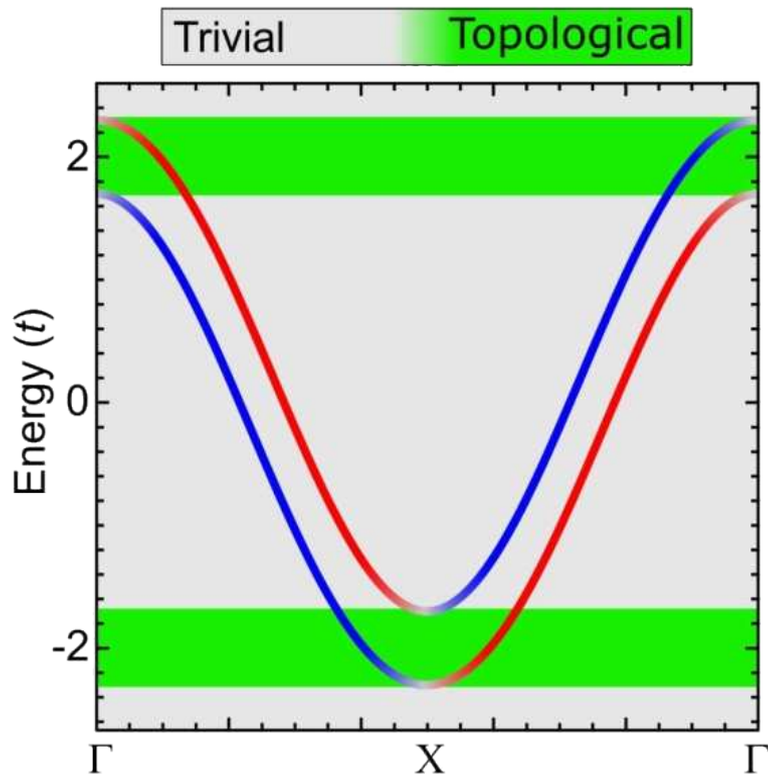


# Artificial moiré topological superconductor

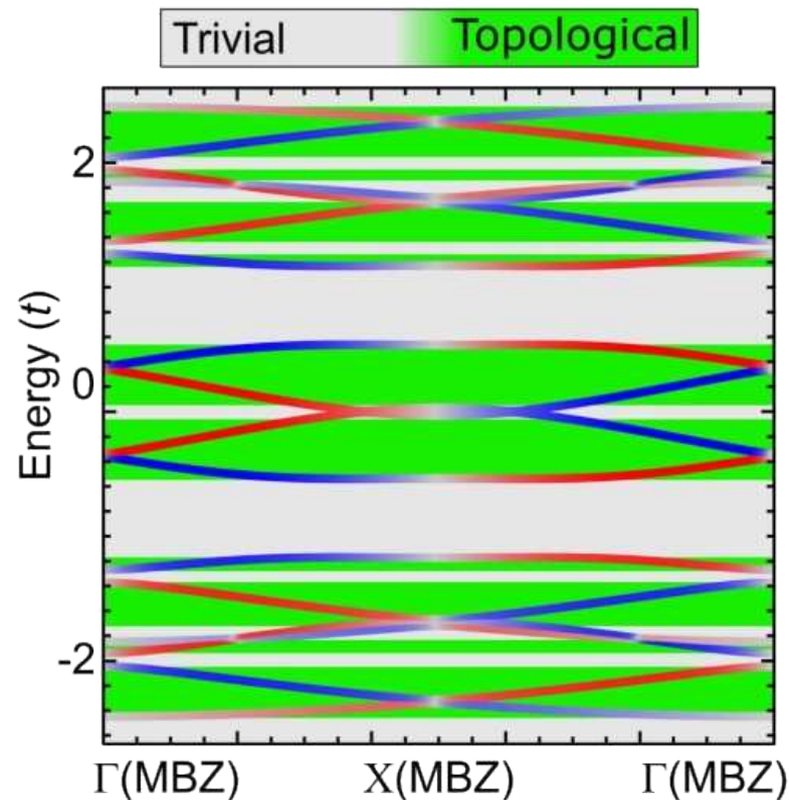


# Artificial moire topological superconductor

## Without moire

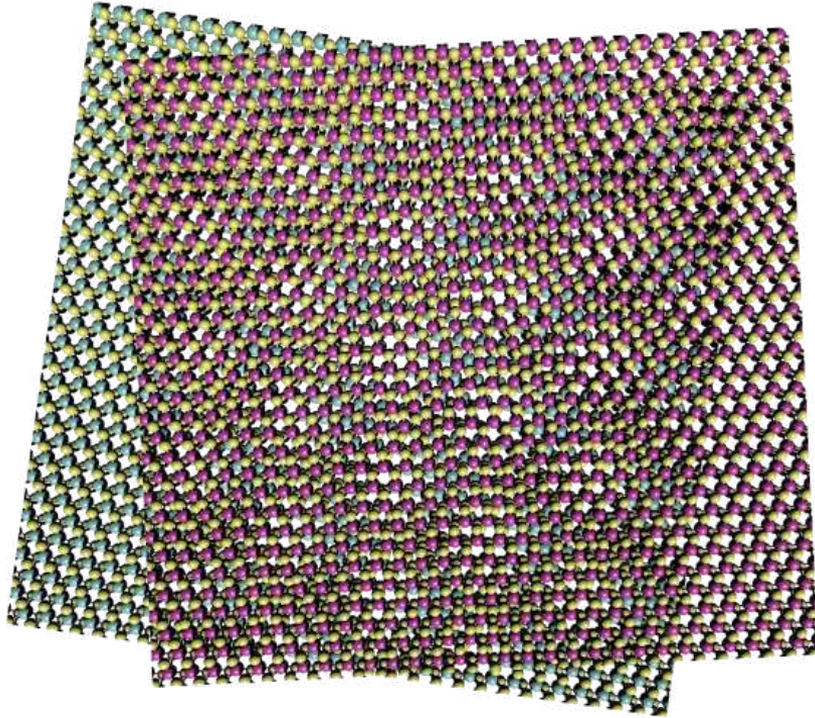


## With moire



# Moire driven gaps in the band structure

In twisted TMDC heterobilayers, the moire modulates the band off set of one layer



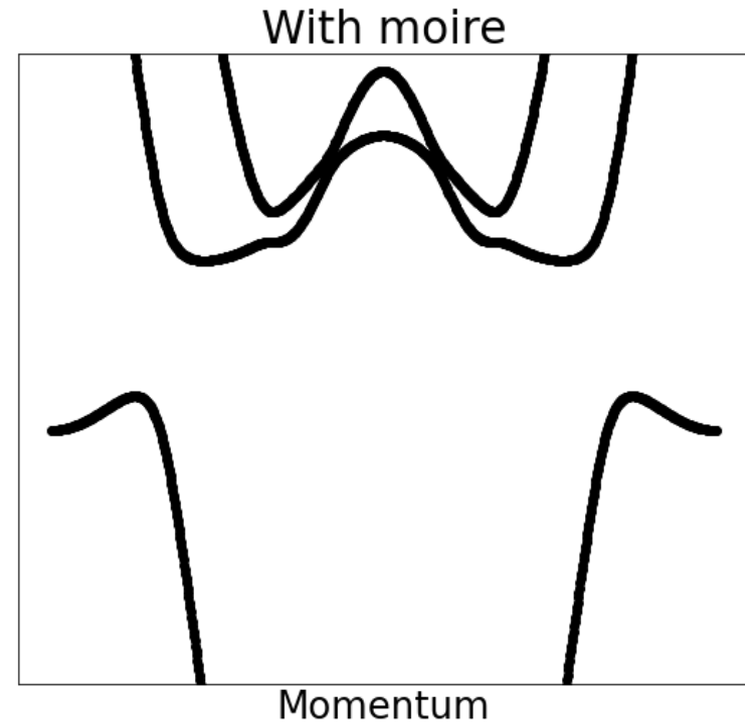
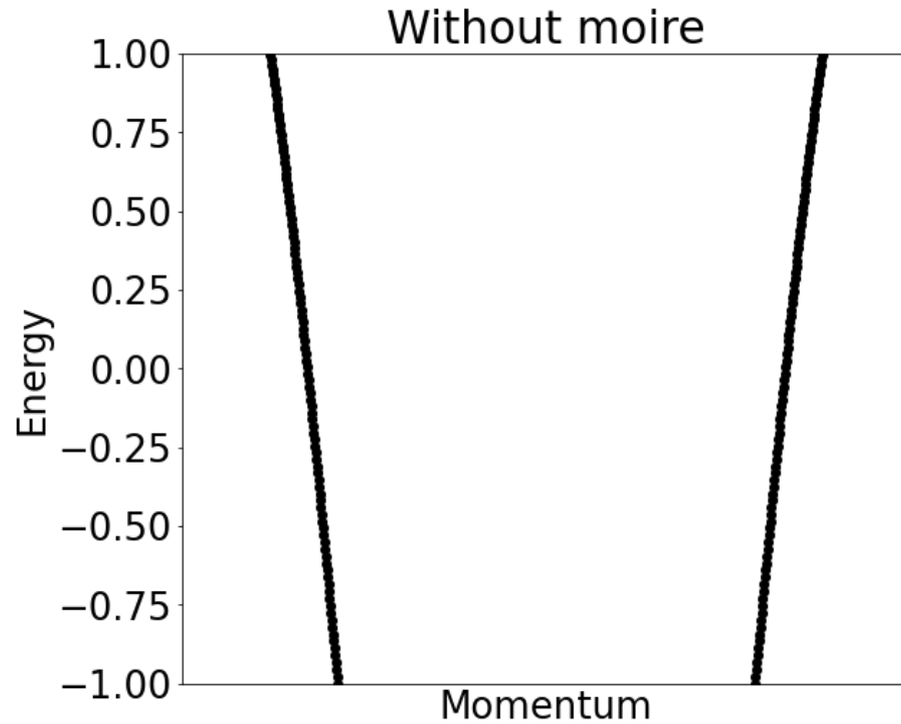
$$H = H_0 + H_J + H_R$$

Kinetic energy

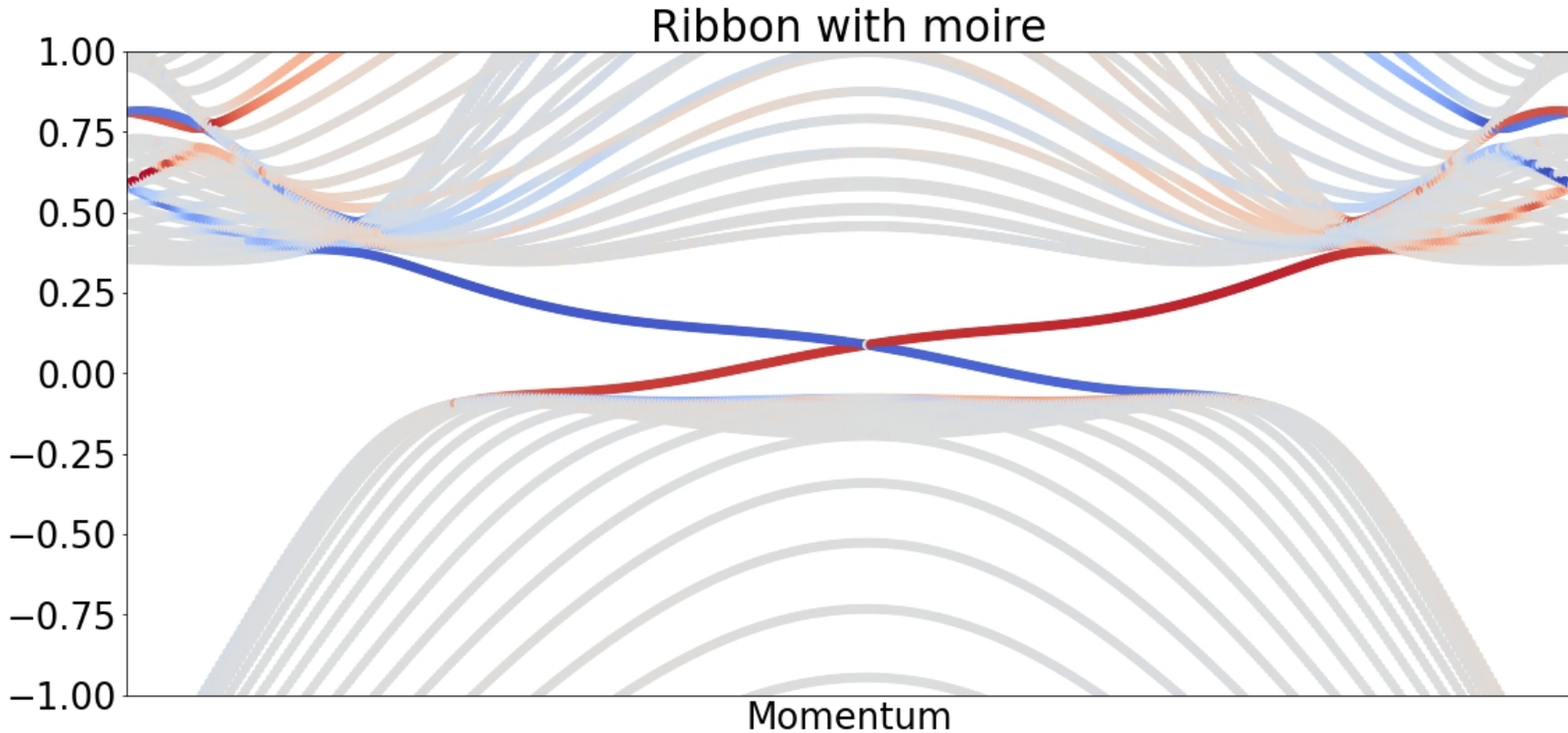
Exchange field

Rashba spin-orbit coupling

# Moire driven gaps in the band structure



# Moire driven gaps in the band structure



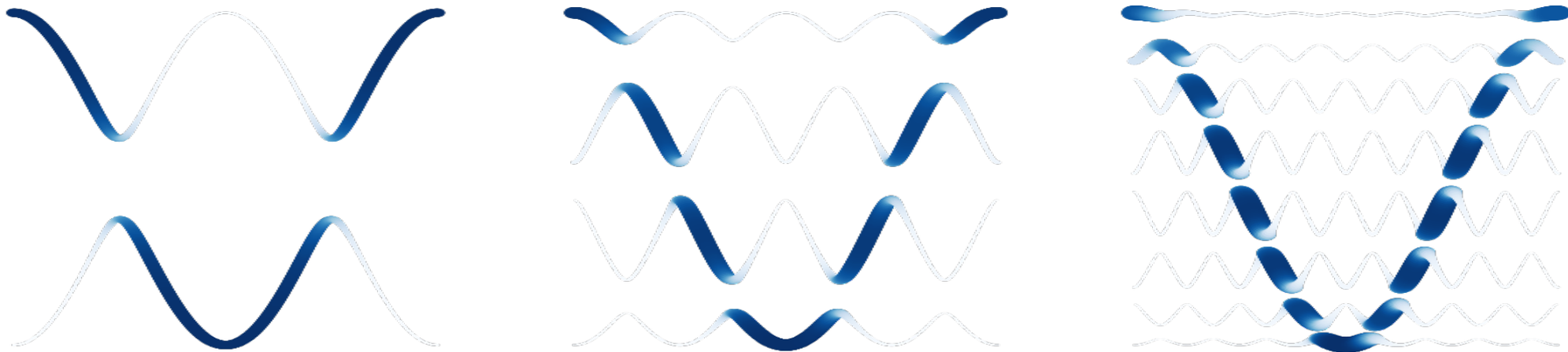
Chiral states appear in a ribbon driven by the moire

# Break

10-15 min break

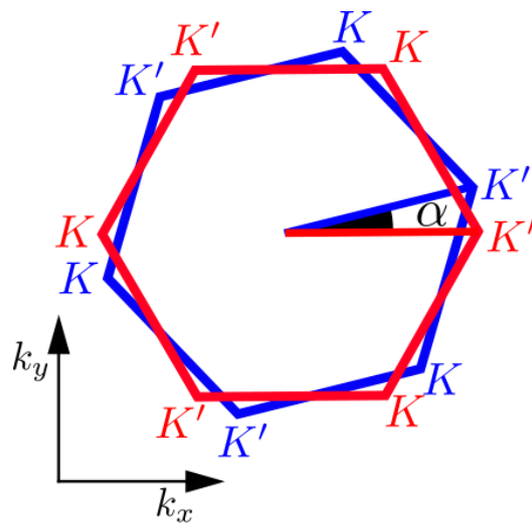
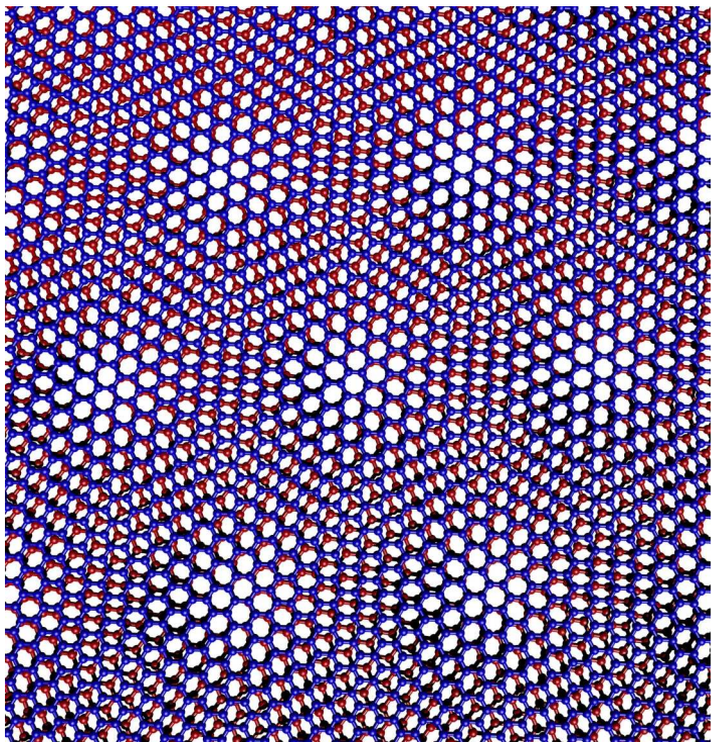
*(optional) to discuss during the break*

Which band structure has more van Hove singularities?



# Electronic structure of twisted graphene multilayers

# Twisted bilayer graphene

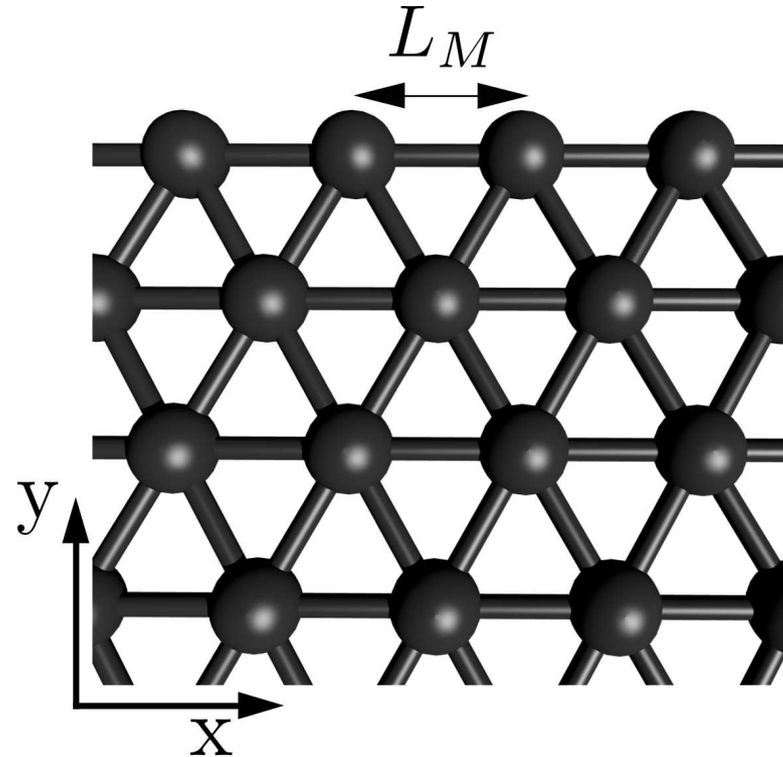
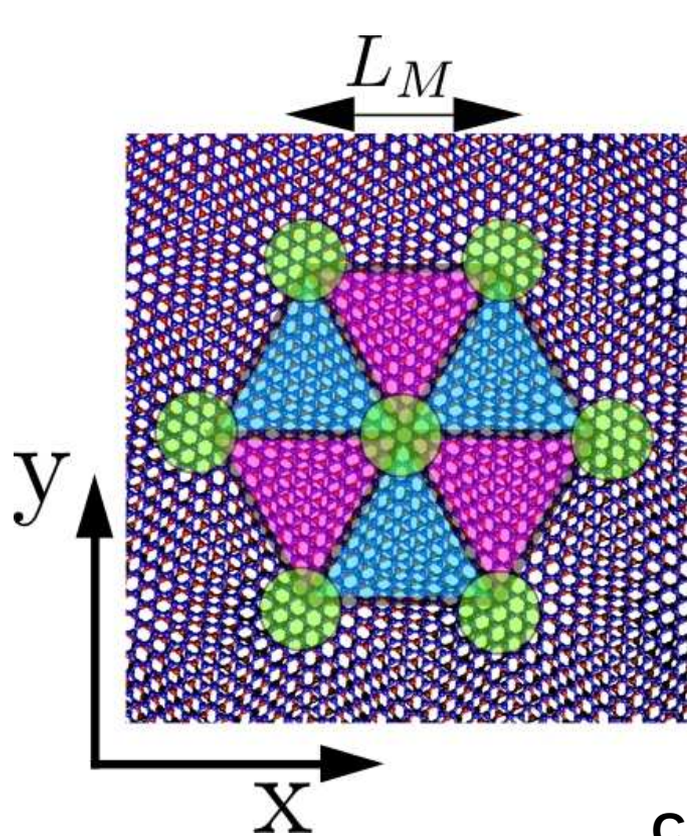


**Additional parameters in the system**

Angle between the layers  $\alpha$

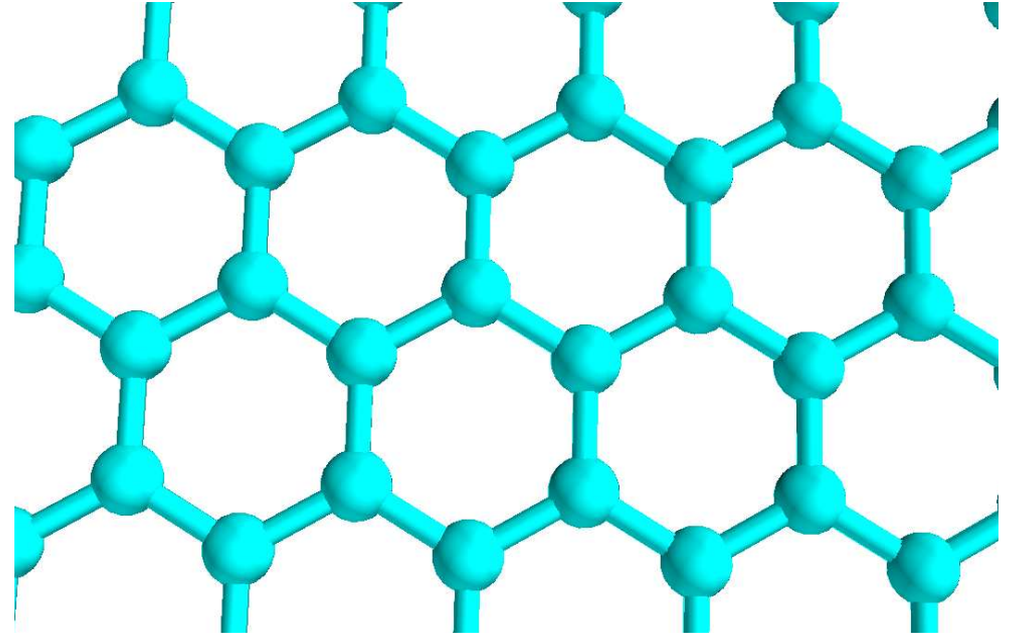
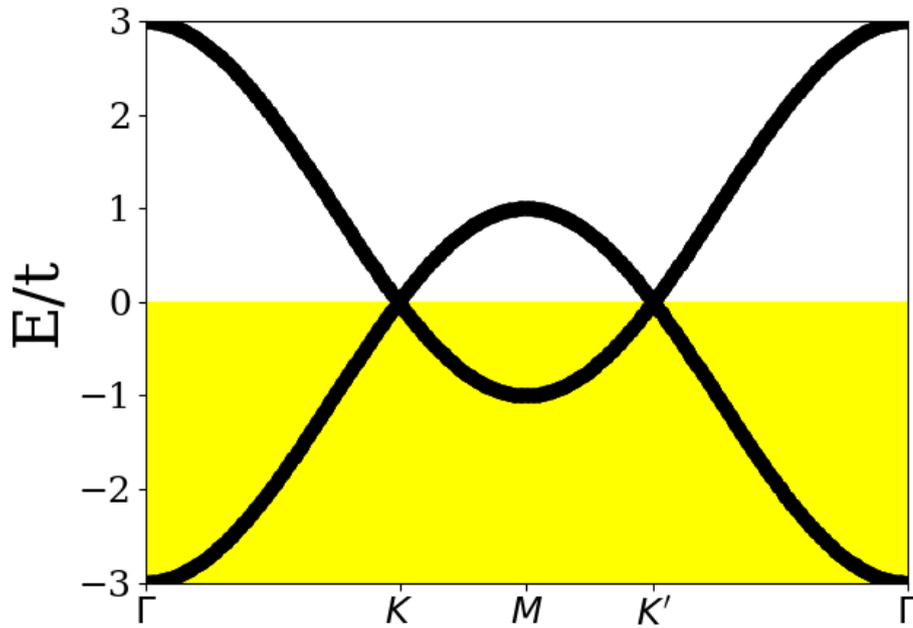
Bias between the layers  $U$

# A new length scale: the moire length



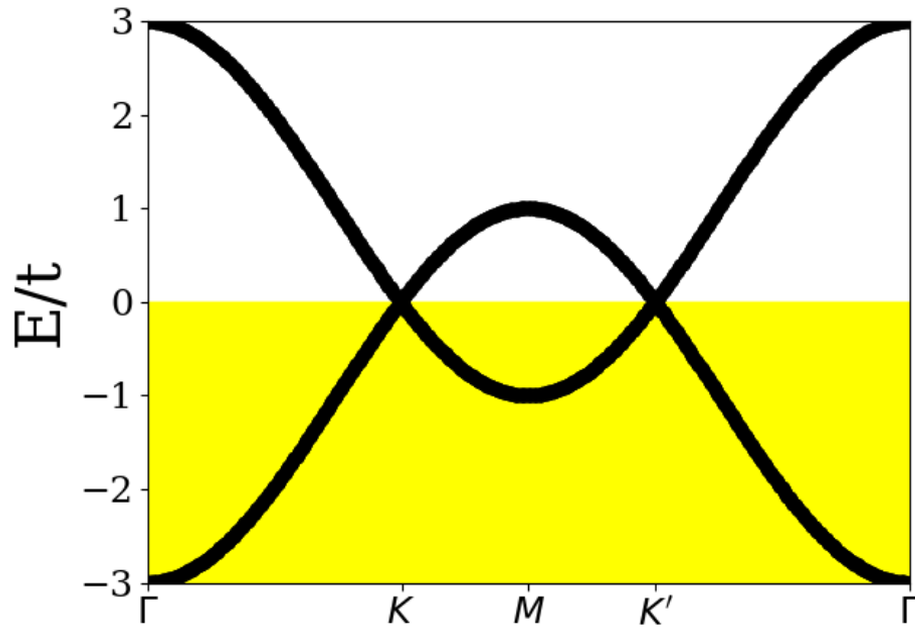
Creating effective lattices with tunable lattice constants

# Structure of graphene



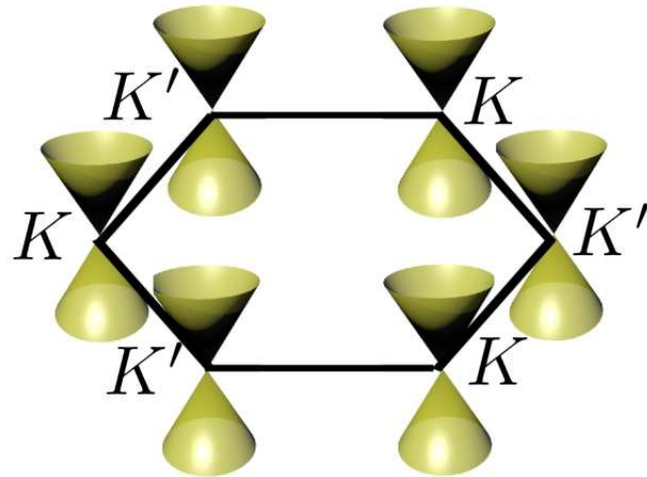
**Minimal model: Single orbital in a honeycomb lattice**

# The electronic structure of graphene

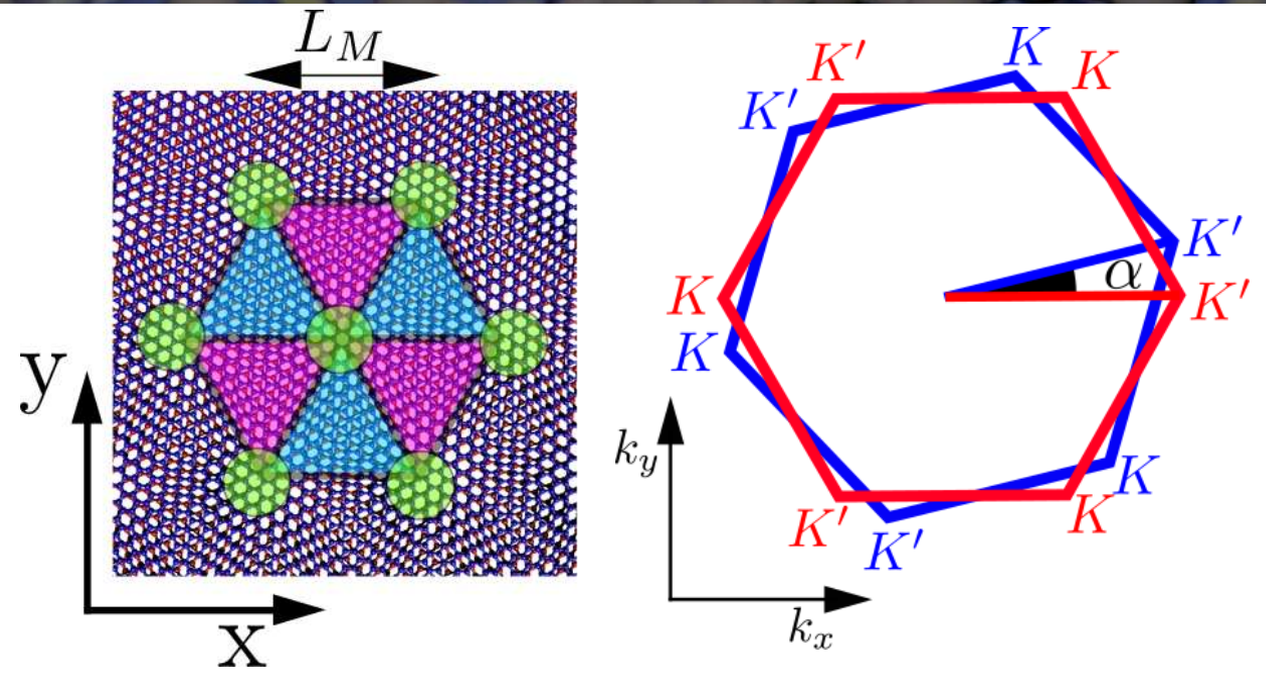


$$H = \begin{pmatrix} 0 & p_x \pm ip_y \\ p_x \mp ip_y & 0 \end{pmatrix}$$

$$H = p_x \sigma_x \pm p_y \sigma_y$$



# Real and reciprocal space in twisted bilayer graphene



Intralayer

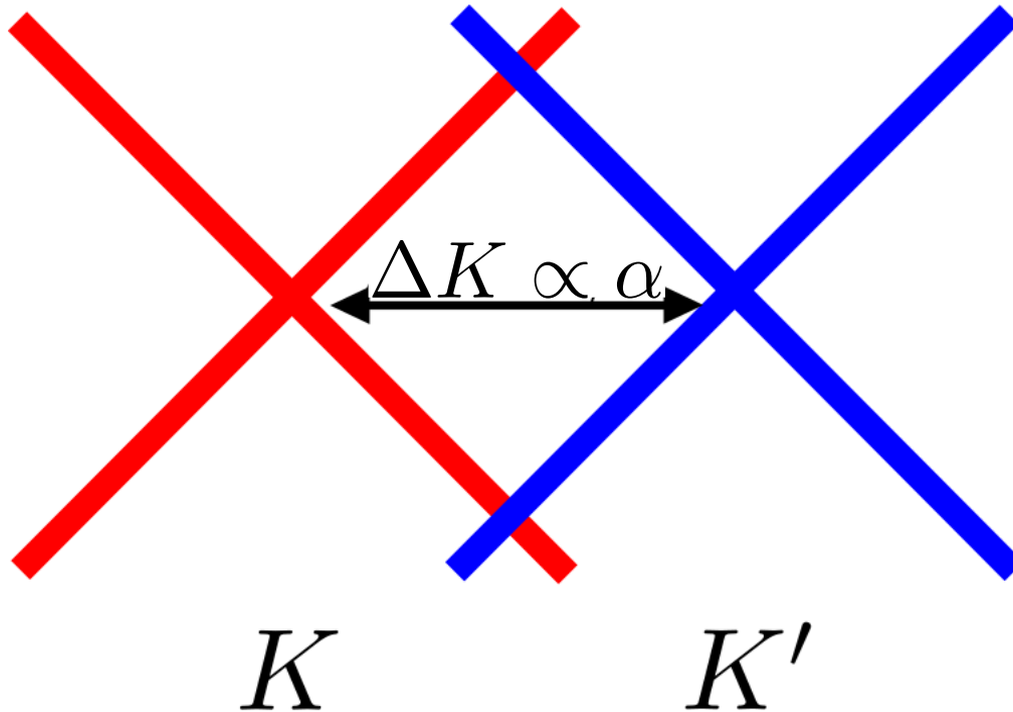
Interlayer

Bias

$$H = t \sum_{\langle ij \rangle} c_i^\dagger c_j + \sum_{ij} \hat{t}_\perp(\mathbf{r}_i, \mathbf{r}_j) c_i^\dagger c_j + U \sum_i \tau_z^{ii} c_i^\dagger c_i,$$

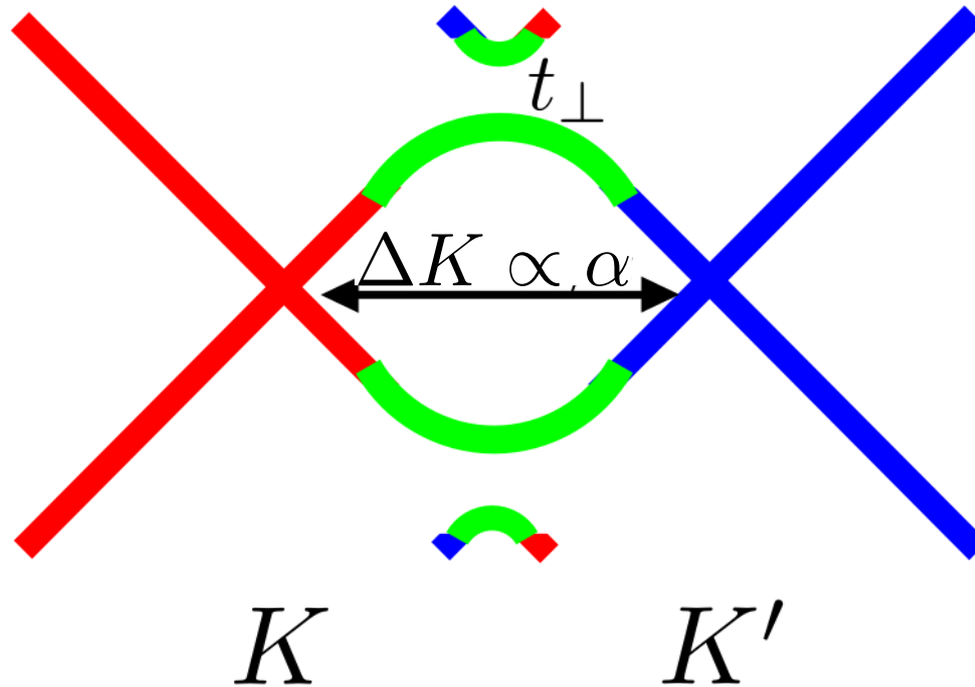
# Velocity renormalization at large angle

$$\bar{v}_F/v_F = 1 - 9[t_{\perp}/v_F\Delta K]$$



# Velocity renormalization at large angle

$$\bar{v}_F/v_F = 1 - 9[t_{\perp}/v_F\Delta K]$$

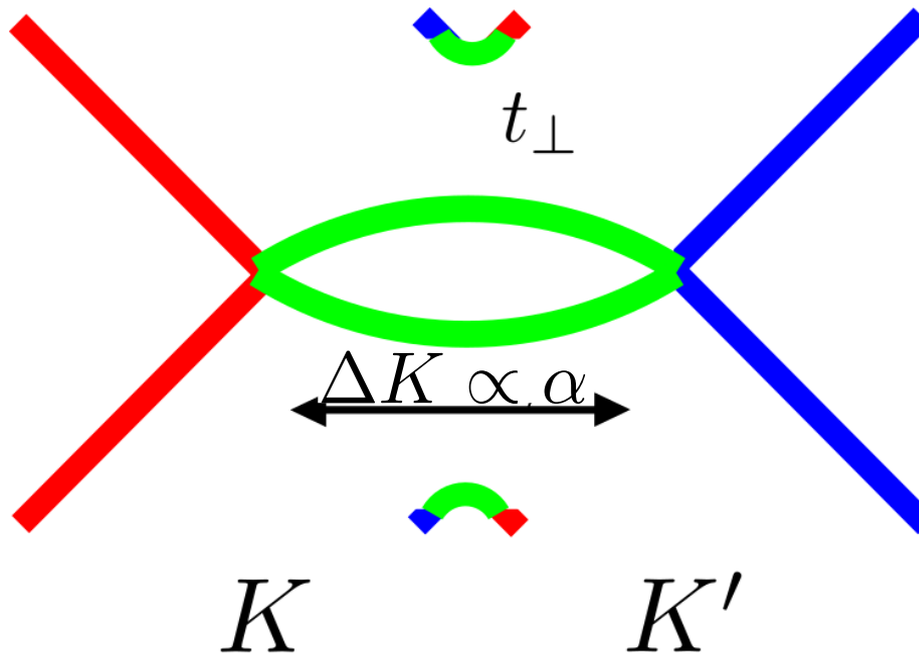


Relevant ratio

$$\frac{t_{\perp}}{\alpha}$$

# Velocity renormalization at large angle

$$\bar{v}_F/v_F = 1 - 9[t_{\perp}/v_F\Delta K]$$

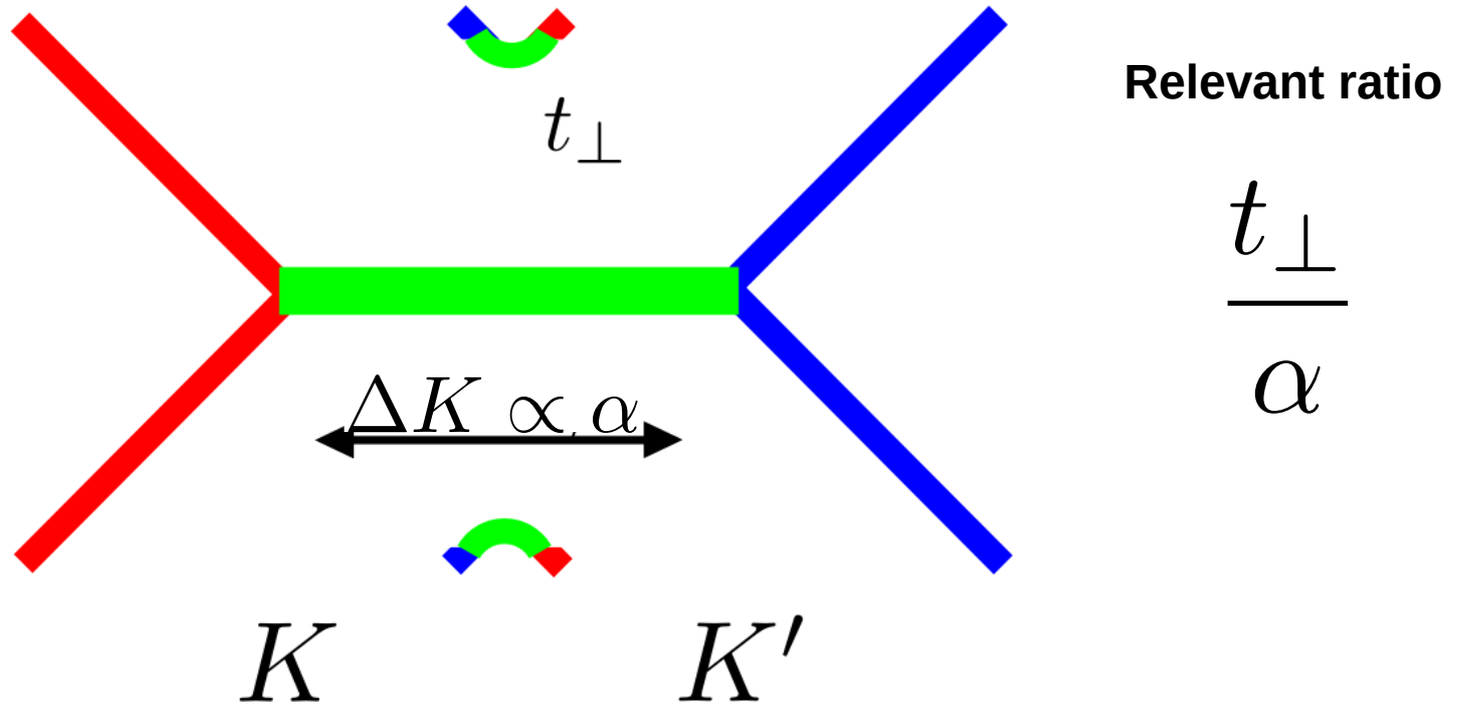


Relevant ratio

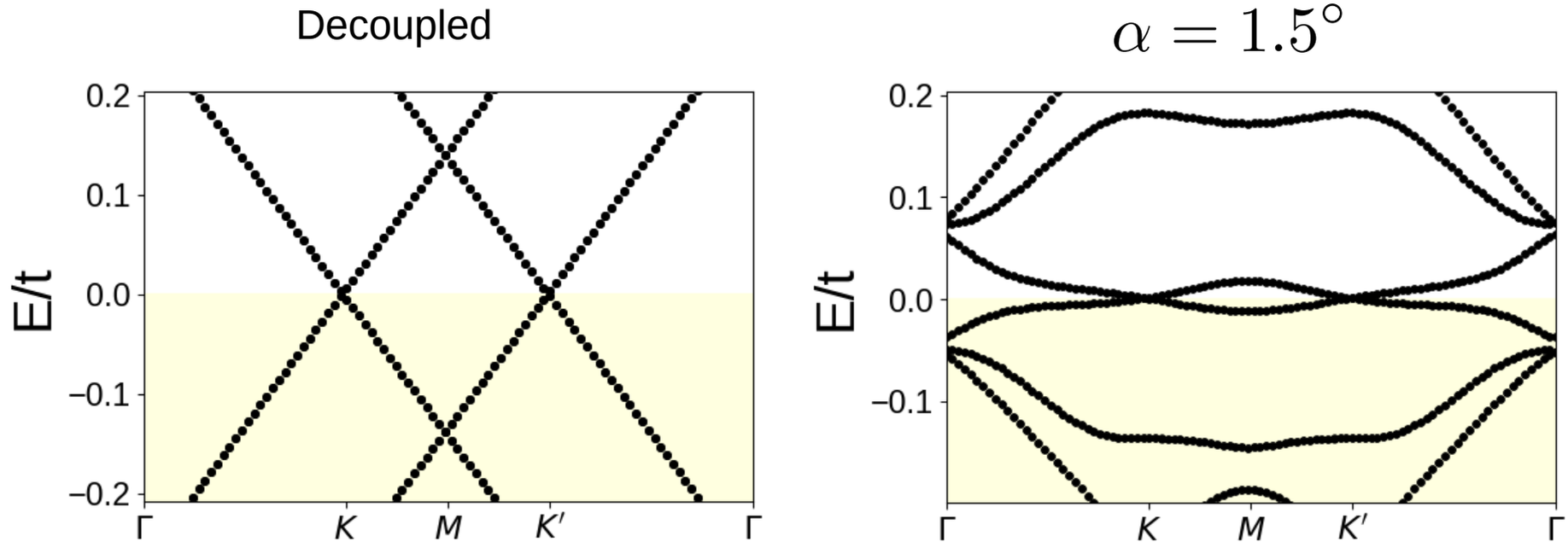
$$\frac{t_{\perp}}{\alpha}$$

# Velocity renormalization

$$\bar{v}_F/v_F = 1 - 9[t_\perp/v_F\Delta K]$$



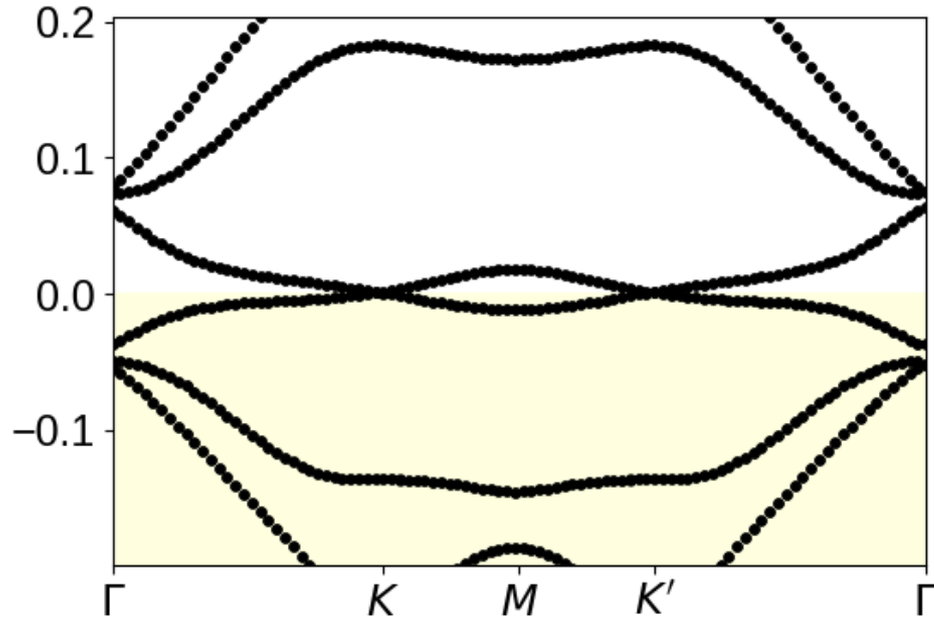
# Band structure of twisted bilayer graphene



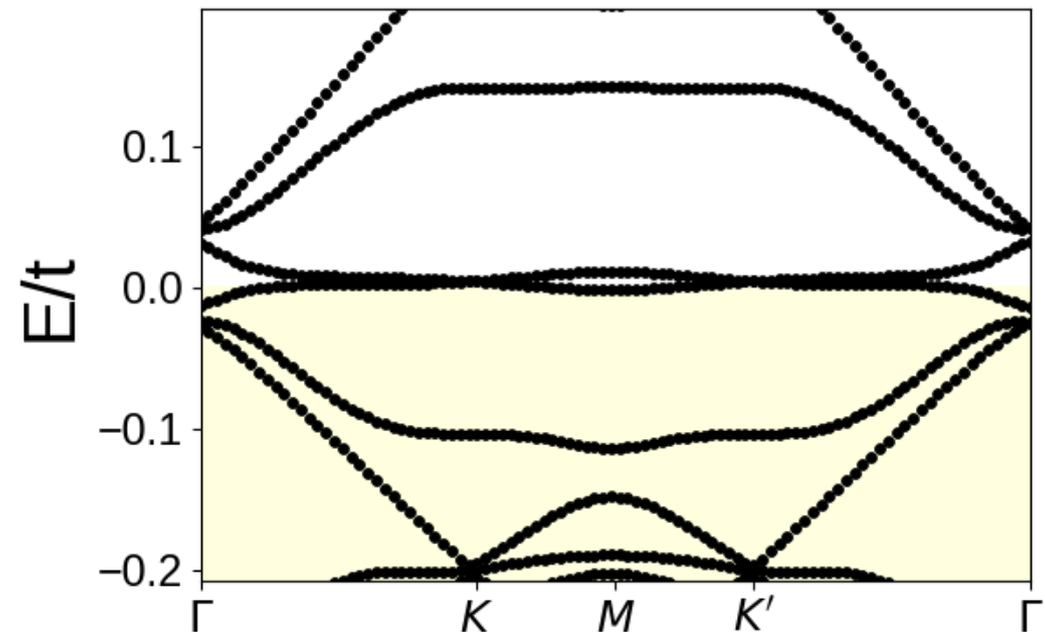
As the angle between layers is decreased, the bands become flatter

# Band structure of twisted bilayer graphene

$$\alpha = 1.5^\circ$$

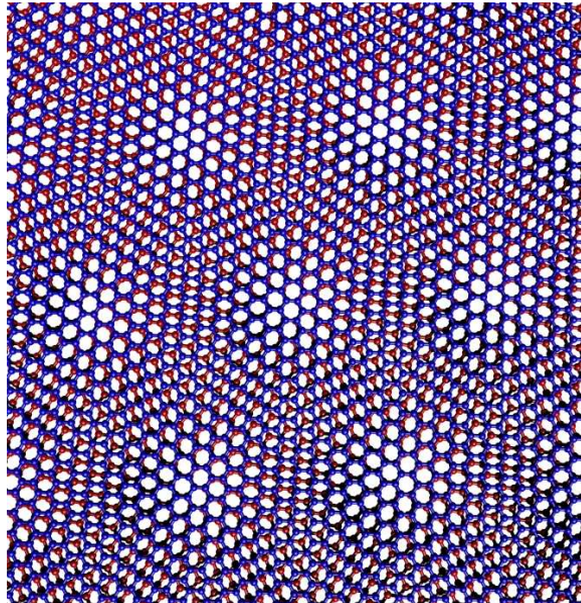


$$\alpha = 1.2^\circ$$

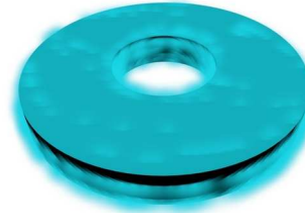


As the rotation angle approached 1 degree, the lowest band becomes flatter

# Correlations in twisted bilayer graphene flat bands

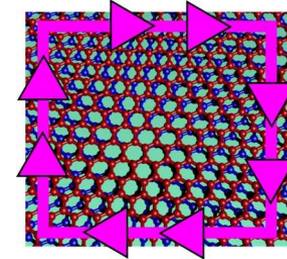


## Superconductivity



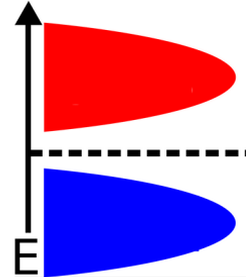
*Nature* 556, 43–50 (2018)

## Chern insulators



*Science* 365, 605-608 (2019)

## Correlated insulators



*Nature* 556, 80–84 (2018)

## Fractional Chern insulators

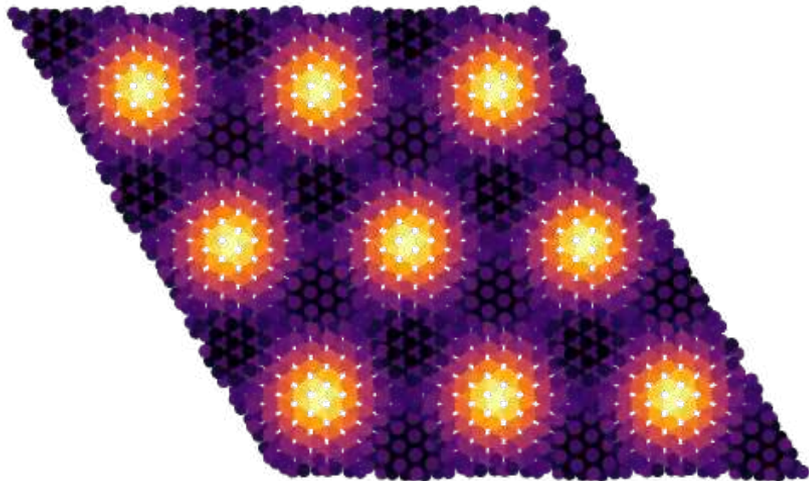


*Nature* 600, 439–443 (2021)

**Correlated states in flat bands give rise to a wide variety of phenomena**

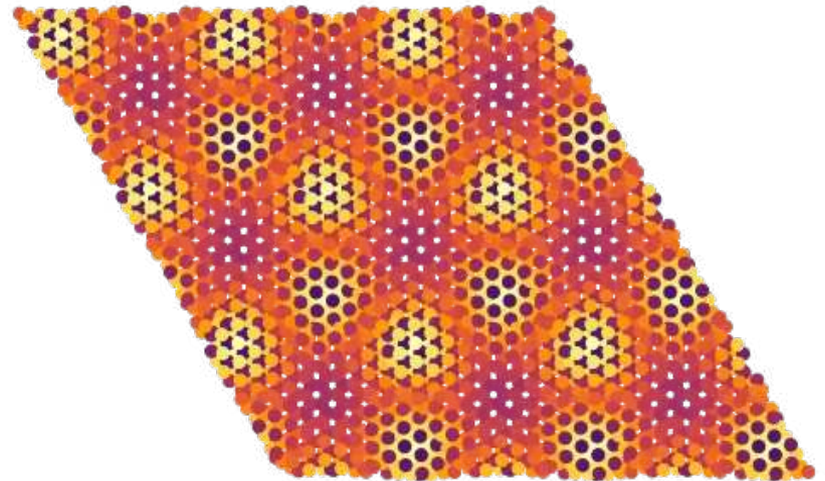
# Distribution of states in the twisted graphene bilayer

$E = 0.0$



*Triangular*

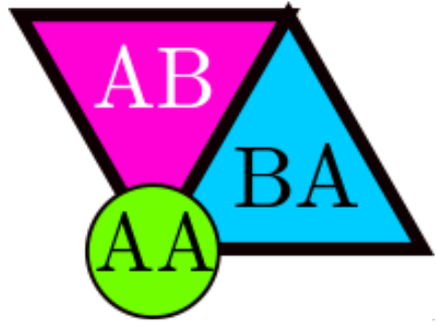
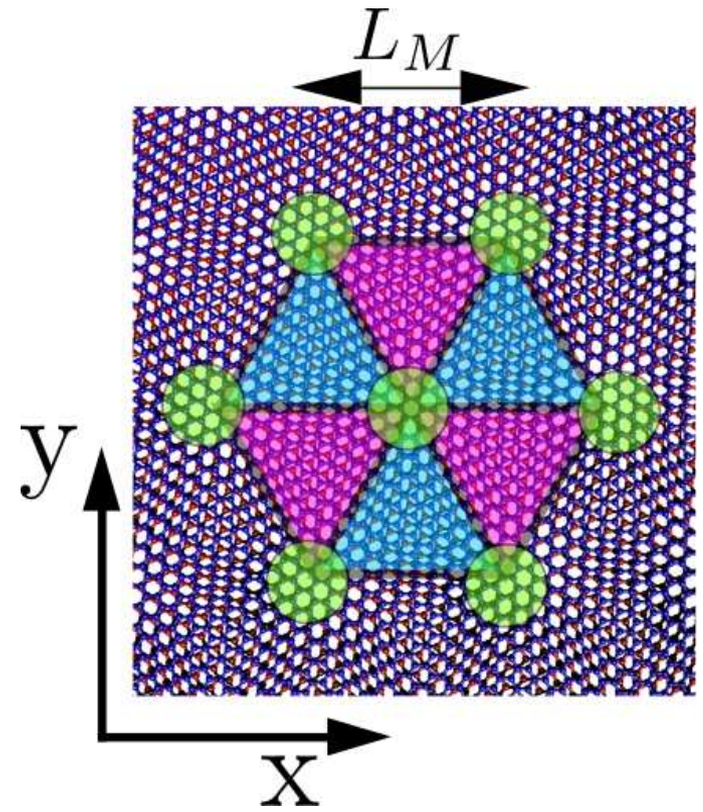
$E = 0.25$



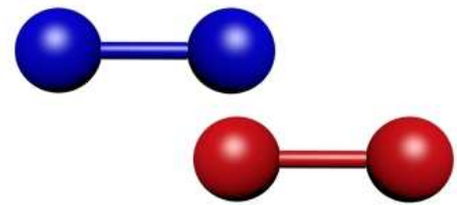
*Hexagonal*

The spatial distribution of the states is highly dependent on the energy

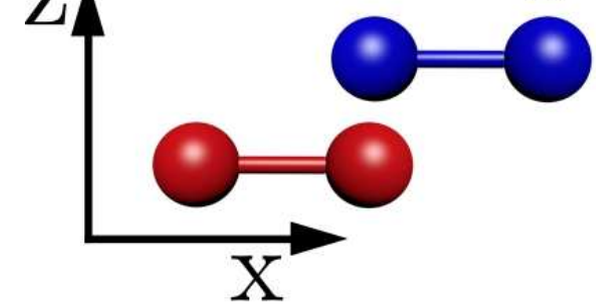
# Stacking of twisted bilayer graphene



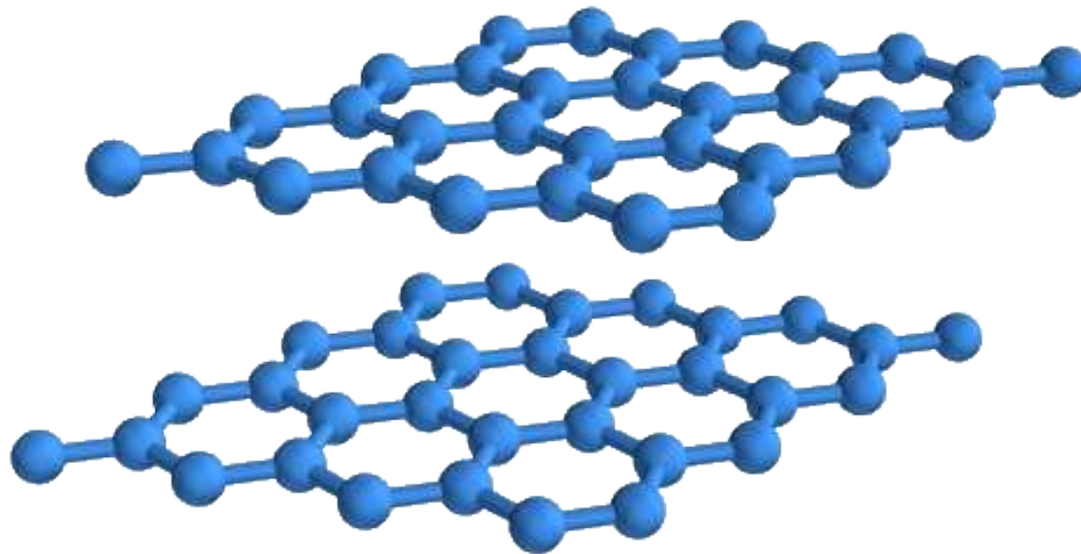
AB stacking



BA stacking



# Bias in AB bilayer graphene



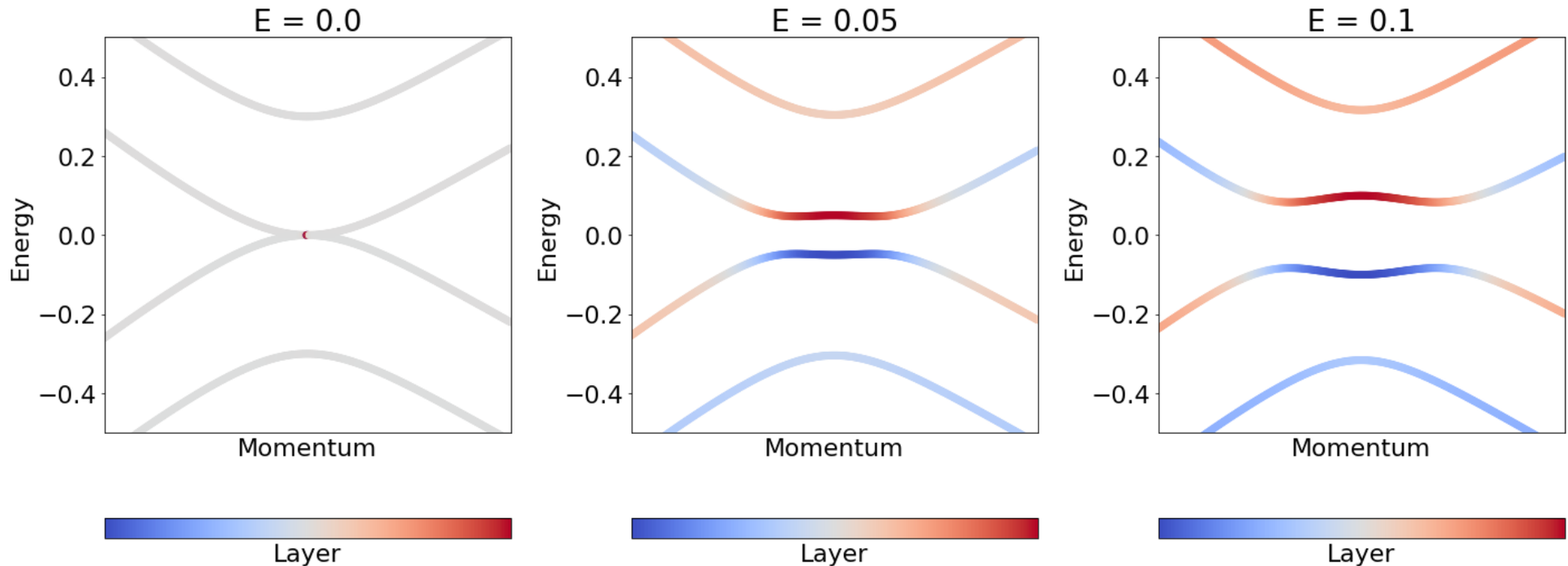
$$V c_n^\dagger c_n$$

$$-V c_n^\dagger c_n$$

Let us look at the impact of a bias in an aligned graphene bilayer

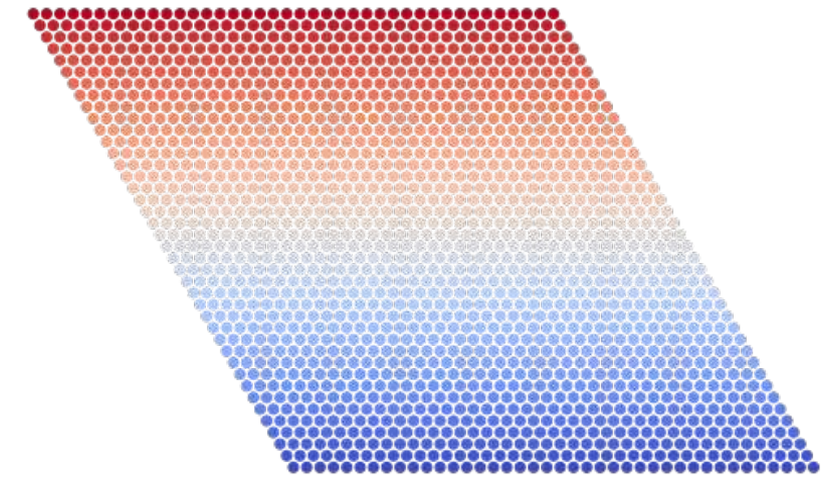
# The electronic structure of bilayer graphene

Graphene bilayers open a gap when an interlayer bias is applied



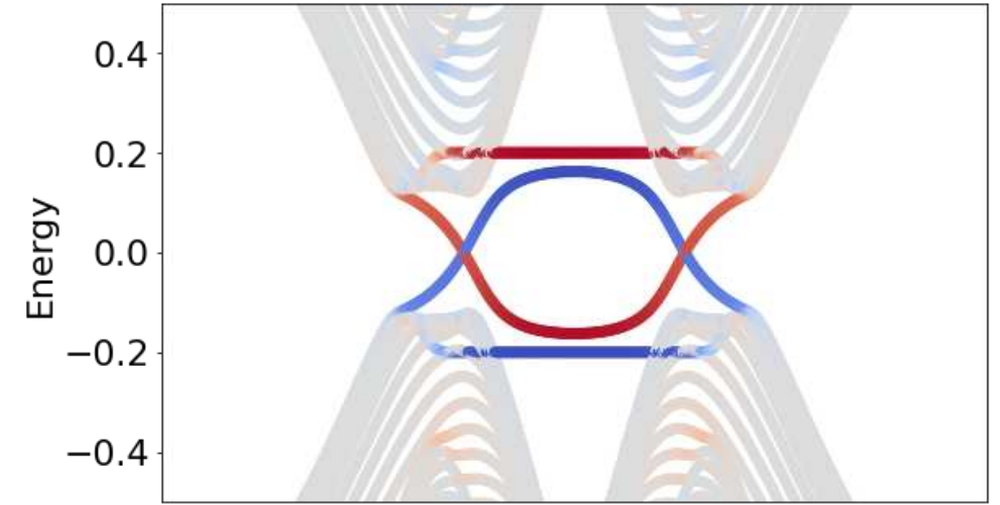
# Biased bilayer graphene, pseudo-helical states

Position operator



edge/bulk/edge

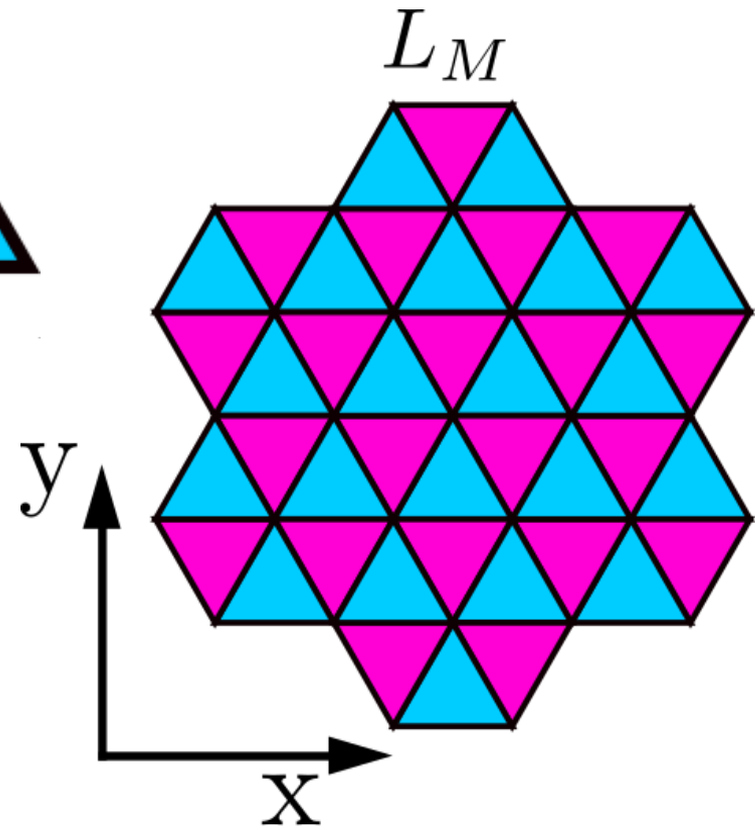
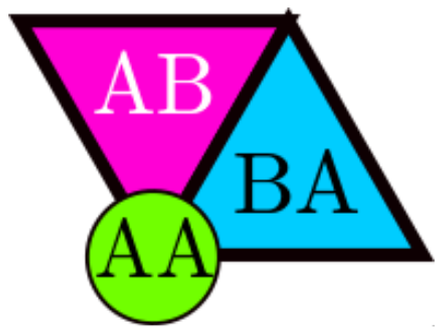
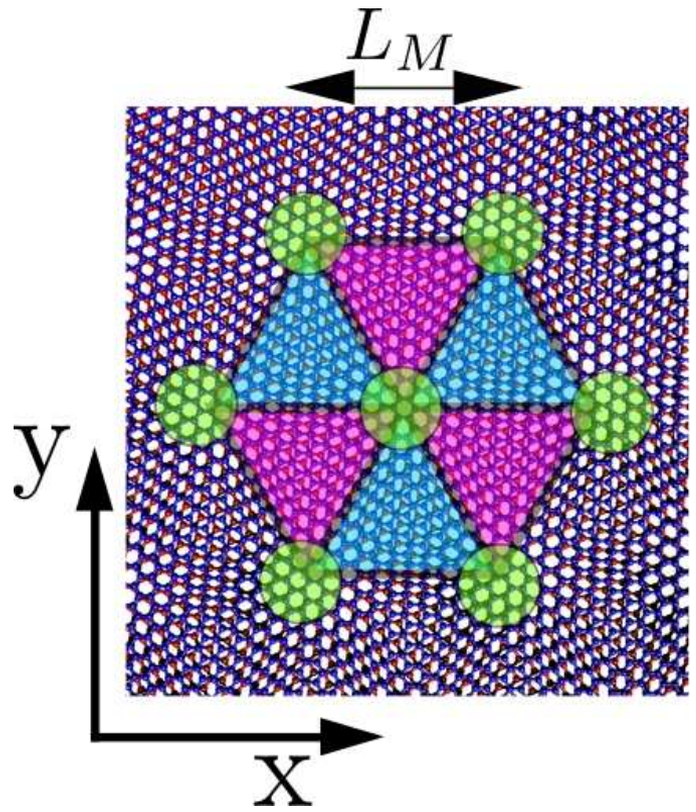
Band structure



edge/bulk/edge

Edge states appear in the presence of a bias between layers

# Stacking of twisted bilayer graphene

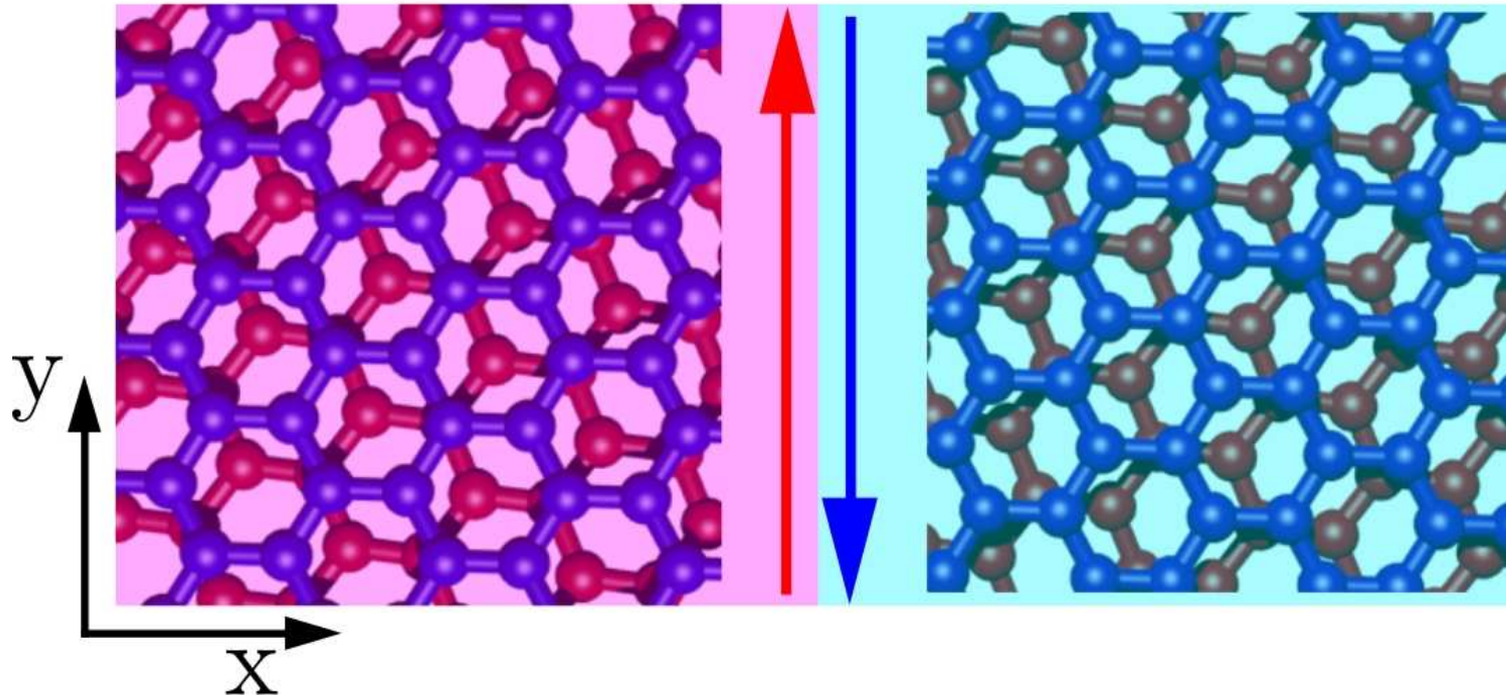


# Interfacial modes in biased TBG

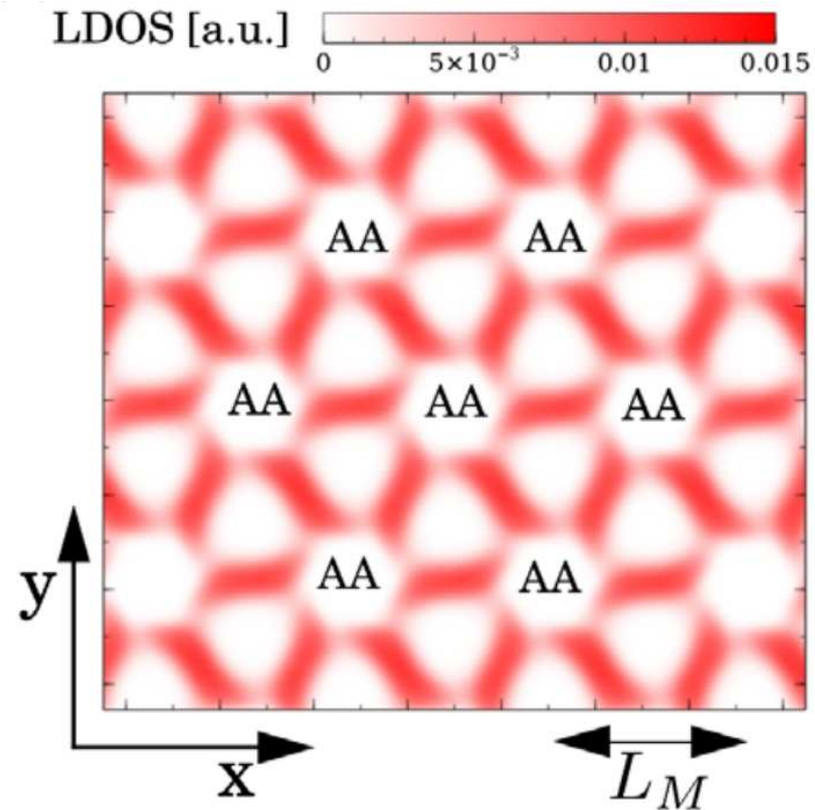
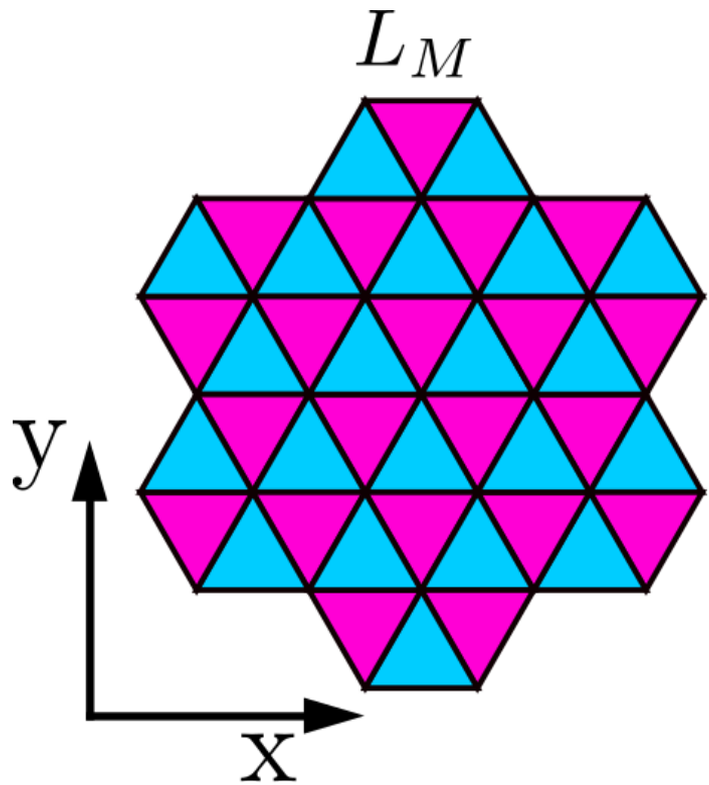
AB stacking

**K** **K'**

BA stacking

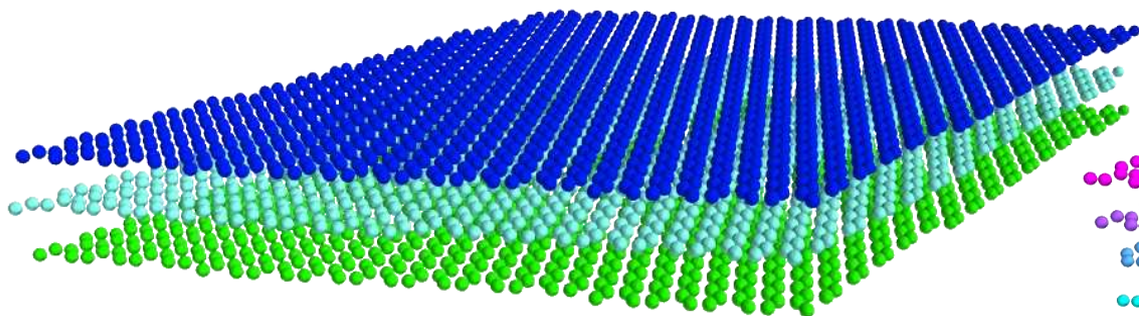


# Helical networks in TBG

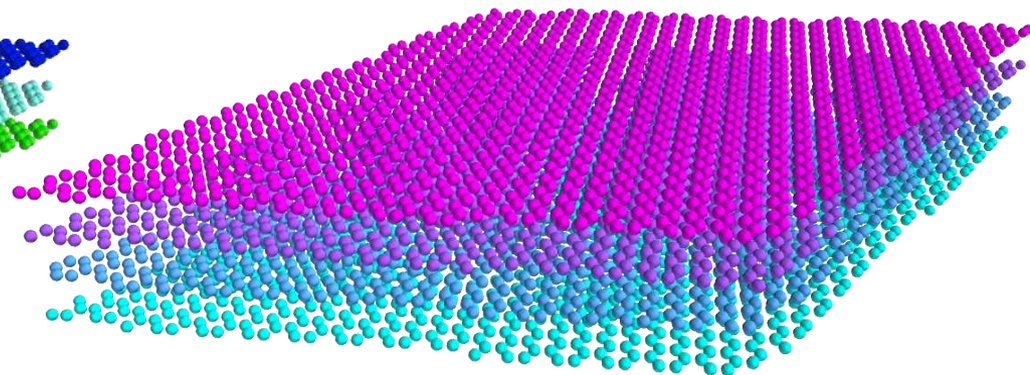


# Other twisted graphene multilayers

Twisted graphene trilayer

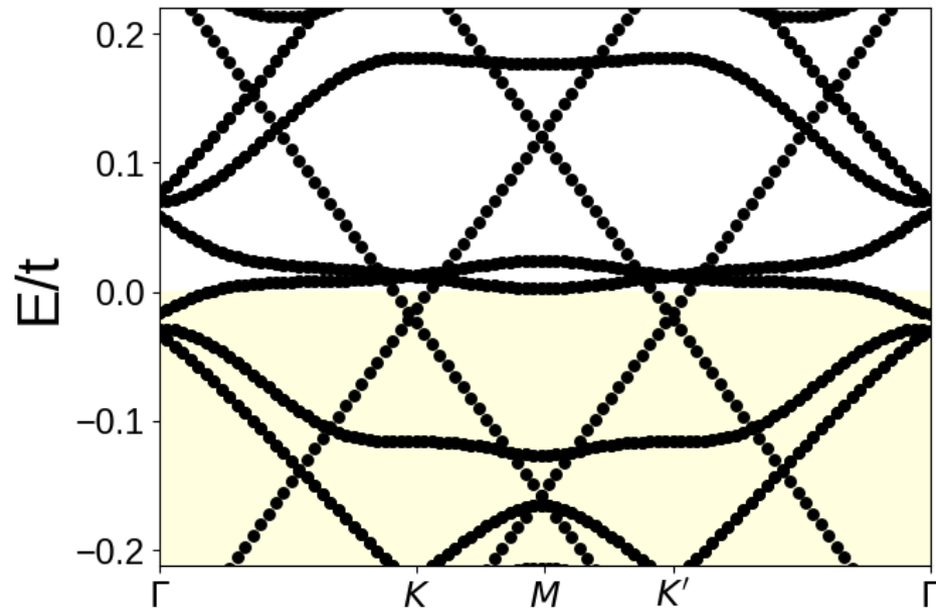


Twisted graphene double bilayer

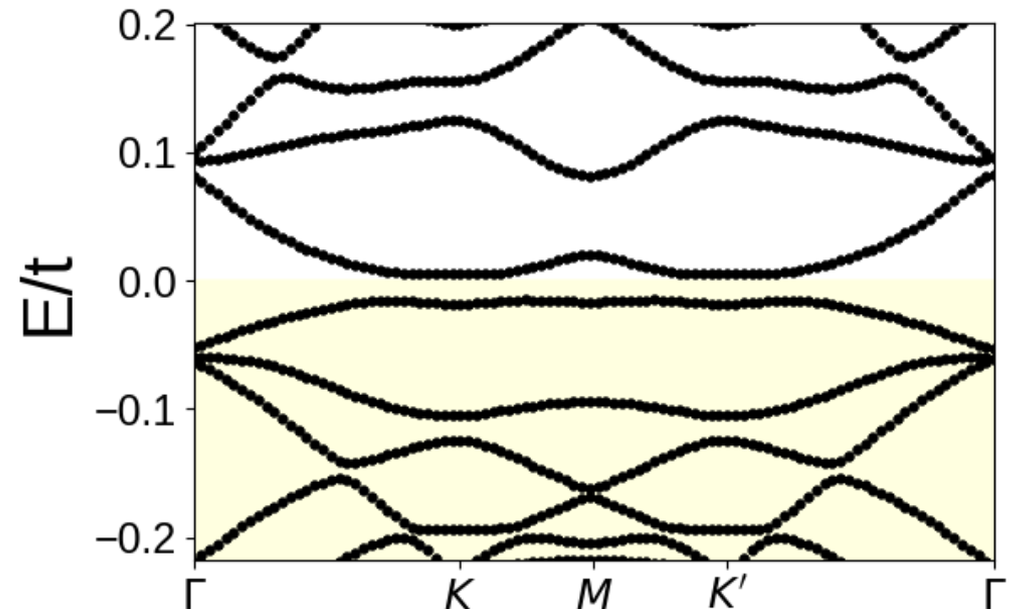


# Other twisted graphene multilayers

Twisted graphene trilayer



Twisted graphene double bilayer



Flat and dispersive moire bands appear in generic twisted graphene multilayers

# For the exercise session this afternoon

## Download the Jupyter-notebook from

[https://github.com/joselado/jyvaskyla\\_summer\\_school\\_2022/blob/main/sessions/session4.ipynb](https://github.com/joselado/jyvaskyla_summer_school_2022/blob/main/sessions/session4.ipynb)

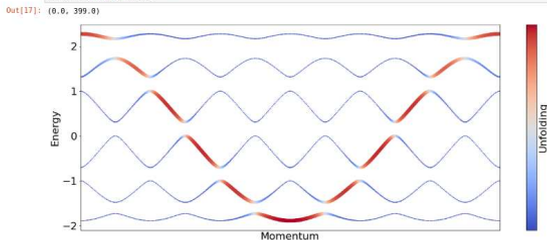
## The tasks during the exercise sessions

*You will see examples with the code*

```
In [17]: # let us now add an impurity in the previous supercell, and see how it leads to anticrossings in the bands
from pyquanta import geometry
g = geometry.Chain(1) # generate chain
N = 6 # size of the supercell
g = g.get_supercell(N, store_primal=True) # generate a supercell, store_primal is required for unfolding
h = g.get_hamiltonian() # and generate the Hamiltonian

# let us define a potential for an impurity
r0 = g.r(0) # location of the impurity
foot = lambda r: 1 - ((r-r0).dot(r-r0))**2 # impurity in site r0
# from pyquanta import potentials; foot = potentials.Impurity(r0,v=1.) # this is equivalent

h.add onsite(foot) # and add the impurity potential
kpath = g.get_kpath()*N # compute in the original Brillouin zone, just by extending the reciprocal vectors
(E, e, c) = h.get_bands(operators="unfold", kpath=kpath) # compute band structure
plt.scatter(E, c, c=c, s=c*8, 5); plt.xlabel("Momentum"); plt.ylabel("Energy"); plt.xticks([]) # plot
plt.colorbar(label="Unfolding", ticks=[]);
plt.xlim(0, 2*np.pi)
```



*You have to modify them, and answer questions*

### Exercise

- Change the size of the supercell. Do the anticrossings appear at the same energies?
- If the strength of the impurity becomes very large, what happens to the electronic structure? Discuss why it has the behavior observed