Graphene, Index Theorem and Topological Degeneracy

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Quantum Information, Physics and Topology

- **Encoding and manipulating QI** in small physical systems is pledged by decoherence and control errors.

- **Error correction** can be employed to resolve this problem by using a (huge) overhead of qubits and quantum gates.

- An alternative method is to employ **intrinsically error protected systems** such as topological ones => properties are described by integer numbers! protected by macroscopic properties: hard to destroy.

- E.g. you can employ system with **degenerate ground states**:
  - Make sure degeneracy is protected by topological properties (V)
  - Make sure degenerate states are **locally indistinguishable** (X)
  - Encode information in these degenerate levels

**TOPOLOGICAL DEGENERACY**
Overview

- **Graphene**: two dimensional layer of graphite –hexagonal lattice of C atoms
  - Fullerene: C60, C70
  - Nanotubes
- Conducting properties of these materials: zero energy modes.
- Can be used as miniaturized elements of circuits.

- **Index theorem** (Atiyah-Singer)
  - Smooth, orientable, compact, Riemannian manifolds, M, with genus, \( g \).
  - Define elliptic operator \( D \) on M. Includes curvature and gauge fields.
  - The **index theorem** relates the **number of zero energy modes** of \( D \) with \( g \).

- **Conductivity** can depend on topology.

- Zero modes provide **degeneracy** of ground state: \( G \)-2 deg.
- **Topological quantum computation**
  - Kitaev’s toric code
  - Honeycomb lattice (same as graphene, but with “real” fermions)
Different geometries of Graphene

Fullerene (C60):

Nanotubes:
Graphene: structure

The Hamiltonian of graphene is given by

\[ H = -t \sum_{<i,j>} a_i^+ a_j = -\frac{t}{2} \sum_{<i,j>} (a_i^+ b_j + b_i^+ a_j) \]

\( a_i \) fermionic modes

Fourier transformation:

\[
H_{\vec{k}} = \begin{pmatrix}
0 & -t(1 + e^{-i\vec{k} \cdot \vec{u}} + e^{-i\vec{k} \cdot \vec{v}}) \\
-t(1 + e^{i\vec{k} \cdot \vec{u}} + e^{i\vec{k} \cdot \vec{v}}) & 0
\end{pmatrix}
\]

\[
E(\vec{k}) = \pm t \sqrt{3 + 2 \cos \vec{k} \cdot \vec{u} + 2 \cos \vec{k} \cdot \vec{v} + 2 \cos \vec{k} \cdot (\vec{u} - \vec{v})}
\]

Fermi points: \( E(k) = 0 \)
Graphene: structure

\[ E(\vec{k}) = \pm t \sqrt{3 + 2 \cos \vec{k} \cdot \vec{u} + 2 \cos \vec{k} \cdot \vec{v} + 2 \cos \vec{k} \cdot (\vec{u} - \vec{v})} \]

Linearise energy \( E(\vec{k}) \) around a conical point,

\[ \vec{k} = \vec{K} + \vec{p} \]

\[ H_{\vec{p}} \approx \pm \frac{3t}{2} \begin{pmatrix} 0 & p_x + ip_y \\ p_x - ip_y & 0 \end{pmatrix} = \pm \frac{3t}{2} \vec{\sigma} \cdot \vec{p} \]

**Relativistic Dirac equation** at the tip of a pencil!

Two types of spinors:

\[ \begin{pmatrix} |K_+, A\rangle \\ |K_+, B\rangle \end{pmatrix}, \quad \begin{pmatrix} |K_-, A\rangle \\ |K_-, B\rangle \end{pmatrix} \]

\( K_{\pm} \) are the Fermi points and A and B are the two triangular sub-lattices

Note: \( \vec{\sigma}^z \) rotation maps to states with the same energy, but opposite momenta
Graphene: curvature

To introduce curvature:
- cut $\pi / 3$ sector and reconnect sites.
This creates a single **pentagon** with no other deformations present.
Results in a **conical configuration**.
To preserve continuity of the spinor field when circulating the pentagon one can introduce two additional fields:

- Spin connection $Q$: \[ \int Q_\mu dx^\mu = -\frac{\pi}{6} \sigma^z \] Mixes A and B components

- Non-abelian gauge field, $A$: \[ \int A_\mu dx^\mu = -\frac{\pi}{2} \tau^y \] Mixes + and – spinors

Resulting 4x4 Dirac equation can be decoupled by simple rotation to a pair of 2x2 Dirac equations ($\kappa=1,2$):

\[ \frac{3t}{2} \sum_{a,\mu} \sigma^a e^\mu_a (p_\mu - iQ_\mu - iA^k_\mu) \psi^k = E \psi^k \]

\[ \int A^k_\mu dx^\mu = \pm \frac{\pi}{2} \]
Index Theorem

Consider finite matrices, \( M, M^+ \) \( V_+ \xrightarrow{M} V_- \), \( V_- \xrightarrow{M^+} V_+ \)

For \( \lambda \neq 0 \), \( M^+M u = \lambda u \Rightarrow (M M^+)M u = \lambda M u \)

Define operator:

\[
Q = \begin{pmatrix} 0 & M^+ \\ M & 0 \end{pmatrix}, \quad Q^2 = \begin{pmatrix} M^+ M & 0 \\ 0 & M M^+ \end{pmatrix}
\]

with eigenvalues +1, -1 for \( V_+, V_- \)

Define operator:

\[
\gamma_5 = \begin{pmatrix} 1 & 0 \\ 0 & -1 \end{pmatrix}
\]

Consider \( V_+, V_- \) the dimension of the null subspace of \( V_+, V_- \)

Then

\[
\text{Tr}(\gamma_5 e^{-tQ^2}) = \sum_{\text{Sp}(M^+M)} e^{-t\lambda^2} - \sum_{\text{Sp}(MM^+)} e^{-t\lambda^2} = v_+ - v_- \equiv \text{index}(Q)
\]

Non-zero eigenvalues cancel in pairs. Expression is \( t \) independent.
Index Theorem

Consider 2-dimensional **Dirac** operator defined over a **compact** surface \( Q = D \)

\[
D = \sigma^x p_x + \sigma^y p_y = \begin{pmatrix}
  0 & (\partial_x - iA_x) - i(\partial_y - iA_y) \\
  (\partial_x - iA_x) + i(\partial_y - iA_y) & 0
\end{pmatrix}
\]

\[
D^2 = -\nabla^2 - \sigma^z F_{xy} = \begin{pmatrix}
  -\nabla^2 - F_{xy} & 0 \\
  0 & -\nabla^2 + F_{xy}
\end{pmatrix}
\]

One can show that

\[
e^{-t(-\nabla^2 + \sigma^z F_{xy})} \bigg|_{r=r'} = \frac{1}{4\pi t} \left[ 1 + \sigma^z F_{xy} t + \left( \frac{1}{3} F_{xy}^2 + \frac{1}{2} \sigma^z \nabla^2 F_{xy} \right) t^2 + \ldots \right]
\]

That gives

\[
Tr[\sigma^z e^{-t(-\nabla^2 - \sigma^z F_{xy})}] = \frac{1}{2\pi} \int F_{xy} d^2 x
\]

Higher order terms in \( t \) have to be checked...
Index Theorem

\[ \text{Tr}(\gamma_5 e^{-tQ^2}) = \sum_{Sp(M^+M)} e^{-t\lambda^2} - \sum_{Sp(MM^+)} e^{-t\lambda^2} = \nu_+ - \nu_\equiv \text{index}(Q) \]

Assume it holds for infinite dimensions (continuum).

\[ \text{Tr}[\sigma^z e^{-t(-\nabla^2 - \sigma^z F_{xy})}] = \frac{1}{2\pi} \int F_{xy} d^2x \]

If \( D \) is defined on compact manifold then RHS is an integer (topological invariant). Open boundary conditions can give a discrepancy caused by boundary terms. Thus, the number of zero modes depends on the gauge field configuration.

Continuous deformations of the gauge field will not change the number of zero modes.

Including surface \textbf{curvature} does not change the above result (only in 2-dims).
Index Theorem

The Index theorem states:

\[
\text{index}(D) = \nu_+ - \nu_- = \frac{1}{2\pi} \int \int F
\]

integer!

The integral is taken over the whole compact surface.

F: field strength of gauge vector potential, A.

For compact manifolds the term on the r. h. s. is an integer. It is a topological number: small deformations does not change its value.

From this theorem you can obtain the least number of zero modes. The exact number is obtained if \( \nu_+ \) or \( \nu_- \) is equal to zero.

[Atiyah and Singer, Ann. of Math. 87, 485 (1968);...]
**Index Theorem: Euler characteristic**

**Euler characteristic** for lattices on compact surfaces:

\[ \chi = V - E + F = 2(1 - g) = \frac{1}{2\pi} \iint R \]

Consider folding of graphene in a compact manifold. The **minimal** violation is obtained by insertion of **pentagons** or **heptagons** that contribute positive or negative curvature respectively. Consider

- \( n_5 \) number of pentagons
- \( n_6 \) number of hexagons
- \( n_7 \) number of heptagons

\[
\begin{align*}
V &= (5n_5 + 6n_6 + 7n_7) / 3 \\
E &= (5n_5 + 6n_6 + 7n_7) / 2 \\
F &= n_5 + n_6 + n_7
\end{align*}
\]

From the Euler characteristic formula:

\[ n_5 - n_7 = 12(1 - g) \]

**Fullerenes:** \( g = 0 \Rightarrow n_5 = 12 \)

“**Nanotubes**”: \( g = 1 \Rightarrow n_5 - n_7 = 0 \)
Index Theorem: Graphene application

\[ \int \int F = \oint A \]

\[ \frac{1}{2\pi} \left( \pm \frac{\pi}{2} \right) (n_5 - n_7) = \pm 3(1 - g) \]

Stokes’s theorem

\[ \text{index}(D) = \nu_+ - \nu_- = \frac{1}{2\pi} \int \int F \]

Thus, one obtains:

\[ \nu_+ - \nu_- = \begin{cases} 
3(1 - g), & \text{for } k = 1 \\
-3(1 - g), & \text{for } k = 2 
\end{cases} \]

Least number of zero modes:

\[ 6 |1 - g| \]
Index Theorem: Graphene application

\[ \text{index}(D) = \nu_+ - \nu_- = 6 |1 - g| \]

C60: \( g=0 \)  
Nanotubes: \( g=1 \)

Zero mode pairs
No zero modes

Ultra-cold Fermi atoms and optical lattices

Single species ultra cold **Fermi atoms** superposed by **optical lattices** that form a hexagonal lattice.

[Duan *et al.* Phys. Rev. Lett. 91, 090402 (2003)]

- Very low temperatures: \( T \approx 0.1T_F \)
- Arbitrary filling factors: e.g. 1/2

See dependence of conductivity on **disorder**, **impurities** and lattice **defects**: e.g. insert pentagons at the edge of the lattice of effect of empty sites.

Similar **index theorem** can be devised for open boundary conditions.

**Measurement** of conductivity in Fermi lattices has already been performed in the laboratory:

Conclusions

- **Index Theorem** for compactified graphene sheets.
- Agrees well with known models of *fullerenes* and *nanotubes*.
- Gives conductivity properties for *higher genus models*:
  - sideways connected nanotubes.
- Predicts **stability** of spectrum under small deformations.

- Relate to **topological models**:
  - obtain **topologically related degeneracy**: $2^6|1-g|$
  - encode and manipulate **quantum information**.
  - apply **reverse engineering** to find new models with specific degeneracy properties.

- Related experiments with **ultra-cold Fermi atoms** can give insight to the properties of graphene. May be easier to implement than solid state setup.

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*Thank you for your attention!*
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