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#### -Introduction

#### Intro

- electronics structure of solids
- Berry phase, Berry connection, Berry curvature
- degenerated system, Dirac equation
- Berry curvature and transport
- relation between Berry curvature and Kubo formula
- time reversal operator, Kramers degeneracy
- spin-orbit coupling of electron
- spin-orbit coupling of light

Electron in periodic potential

# Electron in periodic potential

Electron in periodic potential

Free electron

#### Free electron

Free electron's state is described by Schrödinger equation:

$$H\psi = E\psi$$
$$\left(-\frac{\hbar^2}{2m}\nabla^2 + U(\mathbf{r})\right)\psi = E\psi, \text{ where } U(\mathbf{r}) = 0$$

In case of free electron, the solution of the electron's wavefunction is plane wave,  $\psi = \exp(i\mathbf{k} \cdot \mathbf{r})$ , and the corresponding electron's energy is:

$$E=\frac{\hbar^2|\mathbf{k}|^2}{2m}.$$

Electron in periodic potential

└─ Free electron

# Free electron energy: $E = \frac{\hbar^2 |\mathbf{k}|^2}{2m}$

- Relation between wavelength  $\lambda$  and wavevector **k** is  $\lambda = \frac{2\pi}{|\mathbf{k}|}$ .
- When compared with the classical relation E = p<sup>2</sup>/2m, we obtain relation between the linear momentum p and the wavevector k (de Broglie relation):

$$\mathbf{p} = \hbar \mathbf{k}.$$

#### Comparison with photon:

- photon with energy 1 eV has wavelength 1240 nm  $E = \hbar \omega = \hbar kc = pc$
- free electron with energy 1 eV has wavelength 1.23 nm. Newton (classical):  $E = \frac{1}{2}mv^2 = \frac{p^2}{2m}$ Einstein (relativity):  $E = pc \approx mc^2 + \frac{p^2}{2m}$

Electron in periodic potential

Free electron

#### Dispersion relation of free the electron

Newton (classical):

$$E = \frac{1}{2}mv^2 = \frac{p^2}{2m}$$

Einstein (special relativity):

$$E = \sqrt{p^2 c^2 + (mc^2)^2}$$
$$\approx mc^2 + \frac{p^2}{2m}$$

Dirac equation:

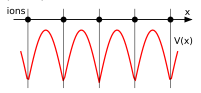
E = cp

Electron in periodic potential

Bloch theorem

#### Electron in periodic potential - Bloch theorem

• Let us assume electron's potential to be periodical:  $U_0(\mathbf{r}) = U_0(\mathbf{r} + \mathbf{T})$ , where **T** is the lattice vector.



- We assume one-electron approximation, i.e. interaction with other electrons (exchange, correlation, Coulomb force) are included as the additional effective potential acting on electron,  $U_{\text{eff}}(\mathbf{r})$ , with the same periodicity as  $U_0(\mathbf{r})$ :  $U(\mathbf{r}) = U_0(\mathbf{r}) + U_{\text{eff}}(\mathbf{r})$ .
- Then, the electron's state is described by a wavefunction ψ fulfilling the Schrödinger equation:

$$\Pi \psi \equiv E \psi$$
$$\left(-\frac{\hbar^2}{2m} \nabla^2 + U(\mathbf{r})\right) \psi = E \psi.$$

Electron in periodic potential

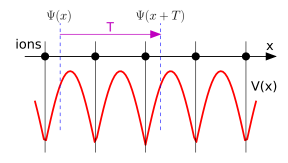
Bloch theorem

#### Bloch theorem: proof I

• All observables must have the same periodicity as the lattice. Hence, the electron probability  $|\psi|^2$  must fulfill:  $|\psi(\mathbf{r})|^2 = |\psi(\mathbf{r} + \mathbf{T})|^2$ . Therefore, we can express  $\psi(\mathbf{r})$  as:

$$\psi(\mathbf{r} + \mathbf{T}) = e^{i\theta(\mathbf{T})}\psi(\mathbf{r}),$$

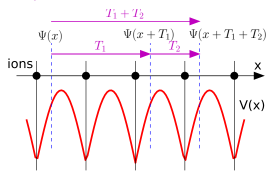
where  $\theta(\mathbf{T})$  is (at this stage arbitrary) phase, as  $|e^{i\theta(\mathbf{T})}|^2 = 1$ 



Electron in periodic potential

Bloch theorem

#### Bloch theorem: proof I



For two translations T<sub>1</sub> and T<sub>2</sub> we have

$$\psi(\mathbf{r} + \mathbf{T}_1 + \mathbf{T}_2) = e^{i\theta(\mathbf{T}_1 + \mathbf{T}_2)}\psi(\mathbf{r}) = e^{i\theta(\mathbf{T}_1)}e^{i\theta(\mathbf{T}_2)}\psi(\mathbf{r})$$

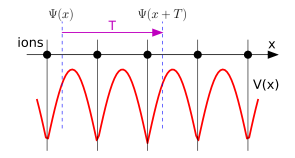
Hence,  $\theta(\mathbf{T}_1 + \mathbf{T}_2) = \theta(\mathbf{T}_1) + \theta(\mathbf{T}_2)$ . Hence, the phase  $\theta(\mathbf{T})$  has form  $\theta(\mathbf{T}) = \mathbf{k} \cdot \mathbf{T}$ , as the only this function fulfills the requirement.

Electron in periodic potential

Bloch theorem

#### Bloch theorem: proof I

■ In final, the electron wavefunction of the periodical potential writes:  $\psi(\mathbf{r} + \mathbf{T}) = e^{i\mathbf{k}\cdot\mathbf{T}}\psi(\mathbf{r})$ . In another words, the translation by a lattice vector  $\mathbf{T}$  is equivalent to multiplaying the wavefunction  $\psi(\mathbf{r})$  by a phase factor  $e^{i\mathbf{k}\cdot\mathbf{T}}$ .



Electron in periodic potential

Bloch theorem

## Bloch theorem: proof II

• Let us assume, that the solution of  $\psi(\mathbf{r})$  is in form:

$$\psi_{\mathbf{k}}(\mathbf{r}) = e^{i\mathbf{k}\cdot\mathbf{r}}u_{\mathbf{k}}(\mathbf{r}).$$

Then, we show that  $u_{\mathbf{k}}(\mathbf{r})$  has the same periodicity as the lattice.

■ In point **r** + **T** the wavefunction has value:

$$\psi(\mathbf{r}+\mathbf{T})=e^{i\mathbf{k}\cdot(\mathbf{r}+\mathbf{T})}u_{\mathbf{k}}(\mathbf{r}+\mathbf{T}),$$

whereas the previous proof provides:

$$\psi(\mathbf{r}+\mathbf{T})=e^{i\mathbf{k}\cdot\mathbf{T}}\psi(\mathbf{r})=e^{i\mathbf{k}\cdot\mathbf{T}}e^{i\mathbf{k}\cdot\mathbf{r}}u_{\mathbf{k}}(\mathbf{r}).$$

By comparing both equations, we see:

$$u_{\mathbf{k}}(\mathbf{r}) = u_{\mathbf{k}}(\mathbf{r} + \mathbf{T})$$

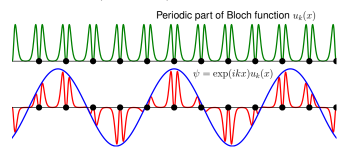
i.e.  $u_{\mathbf{k}}(\mathbf{r})$  has the same periodicity as the lattice.

- Electron in periodic potential
  - Bloch theorem

Wavefunction of the periodic potential is (Bloch theorem):

 $\psi_{\mathbf{k}}(\mathbf{r}) = \exp(i\mathbf{k}\cdot\mathbf{r})u_{\mathbf{k}}(\mathbf{r}),$ 

- $u_{\mathbf{k}}(\mathbf{r})$  has equal periodicity as the lattice,  $u_{\mathbf{k}}(\mathbf{r}) = u_{\mathbf{k}}(\mathbf{r} + \mathbf{T})$ .
- exp(*i***k** · **r**) corresponds to free-electron wave (free-electron propagation). It implies that the electron propagates through the crystal like a free (pseudo-free) particle.

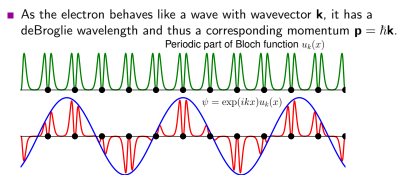


Electron in periodic potential

Bloch theorem

$$\psi_{\mathbf{k}}(\mathbf{r}) = \exp(i\mathbf{k}\cdot\mathbf{r})u_{\mathbf{k}}(\mathbf{r}),$$

• Function  $u_{\mathbf{k}}(\mathbf{r})$  modulates this free-electron-like wave so that the amplitude oscillates periodically form one cell to the next. It does not affect the basic character of  $\psi$ , which is that of a traveling wave. In reality the electron is not free, (it interacts with the lattice, electrons etc), but its propagation has features of a free electron propagation.



Electron in periodic potential

-Fourier transform

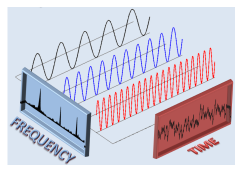
#### 1D Fourier transform

Fourier transform:

$$F(k) = \frac{1}{\sqrt{2\pi}} \int_{-\infty}^{\infty} f(x) \exp(-ikx) dx$$

Inverse Fourier transform:

$$f(x) = rac{1}{\sqrt{2\pi}} \int_{-\infty}^{\infty} F(k) \exp(ikx) \mathrm{d}k$$



Build function f(x) as a sum of harmonic functions  $\exp(ikx)$  having amplitudes F(k).

Electron in periodic potential

-Fourier transform

#### 1D Fourier transform

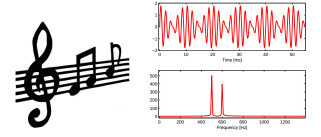
Fourier transform:

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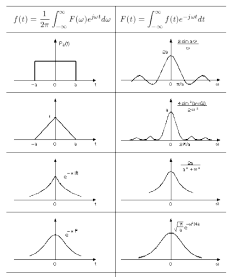
Example: sound in real (time) and reciprocal (frequency) space:



Electron in periodic potential

Fourier transform

Examples of Fourier transforms



http://www.etc.tuiasi.ro/cin/Downloads/Fourier/Fourier.html

Basic properties of 1D Fourier transformations

transformations	
f(t)	$F(\omega)=F.T.(f(t))$
f(at)	$\frac{1}{ a }F\left(\frac{\omega}{a}\right)$
$f^{\star}(t)$	${\sf F}^{\star}(-\omega)$ (conjugation)
$f(t-t_0)$	$F(\omega) e^{-it_0\omega}$
$f(t)e^{i\omega_0 t}$	${\sf F}(\omega-\omega_0)$
$f(t)\cos(\omega t)$	$rac{1}{2}\left[ F(\omega+\omega_0)+F(\omega-\omega_0) ight]$
$\frac{\mathrm{d}^n f(t)}{\mathrm{d} t^n}$	$(i\omega)^n F(\omega)$
$(-it)^n f(t)$	$\frac{\mathrm{d}^n F(\omega)}{\mathrm{d} \omega^n}$

Electron in periodic potential

-Fourier transform

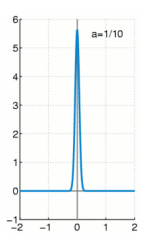
# Dirac $\delta$ -function

#### Definition of $\delta$ -function

$$\delta(x) = \begin{cases} +\infty, & x = 0 \\ 0, & x \neq 0 \end{cases}$$
  
so that 
$$\int_{-\infty}^{\infty} \delta(x) \, dx = 1$$

Properties:

$$\int_{-\infty}^{\infty} f(x) \,\delta(x-a) \mathrm{d}x = f(a)$$
$$\delta(ax) = \frac{\delta(x)}{|a|}$$
$$\int_{-\infty}^{\infty} e^{ikx} \mathrm{d}k = 2\pi\delta(x)$$



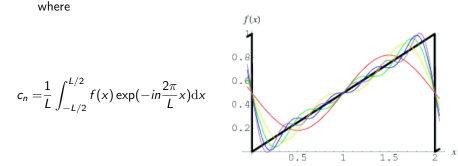
Electron in periodic potential

-Fourier transform

#### Fourier series

Let us have *periodic* function f(x) with period T, f(x) = f(x+L). Then,

$$f(x) = \sum_{n=-\infty}^{\infty} c_n \exp(in\frac{2\pi x}{L}).$$



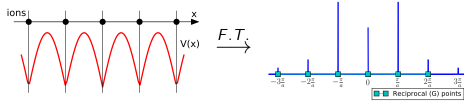
Electron in periodic potential

└─ Fourier transform

# Fourier series and Fourier transformation Periodic function $f(x) = \sum_{n=-\infty}^{\infty} c_n \exp(in\frac{2\pi x}{L}).$

Fourier transform of periodic function f(x) is:

$$F(k) = \frac{1}{\sqrt{2\pi}} \int_{-\infty}^{\infty} f(x) \exp(-ikx) dx = \frac{1}{\sqrt{2\pi}} \sum_{n=-\infty}^{\infty} c_n \int_{-\infty}^{\infty} \exp\left[ix(-n\frac{2\pi}{L}+k)\right]$$
$$= \sum_{n=-\infty}^{\infty} \sqrt{2\pi} c_n \delta(k-n\frac{2\pi}{L}).$$



Periodic function becomes a train of  $\delta$ -functions with equidistant spacing.

Electron in periodic potential

Reciprocal space

## Reciprocal space (k-space)

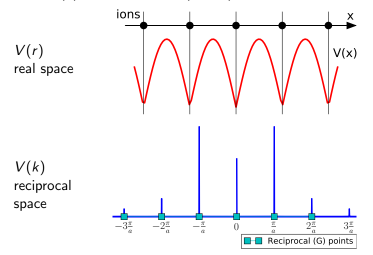
- In the real space, the quantities are expressed as function of positions, e.g. V(r), ψ<sub>k</sub>(r).
- In the reciprocal space, the quantities are expressed as function of wave-vector k or momentum p = ħk.
- Conversion between real and reciprocal space is Fourier transform, e.g. V(k) = F.T.(V(r)), and V(r) = inv.F.T.(V(k)).

Electron in periodic potential

Reciprocal space

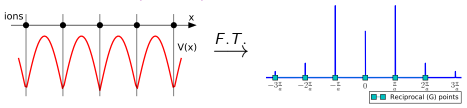
# Reciprocal space (k-space): potential V(x)

Potential  $V(\mathbf{x})$  of the crystal is periodical. Hence, in the reciprocal space,  $V(\mathbf{k})$  consists of train (lattice) of  $\delta$ -functions.



- Electron in periodic potential
  - Reciprocal space

# Reciprocal space (k-space): potential V(x)



- the position of δ-functions are called reciprocal points or G-points, forming reciprocal lattice
- the distance between G-points is given only by periodicity of the function
- $\Rightarrow$  any periodic function with equal periodicity is described by  $\delta$  functions at identical G-points
- $\Rightarrow$  any periodic property of crystal (potential, electron density) is expressed at identical *G*-points
  - $\blacksquare$  the shape of the function is given by amplitudes of the  $\delta\text{-functions}$

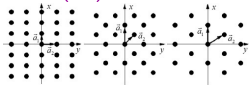
Electron in periodic potential

Reciprocal space

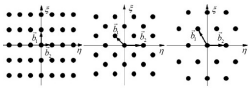
# Reciprocal space of 2D periodic function $f(\mathbf{r})$

- 2-dimensional (2D) periodic function  $f(\mathbf{r}) = f(\mathbf{r} + \mathbf{R}_{\mathbf{u}})$ ,  $\mathbf{u} \in \mathbb{Z}$ , where translation vectors  $\mathbf{R} = u_1 \mathbf{a}_1 + u_2 \mathbf{a}_2$ form *Bravais lattice*
- $\rightarrow$  **a**<sub>1</sub>, **a**<sub>2</sub> called primitive vectors
  - Fourier transform of f(r) consists of 2D lattice of δ-functions
- ightarrow each reciprocal point denotes position of  $\delta$ -function

#### Bravais (real) lattice:



#### **Reciprocal lattice:**



Lin-Wei et al, JOSAA 25, 203 (2008)

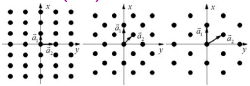
Electron in periodic potential

Reciprocal space

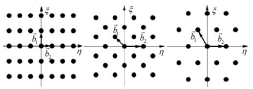
# Reciprocal space of 2D periodic function $f(\mathbf{r})$

- position of reciprocal points  $\mathbf{G}_{\mathbf{m}}$  are determined by relation  $\mathbf{G}_{\mathbf{m}} \cdot \mathbf{R}_{\mathbf{u}} = 2\pi n$ ,  $n \in \mathbb{Z}$ , originating from condition  $\exp(i\mathbf{G}_{\mathbf{m}} \cdot \mathbf{R}_{\mathbf{u}}) = 1$
- primitive vector of reciprocal lattice are b<sub>1</sub>, b<sub>2</sub>, providing G<sub>m</sub> = m<sub>1</sub>b<sub>1</sub> + m<sub>2</sub>b<sub>2</sub>
- similar for 3D reciprocal lattice

#### Bravais (real) lattice:



#### **Reciprocal lattice:**



Lin-Wei et al, JOSAA 25, 203 (2008)

Electron in periodic potential

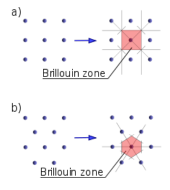
-Brillouin zone

# (First) Brillouin zone

#### **Definitions:**

- Def1: First Brillouin zone is a smallest possible cell in reciprocal space, which by translation can form whole reciprocal space.
- Def2: Any point of the reciprocal space **k** can be reached from the first Brillouin zone **k**<sub>inside\_1st\_Brillouin\_zone</sub> added to a translation vector of the reciprocal lattice **G**<sub>mn</sub>:

 $\mathbf{k} = \mathbf{k}_{\mathrm{inside\_1st\_Brillouin\_zone}} + \mathbf{G}_{\textit{mn}}.$ 



Electron in periodic potential

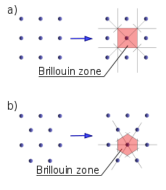
—Brillouin zone

#### Properties of Brillouin zone:

Volume of the (first) Brillouin zone
 V<sub>k</sub> is inversely proportional to
 volume of unit cell of the real
 (direct) lattice V<sub>r</sub>:

$$V_{\mathbf{k}} = \frac{(2\pi)^N}{V_{\mathbf{r}}}$$

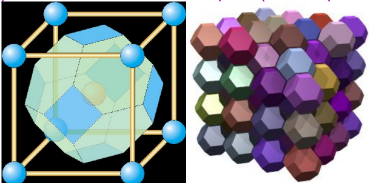
N - dimension of lattice (2D or 3D)

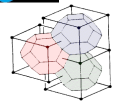


Electron in periodic potential

Brillouin zone

# (First) Brillouin zone of fcc real space (bcc reciprocal)

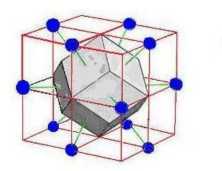




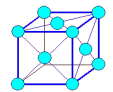
Electron in periodic potential

Brillouin zone

# (First) Brillouin zone of bcc real space (fcc reciprocal)







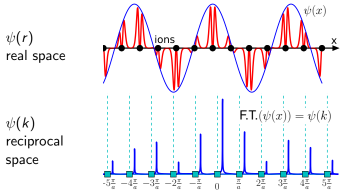
Electron in periodic potential

Brillouin zone

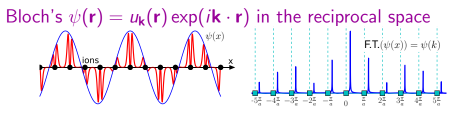
# Bloch's $\psi(\mathbf{r}) = u_{\mathbf{k}}(\mathbf{r}) \exp(i\mathbf{k} \cdot \mathbf{r})$ in the reciprocal space

- $u_{\mathbf{k}}(\mathbf{r})$  periodical part of Bloch's function
- $\exp(i\mathbf{k} \cdot \mathbf{r})$  plane wave.

As  $\psi(\mathbf{r})$  is not periodic in  $\mathbf{r}$ ,  $\psi(\mathbf{k})$  is expressed by train of  $\delta$ -functions shifted by  $\mathbf{k}$  from the reciprocal points as F.T. $(f(x) \exp(ik_0 x)) = F(k - k_0)$ .



- Electron in periodic potential
  - Brillouin zone



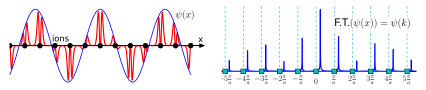
- as ψ(k) is described by a train of δ-functions in the reciprocal space, it is enough to use k from the 1st Brillouin zone
- $\Rightarrow$  reduction to the first Brillouin zone
  - wavevector **k** is a quantum number of the wavefunction:
- $\Rightarrow$  **k** in crystal must be described by integer and hence must be *quantized number*
- $\rightarrow\,$  Pauli principle: No two electrons in an atom/crystal can have identical all quantum numbers.
- $\Rightarrow\,$  Inside whole crystal, there can be only two electrons at each  ${\bf k}\,$  state (spin-up and spin-down)
  - So, how many k-states is in the crystal?

- Electron in periodic potential
  - Brillouin zone

#### Note: conservation of momentum $\mathbf{p} = \hbar \mathbf{k}$ in a crystal

- Energy conservation *E*: due to time-invariant space.
- Momentum conservation **p**: due to translation-invariant space.
- Angular momentum conservation L: due to rotation-invariant

space.

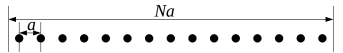


- in crystal, there is no invariance of space in both translation and rotation.
- $\Rightarrow \mbox{ momentum } {\bf p} = \hbar {\bf k} \mbox{ of the electron in crystal is not uniquely} \\ \mbox{ defined, and it can behave as having any value} \\ {\bf p} = \hbar ({\bf k} + {\bf G}_{m}), \mbox{ G}_{m} \mbox{ being any reciprocal vector in the lattice.}$

Electron in periodic potential

└─ Number of **k**-states in the first Brillouin zone

#### How many k-states is in the crystal?



We need to know, how many **k**-states are available inside whole (macroscopic) crystal having N unit cells. Hence, boundary conditions of  $\psi(\mathbf{r})$  on crystal's interface must be described.

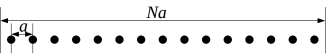
#### Number of k states in the Brillouin zone:

There is exactly as many allowed **k**-vector states in the Brillouin zone as there is number of the unit cells in the crystal.

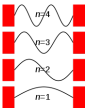
Electron in periodic potential

-Number of **k**-states in the first Brillouin zone

# Boundary conditions of $\psi(\mathbf{r})$ on interface of the crystal:



Describe standing waves inside the 1D box, 1st opt: having boundary conditions  $\psi(x < 0) = \psi(x > Na) = 0$ , where *a* is inter-atomic distance and *N* is a number of atoms in *x*direction. However, difficult to handle analytically.



2nd opt: Cyclic (Born – von Kármán) boundary conditions  $\psi(x) = \psi(x + Na)$ . Not a real (physical) boundary conditions, but accounts correct number of allowed *k*-states.

Electron in periodic potential

└─Number of **k**-states in the first Brillouin zone

#### Cyclic boundary conditions:

Let us demonstrate cyclic boundary conditions in x-direction:

$$\psi(\mathbf{x}) = \psi(\mathbf{x} + N\mathbf{a})$$

For Bloch electron in *x*-direction:

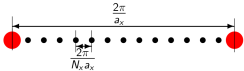
$$\psi(x) = e^{ikx}u_k(x)$$
  
 $\psi(x + Na) = e^{ikNa}e^{ikx}u_k(x) = e^{ikNa}\psi(x)$ 

And hence:  $e^{ikNa} = 1$ , providing  $kNa = 2\pi m$ ,  $m \in \mathbb{Z}$ . Hence, allowed values of k are:

$$k_m = \frac{2\pi}{a} \frac{m}{N}$$

- Electron in periodic potential
  - └─ Number of **k**-states in the first Brillouin zone

Cyclic boundary conditions:



Due to the boundary conditions, not all values of k-vector are allowed. Only allowed values of k are  $(m \in \mathbb{Z})$ :

$$k_m = \frac{2\pi}{a} \frac{m}{N} = G \frac{m}{N},$$

i.e. the reciprocal translation vector  $G = 2\pi/a$  is divided to N parts.

To generalize to 3D and general unit cell (general Brillouin zone):

#### Number of k states in the Brillouin zone:

There is exactly as many allowed **k**-vector states in the Brillouin zone as there is number of the unit cells in the crystal.

Electron in periodic potential

Free electron states

#### Electron in crystal

- Assume crystal has N unit cells in 2D crystal, and Z electrons per unit cell. Hence, whole crystal contains NZ electrons.
- k-vector is a quantum number of electron. Hence, each
   k-vector + spin provides unique quantum numbers to each electron in whole crystal.
- Pauli principle states that on each quantum state (i.e. for each value of k-vector), the k-state can contain only two electrons (with spin-up and spin-down).
- The electrons are filled to the electronic structure starting from smaller energies, i.e. in our case from smaller k-vectors.

#### Free electron approximation:

Let us assume an electron inside periodic lattice. However, we assume wavefunction  $\psi = e^{i\mathbf{k}\cdot\mathbf{r}}$  (which is solution for  $U(\mathbf{r}) = 0$ ), with energy  $E = \frac{\hbar^2 k^2}{2m}$  (free electron approximation).

Electron in periodic potential

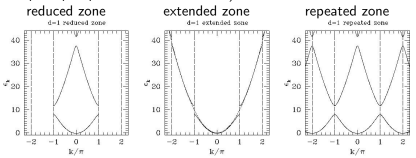
└─ Free electron states

## Reduction to the Brillouin zone

In case of electron with  $\mathbf{k}$ -vector outside Brillouin zone, the  $\mathbf{k}$ -vector can be shifted back to the first Brillouin zone,

$$\mathbf{k} = \mathbf{k}_{\mathrm{inside\_1st\_Brillouin\_zone}} + \mathbf{G}_{mn}$$

In free electron model, electron energy is  $E = \frac{\hbar^2 k^2}{2m}$ . Hence shift of **k** to the Brillouin zone provides electron bands (with principal quantum number n > 0).



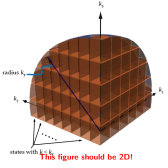
Electron in periodic potential

#### └─ Free electron states

## Free electron in 2D cubic crystal

- 2D volume of the Brillouin zone  $V_k = 4\pi^2/V_r$  contains 2N electrons' k-states (times 2 because of two spins).
- The electron energy depends solely on |k|, the area of constant energy is a sphere in the reciprocal space.
- When all available electrons fill the k-spaces, a circle is formed up to the highest occupied energy (*Fermi level*), with k-vector k<sub>F</sub>.
- This 2D circle occupies reciprocal space  $V_k = \pi k_F^2$ , providing  $\pi k_F^2/(NZ) = (4\pi^2/V_r)/(2N)$ . Hence, the Fermi level of this circle has radius:

$$k_F = \sqrt{\frac{2\pi}{V_r}Z} \equiv \sqrt{\frac{2\pi}{V}NZ}.$$





Electron in periodic potential

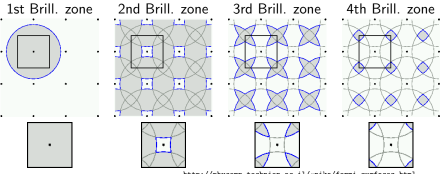
└─ Free electron states

#### Free electron in 2D cubic crystal

*Z* = 4

What states are occupied and what are filled?

square length:  $k_a = 2\pi/a = 6.28/a$  $k_F = \sqrt{2\pi Z/(a^2)} = 5.01/a$  $k_F/(k_a/2) = 1.60 > \sqrt{2}$ 



http://phycomp.technion.ac.il/~nika/fermi\_surfaces.html

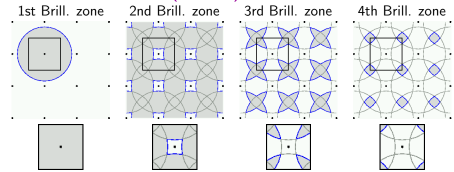
Electron in periodic potential

└─ Free electron states

## Fermi surface

As electrons fill the reciprocal space up to  $E_F$ , they create a boundary in the reciprocal space between filled and empty k-spaces. This boundary is called Fermi surface.

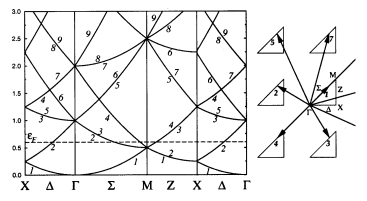
Fermi surface in 2D, cubic (blue line):

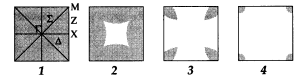


Electron in periodic potential

Free electron states

## Band structure of 2D electrons in cubic crystal





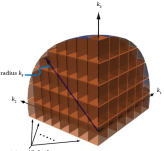
Electron in periodic potential

└─ Free electron states

## Free electron in 3D

- In the volume of the Brillouin zone  $V_k = (2\pi)^3/V_r$  there is 2N electrons' k-states.
- The electron energy depends solely on |k|, the area of constant energy is a sphere in the reciprocal space.
- When all NZ available electrons fill the k-spaces, they form a sphere called Fermi sphere (terminated by Fermi surface), with maximal k-vector k<sub>F</sub>.
- This sphere occupies reciprocal space  $V_k = \frac{4}{3}\pi k_F^3$ , providing  $\frac{4}{3}\pi k_F^3/(NZ) = (8\pi^3/V_r)/(2N)$ . Hence, the Fermi sphere has radius:

$$k_F = \sqrt[3]{\frac{3\pi^2}{V_r}Z} \equiv \sqrt[3]{\frac{3\pi^2}{V}NZ}.$$



states with  $k \leq k_p$ 

 $E_F = \frac{\hbar^2 k_F^2}{2m}$ 

Electron in periodic potential

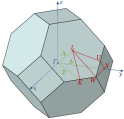
Free electron states

	GROUP		PE	ERI	OD	OIC	TA	BL	ΕC	)F	TH	EE				-		18 VIII
1	1 1.0079			RELATIVE ATOMIC MASS (1)				Metal Semimetal 🔝 Nonmetal				$\sum$			span.ar/per	ounien		2 4.002
	HYDROGEN	GR			OUP IUPAC GROUP CAS			Alkali metal     Alkaline earth metal			16 Chalcogens element		13 IIIA 14 IVA 15 VA 16 VIA-17 VIIA HELK					
_	3 6.941	4 9.0122	ATOMIC N	UMBER 5	10.811		-	insition metal		18 Noble			5 10.811	6 12.011	7 14.007	8 15.999	9 18.998	10 20.1
2	Li	Be		SYMBOL -	B			Lanthanide Actinide		OARD STATE	(25 °C; 101	(Pa)	B	С	N	0	F	Ne
	LITHIUM 11 22.990	BERYLLIUM 12 24.305		/ •	BORON			, 1011100		- yas - liquid	Fe - solid	tic	BORON 13 26.982	CARBON 14 28.086	N/TROGEN 15 30.974	0XYGEN 16 32.065	FLUORINE 17 35.453	NEOR 18 39.
3				ELE	MENT NAME					1	/	_	Al	Si	P	S	~	
-	Na	Mg	3 IIIB	4 IVB	5 /VB	6 VIB	7 VIIB	8	VIIIB -	10	11 18	12 IIB	ALUMINIUM	SLICON	PHOSPHORUS	SULPHUR	CI	ARGO
1			21 44.956	22 47.867	. /				27 58.933				31 69.723	32 72.64	33 74.922	34 78.96	35 79.904	36 83
4	K	Ca	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	Ge	As	Se	Br	Kı
	POTASSIUM	CALCIUM	SCANDIUM	TITANIUM	VANADIUM	CHROMIUM	MANGANESE	IRON	COBALT	NICKEL	COPPER	ZINC	GALLIUM	GERMANIUM	ARSENIC	SELENIUM	BROMINE	KRYPT
		38 87.62	39 88.906	40 91.224	41 92.906	42 95.94	43 (98)	44 101.07	45 102.91	46 106.42	47 107.87	48 112.41	49 114.82	50 118.71	51 121.76	52 127.60	53 126.90	54 131
5	Rb	Sr	Y	Zr	Nb	Mo	Te	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Te	I	X
	RUBIDIUM	STRONTIUM	YTTRIUM	ZIRCONIUM	NIOBIUM		TECHNETIUM		RHODIUM	PALLADIUM	SILVER	CADMUM	INDIUM	TIN	ANTIMONY	TELLURIUM	IODINE	XENO
6		56 137.33	57-71 La-Lu	72 178.49	73 180.95	74 183.84	_	76 190.23	-	78 195.08				82 207.2	83 208.98		85 (210)	86 (2
^	CS	Ba	Lanthanide	HAFNIUM	Ta	W	Re	OSMILM	Ir	Pt	GOLD	Hg	TI	Pb	Bi	POLONIUM	At	RI
		88 (226)	89-103	104 (261)		106 (266)	107 (264)	108 (277)	109 (268)		111 (272)	112 (285)		114 (289)		1 0201101		10.00
7	Fr	Ra	Ac-Lr	IRf	IDb	Sg	IBh	IHIs	MItt	Umm	Winn	Մախ		Uua				
	FRANCIUM	RADIUM	Actinide	RUTHERFORDUV	DUBNIUM	SEABORGIUM	BOHRIUM	HASSIUM	MEITNERIUM	UNUNNILIUM	UNUNUNIUM	UNUNBIUM		UNUNQUADUN				
				LANTHAN	DE												198-2003 EriG.	
Pure Appl. Chem., 73, No. 4, 667-683 (2001)				57 138.91	58 140.12	59 140.91	60 144.24	61 (145)	62 150.36	63 151.96	64 157.25	65 158.93	66 162.50	67 164.93				
Relative atomic mass is shown with five significant figures. For elements have no stable nuclides, the value enclosed in brackets indicates the mass sumther of the locoest-lived			e no stable brackets	La	Ce	Pr	Nd	IPm	Sm	Eu	Gd	Tb	Dy	Ho	Er	Tm	Yb	Lu
isotope of the element / LANTHANUM However three such elements (Th, Pa, and U) ACTINIDE											EUROPIUM GADOLINIUM TERBIUM DYSPROSIUM HOLMIUM ERBIUM THULIUM YTTERBIUM					LUTETR		
do have a characteristic terrestrial isotopic composition, and for these an atomic weight is tabulated.					90 232.04	91 231.04	92 238.03	93 (237)	94 (244)	95 (243)	96 (247)	97 (247)	98 (251)	99 (252)	100 (257)	101 (258)	102 (259)	103 (2
				Ac	Th	Pa	U	Np	Pu	Am	Cm	Bk	Cf	Es	Fm	Md	No	ILI

Electron in periodic potential

Free electron states

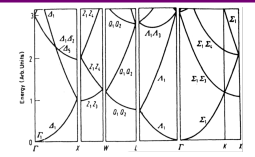
#### Free electron in fcc:

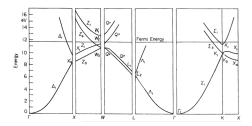


#### Aluminium (fcc):

- 1s<sup>2</sup> 2s<sup>2</sup> 2p<sup>6</sup> 3s<sup>2</sup> 3p<sup>1</sup>
- three unbounded electrons







- Electron in periodic potential
  - Free electron states

#### Aluminium (fcc):

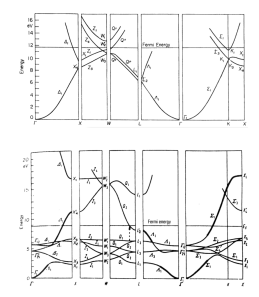
- $\bullet 1s^2 2s^2 2p^6 3s^2 3p^1$
- three unbounded electrons



## Cuprum (fcc):

- [Ar] 3d<sup>10</sup> 4s<sup>1</sup>
- eleven unbounded electrons





Electron in periodic potential

Density of states

## Density of states (DOS):

Density of states (DOS): number of electron's states per energy interval:

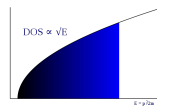
$$g=rac{\mathrm{d}(NZ)}{\mathrm{d}E}.$$

For free elecctron,  $E_F = \frac{\hbar^2 k_F^2}{2m}$ , where  $k_F = \sqrt[3]{\frac{3\pi^2}{V}NZ}$ . Hence:

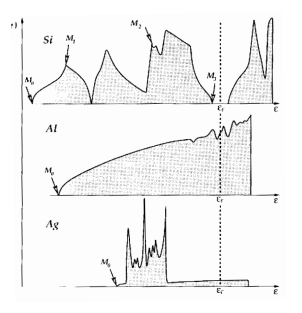
$$NZ = \frac{V}{3\pi^2} \left(\frac{2mE}{\hbar^2}\right)^{3/2}$$

Therefore, Density of states g(E) of free electron is:

$$g(E) = \frac{\mathrm{d}(NZ)}{\mathrm{d}E} = \frac{V}{2\pi^2} \left(\frac{2m}{\hbar^2}\right)^{3/2} \sqrt{E}$$



- Electron in periodic potential
  - Density of states



Si: complicated DOS, contains gap at Fermi level.

Al: nearly free electrons DOS  $\sim \sqrt{E}$ .

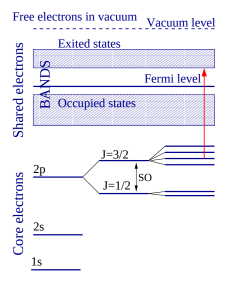
Ag: *d* electrons are burried and the only electrons on the Fermi level are *s*-electrons. Electron in periodic potential

Density of states

## Complete energy levels in solids:

#### Core states:

- Electrons in the spherical potential of the nucleus (bounded, not shared, electrons).
- Approximately, they correspond to electrons in filled electronic shells.
- They provide sharp energies.
- Quantum numbers without spin-orbit coupling: n, l, m = l<sub>z</sub>, s, s<sub>z</sub>
- Quantum numbers with spin-orbit coupling: n, l, s, j, jz



Electron in periodic potential

└─ Density of states

## Complete energy levels in solids:

#### **Occupied states:**

Electrons shared through the crystal. They approximately correspond to valence electrons of the atoms.

They form energy bands.

#### Fermi level:

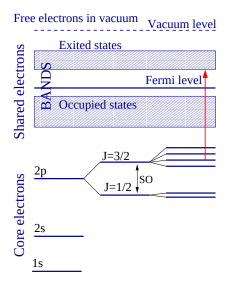
Energy level separating occupied and unoccupied states.

#### Excited states:

Empty energy levels above Fermi level. Electrons can be excited to those states and then they relax back.

#### Vacuum level:

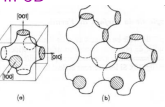
Overcoming this barrier, electrons do not feel periodical potential of the crystal and become free.

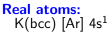


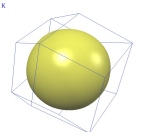
- Electron in periodic potential
  - Fermi surface

## Fermi surface in 3D



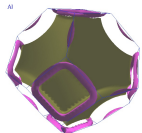








Al (fcc) [Ne] 3s<sup>2</sup> 3p<sup>1</sup>

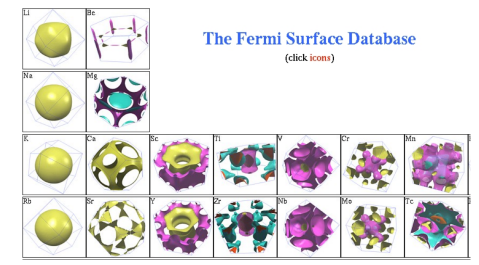


http://www.phys.ufl.edu/fermisurface/

Electron in periodic potential

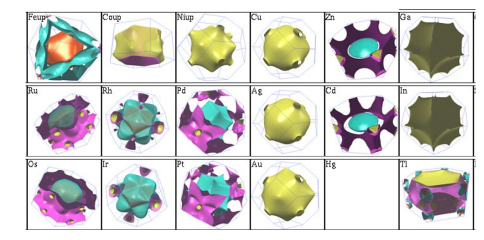
Fermi surface

## Example of some elements' Fermi surfaces.



- Electron in periodic potential
  - Fermi surface

## Example of some elements' Fermi surfaces.



- Electron in periodic potential
  - └─ Fermi surface

## Comments on Fermi surface

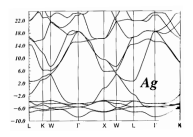
- Fermi surface is formed by free (unbounded or easily excited) electrons. Such electrons are provides e.g. electrical or thermal conductivity.
- Hence, the metal can be defined as 'material having Fermi surface'.
- Semiconductors and insulators do not have Fermi surfaces, because they do not have free electrons on Fermi level. In this case, one Brillouin zone is completely full and next Brillouin zone is completely empty.

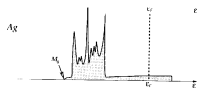
Electron in periodic potential

└─ Fermi surface

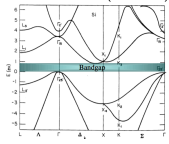
## Basic classification of materials (according to DOS): metal semiconductor/isolator

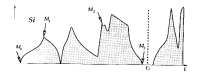
no gap at Fermi level Fermi surface





gap at Fermi level no Fermi surface (filled BZ)





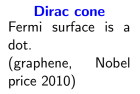
Electron in periodic potential

└─ Fermi surface

## Basic classification of materials (according to DOS):

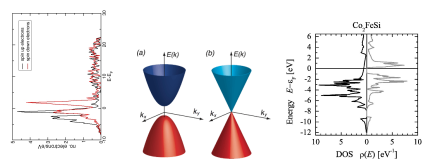
#### ferromagnet

different DOS for up and down electrons Fe(bcc): [Ar]4s<sup>2</sup>3d<sup>6</sup>



#### half-metal

gap only for spin down (or only for spin up)



Berry phase, Berry curvature

# Berry phase, Berry connection, Berry curvature

Berry phase and Barry curvature

Phase in parameters space, Berry connection

## Phase in parameters space : I

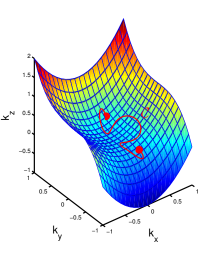
Consider a system described by Hamiltonian depending on time through sets of parametres

$$\mathcal{R}(t) = (R_1(t), R_2(t), \ldots)$$
, i.e.

 $H=H(\mathcal{R}(t))$ 

- adiabatic evolution of the system as R(t) moves slowly along a path C in the parameter space
- parametr space is for example
  - $\rightarrow$  *k*-space
  - $\rightarrow$  direction of magnetic fied *B*, etc.

Xiao et al, arxiv:0907.2021 (2009) Berry (1983)



Berry phase and Barry curvature

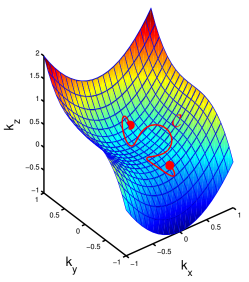
Phase in parameters space, Berry connection

## Phase in parameters space : I

Let us introduce instantaneous orthonormal basis for each  $H(\mathcal{R})$ 

$$H(\mathcal{R}) | n(\mathcal{R}) \rangle = \epsilon_n(\mathcal{R}) | n(\mathcal{R}) \rangle$$

- However, the phase factor of |n(R)⟩ is not determined.
- We require that phase of the basis functions |n(R)⟩ is smooth along path C in the parameter space.



Berry phase and Barry curvature

Phase in parameters space, Berry connection

## Derivation 1 of Berry connection:

For simplicity, we assume the path C is on isoenergy surface  $\epsilon(\mathcal{R})_n = \epsilon_n = \text{const}_n$ . Then, the wavefunction over the path C is

$$|\psi_n(t)
angle = e^{i\gamma_n(\mathcal{R}(t))}e^{-rac{i}{\hbar}\epsilon_n t} |n(\mathcal{R}(t))
angle$$

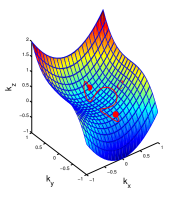
where phase consist of phase  $\gamma_n(\mathcal{R}(t))$  and dynamic phase  $-i\epsilon_n t/\hbar$ .

#### Note:

Generally, when  $\epsilon_n(\mathcal{R})$  changes on the path  $\mathcal{C}$ , the wavefunction becomes

$$|\psi_n(t)\rangle = e^{i\gamma_n(\mathcal{R}(t))}e^{-\frac{i}{\hbar}\int_0^t \mathrm{d}t'\epsilon_n(\mathcal{R}(t'))}|n(\mathcal{R}(t))\rangle$$

where phase consist of phase  $\gamma_n(\mathcal{R}(t))$  and dynamical phase factor.



Berry phase and Barry curvature

Phase in parameters space, Berry connection

## Derivation 1 of Berry connection:

Inserting this wavefunction  $|\psi_n(t)\rangle = e^{i\gamma_n(\mathcal{R}(t))}e^{-\frac{i}{\hbar}\epsilon_n t} |n(\mathcal{R}(t))\rangle$  to the Schrodinger equation

$$i\hbarrac{\partial}{\partial t}\left|\psi_{n}(t)
ight
angle=\mathcal{H}(\mathcal{R}(t))\left|\psi_{n}(t)
ight
angle$$

we got

$$i \frac{\partial \gamma(\mathcal{R})}{\partial \mathcal{R}} |n(\mathcal{R})\rangle + \frac{\partial}{\partial \mathcal{R}} |n(\mathcal{R})\rangle = 0$$

multiplying from left by  $\langle n(\mathcal{R})|$ 

$$\frac{\partial \gamma(\mathcal{R})}{\partial \mathcal{R}} = i \langle n(\mathcal{R}) | \frac{\partial}{\partial \mathcal{R}} n(\mathcal{R}) \rangle = i \mathcal{A}_n(\mathcal{R})$$

where  $\mathcal{A}_n(\mathcal{R})$  is Berry connection or Berry vector potential.  $\overline{\mathcal{A}_n(\mathcal{R}) = i \langle n(\mathcal{R}) | \frac{\partial}{\partial \mathcal{R}} | n(\mathcal{R}) \rangle}$ 

Berry phase and Barry curvature

Phase in parameters space, Berry connection

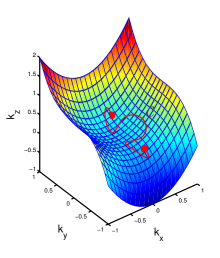
## Derivation 1 of Berry connection:

i.e.  $\mathcal{A}_n(\mathcal{R})$  expresses speed of change of phase on element of the path,

$$\mathcal{A}_n(\mathcal{R}) = \frac{\partial \gamma_n}{\partial \mathcal{R}} = i \langle n(\mathcal{R}) | \frac{\partial}{\partial \mathcal{R}} | n(\mathcal{R}) \rangle$$

Finally, the change of phase  $\gamma_n$  between starting and final point of path can be expressed as a path integral in the parameter space

$$\gamma_n = \int_{\mathcal{C}} \mathrm{d}\gamma_n = \int_{\mathcal{C}} \mathrm{d}\mathcal{R} \cdot \mathcal{A}_n(\mathcal{R})$$



Berry phase and Barry curvature

Phase in parameters space, Berry connection

## Derivation 2 of Berry connection:

Two close eigenwavefunctions  $|n(\mathcal{R})\rangle$  and  $|n(\mathcal{R} + \Delta \mathcal{R})\rangle$  are scalarly multiplied:

$$\langle n(\mathcal{R}) | n(\mathcal{R} + \Delta \mathcal{R}) \rangle \approx 1 + \Delta \mathcal{R} \langle n(\mathcal{R}) | \nabla_{\mathcal{R}} | n(\mathcal{R}) \rangle \\ \approx \exp[-i\Delta \mathcal{R} \cdot \mathcal{A}_n(\mathcal{R})] = \exp[-i\gamma_n] = \exp[-i[\gamma_n(\mathcal{R} + \Delta \mathcal{R}) - \gamma_n(\mathcal{R})]]$$

where Berry connection  $\mathcal{A}_n(\mathcal{R})$  is

$$\mathcal{A}_n(\mathcal{R}) = i \langle n(\mathcal{R}) | \nabla_{\mathcal{R}} | n(\mathcal{R}) \rangle$$

and Berry phase  $\gamma_n$  is expressed again as

$$\gamma_n = \int_{\mathcal{C}} \mathrm{d}\gamma_n = \int_{\mathcal{C}} \mathrm{d}\mathcal{R} \cdot \mathcal{A}_n(\mathcal{R})$$

Weng et al, arxiv:1508.02967 (2015)

Berry phase and Barry curvature

Phase in parameters space, Berry connection

## Berry connection: gauge transformation

Obviously, Berry connection  $\mathcal{A}_n(\mathcal{R})$  is gauge-dependent. We introduce gauge by applying

$$|n'(\mathcal{R})
angle 
ightarrow e^{i\zeta(\mathcal{R})}|n(\mathcal{R})
angle$$

where  $\zeta(\mathcal{R})$  is arbitrary smooth function. Then  $\mathcal{A}_n(\mathcal{R})$  transforms as

$$\mathcal{A}'_{n}(\mathcal{R}) \to \langle n'(\mathcal{R}) | \frac{\partial}{\partial \mathcal{R}} | n'(\mathcal{R}) \rangle = i \langle ne^{-i\zeta} | \frac{\partial}{\partial \mathcal{R}} | e^{i\zeta} n \rangle$$
$$= i \langle ne^{-i\zeta} | \left[ e^{i\zeta} | \frac{\partial}{\partial \mathcal{R}} n \rangle + i | e^{i\zeta} n \rangle \frac{\partial \zeta}{\partial \mathcal{R}} \right]$$
$$= \mathcal{A}_{n}(\mathcal{R}) - \frac{\partial}{\partial \mathcal{R}} \zeta(\mathcal{R})$$

Consequently, phase  $\gamma_n$  between starting and final points changes by  $\zeta(\mathcal{R}(0)) - \zeta(\mathcal{R}(T))$  where  $\mathcal{R}(0)$  and  $\mathcal{R}(T)$  are initial and final points of the path C.

Berry phase and Barry curvature

Phase in parameters space, Berry connection

# Berry connection: gauge transformation in Maxwell equations

$$\mathcal{A}'_n(\mathcal{R}) = \mathcal{A}_n(\mathcal{R}) - \frac{\partial}{\partial \mathcal{R}} \zeta(\mathcal{R})$$

Note analogy with gauge transformation in Maxwell equations:

$$\mathbf{B} = \nabla \times \mathbf{A}, \qquad \mathbf{E} = -\nabla \varphi - \frac{\partial}{\partial t} \mathbf{A}$$

where **A** and  $\varphi$  are magentic vector potential and electric potentials.

Gauge transformation of **A** and  $\varphi$  by arbitrary scalar function  $\Psi(\mathbf{r}, t)$  (called gauge function)

$$\mathbf{A'} 
ightarrow \mathbf{A} + 
abla \Psi, \qquad arphi' 
ightarrow arphi - rac{\partial \Psi}{\partial t}$$

Under this transformation, the **E** and **B** do not change. Note, we assume the gauge phase  $\zeta(\mathcal{R})$  does not depend on time. https://en.wikipedia.org/wiki/Gauge\_fixing

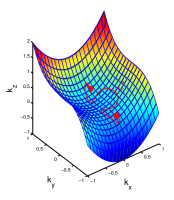
Berry phase and Barry curvature

Phase in parameters space, Berry connection

## Berry connection: gauge transformation

$$egin{aligned} n'(\mathcal{R}) &
ightarrow e^{i\zeta(\mathcal{R})} \ket{n(\mathcal{R})} \ \gamma_n &= \int_{\mathcal{C}} \mathrm{d}\mathcal{R} \cdot \mathcal{A}_n(\mathcal{R}) \end{aligned}$$

Fock 1928: one can always find suitable gauge  $\zeta(\mathcal{R})$  so that accumulated phase  $\gamma_n$  along any path  $\mathcal{C}$  is always zero  $\Rightarrow \gamma_n$  was thought to be unimportant and was usually neglected



Berry phase and Barry curvature

└─ Phase in parameters space, Berry connection

## Berry phase and Berry connection: gauge transformation

Berry 1984:

path C is closed, i.e.  $\mathcal{R}(0) = \mathcal{R}(T)$   $\Rightarrow$  We assume gauge exp $[i\zeta(\mathcal{R})]$  to be single-valued.

 $\Rightarrow$  the gauge must fullfils

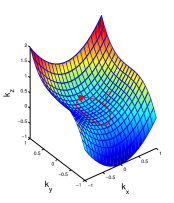
 $\zeta(\mathcal{R}(0)) - \zeta(\mathcal{R}(T)) = 2\pi N$ 

N is integer.

 $\Rightarrow$  Integral of phase over close path is well-defined, and gauge-invariant (with exception  $2\pi N$ )

$$\gamma_n = \oint_{\mathcal{C}} \mathrm{d}\mathcal{R} \cdot \mathcal{A}_n(\mathcal{R}) + 2\pi N$$

being known as Berry phase or geometrical phase.



Berry phase and Barry curvature

—Berry curvature

## Berry curvature I

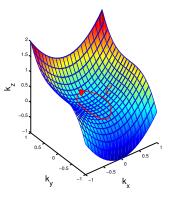
In following, we limit to 3D space of parameters  $\mathcal{R}$ . In case  $\mathcal{R}$  is in 3D, Stokes thorem relates Berry phase as

$$\gamma_n = \oint_{\mathcal{C}} \mathrm{d}\mathcal{R} \cdot \mathcal{A}_n(\mathcal{R}) = \int_{\mathcal{S}} \Omega_n(\mathcal{R}) \cdot \mathrm{d}\mathcal{S}$$

where  $\Omega_n(\mathcal{R})$  is called Berry curvature:

$$\Omega_n(\mathcal{R}) = \nabla_{\mathcal{R}} \times \mathcal{A}_n(\mathcal{R}) = i \left\langle \frac{\partial}{\partial \mathcal{R}} n \right| \times \left| \frac{\partial}{\partial \mathcal{R}} n \right\rangle$$

- Ω<sub>n</sub> is defined for each point of *R*-space, it is property of each point
- Ω<sub>n</sub> is gauge invariant (i.e. well-defined, measureble)



Berry phase and Barry curvature

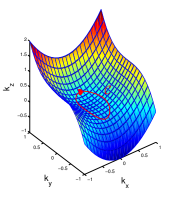
└─ Berry curvature

## Berry curvature in k-space

In solid state physics, parameter space  $\mathcal{R}$  is **k**-space, i.e.  $\mathcal{R} \equiv \mathbf{k}$ . Then, Berry curvature in **k**-space:

$$\Omega_n(\mathbf{k}) = \nabla_{\mathbf{k}} \times \mathcal{A}_n(\mathbf{k}) = i \langle \frac{\partial}{\partial \mathbf{k}} n | \times | \frac{\partial}{\partial \mathbf{k}} n \rangle$$

- Notice that Berry curvature writes in form 'gradient vector-times gradient', which is zero for scalar functions,  $\nabla \psi \times \nabla \psi = 0$
- Here, there is gradient in bra and ket wavefunctions, ⟨∇n| × |∇n⟩ ≠ 0, as ⟨∇n| and |∇n⟩ differ by phase.



Berry phase and Barry curvature

Berry curvature

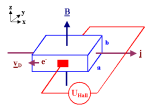
## Berry curvature and transport

 $\Omega_n$  can be viewed as the magnetic field in the k-space

$$egin{aligned} \dot{\mathbf{k}} &= -rac{e}{\hbar} \left( \mathbf{E} + \dot{\mathbf{r}} imes \mathbf{B} 
ight) \ \dot{\mathbf{r}} &= rac{1}{\hbar} rac{\partial \epsilon_n(\mathbf{k})}{\partial \mathbf{k}} - \dot{\mathbf{k}} imes \Omega_n(\mathbf{k}) \end{aligned}$$

 $\Rightarrow$  non-zero Berry curvature is responsible for phenomena in solids, where motion is perpendicular to the applied force

- Anomalous (Hall) efffects, spin-Hall effect, **j** =  $\sigma_{\text{Hall}}$  **M** × **E**
- Nerst effect, spin Nerst effect, j = NM × VT
- magneto-optical Kerr effect, magnetic circular dichroism: off-diagonal permittivity e<sub>xy</sub> ≠ 0



Berry phase and Barry curvature

Berry curvature

## Berry curvature: symmetry considarations

#### Inversion (point) symmetry:

Inversion symmetry is given by symmetry of the Hamiltonian

$$H(\mathbf{k}) = H(-\mathbf{k})$$

leading to symmetry in eigenstates and eigenvalues:

$$\epsilon_n(\mathbf{k}) = \epsilon_n(-\mathbf{k}), \qquad u_n(\mathbf{k}) = u_n(-\mathbf{k})$$

Then, following definition of  $\Omega_n(\mathbf{k}) = i \langle \frac{\partial}{\partial \mathbf{k}} n | \times | \frac{\partial}{\partial \mathbf{k}} n \rangle$ 

$$\Omega_n(\mathsf{k})=\Omega_n(-\mathsf{k})$$

#### Time reversal symmetry:

Under time reversal:  $\Omega_n(\mathbf{k}) = -\Omega_n(-\mathbf{k})$ 

 $\Rightarrow \Omega \neq 0$  only when time or inversion symmetry is broken

└─ Two level degenerate system

## Two level degenerate system

└─ Two level degenerate system

## Two level degenerate system: Hamiltonian

Let us have two-fold degenerate states |a
angle, |b
angle

```
 \begin{array}{l} H_0 \left| a \right\rangle = \epsilon_0 \left| a \right\rangle \\ H_0 \left| b \right\rangle = \epsilon_0 \left| b \right\rangle \end{array}
```

Adding perturbation Hamiltonian  $\Delta H$ , we search for solution  $|u\rangle$ 

$$(H + \Delta H) |u\rangle = \epsilon |u\rangle$$

Assuming the solution is in basis of original wavefunctions,  $|u\rangle = a |a\rangle + b |b\rangle$ , the general form of two-level perturbation Hamiltionan is as (i) *H* must be Hermitian,  $H = H^*$  (ii) without lost of generality, we assume degenerate eigenfrequencies are zeros  $(\epsilon_0 = 0)$ .

$$\Delta H(\mathcal{R}) = \begin{bmatrix} Z & X + iY \\ X - iY & -Z \end{bmatrix} = \begin{bmatrix} Z & R_+ \\ R_- & -Z \end{bmatrix}$$

#### Two level degenerate system: Hamiltonian

$$\Delta H(\mathcal{R}) = \begin{bmatrix} Z & X + iY \\ X - iY & -Z \end{bmatrix} = \begin{bmatrix} Z & R_+ \\ R_- & -Z \end{bmatrix}$$

where  $X(\mathcal{R})$ ,  $Y(\mathcal{R})$ ,  $Z(\mathcal{R})$  depends on path in the parameter space  $\mathcal{R}$ . Schrodinger equation has eigenvector  $|u\rangle = a |a\rangle + b |b\rangle$ 

$$\begin{bmatrix} Z & R_+ \\ R_- & -Z \end{bmatrix} \begin{bmatrix} a \\ b \end{bmatrix} = \epsilon \begin{bmatrix} a \\ b \end{bmatrix}$$

leading to two solutions +, -

$$\epsilon_{\pm} = \epsilon_0 \pm \sqrt{X^2 + Y^2 + Z^2} = \pm R$$
$$|\pm\rangle \equiv |u_{\pm}\rangle = a_{\pm} |a\rangle + b_{\pm} |b\rangle$$
$$\left(\frac{b}{a}\right)_{\pm} = -\frac{Z - \epsilon_{\pm}}{X + iY} = \frac{X - iY}{Z + \epsilon_{\pm}}$$

#### Two distinct meaning of Pauli matrices

$$\tilde{\sigma}_x = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix}$$
  $\tilde{\sigma}_y = \begin{bmatrix} 0 & -i \\ i & 0 \end{bmatrix}$   $\tilde{\sigma}_z = \begin{bmatrix} 1 & 0 \\ 0 & -1 \end{bmatrix}$ 

Pauli matrices as a spin operator S in up-down bases of the spin part of the wavefunction,

$$\chi = \begin{bmatrix} u_{\uparrow} \\ u_{\downarrow} \end{bmatrix}$$

The Hamiltonian of spin in magnetic field is (Zeeman term)

$$H = -\mu \cdot \mathbf{B} = \frac{e}{mc} \mathbf{S} \cdot \mathbf{B}, \qquad \mathbf{S} = \frac{\hbar}{2} \boldsymbol{\sigma} = \frac{\hbar}{2} \begin{bmatrix} \sigma_x \\ \sigma_y \\ \sigma_z \end{bmatrix}$$

Comment:  $\sigma$  denotes *vector* of Pauli matrices.

#### Two distinct meaning of Pauli matrices

$$\tilde{\sigma}_x = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix}$$
  $\tilde{\sigma}_y = \begin{bmatrix} 0 & -i \\ i & 0 \end{bmatrix}$   $\tilde{\sigma}_z = \begin{bmatrix} 1 & 0 \\ 0 & -1 \end{bmatrix}$ 

Pauli matrices as generating matrices of any Hermitian 2x2 matrix. I.e. they are generating (base) matrices of any two-level system

$$\Delta H(\mathcal{R}) = \begin{bmatrix} Z & X + iY \\ X - iY & -Z \end{bmatrix} = X\sigma_x + Y\sigma_y + Z\sigma_z = \mathbf{h} \cdot \boldsymbol{\sigma}$$

- $\rightarrow\,$  'spin-in-magnetic-field' problem is example of the two level degenerate problems.
- $\rightarrow$  appearance of  $\sigma$  matrices in two-level degenerate problem may be confusing, suggesting that problem handles spin.
- $\rightarrow\,$  two level degenerate system is a very common problem, e.g. spin-orbit coupling, graphene etc.

#### Two level degenerate system: spin in magnetic field

Free electron under magnetic field has perturbation Hamiltonian

$$\Delta H = -\boldsymbol{\mu} \cdot \mathbf{B} = \frac{e}{mc} \mathbf{S} \cdot \mathbf{B} = \frac{e}{mc} [S_x B_x + S_y B_y + S_z B_z]$$

where  $\mu$ , **S** are electron's operators of magnetic moment and spin angular momentum, respectively. The spin operators are

$$S_x = \frac{\hbar}{2}\sigma_x$$
  $S_y = \frac{\hbar}{2}\sigma_y$   $S_z = \frac{\hbar}{2}\sigma_z$ 

where in basis of up/down spins, the  $\sigma_{x/y/z}$  are Pauli matrices:

$$\tilde{\sigma}_{x} = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix} \qquad \tilde{\sigma}_{y} = \begin{bmatrix} 0 & -i \\ i & 0 \end{bmatrix} \qquad \tilde{\sigma}_{z} = \begin{bmatrix} 1 & 0 \\ 0 & -1 \end{bmatrix}$$

#### Two level degenerate system: spin in magnetic field

Then, normalized Hamiltonian in up/down spin basis writes:

$$\Delta H \frac{2mc}{e\hbar} = B_x \begin{bmatrix} 0 & 1\\ 1 & 0 \end{bmatrix} + B_y \begin{bmatrix} 0 & -i\\ i & 0 \end{bmatrix} + B_z \begin{bmatrix} 1 & 0\\ 0 & -1 \end{bmatrix}$$
$$\Delta H \frac{2mc}{e\hbar} = \begin{bmatrix} B_z & B_x + iB_y\\ B_x - iB_y & -B_z \end{bmatrix}$$

i.e. it has form of general two-level perturbation form as discussed above.

#### Two level degenerate system: spin in magnetic field

Solving problem, where size/direction of external field is a path parameter,  $\mathcal{R} = \frac{e\hbar}{2mc}\mathbf{B}$ . Then, Berry curvature is

$$m{\Omega}_+ = i rac{\langle + | m{
abla}_{m{\mathcal{R}}} H | - 
angle imes \langle - | m{
abla}_{m{\mathcal{R}}} H | + 
angle}{(\epsilon_+ - \epsilon_-)^2}$$

where  $\nabla_{\mathcal{R}} H$  is vector of  $\sigma$  matrices, namely  $\partial H/\partial x = \sigma_x$ ,  $\partial H/\partial y = \sigma_y$ ,  $\partial H/\partial z = \sigma_z$ .

Rotating coordinate system that  ${f B} \parallel z,$  Berry curvature  $\Omega_+$  is

$$\begin{split} \Omega_{+,x} &= i \left\langle + |\sigma_y| - \right\rangle \left\langle - |\sigma_z| + \right\rangle / 2R^2 = 0\\ \Omega_{+,y} &= i \left\langle + |\sigma_z| - \right\rangle \left\langle - |\sigma_x| + \right\rangle / 2R^2 = 0\\ \Omega_{+,z} &= i \left\langle + |\sigma_x| - \right\rangle \left\langle - |\sigma_y| + \right\rangle / 2R^2 = \frac{1}{2R^2} \end{split}$$

Rotating coordinate system back,

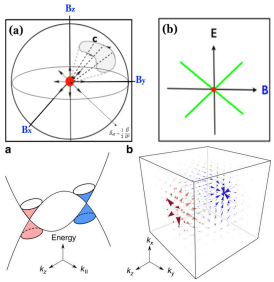
$$\Omega_{\pm}(\mathcal{R}) = \pm rac{\mathcal{R}}{2R^3} = 2\left(rac{mc}{e\hbar}
ight)^2 rac{\mathbf{B}}{B^3}$$

└─ Two level degenerate system

#### Two level degenerate system: Chern charge

$$\Omega_+(\mathcal{R})=rac{\mathcal{R}}{2R^3}\sim rac{\mathsf{B}}{B^3}$$

 $\Rightarrow \text{Berry curvature of} \\ \text{two-level degenerate state} \\ \text{behaves as originating} \\ \text{from monopole source} \\ \text{with strength } \frac{1}{2} \text{ located at} \\ \text{point of degeneracy, at} \\ \text{position } \mathcal{R} = 0 \\ \Rightarrow \text{ degeneracy points serve} \\ \text{as a source or drain of} \\ \text{Berry curvature flux.} \end{cases}$ 



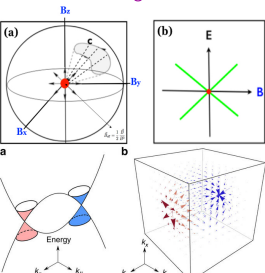
└─ Two level degenerate<u>system</u>

#### Two level degenerate system: Chern charge

Total flux of Berry curvature by close surface around this degeneracy point is

$$\frac{2}{4\pi}\oint_{S}\mathbf{\Omega}\cdot\mathrm{d}\mathbf{S}=\mathbf{N},$$

where N=integer, being called Chern charge, enclosed within the area S. Quantized nature of Chern charge is responsible of quantization of many phenomena (e.g. quantum Hall effect)

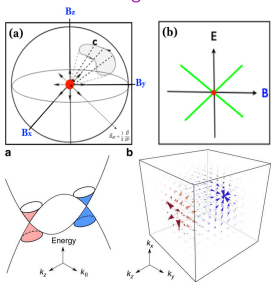


└─ Two level degenerate system

#### Two level degenerate system: Chern charge

The Berry phase (a.k.a geometrical phase) associated with close path C in the vicinity of the monopole of strength 1/2

$$\gamma_{\mathcal{C}} = \int_{\mathcal{S}} \mathbf{\Omega} \cdot d\mathbf{S}$$
  
=  $\frac{1}{2}$  (Solid angle given by  $\mathbf{S}$ 



#### Two level degenerate system: symmetry

Let as assume, in point  $\mathcal{R}_0,$  two-level system become degenerate,  $\epsilon_n\approx \epsilon_m$ 

 Then, Berry curvatures Ω<sub>m</sub>, Ω<sub>n</sub> in the vicinity of R<sub>0</sub> is dominantly determined by (nearly) degenerate states |n⟩, |m⟩ and having opposite signs

$$m{\Omega}_n = -m{\Omega}_m = i rac{\langle n | m{
abla}_{m{R}} H | m 
angle imes \langle m | m{
abla}_{m{R}} H | n 
angle}{(\epsilon_n - \epsilon_m)^2}$$

• Hence, the change of phase in path around degenerate point  $\mathcal{R}_0$  are opposite,  $\gamma_{n,C} = -\gamma_{m,C}$ .

#### Dispersion relation of free the electron

Newton (classical):

$$E = \frac{1}{2}mv^2 = \frac{p^2}{2m}$$

Einstein (special relativity):

$$E = \sqrt{p^2 c^2 + (mc^2)^2}$$
$$\approx mc^2 + \frac{p^2}{2m}$$

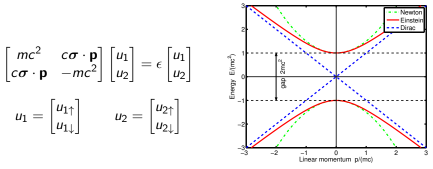
Dirac equation:

E = cp

└─ Two level degenerate<u>system</u>

#### Two level degenerate system: Dirac equation

Dirac equation, describing spin- $\frac{1}{2}$  particle (e.g. electron):



u<sub>1</sub>, u<sub>2</sub>: each two-component wavefunctions (up/down spin)
u<sub>1</sub>, u<sub>2</sub>: solution for particle (m > 0) and antiparticle (m < 0)</li>
for p = 0, gap 2mc<sup>2</sup> between particles and antiparticles

└─ Two level degenerate<u>system</u>

#### Two level degenerate system: effect of mass

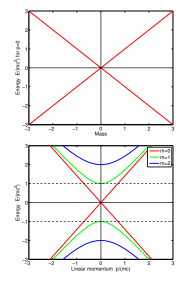
# $\rightarrow$ Effect of the diagonal term (mass)

Here, we keep  $\mathbf{p} = 0$ 

$$H_z = \begin{bmatrix} mc^2 & 0\\ 0 & -mc^2 \end{bmatrix}$$

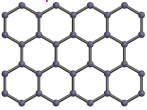
# $\rightarrow$ Effect of the off-diagonal terms (momentum)

$$H_{x} = \begin{bmatrix} mc^{2} & cp \\ cp & -mc^{2} \end{bmatrix}$$
$$H_{y} = \begin{bmatrix} mc^{2} & -icp \\ icp & -mc^{2} \end{bmatrix}$$



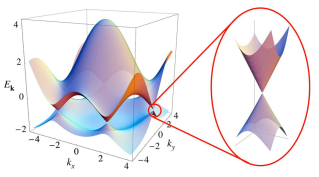
└─ Two level degenerate system

#### Examples of Dirac cones in electronic structure: Graphene



- 2D sheet of carbon
- 2D-Dirac cones
   (along k<sub>x</sub> and k<sub>y</sub>)
- up-down electrons are degenerate
- Fermi surface consists of just points.

Electronic structure of graphene:



http://faraday.fc.up.pt/cfp

### Weyl equation

Dirac equation for massless spin- $\frac{1}{2}$  particles  $(m_0 = 0) \rightarrow$  Weyl equation (in particle physics describing neutrino)

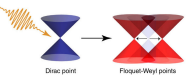
$$\begin{bmatrix} 0 & c\boldsymbol{\sigma} \cdot \mathbf{p} \\ c\boldsymbol{\sigma} \cdot \mathbf{p} & 0 \end{bmatrix} \begin{bmatrix} \varphi_L \\ \varphi_R \end{bmatrix} = |\mathbf{p}| c \begin{bmatrix} \varphi_L \\ \varphi_R \end{bmatrix}$$

which can be rewritten to

$$\begin{aligned} [|\mathbf{p}| + \boldsymbol{\sigma} \cdot \mathbf{p}] \varphi_L(\mathbf{p}) &= 0 \\ [|\mathbf{p}| - \boldsymbol{\sigma} \cdot \mathbf{p}] \varphi_R(\mathbf{p}) &= 0 \end{aligned} \tag{1}$$

solutions: spin helicity is fixed with the direction of motion.

In solids, terminology 'Dirac', 'Weyl' is used for degenerate, non-degenerate spin states on the cone, respectively



#### Two level degenerate system: Maxwell equation

Lossless Maxwell equations describing photon (boson with spin=1)

$$i \begin{bmatrix} 0 & \nabla \times \\ -\nabla \times & 0 \end{bmatrix} \begin{bmatrix} \mathbf{E} \\ \mathbf{B} \end{bmatrix} = \omega \begin{bmatrix} \epsilon & \chi \\ \chi^{\dagger} & \mu \end{bmatrix} \begin{bmatrix} \mathbf{E} \\ \mathbf{B} \end{bmatrix}$$

The main difference between fermions and bosons is time symmetry (fermions:  $T^2 = -1$ , bosons  $T^2 = 1$ , T is time operator)

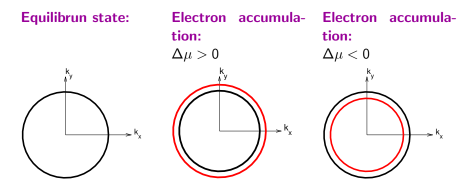
Transport properties

# Transport properties

Transport properties

#### Visualization of non-equilibrum electron states

Sketch of equilibrum electron distribution (i.e. Fermi surface) and examples of non-equilibrum electron distributions



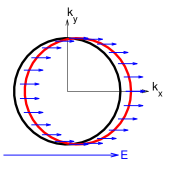
- Transport properties

## Introduction to electric conductivity

- applied electric field E
- all **k**-vectors move in the reciprocal space by  $\mathbf{k} = \mathbf{k}_0 \frac{e\mathbf{E}t}{\hbar}$
- it corresponds to speed of electrons

$$\mathbf{v} pprox rac{\hbar}{m} (\mathbf{k} - \mathbf{k_0}) = rac{e\mathbf{E}}{m} t$$

- i.e. classical acceleration of electrons by electric field
- this electron acceleration continues until electrons get scattered
- note: only electrons from unfilled bands contribute to the charge transfer
- **s** finally, charge current is generated  $\mathbf{j} = e \int_{\mathrm{BZ}} \mathbf{v}$



-Transport properties

## Electron dynamics in electric field

Electric field **E** is an perturbation to the Hamiltonian

$$\Delta H = e\phi(\mathbf{r}) = -e\mathbf{E}\cdot\mathbf{r}; \qquad \mathbf{E} = -\nabla\phi$$

 $\phi$  being electrostatic potential.

- However, this perturbation breaks translational symmetry of the crystal ⇒ Bloch theorem can not be applied.
- To introduce E without breaking time symmetry, E can be introduced by time-varying vector potential A(t),

$$\mathbf{E} = -
abla \phi - rac{\partial}{\partial t} \mathbf{A}$$
 $\mathbf{A}(t) = -\mathbf{E}t$ 

Transport properties

### Electron dynamics in electric field

Then, time dependent Hamiltonian has form

$$H(t) = \frac{1}{2m} [\mathbf{p} + e\mathbf{A}(t)]^2 + V(\mathbf{R})$$
$$= \frac{1}{2m} [\hbar \mathbf{q}]^2 + V(\mathbf{R})$$

•  $\hbar \mathbf{q} = \hbar (\mathbf{k} + e \mathbf{A}(t))$  is (time independent) canonical monentum

 $\Rightarrow~{\bm q}$  is still a good quantum number, with  ${\dot {\bm q}}=0$ 

- $\mathbf{p} = \hbar \mathbf{k} = \hbar (\mathbf{q} \mathbf{A}(t))$  is momentum.
- $\Rightarrow$  change of momentum **k** is simply

$$\dot{\mathbf{k}} = -rac{e}{\hbar}\mathbf{E}$$

- Transport properties

### Electron dynamics in electric field

Substituting  $\dot{\mathbf{k}} = -\frac{e}{\hbar}\mathbf{E}$  to transport equation

$$egin{aligned} \dot{\mathbf{k}} &= -rac{\mathbf{e}}{\hbar} \left( \mathbf{E} + \dot{\mathbf{r}} imes \mathbf{B} 
ight) \ \dot{\mathbf{r}} &= rac{1}{\hbar} rac{\partial \epsilon_n(\mathbf{k})}{\partial \mathbf{k}} - \dot{\mathbf{k}} imes \mathbf{\Omega}_n(\mathbf{k}) \end{aligned}$$

We got speed of Bloch electron in the crystal (in absence of scattering)

$$\mathbf{v}_n(\mathbf{k}) = rac{e\mathbf{E}}{m}t - rac{e}{\hbar}\mathbf{E} imes \mathbf{\Omega}_n(\mathbf{k})$$

- → the first term describes conductivity of the electronds (completly filled bands to not contribute to the conductivity).
   > the second term describes anomalous conductivity is a w + F
- $\rightarrow\,$  the second term describes anomalous conductivity, i.e.  $\mathbf{v}\perp\mathbf{E}$

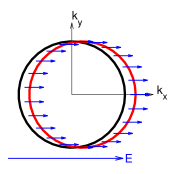
└─ Transport properties

(Ordinary) electric conductivity the speed of electron in solids (the first term):

$$\dot{\mathbf{r}} = rac{1}{\hbar} rac{\partial \epsilon_n(\mathbf{k})}{\partial \mathbf{k}}$$

can be approximately expressed as (assuming parabolic dispersion  $\epsilon \sim (\mathbf{k} - \mathbf{k}_0)^2$ ):  $\mathbf{v} \approx \frac{1}{\hbar} \frac{\partial}{\partial \mathbf{k}} \epsilon(\mathbf{k}) =$  $= \frac{1}{\hbar} \frac{\partial}{\partial \mathbf{k}} \left[ \frac{\hbar^2}{2m} (\mathbf{k} - \mathbf{k}_0)^2 \right]$  $= \frac{\hbar}{m} (\mathbf{k} - \mathbf{k}_0) = \frac{\hbar}{m} \dot{\mathbf{k}} t = \frac{e\mathbf{E}}{m} t$ 

providing just simple acceleration of the electron by electric field



#### Transport properties

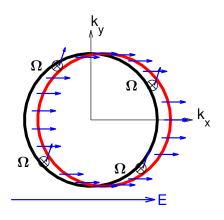
## Anomalous electric conductivity

we add non-zero Berry curvature  $\Omega$  to our sketch of electron states

- Berry curvature on Fermi surafce is a point/line feature.
- here, we assume crystal with:
  - ightarrow time symmetry is broken
  - $\rightarrow\,$  inversion symmetry is present

 $\Omega(\mathsf{k})=\Omega(-\mathsf{k})$ 

- $\label{eq:constraint} \begin{array}{l} \Rightarrow \mbox{ due to equal direction of } \Omega, \mbox{ all} \\ \mbox{ anomalously scattered electrons} \\ \mbox{ are scattered in one direction} \\ \mbox{ (up), } \textbf{v} \sim \textbf{E} \times \Omega \end{array}$
- $\begin{array}{l} \Rightarrow \mbox{ anomalous (Hall) current,} \\ \textbf{j}_{Hall} \perp \textbf{E} \mbox{ is created, due to} \\ \mbox{ break of time symmetry} \end{array}$



└─ Transport properties

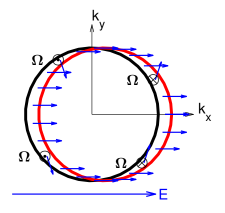
## Anomalous electric conductivity

now, add non-zero Berry curvature, but with broken inversion symmetry

- here, we assume crystal with:
  - ightarrow inversion symmetry is broken
  - $\rightarrow \,$  time symmetry is present

$$\Omega(\mathsf{k}) = -\Omega(-\mathsf{k})$$

- $\Rightarrow \mbox{ due to different direction of } \Omega, \\ \mbox{ there is no net current} \\ \mbox{ perpendicular to } E$
- $\Rightarrow$  anomalous (Hall current),  $j_{\rm Hall}\perp$  E is NOT created, due to presence of time symmetry



Transport properties

#### Visualization of non-equilibrum electron states

Applied electric field **E** in *x*-direction  $\Rightarrow$  charge current is generated

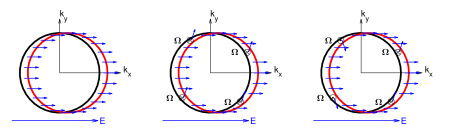
No Berry curvature

#### Berry curvature

no time symmetry inversion symmetry  $\Omega({\bf k}) = \Omega(-{\bf k})$ 

#### Berry curvature

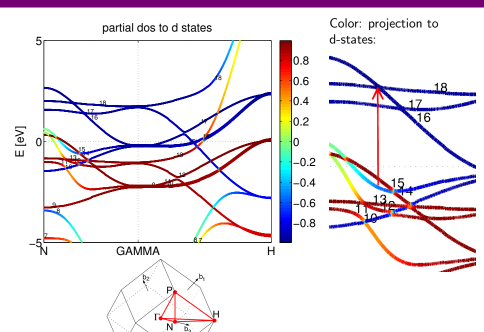
no inversion symmetry time symmetry  $\Omega({\bf k})=-\Omega(-{\bf k})$ 



Examples of Berry curvatures on bcc Fe

# Examples of Berry curvatures on bcc Fe

Examples of Berry curvatures on bcc Fe

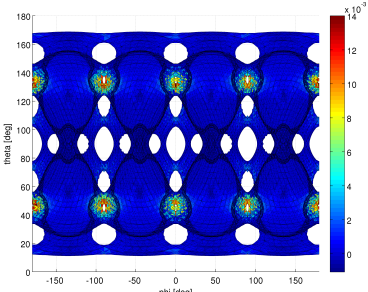


Examples of Berry curvatures on bcc Fe

## isoenergy surface E = 2.23 eV, bands 14,17, color= $\Omega_z$

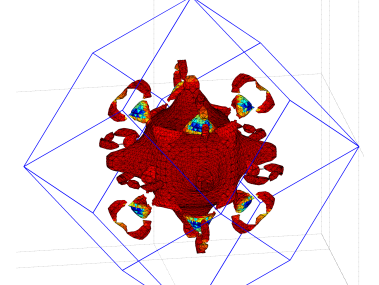
Examples of Berry curvatures on bcc Fe

#### isoenergy surface E = 2.23 eV, bands 14, 17, 2D view



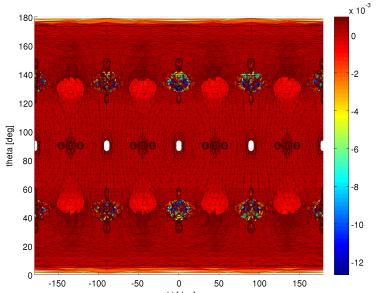
Examples of Berry curvatures on bcc Fe

## isoenergy surface E = 2.35 eV, bands 14,18, color= $\Omega_z$



Examples of Berry curvatures on bcc Fe

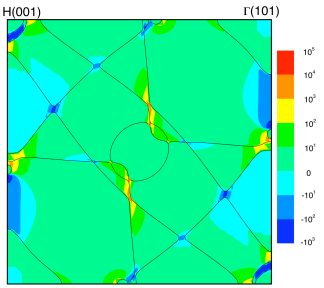
### isoenergy surface E = 2.35 eV, bands 14, 18, 2D view



Examples of Berry curvatures on bcc Fe

#### Integrated Berry curvature on bcc Fe

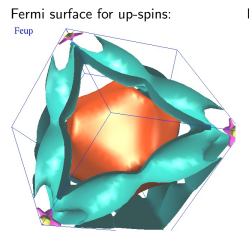
Fermi surface in (010) plane (solid lines) and the integrated Berry curvature  $-\Omega_z(\mathbf{k})$  in atomic units (color map) of fcc Fe. From Yao et al.,PRL, 2004.



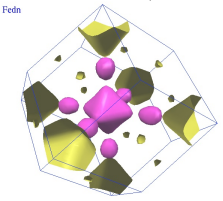
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Examples of Berry curvatures on bcc Fe

#### Fermi surface on bcc Fe



#### Fermi surface for down-spins:



Relation between Berry curvature and Kubo formula

# Relation between Berry curvature and Kubo formula

Relation between Berry curvature and Kubo formula

#### Berry curvature: dc Hall conductivity

Conductivity in x-direction, when field is applied in y-direction:

$$j_{x} = -ev_{x}$$

$$= -e\sum_{n} \int_{BZ} \frac{d^{3}\mathbf{k}}{(2\pi)^{3}} f_{n}(\mathbf{k})v_{x}$$

$$= -\frac{e^{2}}{\hbar} \sum_{n} \int_{BZ} \frac{d^{3}\mathbf{k}}{(2\pi)^{3}} f_{n}(\mathbf{k}) (\mathbf{E} \times \Omega_{n}(\mathbf{k}))_{x}$$

$$= -\frac{e^{2}}{\hbar} \sum_{n} \int_{BZ} \frac{d^{3}\mathbf{k}}{(2\pi)^{3}} f_{n}(\mathbf{k}) E_{y}\Omega_{n,z}(\mathbf{k})$$

where  $f_n(\mathbf{k})$  is Fermi distribution function. Note, amplitude  $(j_x)$  is perpendicular to applied force  $(E_x)$ .

#### Berry curvature: dc Hall conductivity

off-diagonal (Hall conductivity) given by Berry curvature writes:

$$\sigma_{xy} = \frac{j_x}{E_y} = -\frac{e^2}{\hbar} \sum_n \int_{BZ} \frac{d^3 \mathbf{k}}{(2\pi)^3} f_n(\mathbf{k}) \Omega_{n,z}(\mathbf{k})$$

On the other hand, the well-know relation describing conductivity (and light absorption) is Kubo formula:

$$\sigma_{xy} = i \frac{e^2}{m_e^2 \hbar} \sum_n \sum_{n' \neq n} \int_{BZ} \frac{d^3 \mathbf{k}}{(2\pi)^3} \left[ f_n(\mathbf{k}) - f_{n'}(\mathbf{k}) \right] \\ \times \frac{\langle n | p_x | n' \rangle \langle n' | p_y | n \rangle - \langle n | p_y | n' \rangle \langle n' | p_x | n \rangle}{(\epsilon_n - \epsilon_{n'})^2}$$

How to relate both expressions?

#### Relation between Berry curvature and Kubo formula II

First, let us establish identity:

$$\begin{aligned} \epsilon_{n'} \langle \nabla_{\mathbf{k}} n | n' \rangle &= \langle \nabla_{\mathbf{k}} n | H n' \rangle \\ &= \langle n | \nabla_{\mathbf{k}} H | n' \rangle + \langle n | H | \nabla_{\mathbf{k}} n' \rangle \\ &= \langle n | \nabla_{\mathbf{k}} H | n' \rangle + \epsilon_n \langle n | \nabla_{\mathbf{k}} n' \rangle \end{aligned}$$

taking into account  $\langle n|\nabla_{\bf k}n'\rangle=\langle n|\nabla_{\bf k}n'\rangle=\langle n|\nabla_{\bf k}|n'\rangle$  we obtain

$$\langle \nabla_{\mathbf{k}} n | n' \rangle (\epsilon_{n'} - \epsilon_n) = \langle n | \nabla_{\mathbf{k}} H | n' \rangle$$

#### Relation between Berry curvature and Kubo formula II

• applying the identity using complete set of orthonormal basis vectors  $\mathbf{1} = \sum_{m} |m\rangle \langle m|$ 

$$\mathbf{\Omega} = \nabla_{\mathbf{k}} \times i \langle n | \nabla_{\mathbf{k}} n \rangle = i \langle \nabla_{\mathbf{k}} n | \times | \nabla_{\mathbf{k}} n \rangle = i \sum_{m \neq n} \langle \nabla_{\mathbf{k}} n | m \rangle \times \langle m | \nabla_{\mathbf{k}} n \rangle$$

substituting 
$$\langle \nabla_{\mathbf{k}} n | m \rangle (\epsilon_m - \epsilon_n) = \langle n | \nabla_{\mathbf{k}} H | m \rangle$$
  
$$\Omega_n = i \sum_{n \neq m} \frac{\langle n | \nabla_{\mathbf{k}} H | m \rangle \times \langle m | \nabla_{\mathbf{k}} H | n \rangle}{(\epsilon_n - \epsilon_{n'})^2}$$

#### Relation between Berry curvature and Kubo formula II

Taking into account

$$abla_{\mathbf{k}}H = rac{\hbar^2}{m}\mathbf{k} = rac{\hbar}{m}\mathbf{p}$$

Berry curvature can be expressed by matrix elements of momentum:

$$\boldsymbol{\Omega}_n = i \frac{\hbar^2}{m_e} \sum_{n \neq m} \frac{\langle n | \mathbf{p} | m \rangle \times \langle m | \mathbf{p} | n \rangle}{(\epsilon_n - \epsilon_m)^2}$$

- $\rightarrow\,$  this form of Berry curvature corresponds to Kubo formula
- $\rightarrow\,$  this form is used to express Kubo formula by calculations (both analytical and numerical), to avoid gradient of wavefunctions.

└─ Time reversal symmetry

# Time reversal symmetry

#### Time reversal symmetry I

(according F. Haake, Time Reversal and Unitary Symmetries in Quantum Signatures of Chaos)

A classical Hamiltonian system is called time invariant, when for time reversal  $t \rightarrow -t$ , the solutions transforms as (conventional invariance):

$$\begin{array}{cccc} t \rightarrow -t & \mathbf{x} \rightarrow \mathbf{x} & \mathbf{p} \rightarrow -\mathbf{p} & \mathbf{L} \rightarrow -\mathbf{L} & \mathbf{S} \rightarrow -\mathbf{S} \\ \psi \rightarrow T \psi & \end{array}$$

where T is time-reversal operator, being so-called antiunitary, defined as:

$$\langle T\psi | T\phi \rangle = \langle \psi | \phi \rangle^{\star} = \langle \phi | \psi \rangle$$

because

- overlap of two wavefunctions must be preserved
- explicit '*i*' in Schrodinger equation;  $i\hbar\partial\psi/\partial t = H\psi$

#### Time reversal symmetry II

Time reveral operator T is antiunitary  $\Rightarrow$  T<sup>2</sup> is unitary,  $|T^2| = 1$ 

 $\Rightarrow$  time-reversal operator can be written as

$$T = UK$$

where K is complex conjugation and U is unitary operator  $\Rightarrow T^2 = \pm 1$ 

•  $T^2 = 1$  for spinless or spin-integer particles (e.g. photons)

• 
$$T^2 = -1$$
 for spin-1/2 particles (electrons)

Note:

- Hamiltonian is non-dissipative (i.e. conserving phase-space volume according to Liouville theorem)
- $\Rightarrow\,$  following discussions regarding time reversal does not work for dissipative system.

Note: unitary operator:  $U^* = U^{-1}$ , i.e. complex conjugation equals inverse (e.g. U is rotation operator, keeping angles and lengths).

### Spinless particle

Schrodinger equation  $i\hbar \frac{\partial}{\partial t}\psi(\mathbf{x},t) = H\psi(\mathbf{x},t)$ Hamiltonian  $H = \frac{\mathbf{p}^2}{2m} + V(x)$  is called time-reversal invariant when having solution  $\psi(\mathbf{x},t)$  for t, there is another solution  $\psi'(\mathbf{x},t')$  for t' = -t uniquely related to  $\psi(\mathbf{x},t)$ . In case of so-called conventional time reversal

$$t \rightarrow -t$$
  $\mathbf{x} \rightarrow \mathbf{x}$   $\mathbf{p} \rightarrow -\mathbf{p}$   
 $\psi(\mathbf{x}) \rightarrow \psi^{\star}(\mathbf{x}) = K\psi(\mathbf{x})$ 

with K being operator of complex conjugation, with  $K^2 = 1$ . Hence, in case of spinless particle, time reversal operator T equals operator of complex conjugation K, T = K

$$T \ket{\psi} = K \ket{\psi} = \ket{\psi}^*$$

Spin-1/2 particle requires reversal of spin under time reversal $\langle T\phi | T\mathbf{S}\psi \rangle = \langle T\phi | T\mathbf{S}T^{-1} | T\psi \rangle = -\langle \phi | \mathbf{S} | \psi \rangle$ 

providing

$$T\mathbf{S}T^{-1} = -\mathbf{S}$$

 $\Rightarrow$  operator T can not be solely complex conjugation  $\Rightarrow$  T = UK,

### Spin-1/2 particle: derivation of form of *T*-operator

Derivation of T = UK for spin-1/2 particle  $(\mathbf{S} = \frac{\hbar}{2}\boldsymbol{\sigma})$ :

$$T\sigma_i T^{-1} = -\sigma_i$$

U must have general form as (general form of any Hermitian 2x2 matrix)

$$U = \alpha \sigma_x + \beta \sigma_y + \gamma \sigma_z + \delta$$

we get only non-zero term  $\beta=i$  and hence time operator for  ${\rm spin-}1/2$  particles is

$$T = i\sigma_y K = i \begin{bmatrix} 0 & -i \\ i & 0 \end{bmatrix} K = \exp[i\pi\sigma_y/2]K$$

#### Kramers' degeneracy

For any Hamiltonian invariant under time reversal

$$[H, T] = 0$$

its eigenvectors  $|\psi_n\rangle$  and  $|T\psi_n\rangle$  has equal eigenvalues  $E_n$ .

• for  $T^2 = 1$ : we can choose eigenvectors to follow  $|\psi\rangle = |T\psi\rangle$  using combination  $|\psi\rangle = a |\psi'\rangle + aT |\psi'\rangle$ 

• for  $T^2=-1,~|\psi
angle$  and  $|T\psi
angle$  are orthogonal

$$\langle \psi | T\psi \rangle = \langle T\psi | T^2\psi \rangle^* = - \langle T\psi | \psi \rangle^* = - \langle \psi | T\psi \rangle = 0$$

 $\Rightarrow$  all eignevalues of *H* are double degenerate, with eigenvectors  $|\psi\rangle$  and  $T |\psi\rangle \equiv |T\psi\rangle$ , called Kremers' degeneracy.

For example, in case of single-electron Hamiltonian without SOC and without B, spin-up and spin-down states are degenerates.

# Kramers' degeneracy without geometric symmetries I

For  $T^2 = -1$ , we adopt basis

$$|1\rangle, |T1\rangle, |2\rangle, |T2\rangle, ... |N\rangle, |TN\rangle$$

General wavefunction can be written as

$$|\psi\rangle = \sum_{m} a_{mt+} |m\rangle + a_{mt-} |Tm\rangle$$
  
 $T\psi\rangle = \sum_{m} a_{mt+}^{\star} |Tm\rangle - a_{mt-}^{\star} |m\rangle$ 

As  $|T\psi
angle = U\!K\,|\psi
angle$ , U must have form

$$U_{mm} = \begin{bmatrix} 0 & -1 \\ 1 & 0 \end{bmatrix} = -i\sigma_y \qquad U_{mn} = 0 \text{ for } m \neq n$$

#### Kramers' degeneracy without geometric symmetries II

Similarly, Hamiltonian can be written as

$$h_{mn} = \begin{bmatrix} \langle m|H|n \rangle & \langle m|H|Tn \rangle \\ \langle Tm|H|n \rangle & \langle Tm|H|Tn \rangle \end{bmatrix}$$

where the matrix form of Hamiltonian element  $h_{mn}$  can be written using Pauli matrices and four real  $h_{mn}^{(0...3)}$  with

$$h_{mn} = h_{mn}^{(0)} \mathbf{1} - i \mathbf{h}_{mn} \cdot \boldsymbol{\sigma}$$

This is similar form of Hamiltonian we discussed for two-level degenerate system

 $\Rightarrow$  break of time symmetry when Hamiltonian contains terms not fulfilling time reversal symmetry (such as SOC) then in general off-diagonal terms are nonzero, diagonal terms do not equal  $\Rightarrow$  break of time symmetry provides splitting of eigenvalues and non-zero Berry curvature.

#### Kramers' degeneracy without geometric symmetries III

In case of photons,  $T^2 = 1 \Rightarrow$  Hamiltonian element

$$h_{mn} = \begin{bmatrix} \langle m|H|n \rangle & \langle m|H|Tn \rangle \\ \langle Tm|H|n \rangle & \langle Tm|H|Tn \rangle \end{bmatrix}$$

is diagonal,  $h_{mn} = h_{mn}^{(0)} \mathbf{1}$ 

 $\Rightarrow$  break of time symmetry (i.e. Hamiltonian non-invariant under time reversal,  $[H, T] \neq 0$ ) does not provide energy splitting or Berry curvature of photon wavefunction

 $\Rightarrow$  splitting of photon eigenvalues can be obtained by interaction with solids (magnetooptics)

—Spin-orbit interaction of light

# Spin-orbit interaction of light

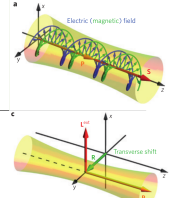
### Spin-orbit interaction of light

K.Y. Bliokh, Nature Photon. 9, 796 (2015) Similar to spin-orbit coupling of electron, there is spin-orbit coupling also for photon (for light beam).

- Spin-orbit interaction of electron: connect spin wavefunction and space wavefunction
- Spin orbit coupling of photon: connects beam trajectory and its polarization state (also know as optical spin-Hall effect).

#### Angular momenta of light I

## Spin angular momentum S: degree of circular polarization (helicity) $\sigma = (-1, 1)$ $\mathbf{S} = \sigma \frac{\mathbf{k}}{k} = \sigma \frac{\mathbf{p}}{p}$



Extrinsic orbital angular momentum  $L^{e\times t}$ : determined by the trajectory of the beam R  $L^{e\times t} = \mathbf{R} \times \mathbf{p}$ 

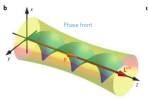
### Angular momenta of light II

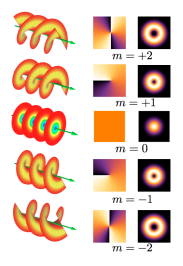
**Intrinsic angular momentum** *L*<sup>*int*</sup>: helical phase front: phase of the beam depends on position inside beam,

approximately

 $E(r, z, \phi) \approx E_0(r, z) \exp[il\phi]$ , where r, z,  $\phi$  are coordinates in cylindrical coordinates.

$$L^{int} = I \frac{\mathbf{k}}{k}, \qquad I \in \mathbb{Z}$$

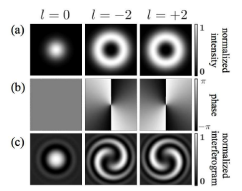




#### Angular momenta of light III

# Intrinsic angular momentum L<sup>int</sup>:

Laguerre-Gaussian modes: •symmetrical beams with non-zero intrinsic angular momentum of light •also called optical vortex with topological charge *I* 



$$u(r, z, \phi) = \exp\left(-ik\frac{r^2}{2R(z)}\right) \exp(-il\phi) \exp(-ikz) \exp(i\psi(z))$$

Helical mode conversion using conical reflector, Optics Express 20, 14064 (2012)

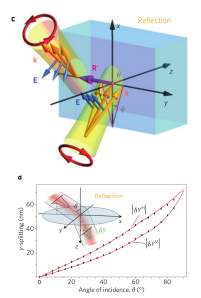
#### Optical spin Hall effect I

Interactions between those three angular momenta of light represents spin-orbit coupling of light. **Example: optical spin Hall effect:** The light beam on reflection displaces (shifts) according to the beam helicity  $\sigma$ . Consequence of total angular momentum conservation:

 $\mathbf{J} = \mathbf{S} + \mathbf{L}^{ext} = \mathbf{S} + \mathbf{R} \times \mathbf{p}$ 

adjusting coordinates that incident beam has R=0 and hence  $\textbf{L}^{ext}=0$   $\textbf{S}-\textbf{S}'\approx \textbf{R}'\times \textbf{p}'$ 

K.Y. Bliokh, Nature Photon. 9, 796 (2015)

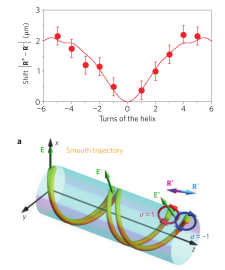


#### Optical spin Hall effect II

 $J = S + L^{ext} = const.$ displacement of beam due to light helicity inside the glass cylinder with gradient of refraction index.

$$\dot{\mathbf{p}} = \nabla n(\mathbf{R})$$
  $\dot{\mathbf{R}} = \frac{\mathbf{p}}{p} - \frac{\sigma}{k_0} \frac{\mathbf{p} \times \dot{\mathbf{p}}}{p^3}$ 

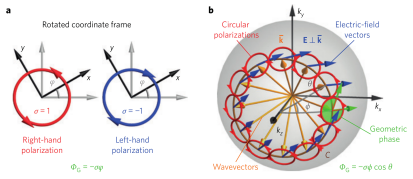
(overdot denotes derivation according to the trajectory,  $\mathbf{p} = \mathbf{k}/k_0$  is dimensionless momentum)



Bliokh Nature Photon. 2, 748 (2008)

#### Optical spin Hall effect III

The relation between light polarization and trajectory can be laso understand in terms of Berry phase:



Phase of the light wave varies as light propagetes inside glass cylinder providing cyclic variation of wavevector direction  ${\bf p}$ 

#### Optical spin Hall effect IV

Berry connection:

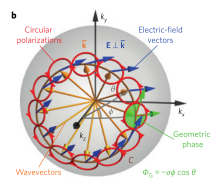
$$\mathbf{A} = -i\mathbf{E}^{\sigma}\cdot\nabla_{\mathbf{k}}\mathbf{E}^{\sigma}$$

Berry curvature:

$$\Omega^{\sigma}(\mathbf{k},\sigma) = 
abla imes \mathbf{A} = \sigma rac{\mathbf{k}}{k^3}$$

Geometric (Berry) phase:

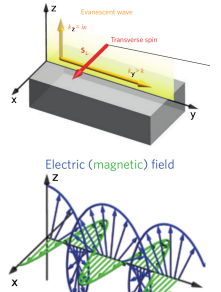
$$\Phi_{G} = \int_{C} \mathbf{A} d\mathbf{k} = \int_{S} \Omega dS_{\mathbf{k}}$$
$$= 2\pi\sigma(1 - \cos\theta)$$



#### Spin transfer by evanescent wave

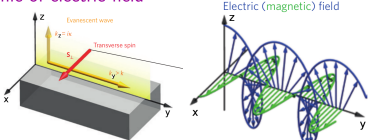
Electrical field of the evanescent wave (propagation along y, recall  $\mathbf{E} \cdot \mathbf{k} = 0$ ):

$$\mathbf{k} = \begin{bmatrix} 0\\k_y\\i\mathcal{K} \end{bmatrix} \qquad \mathbf{E} = \begin{bmatrix} 0\\i(-\mathcal{K}/k_y)E_z\\E_z \end{bmatrix}$$

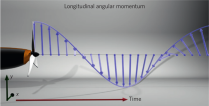


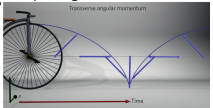
Spin-orbit interaction of light

#### Profile of electric field



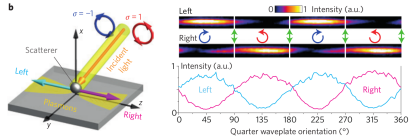
Profiles of electric field in paraxial beam and evanescent beam, providing longitudinal and transverse spin angular momentum.





#### Selection of evanescent wave propagation

As spin of the evanescent wave is given by propagation of spinwave, the polarization (helicity) of incoming beam determines propagation direction of the evanescent wave.



"Remarkably, the universal character of spin-direction locking in evanescent waves can be associated with the quantum spin-Hall effect of photons, which makes it an optical counterpart of the quantum spin-Hall effect of electrons in topological insulators"

-Spin-orbit interaction

# Spin-orbit interaction

└─ Dirac equation

### Spin-orbit coupling: Dirac equation

Spin-orbit coupling term couples spin of the electron  $\sigma = 2\mathbf{S}/\hbar$  with movement of the electron  $m\mathbf{v} = \mathbf{p} - e\mathbf{A}$  in presence of electrical field  $\mathbf{E}$ .

$$egin{aligned} \mathcal{H}_{SOC} = -rac{e\hbar}{4m^2c^2}m{\sigma}\cdot\left[\mathbf{E} imes(\mathbf{p}-e\mathbf{A})
ight] \end{aligned}$$

The maximal coupling is obtained when all three componets are perpendicular each other.

The spin-orbit term can be determined from solution of electron state in relativistic case. The equation describing relativistic electron is called Dirac equation, relativistic analogue of Schrodinger equation.

└─ Dirac equation

#### Dirac equation: introduction I

- Relativity describes nature at high speeds,  $v \approx c$ .
- Relativity unites time and space, described by Lorentz transformation

$$x' = rac{x - vt}{\sqrt{1 - rac{v^2}{c^2}}} \qquad t' = rac{t - rac{vx}{c^2}}{\sqrt{1 - rac{v^2}{c^2}}}$$

⇒ relativistic quantum theory must do the same. Schrodinger equation does not fulfils this, as it it has first derivative in time and second in space.

└─ Dirac equation

#### Dirac equation: introduction II

Relativistic theory expresses total energy of the particle as:

$$W^2 = p^2 c^2 + m_0^2 c^4 \tag{3}$$

Quantum operator substitution:  $\mathbf{p} \rightarrow \hat{\mathbf{p}} = -i\hbar\nabla$ ,  $W \rightarrow \hat{W} = i\hbar\partial/\partial t$ . It follows in Klein-Gordon equation

$$\left(\nabla^2 - \frac{1}{c^2}\frac{\partial^2}{\partial t^2} - \frac{m_0^2 c^2}{\hbar^2}\right)\psi(\mathbf{r}, t) = 0$$
(4)

This Eq. reduces to Eq. (3) for plane wave (free particle)  $\psi(\mathbf{r}, t) = \exp[i(\mathbf{r} \cdot \mathbf{p} - Wt)/\hbar]$ . This condition limits following solutions to particles with spin 1/2, as space-time wavefunction is symmetric, and hence spin-part must be antisymmetric.

└─ Dirac equation

#### Dirac equation: derivation I

- I let as ASSUME, the Dirac equation will have first derivative in time. Then, it must be also in first derivative in space.
- 2 wave function is superposition of N base wavefunctions  $\psi(\mathbf{r}, t) = \sum \psi_n(\mathbf{r}, t)$
- 3 must fulfil Klein-Gordon equation, Eq. (4)

General expression of condition 1:

$$\frac{1}{c}\frac{\partial\psi_i(\mathbf{r},t)}{\partial t} = -\sum_{w=x,y,z}\sum_{n=1}^N \alpha_{i,n}^w \frac{\partial\psi_n}{\partial w} - \frac{imc}{\hbar}\sum_{n=1}^N \beta_{i,n}\psi_n(\mathbf{r},t) \quad (5)$$

Dirac equation

#### Dirac equation: derivation II

When expressed in matrix form ( $\psi$  is column vector,  $\alpha_{i,n}^k$  is  $3 \times N \times N$  matrix,  $\beta_{i,n}$  is  $N \times N$  matrix)

$$\frac{1}{c}\frac{\partial\psi(\mathbf{r},t)}{\partial t} = -\tilde{\boldsymbol{\alpha}}\cdot\nabla\psi(\mathbf{r},t) - \frac{imc}{\hbar}\tilde{\beta}\psi(\mathbf{r},t)$$
(6)

Substituting quantum operators  $\hat{\mathbf{p}} 
ightarrow -i\hbar 
abla$ , we get Dirac equation

$$i\hbar \frac{\partial \psi(\mathbf{r},t)}{\partial t} = \hat{H}\psi(\mathbf{r},t) = (c\tilde{\boldsymbol{\alpha}} \cdot \hat{\mathbf{p}} + \tilde{\beta}mc^2)\psi(\mathbf{r},t)$$
(7)

where matrices  $\tilde{\alpha}$ ,  $\tilde{\beta}$  are unknown.

-Spin-orbit interaction

Dirac equation

#### Dirac equation: non-relativistic limit

When Dirac equation is solved up to order  $1/c^2$ , we get

$$\begin{split} \hat{H} &= \frac{1}{2m} \left( \frac{\hbar}{i} \nabla - e \mathbf{A}(\mathbf{r}) \right)^2 + V(\mathbf{r}) + mc^2 \qquad \text{Unrelativistic Hamiltonian} \\ &- \frac{e\hbar}{2m} \boldsymbol{\sigma} \cdot \mathbf{B} \qquad \text{Zeeman term} \\ &- \frac{e\hbar}{4m^2c^2} \boldsymbol{\sigma} \cdot [\mathbf{E} \times (\mathbf{p} - e\mathbf{A})] \qquad \text{Spin-orbit coupling} \\ &- \frac{1}{8m^3c^2} (\mathbf{p} - e\mathbf{A})^4 \qquad \text{Mass of electron increases with speed} \\ &+ \frac{\hbar^2 e}{8m^2c^2} \nabla^2 V(\mathbf{r}) \qquad \text{Darwin term} \end{split}$$

Darwin term: electron is not a point particle, but spread in volume of size of Compton length  $\approx \hbar/mc$ .

-Spin-orbit interaction

Understanding spin-orbit coupling

#### Spin-orbit coupling: discussion I

Spin-orbit coupling term can be sepaarted into two components:

$$-rac{e\hbar}{4m^2c^2}m{\sigma}\cdot\left[\mathbf{E} imes(\mathbf{p}-e\mathbf{A})
ight]=-rac{e\hbar}{4m^2c^2}m{\sigma}\cdot\left[\mathbf{E} imes\mathbf{p}
ight]+rac{e^2\hbar}{4m^2c^2}m{\sigma}\cdot\left[\mathbf{E} imes\mathbf{A}
ight]=H_{SOC}+H_{AME}$$

AME=Angular magneto-electric

• The electric field  $\mathbf{E} = -\frac{1}{e}\nabla V - \frac{\partial}{\partial t}\mathbf{A}$ 

■ canonical momentum  $\mathbf{p} = -i\hbar\nabla$  (conjugate variable of position;  $\frac{\partial H}{\partial x_i} = -\dot{p}_i$ ,  $\frac{\partial H}{\partial p_i} = \dot{x}_i$ )

• kinetical momentum  $m\mathbf{v} = \mathbf{p} - e\mathbf{A}$  (defines kinetic energy and represents velocity)

-Spin-orbit interaction

Understanding spin-orbit coupling

#### $H_{SOC}$ in spherical potential, static case

$$H_{SOC} = -rac{e\hbar}{4m^2c^2} \boldsymbol{\sigma} \cdot [\mathbf{E} imes \mathbf{p}]$$

Spherical potential  $V(\mathbf{r}) = V(|\mathbf{r}|) = V(r)$ ; static case  $\frac{\partial}{\partial t}\mathbf{A} = 0$ :

$$e\mathbf{E} = -\nabla V(|\mathbf{r}|) = \frac{dV(r)}{dr} \frac{\mathbf{r}}{|\mathbf{r}|}$$

providing:

$$H_{SOC} = \frac{\hbar}{4m^2c^2} \frac{1}{r} \frac{dV}{dr} \boldsymbol{\sigma} \cdot (\mathbf{r} \times \mathbf{p}) = \frac{1}{2m^2c^2} \frac{1}{r} \frac{dV}{dr} \mathbf{S} \cdot \mathbf{L} = \xi \mathbf{S} \cdot \mathbf{L}$$

where spin angular momentum  ${\bf S}=\frac{\hbar}{2}{\pmb \sigma}$  and orbital angular momentum  ${\bf L}={\bf r}\times{\bf p}$ 

-Spin-orbit interaction

└─ Understanding spin-orbit coupling

### Understanding spin-orbit coupling: spherical potential

 spin of the electron creates electron's magnetic moment (in SI)

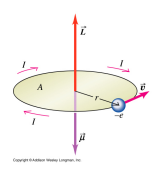
$$\boldsymbol{\mu}_{S} = -\frac{e}{m}\mathbf{S} = -\frac{e}{m}\frac{h}{2}\boldsymbol{\sigma} = -\mu_{b}\boldsymbol{\sigma} = -\frac{2\mu_{B}}{h}\mathbf{S}$$

where μ<sub>B</sub> = eħ/2m is Bohr magneton.
 orbital moment (around atomic core) creates magnetic moment too

$$\boldsymbol{\mu}_L = -\frac{e}{2m} \mathbf{L} = -\frac{\mu_B}{\hbar} \mathbf{L} = -\mu_B I$$

(or can be understood as creating magnetic field  $H_{eff}$  due to current created by electron orbital)

• the mutual static energy of spin and orbital is then  $E_{SO,approx} = -\mu_S \cdot \mathbf{B}_{eff}$  or just electrostatic interaction between both magnetic dipoles.



Understanding spin-orbit coupling

#### Understanding spin-orbit coupling: Lorentz transformation

Electromagnetic field appears different as observing frame is moved. For example, if a charge is moving in the laboratory frame (unprimed), we observe both electric and magnetic fields. In the frame of the moving charge (primed), only electric field is observed and the current and magnetic field are absent. Lorentz transformation of el.-mag. fields between both frames is:

$$\begin{split} \mathbf{E}'_{\parallel} &= \mathbf{E}_{\parallel} & \mathbf{B}'_{\parallel} &= \mathbf{B}_{\parallel} \\ \mathbf{E}'_{\perp} &= \frac{(\mathbf{E} + \mathbf{v} \times \mathbf{B})_{\perp}}{\sqrt{1 - \frac{v^2}{c^2}}} & \mathbf{B}'_{\perp} &= \frac{(\mathbf{B} - \mathbf{v}/c^2 \times \mathbf{E})_{\perp}}{\sqrt{1 - \frac{v^2}{c^2}}} \end{split}$$

where  $\perp$  and  $\parallel$  are relative to the direction of the velocity **v**. I.e. for small speeds,  $\mathbf{E}' = \mathbf{E} + \mathbf{v} \times \mathbf{B}$  and  $\mathbf{B}' = \mathbf{B} - \frac{\mathbf{v}}{c^2} \times \mathbf{E}$ 

Understanding spin-orbit coupling

### Understanding spin-orbit coupling: Lorentz transformation

For electron flying by speed **v** through static electric field **E**, in its frame the electron feels magnetic field  $\mathbf{B}' = -\frac{-\mathbf{v}}{c^2} \times \mathbf{E}$ , which torques/acts on its spin. The Hamiltonian is given by Zeeman interaction

$$H_{SO,E\to B} = -\mu_S \cdot \mathbf{B}' \tag{8}$$

$$= -\left(-\frac{e\hbar}{2m}\sigma\right) \cdot \left(-\frac{1}{c^2}(-\mathbf{v}) \times \mathbf{E}\right)$$
(9)  
$$= -\frac{e\hbar}{2m^2c^2}\sigma \cdot (\mathbf{E} \times \mathbf{p})$$
(10)

which is twice larger compared to  $H_{SOC}$  derived from Dirac equation. Missing half is due to Thomas precession (in case of electron orbiting nucleus, it is the precession of the electron rest frame as it orbits around the nucleus).

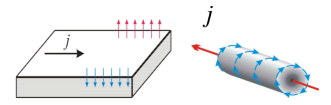
-Spin-orbit interaction

Understanding spin-orbit coupling

### Lorentz transformation: extrinsic spin Hall effect

In laboratory frame, spin-Hall effect provides scattering of electrons on charged impurity along to electron spin.

In electron frame, it can be understood as charge current from impurities, providing magnetic field, according which the electron spin aligns.





**Figure 1.** The Spin Hall Effect. An electrical current induces spin accumulation at the lateral boundaries of the sample. In a cylindrical wire the spins wind around the surface, like the lines of the magnetic field produced by the current. However the value of the spin polarization is much greater than the (usually negligible) equilibrium spin polarization in this magnetic field.

-Spin-orbit interaction

Examples of spin-orbit effects

#### Examples of spin-orbit effects

$$H_{SOC} = -rac{e\hbar}{4m^2c^2} \boldsymbol{\sigma} \cdot \left[ \mathbf{E} imes (\mathbf{p} - e\mathbf{A}) 
ight]$$

Various SOC effects are obtained by different origins of **A** and  $\mathbf{E} = \frac{1}{e} \nabla V - \frac{\partial}{\partial t} \mathbf{A}$ . Examples:

- SOC in spherical potential (already discussed)
- optical spin pumping: excitation of electrons with selective spins in GaAs
- E has contribution originating from interface of two materials: → Rasha effect
- A has contribution of incident light: coupling between angular momentum of light and electron spin (optomagnetic field)

-Spin-orbit interaction

Examples of spin-orbit effects

#### Example: splitting of atomic levels by SOC

Splitting of atomic levels due to spin-orbit coupling (without magnetic field). The energy levels corresponds to different values of the total angular momentum  ${\bf J}$ 

 $\mathbf{J} = \mathbf{L} + \mathbf{S}$ 

$$\mathbf{J} \cdot \mathbf{J} = (\mathbf{L} + \mathbf{S}) \cdot (\mathbf{L} + \mathbf{S}) = \mathbf{L} \cdot \mathbf{L} + \mathbf{S} \cdot \mathbf{S} + 2 \langle \mathbf{L} \cdot \mathbf{S} \rangle$$
$$j(j+1) = l(l+1) + s(s+1) + 2 \langle \mathbf{L} \cdot \mathbf{S} \rangle$$
$$\langle \mathbf{L} \cdot \mathbf{S} \rangle = \frac{1}{2} [j(j+1) - l(l+1) - s(s+1)]$$

For p states, l = 1, s = 1/2 and j = 3/2 (4 electrons) or 1/2 (2 electrons). So, due to spin-orbit coupling (without magnetic field), the energy level of electron splits into two levels.

Thus, the spin-orbit interaction does not lift all the degeneracy for atomic states. To lift this additional degeneracy it is necessary to apply a magnetic field.

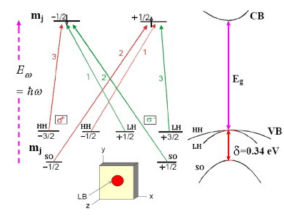
-Spin-orbit interaction

└─ Examples of spin-orbit effects

#### Optical spin orientation

Electron excitation by circularly polarized beam in GaAs excites electrons with selective spins.

- for  $\hbar\omega$  between  $E_g$  and  $E_g + \Delta_{SO}$ , only the light and heavy hole subband are excited. Then for zinc-bland structure (e.g. GaAs), the spin-polarization is  $P_n = -1/2$ .
- Light polarization can also be used to detect spin polarization in semiconductors.



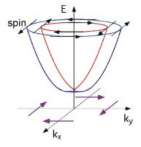
Examples of spin-orbit effects

## Rashba effect I

Rashba Hamiltonian: electric field **E** is created on interface, **E**  $\parallel \hat{z}$ :

$$H_{\text{Rashba}} = \alpha(\boldsymbol{\sigma} \times \mathbf{p}) \cdot \hat{\mathbf{z}}$$

- $\alpha: \text{ Rasba coupling}$
- p: electron's momentum
- $\sigma$ : spin direction (Pauli matrix vector)



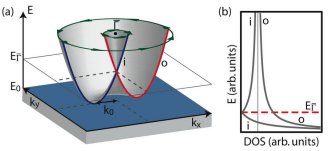
The Rashba effect is a momentum dependent splitting of spin bands in two-dimensional condensed matter systems (heterostructures and surface states). It originates from concurrent appearance of

- spin-orbit coupling
- asymmetry of the potential in the direction  $\hat{z}$  perpendicular to the two-dimensional plane, creating electric field  $\mathbf{E} = E_z \hat{z} = -\frac{1}{e} \nabla V$

-Spin-orbit interaction

Examples of spin-orbit effects

### Rashba effect II



http://www.sps.ch/fr/articles/progresses/

- For  $\mathbf{p} = p_x$  and  $H_{\text{Rashba}} = \alpha(\boldsymbol{\sigma} \times \mathbf{p}) \cdot \hat{\mathbf{z}} \Rightarrow H_{\text{Rashba}} = -\alpha p_x \sigma_y$
- splitting of energy states according to  ${\bf p}$  and  ${\bf \sigma}$  directions.
- **•** max. splitting when z, **p** and  $\sigma$  are perpendicular each other.
- when crystal lacks inversion symmetry, internal electric field E is created.

-Spin-orbit interaction

Examples of spin-orbit effects

#### Optomagnetic field I

according to: Paillard, Proc. of SPIE 9931, 99312E-1 (2016)

$$H_{AME} = -rac{e^2\hbar}{4m^2c^2}m{\sigma}\cdot[\mathbf{E} imes\mathbf{A}]$$

Assume electric field as plane wave

$$\mathbf{E}_{\text{ext}} = -\frac{\partial \mathbf{A}}{\partial t} = \Re \left( \mathbf{E}_0 \exp[i(\mathbf{k} \cdot \mathbf{r} - \omega t)] \right)$$

providing vector potential as  $\mathbf{A} = \Re(-\frac{i}{\omega}\mathbf{E}_0 \exp[i(\mathbf{k} \cdot \mathbf{r} - \omega t)])$ 

- Electric field acting on electron has two contributions,
   E = E<sub>int</sub> + E<sub>ext</sub>, E<sub>int</sub> = -1/e∇V provided by crystal and E<sub>ext</sub> provided by incoming el.-mag. field.
- term [E<sub>int</sub> × A] vanishes as E<sub>int</sub> varies much quicker compared to A (due to a ≪ λ).

-Spin-orbit interaction

Examples of spin-orbit effects

#### Optomagnetic field II

$$H_{AME} = -\frac{e^{2}\hbar}{8m^{2}c^{2}\omega}\boldsymbol{\sigma} \cdot \Re[i\mathbf{E}_{0} \times \mathbf{E}_{0}^{*}] = -\mu_{B} \cdot \mathbf{B}_{OM}$$
$$\mathbf{B}_{OM} = -\frac{\mu_{B}}{\varepsilon_{0}c^{3}\omega\hbar}\boldsymbol{I}\boldsymbol{\sigma}_{\text{helicity}}$$

- $\mu = -\mu_b \sigma$ : electron magnetic moment,  $\mu_b = e\hbar/(2m)$  Bohr magneton
- $\sigma_{\text{helicity}} = \Re[i\mathbf{u} \times \mathbf{u}]$ : helicity of beam, where  $\mathbf{u}$  is beam polarization,  $\mathbf{u} = \mathbf{E}_0 / E_0$

• 
$$I = \frac{c\varepsilon_0}{2}E_0^2$$
 : beam intensity

• direction of  $\mathbf{B}_{OM}$  is determined by helicity of the incident beam  $\sigma_{\mathrm{helicity}}$ 

Note: although  $\mathbf{B}_{OM}$  contributes to magnetization torque by induced light, it is not probably the dominanting term.