Quantum transport in carbon nanotubes

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Carbon nanotubes are a versatile material in which many aspects of condensed matter physics come together. Recent discoveries have uncovered new phenomena that completely change our understanding of transport in these devices, especially the role of the spin and valley degrees of freedom. This review describes the modern understanding of transport through nanotube devices. Unlike in conventional semiconductors, electrons in nanotubes have two angular momentum quantum numbers, arising from spin and valley freedom. The interplay between the two is the focus of this review. The energy levels associated with each degree of freedom, and the spin-orbit coupling between them, are explained, together with their consequences for transport measurements through nanotube quantum dots. In double quantum dots, the combination of quantum numbers modifies the selection rules of Pauli blockade. This can be exploited to read out spin and valley qubits and to measure the decay of these states through coupling to nuclear spins and phonons. A second unique property of carbon nanotubes is that the combination of valley freedom and electron-electron interactions in one dimension strongly modifies their transport behavior. Interaction between electrons inside and outside a quantum dot is manifested in SU(4) Kondo behavior and level renormalization. Interaction within a dot leads to Wigner molecules and more complex correlated states. This review takes an experimental perspective informed by recent advances in theory. As well as the well-understood overall picture, open questions for the field are also clearly stated. These advances position nanotubes as a leading system for the study of spin and valley physics in one dimension where electronic disorder and hyperfine interaction can both be reduced to a low level.

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I. INTRODUCTION AND MOTIVATION

Carbon nanotubes are exceptional materials in many different ways. They are mechanically ultrastrong, the surface is perfectly clean, electrons move ballistically, and they vibrate like guitar strings with record-breaking quality factors. Moreover, by zipping nanotubes open one obtains the other wonder material, graphene. Together with C_{60} buckyballs and diamond, these allotropes of carbon have a central position in nanotechnology. Many of their properties have been studied and reviewed in great detail (Saito, Dresselhaus, and Dresselhaus, 1998).

Nanotube electronic transport properties have been studied since the mid-1990s, first in bulk and since 1997 using individual single-wall nanotubes (Bockrath et al., 1997; Tans et al., 1997). Many of the basic transport properties were quickly discovered, including Coulomb blockade, Fabry-Pérot interference, 1D electronic interactions, Kondo physics, spintronics effects, and induced superconductivity. These properties have all been comprehensively reviewed, with both theoretical (Charlier, Blase, and Roche, 2007) and experimental focus (Biercuk et al., 2008, Schönenberger, 2006).¹ The general understanding in 2008 can be described as "consistent on a coarse scale." On a fine scale the specific properties arising from residual disorder together with the specific, usually unknown, chirality of the nanotube under study were hampering a detailed description. On a coarse scale all nanotubes showed similar transport behavior, but on a fine scale each experimentally studied nanotube was unique.

An important technical advance was a device scheme in which the nanotube was not exposed to any

¹The early generation of nanotube experiments that established basic quantum dot behavior was reviewed by Nygård *et al.* (1999) and Yao, Dekker, and Avouris (2001). Open devices and early attempts to analyze the quantum dot shell structure were described by Liang, Bockrath, and Park (2005) and Sapmaz, Jarillo-Herrero *et al.* (2006). Hybrid devices involving superconducting and ferromagnetic leads have been reviewed by de Franceschi *et al.* (2010) and Cottet *et al.* (2006), respectively, while aspects pertinent to one-dimensional wires were addressed by Deshpande *et al.* (2010). Coupled quantum dots were introduced in Schönenberger (2006) and Biercuk *et al.* (2008), whereas only recent reviews introduce spin-orbit interaction and valley physics (Ilani and McEuen, 2010; Kuemmeth *et al.*, 2010), which are the themes of this review.

fabrication chemicals, thereby retaining pristine material quality (Cao, Wang, and Dai, 2005). Transport experiments on such "ultraclean" nanotubes immediately showed more reproducible detail despite the still unknown chirality. Most importantly, the role of spin-orbit interaction was strikingly uncovered (Kuemmeth *et al.*, 2008). Although this spin-orbit interaction had already been predicted (Ando, 2000), it went unobserved and was therefore largely ignored until 2008. The experimental clarity revealed, however, that detailed understanding of quantum phenomena in carbon nanotubes has to include this effect.

The electronic orbits in nanotubes come in two flavors, known as the *K* and *K'* valleys, that roughly correspond to clockwise and counterclockwise motion around the nanotube. The resulting quantum states form interesting superpositions of spin up and spin down with the *K* and *K'* valleys. Our central aim is to present a coherent description of spin-orbit and K - K' physics in carbon nanotubes. We present the theory on a conceptual level and make references to detailed calculations in the literature. More details of the theoretical background are given in Appendix B. We highlight experimental results that demonstrate the essential concepts most clearly.

Spin-orbit and K - K' physics and their experimental consequences are described in Secs. II and III for nanotubes confined as single quantum dots. The quantum dot geometry allows for a precise, straightforward description of energy eigenstates, which can be probed with well-established techniques of Coulomb blockade spectroscopy. Double quantum dots increase the complexity, with quantum states now described by three numbers: spin (up or down), valley (K or K'), and location (left or right). Since the occupancy of both quantum dots is so easily controlled by gate voltages, the double-dot geometry provides for exquisite experimental control. Section IV describes spin-valley selection rules for tunneling, probed by Pauli blockade experiments. The experimental control in double dots is utilized further in Sec. V describing the realization and operation of qubits employing various choice of basis states.

Sections II–V make use of a simplified model with electronelectron interactions included as a capacitive charging energy. In Sec. VI we extend this picture to include interactions between quantum dot states and the continuum in the leads. Quantum dots strongly coupled to leads show renormalization of the energy states as well as the formation of macroscopic coherence in a Kondo state. Section VI focuses on renormalization and Kondo effects in the specific context of spinorbit and K - K' physics. In Sec. VII we consider interaction effects within quantum dots, which can be extraordinarily strong in the one-dimensional geometry of nanotubes. In quantum dots of a somewhat longer length, this leads to the formation of correlated Wigner molecules.

II. BASICS OF CARBON NANOTUBE DEVICES

A. Structure of carbon nanotubes

Carbon nanotubes consist of one or more concentric cylinders of graphene (Saito, Dresselhaus, and Dresselhaus, 1998). Both multiwall and single-wall carbon nanotubes

(MWCNTs and SWCNTs) can be synthesized and measured, but in this review we discuss only SWCNTs. As well as being simpler, these are the most studied both experimentally and theoretically.

The structure of nanotubes derives from the structure of graphene. A SWCNT is equivalent to a rolled-up strip taken from the two-dimensional honeycomb of carbon atoms that makes up a graphene sheet [Fig. 1(a)]. Since there are infinitely many ways of selecting a strip to roll up, there are correspondingly many different nanotube structures. Each structure is specified by its chiral vector \mathbf{C} , which connects lattice sites on opposite sides of the strip that are superposed

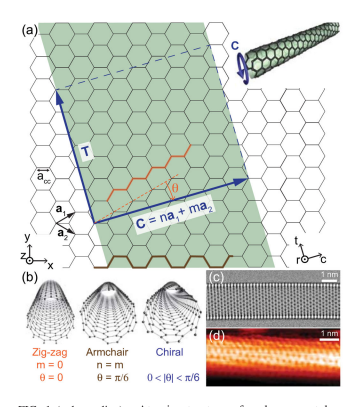


FIG. 1 (color online). Atomic structure of carbon nanotubes. (a) Derivation of nanotube structure from graphene. A singlewall nanotube is equivalent to a rolled-up graphene strip (shaded, with the direction of rolling chosen so that the printed pattern faces outward). The chiral vector C spans the nanotube circumference (inset) and connects lattice sites that are brought together by rolling up. Chiral indices (n, m) completely define the nanotube structure. The unit cell of the nanotube (which is much larger than the unit cell of graphene) is outlined by dashed lines, and the unit vector T is indicated. Graphene coordinates (x, y, z), nanotube coordinates (t, c, r), and the chiral angle θ are also marked. In this example, (n, m) = (6, 2)and $\theta = 13.9^{\circ}$. (b) Nanotubes are divided into three classes according to their chiral indices: zigzag, armchair, or chiral. Zigzag and armchair nanotubes are so called because of the shape of the edge formed by a cut perpendicular to the nanotube axis [see highlighted lines in (a)]. These three nanotubes are (12,0), (6,6), and (6,4). (c), (d) Nanotubes directly imaged by transmission electron microscopy (TEM) [(c), a (28,0) zigzag nanotube] and scanning tunneling microscopy [(d), an unidentified chiral nanotube]. Adapted from Venema et al., 1998, Charlier, Blase, and Roche, 2007, Warner et al., 2011, and Churchill, 2012.

by rolling up into a nanotube. A given structure is labeled by its chiral indices (n, m), which are the coordinates of the chiral vector $\mathbf{C} = n\mathbf{a}_1 + m\mathbf{a}_2$ in terms of the graphene basis vectors \mathbf{a}_1 and \mathbf{a}_2 . From Fig. 1(a), *n* and *m* are integers; to ensure that the same structure is not labeled two different ways, *m* is conventionally taken in the range $-n/2 < m \le n$. Instead of specifying (n, m), a nanotube can also be described by its diameter and chiral angle θ , defined as the angle between **C** and \mathbf{a}_1 .

Two special cases are zigzag structures (m = 0) and armchair structures (n = m), so called because of the arrangement of atoms along a cut normal to the nanotube. Structures not in either category are called chiral [Fig. 1(b)]. Unlike armchair and zigzag structures, chiral nanotubes lack inversion symmetry; the inversion isomer (with $\theta \rightarrow -\theta$) of an (n, m) chiral structure is an (n+m, -m) structure. From the Onsager-Casimir relations, the transport properties of isomer pairs are expected to be similar, but they may differ in their nonlinear conductance in the presence of electron-electron interactions and time-reversal symmetry breaking by a magnetic field (Ivchenko and Spivak, 2002; Sanchez and Büttiker, 2004; Spivak and Zyuzin, 2004; Wei et al., 2005). The differences between isomer pairs, well established in optical measurements (Samsonidze et al., 2004; Peng et al., 2007), are not discussed further here. Some structure parameters and their dependence on chiral indices are given in Table I.

This structure is confirmed by atomic-resolution microscopy. Transmission electron microscopy images the entire cross section, allowing exact chiral indices to be deduced [Fig. 1(c)]. Nanotubes on surfaces can be imaged by scanning tunneling microscopy [Fig. 1(d)], although because of the poor edge resolution, the precise chirality is usually undetermined. Both images confirm the atomic arrangement of Fig. 1(a), with the same atomic spacing $a_{CC} = 0.142$ nm as graphite.

Unfortunately, high-resolution microscopy is usually incompatible with transport measurements and the chiral indices of nanotubes in electronic devices are often unknown. A few experiments have combined transport measurements with structure determination by electron diffraction (Kociak

TABLE I. Summary of structure parameters for an (n, m) nanotube (Baskin and Meyer, 1955; Saito, Dresselhaus, and Dresselhaus, 1998). Vectors are written with respect to the graphene coordinates (x, y) defined in Fig. 1.

Name	Symbol	Value
C-C bond length	$a_{\rm CC}$	0.142 nm
Graphene lattice constant	а	$\sqrt{3}a_{\rm cc} = 0.246$ nm
Graphene basis vectors	$a_{1,2}$	$(\frac{\sqrt{3}}{2}, \pm \frac{1}{2})a$
Graphene reciprocal lattice vectors	${\bf b}_{1,2}$	$\left(\frac{1}{\sqrt{3}},\pm 1\right)\frac{2\pi}{a}$
Graphene Dirac points	\mathbf{K}, \mathbf{K}'	$\pm \frac{\mathbf{b}_2 - \mathbf{b}_1}{3} = (0, \mp 1) \frac{4\pi}{3a}$
Chiral vector	С	$n\mathbf{a}_1 + m\mathbf{a}_2$
		(m, n integer; n > 0;
		$-n/2 < m \le n$)
Chiral angle	θ	$\tan^{-1}\left(\frac{\sqrt{3}m}{2n+m}\right)$
-		$\left(-\frac{\pi}{6} < \theta \le \frac{\pi}{6}\right)$
Nanotube diameter	D	$a\sqrt{n^2+m^2+nm}/\pi$

et al., 2002; Allen *et al.*, 2011). The structure can also be determined using optical Raman or Rayleigh spectroscopy, which is less invasive but does not always give unambiguous chiral indices (Cao *et al.*, 2004; Huang *et al.*, 2005; Deshpande *et al.*, 2009). Most of the results in this review will therefore be from nanotubes of unknown chirality; however, as discussed in the next section, the electronic properties of nanotubes are sufficiently independent of the chiral indices that most of the underlying physics can still be explored.

B. Quantum dots

A basic carbon nanotube electronic device is shown in Fig. 2. The purpose is to allow measurement of the electrical current *I* through a single nanotube (Bockrath *et al.*, 1997; Tans *et al.*, 1997). To achieve this, the nanotube is contacted with metallic source and drain electrodes connected to an external circuit. A third electrode, the gate, coupled capacitively, allows the electrostatic potential to be tuned. Quantum dots are usually measured at low temperature (≤ 1 K) to suppress thermal smearing of transport features.

A nanotube naturally confines electrons to one dimension. In quantum transport experiments, it is common to add longitudinal confinement by introducing tunnel barriers. These barriers can be created by modifying the electrostatic potential using gate voltages, often taking advantage of Schottky barriers induced near the metal contacts in the nanotube (Heinze et al., 2002; Biercuk et al., 2008). The stretch of nanotube between the barriers where electrons are trapped is called a quantum dot. By studying the current through such a quantum dot as a function of bias, gate voltage, and other parameters such as magnetic field, the energy levels of electrons in the nanotube can be deduced. Quantum dot transport spectroscopy has been extensively reviewed by Kouwenhoven et al. (1997), Kouwenhoven, Austing, and Tarucha (2001), and Hanson et al. (2007). Basic concepts needed in this review are explained in Appendix A.

C. Fabrication challenges of gated quantum devices

The realization of clean and tunable quantum dots in carbon nanotubes is not straightforward. Unlike carriers in III-V

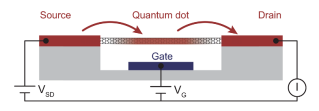


FIG. 2 (color online). Schematic of a basic quantum dot device. The device consists of a nanotube contacted by source and drain electrodes and capacitively coupled to a gate. Tunnel barriers to the source and drain, imposed through the combination of the gate potential and Schottky barriers, define a quantum dot. The current *I* through the device is measured as a function of bias voltage $V_{\rm SD}$ and gate voltage $V_{\rm G}$. Both the number of electrons *N* on the island and the dot energy levels can be adjusted by tuning $V_{\rm G}$.

heterostructures, which are separated from the crystal's surface by an atomically clean buffer layer, the nanotube's π band is composed of atomic *p* orbitals that stick out perpendicular to the surface (see Sec. III). Patterning of gate oxides, mechanical deformation, and contamination from fabrication chemicals can easily induce disorder and irreproducible device characteristics (Bezryadin *et al.*, 1998; Zhu, Brink, and McEuen, 2005).

The characteristics of nanotube quantum dots depend on the band gap, which varies widely between different nanotubes (Sec. III.B). Semiconducting nanotubes (band gap $\gtrsim 0.1 \text{ eV}$) often show poor transport characteristics at low carrier density and low temperature. Presumably, this arises from unintentional localization of carriers into disordered puddles, facilitated by the carriers' relatively large effective mass. Conversely, in quasimetallic nanotubes (band gap $\lesssim 10$ meV), controlled creation of sufficiently opaque barriers by electrostatic potentials is difficult, presumably due to the small effective mass (Sec. III.B.1). Kinks made by atomic force microscope (AFM) manipulation or mechanical templating can be used to locally induce band gaps and backscattering centers (Yao et al., 1999; Bozovic et al., 2001; Postma et al., 2001; Park et al., 2002; Biercuk et al., 2004; Stokes and Khondaker, 2008), resulting in addressable tunnel barriers and Coulomb blockade even at room temperature. A similar effect can occur unintentionally due to disordered mechanical deformations induced by fabrication (Bezryadin et al., 1998).

The largest experimental interest has been attracted by devices between these extremes (narrow-gap nanotubes). Tunable tunnel barriers can then be induced rather easily by electrostatic gates. These nanotubes allow gate-controlled devices that do not uncontrollably break up into disordered puddles, yet their tunnel barriers remain tunable over a wide range, even in the few-charge regime. Most devices can be classified according to whether gate fabrication occurs after nanotube growth (top gating), before growth (bottom gating), or on a separate chip (mechanical transfer method).

1. Top gating

The simplest way to make devices is usually to fabricate electrodes on top of nanotubes. This allows complex devices with many kinds of contact material including normal metals, ferromagnets, and superconductors. After growth or deposition, suitable nanotubes are imaged, and the electrodes are patterned by electron-beam lithography and liftoff. Early single-electron transistors were contacted in this way (Bockrath *et al.*, 1997), as were the first double quantum dots (Mason, Biercuk, and Marcus, 2004). Although cleanliness and fabrication-induced disorder are a concern, devices fabricated this way have demonstrated ambipolar operation and discrete excited states (Biercuk *et al.*, 2005), as well as charge sensing and pulsed gate spectroscopy (Biercuk *et al.*, 2006; Gotz *et al.*, 2008).

Full control of a double quantum dot requires at least five gate electrodes, necessitating thin, high-dielectric constant gate oxides (e.g., atomic-layer-deposited aluminum or hafnium oxide) and densely packed gate arrays (Churchill, Bestwick *et al.*, 2009; Churchill, Kuemmeth *et al.*, 2009).

Such a device is shown in Figs. 3(a) and 3(b), consisting of a fully tunable double quantum dot capacitively coupled via a floating gate to a charge-sensing single quantum dot on the same nanotube. Among other applications, these devices allow measurement of spin relaxation and dephasing (Sec. V). By selectively etching beneath the nanotube, suspended devices can also be fabricated (Leturcq *et al.*, 2009).

2. Bottom gating

A drawback of top gating is that the fabrication process itself can introduce disorder in the nanotube. An alternative is to grow or deposit nanotubes over predefined electrodes, resulting in devices with improved control and cleanliness (Cao, Wang, and Dai, 2005). Early single quantum dots were realized by depositing nanotubes across Pt source and drain electrodes, using the Si/SiO₂ substrate as a back gate (Tans *et al.*, 1997). Similar to graphene devices, where suspending the layer dramatically improved the mobility (Bolotin *et al.*, 2008; Du *et al.*, 2008), suspended nanotubes often show near-ideal transport characteristics, indicating that much of the disorder arises from interactions with the substrate (Steele, Gotz, and Kouwenhoven, 2009; Ilani and McEuen, 2010; Jung *et al.*, 2013).

Motivated by the results of suspended single quantum dots as in Fig. 3(d), more complex contact and gate arrays were developed that can be loaded into the nanotube growth furnace as the last step before cool down and measurements (Kuemmeth *et al.*, 2008; Steele, Gotz, and Kouwenhoven, 2009). Although these devices were of high quality² and resulted in new discoveries, the harsh conditions in the growth reactor greatly restrict the materials and design. The overall device yield is low because a nanotube must grow across contacts and gates by chance.

3. Mechanical transfer

Mechanical transfer attempts to benefit from the best of both approaches, achieving high gate tunability without postgrowth processing. The device chip (without nanotubes) and the growth chip (with nanotubes suspended across trenches) are fabricated separately. Just before measurement, a single nanotube is transferred from the growth chip to the device chip using an aligned stamping process (Wu, Liu, and Zhong, 2010; Pei *et al.*, 2012). By employing piezocontrolled scanning probe microscope manipulators, the transfer is possible in vacuum at cryogenic temperatures (Waissman *et al.*, 2013), allowing the cleanliness of the nanotube to be tested *in situ*. A state-of-the-art example is shown in Figs. 3(e) and 3(f).

D. Nanotube synthesis and isotopic engineering

For research applications nanotubes are readily synthesized in desktop furnaces, using chemical-vapor deposition (usually

²Nanotubes that have never been in contact with solvents, resists, or a substrate are sometimes called "ultraclean" (Deshpande *et al.*, 2009; Steele, Gotz, and Kouwenhoven, 2009; Pei *et al.*, 2012; Pecker *et al.*, 2013; Waissman *et al.*, 2013; Benyamini *et al.*, 2014).

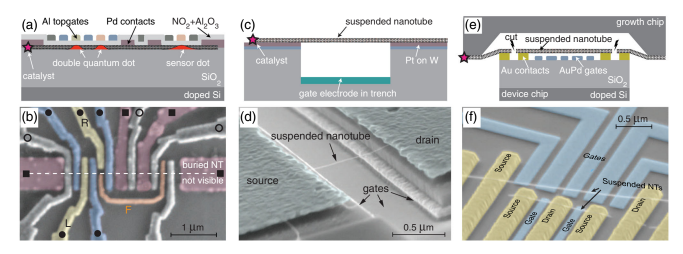


FIG. 3 (color online). (a), (c), and (e) Schematics and (b), (d), and (f) scanning electron micrographs of devices fabricated by different methods. (a), (b) Top gating: Nanotubes are located on a growth chip, and electrodes fabricated afterward. Here a nanotube (not visible) is contacted by metal electrodes (\blacksquare) and covered by a thin gate oxide. Five gates (•) control a double quantum dot, while a floating antenna (F) allows charge sensing via a separate dot on the same nanotube. Other electrodes (\bigcirc) are helper gates. Adapted from Churchill, Kuemmeth *et al.*, 2009. (c), (d) Bottom gating: Trench, contacts, and gates are fabricated from inert materials before synthesis, and nanotubes grown across. Adapted from Steele *et al.*, 2009. (e), (f) Mechanical transfer: Suspended nanotubes are synthesized on a growth chip, while electrodes are patterned on a device chip. By stamping the chips together, a nanotube is transferred to the device. Electrical current can be used to cut the nanotube at specific places. In this complex two-nanotube device, five gates define a single or double quantum dot in the upper nanotube, while a pair of dots in the lower nanotube serve as independent charge sensors. Adapted from Waissman *et al.*, 2013.

from methane, ethanol, or ethylene) in the presence of suitable catalysts (Kong *et al.*, 1999; Kuemmeth *et al.*, 2010). Unlike III-V devices such as GaAs double dots, in which all stable isotopes possess a nuclear magnetic moment, carbon nanotubes allow fabrication of devices with and without nuclear spins in the host material in a straightforward way.

Nanotubes synthesized from natural hydrocarbons consist of 99% ¹²C and ~1% ¹³C. By using isotopically purified ¹³CH₄ or ¹²CH₄, the isotopic composition can be tuned during growth. This not only affects the phonon modes [revealed by Raman spectroscopy (Liu and Fan, 2001)], but also the electron spin properties, because ¹³C possesses a nuclear spin $|\vec{I}| = 1/2$, while ¹²C has $|\vec{I}| = 0$. As discussed in Sec. IV.D, a local spin impurity (such as ¹³C) can flip the spin and/or valley of an electron.

III. CARBON NANOTUBE BAND STRUCTURE

Just as the atomic structure of carbon nanotubes can be derived from that of graphene, the electronic band structure inherits many of its properties from graphene. However, the simple effect of being rolled up drastically modifies the band structure, leading to many effects that are not present in graphene. The most dramatic difference is the introduction of a band gap, which allows electrons in nanotubes to be confined using gate voltages, but a variety of other subtle effects arise.

Briefly, the results are as follows. Although graphene is a semimetal, the formation of a nanotube leads to a confinement band gap (a few hundred meV) for two-thirds of the possible structures. These are known as semiconducting nanotubes. Most of the remaining nominally metallic nanotubes show narrow band gaps ($\sim 10 \text{ meV}$) due to a combination of

curvature and strain. If the band gap is undetectibly small, the nanotube is called quasimetallic, and a metallic nanotube is defined as one for which the band gap is exactly zero. More subtle details of the band structure become evident in a magnetic field, including a magnetic moment associated with the valley degree of freedom, and spin-orbit coupling that is much stronger than in graphene and arises from curvature.

A. From atomic carbon to graphene band structure

To understand nanotube band structure, we begin with the energy levels of atomic carbon. In a free atom, the six electrons occupy the configuration $1s^22s^22p^2$. The outermost atomic shell includes one spherically symmetric *s* orbital and three *p* orbitals p_x , p_y , and p_z [Fig. 4(a)]. Because of twofold spin degeneracy in each orbital, there are therefore eight states in the outermost shell, of which four are occupied.

The 2s - 2p energy splitting is small enough (less than a typical bond energy) that all four outermost orbitals can hybridize to form covalent bonds. For a given structure, the number of 2p orbitals that hybridize with the 2s orbital is determined by symmetry. In graphene, the p_z orbital, oriented perpendicular to the plane, is odd under z inversion and therefore cannot hybridize with the even-parity 2s orbital. No such symmetry protects the p_x and p_y orbitals. This type of hybridization, in which an s orbital is mixed with two p orbitals, is known as sp^2 hybridization.

In graphene, these three orbitals further hybridize across neighboring atoms in the crystal, forming a low-energy (bonding) band σ and a high-energy (antibonding) band σ^* [Fig. 4(b)]. Likewise, hybridization of the p_z orbitals forms bonding and antibonding bands denoted π and π^* , although with smaller bonding energy because the interatomic overlap

is less. In undoped graphene, the electrons exactly fill the bonding bands, with three electrons per atom occupying σ and one occupying π . The σ band remains filled at all times and does not participate in transport. The electrical behavior of nanotubes is therefore determined almost entirely by the properties of the π and π^* bands.

Ignoring spin-orbit coupling, the p_z orbitals do not hybridize with any of the lower-lying states, so the structure of the π and π^* bands follows simply from energy levels in the honeycomb graphene potential. Graphene consists of a rhombus unit cell with a two-atom basis [Fig. 4(c)] and has the hexagonal Brillouin zone shown in Fig. 4(d). The corners of this hexagon in k space are alternately labeled K or K'. Because the three K points are connected by reciprocal lattice vectors, by Bloch's theorem they correspond to equivalent electron states; likewise, the three K' points are equivalent to each other, but not to the K points. States close to the K' point are time-reversal conjugates of those close to the K point.

The band structure that arises from this potential [Fig. 4(e)] has quite unusual properties (Wallace, 1947; Saito, Dresselhaus, and Dresselhaus, 1998; Castro Neto *et al.*, 2009). Although there is no band gap, the π and π^* bands

(a) (b) Free Hybridized 10 π^* σ Energy (eV) vacuum E 2p_x,2p_v,2p_z 2s -10 (c) (e) В π (conduction) (d) band C E (valence) band кo k,

FIG. 4 (color online). (a) Electron orbitals of atomic carbon. Lighter (darker) colors denote regions where the *p*-orbital wave functions are positive (negative). Bond directions in graphene are indicated by gray lines. (b) Schematic energy levels of atomic (left) and sp^2 hybridized (right) carbon. Energies are referenced to E_F , approximated as equal to the negative of the work function. (c) Segment of graphene with the unit cell shaded and the *A* and *B* sublattices marked. (d) First Brillouin zone of graphene in reciprocal space, showing the six symmetry points, labeled *K* or *K'*. (e) Energy bands (σ bands omitted) of graphene close to the Fermi level, showing the six Dirac cones where π and π^* bands touch.

touch only at *K* and *K'*, where the density of states is zero. Since the available electrons exactly fill the π band, these points are where the Fermi level $E_{\rm F}$ intersects the band structure, so that undoped graphene is neither a true metal nor a true semiconductor, but a semimetal. Close to the Fermi surface, the dispersion relation is linear, with a slope that determines the Fermi velocity³ $v_{\rm F} = (1/\hbar) |\nabla_{\bf k} E| \approx$ $8 \times 10^5 \text{ ms}^{-1}$. Expanding about the *K* or *K'* points by writing $\mathbf{k} = \mathbf{K} + \boldsymbol{\kappa}$ or $\mathbf{k} = \mathbf{K}' + \boldsymbol{\kappa}$, and defining $E_{\rm F}$ as the zero of energy, the dispersion relation for $|\boldsymbol{\kappa}| \ll |\mathbf{K}|$ is simply

$$E = \pm \hbar v_{\rm F} |\boldsymbol{\kappa}|,\tag{1}$$

where the + sign applies to electrons and the - to holes. Because this dispersion relation also describes massless Dirac fermions, the points where the bands touch are known as Dirac points, and the nearby bands as Dirac cones. The correspondence of electron states in a nanotube with solutions of a Dirac-like equation is explained in Appendix B. Although we use this correspondence only in a few places in this review, it is theoretically convenient because it allows many effects on nanotube band structure to be derived as perturbations to the Dirac equation.

B. Semiconducting, narrow-gap and metallic nanotubes

1. Theory: The zone-folding approximation

Since the nanotube diameter is usually much larger than the interatomic spacing, the graphene band structure is to a good approximation unperturbed by rolling up into a nanotube except for the imposition of a periodic boundary condition (Hamada, Sawada, and Oshiyama, 1992; Saito, Fujita, and Dresselhaus, 1992). This is known as the "zone-folding approximation." The boundary condition to ensure single valuedness is that $\mathbf{k} \cdot \mathbf{C} = 2\pi p$, where p is an integer, i.e., the component of \mathbf{k} perpendicular to the nanotube axis is $k_c = 2p/D$. The allowed \mathbf{k} values correspond to a series of lines in reciprocal space, known as quantization lines, running at an angle $\pi/3 + \theta$ from the k_x axis [Figs. 5(a) and 5(b)].

The one-dimensional dispersion relation $E(\kappa_{\parallel})$ is a cut along the quantization lines of the two-dimensional graphene dispersion relation. Since it is the branches closest to $E_{\rm F}$ that determine transport properties, we neglect the other branches. The nanotube band gap depends on the minimum separation of the quantization lines from the Dirac points. There are two possible situations. If quantization lines run straight through the Dirac points [Figs. 5(a) and 5(c)], then $E(\kappa_{\parallel})$ is linear near $\kappa_{\parallel} = 0$, giving zero band gap and a metallic nanotube. However, if the lines bypass the Dirac points with separation

³This value is derived from numerical simulations of graphite (Painter and Ellis, 1970; Tatar and Rabii, 1982; Trickey, Müller-Plathe, and Diercksen, 1992) and nanotubes (Mintmire, Dunlap, and White, 1992), which indicate $v_{\rm F} = (7.8 - 9.8) \times 10^5 \text{ ms}^{-1}$ [although interactions may renormalize the value significantly (Kane and Mele, 2004)], as well as nanotube scanning tunneling microscopy (STM) density-of-state measurements (Odom *et al.*, 1998; Wildoer *et al.*, 1998) and ballistic electron resonance experiments (Zhong *et al.*, 2008), which give $v_{\rm F} = (7.9 - 8.7) \times 10^5 \text{ ms}^{-1}$.

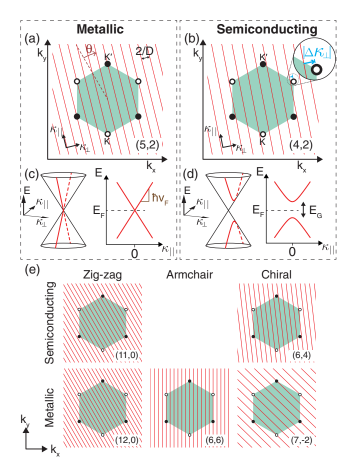


FIG. 5 (color online). The effect of periodic boundary conditions. (a), (b) Requiring that the electron wave function be single valued constrains k to lie on one of the quantization lines in reciprocal space corresponding to integer values of $k_c D/2$. If quantization lines intersect the Dirac points the nanotube is (a) metallic; otherwise it is (b) semiconducting, with minimum quantization line offset $|\Delta \kappa_{\perp}| = 2/3D$. The $\{\kappa_{\perp}, \kappa_{\parallel}\}$ axes in reciprocal space, corresponding to motion around or along the nanotube, are indicated. (c), (d) Dispersion relations (in the lowest-energy one-dimensional band) close to a Dirac point for the two kinds of nanotubes, showing how the offset gives rise to a band gap. The Fermi level for undoped nanotubes is indicated. (e) Examples of quantization lines for several metallic and semiconducting structures of the three types shown in Fig. 1(b). Of the six combinations, armchair semiconducting nanotubes do not exist.

 $|\Delta \kappa_{\perp}|$, the situation is as shown in Figs. 5(b) and 5(d). The dispersion relation gives a pair of hyperbolae with band gap $E_{\rm G} = 2\hbar v_{\rm F} |\Delta \kappa_{\perp}|$, and therefore a semiconducting nanotube. In the zone-folding approximation, the band gap is determined by a simple rule: If n - m is a multiple of 3, the nanotube is nominally metallic (Hamada, Sawada, and Oshiyama, 1992; Saito, Fujita, and Dresselhaus, 1992). Otherwise, it is semiconducting, with band gap $E_{\rm G} = 4\hbar v_{\rm F}/3D \approx 700 \text{ meV}/D$ [nm]. In a collection of nanotubes will therefore outnumber metallic ones by approximately 2:1. Figure 5(e) illustrates how the chiral indices determine whether the quantization lines intersect the Dirac points for various nanotube structures. Examples of both cases are

shown for the three kinds of structures defined in Fig. 1(b), with one exception: Zigzag and chiral tubes can be either semiconducting or metallic, but all armchair nanotubes are metallic.

The nanotube structure also sets the electron dispersion relation and hence the effective mass. The equation of the hyperbola in Fig. 5(d) is (Zhou *et al.*, 2005)

$$E^{\pm}(\kappa_{\parallel}) = \pm \sqrt{\hbar^2 v_{\rm F}^2 \kappa_{\parallel}^2 + E_{\rm G}^2/4}.$$
 (2)

This low-energy dispersion relation is clearly electron-hole symmetric. This is a fragile symmetry, because any charge in the environment oppositely couples to electrons and holes, but it is sometimes reflected in data (Jarillo-Herrero *et al.*, 2004).

The effective mass arises from the curvature of the dispersion relation and for low energy $(|E^{\pm}(\kappa_{\parallel})| \ll E_{\rm G})$ is

$$m_{\rm eff} = \hbar^2 \left(\frac{d^2 E}{d\kappa_{\parallel}^2}\right)^{-1} \approx \frac{E_{\rm G}}{7.3 \text{ eV}} \times m_{\rm e},\tag{3}$$

where m_e is the free electron mass. A band gap of 100 meV corresponds to effective mass ~0.014 m_e , smaller than that in many conventional semiconductors (e.g., in GaAs $m_{eff} = 0.067m_e$). Because small m_{eff} leads to larger longitudinal level spacing, nanotubes with small E_G are often preferred for quantum dot experiments.

2. Valley as a robust quantum number

Just as in graphene, the band structure in nanotubes is characterized by the distinct and time-conjugate valleys K and K'. In graphene the robustness of the valley quantum number is linked to the symmetries of the lattice. Mixing between valleys requires a large transfer of crystal momentum and is therefore weak in a smoothly varying Coulomb potential. This is less obvious in metallic nanotubes, because the two Dirac points sometimes remain well separated in momentum space, and sometimes they merge at $k_t = 0$. In fact, all metallic nanotubes (see Fig. 6) can be divided into two classes (Samsonidze et al., 2003; Marganska et al., 2014): the Dirac points are either well separated in longitudinal momentum space (such nanotubes are known as armchairlike metals) or collapse to the origin of the longitudinal Brillouin zone (zigzaglike metals). For chiral metallic nanotubes, this classification is possible by introducing a helical translational basis vector (Lunde, Flensberg, and Jauho, 2005). For the zigzaglike metals, the two bands at k = 0 are distinct by having different crystal angular momentum (Lunde, Flensberg, and Jauho, 2005), where the angular momentum is defined as the quantum number related to the rotation part of the helical symmetry (White, Robertson, and Mintmire, 1993). In the armchairlike metals, the angular momenta are the same, but their longitudinal crystal momenta differ by $4\pi/3|T|$. Consequently, in both cases valley-valley scattering is suppressed by a difference in crystal angular momenta or crystal longitudinal momenta. Scattering within a valley may also require atomically sharp Coulomb scatterers or lattice imperfections, due to the spinor structure of the solutions to the Dirac equation, which differs between right movers and left movers (Ando and Nakanishi, 1998; Ando, Nakanishi,

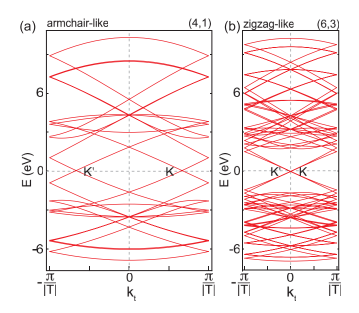


FIG. 6 (color online). Robustness of the vallev index in nanotubes. All subbands of the 1D dispersion relation (corresponding to different quantization lines in Fig. 5) are plotted in the first longitudinal Brillouin zone vs longitudinal wave vector k_t . All nominally metallic nanotubes can be classified as armchairlike or zigzaglike. For armchairlike nanotubes, the two Dirac points are separated in k_t ; for zigzaglike nanotubes, they are separated in crystal angular momentum. Since all metallic nanotubes fall into one of these classes, valley is a good quantum number in a slowly varying potential. (a) Armchairlike (4,1) nanotube. (b) Zigzaglike (6,3) nanotube. Each spin-degenerate band is calculated using a graphene tight-binding model taking account of nearest-neighbor overlap integrals but without spin-orbit coupling. Only states near E = 0 participate in transport. Note that the Brillouin zone in (a) has been plotted wider than in (b), to reflect the different longitudinal length |T| of the unit cell in real space.

and Saito, 1998; McEuen et al., 1999; Roche, Triozon, and Rubio, 2001).

Armchairlike and zigzaglike band structures are exemplified in Fig. 6. The number of subbands equals the number of carbon atoms in the unit cell of the nanotube, spanned by Cand T in Fig. 1(a). Each subband shown is twofold degenerate due to spin and arises from a mapping of the quantization lines in Fig. 5 into the 1D Brillouin zone of the nanotube.

The above discussion applies to narrow-gap nanotubes, i.e., those that would be metallic in the zone-folding approximation but where other perturbations introduce a small band gap (Sec. III.C). For nanotubes that are semiconducting even in the zone-folding approximation (White, Robertson, and Mintmire, 1993; Mintmire and White, 1995), the situation is similar. This can again be seen using helical quantum numbers (White, Robertson, and Mintmire, 1993; Mintmire and White, 1995) as follows: All bands can be classified by their crystal angular momentum, which means that mixing of two bands with different crystal angular momentum is protected (as for zigzaglike metallic tubes). Mixing of bands with the same crystal angular momentum is suppressed by their difference in wave numbers when folded onto the smaller translational Brillouin zone.

Local Coulomb scatterers can flip the valley index (Pályi and Burkard, 2010; Bercioux *et al.*, 2011), and spin-carrying impurities can flip both spin and valley with comparable rates (Pályi and Burkard, 2009). One example is hyperfine coupling to nuclear ¹³C spins, which can cause both spin and valley relaxation (Sec. IV.D). Another example is the local part of the electron-electron interaction, discussed in Sec. VII and Appendix B. In addition, electrical contacts can induce valley scattering due to valley mixing during tunneling (Sec. VI.C).

3. Experiment

Nanotubes of different kinds can be distinguished experimentally by measuring the current as a function of $V_{\rm G}$ at fixed $V_{\rm SD}$, as in Figs. 7(a)–7(c). The gate potential shifts the energy levels up or down and therefore tunes the position of the gap relative to $E_{\rm F}$. Tuning $E_{\rm F}$ into the band gap suppresses the current. This can be seen in Fig. 7(a), where the Fermi level is shifted from the valence band (for $V_{\rm G} \leq 0$) to the band gap (for $V_{\rm G} \gtrsim 0$), showing that the nanotube is semiconducting. A quasimetallic nanotube, by contrast, is one with no dependence on $V_{\rm G}$ [Fig. 7(c)], indicating $E_{\rm G} \ll k_{\rm B}T$, where $k_{\rm B}$ is Boltzmann's constant and T is temperature.

Experimentally, the fraction of nanotubes showing quasimetallic behavior at room temperature is very small ($\leq 1\%$) (Cao, Wang, and Dai, 2005; Churchill, 2012). More common

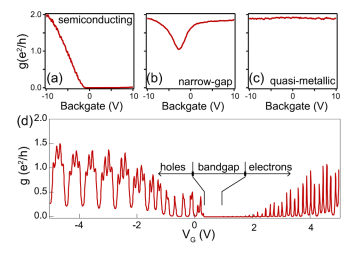


FIG. 7 (color online). Signatures of the band gap in transport. (a)-(c) Room temperature conductance measurements as a function of gate voltage. (a) A semiconducting nanotube has $E_{\rm G} \gg k_{\rm B}T$ and can be tuned between a conducting state (Fermi level in the valence band) and an insulating state (Fermi level in the band gap). Transport via the conduction band is not observed, because it would require a much higher gate voltage. (b) A narrow-gap nanotube $(E_{\rm G} \sim k_{\rm B}T)$ shows transport via both conduction and valence bands. Tuning the Fermi level into the band gap does not completely suppress current at room temperature. (c) A few nanotube devices (< 1%) show no gate dependence of conduction. These could be truly metallic, although it cannot be excluded that this device in fact contains a bundle of nanotubes that screens the gate. (d) Conductance of a single narrow-gap nanotube at 300 mK. Transport is now completely suppressed in the band gap, and the device can be tuned into electron or hole configurations by tuning $V_{\rm G}$. Adapted from Cao, Wang, and Dai, 2005 and Churchill, 2012.

is "narrow-gap" behavior [Fig. 7(b)], where partial current suppression indicates a small band gap $E_{\rm G} \sim k_{\rm B}T$ at room temperature (Ouyang *et al.*, 2001). This interpretation is confirmed by low-temperature experiments [Fig. 7(d)], where precise measurements from Coulomb peak positions frequently give $E_{\rm G} \sim 10 - 100$ meV [e.g., in Fig. 7(d), $E_{\rm G} = 60$ meV]. To explain such small band gaps from circumferential quantization alone requires D = 7 - 70 nm, which might be structurally unstable and is excluded by AFM topography measurements. More likely, nearly all nanotubes that the zone-folding model predicts should be metallic acquire narrow band gaps by perturbations discussed in the next section.

C. Structural origins of the narrow gap

1. Theory

The zone-folding approximation assumes that the allowed electron states in nanotubes are exactly the same as their equivalents in graphene. Perturbations arise if the symmetry of the carbon bonds is broken by changing the overlap between adjacent electron orbitals. One unavoidable example is the curvature of the rolled-up sheet (Blase et al., 1994). This has two effects on the band structure. First, it leads to a small renormalization of the Fermi velocity by at most a few percent, which is insignificant in experiments (Izumida, Sato, and Saito, 2009). More importantly, it displaces the Dirac points in reciprocal space away from K and K' (Kane and Mele, 1997; Izumida, Sato, and Saito, 2009), because it breaks the equivalence of the three bond directions. This shift is parametrized by a displacement vector $\Delta \kappa^{cv}$ [Fig. 8(a)] and is opposite for K and K' because states in the two valleys are time-reversed conjugates of each other (Castro Neto et al., 2009). In semiconducting nanotubes, $|\Delta \kappa^{cv}|$ is much smaller than the offset $|\Delta \kappa_{\perp}|$ arising from quantization and therefore has only a small effect. However, in nominally metallic nanotubes, the shift of the Dirac cones relative to the quantization lines introduces a band gap $E_{\rm G} = 2\hbar v_{\rm F} |\Delta \kappa_{\perp}^{\rm cv}|$, where $\Delta \kappa_{\perp}^{cv}$ is the component of $\Delta \kappa^{cv}$ perpendicular to the nanotube axis [Fig. 8(b)]. (The component parallel to the axis $\Delta \kappa_{\parallel}^{cv}$ has no effect.) This curvature-induced band gap is always much smaller than the quantization energy difference. Unlike the quantization band gap, it depends on the chiral angle. As shown in the inset of Fig. 8(a), the vector $\Delta \kappa^{cv}$ points at an angle of 3θ from the perpendicular. The curvatureinduced band gap is therefore proportional to $\cos 3\theta$; it is calculated to be (Kane and Mele, 1997; Park, Kim, and Chang, 1999; Yang et al., 1999; Yang and Han, 2000; Kleiner and Eggert, 2001; Izumida, Sato, and Saito, 2009; Klinovaja et al., 2011a)

$$E_{\rm G}^{\rm cv} \sim \frac{50 \text{ meV}}{D[\rm nm]^2} \cos 3\theta. \tag{4}$$

For armchair nanotubes $\theta = \pi/6$ [Fig. 1(b)] and therefore $\cos 3\theta = 0$. These are the only nanotubes expected to be truly metallic, if no other perturbation is applied (see Appendix B for more theoretical details).

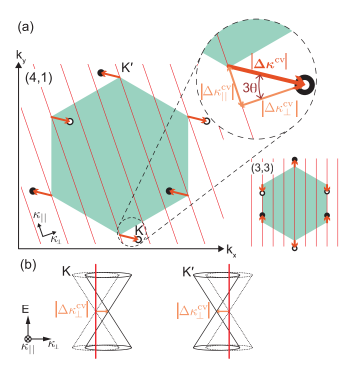


FIG. 8 (color online). Perturbation of the graphene band structure by the curvature in nanotubes. (a) Displacement of the Dirac points away from the corners of the Brillouin zone due to curvature in a (4,1) nanotube. For visibility, the shift has been exaggerated by a factor of 15. Top inset: Decomposition of the displacement vector $\Delta \kappa^{cv}$ near the *K* point into components parallel and perpendicular to the nanotube axis. The shift is at an angle 3θ to the nanotube circumference. Bottom inset: Shift for an armchair nanotube. Because the shift is along the quantization lines, curvature does not lead to a gap in these structures. (b) Dirac cones close to *K* and *K'* valleys with (solid) and without (dotted) curvature effects, showing how horizontal shifts by $\Delta \kappa_{1}^{cv}$ open a band gap in a nominally metallic tube.

A gap of similar magnitude can be opened by strain, in a way that also depends on nanotube chirality (Heyd, Charlier, and McRae, 1997; Kane and Mele, 1997; Rochefort, Salahub, and Avouris, 1998; Yang and Han, 2000). A uniaxial strain ϵ has the same symmetry-breaking effect as curvature (namely, to break C_3), and therefore leads to a Dirac point displacement in the same direction with magnitude

$$|\Delta \kappa^{\epsilon}| = \frac{12\zeta}{1+6\zeta} (1+\lambda)\epsilon/a_{\rm CC},\tag{5}$$

where $\lambda \approx 0.2$ is the Poisson ratio and $\zeta \approx 0.066$ is a parameter related to the carbon-carbon bond force constants (Nisoli *et al.*, 2007; Huang *et al.*, 2008). A torsional strain γ displaces the Dirac points by an amount $|\Delta \kappa^{\gamma}| \approx \gamma$ at an angle $\pi/2 - 3\theta$ from the perpendicular (Yang and Han, 2000). The uniaxial band gap is therefore proportional to $\cos 3\theta$, while the torsional band gap is proportional to $\sin 3\theta$. A third type of strain, nanotube bending, has no first-order effect on the band gap for any structure (Kane and Mele, 1997). Numerical estimates of these effects are given in Table III.

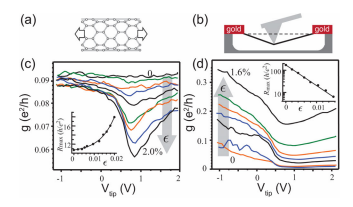


FIG. 9 (color online). Contribution of strain to the band gap. (a) Distortion of the lattice by tensile strain. (b) The experimental setup. An AFM tip is used simultaneously to gate and tension a suspended nanotube. (c) Conductance of an initially metallic nanotube as a function of gate voltage measured for several values of strain ϵ , showing an increasing band gap with ϵ . Inset: Maximum resistance as a function of ϵ , fitted assuming thermally activated conductance across a band gap proportional to ϵ . The fit yields the strain sensitivity $dE_G/d\epsilon = 35 \text{ meV}/\%$, where ϵ is expressed as a percentage. (d) Similar data for a semiconducting nanotube. In this case, the band gap is found to decrease linearly with ϵ , with fitted $dE_G/d\epsilon = -53 \text{ meV}/\%$. Adapted from Minot *et al.*, 2003.

2. Experiment

The existence of a narrow gap in nominally metallic nanotubes was first shown in density-of-states measurements by scanning tunneling microscopy (Ouyang *et al.*, 2001). For zigzag nanotubes, a diameter dependence $E_G =$ 39 meV/D (nm)² was found in close agreement with theoretical expectations. Isolated armchair nanotubes showed no band gap, consistent with the expected cos 3θ dependence (and implying negligible torsion for nanotubes lying on a surface). Many transport experiments have since found band gaps of this order of magnitude, although usually without identifying the chirality.

In transport measurements, quasimetallic nanotubes typically show band gaps a few times larger than expected from curvature alone, suggesting a significant contribution from strain. The sensitivity of the band gap to uniaxial strain has been measured by using an AFM tip to apply tension to suspended nanotubes (Minot et al., 2003). By varying the applied force, it was possible both to induce a band gap where none had been present before and to decrease the band gap in a semiconducting nanotube (Fig. 9). From the variation of conductance with strain, it was possible to deduce $dE_{\rm G}/d\epsilon =$ +35 meV/% for the metallic and $dE_G/d\epsilon = -53$ meV/% for the semiconducting nanotube, both with unknown chirality, where ϵ is expressed as a percentage elongation of the nanotube. Both values are comparable with that expected from Eq. (5), $dE_G/d\epsilon = 51 \text{ meV}/\% \times \cos 3\theta$. Similar results, including confirmation of the $\cos 3\theta$ dependence, have been obtained by optical methods (Huang et al., 2008).

D. Longitudinal confinement and quantum dot energy shells

Different modes of the longitudinal wave function in a quantum dot of length L lead to different confinement

energies $E_{\rm conf}$. The mode spectrum can be complex, depending on the band gap, boundary conditions, and interactions. The confinement can be classified as atomically sharp or nonsharp (McCann and Fal'ko, 2004), with the latter further subdivided into hard-wall or soft-wall cases depending whether the potential rises over a shorter or a longer distance than the dot length. An additional complication arises from the fact that bound states are formed from right- and left-moving Dirac particles that do not necessarily have the same group velocity in the unconfined nanotube. In Fig. 6 right movers and left movers within a valley travel at different speeds. This effect arises away from $E_{\rm F}$ due to trigonal warping of flat graphene. If curvature of the nanotube is also taken into account (not shown in Fig. 6), a stronger asymmetry arises. This happens already at $E_{\rm F}$ and directly affects how standing waves are constructed. For example, if the confinement is sharp, then the discrete eigenstates of the quantum dot will be superpositions involving both valleys (Izumida, Vikström, and Saito, 2012). We mention two simple limiting cases [Fig. 10(a)]. Electrons (or holes) with low enough energy sample only the region near the potential minimum where confinement is parabolic ("soft-wall confinement"). If the energy is also much less than $E_{\rm G}$, so that by Eq. (3) the

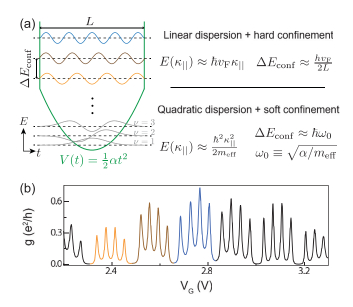


FIG. 10 (color online). (a) Schematic energy levels in the same quantum dot potential for two limits of dispersion and confinement discussed in the text. (Electron correlations beyond the constant-interaction model, which change the potential depending on occupation, are not taken into account.) Superimposed on the level diagram are wave function envelopes $\psi(t)$ for several electron shells. Mode index ν is indicated for the first three shells. (b) Sequence of Coulomb blockade peaks measured in a quantum dot of length $L \approx 760$ nm. Peaks corresponding to three successive shells are colored to illustrate the connection to different wave functions. Although the four electrons that fill each shell occupy states of similar energy, extra energy ΔE is needed to populate a higher shell, leading to fourfold periodic peak spacing. (The connection between colored shells and particular longitudinal modes is only schematic, because absolute shell numbers cannot be deduced from these data.) Adapted from Sapmaz, Jarillo-Herrero, Kong et al., 2005.

electron behaves as a massive particle (e.g., in a sufficiently large few-electron dot), the spectrum is harmonic with mode spacing $\Delta E_{\text{conf}} = \hbar \omega_0$, where ω_0 is the harmonic frequency. Conversely, in a many-electron-quantum dot the kinetic energy may both be large enough to reach the hard walls of the potential well and be in the linear part of the dispersion relation (2), so that the velocity is $v_{\rm F}$, independent of energy. The longitudinal modes then take on a sinusoidal form [Fig. 10(a)]. The modes are again regularly spaced in energy, but now with $\Delta E_{\rm conf} = h v_{\rm F}/2L$ (Tans *et al.*, 1997). This regular spacing, first observed by Liang, Bockrath, and Park (2002), suggests that this picture is accurate in at least some real devices. If neither of these cases applies, or if the potential is strongly disordered, the mode spacing need not be regular. For example, when the electrons behave as massive particles in a hard-wall potential, the confinement energy is given by $E_{\rm conf} = \nu^2 h^2 / \hat{8m}_{\rm eff} L^2$, with $\nu = 1, 2, 3, \dots$

A set of states with the same mode index ν is called a shell. As explained in the next section, each state is characterized by twofold spin and valley quantum numbers, and thus the number of single-particle states per shell is four. In the socalled constant-interaction model (Appendix A), the quantum dot states are filled in order of increasing energy, so that $\Delta E_{\rm conf}$ contributes to the Coulomb peak spacing only for every fourth electron. This is evident in the ground-state spectroscopy data of Fig. 10(b). The regular shell spacing shows that single-particle energy levels, in combination with the constant-interaction model, are a good approximation.

E. Orbital magnetic moment

Because each state in the K valley has a time-reversed conjugate in the K' valley, time-invariant perturbations such as curvature and strain do not break the degeneracy between them. This degeneracy can, however, be broken by a magnetic field. Intuitively, this can be seen by associating each state in the nanotube with a direction of circulation and hence a valley-dependent magnetic moment. This section shows how this orbital magnetic moment arises and is evident in the energy levels.

1. Theory

The orbital effect of a magnetic field **B** on an electron with charge -e is captured by modifying the bare-electron Schrödinger equation so that the momentum operator **p** is replaced by $\mathbf{p} + e\mathbf{A}$, where $\mathbf{A}(\mathbf{r})$ is the vector potential and $\mathbf{B} = \nabla \times \mathbf{A}$ (Merzbacher, 1998). So long as $\mathbf{A}(\mathbf{r})$ varies slowly on the scale of the lattice potential, the effect of this replacement on an electron confined in a closed loop is to add an Aharonov-Bohm phase to its eigenfunctions: if $\psi_0(\mathbf{r})$ is an eigenstate at $\mathbf{A} = 0$, then

$$\psi_{\mathbf{A}} = \exp\left(ie\frac{\mathbf{A}(\mathbf{r})\cdot\mathbf{r}}{\hbar}\right)\psi_{0} \tag{6}$$

is an eigenstate at finite **A** with the same energy (Luttinger, 1951; Hofstadter, 1976). In other words, the finite-field dispersion relation $E_{\mathbf{A}}(\boldsymbol{\kappa})$ is related to the zero-field dispersion relation $E_0(\boldsymbol{\kappa})$ by

Here the field-induced shift is $\Delta \kappa_{\perp}^{B} = \Delta \kappa_{\perp}^{B} \hat{\mathbf{C}}$, with (Ajiki and Ando, 1993; Lu, 1995)

$$\Delta \kappa_{\perp}^{B} = \frac{eA}{\hbar} = \frac{eD}{4\hbar} B_{\parallel}, \qquad (8)$$

where B_{\parallel} is the component of **B** along the nanotube axis **T**, *A* is the corresponding component of **A**, and $\hat{\mathbf{C}}$ is the direction of the chiral vector. The quantization condition, however, is unchanged.

The consequences for the band structure are shown in Fig. 11. For a true metallic nanotube [Figs. 11(a) and 11(b)], the Dirac cones are shifted horizontally away from the quantization lines, opening a band gap $E_G^B = 2\hbar v_F |\Delta \kappa_{\perp}^B| =$ $v_{\rm F}eD|B_{\parallel}|/2$. If the nanotube already has a band gap, the effect of the magnetic field is opposite for the two valleys [Figs. 11(c) and 11(d)]. In one valley, the electron energy is initially reduced by $v_{\rm F}eDB_{\parallel}/2$; in the other, it is increased by the same amount. At a field $B_{\parallel} = B_{\text{Dirac}} = E_{\text{G}}/ev_{\text{F}}D$, one of the Dirac cones crosses a quantization line and the band gap vanishes. Increasing B_{\parallel} beyond B_{Dirac} causes the band gap to increase again. For a true semiconducting nanotube, B_{Dirac} can be as large as ~ 100 T and is usually outside the experimental range, but for quasimetallic nanotubes B_{Dirac} can be just a few tesla. Because the effective mass depends on a band gap [Eq. (3)], $m_{\rm eff}$ can be tuned by a magnetic field.

The ground-state energies are plotted in Figs. 11(e)–11(g) as a function of magnetic field. Each zero-field level is twofold split, with slopes $dE/dB_{\parallel} = \pm Dev_{\rm F}/4$. This linear splitting allows each state to be assigned a magnetic moment $\pm \mu_{\rm orb}$, which has a straightforward physical interpretation [Fig. 11(i)]: Electron states with positive (negative) magnetic moment correspond to clockwise (counterclockwise) circulation of electrons around the nanotube. In this interpretation, the direction of circulation for the first electron switches as the field is swept through $B_{\rm Dirac}$. A similar picture applies in the valence band. The orbital moment is related to the band structure by

$$\mu_{\rm orb} \equiv \frac{dE}{dB_{\parallel}} = \frac{eD}{4\hbar} \left| \frac{\partial E^{\rm Dirac}(\kappa_{\perp}, \kappa_{\parallel})}{\partial \kappa_{\perp}} \right|,\tag{9}$$

where $E^{\text{Dirac}}(\kappa_{\perp},\kappa_{\parallel}) = \pm \hbar v_{\text{F}} \sqrt{\kappa_{\perp}^2 + \kappa_{\parallel}^2}$ is the two-dimensional energy function describing the Dirac cone. For a low-energy electron ($\kappa_{\parallel} \approx 0$), this has the value

$$\mu_{\rm orb}^0 = Dev_{\rm F}/4. \tag{10}$$

To emphasize the analogy with Zeeman spin splitting, an orbital *g* factor $g_{\text{orb}} \equiv \mu_{\text{orb}}/\mu_{\text{B}}$ is sometimes defined, where μ_{B} is the Bohr magneton. For a nanotube with a band gap, the magnetic energy in a parallel field $B_{\parallel} < B_{\text{Dirac}}$ is then

$$E_{\rm mag} = (\mp g_{\rm orb} \tau + \frac{1}{2} g_{\rm s} s) \mu_{\rm B} B_{\parallel}, \qquad (11)$$

where $g_s \approx 2$ is the spin g factor, the -(+) sign applies for electrons (holes), and the valley and spin quantum numbers are denoted by $\tau = \{+1, -1\}$ for $\{K, K'\}$ and $s = \{+1, -1\}$

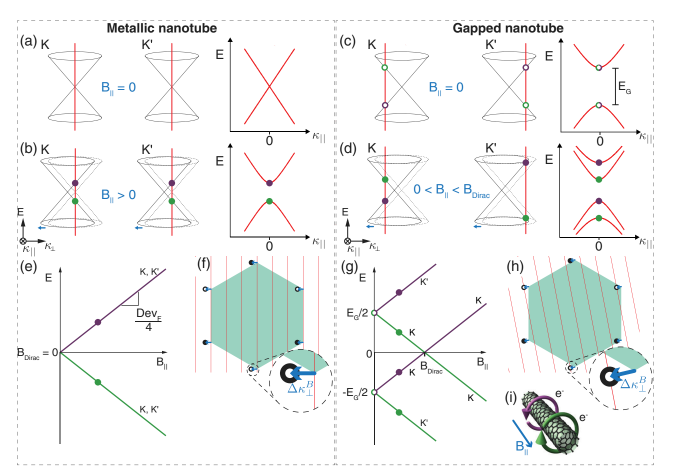


FIG. 11 (color online). Dependence of the band structure on a parallel magnetic field. (a), (b) Left: Dirac cones and quantization lines for a nanotube that is metallic at zero field, (a) without and (b) with magnetic field. Arrows in (b) mark the shift from zerofield (dotted) to finite-field (solid) Dirac cones. A field-induced horizontal shift $\Delta \kappa_{\perp}^{B}$ opens a band gap between the conduction band (darker circles) and valence band (lighter circles). Right: Corresponding one-dimensional electron dispersion relations. (c), (d) The same plots for a nanotube with zero-field gap $E_{\rm G}$. A magnetic field shifts one Dirac point toward the quantization line and one away, lifting valley degeneracy. (e) Conduction-band (darker line) and valence-band (lighter line) edges as a function of magnetic field for a metallic nanotube. In the conduction band both valleys increase in energy with field, corresponding to a negative magnetic moment $\mu_{\rm orb} = -Dev_{\rm F}/4$ or counterclockwise circulation. The valence band decreases in energy, corresponding to a positive magnetic moment and clockwise circulation. (f) Shift of the Dirac points perpendicular to the quantization lines for a zerogap nanotube. (g) Band edges for a gapped nanotube. In the conduction band, K(K') states move with positive (negative) magnetic moments. In the valence band, K'(K) states move with positive (negative) magnetic moments. (h) Shift of the Dirac points for $B_{\parallel} = B_{\rm Dirac}$, showing how one set of Dirac points is shifted onto the quantization lines. (i) Electron circulation directions corresponding to upmoving and downmoving states in (e) and (g).

for $\{\uparrow, \downarrow\}$, with the spin axis, along *t*, being parallel to the nanotube.⁴

2. Experiment

The orbital energy splitting can be seen in Coulomb ground-state spectroscopy as a function of magnetic field (Fig. 12), which shows the contribution E_{mag} to the single-particle energy levels E_{N} (Minot *et al.*, 2004). Ignoring spinorbit coupling (to be discussed in Sec. III.F), the first four electrons fill the four lowest states in order of energy: $K \downarrow$, $K \uparrow$, $K' \downarrow$, and $K' \uparrow$. Subsequent electrons must enter a higher shell of the dot, but repeat the fourfold filling sequence for spinvalley states. The expected pattern of ground-state energies is therefore alternating pairs with positive and negative magnetic moments. Typical data are shown in Fig. 12 for the first three hole shells. The measured magnetic moments $\mu_{\rm orb} \sim$ 0.9 meV/T are of the expected magnitude for orbital coupling with $D \approx 4.5$ nm. In terms of orbital g factors, this corresponds to $g_{\rm orb} \approx 16$, much larger than $g_{\rm s} = 2$, qualitatively confirming the picture in the previous section. For a quantitative comparison with theory, an independent measurement of D is necessary. This was achieved using an AFM for a nanotube with $D = 2.6 \pm 0.3$ nm, for which $\mu_{\rm orb} =$ 0.7 ± 0.1 meV/T was measured, in fair agreement with the value $\mu_{\rm orb} = 0.52 \pm 0.06$ meV/T expected from Eq. (10) (Minot *et al.*, 2004).

As seen from Fig. 11, a nanotube that is semiconducting at zero field becomes metallic in a parallel field with magnitude

⁴Our convention for assigning valley labels is that conductionband states decreasing (increasing) in energy with increasing B_{\parallel} are labeled *K* (*K*').

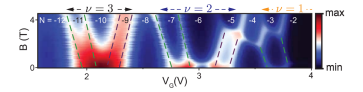


FIG. 12 (color online). Conductance of a nanotube quantum dot as a function of gate voltage and magnetic field, allowing groundstate spectroscopy of the first 12 hole states. (The first two peaks of the $\nu = 1$ shell are not visible on this color scale.) Occupation numbers *N* for holes 1–12 and shell numbers ν are indicated. The orbital magnetic moment leads to a field-dependent shift of the conductance peaks, marked by dashed lines. From the slope of these lines, μ_{orb} values of 0.90, 0.80, and 0.88 meV/T for shells 1–3 can be deduced, an order of magnitude larger than the spin magnetic moment. As well as the linear shift of peak positions, other features are seen: the complex network of lines above 2 T on the right-hand side reflects energy crossings between different shells, while the barely resolved low-field anticrossing at $V_G \sim$ 3 V probably reflects spin-orbit coupling not recognized at the time (Sec. III.F). Adapted from Jarillo-Herrero *et al.*, 2005a.

 $|B_{\parallel}| = B_{\text{Dirac}}$. This peculiar metal-semiconductor transition is specific to the cylindrical form of nanotubes and in fact recurs periodically with every flux quantum $\Phi_0 = h/e$ that threads the cross section. For a semiconducting nanotube, the gap closes twice per period, at flux equal to $\Phi \equiv \pi D^2 B_{\parallel}/4 =$ $\Phi_0/3$ and $\Phi = 2\Phi_0/3$, where the open and filled circles, respectively, in Fig. 11(h) cross quantization lines. The semiconducting gap reopens completely at $\Phi = \Phi_0$. Figure 13 shows magnetoconductance of a semiconducting nanotube for which an AFM determined $D \approx 8$ nm. For this diameter, the expected B_{Dirac} is 27 T, which is accessible at dedicated facilities. The nanotube has low conductance at zero field when $E_{\rm F}$ is tuned into the gap. At 22 T the conductance is maximal, likely since the band gap is reduced to a smaller value. The band gap reopens to a maximum near 37 T before closing again as expected. The inset curve is calculated for

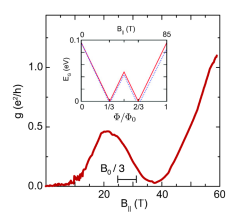


FIG. 13 (color online). Magnetoconductance of a semiconducting nanotube near the charge neutrality point measured at 3.1 K. The observed maximum is slightly below the expected value (indicated by the bar labeled $B_0/3$). The inset shows a calculated energy gap for a (95,15) semiconducting nanotube (D = 8.1 nm) vs parallel magnetic field. Solid and dashed lines are without and with Zeeman effect. Adapted from Jhang *et al.*, 2011.

D = 8.1 nm and predicts gap minima at 27 and 55 T. The observed conductance maximum, corresponding to the first gap closing, occurs at a somewhat lower field of 22 T, which is attributed to strain.

Equation (10) assumed an electron with zero longitudinal momentum. For electrons confined in a quantum dot, g_{orb} is reduced, for the following reason (Jespersen, Grove-Rasmussen, Flensberg *et al.*, 2011). As seen from the insets of Fig. 14, the partial derivative in Eq. (9) decreases with increasing κ_{\parallel} ; the larger κ_{\parallel} , the smaller the fraction of $v_{\rm F}$ directed around the nanotube and hence the smaller μ_{orb} . Because of confinement the shells participating in transport consist of superpositions of states with $|\kappa_{\parallel}| \neq 0$. (This can be seen for the sequence of shells in Fig. 10.) The total orbital moment therefore decreases with increasing confinement energy (Fig. 14), with predicted scaling (Jespersen, Grove-Rasmussen, Flensberg *et al.*, 2011):

$$g_{\rm orb} = \frac{g_{\rm orb}^0}{\sqrt{1 + (2E_{\rm conf}/E_{\rm G}^0)^2}},$$
 (12)

where $g_{\rm orb}^0$ is the unconfined value derived from Eq. (10),

$$g_{\rm orb}^0 = \frac{e v_{\rm F} D}{4\mu_{\rm B}},\tag{13}$$

where E_G^0 is the band gap at zero magnetic field without spinorbit coupling. Figure 14 shows a series of measured g_{orb} values as a quantum dot was tuned across the electron-hole transition using a gate voltage. As expected from Eq. (12), g_{orb} is maximized close to the transition, where electrons and holes can occupy the lowest-energy confined states, but reduced as

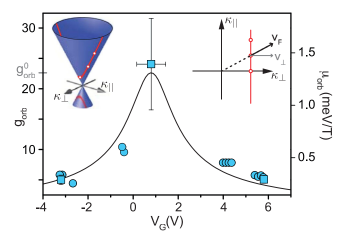


FIG. 14 (color online). Effect of confinement on magnetic moment. Main plot: Experimentally determined $g_{\rm orb}$ and $\mu_{\rm orb}$ (points) as a function of gate voltage, together with a fit by Eq. (12). The fit assumes a constant dot length and linear dependence of $E_{\rm conf}$ on $V_{\rm G}$. Left inset: Dirac cone showing quantization line and points corresponding to κ_{\parallel} values for three longitudinal energy levels. Right inset: View of the Dirac cone from above, giving a physical explanation for the reduction of $\mu_{\rm orb}$. Regardless of $\kappa_{\parallel}, \kappa_{\perp}$, the velocity (black arrow) of the electron is $v_{\rm F}$ directed away from the origin. Higher energy states, with larger κ_{\parallel} , have smaller perpendicular components v_{\perp} and therefore smaller $g_{\rm orb}$. From Jespersen, Grove-Rasmussen, Flensberg *et al.*, 2011.

the quantum dot occupation is increased. The data are well fitted by Eq. (12) assuming $E_{\text{conf}} \propto V_{\text{G}} - 0.8 \text{ V}$, which is reasonable if the length of the quantum dot is independent of gate voltage.

The orbital moment deduced from Fig. 14 corresponds to Eq. (13) with $D \approx 6$ nm. This is surprisingly large for the chemical-vapor-deposited nanotube used in the experiment, for which $D \lesssim 3$ nm is expected. Other experiments have measured a range of values for $g_{\rm orb}$; while some have obtained similarly large values (Jarillo-Herrero *et al.*, 2005a; Kuemmeth *et al.*, 2008; Steele *et al.*, 2013), other results are consistent with smaller-diameter nanotubes (Minot *et al.*, 2004; Makarovski *et al.*, 2007; Churchill, Bestwick *et al.*, 2009). Although the discrepancy is not large, it is possible that expectations for either the nanotube diameter or the orbital $v_{\rm F}$ need to be revised. Measurements on devices with independently measured diameters allow the discrepancy to be explored.

F. Spin-orbit coupling

On first consideration, it might be expected that carbon, as the second lightest of all semiconductors, should have negligible spin-orbit coupling. Indeed, spin-orbit coupling is comparatively weak in free carbon $[{}^{3}P_{0} - {}^{3}P_{1}$ splitting ~2 meV (Kramida, Ralchenko, and Reader, 2013)], and almost completely suppressed near the Dirac points in flat graphene (Huertas-Hernando, Guinea, and Brataas, 2006; Min *et al.*, 2006). However, it was realized by Ando (2000) that the suppression relies on the symmetry of graphene. In a nanotube, this symmetry is broken by curvature, leading to a coupling up to a few meV between the spin and orbital moment of electrons. This coupling, first detected by Kuemmeth *et al.* (2008), is the key to electrically controlling spins in nanotubes. This section explains in detail how it arises and how it is measured.

1. Origin of spin-orbit interaction in nanotubes

In atomic carbon, coupling between the total spin S and orbital angular momentum L adds a term to the Hamiltonian:

$$H_{\rm SO}^{\rm atomic} = \Delta_{\rm SO}^{\rm atomic} \mathbf{L} \cdot \mathbf{S},\tag{14}$$

where $\Delta_{\text{SO}}^{\text{atomic}}$ is the atomic spin-orbit strength (From the atomic ${}^{3}P_{0} - {}^{3}P_{1}$ splitting quoted above, $\Delta_{\text{SO}}^{\text{atomic}} \sim 4 \text{ meV.}$) The effect of this coupling is to mix single-particle states with opposite spin from different orbitals, such as $|p_{z}\uparrow\rangle$ and $|p_{x}\downarrow\rangle$. Whether this leads to spin-orbit coupling in the band structure depends on how it affects hybridization between orbitals in different atoms, which in turn depends on the crystal structure.

The contrasting situations in flat and curved graphene are illustrated in Fig. 15, which shows the atomic orbitals for two adjacent atoms A and B. Any effect on the band structure arises through the combination of intra-atomic spinorbit coupling and interatomic hopping. In flat graphene [Fig. 15(a)], symmetry forbids direct hopping from a p_x state on one atom to a p_z state on another because p_x and p_z orbitals have opposite parity under z inversion. Therefore atomic spin-orbit coupling between, e.g., $|p_z^{+}\uparrow\rangle$ and $|p_x^{+}\downarrow\rangle$

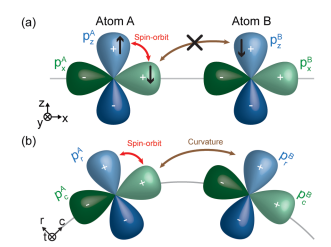


FIG. 15 (color online). How curvature enhances spin-orbit coupling. (a) Atomic spin-orbit coupling and interatomic hopping in flat graphene. The p_z and p_x orbitals of two adjacent atoms are shown, with the sign of the wave function in each lobe marked. Intra-atomic spin-orbit coupling mixes opposite-spin states involving different p orbitals in the same atom, e.g., $|p_z^A\uparrow\rangle$ and $|p_x^A\downarrow\rangle$. However, this does not mix spin states in the band structure, because hopping between different p orbitals is forbidden by symmetry. (For example, the hopping contributions between p_x^A and p_z^B marked by a double arrow exactly cancel.) Consequently, there is no coupling between states such as $|p_z^A\uparrow\rangle$ and $|p_z^B\downarrow\rangle$ and therefore no bulk spin-orbit coupling. (b) In a nanotube, curvature breaks the up-down symmetry, meaning that direct hopping between different orbitals on adjacent atoms becomes possible. The combination of atomic spin-orbit coupling and interatomic hopping therefore mixes opposite-spin states on adjacent atoms (e.g., $|p_r^A\uparrow\rangle$ and $|p_r^B\downarrow\rangle$), leading to a spin-orbit splitting of the π band.

states does not introduce any non-spin-conserving hybridization between $|p_z^A\uparrow\rangle$ and $|p_z^B\downarrow\rangle$, and thus spin-orbit coupling in the π band is second order and in practice negligible.

This situation is changed in the presence of curvature, which breaks the *z* inversion symmetry on which the above suppression relies [Fig. 15(b)]. To understand this, it is convenient to work in the curved coordinate basis $\{r, c, t\}$ labeling radial, circumferential, and axial directions, so that the π band is composed predominantly of hybridized p_r orbitals. Since the p_c^A and p_r^B orbitals are not orthogonal, hopping between them is allowed, leading to an indirect hybridization between $|p_r^A \uparrow \rangle$ and $|p_r^B \downarrow \rangle$ and consequently a spin-orbit coupling in the π band.

As a result of this spin-orbit coupling, the effective hopping matrix element between p_r^A and p_r^B now contains both a direct and a spin-flip term. The interference between these terms causes a spin precession about the y axis, and a corresponding splitting of the two spin states within a given valley as though by a magnetic field \mathbf{B}_{SO} directed along the nanotube. The spin-orbit splitting is defined as the Zeeman splitting due to this field $\Delta_{SO} = g_s \mu_B B_{SO}$.

Figures 16(a) and 16(b) show the consequences of spinorbit coupling for the band edges. Without spin-orbit coupling [Fig. 16(a)], the zero-field levels are fourfold degenerate, but are split in a magnetic field through a combination of Zeeman

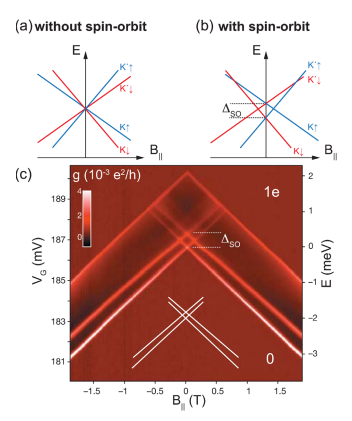


FIG. 16 (color online). Observation of spin-orbit coupling in a nanotube. (a), (b) Expected spectra without and with spin-orbit coupling. (c) Conductance as a function of magnetic field and gate voltage across the 0-1*e* transition at $V_{SD} = 2$ mV, allowing high-bias spectroscopy of the lowest four one-electron states. From line slopes (inset), the spin and orbital magnetic moments can be deduced, allowing assignment of spin and valley quantum numbers. The data are clearly consistent with (b) rather than (a). Adapted from Kuemmeth *et al.*, 2008.

and orbital coupling. [This figure differs from Fig. 11(g) by the inclusion of Zeeman spin splitting.] Spin-orbit coupling splits each fourfold degenerate level at B = 0 into a pair of twofold degenerate levels [Fig. 16(b)]; each element of the pair comprises a Kramers doublet, as required by time-reversal symmetry. The sign of Δ_{SO} determines whether parallel or antiparallel alignment of spin and valley magnetic moments is favored. For $\Delta_{SO} > 0$, the magnetic moments of spin and valley of the lowest (highest) edge of the conduction (valence) band add, whereas they subtract for $\Delta_{SO} < 0$ (see Fig. 18 for examples). For $\Delta_{SO} > 0$, as drawn here, spin-orbit coupling favors alignment of the spin and valley magnetic moments. The lower doublet therefore comprises the states $\{K'\uparrow, K\downarrow\}$ for which both magnetic moments have the same sign, while the upper doublet comprises the states $\{K'\downarrow, K\uparrow\}$.

2. The discovery of nanotube spin-orbit coupling

Figure 16(c) shows excited-state spectroscopy of the first electron shell of an ultraclean nanotube as a function of the magnetic field (Kuemmeth *et al.*, 2008). The positions in gate voltage of the first four conductance peaks provide a map of the lowest four energy levels, as explained in Appendix A.2 [which also explains how to convert from gate voltage (left

axis) to energy (right axis)]. The dependence of the energy levels on magnetic field arises from the combination of valley and spin magnetic moments in each level, as in Fig. 11(g), but taking account of the spin magnetic moment. From the line slopes, valley and spin quantum numbers can be assigned to each level [as in Figs. 16(a) and 16(b)].

The data are clearly consistent with Fig. 16(b) rather than with Fig. 16(a). The key signature of spin-orbit coupling is the separation of the four spin-valley levels at zero field into two doublets. The magnitude of the splitting gives the spinorbit coupling strength $\Delta_{SO} = 0.37$ meV, corresponding to $B_{SO} = 3.1$ T. From the spin and valley assignments deduced from the line slopes, it is clear that in this case spin-orbit coupling favors states with parallel spin and valley magnetic moments.

3. Different types of nanotube spin-orbit coupling

Detailed calculations of the spin-orbit coupling reveal that there are actually two terms in the spin-orbit Hamiltonian,

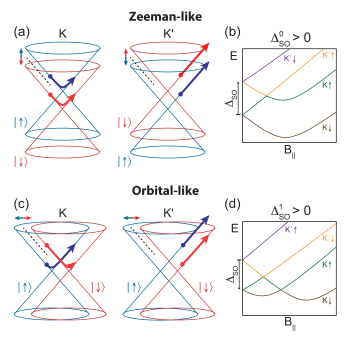


FIG. 17 (color online). Two types of spin-orbit coupling. (a), (b) Zeeman-like coupling (Izumida, Sato, and Saito, 2009) leads to a spin-dependent vertical shift of the band structure, equivalent to a Zeeman splitting (a) that is opposite in the two valleys. (b) Energy levels as a function of B_{\parallel} . Going beyond Fig. 18, a residual band gap (induced, for example, by level quantization) is assumed, leading to rounding of the band minima. (c), (d) Orbital-like coupling (Ando, 2000) leads to a horizontal shift of the band structure (c) equivalent to a spindependent magnetic field coupling to $\mu_{\rm orb}$. (d) Corresponding energy levels as a function of B_{\parallel} . A signature to distinguish these two couplings comes from the minima in (b) and (d); whereas Zeeman-like coupling leads to minima in the first and second energy levels, orbital-like coupling leads to two minima in the first level. In (a) and (c), thick arrows indicate evolution of the state energies in the Dirac cones as B_{\parallel} is increased. Spectra (b) and (d) are calculated from Eq. (20) using $E_{\rm G} = 4$ meV, $E_{\rm conf} = 1$ meV, $\mu_{\rm orb}^0 = 0.9$ meV/T, and $\Delta_{\rm SO}^0, \Delta_{\rm SO}^1 = 0$ or 2 meV.

corresponding to Zeeman-like and orbital-like coupling (Fig. 17). The Zeeman-like contribution, characterized by a parameter Δ_{SO}^0 , can be visualized as a vertical shift of the Dirac cones that is opposite for the two spin directions [Fig. 17(a)]. This is equivalent to an effective Zeeman shift of each spin state given by

$$\Delta E^{\rm SO,Z}(\tau,s) = \Delta_{\rm SO}^0 \tau s, \tag{15}$$

where s and τ are defined at the end of Sec. III.E.1.

The orbital-like contribution to the Hamiltonian gives rise to a horizontal shift of the Dirac cones, similar to the curvature-induced shift discussed in Sec. III.C, but opposite for the two spin directions [Fig. 17(c)]. This is equivalent to a spin-dependent magnetic flux coupling to the orbital moment. The horizontal shift of each cone is

$$\Delta \kappa_{\perp}^{\rm SO,orb}(s) = -s \frac{\Delta_{\rm SO}^{\rm l}}{\hbar v_{\rm F}},\tag{16}$$

where Δ_{SO}^1 parametrizes the strength of the coupling. In this case, the hole energy levels are no longer simply the negative of the electron energy levels at finite field, and the electron-hole symmetry is broken (Kuemmeth *et al.*, 2008).

Equations (15) and (16) can be combined with the graphene dispersion relation (1) and the magnetic field response (11) to give the Hamiltonian in terms of the spin and valley quantum numbers, including spin-orbit and magnetic coupling:

$$H = \left[\hbar v_{\rm F} \left(\tau \Delta \kappa_{\perp} + s \frac{\Delta_{\rm SO}^1}{\hbar v_{\rm F}} \right) + \mu_{\rm orb}^0 B_{\parallel} \right] \tau \sigma_1 + \hbar v_{\rm F} \kappa_{\parallel} \sigma_2 + \tau s \Delta_{\rm SO}^0 + \frac{g_{\rm s} \mu_B}{2} \mathbf{s} \cdot \mathbf{B}, \qquad (17)$$

where **s** is the spin operator (with eigenvalues $s = \pm 1$). Writing this Hamiltonian required the use of the Pauli operators $\{\sigma_1, \sigma_2\}$, which act in a two-dimensional sublattice space describing the wave function amplitude on the two sublattices of Fig. 4. These are described in more detail in Appendix B. In Eq. (17), the horizontal and vertical shifts due to the two spin-orbit terms are evident.

The existence of two forms of spin-orbit coupling goes beyond the simple picture of Sec. III.F.1. The orbital-like contribution can be understood as a Rashba-type coupling, arising from the broken reflection symmetry about the graphene plane. A curvature-induced displacement of the orbitals gives rise to a radial electric field, which circulating electrons experience as a magnetic field proportional to the azimuthal component of their momentum. It is similar to the Rashba coupling predicted by Kane and Mele (2005) for graphene in an electric field and is equivalent to a horizontal shift of the dispersion relation of the form in Eq. (16).

The Zeeman-like contribution also comes from a lack of reflection symmetry through the nanotube surface. In contrast to the orbital-like contribution, which is caused by the homogeneous part of the radial electric field, the Zeemanlike contribution comes from variation of the electric field within the graphene unit cell. In a tight-binding picture, this contribution can be thought of as curvature-induced spin-orbit scattering between next-nearest neighbors (e.g., from one A site to another), whereas the orbital-like contribution comes from scattering between nearest neighbors (between an A and a B site). The perturbation theory leading to these separate effects is outlined in Appendix B.

The values of the coefficients Δ_{SO}^0 and Δ_{SO}^1 depend on the structure of the nanotube. The theoretical values are estimated as

$$\Delta_{\rm SO}^0 \approx -\frac{0.3 \text{ meV}}{D \text{ (nm)}} \cos 3\theta$$
 (Zeeman-like), (18)

$$\Delta_{\rm SO}^1 \approx -\frac{0.3 \text{ meV}}{D \text{ (nm)}} \quad \text{(orbital-like)}. \tag{19}$$

These values are quite sensitive to the method of computation. The first calculations from Ando (2000), later refined by Yanik et al. (2004) and Huertas-Hernando, Guinea, and Brataas (2006)] considered the modification of hopping amplitudes by atomic spin-orbit coupling which gives rise to the orbital-like contribution. Later work (Izumida, Sato, and Saito, 2009) calculated the spin-orbit correction in more detail using a nonorthogonal tight-binding calculation that incorporated the spin degree of freedom and used four orbital states per atom. This work was the first to predict the Zeeman-like contribution. The parameters Δ_{SO}^0 and Δ_{SO}^1 were estimated by Izumida, Sato, and Saito (2009) in two ways: by nearestneighbor tight binding using density-functional theory potentials, and by fitting to a full numerical model, from which Eqs. (18) and (19) are taken. They have also been calculated by an extended tight-binding Slater-Koster method (Chico, López-Sancho, and Munoz, 2009) and using densityfunctional theory combined with atomic spin-orbit coupling and tight binding (Zhou, Liang, and Dong, 2009). A similar calculation to Izumida, Sato, and Saito (2009) was performed by Jeong and Lee (2009), corroborating these results. The coefficients have also been estimated by Klinovaja et al. (2011a), who included the effects of external electric fields. These different methods differ quantitatively by a factor up to ~3, but the $\cos 3\theta$ dependence is dictated by symmetry.

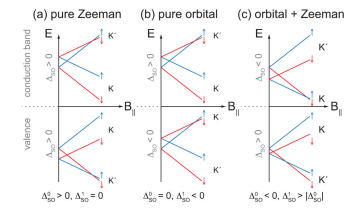


FIG. 18 (color online). Dependence of band edges on B_{\parallel} for different types of spin-orbit coupling for (a) Zeeman-like, (b) orbital-like, and (c) mixed spin-orbit. At finite field, electron-hole symmetry is broken by orbital-like but not by Zeeman-like coupling; at zero field, a combination of both is required.

Combining Eqs. (2), (8), (15), and (16) shows that both spin and valley are good quantum numbers in a magnetic field B_{\parallel} directed along the nanotube. The corresponding eigenenergies (Jespersen, Grove-Rasmussen, Paaske *et al.*, 2011) are, again assuming a flat potential as in Eq. (12),

$$E_{\tau,s}^{\pm}(B_{\parallel}) = \pm \sqrt{\left(-\tau \frac{E_{\rm G}^{0}}{2} + \mu_{\rm orb}^{0} B_{\parallel} + s \Delta_{\rm SO}^{1}\right)^{2} + E_{\rm conf}^{2}} + s \tau \Delta_{\rm SO}^{0} + \frac{s g_{\rm s} \mu_{\rm B} B_{\parallel}}{2}, \qquad (20)$$

where E_G^0 is the band gap at zero field without spin-orbit coupling and the upper (lower) sign refers to the conduction (valence) band. In the limit $E_G^0 \gg \mu_{orb}^0 |B_{\parallel}|, |\Delta_{SO}^1|$, this gives for the combined zero-field splitting

$$\Delta_{\rm SO} = 2 \left(\Delta_{\rm SO}^0 \mp \Delta_{\rm SO}^1 \frac{g_{\rm orb}}{g_{\rm orb}^0} \right). \tag{21}$$

The sign of Δ_{SO} can be deduced from the spectrum as follows (Fig. 18): If the two *K'* states converge with increasing field and eventually cross (as in Fig. 16), then $\Delta_{SO} > 0$; if it is the *K*

TABLE II. Summary of spin-orbit parameters measured in the literature, compared with the largest theoretical values predicted by Eq. (21). Unless otherwise noted, *D* is measured from the orbital magnetic moment according to Eq. (12). The spread of Δ_{SO} values is large, but in many cases the measured values considerably exceed the largest possible predictions or have the wrong sign. Other measurements (Pei *et al.*, 2012) on the same devices are not included. Also not included are earlier measurements subsequently reinterpreted as indicating spin-orbit interaction with $\Delta_{SO} \sim \Delta_{KK'}$ (Jarillo-Herrero *et al.*, 2005a; Makarovski *et al.*, 2007).

Reference	D	$\Delta_{ m SO}^{ m min}$	$\Delta_{ m SO}^{ m max}$	$\Delta_{ m SO}$
		(The	eory)	(Experiment)
	(nm)	(µeV)	(µeV)	(µeV)
Kuemmeth et al. (2008)	5.0	-120	120	370 (1 <i>e</i>)
		-240	0	-210(1h)
Churchill,	1.5	-400	400	170 (1 <i>e</i>)
Kuemmeth et al. (2009)				
Jhang et al. (2010)	1.5^{a}	-800	400	$\pm 2500^{b}$
Jespersen, Grove-Rasmussen, Paaske <i>et al.</i> (2011)	2.9 ^c	-210	210	150 (Many <i>e</i>)
		-410	0	75 (Many h)
Lai, Churchill, and	1.3 ^c	-920	460	· · · ·
Marcus (2014)				
				(Many e/h)
Steele et al. (2013)				
Device 1	7.2	-80	80	3400 (1 <i>e</i>)
Device 2	6.8	-90	90	1500 (1e)
Device 3	4.1	-150	150	
	3.7	0	-320	1300 (1 <i>h</i>)
Cleuziou et al. (2013)				
Device 1				-240
				(h, few shells)
Device 2				-340 (1e, 3e)
Schmid et al. (2015)				-350 (17 <i>e</i>)

^aAFM measurement.

^bInferred from the magnetoresistance of an open CNT quantum wire.

^cDerived from many-carrier regime but ignoring suppression of g_{orb} by confinement [Eq. (12)]. These values of *D* are therefore likely to be underestimates.

states that cross, then $\Delta_{SO} < 0$ (Bulaev, Trauzettel, and Loss, 2008; Kuemmeth *et al.*, 2008).

From Eqs. (18), (19), and (21), four predictions can be derived:

- (1) Spin-orbit coupling depends on chirality and diameter; hence different devices should display different coupling.
- (2) The different terms lead to different behavior when fields comparable to B_{Dirac} are applied (see Fig. 11). This can be seen by plotting the first four energy levels as a function of B_{\parallel} across the Dirac point. For orbital coupling (and assuming a residual band gap remains), the lowest-energy level has a pair of minima [Fig. 17(d)], corresponding to the Dirac points crossing the quantization lines for spin up and spin down. For Zeeman-like coupling, the two minima occur in the first and second energy levels, as in Fig. 17(b).
- (3) The orbital-like term contributes with opposite sign to Δ_{SO} for electrons and holes, thereby breaking electron-hole symmetry (because the hole energy levels are no longer a mirror image of the electron

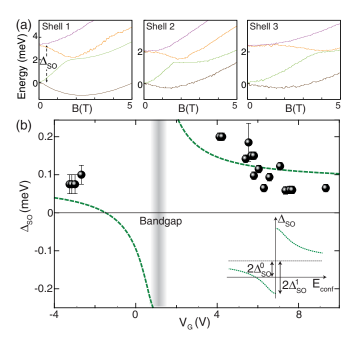


FIG. 19 (color online). Experimentally distinguishing orbitaltype and Zeeman-type spin-orbit coupling contributions. (a) Measured magnetic field dependence of the first 12 electron energy levels, split across three shells, in a narrow-gap nanotube, derived from low-bias spectroscopy. (The vertical offsets of each trace, which this experimental technique leaves undetermined, have been chosen so that adjacent levels align.) The levels correspond more closely to Fig. 17(b) than Fig. 17(d) (and the colors have been chosen accordingly), indicating predominantly Zeeman-like coupling in this device. (b) Δ_{SO} as a function of V_G in a separate device. The dashed lines are fits to theory taking account of the different dependence of the two contributions on orbital moment. Inset: Expected dependence of Δ_{SO} on E_{conf} , showing the strengths of the two contributions. Δ_{SO}^0 and Δ_{SO}^1 parametrize the Zeeman-like and orbital-like contributions, respectively. Adapted from Jespersen, Grove-Rasmussen, Paaske et al., 2011 and Steele et al., 2013.

levels.) The Zeeman-like term by itself preserves electron-hole symmetry [Fig. 18(b)].

(4) The orbital contribution leads to a smaller energy shift for higher energy shells in the same way that an orbital magnetic field does (Sec. III.E). Thus Δ_{SO} depends on density in the same way as g_{orb} [Eq. (21)].

Evidence for prediction (1) comes from measurements on several devices, presumably with a distribution of structures. The spin-orbit parameters are indeed found to take different values (Table II), although it has not yet been possible to determine chiral indices in the same device for comparison. The spread of Δ_{SO} is in fact larger than expected from Eqs. (18) and (19), as discussed in Sec. III.H.

Testing prediction (2) experimentally requires a device where B_{Dirac} lies at an accessible field. Figure 19(a) shows ground-state spectroscopy of the first four Coulomb peaks in a device where this was achieved (Steele *et al.*, 2013). The pattern of energy levels clearly resembles Fig. 17(b) more than Fig. 17(d), suggesting that the coupling in this device is predominantly Zeeman like.

Prediction (3) was confirmed in the first measurements by Kuemmeth *et al.* (2008), who found Δ_{SO} to be different in both magnitude and sign for the first electron and first hole. Further confirmation, and a test of prediction (4), was obtained in a different device where Δ_{SO} could be measured across several shells of electrons and holes (Jespersen, Grove-Rasmussen, Paaske *et al.*, 2011). As seen in Fig. 19(b), Δ_{SO} decreases with higher $|E_{conf}|$, qualitatively consistent with Eq. (21) assuming an orbital contribution. For the shells measured in this device, Δ_{SO} did not change sign between electron and holes, providing further evidence of a Zeemantype contribution.

4. Uniform electric fields

As well as intrinsic spin-orbit coupling from the nanotube structure, an extrinsic coupling is predicted due to electric fields (Klinovaja *et al.*, 2011a). This is a form of the Rashba effect and leads to a shift of the Dirac cones in κ_{\parallel} by an amount

$$\Delta \kappa_{\parallel}^{\rm SO,R} = \frac{eE\xi}{\hbar v_{\rm F}} \tau s_{\perp}, \qquad (22)$$

where *E* is the electric field (perpendicular to the nanotube) and s_{\perp} is the spin component perpendicular to both the

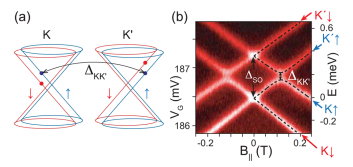


FIG. 20 (color online). (a) Example of a pair of states mixed by intervalley scattering, which couples states with equal spin in opposite valleys. (b) Close-up of Fig. 16. Intervalley scattering is manifest as an anticrossing between $|K\uparrow\rangle$ and $|K'\uparrow\rangle$ levels, with strength $\Delta_{KK'}$. Adapted from Kuemmeth *et al.*, 2008.

nanotube and *E*. The parameter ξ , which governs the strength of this effect, is uncertain because it depends on several numerically calculated band-structure parameters, but is estimated as $\xi \approx 2 \times 10^{-5}$ nm (Klinovaja *et al.*, 2011a). This Rashba-like coupling has not yet been observed, but in principle allows for all-electrical spin manipulation (Bulaev, Trauzettel, and Loss, 2008; Klinovaja *et al.*, 2011b).

G. Intervalley scattering

In quantum dots (i.e., confined states), magnetospectroscopy as in Fig. 16 allows measurement of a phenomenological parameter $\Delta_{KK'}$ that governs the strength of avoided level crossings between opposite valley states. This was observed by Kuemmeth et al. (2008) in a suspended oneelectron quantum dot for which $\Delta_{KK'} = 65 \ \mu eV$ was found [Fig. 20(b)]. Although this splitting indicates scattering between states in different valleys with the same spin, it does not reveal a specific mechanism. Experiments performed on top-contacted nanotubes showed $\Delta_{KK'}$ as small as 25 μ eV (Churchill, Kuemmeth et al., 2009), and Jespersen, Grove-Rasmussen, Paaske et al. (2011) reported significant fluctuations within a device $\Delta_{KK'} = 75 - 700 \,\mu\text{eV}$ with no obvious correlation to gate voltage or occupation number. Grove-Rasmussen et al. (2012) investigated the suppression of K - K' mixing by application of a parallel magnetic field. Although $\Delta_{KK'}$ has been used in other works as an empirical fitting parameter (Pei et al., 2012; Lai, Churchill, and Marcus, 2014), the microscopic origin has not been investigated. In particular, all experiments involved finite tunneling to the source or drain electrode, and hence the intrinsic valley coupling in closed quantum dots has yet to be measured.

Historically, we suspect that electrical disorder on the scale of the interatomic spacing, leading to comparatively large $\Delta_{KK'}$, was the main reason for the decade-long delay between the first nanotube quantum dots and the discovery of spinorbit coupling. As evident below, $\Delta_{KK'}$ values larger than Δ_{SO} obscure the signatures of spin-orbit coupling. It was only the development of low-disorder fabrication techniques that allowed such a delicate effect to be identified.

1. Putting it all together

Figure 21 shows calculated spectra in a single-electron shell as a function of magnetic field in both perpendicular and parallel directions for a range of values of Δ_{SO} and $\Delta_{KK'}$. This calculation proceeds by first working out the eigenenergies $E_{\tau,s}^{\pm}$ for a magnetic field directed along the nanotube and in the absence of disorder. Under the assumption that there is no mixing of electron and hole shells, i.e., in the limit

$$E_{\rm G}^0 \gg |\Delta_{\rm SO}^1|, \mu_{\rm orb}^0 |B_{\parallel}|, \qquad (23)$$

Eq. (20) reduces to

$$E_{\tau,s}^{\pm} \approx E_0^{\pm} + s\tau \frac{\Delta_{\rm SO}}{2} + \left(\mp \tau g_{\rm orb} + \frac{1}{2} sg_{\rm s}\right) \mu_{\rm B} B_{\parallel}, \qquad (24)$$

where $E_0^{\pm} = \pm \sqrt{(E_G^0)^2/4 + E_{\text{conf}}^2}$ is the energy without magnetic field or spin-orbit coupling.

Introducing disorder or tilting the field relative to the nanotube axis by angle Θ (so that $B_{\parallel} = B \cos \Theta$) mixes the eigenstates. In the basis $(K\uparrow, K'\downarrow, K\downarrow, K'\uparrow)$, the Hamiltonian is

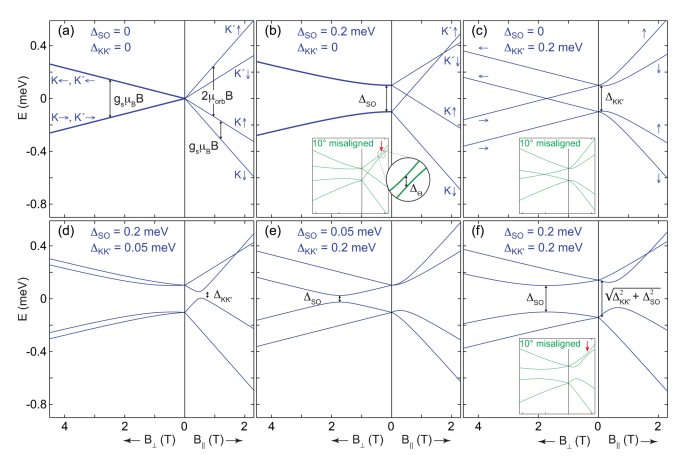


FIG. 21 (color online). (a)–(f) Single-particle energy levels as a function of magnetic field in perpendicular and parallel directions for various strengths of spin-orbit coupling and intervalley scattering, taking $\mu_{orb} = 0.2 \text{ meV/T}$. (a) No spin-orbit or intervalley scattering, (b) spin-orbit only, (c) intervalley scattering only, and (d)–(f) combined spin-orbit and intervalley scattering. In selected panels, insets show the same spectra for field axes misaligned by $\Theta = 10^{\circ}$ from the nanotube; this introduces an anticrossing (marked by arrows) between the two highest levels. Selected energy splittings mentioned in the text are marked. Double-thickness lines indicate degenerate levels.

$$H = \begin{pmatrix} E_{1,1}^{\pm} & 0 & 0 & \Delta_{KK'}/2 \\ 0 & E_{-1,-1}^{\pm} & \Delta_{KK'}/2 & 0 \\ 0 & \Delta_{KK'}/2 & E_{1,-1}^{\pm} & 0 \\ \Delta_{KK'}/2 & 0 & 0 & E_{-1,1}^{\pm} \end{pmatrix} + \frac{g_{\rm s}\mu_{\rm B}B}{2} \begin{pmatrix} 0 & 0 & \sin\Theta & 0 \\ 0 & 0 & \cos\Theta & 0 \\ \sin\Theta & 0 & 0 & 0 \\ 0 & \sin\Theta & 0 & 0 \end{pmatrix}. \quad (25)$$

Figure 21 shows the numerically calculated eigenstates of this Hamiltonian. With no spin orbit or disorder [Fig. 21(a)], the four states are degenerate at zero field, being split through a combination of orbital and Zeeman coupling. Pure spin-orbit coupling [Fig. 21(b)] splits the zero-field quadruplet into two doublets; note that in small perpendicular field, Zeeman coupling is ineffective, because the spin states are locked to the valley, and valleys are not coupled. Pure disorder [Fig. 21(c)] suppresses orbital coupling at low field, but preserves the Zeeman splitting in both field directions. In cases where both terms are finite but one dominates, the smaller parameter leads to anticrossings [Figs. 21(d)]

and 21(e)). Finally, if the two terms are of comparable finite magnitude, a complex spectrum emerges showing a mixture of effects [Fig. 21(f)]. A small misalignment of the nanotube relative to the field axes, illustrated in the insets, introduces an anticrossing between the upper two levels. The magnitude of the anticrossing is $\Delta_{\Theta} = |\Delta_{SO}| \tan \Theta$.

This picture is confirmed in Fig. 22 for a device with comparable $\Delta_{KK'}$ and Δ_{SO} . The energy levels were measured by cotunneling spectroscopy, which maps out energy differences between ground and excited states [see Jespersen, Grove-Rasmussen, Paaske et al. (2011) for a discussion of this experimental technique]. The resonant transitions appear as peaks or dips in d^2I/dV_{SD}^2 whenever $eV_{\rm SD}$ is equal to the difference of energy levels. The measured transitions [Fig. 22(a)] as a function of perpendicular and parallel fields agree well with the predicted level differences [Fig. 22(b)] from the spectrum [Fig. 22(c)], calculated in the same way as in Fig. 21 assuming $\Delta_{KK'} = 3\Delta_{SO}$. In particular, the curvature of energy levels in the parallel field due to $\Delta_{KK'}$ mixing and the anticrossing in the perpendicular field due to Δ_{SO} are seen. From similar data the parameters Δ_{SO} and $\Delta_{KK'}$ can be precisely measured over a wide range of electron and hole occupation (Jespersen, Grove-Rasmussen, Paaske et al., 2011).

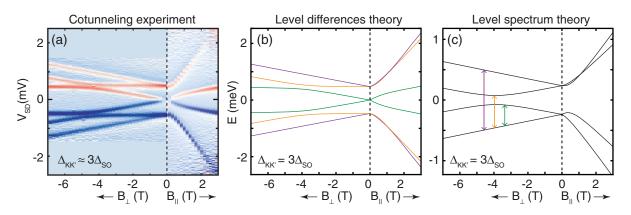


FIG. 22 (color online). Energy levels for comparable spin-orbit coupling and intervalley scattering. (a) Measurements of the four states of an electron shell by cotunneling spectroscopy (see text). The second derivative d^2I/dV_{SD}^2 is plotted as a function of V_{SD} and the magnetic field. (b) Calculated transition energies to the first three excited states assuming $\Delta_{KK'} = 3\Delta_{SO}$, showing good agreement with the peaks and dips in (a). (c) Energy levels corresponding to the transitions (marked with arrows) in (b). Adapted from Jespersen, Grove-Rasmussen, Paaske *et al.*, 2011.

We conclude this section by summarizing, with best numerical estimates, the various nanotube band-structure parameters discussed in the text (Table III). Experimentally, one set of parameters usually suffices to characterize an entire shell, implying that these parameters are not strongly affected by the addition of a few extra electrons. This is as expected from the constant-interaction model, which assumes all interactions can be parametrized by a single constant capacitance (Appendix A).

H. Open questions

Comparison of theoretical and experimental spin-orbit coefficients shows serious discrepancies. As shown in Table II different devices give unexpectedly large variation. With the diameters inferred from μ_{orb} , Eqs. (18) and (19)

predict that $|\Delta_{SO}|$ should range up to ~900 μ eV. Instead, values as large as 3.4 meV have been reported, with several devices yielding results up to 16 times larger than expected (Jhang *et al.*, 2010; Steele *et al.*, 2013). Other experiments have found Δ_{SO} within the expected range (Churchill, Kuemmeth *et al.*, 2009; Cleuziou *et al.*, 2013; Lai, Churchill, and Marcus, 2014). Furthermore, in some cases the calculations even predict the wrong sign for both couplings Δ_{SO}^0 and Δ_{SO}^1 (Kuemmeth *et al.*, 2008; Jespersen, Grove-Rasmussen, Paaske *et al.*, 2011).

This is clearly an open question for both theory and experiment. One explanation might be uncertainty in the tight-binding overlap integrals, which enter Eqs. (18) and (19) as empirical input parameters. Alternatively, electron interactions (Sec. VII.B.2) may play a role (Rontani, 2014). Another possibility is that some other symmetry breaking

TABLE III. Summary of nanotube quantum dot energy parameters. For the numerical values, representative estimates are given, based on experiments for the last two lines and theory elsewhere. A value $v_F = 8 \times 10^5 \text{ ms}^{-1}$ is assumed.

Quantity	Expression	Value	Reference
Confinement band gap	$4\hbar v_{\rm F}/3D$	0.70 eV/D (nm)	Charlier, Blase, and Roche (2007)
Curvature band gap	$\sim \frac{\hbar v_{\rm F} a_{\rm CC}}{2D^2} \cos 3\theta$	~50 meV/[D (nm)] ² × cos 3 θ	Izumida, Sato, and Saito (2009)
Strain band gap	$\frac{2\hbar v_{\rm F}}{a_{\rm CC}} \frac{12\zeta}{1+6\zeta} (1+\lambda)\epsilon\cos 3\theta$	51 meV $\times \epsilon(\%) \times \cos 3\theta$	Huang et al. (2008)
Torsion band gap	$2\hbar v_{\rm F}\gamma\sin 3\theta$	0.018 meV $\times \gamma$ (°/ μ m) $\times \sin 3\theta$	Yang and Han (2000)
Effective mass	$m_{\rm eff} = E_{\rm G}/2v_{\rm F}^2$.	$0.14m_{\rm e} \times E_{\rm G} ~({\rm eV})$	
Orbital magnetic moment	$\mu_{ m orb}^0 = Dev_{ m F}/4$	$0.20 \text{ meV/T} \times D \text{ (nm)}$	Ajiki and Ando (1993)]
	$g_{ m orb}^0\equiv\mu_{ m orb}^0/\mu_{ m B}$	$3.5 \times D \text{ (nm)}$	
Zeeman magnetic moment	$\frac{1}{2}g_{s}\mu_{B}$	58 µeV/T	
Spin-orbit coupling	$ au s(\Delta_{ m SO}^0+\Delta_{ m SO}^1\sigma_1)$	$\Delta_{\rm SO}^0 \approx \frac{-0.3 \text{ meV}}{D(\text{nm})} \cos 3\theta$	Izumida, Sato, and Saito (2009)
	(see Appendix B)	$\Delta_{\rm SO}^1 \approx \frac{-0.3 \text{ meV}}{D(\text{nm})}$	Izumida, Sato, and Saito (2009)
		$\Delta_{\rm SO} \equiv 2(\Delta_{\rm SO}^0 \mp \Delta_{\rm SO}^1 g_{\rm orb} / g_{\rm orb}^0)$	
Electric field spin splitting	$eE\xi$	$\sim 20 \ \mu \text{eV} \times E \ (\text{V nm}^{-1})$	Klinovaja <i>et al.</i> (2011a)
Intervalley scattering	$\Delta_{KK'}$	$\Delta_{KK'} \ge 60 \ \mu eV$ (typical)	Kuemmeth et al. (2008)
Longitudinal mode spacing (high-energy limit)	$\Delta E_{\rm conf} = \frac{hv_{\rm F}}{2L}$	$1.7 \text{ meV}/L (\mu \text{m})$	Tans et al. (1997)

between inside and outside the nanotube is responsible for the observed couplings, such as gate dielectric or adsorbates. For example, hydrogen adsorbed onto graphene is known to enhance the spin-orbit coupling (Balakrishnan *et al.*, 2013). In nanotube devices, adsorbed water affects the current-voltage characteristics (Kim *et al.*, 2003), although it is uncertain whether this is directly by modifying the band structure (Na *et al.*, 2005) or through another mechanism such as gathering ions from the environment (Sung *et al.*, 2006).

A possibly related effect is that diameters inferred from measured μ_{orb} are sometimes unexpectedly large. Whereas chemical-vapor deposited nanotubes are expected to have $D \lesssim 3$ nm, values of $\mu_{\rm orb}$ corresponding to $D \sim 5$ nm have been measured (Jarillo-Herrero et al., 2004, 2005a; Minot et al., 2004; Kuemmeth et al., 2008). However, other devices yield values in the expected range (Minot et al., 2004; Makarovski et al., 2007; Deshpande and Bockrath, 2008; Churchill, Kuemmeth et al., 2009). Measurements on nanotubes with known chirality should help clarify these discrepancies. Interestingly, the prediction of complete band closing at B_{Dirac} (Fig. 11) is not borne out by experiments, which typically find minimal band gaps $E_{\rm G} \approx 10 - 100$ meV. This hints at physics beyond the single-particle picture discussed here, such as formation of a Mott gap (Deshpande et al., 2009) or an excitonic insulator (Rontani, 2014).

A separate set of questions addresses how to take advantage of the spin-orbit interaction. Ideas in this direction such as spin filtering and detection (Braunecker, Burset, and Levy Yeyati, 2013; Mazza *et al.*, 2013), generation of helical states (Klinovaja *et al.*, 2011b), and spin-orbit mediated spin control (Bulaev, Trauzettel, and Loss, 2008; Flensberg and Marcus, 2010) have already emerged (the latter is the topic of Sec. V.C.3).

IV. DOUBLE QUANTUM DOTS AND PAULI BLOCKADE

Transport through a single quantum dot involves an electron from the Fermi sea in one lead that tunnels via a discrete quantum state to an empty state in the other lead. For two dots in series an additional tunneling event occurs which involves a transition from one particular initial quantum state to a particular final state. This dot-to-dot transition is sensitive to selection rules, which determine the transition probability. The selection rules for nanotubes are based on the spin and valley quantum numbers. Whether they are obeyed in an experiment depends on to what extent spin and valley are good quantum

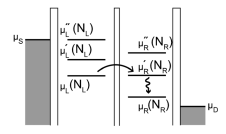


FIG. 23. Transport through double quantum dots involves a dotto-dot transition with a probability reflecting selection rules. The strictness of spin and valley selection rules depends on interactions that mix the spin or valley states, e.g., spin-orbit coupling, hyperfine interactions, and disorder.

numbers and how this is affected by spin-orbit coupling, hyperfine interaction, or disorder. This sensitivity makes double quantum dots a versatile platform for studying quantum states and relaxation processes in nanotubes (Fig. 23).

A. Role of band gap and electron-hole symmetry in charge stability diagrams

1. Theory

A double quantum dot formed by two dots in series defined within the same nanotube has similarities and differences to double dots defined in conventional semiconductors reviewed by Wiel *et al.* (2002) and Ihn (2010). Analogous to Fig. 53, the device can be modeled by an electric circuit [Fig. 24(a)]. If each of the tunnel barriers is sufficiently opaque, $\Gamma_{L,M,R} \ll E_C$, then the charge within each dot is quantized and the number of electrons $N_{L,R}$ can change only at specific

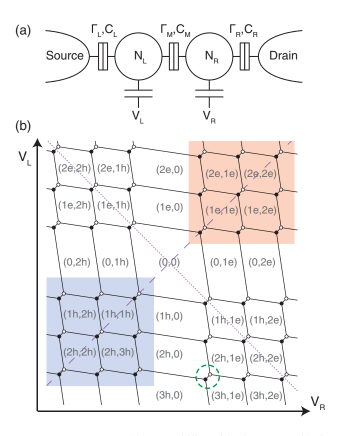


FIG. 24 (color online). Charge stability with electrons and holes (a) Circuit model of a series double quantum dot. Each of the three tunnel barriers is characterized by a tunnel rate Γ_i and a capacitance C_i . (b) Stability diagram for a weakly tunnel-coupled double dot. Pairs of triple points define a honeycomb pattern. The neutral Coulomb valley is largest, which can be understood by considering the $(1h, 0) \rightarrow (1e, 0)$ transition: adding two electrons to the left dot requires lowering its potential by an electrostatic energy (charging energy) and a kinetic energy (band gap), $\Delta V_L \propto 2E_C + E_G$. Axes of approximate spatial and electronhole symmetry are denoted by dashed and dotted lines, respectively. The top right shaded region is most similar to conventional double dots, in which transport and occupation involve only electronlike carriers. The dashed circle marks a transition with particularly strong spin-valley blockade (cf. Figs. 32 and 33).

gate voltages. A graph of the equilibrium charge configuration $(N_{\rm L}, N_{\rm R})$ as a function of gate voltages is called a stability diagram [Fig. 24(b)]. The size of each region is a measure of the addition energy [Eq. (A6)], horizontally for adding an electron in the right dot and vertically for an addition to the left dot. In most semiconductor dots either the electron or hole region is accessible. In narrow-gap nanotubes the gate coupling can be sufficient to cross the band gap, evident as larger honeycombs, and enter both regions. Because of the approximate electron-hole symmetry, a similar honeycomb pattern is expected in all regions of the stability diagram. Interestingly, the interdot tunnel barrier in the (p, n) and (n, p) regions is formed by a pn junction.

2. Experiment

The electron-hole stability diagram expected from Fig. 24 is most easily observed in narrow-gap nanotubes. Full control over the basic parameters of a double quantum dot, namely, charge occupation (N_L, N_R) and tunnel couplings $(\Gamma_L, \Gamma_M,$ and $\Gamma_{\rm R}$), requires at least five gate electrodes, and so the full charge stability diagram is at least five dimensional. A twodimensional cut is shown in Fig. 25. Here the conductance g through the device is plotted as a function of control parameters $V_{\rm R}$ and $V_{\rm L}$. Figure 25 demonstrates that an actual device can show a stability diagram that is strikingly different from the diagram of Fig. 24, characteristic of the weakcoupling regime. In Fig. 25, the middle gate voltage was intentionally chosen such that electron-hole double dots and single dots with electron or hole filling were demonstrated within the same device. In this regime, cotunneling processes give a significant contribution to transport, and hence boundaries between Coulomb valleys as well as triple points show up as conductance features. Note both the spatial and electronhole symmetry displayed by the data (mirror symmetry about

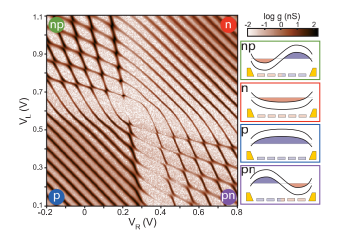


FIG. 25 (color online). Conductance through a narrow-gap nanotube suspended over five gate electrodes that allow independent control over N_L , N_R , Γ_L , Γ_M , and Γ_R . For the gate configuration shown, the nanotube is neutral in the center of the plot $(N_L, N_R) = (0, 0)$, forms a double quantum dot in the heteropolar regions (top left, bottom right), and forms a single quantum dot in the homopolar regions (top right for electrons, bottom left for holes). Insets visualize the charge distribution along the nanotube for electrons and holes. Adapted from Waissman *et al.*, 2013, device shown in Fig. 3(f).

the $+45^{\circ}$ and -45° diagonal, respectively), attesting the cleanliness and tunability of suspended devices as in Fig. 3(f).

If the tunnel rates are too small to measure conductance, the charge stability diagram can be studied by charge sensing. Such capacitive sensing techniques are useful for the readout of pulsed-gate experiments on closed double dots and qubits (Churchill, Kuemmeth *et al.*, 2009), as well as for investigating the quantum capacitance associated with electron interactions and correlations (Ilani *et al.*, 2006).

Many other regimes are possible in double quantum dots. For example, a sizable longitudinal level spacing in each dot can result in an overall eight-electron shell structure (Jørgensen *et al.*, 2008), whereas strong interdot tunneling lifts charge quantization within each dot, which can be interpreted as the formation of delocalized molecular states (Gräber *et al.*, 2006). Devices with ambipolar charge stability and a high degree of tunability have been used to study many phenomenona, including Wigner crystallization (Pecker *et al.*, 2013), Klein tunneling (Steele, Gotz, and Kouwenhoven, 2009), and tunable electron-phonon coupling (Benyamini *et al.*, 2014).

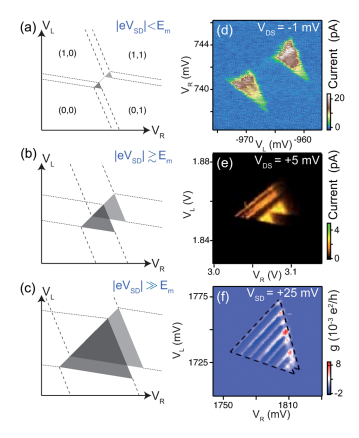


FIG. 26 (color online). (a)–(c) Dependence of bias triangles on $V_{\rm SD}$ (assumed negative). Dashed (dotted) lines indicate alignment of the electrochemical potential of the right (left) dot with the Fermi level in the right (left) lead. Sequential tunneling from source to drain is allowed only in the shaded regions, which expand with increasing $|V_{\rm SD}|$. The discrete density of states within each dot additionally restricts the current (not shown). (d)–(f) Representative bias triangles for increasing $V_{\rm SD}$. Excited-state lines in (e) and (f) identify $\Gamma_{\rm M}$ as the rate-limiting tunnel barrier. Adapted from Sapmaz, Meyer *et al.*, 2006, Churchill, 2012, and Jung *et al.*, 2013.

B. Spectroscopy of energy levels in bias triangles

Increasing the source-drain bias $V_{\rm SD}$ across a double dot allows nonlinear conductance to be probed, providing spectroscopic information on the energy levels. Applying a bias large enough to overcome Coulomb blockade expands the triple points to finite-bias triangles. These triangles serve as a powerful experimental tool to reveal a variety of quantum effects in nanotubes. Figures 26(a)-26(c) show their evolution with increasing $V_{\rm SD}$. Pairs of bias triangles start to overlap once $V_{\rm SD}$ becomes larger than the mutual charging energy $E_{\rm m}$ of the double dot.⁵ The finite bias breaks the left-right symmetry of a nominally symmetric device, yielding triangles pointing along the diagonal with a direction depending on the sign of $V_{\rm SD}$. In Figs. 26(a)-26(c) the sign of $V_{\rm SD}$ has been chosen so that electrons flow from the left contact (source) to the right contact (drain).

Under appropriate conditions, excited states (in either dot) are evident as discrete lines within a bias triangle. If interdot tunneling is the rate-limiting process, these lines appear parallel to the base of the triangles, but only if both tunnel rates to the leads remain at the same time smaller than level spacing and bias ($\Gamma_{\rm M} \ll \Gamma_{\rm L}, \Gamma_{\rm R} < \Delta_{\rm ls}, |eV_{\rm SD}|$). From the line separation, measured from the base of the triangles, the corresponding excitation energies can be deduced. Examples from three different devices [Figs. 26(d)–26(f)] show the expansion of the bias triangles with $|V_{\rm SD}|$ and the appearance of excited-state lines.⁶

C. Pauli blockade involving spin and valley

1. Motivation

The dot-to-dot transitions in conventional double dots are strongly regulated by selection rules. These selection rules arise from the Pauli exclusion principle and can provide insight into the robustness of quantum numbers in the two dots and during interdot tunneling. Since in nanotubes both spin and valley can form approximate good quantum numbers even in the presence of spin-orbit coupling, the manifestations of Pauli blockade and Pauli rectification are more complex than in conventional semiconductors. In this section, we briefly review Pauli blockade in conventional semiconductors with only twofold spin degeneracy to establish useful terminology. Next, we extend the model by adding twofold valley degeneracy to illustrate the persistence of Pauli blockade beyond spin blockade. In order to make a connection to actual nanotube experiments, we then discuss the main effects of spin-orbit coupling and electron-electron interactions on twoelectron states within a quantum dot. Finally, we present experimental evidence for Pauli blockade in nanotubes and discuss the roles of spin, valley, and hyperfine coupling.

2. State counting and Pauli blockade

Pauli blockade is well established in conventional quantum dots containing a total of two electrons (Ono *et al.*, 2002;

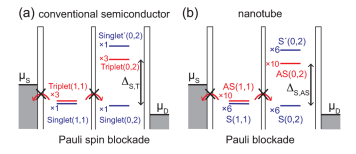


FIG. 27 (color online). Pauli rectification in conventional double dots and carbon nanotubes (theory). (a) In a conventional double dot without spin-orbit coupling, the spin-triplet (0,2) state involves an excited single-particle orbital of the right dot and hence is higher than the spin-singlet (0,2) state by energy $\Delta_{S,T}$. Near the (1,1)-(0,2) degeneracy, one of the spin-triplet states Triplet (1,1) can by chance become occupied. For sufficiently large bias $(\mu_{\rm S} - \mu_{\rm D} \gg k_{\rm B}T)$, it is long lived, resulting in a suppression of current compared with under opposite bias (not shown). (b) If each dot participates with one longitudinal singleparticle shell, then 16 different (1,1) states are possible in carbon nanotubes, colored here according to their longitudinal symmetry (S or AS). If the right dot participates with the lowest longitudinal single-particle shell, then only six different (0,2) states are possible. Other (0,2) states are higher in energy by $\Delta_{S,AS}$ (see Fig. 28). Similar to (a), sufficiently large bias $\mu_{\rm S} - \mu_{\rm D}$ can populate the long-lived (1,1) states [denoted AS(1,1)], leading to a suppression of current.

Hanson *et al.*, 2007). It relies on the fact that the (0,2) ground state is nondegenerate (it is a spin singlet) and is well separated in energy from the lowest spin-triplet states, as illustrated in Fig. 27(a). The spin-triplet states are antisymmetric in their orbital degree of freedom with respect to electron transposition and hence necessarily involve an excited single-particle state. Therefore, the energy cost

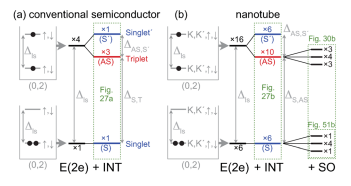


FIG. 28 (color online). Lowest quantum states of (0,2) for (a) conventional semiconductors and (b) nanotubes in the presence of interactions (INT) and spin-orbit coupling (SO). For weak electron interactions $(\Delta_{AS,S'} \ll \Delta_{ls})$, the multiplet splitting $\Delta_{S,T}$ or $\Delta_{S,AS}$ is approximately the single-particle level spacing Δ_{ls} , and Pauli blockade between Triplet (1,1) and Singlet (0,2) or AS(1,1) and S(0,2) can be expected. The states *S*, *AS*, and *S'* are part of the basis states of the matrix shown in Fig. 29. Spin-orbit interaction splits the two-electron multiplets into three. This is shown for *S* and *AS* and further discussed in Fig. 30. Additional effects due to intervalley exchange are discussed in Fig. 51.

⁵Details can be found in Wiel et al. (2002) or Ihn (2010).

⁶Figure 26 uses $V_{\rm DS}$ instead of $V_{\rm SD}$ for data with a different source-drain or left-right convention from Fig. 24(a).

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 $\Delta_{S,T}$ to form the Triplet⁷ (0,2) states is approximately the single-particle level spacing in the right dot modified by electron interactions [cf. discussion of $\Delta_{AS,S'}$ in Fig. 28(a)]. This is in contrast to the Triplet (1,1) states, which for small interdot tunneling are nearly degenerate with the Singlet (1,1). If the (0,2) splitting $\Delta_{S,T}$ is larger than temperature, then a Triplet (1,1) state cannot easily transition into the (0,2) state. If the applied source-drain bias is also larger than temperature, then the Triplet (1,1) is long lived, and its occupation suppresses current flow due to Coulomb blockade. This effect is known as spin blockade and manifests itself in current rectification (Pauli rectification).

In nanotubes, the two-electron spectrum is richer due to the two valleys. Ignoring spin-orbit coupling for clarity, it is schematically shown in Fig. 27(b). Anticipating the breakdown of spin-singlet and spin-triplet terminology due to spinorbit coupling, we have labeled the states according to the symmetry of the longitudinal quantum numbers under electron exchange (*S* or *AS* as defined below). As seen, Pauli blockade is nevertheless possible in nanotubes. Even in the presence of spin-orbit coupling and small valley scattering, the projections of spin *s* and valley τ in Eq. (25) can remain (approximate) good quantum numbers, for example, in the presence of a parallel magnetic field.⁸ As in Fig. 27(a), interactions alter the spectrum, and the splitting $\Delta_{S,AS}$ between the symmetric ground states and antisymmetric excited states is given by the level spacing modified by an interaction energy.

Figure 28 shows a simple state-counting argument leading to the degeneracies indicated in Figs. 27(a) and 27(b) for the (0,2) states. For conventional semiconductors we consider two spin-degenerate levels separated by a level spacing Δ_{ls} . The nondegenerate ground state, Singlet (0,2), is formed by two electrons occupying the lower level (lower panel). If each level is occupied by only one electron, then four degenerate states are possible (upper panel). Exchange interactions (INT) result in an energy splitting $\Delta_{AS,S'}$ between the Singlet' (0,2) and Triplet (0,2) states, thereby reducing $\Delta_{S,T}$ slightly (Kouwenhoven, Austing, and Tarucha, 2001).

The case of two fourfold degenerate nanotube shells is shown in Fig. 28(b). The lower shell can be occupied by two electrons in six ways, while 16 different states are possible with one electron in each shell.⁹ Analogous to conventional semiconductor quantum dots and to lowest order, electron interactions (INT) split the 16 states into 10 longitudinal antisymmetric (lower energy) and six longitudinal symmetric states (higher energy). In the framework of first-order perturbation theory, this can be understood by calculating the (longrange) exchange integral associated with each two-electron basis state and noticing that it differs between AS(0, 2) states and S'(0, 2) states (Appendix B.8). The complexity of the nanotube energy spectrum compared to conventional semiconductors is further revealed by turning on spin-orbit coupling within each multiplet (SO), as shown by the rightmost panel of Fig. 28(b) and discussed further.

State-counting arguments similar to those presented for (0,2) lead to 6 + 10 = 16 symmetric and antisymmetric states in the (1,1) configuration. Formally, this is accomplished by redefining the excited shell of the right dot as the lowest shell in the left dot: $AS(0,2) \rightarrow AS(1,1), S'(0,2) \rightarrow S(1,1)$. In the limit of vanishing interdot tunneling, interactions in the (1,1) regime can be neglected ($\Delta_{AS,S'} \rightarrow 0$).

3. Symmetric and antisymmetric multiplets in nanotubes: Theory

To gain more insight into the two-electron states in the presence of spin-orbit interaction,¹⁰ we first consider the quantum numbers relevant for the lowest longitudinal symmetric multiplet S(0, 2), i.e., two electrons occupying the lowest shell in the right quantum dot. The valley, spin (and longitudinal) quantum numbers are listed in Table IV with the states organized according to their energies. The two-electron energy within the constant-interaction model is found by adding the single-particle energies and a charging energy $E_{\rm C}$:

$$E = E(\nu, \tau, s) + E(\nu', \tau', s') + E_{\rm C}, \tag{26}$$

where $\nu = \nu' = 1$ specifies the lowest shell in the right dot and τ, τ' and s, s' are the valley and spin quantum numbers of the two electrons. The states are calculated by the method of Slater determinants, but taking account of the fact that ν, τ , and *s* are coupled.

Our conventions for labeling two-electron quantum states are stated below the three table headers. For example, consider the S(0, 2) state denoted $K \downarrow, K' \uparrow$, which is spin and valley unpolarized. Written out explicitly, the energy and the state are¹¹

$$E_{K\downarrow,K\uparrow}^{S(0,2)} = -\Delta_{\rm SO} + E_{\rm C}, \qquad (27)$$

$$|\psi_{K\downarrow,K'\uparrow}^{S(0,2)}\rangle = \frac{1}{\sqrt{2}}(|1K\downarrow\rangle_1|1K'\uparrow\rangle_2 - |1K'\uparrow\rangle_1|1K\downarrow\rangle_2), \quad (28)$$

where the subscripts on the right of Eq. (28) refer to electron 1 or 2, and the three labels in a single-particle state $|\nu\tau s\rangle$ are the longitudinal shell, valley, and spin quantum numbers associated with that electron. The states in Table IV are classified as longitudinally symmetric or antisymmetric according to whether the wave function remains the same or changes sign under the interchange $\nu \leftrightarrow \nu'$. Thus the example in Eq. (28) is symmetric ($\nu = \nu'$). We do not decompose $|\nu\tau s\rangle$ further into a product of longitudinal, valley, and spin wave functions, as this is strictly correct only at B = 0 and in the absence of spinorbit coupling.¹² In general, the longitudinal wave function depends on both τ and s (Weiss *et al.*, 2010).

⁷We capitalize Triplet and Singlet whenever we refer to specific spin-singlet and spin-triplet states indicated in Fig. 27.

^bEven in the absence of good quantum numbers Pauli rectification behavior can still occur, due to Kramers degeneracy. This was theoretically exemplified for double dots with strong spin-orbit coupling by Danon and Nazarov (2009).

⁹The multiplet S''(0, 2) representing two electrons in the upper shell is not shown in Fig. 28(b).

¹⁰Related theoretical treatments are found in Weiss *et al.* (2010) von Stecher *et al.* (2010), and Reynoso and Flensberg (2011, 2012).

¹¹See Appendix B for a more thorough derivation of the basis used.

¹²This decomposition can be illustrative [see, e.g., Pei *et al.* (2012) and Pecker *et al.* (2013), but is unsuitable for accurate calculations.

TABLE IV. Two-electron states in the right quantum dot (normalization factors omitted). The three sections list quantum numbers for the three lowest two-electron multiplets (*S*, *AS*, and *S'*) at zero magnetic field. Explicit expressions for the states corresponding to particular quantum numbers in this notation are given below the table headers. Each ket $|\nu \tau s\rangle_i$ represents a single-particle longitudinal, valley, and spin state of the *i*th electron. The multiplets are classified as symmetric or antisymmetric according to their behavior under the interchange $\nu \leftrightarrow \nu'$. The energy splittings $\Delta_{S,AS}$ and $\Delta_{AS,S'}$ are defined in Fig. 28.

Longitudinally symmetric states $S(0,2)$ $\tau s, \tau' s' \equiv 1\tau s\rangle_1 1\tau' s'\rangle_2 - 1\tau' s'\rangle_1 1\tau s\rangle_2$			
Designation $\tau s, \tau' s'$	Energy		
$K\downarrow,K'\uparrow$	$-\Delta_{\rm SO} + E_{\rm C}$		
$\begin{array}{l} K \downarrow, K \uparrow \\ K \uparrow, K' \uparrow \\ K \downarrow, K' \downarrow \\ K' \uparrow, K' \downarrow \end{array}$	E _C		
$K\!\uparrow,K'\!\downarrow$	$\Delta_{\rm SO} + E_{\rm C}$		

Longitudinally antisymmetric states AS(0,2) $\tau s, \tau' s' \equiv |1\tau s\rangle_1 |2\tau' s'\rangle_2 - |2\tau s\rangle_1 |1\tau' s'\rangle_2$

	$+ 1\tau's'\rangle_1 2\tau s\rangle_2 - 2\tau's'\rangle_1 1\tau s\rangle_2$
Designation $\tau s, \tau' s'$	Energy
$egin{array}{c} K\downarrow, K\downarrow\ K\downarrow, K'\uparrow\ K'\uparrow, K'\uparrow \end{array}$	$-\Delta_{\rm SO} + E_{\rm C} + \Delta_{\rm S,AS}$
$egin{array}{c} K \downarrow, K \uparrow \ K \uparrow, K' \uparrow \ K \downarrow, K' \downarrow \ K' \uparrow, K' \downarrow \end{array}$	$E_{ m C}+\Delta_{ m S,AS}$
$egin{array}{c} K\uparrow, K\uparrow\ K\uparrow, K'\downarrow\ K'\downarrow, K'\downarrow \end{array}$	$\Delta_{ m SO}+E_{ m C}+\Delta_{ m S,AS}$

Longitudinally symmetric states S'(0,2) $\tau s, \tau' s' \equiv |1\tau s\rangle_1 |2\tau' s'\rangle_2 + |2\tau s\rangle_1 |1\tau' s'\rangle_2$

	$- 1\tau's'\rangle_1 2\tau s\rangle_2- 2\tau's'\rangle_1 1\tau s\rangle_2$
Designation $\tau s, \tau' s'$	Energy
$K \downarrow, K' \uparrow$	$-\Delta_{ m SO}+E_{ m C}+\Delta_{ m S,AS}+\Delta_{ m AS,S'}$
$egin{array}{c} K \downarrow, K \uparrow \ K \uparrow, K' \uparrow \ K \downarrow, K' \downarrow \ K' \uparrow, K' \downarrow \end{array}$	$E_{\rm C} + \Delta_{ m S,AS} + \Delta_{ m AS,S'}$
$K\uparrow,K'\downarrow$	$\Delta_{\rm SO} + E_{\rm C} + \Delta_{\rm S,AS} + \Delta_{\rm AS,S'}$

Next we consider two-electron states with one electron in the lowest shell ($\nu = 1$) and one electron in the first excited shell ($\nu' = 2$). By a similar procedure, sixteen distinct Slater determinants can be written. However, some of them are coupled by electron interactions ["INT" in Fig. 28(b)], resulting in the splitting $\Delta_{AS,S'}$ and modifying $\Delta_{S,AS}$ (see Sec. VII.B). The eigenstates of the electron interactions are the ten antisymmetric states [AS(0, 2)] and six symmetric states [S'(0, 2)] formed by a linear combination of the Slater determinants. These eigenstates are listed with their energies in Table IV.

As for S(0, 2), the energies at B = 0 are found by adding the single-particle energies, charging energy, and interaction energy. However, the expressions for the explicit states now contain four terms. For the spin and valley polarized states $(\tau s, \tau s)$, these four terms reduce to a simple two-term Slater determinant, but for the remaining six AS(0,2) and six S'(0,2) states this is not the case due to the interactions $\Delta_{AS,S'}$. This tells us that in general Eq. (26) is too simple to accurately predict the spectrum. In particular, it does not take magnetic fields into account, nor differences in spin-orbit coupling and valley scattering between different shells, not to mention short- or long-range interactions discussed in Sec. VII. As an example, the energy and quantum state of the spin-valley-unpolarized AS(0,2) state denoted by $K \downarrow, K' \uparrow$ are

$$E_{K\downarrow,K'\uparrow}^{AS(0,2)} = -\Delta_{\rm SO} + E_{\rm C} + \Delta_{S,AS},\tag{29}$$

$$\begin{split} |\psi_{K\downarrow,K'\uparrow}^{AS(0,2)}\rangle &= \frac{1}{2} (|1K\downarrow\rangle_1 |2K'\uparrow\rangle_2 - |2K\downarrow\rangle_1 |1K'\uparrow\rangle_2 \\ &+ |1K'\uparrow\rangle_1 |2K\downarrow\rangle_2 - |2K'\uparrow\rangle_1 |1K\downarrow\rangle_2). \end{split}$$
(30)

The longitudinal antisymmetry of this state is easily seen by comparing terms in the same row of Eq. (30), while comparing between rows shows the symmetry of the spin-valley part. The spin-valley-unpolarized S'(0, 2) state (also denoted $K \downarrow, K' \uparrow$) involves the same single-particle states, but is obtained by changing the longitudinal symmetry, i.e., by changing the sign on the second and third terms in Eq. (30).

All relevant two-electron states in (1,1) can also be constructed from Table IV, simply by identifying S(1,1) with S'(0,2), and AS(1,1) with AS(0,2). This works by assigning $\nu = 1(2)$ to the lowest shell in the right (left) dot and setting level spacing and interactions to zero. We call antisymmetric (1,1) states "blocked" states, because they vanish formally when setting $\nu = \nu' = 1$. Physically, this means that these states cannot be converted into (0,2) charge states without involving a higher orbital in the right dot or changing their spin or valley configuration. The (1,1) states generated in this fashion [along with the (0,2) states] from Table IV are useful basis states for a tunnel-coupled double dot at B = 0, assuming that the spin-orbit coupling in the two dots is identical.¹³

Having introduced all relevant (1,1) and (0,2) multiplets, a useful overview of the states and their mutual coupling is obtained by setting out the Hamiltonian in matrix form. Figure 29(a) shows schematic matrices for a conventional semiconductor double dot in the spin-singlet or spin-triplet basis, and Fig. 29(b) shows a nanotube double quantum dot in the basis of the longitudinal symmetric or antisymmetric multiplets. These matrices can be divided into submatrices coupling manifolds of particular symmetry. In conventional semiconductors all submatrices are diagonal even when interactions are included. Tunnel coupling between states of the same singlet or triplet character is reflected (+ symbols) in the submatrices between (1,1) and (0,2) states. In

¹³In (1,1), the splitting between symmetric states S(1,1) and antisymmetric states AS(1,1) vanishes for small interdot tunneling $\Delta_{AS,S'}^{(1,1)} \rightarrow 0$, and hence a different basis would be used, appropriate for the regime $|\Delta_{SO}^{left} - \Delta_{SO}^{right}| \gg \Delta_{AS,S'}^{(1,1)}$. Similarly, different orbital moments in left and right dots require a different basis at sufficiently large magnetic field.

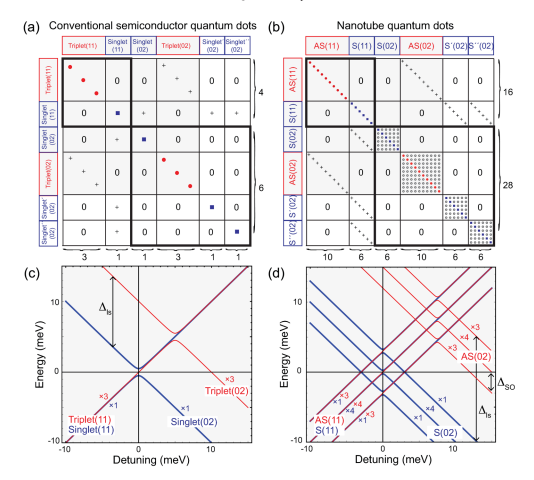


FIG. 29 (color online). (a) Matrix representing the two-electron Hamiltonian for a conventional semiconductor double dot, assuming a single orbital on the left and two orbitals on the right. The basis states are spin-singlet and spin-triplet states as indicated, and symbols indicate nonzero matrix elements. The submatrices in the (1,1) and (0,2) subspaces are all diagonal, and interdot tunneling (+) leaves singlet and triplet states uncoupled. To lowest order, interactions modify the diagonal elements between Triplet (0,2) and Singlet' (0,2)eigenstates (\blacksquare and \bullet), corresponding to the interaction energy $\Delta_{AS,S'}$ discussed in the text. For completeness, we include the higher-lying Singlet" (0,2). (b) The analogous matrix for a nanotube double quantum dot with spin-orbit interaction, assuming a single longitudinal shell on the left and two shells on the right. The basis states are those of Table IV; hence longitudinally symmetric S(1,1) and antisymmetric AS(0,2) states are uncoupled. Most submatrices are still diagonal, but interactions are expected to result in significant off-diagonal matrix elements (\bigcirc) within the (0,2) sectors. For example, Sec. VII shows that valley backscattering leads to off-diagonal elements between spin-valley-unpolarized states within the S(0,2) subspace. Without going into details of the interactions, all offdiagonal elements marked \bigcirc are in principle allowed, as drawn here. In some cases, a detailed analysis shows that particular interaction matrix elements vanish. (c) Energy vs detuning of the shaded states in (a), neglecting interactions. Zero detuning corresponds to the degeneracy between the ground states of (0,2) and (1,1). Avoided crossings due to interdot tunneling occur for states with the same symmetry. The degeneracy of the levels is indicated. Numerical parameters in this plot were chosen for illustration and differ in actual devices. (d) The same plot for a nanotube, assuming identical spin-orbit coupling (Δ_{SO}) in all shells. Interdot tunneling now leads to anticrossings between states with the same longitudinal symmetry. The effect of interactions on S(0, 2), and AS(0, 2) is neglected in this plot; see the discussion of Figs. 30, 31, and 51 for details.

nanotubes, the overall structure is similar, but the number of states in each multiplet is increased and the diagonal elements now include spin-orbit coupling. Again, states with identical symmetry are coupled by diagonal tunnel matrices, and weak interactions (denoted $\Delta_{AS,S'}$) appear as diagonal elements that shift the energies of multiplets with respect to each other. However, off-diagonal elements are allowed in the (0,2) multiplets (\bigcirc symbols), although arguments can be made that these are small (Appendix B). Interactions within the S(0, 2) multiplet (not included in Table IV) appear as diagonal and off-diagonal matrix elements and are further discussed in Sec. VII.

Plots of energy versus detuning for the low-energy states (Singlet, Triplet, and *S*, *AS*) are shown in Figs. 29(c) and 29(d) neglecting interactions $(\Delta_{AS,S'} \rightarrow 0)$. The more complex spectrum for the nanotube is clearly revealed. Valley mixing (assumed zero in this figure) would lead to additional avoided crossings (off-diagonal elements in the matrix) between states with different valley quantum numbers. The nonavoided crossings between the *AS*(1, 1) and *S*(0, 2) states signify long-lived (1,1) states that give rise to a Pauli blockade.

Tunneling from a blocked AS(1, 1) state to a S(0, 2) state requires a change of the longitudinal symmetry. For some states this may simply involve dephasing between the left and

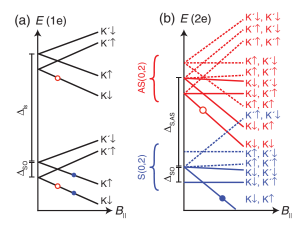


FIG. 30 (color online). Magnetic field dependence of the 16 lowest (0,2) states [cf. Fig. 28(b)]. (a) Spectrum of the two lowest single-particle shells of the right dot, vs parallel magnetic field B_{\parallel} . There are six ways to fill the lowest shell with two electrons, each corresponding to a symmetric state in (b). The magnetic moment of each combination is simply the sum of the involved single-particle spin and orbital magnetic moments, shown here for two examples (colored dots). (b) Magnetic field dependence of the ten antisymmetric and six symmetric states. States that involve the ground state of the lowest shell $K \downarrow$ are plotted with solid lines.

right single-particle states. For other states it also necessitates a change in quantum numbers of at least one electron. In particular, we distinguish lifting of Pauli blockade by

- · dephasing only,
- valley flips,
- spin flips, and
- spin-and-valley flips.

Inspection of all ten *AS* states reveals that a single flip in one of the dots can result in an unblocked double-dot configuration. For some states it involves a spin flip, and for others a valley flip (see examples in Appendix B.10). However, these single flip processes do not conserve energy because of spin-orbit coupling and therefore cannot lift the blockade. Therefore, simultaneous spin and valley flips can become the rate-limiting process ("spin-valley blockade").

To investigate which type of blockade is observed in experiments, careful identification of the involved states and quantum numbers is required. This is best facilitated by application of a magnetic field. In Fig. 30(a), the energies of the two lowest single-particle shells are plotted against a parallel magnetic field. The two-electron magnetic moments are obtained by summing the one-electron magnetic moments [Fig. 30(b)]. As an example, the filled circles in Fig. 30(a) indicate the two single-particle energies summed to give the energy of the S(0, 2) state $K \downarrow, K \uparrow$ as in Fig. 30(b). Similarly, the empty circles indicate the two-single-particle energies combining to give the AS(0,2) state denoted by $K \downarrow, K \downarrow$. The field dependence of the remaining states is found in a similar fashion. The states can be divided into pairs of valleypolarized, spin-polarized, and spin-valley-unpolarized states. Although all degeneracies of the two multiplets are lifted at finite field, and spin and valley quantum numbers can in principle be assigned based on the observed magnetic

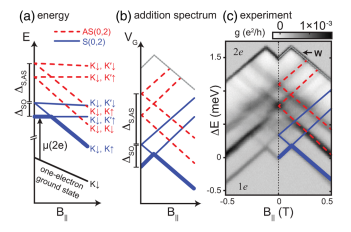


FIG. 31 (color online). Addition spectroscopy of two-electron states in a nanotube single quantum dot, assuming that the oneelectron dot starts in its ground state $K \downarrow$. (a) Two-electron spectrum E(2e) from Fig. 30(b) as a function of the parallel magnetic field. States that do not involve $K \downarrow$ are omitted. (b) Expected 1*e*-2*e* addition spectrum from (a). (c) Measured addition spectrum, with lines from (b) superimposed. The top-most conductance feature (*w*) results from the finite-bias window of 1.7 mV used during measurement (see Fig. 54). The separation between solid and dashed lines is surprisingly small (0.85 meV) compared to the single-particle level spacing (7.8 meV), indicating that $\Delta_{S,AS} \ll \Delta_{Is}$ due to electron-electron interactions. From Pecker *et al.*, 2013.

moment,¹⁴ the spectroscopic intensity associated with each state can be very different, depending how exactly the spectrum is measured. The solid lines in Fig. 30(b) indicate the available two-electron states within the S(0, 2) and AS(0, 2) multiplets, given that one electron occupies the $K\downarrow$ state. These two-electron states are relevant when measuring the addition spectrum given that the first electron is in its finite-field ground state $K\downarrow$.

4. Symmetric and antisymmetric multiplets in nanotubes: Experiment

The two-electron spectrum can be measured by high-bias spectroscopy near the (0,1)-(0,2) transition. Assuming that the device starts in the (0,1) ground state $(K\downarrow)$, then exactly three states of the symmetric two-electron multiplet can be reached and four states of the antisymmetric multiplet [Fig. 30(a)]). All other two-electron states require a higher-order process, in which the incoming electron also promotes the resident electron to change its spin or valley quantum numbers. The expected addition spectrum is shown in Fig. 31(b), and the corresponding data in Fig. 31(c). Although the qualitative agreement is quite good, the data show a surprisingly small multiplet splitting $\Delta_{S,AS} = 0.85$ meV, significantly less than the measured single-particle level spacing of 7.8 meV. This is due to strong electron-electron correlations (Sec. VII) that differ drastically from the weak exchange interactions typical in GaAs dots. This is of practical importance because a small

¹⁴This requires that interactions have lifted the degeneracy with the S'(0, 2) multiplet not included in Fig. 30(b).

splitting $\Delta_{S,AS}$ makes it difficult to observe Pauli blockade in nanotubes.

5. Pauli blockade in nanotubes: Experiment

Generally interdot transitions that are forbidden by spin or valley selection rules are of particular interest, because measurements of leakage current or double-dot charge state then illuminate spin dynamics and spin-valley relaxation processes. We return to this topic in Sec. V. Although aspects of Pauli blockade in nanotubes have been observed by multiple groups, this phenomenon is experimentally less generic than in GaAs. A detailed understanding of the data is complicated by the large number of states involved (see Fig. 29), and often a lack of knowledge of critical device parameters such as differences in intervalley scattering or spin-orbit coupling between left and right dots, and the strength of electron-electron correlations. All these were neglected in Fig. 29.

Even the simplest manifestation of Pauli blockade, namely, Pauli rectification in a dc transport experiment, can be obscured by other effects such as strong electron-electron correlations. In Fig. 27 we outlined the double-dot energy levels for the $(1, 1) \rightarrow (0, 2)$ transition and noted the importance of sufficiently large $\Delta_{S,AS}$. In the (n, n) and (p, p)regimes, $\Delta_{S,AS}$ is limited by the level spacing from the longitudinal quantization Δ_{ls} and likely significantly reduced by correlation effects $\Delta_{AS,S'}$. In order to keep $\Delta_{S,AS}$ as large as possible, the band gap of the nanotube can be used as an effective large "level spacing," making the observation of Pauli blockade more robust against interaction effects. The transition between (3h, 1e) and (2h, 0) [dashed circle in Fig. 24(b)] is one example where the level spacing is enhanced by the band-structure gap E_{G} .

As shown in Fig. 32, rectification behavior for such a transition is observed for detunings as high as the applied bias voltage (± 10 mV). This is larger than the spin-orbit splitting and estimated level spacing in this device (Pei *et al.*, 2012). This observation of strong current suppression up to high bias can be linked to the advantageous use of the band gap [see Fig. 33(a)].

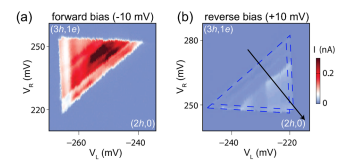


FIG. 32 (color online). Pauli rectification (experiment). Current in the bias triangles measured in (a) forward and (b) reverse directions near the (3h, 1e)-(2h, 0) transition. The asymmetry due to Pauli rectification is clearly observed and persists up to ± 10 meV, i.e., detunings much larger than Δ_{SO} (= 1.6 meV in this device). The arrow defines the detuning axis used in Fig. 33(c). Adapted from Pei *et al.*, 2012.

In the blocked bias triangle of Fig. 32(b), a small increase of leakage current is observed at a detuning of approximately $2 \times \Delta_{SO} = 3.2$ meV. This can be interpreted as a weak lifting of Pauli blockade, but a quantitative understanding of this leakage current, and identification of the corresponding relaxation rates, has not been reached. We speculate that it is necessary to include interaction effects beyond the constant-interaction model to explain such features.

6. Spin-valley blockade

Information about the role of spin and valley quantum numbers in the Pauli rectification of Fig. 32 can be obtained by applying a parallel magnetic field. If this induces an energy-level splitting larger than the interdot tunnel coupling or the intervalley scattering, we expect orbital and Zeeman couplings to restore valley and spin quantum numbers within the two-electron states. In turn we can associate these with the quantum numbers of single-particle levels.

In Fig. 33(c), the leakage current is measured as a function of detuning [defined in Fig. 32(b)] and B_{\parallel} . The base of the bias triangle (detuning $\delta \sim 0$) corresponds to the ground-state-to-ground-state transition between (3h, 1e) and (2h, 0). Energy conservation imposes spin and valley selection rules for this transition, and therefore the leakage current provides information about the relaxation of these selection rules.

Figure 33(a) shows the assignment of single-particle quantum numbers to the highest (lowest) longitudinal shell in the valence (conduction) band of the left (right) quantum dot. These quantum numbers were inferred from the magnetic field dependence of the stability diagram.¹⁵ Based on the single-particle picture, the ground state of (2h, 0) is expected to make a transition from $K'\uparrow$, $K\downarrow$ to $K'\uparrow$, $K'\downarrow$ as a function of the parallel magnetic field.¹⁶ This implies that at low field the ground-state-to-ground-state transition requires a spin flip of the right electron, whereas at higher magnetic field it requires a spin flip and a valley flip [compare the two arrows in Fig. 33(b)]. Therefore, if valley is a good quantum number and conserved during interdot tunneling, then one expects the ground-state leakage current at high field to be smaller than at low field. Indeed, this is seen in Fig. 33(c) by comparing conductance features marked by line 1 and line 2. We mention that the (3h, 1e) ground state is not a blocked state in the sense of Sec. IV.C.3: We can write it as a Slater determinant of one K' electron in the left (L) orbital and one K electron in the right (R) orbital, $|LK'\uparrow\rangle_1 |RK\uparrow\rangle_2 - |RK\uparrow\rangle_1 |LK'\uparrow\rangle_2$, which does not vanish under the substitution $R \rightarrow L$ (i.e., $\nu = \nu'$). We therefore expect that relaxation to the (2,0) charge state is allowed without involvement of higher orbitals or change of

¹⁵To avoid confusion, we show quantum numbers of electronic states, even for the valence band. Other publications may consider the absence of a $K'\uparrow$ electron in the valence band as a $K'\downarrow$ hole, due to conservation of angular momentum. Quantum numbers in Pei *et al.* (2012) and Steele *et al.* (2013) are not consistent with our identification.

¹⁶Table IV should not be used to generate these states, because the conduction band and the valence band differ in their assignment of K and K' (Fig. 18).

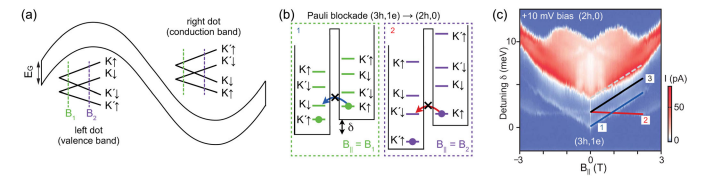


FIG. 33 (color online). Different levels of Pauli blockade involving spin and valley quantum numbers for the charge transition $(3h, 1e) \rightarrow (2h, 0)$. (a) Single-particle electron levels vs the magnetic field, sketched for the left and right sides of a *pn* double dot. (b) At low field [marked B_1 in (a)] the ground-state-to-ground-state transition requires a spin flip. At higher field (marked B_2) the ground-state transition requires a spin flip and a valley flip. The detuning δ between left and right dots is marked. Because of the different magnetic moments of the single-particle states, the value of δ at which each transition becomes resonant depends on magnetic field. (c) Measurement of the leakage current vs B_{\parallel} . Some of the features can be identified within the single-particle representation of (b) (lines 1 and 2). The slight reduction of the ground-state leakage current above 1 T indicates that a spin and valley flip (line 2) is slightly less frequent than a spin-only flip (line 1), and is evidence for a spin-valley blockade. The absence of a simple charge relaxation ($RK\uparrow$) \rightarrow ($LK\uparrow$) (line 3) and the onset of leakage current at $\delta \sim 2\Delta_{SO}$ (dashed line) are currently not understood. Adapted from Pei *et al.*, 2012.

spin or valley configuration. The absence of this charge relaxation [line 3 in (c)] is not understood.

In summary, these data suggest that both spin and valley can contribute to Pauli blockade, but a quantitative understanding of the leakage current and relaxation rates has not been reached. We speculate that several mechanisms contribute, such as disorder, hyperfine coupling, spin-phonon coupling, or bend and spin-orbit mediated relaxation.

D. Lifting of Pauli blockade by hyperfine coupling

In this section we investigate double dots where Pauli blockade is partially lifted due to hyperfine interaction between the electron spin and the ¹³C nuclear spins.

1. Theory

Hyperfine interaction with disordered nuclear spins, such as ¹³C isotopes, couples different spin states by flip-flop processes and, in addition, different valley states, because of the atomically sharp length scale. It is therefore expected that hyperfine coupling generically lifts Pauli blockade and results in spin relaxation and spin dephasing processes. This

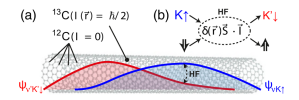


FIG. 34 (color online). (a) Two localized electronic wave functions with different valley, longitudinal, and spin quantum numbers. Hyperfine interaction (HF) with a ¹³C nuclear spin (fat arrow) can scatter between them, because of the local nature of the scattering vertex (b). This mechanism allows lifting of spinand/or valley-based Pauli blockade, if conditions of energy conservation are fulfilled.

mechanism of spin relaxation was considered by Semenov, Kim, and Iafrate (2007), whose numerical estimates predicted a spin relaxation time of ~1 s. Relevant for quantum dot experiments in nanotubes, Yazyev (2008) and Fischer, Trauzettel, and Loss (2009) inspected the role of dipolar and Fermi contact interaction in *sp*-hybridized nanostructures, resulting in an interesting interplay between isotropic and anisotropic hyperfine interactions.

In the tight-binding picture of Pályi and Burkard (2009), the hyperfine interaction with ¹³C is modeled by a matrix element that arises on the site of each nuclear spin (Fig. 34). Because it acts locally at each atomic lattice site that contains a ¹³C, the hyperfine interaction couples not only to the electron spin, but also mixes the valley index (cf. discussion of valley scattering in Fig. 6). The strengths of the valley-conserving and valleymixing parts of the effective hyperfine coupling were estimated to have the same order of magnitude. Pauli blockade, even in its strongest form protected by spin and valley (Sec. IV.C.6), can therefore be lifted by the presence of ¹³C atoms. Pályi and Burkard (2009) considered the situation where valley scattering is dominated by hyperfine coupling. Ignoring spin-orbit coupling, they showed that the leakage current in the Pauli blockade regime of a double quantum dot is only strongly suppressed if both valley and spin splittings are larger than the hyperfine coupling.

The other limit, where valley scattering is dominated by disorder-induced valley scattering, was considered subsequenctly (Pályi and Burkard, 2010). Ignoring hyperfine coupling, but assuming strong spin-orbit interaction (larger than the splittings due to disorder and interdot tunneling), it was predicted that the current in the Pauli blockade regime can show a dip at low fields. Although similar to experimental data discussed in Fig. 35(d), the amplitude of the predicted dip is orders of magnitude smaller than observed. In the theory, the low-field dip occurs because a difference in the valley coupling splittings diminishes the matrix element for tunneling, while at high fields the valley mixing is suppressed.

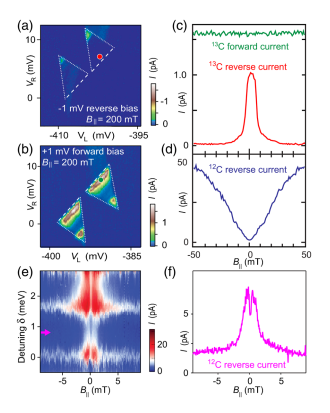


FIG. 35 (color online). (a) Current through a top-gated multielectron double dot formed in a narrow-gap ¹³C nanotube. Near the base of the bias triangles (ground-state-to-ground-state transition) no current is observed for reverse bias. (b) Current at forward bias. (c) Magnetic field dependence of current in reverse and forward bias measured near the base of a triangle [dots in (a) and (b)]. The peak in reverse current at $B_{\parallel} = 0$ is attributed to hyperfine-mediated relaxation. (d) In a similar device formed in a predominantly ¹²C nanotube, the opposite magnetic field dependence is observed: The reverse current shows a minimum at $B_{\parallel} = 0$, presumably a consequence of Van Vleck cancellation. (e) Leakage current in a different device with natural abundance of ¹³C vs detuning and B_{\parallel} . This peak, measured in the spin-valley blockade regime of Fig. 33(c), is narrower than in (c), possibly due to the smaller concentration of ¹³C. (f) Horizontal cut through (e) at location of the arrow, showing a small dip at $B_{\parallel} = 0$. (a)–(d) Adapted from Churchill, Bestwick et al., 2009. (e), (f) From Pei et al., 2012.

2. Experiment

Figure 35 shows electron transport through weakly tunnelcoupled, Pauli-blocked double dots. Figures 35(a) and 35(b) show the asymmetry in forward and reverse bias for a nanotube enriched with ¹³C, similar to the Pauli-blocked ¹²C device discussed in Fig. 32. Figure 35(c) compares the magnetic field dependence of the reverse-bias leakage current near the base of the triangle with the forward current in the same tuning. Whereas the forward current is independent of applied magnetic field (indicating that the rate-limiting tunnel barrier $\Gamma_{\rm M}$ is independent of the magnetic field), the reverse leakage current is strongly suppressed only above a characteristic magnetic field scale $B_{\rm C} \sim 6$ mT [Fig. 35(c)].

Churchill, Bestwick *et al.* (2009) attributed the current peak at B = 0 to spin relaxation via electron-nuclear flip-flops,

similar to the situation in GaAs double dots and InAs nanowires (Koppens *et al.*, 2005; Nadj-Perge, Frolov, van Tilburg *et al.*, 2010). Because of the mismatch of electron and nuclear magnetic moments, these energy-conserving flip-flop processes are expected to be strongly suppressed once the difference in Zeeman splitting exceeds the strength of the hyperfine coupling B_{nuc} . Estimating the number of nuclei N, electron a factor $a \sim 2 \gg a$ and assuming uniform cou-

electron g factor $g_e \sim 2 \gg g_{nuc}$, and assuming uniform coupling to a Gaussian-distributed Overhauser field $g\mu_B B_{nuc} = \mathcal{A}/\sqrt{N}$, an effective hyperfine coupling constant $\mathcal{A} \sim (1-2) \times 10^{-4}$ eV can be estimated. This is 2 orders of magnitude larger than predicted for nanotubes (Yazyev, 2008; Fischer, Trauzettel, and Loss, 2009) or measured in fullerenes (Pennington and Stenger, 1996).

This puzzling result stimulated theoretical work by Coish and Qassemi (2011), who examined the role of thermally activated spin-flip cotunneling in lifting spin blockade. Their theory does not require large hyperfine coupling constants but predicts a peak width (set by temperature, here approximately 100 mK) that is too large. Matching the width in Fig. 35(c) would require a temperature below 10 mK for g = 2. This large value of the hyperfine interaction remains unexplained. It is, however, consistent with a subsequent measurement of the dephasing time in a ¹³C double-dot device (Churchill, Bestwick *et al.*, 2009), discussed in the next section.

A strikingly different field dependence of the leakage current through a Pauli-blocked ¹²C double dot¹⁷ is shown in Fig. 35(d): The current shows a minimum at B = 0, whose width depends on the interdot tunnel coupling. Such behavior was seen in both ¹²C and ¹³C devices, particularly for stronger interdot tunneling. This is at first sight consistent with the predictions of Pályi and Burkard (2010), which give a peak or a dip depending on specific device parameters. However, the observed ratio of low and high field currents was 50, rather than 1.5 as predicted.¹⁸

An alternative explanation of the large dip is spin relaxation mediated by phonon and spin-orbit interaction. Because spinorbit coupling is even under time reversal, one-phonon processes cannot mediate a coupling between time-conjugate states (so-called Van Vleck cancellation, similar to electric dipole transitions). This leads to suppressed spin relaxation near B = 0 as discussed in Sec. V.A (Van Vleck, 1940; Khaetskii and Nazarov, 2001). However, this theory does not lead to a good quantitative fit to the observed dips.

Figure 35(e) shows the leakage current of a different device, namely, the one presented in Fig. 33(c), zoomed in to low fields and small detuning. At first sight, the magnetic field dependence resembles that of Fig. 35(c), with a peak width that is approximately 10 times smaller. Noting that this device had a natural abundance of ${}^{13}C$ (~1%), and that the effective hyperfine coupling scales with the square root of the ${}^{13}C$ concentration, these data corroborate with the hyperfine coupling measured by Churchill, Bestwick *et al.* (2009). On second sight, a small splitting in the leakage current is

¹⁷That is, with natural abundance ¹³C.

¹⁸The observed peak width in the regime of Fig. 35(c) cannot be explained by this theory either, because it did not depend on interdot tunneling, whereas the predicted peak width does.

evident near B = 0 [Fig. 35(f)], indicating that a full understanding of this system has not yet been reached. As discussed by Pei *et al.* (2012), the splitting of the peak could arise from combinations of exchange interaction, spin-orbit coupling, and hyperfine coupling.

Looking beyond nanotubes, we note that dips and peaks of the leakage current at B = 0 have been observed and discussed in InAs nanowires (Pfund *et al.*, 2007; Nadj-Perge, Frolov, van Tilburg *et al.*, 2010) and silicon quantum dots (Lai *et al.*, 2011; Yamahata *et al.*, 2012).

E. Open questions

- The reproducibility of quantum properties and the variability of device characteristics among nanotubes of identical chirality has yet to be established. It is experimentally unverified to what extent right-handed and left-handed species (i.e., inversion isomers of chiral nanotubes) display the same properties. Similarly, the robustness of the valley index (isospin) for armchairlike and zigzaglike nanotubes has not been checked experimentally.
- Although Pauli blockade has been observed in different device geometries by several groups, it is not as well established as in III-V double dots. Several nanotube double dots showed no Pauli blockade,¹⁹ which can be attributed to differences in dielectric surrounding and interaction effects or disorder.
- Relaxation times associated with valley, spin, or combined spin-valley relaxation in single and double quantum dots have not yet been measured systematically, including as a function of a magnetic field. Existing experiments are discussed in Sec. V.
- The type of hyperfine interaction (Fermi contact term versus dipolar) has not been studied experimentally. The unexplained strength of hyperfine interaction inferred from one study by Churchill, Bestwick *et al.* (2009) remains to be confirmed. The lifting of Pauli blockade near B = 0 may have alternative explanations, but we are not aware of any that are consistent with experimental conditions. The short dephasing time measured in ¹³C devices (Churchill, Kuemmeth *et al.*, 2009) is consistent with a large hyperfine coupling, but may originate from mechanisms unrelated to a hyperfine coupling. Laird, Pei, and Kouwenhoven (2013) measured a comparatively short dephasing time in predominantly ¹²C nanotubes.
- In Appendix B.10 we argue that Pauli blockade protected by "spin and valley" strictly speaking does not exist. However, a single spin flip or a valley flip does not conserve energy due to spin-orbit coupling, possibly making spin and valley flips the dominant relaxation process. This underlines the importance of understanding both spin and valley for any quantum device based on carbon nanotubes.

V. SPIN-VALLEY COHERENCE

By studying the decay of spin and valley states, we can use them as delicate probes of their environment. In this section, we discuss different ways for these states to decay and show how they can be used as quantum bits. We focus especially on the interactions of electron spins with phonons and with magnetic nuclei.

We discuss three distinct decay processes of quantum states, known as relaxation, dephasing, and decoherence. Relaxation (characterized by time T_1) describes the equilibration of population between two quantum states. Dephasing (characterized by time T_2^*) describes the loss of phase information in a quantum superposition. The main mechanism by which this happens is through fluctuations of the quantum energy splitting leading to accumulation of random phases. Decoherence describes the loss of phase information when slowly varying fluctuations are removed by dynamical decoupling. For the simplest decoupling scheme, Hahn echo (Sec. V.C.4), this is characterized by a time T_{echo} which is generally longer than T_2^* . For a fuller discussion of spin qubits in single and double quantum dots, see Hanson et al. (2007) and Ihn (2010), which extensively reference the many experiments in other materials (principally GaAs).

A. Spin and valley coupling to phonons

1. Theory

Because the mechanical motion of nanotubes perturbs the confining potential of quantum dots, it couples distinct electron charge states. Through the spin-orbit interaction, spin-valley states are also coupled to mechanical motion. This is most clearly evident as a relaxation channel for spin-valley states; excited states can decay by phonon emission, with a rate that depends on the coupling strength and the phonon density of states.

There are four types of phonon mode in nanotubes: radial breathing, twist, longitudinal, and bending modes (Mariani and von Oppen, 2009). These couple to spin-valley states through two general coupling mechanisms. Deformationpotential coupling perturbs the band structure and, combined with spin-orbit coupling, induces spin flips (Bulaev, Trauzettel, and Loss, 2008). Deflection coupling changes the alignment of the nanotube to the magnetic field, thereby coupling spin and valley through the anisotropy of the valley magnetic moment (Borysenko, Semenov, and Kim, 2008; Rudner and Rashba, 2010). Although deformation-potential coupling is present for all four mode types, deflection coupling arises only from bending modes. Nevertheless, deflection coupling is calculated to be the dominant mechanism for phonon-mediated spin relaxation at low energy (Rudner and Rashba, 2010).

Considering these mechanisms, several statements can be made about the expected T_1 between different valley-spin states as a function of magnetic field:

(1) Relaxation between time-conjugate states is suppressed at low magnetic fields due to Van Vleck cancellation (Khaetskii and Nazarov, 2001). This is a consequence of time-reversal symmetry and applies to relaxation within a Kramers doublet.

¹⁹See, for example, Mason, Biercuk, and Marcus (2004), Gräber *et al.* (2006), Sapmaz, Meyer *et al.* (2006), Jørgensen *et al.* (2008), and Jung *et al.* (2013).

- (2) Relaxation between non-time-conjugate states occurs fastest when they are close together in energy. The reason is that the dispersion relation for bending-mode phonons ω(k) ∝ k² leads to a density of states dk/dω ∝ 1/√ω which is maximal at ω → 0. This is in contrast to higher-dimensional systems, where the density of states is constant or increases with energy (Bulaev, Trauzettel, and Loss, 2008; Rudner and Rashba, 2010).
- (3) The relaxation rate between two states is a nonmonotonic function of their energy splitting, owing to interference between different contributions to the electron-phonon coupling. Interference is predicted between contributions from discrete and continuous phonon modes, as well as due to the match or mismatch of phonon wavelength with the wavelength of a confined electron. These interference oscillations should be evident in the dependence of T_1 on a magnetic field (Bulaev, Trauzettel, and Loss, 2008).

2. Experiment

Spin-valley relaxation has been measured in the device of Figs. 3(a) and 3(b). This device, fabricated from a ¹³C

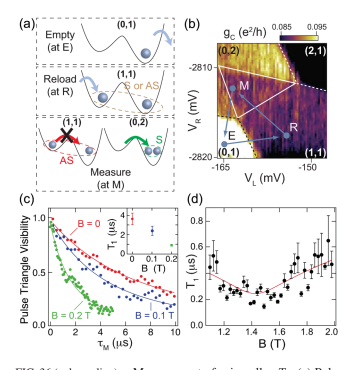


FIG. 36 (color online). Measurement of spin-valley T_1 . (a) Pulse cycle for measuring relaxation from *AS* to *S* states (see text). (b) Stability diagram close to the (1,1)-(0,2) transition with pulse cycle $E \rightarrow R \rightarrow M \rightarrow E$ and $\tau_M = 0.5 \ \mu$ s. Color represents time-average charge-sensor conductance. Inside the "pulse triangle" (marked), Pauli blockade leads to an occupancy of (1,1) for metastable *AS* states, yielding a reduced sensor conductance. (c) Decay of pulse triangle visibility (points) as a function of τ_M , measured for three magnetic field values. Lines are fits from which T_1 values (inset) are extracted. (d) Points: T_1 as a function of *B* close to the upper spin-orbit anticrossing $(K'\uparrow - K'\downarrow)$. Line: fit of the form $T_1 \propto \sqrt{\Delta E}$. Adapted from Churchill, Kuemmeth *et al.*, 2009.

nanotube, incorporates a double quantum dot and a nearby charge sensor, coupled via a floating antenna, whose conductance is sensitive to the charge occupancy of the double dot. Relaxation is studied by preparing an AS two-electron state, Pauli blocked in (1,1), and using the charge sensor to monitor the time to decay to an unblocked S state (Churchill, Kuemmeth *et al.*, 2009).

The two-electron state is manipulated using a cycle of gate voltage pulses applied to gates L and R [Fig. 3(b)], to switch the dot potentials between different configurations [Figs. 36(a) and 36(b)]. The cycle (Johnson et al., 2005) begins with the device configured at point E in gate space, where tunneling to the leads prepares the (0,1) configuration. The device is then pulsed to point R in (1,1), where an electron is reloaded into the left dot. Because $\Delta_{S,AS}$ is small in the (1,1) configuration [left side of Fig. 29(d)]), the two-electron state after reloading can be either S or AS. For readout, the device is quickly pulsed to point M, corresponding to the right side of Fig. 29(d), where the ground-state configuration is S(0,2). Here a symmetry-to-charge conversion occurs; if the prepared twoelectron state was S, the left electron will tunnel to the right dot, leading to (0,2) occupancy. However, for AS states, Pauli blockade enforces occupancy (1,1). This persists for a time $\sim T_1$, until spin-valley relaxation²⁰ causes the AS state to decay to an S state, whereupon the device relaxes to (0,2) occupancy.

The time-average charge-sensor conductance g_C is monitored with this pulse cycle applied continually. The duration τ_M of the third step is chosen to be much longer than that of the others, so that g_C predominantly reflects the average occupancy at M. For $\tau_M \ll T_1$, relaxation of the blocked states is negligible, resulting in a large admixture of (1,1) occupancy and corresponding reduced g_C in the "pulse triangle" region of the stability diagram [Fig. 36(b)]. For $\tau_M > T_1$, this admixture is reduced. By fitting the pulse triangle visibility [defined as the difference between measured g_C and the value expected for (0,2), normalized to unity at $\tau_M \rightarrow 0$] as a function of τ_M , the time T_1 can be deduced [Fig. 36(c)].

As a function of the magnetic field directed approximately along the nanotube, T_1 is observed to decrease initially, consistent with prediction (1) above [Fig. 36(c) inset]. However, T_1 shows a minimum at $B \approx 1.4$ T [Fig. 36(d)], where the two K' states with opposite spin approach each other [as in Fig. 21(d)]. This is consistent with prediction (2) above, assuming that during step R an electron is sometimes loaded into a K' state. Neglecting substrate interaction, the relaxation rate is expected to be proportional to the phonon density of states in the nanotube, giving $T_1 \propto 1/\sqrt{\Delta E}$, where ΔE is the energy difference between the two K' states; taking the proportionality constant as a fit parameter and using the measured field misalignment and Δ_{SO} for this device to calculate ΔE , this prediction is found to be in good agreement with the data (Churchill, Kuemmeth *et al.*, 2009).

The interference oscillations of prediction (3) have not yet been reported. One reason may be that they are sensitive to the confinement potential. Whereas hard-wall confinement

²⁰We define spin-valley relaxation as relaxation between an AS and an S state. As discussed in Sec. IV.C.3, this can involve a flip of spin, valley, both, or neither (i.e., dephasing only).

should give rise to sharp interference maxima of T_1 due to strongly varying overlap of electron and phonon wave functions with energy, a soft confinement typical of fewelectron devices is expected to lead to less pronounced maxima (Bulaev, Trauzettel, and Loss, 2008).

B. Hyperfine mixing of spin states

1. Theory

As well as electron-phonon interaction, a major influence on spin evolution in semiconductors is hyperfine interaction with uncontrolled lattice nuclear spins (Hanson *et al.*, 2007). An electron in a quantum dot interacts with all of the nuclei with which its wave function overlaps; this is equivalent to an effective Zeeman field B_{nuc} that fluctuates slowly about zero due to nuclear spin diffusion. Approximating an equal overlap with all nuclei in the quantum dot, each root-mean-square component of this field is $B_{nuc} = \sqrt{N_{13}} \mathcal{A}/N_{\Sigma}g_{s}\mu_{B}$, where \mathcal{A} is the hyperfine constant and N_{Σ} is the number of nuclei in the dot of which N_{13} are ¹³C. As discussed in Sec. IV, the local nature of the hyperfine interaction can cause spin relaxation. Here we explain that this uncontrolled field also constitutes a major source of spin dephasing (Taylor *et al.*, 2007), and its strength can be deduced by measuring T_2^* .

2. Experiment

Electron spin dephasing was studied in the same device of Fig. 3(b). This was synthesized using 99% 13 C, making

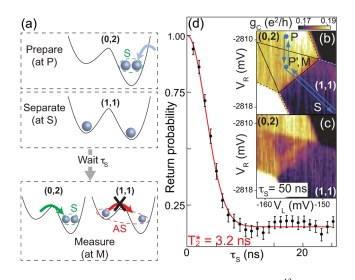


FIG. 37 (color online). Measurement of T_2^* in a ¹³C double quantum dot. (a) Pulse cycle to measure mixing between *S* and *AS* states (see text). (b) Stability diagram close to the (1,1)-(0,2) transition, measured via time-averaged charge-sensor conductance. Gate settings at the three steps of the pulse cycle are indicated *P*, *S*, and *M*. (The step at *P'*, not discussed here, was inserted to reduce pulse overshoot.) The triangle marks the region where *AS* states are Pauli blocked. (c) Stability diagram with pulses applied and $\tau_S = 50$ ns. The triangle of reduced conductance indicates mixing of *S* and *AS* states leading to higher probability of (1,1) during the measurement part of the cycle. (d) Points: Return probability as a function of τ_S , deduced from g_C . Line: Gaussian fit giving $T_2^* = 3.2$ ns. Adapted from Churchill, Kuemmeth *et al.*, 2009.

hyperfine effects particularly strong. Dephasing was detected through the mixing of S and AS states with the two electrons in separate quantum dots (Churchill, Kuemmeth *et al.*, 2009). Because the random hyperfine field is in general different between the dots, the separated spins precess at different rates, mixing S and AS states.

The pulse scheme for this experiment [Figs. 37(a)-37(c)] first configures the device in (0,2) at point *P*, where large $\Delta_{S,AS}$ causes tunneling to the leads to prepare an *S* state. By tilting the potential into (1,1) [point *S*, corresponding to the left of Fig. 29(d)], the electrons are separated for a time τ_S , during which precession in the hyperfine effective field can mix *S* and *AS* states. Finally, the gate voltages are pulsed back toward a (0,2) configuration [point *M*, corresponding to the right of Fig. 29(d)] for measurement. As in Fig. 36, *S* states relax to (0,2) occupancy, whereas *AS* states remain blocked in (1,1). From the sensor conductance g_C in the measurement configuration, averaged over many cycles, the probability of return to (0,2), and hence the degree of *S*-*AS* mixing during the separation step can be deduced (Petta *et al.*, 2005).

As a function of τ_s , the return probability $P(\tau_s)$ decays with characteristic time $T_2^* = 3.2$ ns [Fig. 37(d)], saturating at a value $P(\infty) \approx 0.17$. Attributing the observed T_2^* solely to the difference of the hyperfine effective field between dots, the effective hyperfine field is given by $B_{\text{nuc}} = \hbar/g_s \mu_B T_2^* = 1.8 \text{ mT}$. This is within a factor 2 of the value deduced from Fig. 35 ($B_{\text{nuc}} = 3.5 \text{ mT}$). However, this apparent agreement again suggests a hyperfine constant \mathcal{A} that is 2 orders of magnitude larger than expected theoretically (Yazyev, 2008; Fischer, Trauzettel, and Loss, 2009).

The long- $\tau_{\rm S}$ saturation value of the return probability $P(\infty)$ should reflect the level diagram of Fig. 29(d). Assuming a large longitudinal level spacing, so that only one shell in each dot needs to be considered, the (1,1) configuration allows 16 states, but the lowest manifold in the (0,2) configuration only six. If mixing is fully incoherent, the saturation probability will then be $P(\infty) = 6/16 = 0.375$. If mixing is coherent, $P(\infty)$ will generally be higher.²¹ The case of a clean nanotube ($\Delta_{KK'} = 0$ but including spin-orbit coupling) was analyzed by Reynoso and Flensberg (2011), who calculated the value for a range of specific cases. Depending on whether the system is prepared in its ground state, whether passage through the anticrossings in Fig. 29(d) is adiabatic, and depending on the strength of the magnetic field, $P(\infty)$ can be enhanced as high as unity. Including valley mixing makes the situation even more complicated because $\Delta_{KK'}$ may differ between the dots. This gives rise to new avoided crossings in the level scheme of Fig. 29(d). The speed at which these crossings are passed, set by the detuning sweep rate, is of critical importance (Ribeiro et al., 2013; Ribeiro, Petta, and Burkard, 2013). In the simplest case, where the ground state is always prepared and the first crossing is adiabatic, a value $P(\infty) = 1/3$ is predicted, with corrections due to nonadiabaticity always positive (Reynoso and Flensberg, 2012). The

²¹For example, in conventional semiconductors incoherent mixing gives $P(\infty) = 1/4$ but coherent mixing in a random Overhauser field gives $P(\infty) = 1/3$ or 1/2 depending on the magnetic field (Taylor *et al.*, 2007).

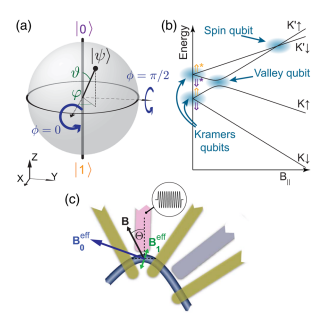


FIG. 38 (color online). (a) Bloch sphere representation of a generic qubit. The qubit state $|\psi\rangle$ is represented by a point with polar coordinates (ϑ, φ) , so that north and south poles correspond to the basis states $|0\rangle$ and $|1\rangle$. In the rotating frame, with Cartesian coordinates (X, Y, Z), the Rabi rotations driven by microwave bursts with phase $\phi = 0, \pi/2$ are marked. (b) Four possible qubits in the spin-orbit coupled energy levels. (c) Qubit device, driven by an oscillating gate voltage. The applied and effective magnetic fields are indicated. Adapted from Flensberg and Marcus, 2010.

measured $P(\infty) = 0.17$ is therefore lower than all theoretical predictions.

C. Qubits

1. Qubit states and the Bloch sphere

A quantum bit, or qubit, is a two-level system that can be controlled in a quantum coherent way (Nielsen and Chuang, 2000). An intuitive way to represent the state of a qubit is as a point on the surface of the Bloch sphere [Fig. 38(a)]. With two orthogonal states of the qubit (for example, two spin states) assigned as the basis states $|0\rangle$ and $|1\rangle$, any superposition can be written

$$|\psi
angle = \cosrac{artheta}{2}|0
angle + e^{iarphi}\sinrac{artheta}{2}|1
angle,$$

where the parameters ϑ and φ are polar coordinates representing that state. Any unitary single-qubit operation then corresponds to a rotation about the origin.

How can such rotations be achieved? One technique is by resonant driving at a frequency corresponding to the energy splitting between qubit states (Hanson *et al.*, 2007). In the case of a spin qubit with gyromagnetic ratio g in static magnetic field **B**₀, a time-varying field **B**₁ cos $(2\pi ft + \phi)$ at driving frequency $f = g\mu_{\rm B}B_0/h$ and perpendicular to **B**₀ induces transitions with Rabi frequency $f_R = g\mu_{\rm B}|B_1|/2h$. In a reference frame rotating with frequency f about the Z axis, the qubit state then precesses at a rate $f_{\rm R}$ about an axis in the XY plane set by the phase ϕ of the driving field. One can achieve arbitrary rotations by concatenating bursts with appropriate phases.

2. Valley, spin, and Kramers qubits

This section identifies various two-level subspaces in the spectrum of a generic, spin-orbit coupled carbon nanotube that can serve as qubits and discusses how they differ in terms of their quantum numbers, ease of operation, and immunity to electrical or magnetic noise. The reason for focusing on these particular subspaces is that the splitting can be made small enough to allow resonant qubit manipulation using microwave fields with frequencies in the range $f \lesssim 40$ GHz that can readily be generated experimentally.

At low field, either Kramers doublet can be used as a qubit subspace, with the two basis states denoted $\{\uparrow, \downarrow\}$ (for one carrier in the shell) and $\{\uparrow^*, \downarrow^*\}$ (for two carriers in the shell) to emphasize the isomorphism with a spin qubit. In a parallel magnetic field and neglecting disorder, these are the eigenstates $\{K'\uparrow, K\downarrow\}$ and $\{K'\downarrow, K\uparrow\}$, but for general field direction or with disorder, they become entangled states of spin and valley. Experimental manipulation of these qubits, known as Kramers qubits or valley-spin qubits, is described in Secs. V.C.3 and V.C.4 (Rohling and Burkard, 2012). Kramers degeneracy guarantees that by reducing the magnetic field the qubit splitting can be made as small as desired.

Two other qubits can be defined that have not yet been realized experimentally. A pure spin qubit can be defined between the two states that cross at $B_{\parallel} = B_{\rm SO}$, i.e., $K' \uparrow$ and $K' \downarrow$ in Fig. 38. For a magnetic field aligned with the nanotube, the energy splitting vanishes at the crossing. However, even quite small misalignment Θ leads to an appreciable splitting Δ_{Θ} . For example, with $\Delta_{\rm SO} = 1$ meV and $\Theta = 1^{\circ}$, the minimum driving frequency is $f = \Delta_{\Theta}/h \approx 4.2$ GHz. Alternatively a pure valley qubit can be defined at the anticrossing between states of the same spin ($K' \uparrow$ and $K \uparrow$ in Fig. 38). However, the minimum driving frequency, set by $\Delta_{KK'}$, can again be substantial, with a typical experimental value $\Delta_{KK'} = 60 \ \mu \text{eV}$ leading to minimum $f = \Delta_{KK'}/h \sim 15$ GHz. More importantly, it may be difficult to control $\Delta_{KK'}$ experimentally.

These various qubits can be manipulated using timevarying electric and magnetic fields. Techniques for driving single-qubit operations have been developed extensively in GaAs quantum dots and many should be applicable to nanotubes (Hanson et al., 2007). Several schemes have been proposed. The conceptually simplest is to use an alternating magnetic field as in Sec. V.C.1. This should work for the spin and Kramers qubits, but not for the valley qubit, because it has no magnetic moment in the perpendicular direction. Since time-varying magnetic fields are hard to generate in nanostructures, schemes have been suggested based on timevarying electric fields. The common principle is that moving the electron back and forth leads to an effective magnetic field mediated by spin-orbit coupling. For example, the Rashbalike coupling discussed in Sec. III.F.4 is equivalent to a momentum-dependent perpendicular magnetic field and can be used to manipulate both spin and Kramers qubits (Klinovaja et al., 2011a). The corresponding Rabi frequency is however rather low ($f_{\rm R} \sim 5$ MHz), making the driving

TABLE V. Comparison of qubit types from Fig. 38. The expected resonant driving mechanisms are marked by a \checkmark ; the mechanisms considered are an oscillating magnetic field and an oscillating electric field E_{ac} combined with extrinsic Rashba spin-orbit coupling, a bend, inhomogeneous static disorder, and inhomogeneous static magnetic field. Only the bend-mediated Kramers qubit (E) has been clearly demonstrated experimentally (Laird, Pei, and Kouwenhoven, 2013).

	$E_{\rm ac}$ combined with				
Qubit	$B_{\rm ac}$	Rashba	Bend	Disorder	Inhomogeneous B
Kramers	\checkmark	\checkmark	$\checkmark^{(E)}$	\checkmark	\checkmark
Spin	\checkmark	\checkmark	\checkmark		\checkmark
Valley				\checkmark	\checkmark^{a}

^aIf *B* varies on a sublattice length scale, e.g., from magnetic impurities.

inefficient in the presence of decoherence. A stronger coupling can be achieved in the presence of a bend, which gives a position-dependent effective magnetic field (Flensberg and Marcus, 2010) and mediates the only coherent control so far clearly achieved in nanotubes (see next section). Similarly, coupling to a true inhomogeneous magnetic field has been suggested (Széchenyi and Pályi, 2014). Another proposal relies on inhomogeneity of the disorder parameter $\Delta_{KK'}$. Although random, if this inhomogeneity is static it should allow driving of both Kramers and valley qubits (Pályi and Burkard, 2011; Széchenyi and Pályi, 2014). These possibilities are summarized in Table V.

As well as the driving mechanism, important considerations are dephasing and decoherence, which limit the coherence time and reduce the fidelity of gate operations. In general, a qubit suffers decoherence through every channel by which it can be driven. As discussed in Secs. V.A and V.B, the dominant decoherence sources in nanotubes are expected to be random time-varying electric fields (e.g., from gate noise and nearby charge switchers) and hyperfine coupling to ¹³C spins, which acts as an effective time-varying magnetic field on individual lattice sites. The coherence properties depend on the strength and power spectrum of the various noise sources.

3. Electrically driven spin resonance in nanotubes

We now focus on the Kramers qubit, which has been experimentally demonstrated. Using bends to mediate qubit control was proposed by Flensberg and Marcus (2010) and realized by Laird, Pei, and Kouwenhoven (2013). It relies on the anisotropic splitting of the Kramers doublets with a magnetic field [Fig. 21(d)]. Each qubit can be regarded as an effective spin 1/2, with spin vector s^* whose components, just like those of the real spin, have eigenvalues defined as ± 1 . Unlike the real spin, the Zeeman splitting of this effective spin depends on the field angle; the parallel and perpendicular components of the **g** tensor are

$$g_{\parallel} = g_{\rm s} \mp \frac{g_{\rm orb} \Delta_{\rm SO}}{\sqrt{\Delta_{KK'}^2 + \Delta_{\rm SO}^2}},\tag{31}$$

$$g_{\perp} = \frac{g_{\rm s} \Delta_{KK'}}{\sqrt{\Delta_{KK'}^2 + \Delta_{\rm SO}^2}},\tag{32}$$

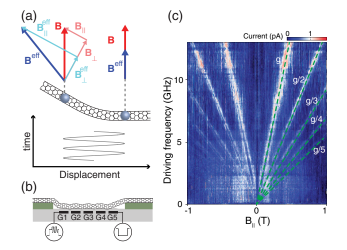


FIG. 39 (color online). (a) Principle of bend-mediated EDSR. An electron driven across a bend experiences a time-varying effective magnetic field which induces coherent qubit precession. (b) Schematic of bent nanotube double quantum dot. Gates G1–G5 define the confinement potential and carry the time-dependent manipulation voltages. (c) Pauli blockade leakage current in a highly disordered device ($\Delta_{KK'} \gg \Delta_{SO}$), showing resonance lines at $g \approx 2$ and at subharmonics. To make the resonance clearer, the mean current at each frequency is subtracted. Adapted from Pei *et al.*, 2012 and Laird, Pei, and Kouwenhoven, 2013.

where upper and lower signs correspond to starred and unstarred doublets, respectively. The effective Zeeman Hamiltonian is then

$$H_{\rm eff} = \frac{1}{2}\mu_{\rm B}\mathbf{s}^* \cdot \mathbf{B}_{\rm eff},\tag{33}$$

where \mathbf{B}_{eff} is an effective magnetic field, defined as the tensor product $\mathbf{B}_{eff} \equiv \mathbf{g} \otimes \mathbf{B}$, about which \mathbf{s}^* precesses. A geometric interpretation of Eq. (33) is shown in Fig. 39(a). When **B** is applied perpendicular to the nanotube (right side of the figure), \mathbf{B}_{eff} is parallel to **B**. However, when **B** is applied at an angle because the nanotube is bent (left side of the figure), the parallel and perpendicular components couple differently, leading to a tilted \mathbf{B}_{eff} .

By applying a microwave electric field, a quantum dot can be driven back and forth across the bend, experiencing an effective magnetic field that contains both a static component and a perpendicular oscillating component.²² Thus the electric field drives transitions between the two qubit states. Because these two states do not have the same spin, this is a form of electrically driven spin resonance (EDSR).

Detection of EDSR is by measuring the current through a double quantum dot configured in a Pauli-blocked configuration [Fig. 39(c)]. With microwaves applied, a peak in current is observed at the resonance condition $f = g\mu_{\rm B}B/h$, indicating spin mixing by EDSR. The measured value $g \approx 2$ presumably arises because this device was highly disordered ($\Delta_{KK'} \gg \Delta_{\rm SO}$) and/or in the many-carrier limit, consistent

²²Because the confinement energy is usually much larger than the effective Zeeman splitting, to a good approximation the electron experiences the average B_{eff} over the entire dot.

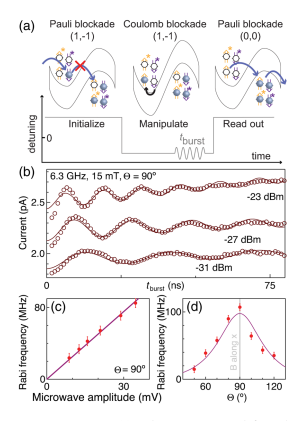


FIG. 40 (color online). (a) Pulse sequence used for coherent qubit manipulation. After initialization in a Pauli-blocked qubit state, the device is pulsed into Coulomb blockade to allow qubit manipulation and returned to Pauli blockade for readout. Electron tunneling occurs only if a qubit was flipped during the manipulation step. Repeated with a period $\sim 1 \ \mu s$, this leads to a current proportional to the qubit flip probability during the manipulation step. (b) Current (points) as a function of microwave burst duration for various applied powers, showing coherent Rabi rotations. The curves are fits to a model assuming a slowly varying random qubit detuning due to, e.g., charge noise. Traces are offset for clarity. (c) Rabi frequency as a function of microwave amplitude, with fit showing the expected proportionality. (d) Rabi frequency at constant microwave amplitude and power, showing a dependence on field angle Θ consistent with bend-mediated EDSR. Adapted from Laird, Pei, and Kouwenhoven, 2013.

with irregular Coulomb spacings seen in transport data (not shown). As well as the main resonance, a series of subharmonics at integer frequency fractions are seen, due to anharmonicity of the confinement potential or disorder (Nowak, Szafran, and Peeters, 2012; Széchenyi and Pályi, 2013). The resonant current increases with field as expected from Eq. (33) because $\mathbf{B}_{\rm eff}$ and hence $f_{\rm R}$ becomes larger.

4. Qubit manipulation and characterization

The data of Fig. 39 show only state mixing. Quantum coherence is demonstrated by measuring Rabi oscillations (Hanson *et al.*, 2007). This was achieved in a less disordered device using a pulsed measurement protocol that shuts off the current while microwaves are applied [Fig. 40(a)]. The scheme operates near the $(1, -1) \rightarrow (0, 0)$ charge transition,

where the qubit states participating in transport are (\uparrow, \downarrow) . Gate voltage pulses are used to adjust the detuning between two configurations, one Pauli blocked where electron tunneling is selective on the qubit state, and one Coulomb blocked where all tunneling is forbidden (Koppens et al., 2006). The sequence has three stages. An initialization stage at a Pauli blockade configuration loads with high probability a parallel two-qubit state (e.g., $\Downarrow \Downarrow$). The device is then configured in a Coulomb blockade, where a qubit manipulation microwave burst is applied, possibly flipping one of the qubits. During this step, tunneling is energetically suppressed regardless of the spin state. Finally the configuration is returned to Pauli blockade. If no qubit flip occurred during the manipulation stage, the state remains blocked. However, if a qubit (in either dot) was flipped, tunneling will occur based on the overlap of the electron state on the left with the empty state on the right. Repeating this cycle many times, the time-average current is proportional to the qubit flip probability during the manipulation stage.

As a function of burst duration, this current is observed to oscillate [Fig. 40(b)], indicating coherent rotations between qubit states at Rabi frequency $f_{\rm R}$. The fitted $f_{\rm R}$ is proportional to the driving microwave amplitude [Fig. 40(c)], consistent with a harmonic confinement potential and a smooth bend. The dependence of $f_{\rm R}$ on field angle Θ is consistent with bend-mediated EDSR coupling (Flensberg and Marcus, 2010) but not with, e.g., Rashba-mediated coupling, suggesting that the bend is indeed the dominant EDSR mechanism in this device.

The qubit is characterized further by determining the coherence time T_{echo} (Hanson *et al.*, 2007) which characterizes how long a superposition can be preserved by the use of a Hahn echo pulse. The coherence time T_{echo} is measured by a Ramsey fringe experiment (Fig. 41), which consists of (1) a

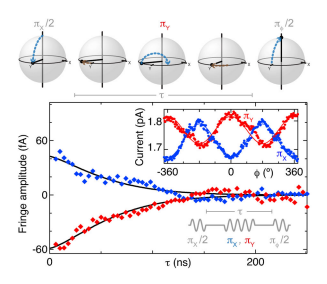


FIG. 41 (color online). Hahn echo amplitude (symbols) as a function of τ plotted for two phases of the central burst, together with fits (lines) of the form $e^{-(\tau/T_{echo})^{\alpha}}$, yielding $T_{echo} = 65$ ns. Inset: example fringes as a function of ϕ . The manipulation sequence is sketched at bottom right, and Bloch spheres along the top illustrate the resulting state evolution. Adapted from Laird, Pei, and Kouwenhoven, 2013.

 $\pi/2$ rotation about X to create a state on the equator, (2) a wait of duration τ , with a π rotation about X or Y inserted half way, and (3) a $\pi/2$ rotation with phase ϕ . Neglecting decoherence, the three rotations interfere to give a qubit flip probability proportional to $1 \pm \cos \phi$ (Fig. 41 inset). However, for $\tau \gg T_{echo}$, phase information is lost during the wait step, and the qubit flip probability is 1/2 independent of ϕ . By fitting the decay of fringe amplitude versus τ , the decay time $T_{echo} \approx 65$ ns is measured (Fig. 41). This coherence time T_{echo} is distinct from the dephasing time T_2^* discussed in the previous section because the π rotation during step (2) makes it insensitive to slow fluctuations of the qubit splitting. The dephasing time of this qubit was measured by a similar method (not shown) and found to be $T_2^* \approx 8$ ns (Laird, Pei, and Kouwenhoven, 2013).

Both T_2^* and T_{echo} are quite short compared with some other semiconductor spin qubits, such as GaAs (Petta et al., 2005; Greilich et al., 2006; Koppens, Nowack, and Vandersypen, 2008; Bluhm et al., 2011), Si (Maune et al., 2012; Pla et al., 2012), or diamond (de Lange *et al.*, 2010), but similar to results in InAs and InSb nanowires (Nadj-Perge, Frolov, Bakkers, and Kouwenhoven, 2010; van den Berg et al., 2013). The measured T_2^* is approximately consistent with hyperfine dephasing, given the unexpectedly large coupling discussed in Sec. V.B.2 (Churchill, Bestwick et al., 2009), but for T_{echo} to be limited in the same way, nuclear spin diffusion would have to be much faster than, e.g., in GaAs. This would be surprising, given the one-dimensional geometry and low density of nuclear spins in a nanotube. Fuller consideration of mechanisms led to the tentative conclusion that T_{echo} and perhaps T_2^* are limited by charge noise (Laird, Pei, and Kouwenhoven, 2013). To explain the ineffectiveness of the echo by extending the coherence time (even with longer decoupling sequences), this charge noise would need a broad spectral range, in contrast to the 1/f spectrum expected for charge switchers in the substrate (Cywiński et al., 2012).

D. Open questions

Although the main results presented in this section are understood, there are still significant unresolved questions. In all these experiments, it is hard to convincingly identify the precise spin-valley states between which transitions occur. Whereas the experiment of Fig. 36 is sensitive to all forms of relaxation between AS and S states, which may be expected in general to have different rates, a single T_1 value appears sufficient to fit each decay curve in Fig. 36(c). Likewise, in the experiment of Fig. 37, it is not clear how the S and AS state populations are redistributed by dephasing, and this is probably reflected in the unexplained $P(\infty)$ value discussed in Sec. V.B.2.

A related mystery comes from the measured EDSR spectra. Although in a disordered many-carrier device, expected resonances with $g \approx 2$ are observed [Fig. 39(c)], the spectrum in a cleaner device is complex and not understood (Laird, Pei, and Kouwenhoven, 2013; Li *et al.*, 2014). Whether the unexplained features relate to the short T_{echo} is not known.

An exciting area opened by this work is the possibility to combine the spin degree of freedom with mechanical and optical degrees in clean, suspended nanotubes. There has already been progress in engineering quantized phonons in nanotubes and studying their interactions with the charge on quantum dots (Sazonova et al., 2004; Sapmaz, Jarillo-Herrero, Blanter, and van der Zant, 2005; Huttel et al., 2009; Lassagne et al., 2009; Steele et al., 2009; Benyamini et al., 2014). Evidence for the discreteness of longitudinal stretching phonon modes comes from a Frank-Condon blockade in suspended nanotubes (Sapmaz, Jarillo-Herrero, Blanter, and van der Zant, 2005; Leturcq et al., 2009), discussed theoretically by Sapmaz et al. (2003), Flensberg (2006), and Mariani and von Oppen (2009). By tuning the discrete phonon modes away from resonance with qubit splittings, long qubit lifetimes may be achievable. On the other hand, when the qubit splitting is nearly resonant with a discrete phonon mode, coherent energy exchange should be possible between them in a solidstate analog of cavity quantum electrodynamics (Pályi et al., 2012). Strong spin-phonon coupling in suspended nanotubes may also enable an enhanced sensing of nanotube motion (Ohm et al., 2012). Finally, Galland and Imamoğlu (2008) and Li and Zhu (2012) theoretically investigated spin-based mechanics and quantum optics. Using a combination of magnetic fields and optical pumping, they predicted highfidelity all-optical control of electron spins, phonon-induced transparency, and applications in quantum communication.

Finally, we compared this work with the large body of experiments on spin relaxation, dephasing, and diffusion in ensembles of single-walled carbon nanotubes measured via electron spin resonance (ESR) and EDSR spectroscopy at higher temperatures 4-300 K (Petit et al., 1997). These techniques focus on resonances that appear within a few percent of g = 2 at several tesla. These results seem to contradict the understanding gained from quantum transport experiments because the g factor is expected to be highly anisotropic in clean nanotubes. Therefore the debate whether such resonances reflect intrinsic spin properties of carbon nanotubes (Dora et al., 2008; Kombarakkaran and Pietraß, 2008) or defects (Rice et al., 2013) has yet to be settled. Indeed, sufficiently purified nanotubes where removal of the catalyst was confirmed by TEM were found to not yield an ESR signal (Zaka et al., 2010). The level structure of Fig. 21 does not provide a reason why randomly oriented assemblies of nanotubes should yield resonances at $g \approx 2$. The only limit where $g \approx 2$ may occur is for samples in which $|\Delta_{KK'}|$ exceeds both $|\Delta_{SO}|$ and the orbital magnetic field splitting, which is not the case for clean, intrinsic nanotubes.

VI. VALLEY PHYSICS IN OPEN QUANTUM DOTS

A. Transport in open regime

Previous sections focused on closed quantum dots where low-transparency barriers ensure that transport occurs by sequential tunneling via strongly confined quantum states. However, nanotube devices with a range of transparencies can be fabricated, allowing the transition from closed to open transport regimes to be studied (Nygård and Cobden, 2001; Cao, Wang, and Dai, 2005; Liang, Bockrath, and Park, 2005; Grove-Rasmussen, Jørgensen, and Lindelof, 2007). Valley and spin physics again play important roles for highly transmitting devices where the quantum dot states are hybridized

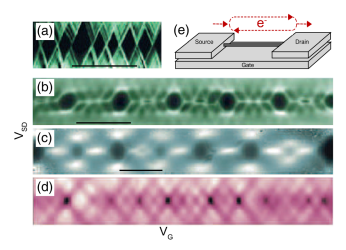


FIG. 42 (color online). Quantum transport from closed to open devices. (a)–(d) Plots of dI/dV_{SD} as a function of gate voltage V_G and bias V_{SD} for four nanotube devices with average conductances $g \sim 0.01, 0.5, 1.5, 3e^2/h$ [schematic in (e)]. All data were taken at 1.5 K and zero field. High (low) conductance is shown by light (dark) colors. Black bars indicate the gate voltage range for the addition of four additional electrons. (e) Device schematic. Dashed paths indicate the origin of Fabry-Pérot resonances induced by reflections at the contacts. Adapted from Liang, Bockrath, and Park, 2005.

with the Fermi seas in the leads. Here we provide an overview of transport mechanisms in open devices and then focus on phenomena involving valley physics: valley and SU(4) Kondo effects (Sec. VI.B) and level renormalization (Sec. VI.C).

Figures 42(a)-42(d) show transport data from four devices with varying contact transparency. All devices exhibit metallic characteristics at room temperature (not shown) with average conductances ranging from around $0.01e^2/h$ to $3e^2/h$, i.e., approaching the maximum conductance of $4e^2/h$ for a single nanotube. The first, low-conductance device (a) behaves as a closed quantum dot. For the second device (b) with average $g \sim 0.5e^2/h$, i.e., close to the conductance quantum, the fourfold periodicity due to valley and spin degeneracy leads to clusters of four peaks (Sec. III.D). The enhanced background conductance reflects cotunneling processes enabled by the stronger coupling. The most extraordinary features are the horizontal (gate-independent) ridges of high conductance, e.g., occurring near zero bias in a large fraction of the Coulomb diamonds. These resonances are due to higher-order tunneling processes, including Kondo physics, described in Sec. VI.B.

For the next devices (c) and (d) the quantum dot features are smeared out as the increased coupling to leads allows for charge fluctuations on the nanotube. However, gateperiodic patterns remain and in the highly transmitting device (d) a distinct pattern of low-conductance lines dominates the spectroscopy plot. Here, in the simplest picture, mode reflections at the contacts [Fig. 42(e)] give rise to interference in transmission, the so-called Fabry-Pérot resonances (Liang *et al.*, 2001). The interference pattern appearing in bias spectroscopy plots is similar to universal conductance fluctuations (UCF) in other mesoscopic systems (Nazarov and Blanter, 2009; Ihn, 2010). However, the randomness that usually characterizes UCF is replaced by nearly perfect periodicity for one-dimensional, ballistic nanotube resonators.

B. Spin, valley, and SU(4) Kondo effects in nanotubes

1. Theory and background

Transport in nearly closed quantum dots can be described in terms of first-order sequential tunneling. With stronger coupling to the leads higher-order processes involving virtual intermediate states become relevant [Fig. 43(a)]. Initially, the system is in a state of Coulomb blockade (left). Higher-order fluctuations can permit tunneling of the trapped electron to the right lead, while a second electron from the source enters the dot. Effectively, one electron charge -e has been transferred from source to drain (right diagram) via an intermediate state (middle) that is classically forbidden due to energy conservation and Coulomb blockade. Such a process is called elastic cotunneling (Averin and Nazarov, 1992; Pustilnik and Glazman, 2004; Ihn, 2010). In Fig. 43(a) the spin on the dot is flipped as permitted by the spin degeneracy of the level at zero field. The nontrivial result of such higher-order spin-flip transitions is the appearance of a new correlated ground state for the combined lead-dot system. This is an effective singlet state between an unpaired spin and the Fermi sea of electrons in the leads. It forms a highly transmitting channel between source and drain at the Fermi energy, known as a Kondo resonance, leading to the breakdown of the Coulomb blockade (Nazarov and Blanter, 2009; Ihn, 2010; Heikkila, 2013).

The Kondo state exists only at low temperatures where coherence is preserved. The transition temperature from the Coulomb blockade to transport resonance is denoted as the Kondo temperature $T_{\rm K}$. The energy scale $k_{\rm B}T_{\rm K}$ can be considered as the binding energy of the many-body singlet state formed from the quantum dot spin and a screening cloud of electrons in the lead. Kondo resonances appear in Fig. 42(b) as ridges of high conductance near zero bias. The corresponding charge states can be identified as the odd occupancy states where the quantum dot holds an unpaired electron. (The apparent resonances in some of the even diamonds are discussed later.) Several diagnostics can verify an underlying Kondo mechanism; the resonance should be suppressed by increasing temperature, bias voltage, or an external magnetic field that breaks the necessary level degeneracy.

Quantum dot Kondo resonances were first discovered in two-dimensional semiconductor devices (Cronenwett, Oosterkamp, and Kouwenhoven, 1998; Goldhaber-Gordon, Shtrikman et al., 1998; Simmel et al., 1999) but observed shortly afterward in carbon nanotubes (Nygård, Cobden, and Lindelof, 2000; Buitelaar et al., 2002; Liang, Bockrath, and Park, 2002; Babić, Kontos, and Schönenberger, 2004). For a more detailed account of Kondo physics and the relation to quantum dots in general see other reviews (Kouwenhoven and Glazman, 2001; Pustilnik and Glazman, 2004; Zaránd, 2006; Grobis et al., 2007) and textbooks (Nazarov and Blanter, 2009; Ihn, 2010; Heikkila, 2013). In all cases, the spin Kondo effect leads to an enhanced conductance at low temperatures. For nanotubes $T_{\rm K}$ is typically 1–10 K.

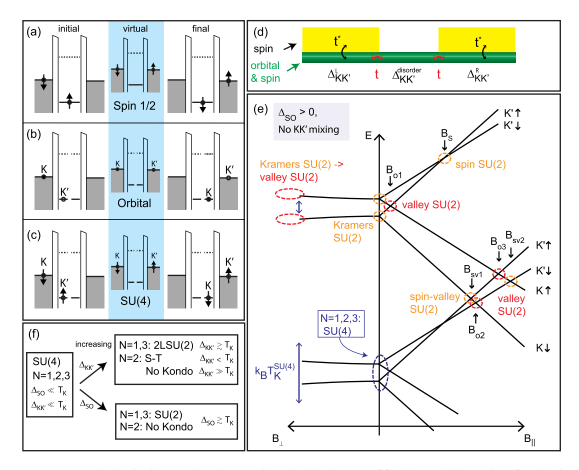


FIG. 43 (color online). Kondo physics in nanotubes due to spin and valley degrees of freedom. (a) Schematics of an elastic cotunneling process resulting in a spin flip on the quantum dot. Such higher-order processes give rise to the spin Kondo effect. (b) The Kondo effect can also occur if another quantum number is present in the leads and the dot, e.g., the valley quantum number as shown here. (c) In nanotubes valley and spin quantum numbers may additionally give rise to the SU(4) Kondo effect involving both degrees of freedom. (d) Device schematic showing valley mixing in the lead $(\Delta_{KK'}^{L,R})$ and the dot $(\Delta_{KK'}^{disorder})$. For SU(4) Kondo effects to be observed these should be small and valley-conserving tunneling (*t*) from nanotube leads is required. (e) Energy vs parallel and perpendicular magnetic fields for two conduction-band shells with spin-orbit interaction $(\Delta_{SO} > 0)$. Dashed ellipses and related *B* fields indicate degeneracies resulting in Kondo phenomena, e.g., SU(4) and various SU(2) Kondo effects. The SU(4) Kondo effect can be observed (lower shell) when the related energy scale is much larger than the zero-field (spin-orbit) splitting. (f) Flow diagram at B = 0 for fillings N = 1 - 3 showing that valley mixing reduces the SU(4) Kondo effect to SU(2) or no Kondo effect. The intervalley mixing parameter $\Delta_{KK'}$ represents all relevant mixing terms. (a)–(c) Adapted from Jarillo-Herrero *et al.*, 2005b.

We will not dwell on the details of the spin Kondo effect but rather point out that in principle any doubly (or higher) degenerate localized state with identical quantum numbers in the leads could mediate Kondo-like resonances. The origin of the degeneracy does not need to be spin; for example, transport resonances could be induced by an orbital level degeneracy on the dot [Fig. 43(b)] (Jarillo-Herrero *et al.*, 2005b). In the context of nanotubes the valley degree of freedom comes to mind. Next we discuss such a valley Kondo effect.

Both the ordinary spin-1/2 Kondo effect and the twofold valley Kondo effect reflect SU(2) symmetry. However, for ideal nanotube dots the concomitant existence of valley and spin freedom leading to an approximate fourfold degeneracy could potentially lead to a Kondo effect described by the higher SU(4) symmetry class (Choi, López, and Aguado, 2005). This situation is absent in most other quantum dots, which lack the spatial symmetry that

naturally leads to valley degeneracy in nanotubes.²³ Figure 43(c) shows an example of states involved in this scenario where both the valley and spin quantum numbers can be exchanged during cotunneling. In order to probe the SU(4) Kondo effect, it is essential that transitions between all four states are possible.

Even if valley degeneracy exists in the nanotube, this does not ensure that the valley quantum number is also present in

 $^{^{23}}$ SU(4) Kondo physics has been studied in vertical quantum dots (Sasaki *et al.*, 2004), parallel double quantum dots (Wilhelm *et al.*, 2002; Holleitner *et al.*, 2004; Okazaki, Sasaki, and Muraki, 2011; Keller *et al.*, 2013), and single dopants (Tettamanzi *et al.*, 2012). In the vertical quantum dots, an orbital degree of freedom also exists in both the leads and the dot because they share similar symmetry. To study SU(4) physics in two parallel dots, however, requires a device with identical intra-Coulomb and inter-Coulomb energy.

the leads. Figure 43(d) shows a nanotube quantum dot coupled to metallic leads. The double arrows indicate that electrons need to enter a nanotube lead segment (t^*) before tunneling onto the quantum dot (t) to allow for SU(4) Kondo physics since this effect requires valley-conserving tunneling. As discussed later, K-K' mixing during tunneling ($\Delta_{KK'}^{\text{tunneling}}$), on the dot ($\Delta_{KK'}^{\text{disorder}}$), and in the lead segments ($\Delta_{KK'}^{L,R}$) must all be weak, as must spin-orbit coupling.

To link the different Kondo phenomena to the nanotube level structure, the spectrum of two shells is plotted in Fig. 43(e), including spin-orbit interaction but not valley mixing. Dashed ellipses indicate degeneracies that can lead to Kondo physics. When the SU(4) Kondo energy scale is much larger than the spin-orbit splitting, the SU(4) Kondo effect can be observed at zero field for occupations N = 1, 2, or 3 as depicted for the lower shell. The opposite case of small SU(4) Kondo temperature is shown for the upper shell, resulting in SU(2) Kondo effects for the two split Kramers doublets (dashed circles at zero field) (Fang, Zuo, and Luo, 2008, 2010; Galpin *et al.*, 2010).

Figure 43(e) also points to possible Kondo effects in a parallel magnetic field. Two intrashell SU(2) Kondo effects are illustrated for the upper shell: a valley SU(2) Kondo effect at half filling and a spin SU(2) Kondo effect for three electrons in the shell. Intershell valley Kondo effects can arise from level crossings between two different shells. Taking account of valley mixing within the shells modifies the energy diagram (Fig. 21), but does not qualitatively change the intershell valley degeneracies (Jarillo-Herrero et al., 2005b; Grove-Rasmussen et al., 2012). However, the intrashell valley degeneracies for parallel and perpendicular fields are split, and when valley mixing dominates, a spin-orbit split degeneracy (not shown) emerges at finite perpendicular field giving rise to a singlet-triplet $(S-T_{-})$ Kondo effect (Nygård, Cobden, and Lindelof, 2000; Pustilnik, Avishai, and Kikoin, 2000; Quay et al., 2007). The zero-field SU(4) Kondo effect survives as long as the Kramers doublet splitting due to valley mixing and spin-orbit coupling is much smaller than the Kondo temperature (Borda et al., 2003; Fang, Zuo, and Luo, 2008; Galpin et al., 2010).

As shown in Figs. 43(e) and 43(f), the SU(4) Kondo effect can be observed for filling N = 1 - 3 (Anders *et al.*, 2008), but only if the valley quantum numbers are conserved during tunneling, i.e., no valley mixing. If this is not the case, the SU(4) Kondo effect reduces to two-level (2L) spin SU(2)(Choi, López, and Aguado, 2005; Lim et al., 2006; Büsser and Martins, 2007; Büsser et al., 2011) and singlet-triplet (S-T) (Eto and Nazarov, 2000; Sasaki et al., 2000; Izumida, Sakai, and Tarucha, 2001) Kondo effects for N = 1, 3 and N = 2, respectively.24 For the 2L SU(2) Kondo effect, the Kondo temperature and maximum conductance $q = 2e^2/h$ at the center of the N = 1, 3 Coulomb diamond is the same as in the case of the SU(4) Kondo effect, and the two effects are therefore to be distinguished either by magnetic field spectroscopy (Choi, López, and Aguado, 2005; Jarillo-Herrero et al., 2005b) or by the overall shape of the N = 1 - 3

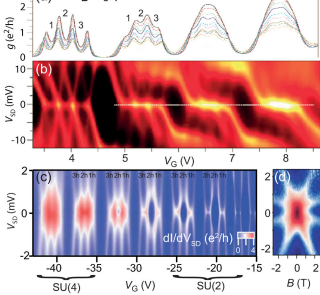
 $V_{\rm G}({\rm V})$ SU(2) SU(4) FIG. 44 (color online). Kondo physics near zero field. (a) Conductance vs gate voltage of a narrow-gap nanotube for four electron shells I-IV. For shells I and II the occupancy in each diamond is indicated. More positive V_{G} increases the tunnel couplings, tuning the dot from the Kondo to the mixed-valence regime. Different traces correspond to temperatures from 1.3 to 15 K. (b) Differential conductance in the same device as a function of $V_{\rm G}$ and $V_{\rm SD}$ at 3.3 K. (c) Similar data for a different device showing spin-orbit split SU(2) and SU(4) Kondo physics at low and high hole fillings, respectively. (d) Magnetic field splitting of N = 1 Kondo resonance into four peaks, indicating SU(4) Kondo physics. (a), (b) Adapted from Makarovski, Liu, and Finkelstein, 2007. (c) From Cleuziou et al. (2013). (d) From

conductance versus gate voltage (Makarovski and Finkelstein, 2008). Similarly, in the center of the N = 2 Coloumb diamond, both the singlet-triplet and SU(4) Kondo effects have the same theoretical maximum conductance $g = 4e^2/h$ (Jarillo-Herrero *et al.*, 2005b). However, for sufficiently strong valley mixing (or spin-orbit interaction), a zero-field Kondo effect is absent for N = 2, as indicated in Fig. 43(f).

2. Experiment

Jarillo-Herrero et al., 2005b.

Unconventional Kondo physics in nanotubes was first experimentally studied by Jarillo-Herrero *et al.* (2005b) and later by Jarillo-Herrero *et al.* (2005a), Makarovski, Liu, and Finkelstein (2007), Makarovski *et al.* (2007), Makarovski and Finkelstein (2008), Cleuziou *et al.* (2013), and Schmid *et al.* (2015).²⁵ For all but the last two experiments, the spin-orbit interaction was thought to be negligible and thus



B = 0 T

Ш

²⁴The analysis in terms of singlet-triplet physics assumed zero spin-orbit interaction.

²⁵Other types of unconventional Kondo phenomena such as the two-impurity Kondo effect have been studied in nanotube systems (Chang and Chen, 2009; Bomze *et al.*, 2010; Chorley *et al.*, 2012). Equipping nanotubes with superconducting or ferromagnetic leads also gives rise to new Kondo systems (Buitelaar, Nussbaumer, and Schönenberger, 2002; Hauptmann, Paaske, and Lindelof, 2008), but that is beyond the scope of this review.

not included in the Kondo analysis (Choi, López, and Aguado, 2005; Lim *et al.*, 2006; Büsser and Martins, 2007; Anders *et al.*, 2008; Mizuno, Kim, and Martins, 2009). However, more recent theory includes spin-orbit interaction and reinterprets the early data (Fang, Zuo, and Luo, 2008, 2010; Galpin *et al.*, 2010). Our aim is to present data relevant for the spectrum of Fig. 43(e), starting with zero magnetic field Kondo physics followed by finite-field phenomena for one and two shells.

The ideal devices for studying an SU(4) Kondo effect are quantum dots with tunable couplings to the leads. Such devices can be realized in narrow-gap nanotubes where the conductance typically increases as carriers are added. Figures 44(a) and 44(b) show the linear conductance dI/dV_{SD} versus gate voltage and the corresponding charge stability diagram. A regular shell structure with zero-bias Kondo ridges for electron filling one, two, and three is observed at low gate voltages. At stronger electrode coupling, the different charge states fully hybridize (Makarovski, Liu, and Finkelstein, 2007; Makarovski and Finkelstein, 2008) and a single merged peak is observed for the two rightmost shells in Fig. 44(a). In this mixed-valence regime the lifetime broadening is comparable to or larger than the charging energy and the single-electron transport features of the stability diagram merge into broad ridges. The observed behavior in both the Kondo and mixed-valence regimes has been reproduced by numerical renormalization group (NRG) calculations within an SU(4) Anderson model for various lead couplings, supporting this interpretation (Anders et al., 2008).

Figure 44(c) systematically studies the transition from SU(4) to SU(2) Kondo physics in a clean narrow-gap device (Cleuziou *et al.*, 2013). The crossover is measured across seven shells, thereby varying both the spin-orbit interaction (decreases with filling, see Fig. 19) and the lead coupling (increases with filling). A transition from $\Delta_{SO} > T_K^{SU(4)}$ to $\Delta_{SO} < T_K^{SU(4)}$ versus hole filling, i.e., SU(2) to SU(4), can thus be realized in accordance with the lower arrow in the flow diagram of Fig. 43(f).

The SU(4) Kondo effect was also reported by Jarillo-Herrero *et al.* (2005a, 2005b) studying the N = 1 zero-bias peak. The temperature dependence (0.3-10 K) of the conductance peaks was consistent with the expected empirical scaling formula (Goldhaber-Gordon, Göres et al., 1998) yielding high Kondo temperatures (around 10 K). Figure 44(d) shows the behavior of the Kondo resonance in a (parallel) magnetic field. The splitting into four peaks (ideally six if the Zeeman splittings are fully resolved) indicates SU(4) Kondo physics, since only two peaks would emerge if valley mixing induced by tunneling or disorder were dominant as for the 2L SU(2) Kondo effect (Choi, López, and Aguado, 2005). A zero-bias ridge observed for N = 2 in the same work was analyzed in terms of singlet-triplet Kondo physics, but is also consistent with an SU(4) Kondo effect (Büsser and Martins, 2007).

Next we turn to Kondo phenomena at finite B_{\parallel} within a single shell [upper part of Fig. 43(e)] (Fang, Zuo, and Luo, 2008, 2010; Galpin *et al.*, 2010). Recent experiments on the addition of the first four electrons in a device with clearly identified spin-orbit interaction give hints of the valley and finite-field spin SU(2) Kondo effects (Cleuziou *et al.*, 2013).

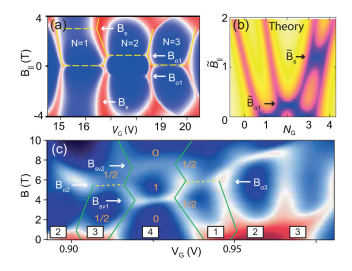


FIG. 45 (color online). Kondo physics at finite field. (a) Differential conductance vs gate voltage and parallel magnetic field for the first three electrons in the conduction band. Dashed lines indicate faint Kondo resonances: Kramers (for N = 1, 3 at $B_{\parallel} = 0$, valley (for N = 2 at $B_{\parallel} = B_{01}$), and spin SU(2) (for N = 1 at $B_{\parallel} = B_{\rm s}$). (b) Theoretical linear conductance vs gate voltage filling (expressed as effective filling $N_{\rm G}$) and normalized parallel field \tilde{B}_{\parallel} . The calculation assumed $|\Delta_{SO}| > T_{K}^{SU(4)}$ as in the top shell of Fig. 43(e). (c) Differential conductance as a function of gate voltage V_{G} and parallel magnetic field B at 30 mK. Boxed numbers indicate the occupation of the topmost shell and unboxed numbers indicate the ground-state spin (0, 1/2, 1). Solid lines highlight the motion of the Coulomb blockade peaks. Dashed lines mark valley Kondo effects [compare white arrows to Fig. 43(e)]. (a)–(c) Adapted from Cleuziou et al., 2013, Galpin et al., 2010, and Jarillo-Herrero et al., 2005b, respectively.

Figure 45(a) shows the linear conductance versus parallel magnetic field and gate voltage, with the electron filling and the possible Kondo ridges indicated. Compared to the model above, the finite-field spin SU(2) Kondo effect at B_s for odd filling appears for N = 1 instead of N = 3. [Whether it is the upper or lower two states that crosses at high field depends on carrier type and the sign of the spin-orbit interaction (see Fig. 18).] The N = 2 valley Kondo effect at B_{o1} is clearly visible close to the Coulomb peaks, mimicking the behavior of the Kramers Kondo effect at zero field. Because of small tunnel coupling in this device, valley or spin Kondo physics cannot be unambiguously identified from the data.

Similar data with strong Kondo ridges were already observed before the identification of nanotube spin-orbit interaction (see Fig. 12, shell around $V_{\rm G} \sim 3$ V), but the Kondo ridge at half filling and finite *B* was interpreted in terms of singlet-triplet Kondo physics²⁶ (Jarillo-Herrero *et al.*, 2005a). Later NRG modeling [Fig. 45(b)] suggested that the data could also be explained in terms of the spin-orbit

²⁶Depending on the exchange interaction, the triplet state occupying two different valleys may become the ground state. In a parallel magnetic field the singlet state occupying two downmoving valleys thus crosses the almost degenerate triplet states, giving rise to a Kondo effect (assuming negligible Zeeman effect).

energy spectrum²⁷ (Galpin *et al.*, 2010). The calculated color plot thus represents the ideal Kondo behavior for the upper shell in Fig. 43(e). Other experiments with dominating valley mixing ($\Delta_{KK'} > \Delta_{SO}$) also identify the spin SU(2) Kondo effect at $B_{\parallel} = B_s$ (Jespersen, Grove-Rasmussen, Paaske *et al.*, 2011; Schmid *et al.*, 2015).

Finally we address the (intershell) valley Kondo effect. It can be found at particular fields (B_{o2}, B_{o3}) that induce valley degeneracy but not spin degeneracy [Fig. 43(e)]. Figure 45(c) shows the (parallel) magnetic field evolution of the conductance for a nanotube quantum dot that has strong Kondo resonances at zero field. At finite fields additional resonances appear at B_{o2} and B_{o3} , consistent with valley degeneracies inducing a spinless valley SU(2) Kondo effect. The appearance of this Kondo resonance indicates that tunneling preserves the valley symmetry, which for this device is less obvious because of the metal deposited on top of the nanotube leads.

In several of the experiments above a Kondo effect arising from inelastic transitions gave rise to conductance peaks at finite bias. Such Kondo enhancement of inelastic cotunneling thresholds was observed and modeled in Paaske *et al.* (2006) while a recent experiment revealed that certain cotunneling thresholds are not observed in the strong coupling regime (Schmid *et al.*, 2015). They identify the two states involved in the relevant cotunneling processes as particle-hole symmetric²⁸ and show theoretically that these processes do not give rise to Kondo correlations (even though cotunneling is allowed).

C. Level renormalization

The most visible consequence of increasing electrode couplings in Fig. 42 is a broadening of the energy levels, but this hybridization is also accompanied by level shifts. These shifts are named tunnel renormalization, since the effect stems from quantum charge fluctuations (cotunneling events) that are particularly relevant when the tunnel couplings are large. Nanotube quantum dots have turned out to be ideal to observe such level shifts, in particular, when the two doublets are differently coupled to the leads (Holm *et al.*, 2008).

1. Theory

We consider a spinless nanotube shell model without any internal couplings and assume metallic leads (no valley quantum number). Since $|K\rangle$ and $|K'\rangle$ states are time-reversed partners, the tunnel couplings are equal $t = |t_K| = |t_{K'}|$. Introducing a complex valley-mixing term $\Delta_{KK'} = |\Delta_{KK'}|e^{i\phi_0}$ results in new eigenstates which have phase dependent tunnel couplings $t(1\mp e^{i\phi_0})$ (Grove-Rasmussen *et al.*, 2012). Figure 46(a) shows the energy versus parallel

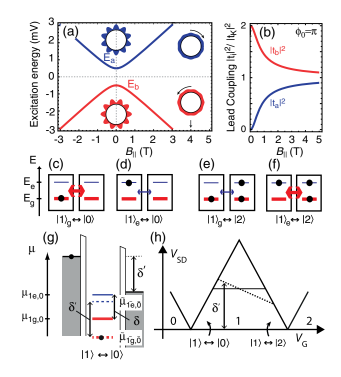


FIG. 46 (color online). Level renormalization. (a) Energy vs parallel magnetic field of two valley states with coupling $\Delta_{KK'}$ that splits the states at zero field. Insets show the electron density distributions at zero and high fields. (b) Lead couplings vs parallel magnetic field of the two eigenstates in (a). (c)-(f) Cotunneling processes for the two doublets of a singly occupied shell near the 1 to 0 (c), (d) and 1 to 2 (e), (f) charge degeneracy points. The thickness of the arrows indicates the tunnel couplings. (g) Electrochemical potentials $\mu_{1q,0} = E_{1q}$ – E_0 (solid lines) and renormalized versions $\tilde{\mu}$ (dashed lines). Here E_{1q} and E_0 are the energies of the one-electron ground state (1q) and the empty-dot state (0). The lead potentials are set at the threshold for inelastic cotunneling; renormalization enhances the threshold energy δ' compared to the original doublet splitting δ . (h) Schematic charge stability diagram for the one-electron diamond. The resulting level renormalizations induce a gate-dependent inelastic cotunneling threshold (dashed line) in contrast to the single-particle prediction δ (solid line). Adapted from Grove-Rasmussen et al., 2012.

magnetic field for the new eigenstates that arise when the valley states are coupled. The schematic insets show that the electron probability distributions for the two eigenstates are different at zero field and thus the tunneling amplitude may be different depending on the exact site of a microscopic contact. This picture of tunneling into one particular atomic site with different probability for the two wave functions is probably too simple for a real device but is adopted here as a minimal model. In Fig. 46(b) the lead couplings for the two states are plotted for the case $\phi_0 = \pi$. At zero field the two new eigenstates have different couplings: the ground state doubles its coupling to the leads compared to the original states while the excited state decouples. The model also makes an important prediction about the magnetic field dependence of the couplings. At high parallel magnetic fields the ground- and excited-state couplings become equal, since the eigenstates are close to the original valley states $|K\rangle$

²⁷While this is a likely interpretation, some caution should be applied since level crossings with a different shell complicate the finite-field spin SU(2) identification.

²⁸The work considers a shell with both spin-orbit interaction and valley mixing. The eigenstates are therefore superpositions of all four basis states $|\tau\sigma\rangle$. A simple example of a particle-hole symmetric pair is $\{|K\downarrow\rangle, |K'\uparrow\rangle\}$.

and $|K'\rangle$ whose probability distributions are equal at high fields [Fig. 46(a) inset].

We now examine the effect of the difference in lead couplings on cotunneling. Consider two states with different tunnel couplings as shown in Figs. 46(c) and 46(d) for a singly occupied dot $|1\rangle$. The ground state experiences stronger cotunneling charge fluctuations via the zero state $(|1\rangle_a \leftrightarrow |0\rangle)$ than the excited state $(|1\rangle_e \leftrightarrow |0\rangle)$ does. The renormalization shift increases with the coupling of the levels and thus the more strongly coupled state has the larger energy shift [Fig. 46(g)]. Consequently, the threshold for inelastic cotunneling transport will increase (from δ to δ') near the $|1\rangle \leftrightarrow |0\rangle$ charge transition (left side of the diamond). In the stability diagram Fig. 46(h) the threshold will correspondingly be shifted to higher bias. In contrast, on the right side of the one-electron diamond, the dominant cotunneling processes involve the two-electron state $(|1\rangle \leftrightarrow |2\rangle)$. The situation is opposite here since the excited state $|1\rangle_e$ experiences

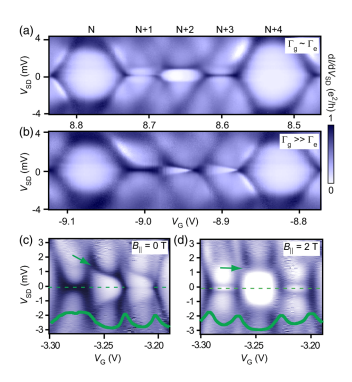


FIG. 47 (color online). Tunnel renormalization in nanotube stability diagrams. (a) Stability diagram for a shell with similar lead couplings to ground and excited states ($\Gamma_q \sim \Gamma_e$). Kondo ridges are observed for one and three electrons and inelastic cotunneling thresholds are gate independent. Occupation numbers are marked above the plot. (b) Stability diagram for a different shell. Unlike in (a), the Kondo ridge appears only for the N + 1 diamond, and the cotunneling thresholds in the N + 2 and N+3 diamonds are gate dependent. This is consistent with cotunneling provided that the doublet lead couplings are asymmetric ($\Gamma_a \gg \Gamma_e$ deduced from thresholds for N + 2 and N + 3). (c) Diagram at zero field for a different device. Again, a gatedependent cotunneling threshold (arrow) indicates asymmetric couplings. (d) At $B_{\parallel} = 2$ T the threshold is gate independent, consistent with equalization of the lead couplings by the magnetic field as discussed in the text. Adapted from Holm et al., 2008 and Grove-Rasmussen et al., 2012.

stronger fluctuations and level shifts than the ground state $|1\rangle_g$ [Figs. 46(e) and 46(f)]. Thus the cotunneling threshold will be reduced compared to the original level in this gate range. The resulting stability diagram is sketched in Fig. 46(h). The models above are easily extended to include spin and higher fillings, but the conclusion remains the same: tunneling-induced level renormalization can result in gate-dependent cotunneling features within the charge stability diamonds.

2. Experiment

Figure 47(a) shows a stability diagram with shell filling identified by the characteristic pattern of three small Coulomb diamonds followed by a large one and Kondo ridges for odd occupancies. However, Fig. 47(b) shows that this regular behavior is not observed for all shells. In this case, the Kondo effect is evident only for one electron, and the inelastic cotunneling thresholds²⁹ are seen to have a marked gate dependence for two and three electrons where arguments similar to the model above hold [compare Fig. 46(h)]. The absence of the Kondo effect for N = 3indicates that the ground-state doublet is more strongly coupled to the leads than the excited-state doublet. The different behavior for different shells, such as between Figs. 47(a) and 47(b), can be understood as due to valley mixing with different phases for each shell and tunnel renormalization.

The effective doublet splitting is set by both spin-orbit and valley coupling $\delta = \sqrt{\Delta_{SO}^2 + \Delta_{KK'}^2}$, while in the leadcoupling model of Fig. 46(b) the origin of the coupling asymmetry was the valley coupling alone. The validity of this model can be tested by examining its prediction for the lead tunnel couplings in a magnetic field. Figure 47(c) shows a stability diagram of a shell whose behavior is similar $(\Gamma_q > \Gamma_e)$ to that of Fig. 47(b). The corresponding stability diagram in parallel magnetic field $B_{\parallel} = 2$ T is shown in Fig. 47(d). As predicted qualitatively by the model, the gate dependence of the cotunneling threshold disappears at large field where the lead couplings become equal, consistent with the observed widths of the Coulomb peaks within the shell [see cut in Fig. 47(d)]. This observation indicates that the origin of the asymmetry is related to valley mixing (Grove-Rasmussen et al., 2012).

Renormalization effects have also been established in devices with ferromagnetic leads, where the difference in spin density of states of the electrodes leads to different effective tunnel coupling (Martinek *et al.*, 2003, 2005; Pasupathy *et al.*, 2004; Hauptmann, Paaske, and Lindelof, 2008). Furthermore, if valley mixing originates from cotunneling (not only disorder as described above) different gate-dependent shifts are predicted with the possibility of inelastic cotunneling lines crossing inside the Coulomb diamond (Kiršanskas, Paaske, and Flensberg, 2012).

²⁹For the one-electron case the inelastic cotunneling threshold is not visible due to the Kondo effect present at zero bias.

D. Open questions

Several experiments have found features consistent with Kondo correlations originating from the additional valley degree of freedom. A complete study of the SU(4) Kondo effect would, however, call for control of the valley-mixing parameters [Fig. 43(d)] to tune the valley Kondo correlations. Valley mixing also plays a role for the intershell valley Kondo experiment, where incipient Kondo correlations enhancing finite-bias features are analyzed (Jarillo-Herrero *et al.*, 2005); Paaske *et al.*, 2006). Such correlations are generally more difficult to quantify. Clearer experimental results may be possible in tunable gate-defined dots with weak valley mixing, where the leads are nanotube segments rather than metal.

VII. CORRELATED-ELECTRON EFFECTS

A. Introduction

Electron correlation effects due to Coulomb interaction can be strong in carbon nanotubes. One reason is the onedimensional nature of the confinement: electrons in two or three dimensions can minimize their Coulomb repulsion by moving out of each other's way. Electrons in nanotubes do not have this freedom and instead tend to develop strong correlations. Electrons in nanotubes also see an environment with a low dielectric constant, which for suspended nanotubes approaches the free-space value $\varepsilon = 1$. This is in contrast to electrons in semiconductors, where electric fields can be screened by the large dielectric constant of the host material. These properties suggest that nanotubes are an interesting system for studying electron correlations.

The strength of Coulomb interactions between particles with parabolic dispersion is characterized by an interaction parameter r_s , defined as the ratio of the average interparticle spacing l to the Bohr radius a_0 :

$$r_{\rm s} = \frac{l}{a_0} = \frac{m_{\rm eff} e^2 l}{\varepsilon \hbar^2} \approx \frac{E_{\rm C}}{E_{\rm K}}.$$
 (34)

Within a numerical constant of order unity, r_s is also the ratio of the Coulomb interaction energy $E_C = e^2/\epsilon l$ to the kinetic energy $E_K = \hbar^2/ml^2$. At high densities (small r_s), kinetic energy dominates and the single-particle approximation can be used for the electronic states. At low densities (large r_s), kinetic energy is quenched and the Coulomb interaction dominates the physics.

For semiconducting nanotubes at sufficiently low densities, the Fermi energy lies in the parabolic region of the dispersion relation (cf. Fig. 5) and r_s is a useful characterization of the strength of electron interactions. A typical Coulomb interaction energy in a nanotube for $\varepsilon = 1$ and l = 100 nm is $E_{\rm C} = 13$ meV. To estimate the kinetic energy, it is important to note that the effective mass is strongly dependent on the band gap. For a semiconducting nanotube with a band gap of 210 meV ($m_{\rm eff} = 0.029$), $E_{\rm K} = 260 \ \mu \rm eV$ (Bohr radius $a_0 \approx 2$ nm). This implies that two electrons in a 200 nm long quantum dot in such a nanotube corresponds to very strong interactions ($r_{\rm s} = 50$).³⁰ Advances in making clean suspended nanotubes have enabled the study of quantum dots at very low density with very low electronic disorder. Using these new devices, the question of electron interaction and correlation effects in nanotubes is being revisited from the ground up. In the fewelectron regime in the clean limit, a clear understanding of the simplest case of two electrons is beginning to emerge.

B. Interactions in two-electron nanotube quantum dots

The Coulomb interaction does not directly exert a force between spins. For example, a spin-spin exchange coupling does not arise from a direct interaction between spins, but instead from a combination of the Pauli exclusion principle with the Coulomb repulsion between orbital wave functions. Thus, to approach Coulomb interaction phenomena, it is important to start by considering the properties of nanotube electronic wave functions.

The spatial wave function in a nanotube has several degrees of freedom: the position x along the nanotube axis,³ а subband index from the quantization around the circumference, and a valley index τ specifying which of the two valleys (K, K') is occupied by an electron. In nanotubes, it is convenient to separate the 1/r Coulomb interaction into longrange and short-range components. A natural length scale for this separation is the nanotube diameter. This separation considerably simplifies the treatment of the Coulomb interaction in nanotubes (Wunsch, 2009). A first approximation is that neither the short-range nor long-range component mixes states from different subbands: this is justified by the large subband spacing, on the order of eV. A second approximation is that only the short-range component of the Coulomb interaction mixes states from different valleys: this is justified since intervalley scattering requires a large momentum shift (Mayrhofer and Grifoni, 2008; Wunsch, 2009; Weiss et al., 2010; Secchi and Rontani, 2013). The third approximation is that the envelope function $\psi(x)$ describing the position of the electron along the nanotube axis is independent of spin and valley. Although this is not true in general (cf. Appendix B.6), it is a reasonable approximation for smooth confinement potentials and large quantum dots (Wunsch, 2009). Within these approximations, we can treat the long-range and shortrange components of the Coulomb interaction separately: the long-range component couples to $\psi(x)$, while the short-range component couples to the valley degree of freedom.

Indications of Wigner correlations in multihole quantum dots were identified already in early works (Deshpande and Bockrath, 2008). In this review, we follow a pedagogical approach to the topic. We start with long-range interactions, presenting a pedagogical model for the Wigner molecule in one dimension, reviewing calculations performed for nanotubes, and discussing experimental results demonstrating a two-electron Wigner molecule. We then focus on effects from the short-range interaction. At the end of the section, we give an outlook toward interactions in the many-electron and many-hole regimes.

³⁰Taking $l \approx 100 \text{ nm} = 200 \text{ nm}/2$ electrons.

³¹For simplicity, in this section we use notation x for the longitudinal coordinate instead of t as used elsewhere in the review.

1. Long-range interactions and Wigner molecules

Our simple model focuses on the spatial two-electron wave function, assuming that antisymmetry of the total wave function is provided by appropriate symmetries in the spin and valley degrees of freedom. Approximating the dispersion relation of a semiconducting nanotube by a parabola, we write down a Schrödinger equation with an effective mass $m_{\text{eff}} = E_{\text{gap}}/2v_F^2$:

$$-\frac{\hbar^2}{2m_{\rm eff}}\frac{\partial^2\Psi}{\partial x_1^2} - \frac{\hbar^2}{2m_{\rm eff}}\frac{\partial^2\Psi}{\partial x_2^2} + V_0(x_1)\Psi + V_0(x_2)\Psi + V_{\rm C}(x_1,x_2)\Psi = E\Psi, \quad (35)$$

where $\Psi(x_1, x_2)$ is the two-electron spatial wave function, $V_0(x)$ is the external confining potential, and $V_C(x_1, x_2)$ is the Coulomb interaction between electrons. To restrict ourselves to the long-range component of V_C , we include a cutoff in the 1/r Coulomb potential as follows:

$$V_{\rm C}(x_1, x_2) = \frac{1}{4\pi\varepsilon} \frac{e^2}{\sqrt{(x_1 - x_2)^2 + d^2}}$$
(36)

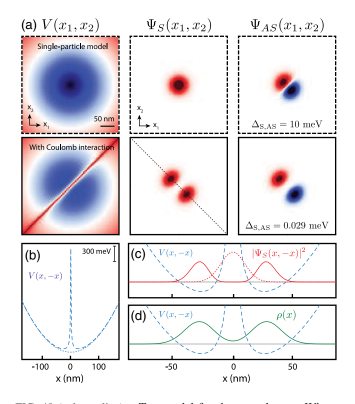


FIG. 48 (color online). Toy model for the two-electron Wigner molecule in one dimension. (a) Two-electron potential $V(x_1, x_2)$ (left panels), symmetric ground-state wave function $\Psi_S(x_1, x_2)$ (middle panels), and antisymmetric excited-state wave function $\Psi_{AS}(x_1, x_2)$ (right panels). Upper panels: no Coulomb interaction $(\varepsilon = \infty)$; lower panels: $\varepsilon = 1$. (b) Line cuts of $V(x_1, x_2)$ along $x_1 = -x_2$ with (dashed) and without (dotted) Coulomb interactions. (c) Line cut of the ground-state two-electron probability density $|\Psi(x, -x)|^2$ along $x_1 = -x_2$ in the single-particle model (dotted line) and the Wigner molecule (solid line). Dashed line: V(x, -x). The suppression of $|\Psi(x, -x)|^2$ near x = 0 is an indication of strong correlations. (d) Electron density $\rho(x) = \int |\psi(x, x_2)|^2 dx_2$ in the Wigner molecule limit.

with *d* chosen as the nanotube diameter. It is instructive to combine the confinement terms in a two-electron electrostatic potential $V(x_1,x_2) = V_0(x_1) + V_0(x_2) + V_C(x_1,x_2)$. The problem of two interacting electrons in one dimension is then formally equivalent to that of a single electron confined in a two-dimensional potential $V(x_1, x_2)$.

This problem can be solved on a desktop computer by exact diagonalization (Jauregui, Häusler, and Kramer, 1993; Szafran et al., 2004; Balder, 2008). The results are shown in Fig. 48, taking $m_{\rm eff} = 0.03 \ m_{\rm e}$ (corresponding to $E_{\rm G} = 210 \ {\rm meV}$ and $D \approx 3.3$ nm) and $V_0(x) = \frac{1}{2}m_{\text{eff}}\omega^2 x^2$ with $\hbar\omega = 10$ meV (confinement length ≈ 30 nm). These parameters correspond to $r_s \approx 18$ (Balder, 2008). The results with ($\varepsilon = 1$) and without $(\varepsilon = \infty)$ Coulomb interaction are shown in Fig. 48(a). Including interactions [lower panels in (a)], the Coulomb repulsion can be seen in $V(x_1, x_2)$ as a diagonal line along $x_1 = x_2$. With these parameters, typical for nanotubes, the Coulomb interaction dominates over the confinement potential. In response, the single lobe of $\Psi_{S}(x_1, x_2)$ in the singleparticle model splits into two well-separated lobes, pushing the two electrons away from each other to minimize Coulomb repulsion [Figs. 48(c) and 48(d)]. In this state, the twoelectron probability density $|\Psi(x_1, x_2)|^2$ goes rapidly to zero along the $x_1 = x_2$ line, indicating the formation of a Wigner molecule. The quantity $|\Psi(x, -x)|^2$ in Fig. 48(c) can be viewed as a two-particle correlation density,³² and its suppression near x = 0 corresponds to the formation of a "correlation hole" [Fig. 48(c)].

An important property of the Wigner molecule state is a strong suppression of the splitting $\Delta_{S,AS}$ between the spatially symmetric ground state $\Psi_S(x_1, x_2)$ and the spatially antisymmetric excited state $\Psi_{AS}(x_1, x_2)$. Without interactions, $\Delta_{S,AS} = \Delta_{ls} = 10$ meV, where $\Delta_{ls} = \hbar\omega$ is the single-particle level spacing. Including interactions, the splitting drops dramatically to $\Delta_{S,AS} = 29 \ \mu$ eV, smaller by a factor ~300. An explanation can be seen by plotting the two-electron wave functions $\Psi(x_1, x_2)$ [lower panels of Fig. 48(a)]. In the Wigner molecule regime, the ground-state wave function $\Psi_S(x_1, x_2)$ deforms to minimize the Coulomb energy, gaining kinetic energy and reducing $\Delta_{S,AS}$ until the symmetric ground state and antisymmetric excited state are nearly degenerate.

Until now, the discussion has been quite general and applies equally to other one-dimensional systems. The relevance for carbon nanotubes becomes clear when looking at the magnitude of such effects. Exact diagonalization calculations for nanotubes (Roy and Maksym, 2009, 2012; Secchi and Rontani, 2009, 2010, 2012, 2013; Wunsch, 2009) arrive at the same conclusions as our toy model: the two-electron dot is strongly correlated, forming a Wigner molecule. The results of such a calculation are shown in Fig. 49. The input parameters are a 100 nm square-well confinement potential with a 50 meV barrier height, a nanotube diameter of 5 nm, a band gap of 90 meV, and an effective mass of $0.009m_e$. To illustrate the transition from the single-particle limit to the Wigner molecule limit, the strength of the Coulomb interaction is tuned by changing the environmental dielectric constant ε ,

³²Integrating $|\Psi(x_1, x_2)|^2$ along the diagonal $x_1 = x_2$ in Fig. 48(a) yields the two-particle correlation function $g(r) = \int |\Psi(r/2 + X, -r/2 + X)|^2 dX$.

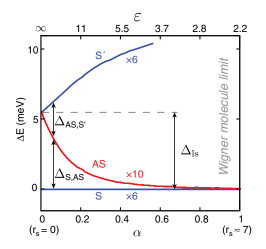


FIG. 49 (color online). Exact calculation for two electrons in a nanotube quantum dot. Vertical axis: energies of the antisymmetric and symmetric excited multiplets relative to the symmetric ground multiplet. Along the horizontal axis, the dielectric constant is changed to tune the interaction strength. Numbers (6, 10, and 6) indicate the degeneracy of each multiplet. Spin-orbit coupling is not included ($\Delta_{SO} = 0$) to illustrate only effects from the long-range interactions. Increasing $\alpha = 2.2/\varepsilon$, there is a transition from the single-particle limit (left) to the Wigner molecule limit (right). In the Wigner molecule, the splitting $\Delta_{S,AS}$ becomes exponentially small. Adapted from Wunsch, 2009.

parametrized by an effective fine structure constant $\alpha = e^2/\epsilon \hbar v_F = 2.2/\epsilon$ [so that $r_s = (\alpha/2.2)me^2l/\hbar^2$]. With no long-range interactions ($\alpha = 0$), the excited multiplet is 16-fold degenerate ($\Delta_{AS,S'} = 0$) and separated from the ground-state multiplet by the single-particle level spacing ($\Delta_{S,AS} = \Delta_{ls}$). With strong interactions ($\alpha = 1$), the antisymmetric multiplet is pushed down in energy relative to the ground state, with $\Delta_{S,AS}$ becoming exponentially small. The two-electron state is deep in the Wigner molecule limit for $r_s \sim 7$, and already strongly correlated for intermediate r_s .

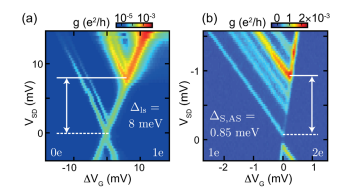


FIG. 50 (color online). Observation of a Wigner molecule. (a) Coulomb blockade spectroscopy of the 0-1 electron transition from which a single-particle level spacing $\Delta_{ls} = 8$ meV is extracted. (b) Spectroscopy of the 1-2 electron transition: analysis of the magnetic field dependence of the levels allows identification of the excited state corresponding to splitting $\Delta_{S,AS}$. The value $\Delta_{S,AS} = 0.85$ meV is extracted, 10 times smaller than Δ_{ls} , indicating that the two-electron quantum dot is in the Wigner molecule regime with $r_s \approx 1.6$ ($\alpha \approx 0.5$ in Fig. 49). Adapted from Pecker *et al.*, 2013.

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Summarizing these predictions, Wigner correlations in a twoelectron quantum dot cause a collapse of $\Delta_{S,AS}$.

Experiments with clean nanotubes have provided clear evidence of a Wigner molecule. Specifically, Pecker *et al.* (2013) compared a one-electron to a two-electron quantum dot in the same device (Fig. 50). A crucial step was a careful analysis of the magnetic field dependence of the excitation lines (Fig. 31). This allowed them to determine which splitting in Fig. 50(a) corresponds to Δ_{Is} , and which splitting in Fig. 50(b) corresponds to $\Delta_{S,AS}$. They found $\Delta_{Is} = 8 \text{ meV}$ and $\Delta_{S,AS} = 0.85 \text{ meV}$. The tenfold suppression of $\Delta_{S,AS}$ compared to Δ_{Is} indicates that the two-electron quantum dot indeed forms a Wigner molecule.

2. Short-range interactions and intervalley exchange

In this section, we consider effects of the short-range Coulomb interaction on the spectrum of a two-electron quantum dot. As seen in the toy model, the long-range interaction distorts the longitudinal wave functions, reducing the splitting $\Delta_{S,AS}$ between the symmetric and antisymmetric multiplets. Because the long-range interaction does not couple to valleys, it does not change the spin or valley level structure inside a multiplet. In contrast, the short-range Coulomb interaction can induce valley scattering and thus changes the splitting between different valley and spin states.

Several exact diagonalization calculations have been performed accounting for Wigner molecule effects, spin-orbit coupling, and short-range Coulomb interactions (Secchi and Rontani, 2009, 2010, 2012, 2013; Wunsch, 2009). The resulting spectrum is shown in Fig. 51. In the calculation, α is set to zero to artificially suppress long-range interactions while keeping the short-range on-site energy³³ U_0 constant.³⁴ Figure 51(a) shows the spectrum as a function of α at zero magnetic field. The colored (gray) lines represent the *S* (*AS*) multiplet of Fig. 49. For $\alpha = 1$, the dot is in the Wigner molecule regime due to long-range interactions, and the gray lines in Fig. 51(a) become nearly degenerate with the colored lines. Figures 51(b) and 51(c) show the B_{\parallel} dependence of the spatially symmetric multiplet for $\alpha = 0$ and $\alpha = 1$.

The influence of short-range interactions can be most clearly seen for $\alpha = 0$ [Fig. 51(b)]. The first effect is to lift the degeneracy of the spin-polarized and valley-polarized states. In the single-particle model (and also at $\alpha \gtrsim 1$, see below), these four states are degenerate at B = 0. With strong short-range interactions, the energy of the valley-polarized doublet is raised with respect to the spin-polarized doublet. This is because a positive U_0 used in the calculation penalizes double occupancy of atomic sites, which by symmetry considerations does not occur for spin-polarized states (see Appendix B.9 for details). This results in a new splitting Δ_{VBS}

³³The Coulomb potential for the short-range interaction is taken with the same form as Eq. (36), but with cutoff *d* chosen such that $V_{\rm C}(x_1 = x_2) = U_0$, where $U_0 = 15$ eV is the charging energy associated with putting two electrons on the same p_z orbital. Such a potential is also referred to as the Ohno potential (Mayrhofer and Grifoni, 2008).

³⁴Physically, this could be achieved by placing the nanotube close to a metallic gate or dielectric slab, such that the distance to the surface is large compared to the diameter but small compared to the nanotube length.

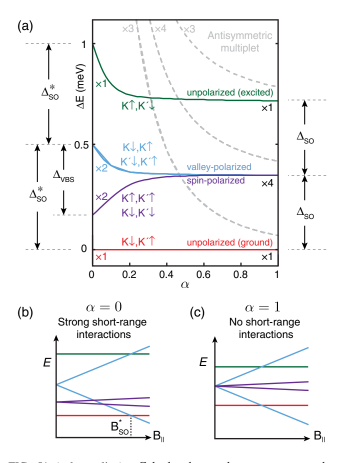


FIG. 51 (color online). Calculated two-electron quantum dot energies including short-range interactions. (a) Energies relative to the ground state of the spatially symmetric multiplet (solid lines) and the spatially antisymmetric multiplet (dashed lines). The parameter α characterizes the strength of the long-range interactions: $\alpha = 1$, Wigner molecule regime; $\alpha = 0$: no longrange correlations. The valley-polarized, spin-polarized, and unpolarized (ground and excited) levels are labeled, and the degeneracy of each set of lines is indicated. For $\alpha = 1$, the dashed lines become degenerate with the solid lines, as in Fig. 49 without spin-orbit coupling and short-range interactions. For $\alpha = 0$, longrange correlations are suppressed, the single-particle wave functions begin to overlap, and short-range interactions become stronger. Here new splittings Δ_{VBS} and Δ_{SO}^* appear in the valleyspin structure of the multiplet. (b), (c) Magnetic field dependence of the ground-state multiplet for (b) $\alpha = 0$ and (c) $\alpha = 1$. Shortrange interactions in (b) break the degeneracy between the valleypolarized states and spin-polarized states at B = 0. For $\alpha = 1$, the valley and spin structure of the multiplet is the same as in the single-particle model, even though the longitudinal wave functions $\Psi(x_1, x_2)$ are highly correlated. States are labeled as in Table IV, except that the states used in the calculation are simplified such that they are independent of longitudinal $\Psi(x_1, x_2)$. (a) Adapted from Wunsch, 2009.

within the S(0,2) multiplet, which can be viewed as an effective intervalley exchange energy.³⁵ For the AS(0,2)

multiplet, the effect of short-range Coulomb interaction is predicted to be much smaller (Secchi and Rontani, 2013), indicating that the longitudinal symmetry of the two-electron wave function plays an important role (Appendix B.9).

The second effect of the short-range interaction is seen in Fig. 51 as an increase in the splitting between the unpolarized ground state and the unpolarized excited state. The splitting is now equal to $2\Delta_{SO}^* \equiv 2\sqrt{\Delta_{SO}^2 + \Delta_{VBS}^2}$ instead of $2\Delta_{SO}$. An important consequence is that the two-electron ground state becomes valley polarized at a higher magnetic field than without short-range interactions [the crossing field B_{SO}^* is higher in Fig. 51(b) than in Fig. 51(c)]. In the Wigner molecule regime ($\alpha = 1$) the effects of short-range interactions are suppressed: Because of strong correlations, the two electrons have little overlap irrespective of their longitudinal symmetry, thereby suppressing double occupancy of the same atomic site. Similar intervalley exchange effects have also been seen in other calculations (Mayrhofer and Grifoni, 2008; Secchi and Rontani, 2009, 2010, 2012, 2013; von Stecher et al., 2010).

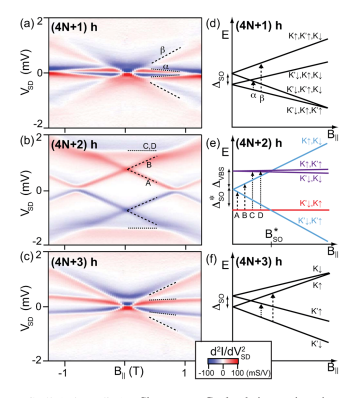


FIG. 52 (color online). Short-range Coulomb interactions in a many-hole quantum dot (estimated as containing 10–40 holes). (a)–(c) Cotunneling spectroscopy of different charge configurations. (d)–(f) Expected spectra from a model with short-range Coulomb scattering. Quantum numbers of occupied electronic states in the valence band are indicated (cf. Fig. 18). Lines in (a)–(c) measure length of the arrows in (d)–(f). Colors in (e) correspond to those used in Fig. 51. The signature of interactions can be seen by comparing the spectra of different charge configurations: the valley and spin spectra of the 4N + 1 and 4N + 3 configurations are consistent with those predicted by the single-particle model, while the spectrum of the 4N + 2 configuration is modified by short-range Coulomb interaction. Adapted from Cleuziou *et al.*, 2013.

³⁵Note that Fig. 51 plots the energies with respect to the ground state, explaining why spin-polarized states appear to be lowered with respect to valley-polarized states. The abbreviation VBS denotes "valley backscattering," a name used for the term in the Coulomb scattering matrix that gives rise to an intervalley exchange splitting.

Experiments by Cleuziou *et al.* demonstrated clear intervalley exchange effects in the spectra of clean nanotubes in the many-hole regime (Fig. 52). The key observation was that the 4N + 1 and 4N + 3 charge states showed a spectrum consistent with shell filling of the single-particle valley and spin levels, whereas a different spectrum was observed for the 4N + 2 charge state (Cleuziou *et al.*, 2013). This difference between odd and even filling of the nanotube multiplet is a clear signature that can only be explained including short-range interactions.

The spectrum measured for 4N + 2 filling is similar to that of Fig. 51(b), except for the sign of the exchange interaction. In particular, theoretical treatments of intervalley exchange as outlined above predict that the valley-polarized states are raised in energy above the spin-polarized states [as in Fig. 51(b)]. In contrast, the observed spectra (Cleuziou *et al.*, 2013) show the opposite [Fig. 52(e)]. This may indicate that the exchange integral implied from the experimental data has the opposite sign compared to that predicted by theory so far. A similar observation was also reported for electrons (Pecker *et al.*, 2013). The reason for this sign difference is not understood, but suggests that the validity of theoretical approximations, as well as alternative mechanisms, should be carefully checked.

As discussed previously, intervalley exchange is not expected to play a significant role in the few-electron regime, since the long-range Wigner correlations will suppress shortrange Coulomb interactions. The device of Fig. 52 was in the many-hole regime: a possible explanation of why Δ_{VBS} is so large is that the long-range Coulomb interaction was screened by the holes in nearby shells. However, a large Δ_{VBS} was also reported in the few-electron regime by both Pecker et al. (2013) ($\Delta_{\text{VBS}} = 0.2 \text{ meV}$) and Cleuziou *et al.* (2013) $(\Delta_{\text{VBS}} = 1.56 \text{ meV} \text{ calculated from the observed } \Delta_{\text{SO}}^*).$ Estimates for uncorrelated states based on first-order perturbation theory predict Δ_{VBS} to be hundreds of μeV (Appendix B.7). Taking into account correlations predicts a much smaller value, $1 - 10 \mu eV$ (Secchi and Rontani, 2009, 2010, 2012, 2013; Wunsch, 2009; Pecker et al., 2013). It is an open question why $\Delta_{\rm VBS}$ is so large in these experiments.

Finally, the large Δ_{VBS} splitting seen in excited-state spectroscopy has important implications for ground-state spectroscopy, in which the magnetic field dependence of Coulomb peak position in gate voltage is used to infer the spin-orbit splitting (Churchill, Kuemmeth et al., 2009; Steele et al., 2013). In particular, the magnetic field where kinks occur in the ground-state chemical potentials is now given by $B_{\rm SO}^* = \Delta_{\rm SO}^* / 2\mu_{\rm orb}$ instead of by $B_{\rm SO} = \Delta_{\rm SO} / 2\mu_{\rm orb}$. This raises the possibility that the large spin-orbit interaction reported by Steele et al. (2013) extracted from the ground-state energies could be due to a large Δ_{SO}^* and Δ_{VBS} instead of a large Δ_{SO} . The intervalley exchange splitting required to match the experimental data, however, would be $\Delta_{VBS} \sim 3$ meV, even larger than the already unexpectedly large values reported from excitation spectra (Cleuziou et al., 2013; Pecker et al., 2013).

C. Beyond Wigner molecules: Correlation effects of many electrons in quantum dots

The search for strong correlation effects with many electrons inspired some of the very first experiments on nanotubes. These early experiments were motivated by the predictions of Luttinger-liquid theory (Luttinger, 1963; Imambekov, Schmidt, and Glazman, 2012) and focused on power-law behavior of the conductance (Bockrath *et al.*, 1999; Yao *et al.*, 1999). Detailed understanding of these experiments was hampered, however, by the large disorder present in these devices, leading to localization and Coulomb blockade at low temperatures. Dynamical Coulomb blockade also leads to power-law dependence on bias and temperature (Ingold and Nazarov, 1992). A clearer signature for a Luttinger liquid would be spin-charge separation, giving spin and charge modes with different velocities. Evidence for such spin-charge separation has been seen in GaAs wires (Auslaender *et al.*,

2005), but has not been reported for nanotubes.

Correlation effects for two electrons in a carbon nanotube quantum dot are now fairly well understood theoretically, and well established experimentally. A natural question is then: what happens if more electrons or holes are added to the quantum dot? One route could be to probe the electron density itself in the quantum dot using an STM (Ziani, Cavaliere, and Sassetti, 2013). Another route is detailed spectroscopy of suspended nanotube quantum dots in the multielectron and multihole regime. From an experimental point of view, there have been suggestions of correlation effects in clean manyhole quantum dots (Deshpande and Bockrath, 2008). Studying the ground-state spin and valley filling of a quantum dot as a function of hole number and magnetic field, regions of strong spin and valley polarization were observed that are not easily explained by a single-particle picture. A possible explanation of this strong spin and valley polarization is the formation of a many-hole Wigner crystal in which the kinetic energy is quenched by interactions, similar to the suppressed Δ_{SAS} in the two-electron Wigner molecule. An exciting next step will be to use low-disorder nanotubes with multiple gates to perform detailed spectroscopy with tunable confinement, exploring the transition from the well-established two-electron Wigner molecule to the regime where many-electron Wigner crystals and Luttinger-liquid-like correlations may occur.

D. Open questions

- Why is the observed intervalley exchange splitting Δ_{VBS} so large? Even with a completely screened long-range interaction, calculations predict $\Delta_{VBS} \lesssim \Delta_{SO}$, while the opposite has been observed in some experiments.
- Why does Δ_{VBS} have the wrong sign in experiments? A possible scenario is a superexchange mechanism for the short-range interaction. In superexchange, a net exchange of two electrons is achieved by two separate exchange processes with electrons in a third orbital. The third orbital could correspond to a state in a different shell, a state in the valence band, or a state in one of the higher subbands. Superexchange can have the opposite sign compared to direct exchange.
- Why are not more devices deep in the Wigner molecule limit? Why is shell-filling theory so effective for describing so many devices?
- Are there qualitatively new predictions from theories that take into account the dependence of the spatial wave function on *τ*, *s*, and magnetic field (see Appendix B.6)?

• How do the well-understood strong correlations of the two-electron Wigner molecule extend to quantum dots with more electrons? Are there clear experimental signatures to look for?

VIII. CONCLUSIONS AND OUTLOOK

The chirality is an unknown parameter in all the experiments discussed in this review. Although optical techniques have been developed to determine chirality on specially made, long nanotubes (Amer et al., 2013; Liu et al., 2013), these characterizations are unfortunately incompatible with the short length of nanotube devices in the quantum regime. Therefore the growth of chirality-specific nanotubes remains an important challenge, although its realization may not be expected soon. Of course, several important (band-)structure parameters can be inferred from transport, such as the gap and the diameter. Uncertainties nevertheless arise since a measured gap may be affected by electron-electron interactions, or even mundane issues like surface coverage with water (Elias et al., 2009). Such band-gap modifications are interesting subjects by themselves but then knowledge of the bare band gap, and thus the chirality, would be an indispensable input parameter.

Fortunately, the general physics discussed in this review does not depend on the precise numerical values of the bandstructure parameters. Most of this physics is a consequence of the simple fact that electrons in nanotubes live on a hexagonal lattice that is confined to a one-dimensional, tubular geometry. We stress that in this sense nanotubes are unique solid-state structures. Given this structure the spin and valley phenomena described in this review represent general physics independent of the precise chirality.

The inclusion of spin-orbit interaction completes our picture of the noninteracting, single-particle physics in nanotubes. Until 2008, this effect was widely thought to be negligible, but it has turned out to be highly relevant in many experiments. It is also important for applications involving coherent control of quantum states. The emerging picture of the role of interactions has also become much clearer in recent years since ultraclean devices have become available. We have described in detail the case of two interacting electrons. The study of many interacting electrons (Deshpande and Bockrath, 2008) is a quest full of interesting challenges with its holy grail being the formation of a long Wigner crystal.

The theoretical understanding of the single-particle physics has been important to obtain a complete picture of the allowed spin-orbit terms. There are still theoretical challenges relating to a quantitative microscopic understanding of the effects originating from the cylindrical geometry. As part of this, renormalization by long-range Coulomb interactions of the curvature-induced gap and the spin-orbit coupling has not been investigated.

Quantum states are best defined in closed systems and this review has therefore mainly focused on quantum dots whose coupling to leads is weak (but large enough to allow a measurable current). Quantum dot states change when the coupling is increased, and qualitatively new phenomena can arise such as various Kondo effects and possibly even quantum phase transitions (Mebrahtu *et al.*, 2012). Great potential for new experiments arises when the leads are given interesting properties. For example, superconducting leads can induce superconducting correlations that are restricted by the special spin-valley quantum numbers of nanotubes. One can imagine Josephson junctions with the junction consisting of a nanotube with multiple bends. Experiments using superconducting contacts performed so far indeed indicate a rich research direction (de Franceschi et al., 2010). Also in such hybrid nanotube devices, spin-orbit interactions may play a role, e.g., leading to modified Josephson currents (Lim, López, and Aguado, 2011) or new detection schemes (Braunecker, Burset, and Levy Yeyati, 2013). Superconducting contacts to ultraclean nanotubes have also recently been demonstrated (Schneider et al., 2012), opening the possibility of studying proximity-induced superconductivity with exceptionally low electronic disorder. Magnetic materials could be used as contacts to explore spintronics confined to one dimension (Sahoo et al., 2005).

The field of quantum computing has been inspirational for the development of all kinds of qubit devices. This review discussed various types of nanotubes. The coherent evolution in qubits can be used as an extremely sensitive probe of the environment. Nanotube qubits could be used to study in detail nuclear spins or mechanical vibrations. The holy grail here could be the realization of a coherent coupling between spin and motion. Another inspiration from the field of qubits is the use of nanotubes as on-chip charge sensors (Biercuk et al., 2006; Gotz et al., 2008; Churchill, Bestwick et al., 2009), or as a component in circuit quantum electrodynamics (Delbecq et al., 2011). If long coherence times could be achieved, there is potential for a dramatically improved charge sensor (Dial et al., 2013). Nanotubes have the advantage for sensing that short quantum dots show Coulomb blockade up to room temperature (Postma et al., 2001).

We hope this review inspires new generations of experiments taking advantage of the unique properties of carbon nanotubes. Similar to the evolution of quantum Hall physics in GaAs in the 1980s and 1990s, the remarkable quality of devices that can now be achieved using ultraclean techniques indicates that nanotubes will remain an exciting playground for exploring rich new physics for years to come.

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APPENDIX A: TRANSPORT SPECTROSCOPY IN QUANTUM DOTS

Here we give a basic overview of quantum dots and transport spectroscopy, focusing on techniques by which data in this review are derived.

1. Coulomb blockade and the constant interaction model

An electrical schematic of a quantum dot as in Fig. 2 is shown in Fig. 53. If both the tunnel rates to the leads $\Gamma_{S,D}$ and the thermal energy k_BT are less than the charging energy $E_C = e^2/C$, where C is the total dot capacitance to the outside world, then the electron occupation becomes constrained to take an integer value. The equilibrium occupation can be adjusted by tuning V_G .

This quantization of the dot occupation strongly modifies the transport characteristics of this circuit. With all other parameters held fixed, any change of the dot occupation away from equilibrium increases the electrostatic energy of the system. This causes a suppression of the current known as Coulomb blockade. However, for particular values of $V_{\rm G}$, the suppression can be lifted. This allows the energy levels of the quantum dot to be mapped out by measuring the device conductance as a function of $V_{\rm G}$ [for extensive reviews, see Kouwenhoven *et al.* (1997), Kouwenhoven, Austing, and Tarucha (2001), and Hanson *et al.* (2007)].

Coulomb blockade is lifted whenever each step of electron tunneling through the device is energetically favorable. To understand how the observed conductance features relate to the energy levels of the device, we introduce the electrochemical potential $\mu(N)$ of the dot for occupation N, defined as the difference in energy U between N-electron and (N - 1)-electron ground states:

$$\mu(N) = U(N) - U(N - 1).$$
(A1)

Correspondingly, the electrochemical potentials of the leads are defined as the energy to add an additional electron at the Fermi level; with a bias V_{SD} applied to the source as in Fig. 2(a),

$$\mu_{\rm S} = E_{\rm F} - eV_{\rm SD},\tag{A2}$$

$$\mu_{\rm D} = E_{\rm F},\tag{A3}$$

where $E_{\rm F}$ is the Fermi energy.

The electrochemical potential is in general related in a complicated way to the single-particle energy levels, because each electron added electrostatically perturbs the energies of the electrons already on the dot. However, the relationship becomes simpler in the *constant-interaction model*, which makes two assumptions. First, all Coulomb interactions, both between electrons on the dot and between the dot and the environment, are parametrized by a single constant capacitance C, which is the sum of capacitances C_S , C_D , C_G to the

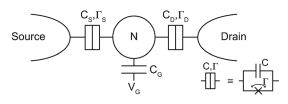


FIG. 53. Electrical model of a single quantum dot: a conducting island is coupled with tunnel rates Γ_S and Γ_D to source and drain, and with capacitance C_S , C_D , and C_G to source, drain, and gates.

source, drain, and gate.³⁶ Second, the single-particle energy levels are assumed to be independent of these interactions, and therefore not changed by adding additional electrons. Under these assumptions, the dot energy is

$$U(N) = \frac{\left[-e(N-N_0) + C_{\rm S}V_{\rm S} + C_{\rm D}V_{\rm D} + C_{\rm G}V_{\rm G}\right]^2}{2C} + \sum_{i=1}^{N} E_i,$$
(A4)

where N_0 is the occupancy with no voltages applied (set by fixed charges in the environment, e.g., substrate charges, and not necessarily quantized) and E_i are the single-particle energy levels. The first term is the electrostatic energy stored in the dot capacitances, while the second is the sum of the single-particle confinement energies.

In this approximation, the electrochemical potential is

$$\mu(N) = \left(N - N_0 - \frac{1}{2}\right) E_{\rm C} - \frac{E_{\rm C}}{e} (C_{\rm S} V_{\rm S} + C_{\rm D} V_{\rm D} + C_{\rm G} V_{\rm G}) + E_N,$$
(A5)

where $E_{\rm C} = e^2/C$ is the charging energy. The electrochemical potential increases for successive values of *N*, forming a ladder of levels as shown in Fig. 54. The separation between adjacent levels is the addition energy:

$$E_{\rm add}(N) = \mu(N) - \mu(N-1)$$
$$= E_{\rm C} + \Delta E(N) \tag{A6}$$

and includes both an electrostatic term $E_{\rm C}$ and the quantum energy level spacing $\Delta E(N) \equiv E_N - E_{N-1}$. From Eq. (A5), changing the gate voltage moves the entire ladder of electrochemical potentials up or down.

2. Low-bias spectroscopy

The condition that both tunneling events be energetically favorable is equivalent to saying that the chemical potential must decrease at each step.³⁷ In other words, some N must exist for which

$$\mu_{\rm S} > \mu(N) > \mu_{\rm D}.\tag{A7}$$

Consider first the situation of low bias, where $eV_{SD} \ll \Delta E$, E_{add} . The corresponding level diagram in the blockaded case [Fig. 54(a)] shows that no ground-state chemical potential satisfies Eq. (A7), so no current flows. However, by increasing V_G to lower the ladder of electrochemical potentials, the blockade can be lifted [Fig. 54(b)]. As a function of V_G , the current shows a series of Coulomb peaks, with each valley between the peaks corresponding to the

³⁶In devices with more than one gate, additional capacitances must be added to the model in a straightforward way.

³⁷Thermal excitations in the leads relax this constraint, broadening the Coulomb peaks. Here this effect is ignored, which is permissible if the temperature is less than both ΔE and $E_{\rm C}$.

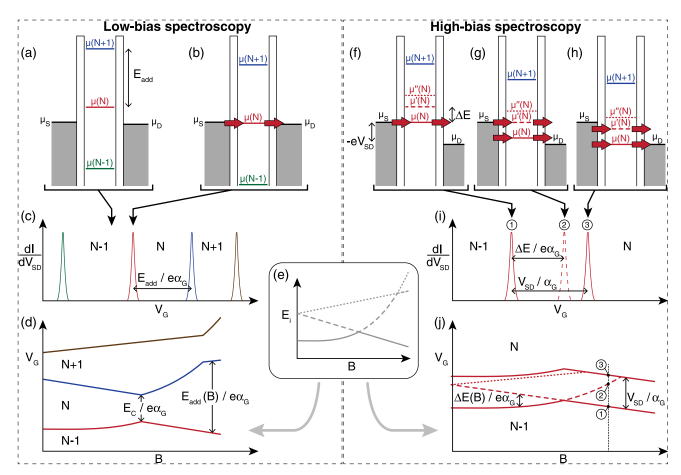


FIG. 54 (color online). Two techniques of electron transport spectroscopy. In low-bias spectroscopy (a)–(d), at most a single quantum dot level falls within the bias window set by the lead electrochemical potentials. (a) With no ground-state level in the bias window, levels are filled up to μ_D , and the electron number is fixed (in this case, at N - 1) due to Coulomb blockade. (b) With one of the levels $\mu(N)$ in the bias window, the electron number fluctuates between N and N - 1, leading to conductance through the dot. (c) Sweeping V_G moves each level in turn through the bias window, leading to a series of current peaks. From the peak spacing, the addition energy E_{add} for each electron number can be deduced, allowing the ground-state energy to be studied. (d) Transport peak evolution measured using this technique, where the energy levels E_i are assumed to vary with magnetic field B as shown in (e). In high-bias spectroscopy (f)–(j), more than one level can fall within the bias window, allowing excited-state energies to be measured. Sweeping V_G through the $N - 1 \leftrightarrow N$ transition, a series of transport peaks appear in dI/dV_{SD} . (f) The current first appears when $\mu(N)$ crosses μ_S . (g) When $\mu'(N)$ crosses μ_S , additional current can flow through an excited state of the dot. (h) The last peak in the series occurs when $\mu(N)$ crosses μ_D , leaving the bias window. (i) The corresponding series of transport peaks, with the spacings corresponding to ΔE and V_{SD} marked. When $\mu(N) < \mu_D$, Coulomb blockade is reestablished with the dot in its N-electron ground state; there are therefore no peaks corresponding to states with excitation energies larger than eV_{SD} , such as that marked by the dotted line in (d)–(f). (j) Sketch of the peak evolution measured using this technique for the same underlying energy levels as shown in (e). The V_G trace in (i) corresponds to the vertical dashed line in (j).

Coulomb blockade with a different fixed occupation N [Fig. 54(c)]. For each N, the peak separation in V_G is equal to $E_{add}(N)/e\alpha_G$, where $\alpha_G = C_G/C$ is the lever arm that characterizes the coupling of the gate to the dot. By measuring the Coulomb peak positions in gate space as a function of some external parameter, the evolution of energy levels can be deduced.

3. High-bias spectroscopy

Low-bias transport is sensitive only to the ground-state energy of the device. With a bias larger than the single-particle level spacing, $e|V_{SD}| > \Delta E$, the excited states can also be populated. To interpret the resulting transport features, it is necessary to consider the corresponding electrochemical potentials. We define the first excited-state electrochemical potential as

$$u'(N) = U'(N) - U(N-1)$$
(A8)

$$= \mu(N) + \Delta E(N+1), \tag{A9}$$

where $U'(N) = U(N) + \Delta E(N+1)$ is the first excited state energy of the *N*-electron dot. Higher excited states can be defined the same way (Kouwenhoven, Austing, and Tarucha, 2001). As illustrated in Figs. 54(f)–54(h), transport can proceed via both the ground state and excited states within the bias window. As V_G is increased from the N - 1 electron valley, the current first increases when $\mu(N)$ crosses μ_S [Fig. 54(f)]. With a further increase in V_G , $\mu'(N)$ enters the bias window [Fig. 54(g)]. This allows transport via the first excited state, which continues until $\mu(N)$ crosses μ_D [Fig. 54(h)]. For more positive values of V_G , Coulomb blockade is reestablished, and transport is blocked through both ground and excited states.

This series of resonances between dot and lead electrochemical potentials is usually seen by plotting the conductance $dI/dV_{\rm SD}$ as a function of $V_{\rm G}$, which results in a series of peaks as each transport channel is opened or closed [Fig. 54(i)].³⁸ From the peak spacings, the excited-state energies can be read off as shown. This technique also gives a convenient way to measure $\alpha_{\rm G}$; the gate voltage separation of the first and last peaks corresponds to shifting $\mu(N)$ from $\mu_{\rm S}$ to $\mu_{\rm D}$, and is therefore equal to $V_{\rm SD}/\alpha_{\rm G}$, where $V_{\rm SD}$ is set in the experiment.³⁹ These two complementary spectroscopy methods allow the energy levels of the quantum dot to be measured as a function of various experimental parameters.

APPENDIX B: THEORETICAL BACKGROUND

This Appendix introduces the theoretical background of the review. The focus will be pedagogical and we will not attempt to refer to relevant theoretical papers. For this see more theoretical reviews and the main text. We derive the Dirac equation used extensively in the main text. Physical pictures of the two types of spin-orbit coupling (orbital- and Zeemanlike terms) as well as the curvature-induced band gap are given. Finally, we address the form of the single-particle and two-particle wave functions.

1. Graphene band structure near the Dirac points

Since we are interested in the behavior close to the Fermi level, we focus on the band structure near the two Dirac points. This will be done in two ways: first a simple $\mathbf{k} \cdot \mathbf{p}$ calculation is applied to show that the spectrum can be derived from symmetry arguments alone. This is confirmed using a tight-binding calculation, which is also easier to generalize to the case with broken symmetry, as in a nanotube.

a. The $\mathbf{k} \cdot \mathbf{p}$ derivation

The $\mathbf{k} \cdot \mathbf{p}$ calculation for graphene (DiVincenzo and Mele, 1984; Marder, 2000) uses the fact that the potential has a unit

cell of two carbon atoms and is invariant under translations $\mathbb{T}_{\mathbf{a}_{1,2}}$ by lattice vectors $\mathbf{a}_{1,2}$ as well as under rotation $\mathbb{R}_{2\pi/3}$ by angle $2\pi/3$ about a lattice symmetry point (C_3 symmetry). At the Dirac points **K** and **K'**, we define Bloch states, denoted $\Psi_A^{\mathbf{K}(\mathbf{K}')}(\mathbf{r})$ and $\Psi_B^{\mathbf{K}(\mathbf{K}')}(\mathbf{r})$, which are eigenstates of the translation operator:

$$\mathbb{T}_{\mathbf{a}_{1,2}}\Psi_{A(B)}^{\mathbf{K}}(\mathbf{r}) = e^{i\mathbf{K}\cdot\mathbf{a}_{1,2}}\Psi_{A(B)}^{\mathbf{K}}(\mathbf{r}), \tag{B1}$$

and likewise for **K**'. They are degenerate because of the inversion symmetry (which interchanges *A* and *B* and **k** and $-\mathbf{k}$). The relative phase between two sites separated by the unit vector \mathbf{a}_1 is given by

$$e^{i\mathbf{K}\cdot\mathbf{a}_1} = e^{-i2\pi/3} = \chi^*,\tag{B2}$$

and by χ for **K**'. Symmetry allows the functions $\Psi_{A,B}^{\mathbf{K}}(\mathbf{r})$ to be chosen as eigenstates of the rotations $\mathbb{R}_{2\pi/3}$ around a center of a hexagon in the following way:

$$\mathbb{R}_{2\pi/3}\Psi_A^{\mathbf{K}} = \chi \Psi_A^{\mathbf{K}}, \qquad \mathbb{R}_{2\pi/3}\Psi_B^{\mathbf{K}} = \chi^* \Psi_B^{\mathbf{K}}. \tag{B3}$$

(At the **K**' point, one should replace χ by χ^* .)

We now derive an effective Hamiltonian using $\mathbf{k} \cdot \mathbf{p}$ perturbation theory. Expanding the Bloch Hamiltonian $H_{\mathbf{k}}$ around the Dirac points, we write $\mathbf{k} = \mathbf{K} + \kappa$ and

$$H_{\mathbf{k}} \approx H_{\mathbf{K}} + H_{\boldsymbol{\kappa}}^{\tau},\tag{B4}$$

where $\tau = 1$ (-1) for the *K* (*K'*) point. At *K* the functions $\Psi_A^{\mathbf{K}}$ and $\Psi_B^{\mathbf{K}}$ are degenerate and we define our energy scale so that $\langle \Psi_A^{\mathbf{K}} | H_{\mathbf{K}} | \Psi_A^{\mathbf{K}} \rangle = 0$. The correction to the Hamiltonian is

$$H_{\kappa}^{\tau} \equiv \frac{\hbar}{m} \boldsymbol{\kappa} \cdot \mathbf{p}, \qquad (B5)$$

with $\mathbf{p} = -i\hbar \nabla$ being the momentum operator. In the $\{\Psi_A^{\mathbf{K}}(\mathbf{r}), \Psi_B^{\mathbf{K}}(\mathbf{r})\}$ basis, one can now find the matrix elements of the momentum operator $\mathbf{p}_{CC'}^{\mathbf{K}} = \langle \Psi_C^{\mathbf{K}} | \mathbf{p} | \Psi_{C'}^{\mathbf{K}} \rangle$, where *C*, *C'* both take the values *A* or *B*. Each matrix element $\mathbf{p}_{CC'}^{\mathbf{K}}$ is a vector in the *x*-*y* plane of the graphene sheet. The *AB* component follows from the rotational symmetry of the wave functions:

$$\mathbb{R}_{2\pi/3}\mathbf{p}_{AB}^{\mathbf{K}} = \chi^* \mathbf{p}_{AB}^{\mathbf{K}}, \qquad \mathbb{R}_{2\pi/3}\mathbf{p}_{AB}^{\mathbf{K}'} = \chi \mathbf{p}_{AB}^{\mathbf{K}'}. \tag{B6}$$

The eigenvectors in Eq. (B6) are

$$\mathbf{p}_{AB}^{\mathbf{K}} \propto \begin{pmatrix} 1 \\ -i \end{pmatrix}, \qquad \mathbf{p}_{AB}^{\mathbf{K}'} \propto \begin{pmatrix} 1 \\ i \end{pmatrix}.$$
 (B7)

The diagonal elements $\mathbf{p}_{AA}^{\mathbf{K}}$ and $\mathbf{p}_{BB}^{\mathbf{K}}$ vanish, since they obey an analogous equation to Eq. (B6), but with χ replaced by 1, which has only zero solutions.

The low-energy Hamiltonian (B5) is thus

$$H_{\kappa}^{\tau} \propto \begin{pmatrix} 0 & \tau \kappa_{x} - i \kappa_{y} \\ \tau \kappa_{x} + i \kappa_{y} & 0 \end{pmatrix} = \hbar v_{F} (\tau \kappa_{x} \sigma_{1} + \kappa_{y} \sigma_{2}), \quad (B8)$$

³⁸Although the most common situation is for all peaks to have positive dI/dV_{SD} as drawn, the excited-state peaks can be negative if there is a strong difference of tunnel coupling between different states (Weinmann, Häusler, and Kramer, 1995).

³⁹This discussion ignores processes whereby tunneling off the quantum dot leaves it in an N - 1 electron excited state, which lead to additional conductance peaks not shown in Fig. 54(g). These additional peaks, which are not relevant to data discussed in this review, are clearly distinguished from those in Fig. 54(i) in a two-dimensional plot of conductance versus V_{SD} and V_G and can usually be eliminated by appropriate tuning of the tunnel barriers. For a full discussion, see Kouwenhoven, Austing, and Tarucha (2001).

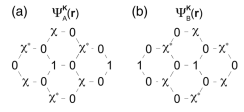


FIG. 55. Examples of orthonormal spatial eigenfunctions satisfying both Eqs. (B1) and (B3) at the *K* point for the minimal tight-binding model. The site coefficients for a one-orbital description of the eigenfunctions are shown, with $\chi = e^{i2\pi/3}$. In the general case, the Ψ_A^K and Ψ_B^K wave functions have the same rotational symmetry as in the figure. The eigenfunctions at the *K'* point are the same but with χ replaced by χ^* .

where the Fermi velocity v_F was introduced as a phenomenological parameter, to match Eq. (1). Here σ_1, σ_2 are the usual Pauli matrices, now working in the A/B sublattice. To describe the Hilbert space spanned by the valley index τ and A/B sublattice spinor some authors use the terms isospin and pseudospin. However, there is no established convention as to which is which; see opposite definitions, for example, in Kane and Mele (1997) and McCann *et al.* (2006).

b. The tight-binding derivation

In the tight-binding approach the starting point is the Bloch functions of *A* and *B* sublattices:

$$\Psi_{A/B}^{\mathbf{k}}(\mathbf{r}) = \sum_{n} e^{i\mathbf{k}\cdot\mathbf{R}_{n}}\varphi_{A/B}(\mathbf{r}-\mathbf{R}_{n}), \qquad (B9)$$

where $\varphi_{A/B}$ are the local basis functions⁴⁰ for atomic orbitals at sites *A* and *B*, respectively (see Fig. 55). Neighboring orbitals hybridize via bonds along direction δ_i with overlap matrix elements t_i , with i = 1, 2, 3. The Hamiltonian overlap of Ψ_A^k and Ψ_B^k is therefore

$$\langle \Psi_A^{\mathbf{k}} | H | \Psi_B^{\mathbf{k}} \rangle = \sum_i t_i e^{i \mathbf{k} \cdot (\boldsymbol{\delta}_i - \boldsymbol{\delta}_1)}. \tag{B10}$$

For graphene, rotational symmetry means all bonds are equivalent $(t_i = t)$, and expanding around the *K* point gives (using the definitions in Table I)

$$\langle \Psi_A^{\mathbf{k}} | H | \Psi_B^{\mathbf{k}} \rangle = t \sum_i e^{i\mathbf{k} \cdot (\delta_i - \delta_1)} = \frac{it}{\sqrt{3}} (\kappa_x - i\kappa_y).$$
(B11)

To be consistent with the $\mathbf{k} \cdot \mathbf{p}$ result (B8), we then make a transformation of the phase of the basis states so that $\Psi_B^{\mathbf{k}} \rightarrow -i\Psi_B^{\mathbf{k}}$, which recovers the Hamiltonian (B8).

c. Graphene wave functions

Solving for the eigenenergies and eigenstates of the lowenergy Hamiltonian (B8)

$$H^{\tau}_{\kappa}\psi^{\tau}_{\kappa} = E_{\kappa}\psi^{\tau}_{\kappa}, \qquad (B12)$$

one finds that the energies are $E_{\kappa} = \pm \hbar v_{\rm F} \sqrt{\kappa_x^2 + \kappa_y^2}$, with the upper (lower) sign corresponding to the conduction (valence) band. The corresponding eigenstates are "spinors" in A/B space:

$$\psi_{\kappa}^{\tau} = \begin{pmatrix} F_A^{\tau}(\kappa) \\ F_B^{\tau}(\kappa) \end{pmatrix} = \frac{1}{\sqrt{2}} \begin{pmatrix} \pm \frac{\tau \kappa_x + \tau \kappa_y}{\sqrt{\kappa_x^2 + \kappa_y^2}} \\ 1 \end{pmatrix}.$$
(B13)

The spinor gives the weights of the sublattice Bloch states (B9). The wave function near *K* is thus

$$\Psi_{\mathbf{K}+\boldsymbol{\kappa}}(\mathbf{r}) = F_A^{\mathbf{K}}(\boldsymbol{\kappa})\Psi_A^{\mathbf{K}+\boldsymbol{\kappa}}(\mathbf{r}) + F_B^{\mathbf{K}}(\boldsymbol{\kappa})\Psi_B^{\mathbf{K}+\boldsymbol{\kappa}}(\mathbf{r}), \qquad (B14)$$

which can be separated into a fast and a slow part:

$$\Psi_{\mathbf{K}+\boldsymbol{\kappa}}(\mathbf{r}) \approx e^{i\boldsymbol{\kappa}\cdot\mathbf{r}}[F_A^{\mathbf{K}}(\boldsymbol{\kappa})\Psi_A^{\mathbf{K}}(\mathbf{r}) + F_B^{\mathbf{K}}(\boldsymbol{\kappa})\Psi_B^{\mathbf{K}}(\mathbf{r})], \quad (B15)$$

where the plane wave part $e^{i\kappa \cdot \mathbf{r}}$ is the so-called envelope function. Of course, a similar relation holds near **K**'.

2. Nanotubes: Graphene on a cylinder

One must take care when transforming from the graphene coordinates to the nanotube cylindrical coordinates. Previously we used the coordinate system defined in Table I, i.e., the horizontal bonds in Fig. 55 are along our *x* direction. When changing to the coordinate system $(k_{\perp}, k_{\parallel})$, the coordinates are rotated as

$$\binom{\kappa_x}{\kappa_y} = \begin{pmatrix} \cos\vartheta & \sin\vartheta \\ -\sin\vartheta & \cos\vartheta \end{pmatrix} \binom{\kappa_\perp}{\kappa_\parallel}, \qquad (B16)$$

where ϑ is the angle between **C** and $\hat{\mathbf{x}}$, or $\vartheta = \pi/6 - \theta$ in terms of the chiral angle θ . In these coordinates, Eq. (B8) becomes

$$H_{\kappa}^{\tau} = \hbar v_F \begin{pmatrix} 0 & e^{i\tau\vartheta} (\tau \kappa_{\perp} - i\kappa_{\parallel}) \\ e^{-i\tau\vartheta} (\tau \kappa_{\perp} + i\kappa_{\parallel}) & 0 \end{pmatrix}. \quad (B17)$$

The coordinate rotation thus creates a phase factor $e^{i\tau\vartheta}$, which can be removed by a unitary transformation. Doing this, we get the nanotube Hamiltonian in cylindrical coordinates:

$$H^{\tau}_{\text{CNT},\boldsymbol{\kappa}} = U_{\vartheta} H U^{\dagger}_{\vartheta} = \hbar v_F (\tau \kappa_{\perp} \sigma_1 + \kappa_{\parallel} \sigma_2), \qquad (B18)$$

where the unitary operator U_{ϑ} is

$$U_{\vartheta} = \begin{pmatrix} 1 & 0\\ 0 & e^{i\tau\vartheta} \end{pmatrix}. \tag{B19}$$

⁴⁰These functions are not orthogonal. The overlap matrix should therefore in principle be included when solving Schrödinger's equation (B12). However, it turns out to be higher order in κ and we neglect it here.

The Hamiltonian (B18) has eigenenergies $E(\kappa_{\parallel}, \kappa_{\perp}) = \pm \sqrt{\kappa_{\parallel}^2 + \kappa_{\perp}^2}$.

When graphene is rolled up to a nanotube (Fig. 1), the wave functions are restricted by periodic boundary conditions $[\Psi_{\mathbf{K}+\boldsymbol{\kappa}}(\mathbf{r}) = \Psi_{\mathbf{K}+\boldsymbol{\kappa}}(\mathbf{r}+\mathbf{C})]$, implying that

$$e^{i(\mathbf{K}+\boldsymbol{\kappa})\cdot\mathbf{C}} = 1. \tag{B20}$$

The condition for a metallic nanotube is that the spectrum of the Hamiltonian (B18) is gapless, i.e., $\kappa_{\perp} = 0$. This happens when $\mathbf{K} \cdot \mathbf{C} = 2\pi M$, where *M* is an integer. From the definitions in Table I, this is equivalent to n - m = 3M (Fig. 5).

3. The curvature-induced gap

Deformation of graphene due to nanotube curvature causes the overlap matrix elements to depend on direction, breaking the C_3 rotational symmetry. Because the effect on the Dirac cones is to shift them in momentum space (Kane and Mele, 1997), graphene remains ungapped when curved. However, in a nominally metallic nanotube, this shift opens a band gap (Fig. 8). To understand these effects, we return to the degenerate subspace $\{\Psi_A^K, \Psi_B^K\}$ and calculate the correction to first order in the curvature perturbation H_{cv} . First, we see the diagonal matrix elements are equal,⁴¹ $\langle \Psi_A^K | H_{cv} | \Psi_A^K \rangle =$ $\langle \Psi_B^K | H_{cv} | \Psi_B^K \rangle$. Next, using the tight-binding wave functions, we find the off-diagonal elements are

$$\langle \Psi_A^{\mathbf{K}} | H_{\mathrm{cv}} | \Psi_B^{\mathbf{K}} \rangle \propto \delta t_1 + \delta t_2 \chi + \delta t_3 \chi^2,$$
 (B21)

where δt_i is the curvature correction to the hopping matrix elements for the three bonds. Since δt is zero for a bond parallel to the nanotube axis and maximal for one perpendicular to it, the leading angular dependence is $\delta t_i \propto \cos^2 \eta_i$, where η_i is the angle between the bond *i* and the chiral vector. Equation (B21) then evaluates to

$$\langle \Psi_A^{\mathbf{K}} | H_{cv} | \Psi_B^{\mathbf{K}} \rangle \propto \sum_{p=-1}^{1} \cos^2(\vartheta + 2p\pi/3) (\chi^*)^p = \frac{3}{4} e^{-i2\vartheta},$$
(B22)

with χ replaced by χ^* in the K' valley. Applying the same transformation that led from Eq. (B11) to Eq. (B8) and changing the phase as explained below Eq. (B11), we obtain the curvature correction in cylindrical coordinates:

$$H_{\text{CNT,cv}} = \hbar v_F \Delta \kappa^{\text{cv}} \begin{pmatrix} 0 & e^{i3\tau\theta} \\ e^{-i3\tau\theta} & 0 \end{pmatrix}$$
$$= \hbar v_F \Delta \kappa^{\text{cv}} [\cos(3\theta)\sigma_1 - \tau\sin(3\theta)\sigma_2]. \quad (B23)$$

When this is added to the Hamiltonian (B18), it is clear that both κ_{\perp} and κ_{\parallel} are shifted. However, the shift in κ_{\parallel} is unimportant and can be absorbed into the longitudinal

4. Spin-orbit coupling

We now include the effect of spin-orbit coupling. Special relativity tells us that an electric field is experienced by a moving electron as a magnetic field. The resulting spin-orbit interaction (SOI) Hamiltonian is given by

$$H_{\rm SOI} = \alpha(\mathbf{E} \times \mathbf{p}) \cdot \mathbf{s},\tag{B24}$$

where **E** is the electric field and α is a constant derived from relativistic quantum mechanics. Each carbon ion contributes to **E**, giving rise to matrix elements between π orbitals and inplane orbitals, which in turn are coupled by curvature. To describe this based on microscopic parameters, one must start from the known atomic spin-orbit coupling of carbon and the sp^2 tight-binding parameters. However, since we do not aim to determine the size of the effect, we use a simpler approach, namely to introduce phenomenological parameters for the SOI-induced coupling between π orbitals in curved graphene. In this approach, the perturbation (B24) gives matrix elements in the Bloch basis (B14).

As explained in Sec. III.F.1, broken symmetry in the nanotube allows the first-order matrix elements to be nonzero. Consider the simplest way to break mirror symmetry, namely a constant radial electric field \mathbf{E}_r . We need to calculate the matrix elements of the spin-orbit Hamiltonian (B24) in the $\{\Psi_A^{\mathbf{K}}(\mathbf{r}), \Psi_B^{\mathbf{K}}(\mathbf{r})\}$ basis. Taking the cross product in Eq. (B24) and projecting to the parallel direction $\hat{\mathbf{T}}$ gives

$$E_r(\hat{\mathbf{z}} \times \mathbf{p}_{AB}^{\mathrm{K}})_{\parallel} = \hbar v_{\mathrm{F}} E_r \hat{\mathbf{T}} \cdot (i\hat{\mathbf{x}} + \hat{\mathbf{y}}) = \hbar v_{\mathrm{F}} E_r e^{i\vartheta}, \quad (B25)$$

where we used the matrix element of **p** from Eq. (B7) and the rotation (B16). This also holds for the K' valley, if $e^{i\vartheta}$ is replaced by $e^{-i\vartheta}$. After the unitary transformation (B19), we obtain the form of the orbital-like spin-orbit interaction in cylindrical coordinates:

$$H_{\rm SOI, orbital-like} = \Delta_{\rm SO}^1 \begin{pmatrix} 0 & 1\\ 1 & 0 \end{pmatrix} \tau s = \Delta_{\rm SO}^1 \sigma_1 \tau s, \qquad (B26)$$

where Δ_{SO}^1 is a phenomenological parameter. As mentioned, this derivation is based only on symmetry and gives no information about the magnitude of Δ_{SO}^1 , except that it is linear in the atomic SOI and inversely proportional to the nanotube diameter. Thus the orbital-like SOI has an easy physical interpretation: it is caused by a Rashba effect, because it is proportional to the azimuthal momentum and the (mean) radial electric field. This is the SOI that was originally derived for graphene (Kane and Mele, 1997) and for nanotubes (Ando, 2000).

It was realized later that one more term is allowed by the reduced symmetry (see Sec. III.F.3). This term comes from the diagonal matrix elements:

$$B_{AA,\parallel}^{\mathbf{K}} = \langle \Psi_A^{\mathbf{K}} | (\mathbf{E} \times \mathbf{p})_{\parallel} | \Psi_A^{\mathbf{K}} \rangle, \qquad (B27)$$

⁴¹This is because the combination of inversion and time-reversal symmetries rules out a σ_z term in the Hamiltonian.

and the identical expression for the *B* sublattice. For a constant electric field (or one obeying the C_3 symmetry) $B_{AA,\parallel}^{\mathbf{K}}$ is zero. When the rotational symmetry is broken, it can be nonzero. It turns out that $B_{AA,\parallel}^{\mathbf{K}}$ depends on the chirality. We study two special cases: armchair and zigzag nanotubes.

Armchair nanotubes have mirror planes perpendicular to the nanotube axis through an *A* atom. Therefore the curvatureinduced electric field has the same symmetry and so does $(\mathbf{E} \times \mathbf{p})_{\parallel} = E_r p_{\perp}$. Using the fact that the wave function transforms under reflection as $\Psi_A^{\mathbf{K}} \to (\Psi_A^{\mathbf{K}})^*$, as evident from Fig. 55, we have for armchair nanotubes

$$B_{AA,\parallel}^{\mathbf{K}} = \langle (\Psi_A^{\mathbf{K}})^* | (\mathbf{E} \times \mathbf{p})_{\parallel} | (\Psi_A^{\mathbf{K}})^* \rangle = -(B_{AA,\parallel}^{\mathbf{K}})^*, \quad (B28)$$

and therefore purely imaginary. On the other hand, since the operator $(\mathbf{E} \times \mathbf{p})_{\parallel}$ is Hermitian, we must also have $B_{AA,\parallel}^{\mathbf{K}} = (B_{AA,\parallel}^{\mathbf{K}})^*$. Hence for an armchair nanotube $B_{AA,\parallel}^{\mathbf{K}} = 0$.

Zigzag nanotubes have no mirror plane through an atom and perpendicular to the axis. Instead, there is a mirror plane along the axis, and since $(\mathbf{E} \times \mathbf{p})_{\parallel}$ changes sign under this symmetry, there is no cancellation as for the armchair case.

We conclude that there is a spin-orbit-interaction contribution diagonal in pseudospin of the form

$$H_{\rm SOI,Zeeman-like} = \Delta_{\rm SO}^0 \tau s, \tag{B29}$$

where Δ_{SO}^0 depends on chirality and therefore has leading harmonic $\Delta_{SO}^0 \propto \cos 3\theta$. This is the Zeeman-like term of Eq. (18), which thus comes from intrasublattice matrix elements, i.e., from next-nearest neighbor couplings.

5. Final form of the Hamiltonian

The final low-energy Hamiltonian is the sum of the metallic nanotube Hamiltonian, the curvature term, and the two spinorbit terms:

$$H_{\kappa} = \hbar v_{\rm F} (\kappa_{\parallel} \sigma_2 + \Delta \kappa_{\perp} \sigma_1) + \Delta_{\rm SO}^0 \tau s + \Delta_{\rm SO}^1 \sigma_1 \tau s, \quad (B30)$$

where both $\Delta \kappa_{\perp}$ and Δ_{SO}^{0} are proportional to $\cos 3\theta$. In addition, a magnetic field gives rise to an Aharonov-Bohm phase, as well as the usual Zeeman term:

$$H_B = \hbar v_{\rm F} \tau \Delta \kappa_{\perp}^B \sigma_1 + \frac{1}{2} g_s \mu_B \mathbf{s} \cdot \mathbf{B}$$
(B31)

with $\Delta \kappa_{\perp}^{B} = eDB_{\parallel}/4\hbar$. The spectrum can be found by simple diagonalization of the Hamiltonian, giving the expression in Eq. (20) for $E_{\tau,s}^{\pm}(B_{\parallel})$ for the case of **B** directed along the nanotube. In that equation the gap is $E_{G}^{0} = -2\hbar v_{\rm F}\Delta\kappa_{\perp}$ and the confinement energy is $E_{\rm conf} = \hbar v_{\rm F}\kappa_{\parallel}$. The convention used here has for the time-reversal operator $\mathcal{T} = i\tau_{x}s_{y}\mathcal{K}$, where \mathcal{K} denotes complex conjugation. Time reversal therefore transforms $s \rightarrow -s$, $\tau \rightarrow -\tau$, $\sigma_{2} \rightarrow -\sigma_{2}$, and $k_{\parallel} \rightarrow -k_{\parallel}^{42}$.

6. Single-particle quantum dot states

Next we discuss the single-particle states in a quantum dot. We assume a parallel magnetic field (making the spin projection s along the axis a good quantum number) and a potential that is flat in the middle of the dot. The wave function is thus a superposition of right-moving and left-moving waves. In principle, both plane wave solutions close to \mathbf{K} and \mathbf{K}' should be included, because the dot terminations can mix the two valleys. However, we approximate the valley index a good quantum number, assuming a smooth confining potential, and later include mixing via a matrix element between the valley-polarized states, as explained in Sec. III.B. The standing wave is then of the form

$$\Psi_{\text{dot},s}(t,c) = (Ce^{i\kappa_{\parallel}t} + De^{-i\kappa_{\parallel}t})u^{\mathbf{K}}(t,c)|s\rangle, \qquad (B32)$$

where $u^{\mathbf{K}}(t, c)$ is the periodic part of the Bloch wave function, and where the relation between *C* and *D* is determined by the reflection coefficients r_{L} and r_{R} at the ends of the dot, because the condition for a bound state is $C = r_{\mathrm{L}}D = r_{\mathrm{L}}r_{\mathrm{R}}C$. Importantly, the envelope wave function depends on valley and spin, because κ_{\parallel} depends on both through the energy

$$\hbar v_{\rm F} \kappa_{\parallel} = \pm \sqrt{(E - s\tau \Delta_{\rm SO}^0 - E_Z)^2 - (s\Delta_{\rm SO}^1 - \tau E_{\rm G}^0/2 + E_B)^2}.$$
(B33)

(Here $E_B = ev_F DB_{\parallel}/4$.) For B = 0, we thus have two Kramers doublets separated by an energy Δ_{SO} , as explained in the main text. We emphasize that because of the spin-orbit induced difference in κ_{\parallel} , the two doublets cannot be written as product states of longitudinal, valley, and spin components.

In the main text, we refer to the single-particle states as $|\nu\tau s\rangle$, where ν is a quantum number that labels the solutions to the above problem. Two-electron states are then built from the two-electron Slater determinants: $(|\nu\tau s\rangle_1 |\nu'\tau' s'\rangle_2 - |\nu\tau s\rangle_2 |\nu'\tau' s'\rangle_1)/\sqrt{2}$ as in Table IV. These states do not take into account the fact that the longitudinal wave functions are easily distorted by Coulomb interactions, resulting in highly correlated two-electron states (cf. Sec. VII).

7. Two-electron states and exchange interaction

In a two-electron quantum dot the question of interaction corrections to the simple single-particle filling arises. We discuss these corrections using first-order perturbation theory. The two-electron matrix element between four states ψ_a, ψ_b, ψ_c , and ψ_d has the form

$$V_{abcd} = \langle a, b | V | c, d \rangle = \int d\mathbf{r} \int d\mathbf{r}' \rho_{ad}(\mathbf{r}) V(\mathbf{r}, \mathbf{r}') \rho_{bc}(\mathbf{r}'),$$
(B34)

where

$$\rho_{ab}(\mathbf{r}) = \langle \psi_a | \mathbf{r} \rangle \langle \mathbf{r} | \psi_b \rangle, \tag{B35}$$

and $V(\mathbf{r}, \mathbf{r}')$ is the Coulomb interaction potential. The state $|a, b\rangle$ is a simple product state $|a, b\rangle = |a\rangle|b\rangle$ and not an antisymmetrized state as in the headers of Table IV.

⁴²Some use a convention where $\sigma_i \rightarrow \tau \sigma_i$, i = 1, 2. This is equivalent to a unitary transformation $H \rightarrow UHU^{\dagger}$, with $U = e^{i\pi(1-\tau)\sigma_3/4}$. This also changes the definition of the time-reversal operator to $T \rightarrow is_2\sigma_3\tau_1\mathcal{K}$, such that time reversal corresponds to a sign change of σ_1, s, τ and k_{\parallel} .

This matrix element between two Bloch states in Eq. (B14) is

$$\rho_{\mathbf{k}_1\mathbf{k}_2}(\mathbf{r}) = \frac{1}{N_A} \sum_{\mathbf{R}_n} [|\varphi_A(\mathbf{r})|^2 + |\varphi_B(\mathbf{r})|^2] e^{-i(\mathbf{k}_1 - \mathbf{k}_2) \cdot \mathbf{R}_n}, \quad (B36)$$

where N_A is a normalization equal to the number of unit cells and where we have neglected contributions from the overlap of neighboring atoms. The Coulomb matrix element obtained by inserting Eq. (B35) into (B34) naturally separates into a short- and a long-range part $V = V^{\text{SR}} + V^{\text{LR}}$. The two contributions are

$$V_{\mathbf{k}_{1}\mathbf{k}_{2}\mathbf{k}_{3}\mathbf{k}_{4}}^{\text{SR}} \approx V_{0}^{\text{SR}} \delta_{\mathbf{k}_{1}+\mathbf{k}_{2}-\mathbf{k}_{3}-\mathbf{k}_{4}}, \tag{B37a}$$

$$V_{\mathbf{k}_{1}\mathbf{k}_{2}\mathbf{k}_{3}\mathbf{k}_{4}}^{\mathrm{LR}} \approx \frac{1}{\mathcal{V}} \int_{\mathcal{V}} d\mathbf{r} \int_{\mathcal{V}} d\mathbf{r}' V(r-r')$$
$$\times e^{i(\mathbf{k}_{1}-\mathbf{k}_{4})\cdot\mathbf{r}+i(\mathbf{k}_{2}-\mathbf{k}_{3})\cdot\mathbf{r}'}, \qquad (B37b)$$

where $V_0^{\text{SR}} = 2U_0/N_A$ and \mathcal{V} is a normalization area of the nanotube surface. For a quantum dot of length *L* and diameter *D*, the short-range part is $V^{\text{SR}} \approx U_0/[60LD/(\text{nm})^2]$, which for parameters L = 300 nm, D = 2 nm, and $U_0 = 10$ eV gives $V_0^{\text{SR}} = 0.3$ meV.

8. Exchange integrals due to long-range Coulomb interaction

Rotational symmetry can be used to show that the long-range part vanishes except when $(\mathbf{k}_1 + \mathbf{k}_2 - \mathbf{k}_3 - \mathbf{k}_4) \cdot \mathbf{C} = 0$, which in terms of valley index means $\tau_1 + \tau_2 = \tau_3 + \tau_4$ (Weiss *et al.*, 2010). This valley selection rule contrasts with that for spin, which is $s_1 = s_4$ and $s_2 = s_3$. Separating both the wave vectors and the coordinates into a transverse part and a longitudinal part, it becomes evident that the rapid oscillations of the Bloch wave functions make it a good approximation to ignore terms off diagonal in valley indices, because

$$V_{\tau_1,\tau_2,\tau_3,\tau_4}^{\mathrm{LR}} \propto \int d\varphi d\varphi' e^{-iM(\tau_1-\tau_4)(\varphi-\varphi)} V(\varphi-\varphi'), \quad (B38)$$

where the condition $\tau_1 + \tau_2 = \tau_3 + \tau_4$ was used and where $M = \mathbf{K} \cdot \mathbf{C}/2\pi$ is an integer. Therefore, it is generally true that for narrow-gap nanotubes, which have $M \neq 0$, the Coulomb matrix element is strongly suppressed.

Turning to the case of two electrons in a single dot, occupying single-particle states $|\nu \tau s\rangle$ in symmetric or antisymmetric combinations as discussed in Sec. IV, the matrix elements between such orbitals therefore obey

$$\langle 1\tau_1 s_1, 2\tau_2 s_2 | V^{\text{LR}} | 3\tau_3 s_3, 4\tau_4 s_4 \rangle \propto \delta_{s_1, s_4} \delta_{s_2, s_3} \delta_{\tau_1, \tau_4} \delta_{\tau_2, \tau_3}.$$
(B39)

This equation can be applied to calculate the long-range interaction energies of the states in Table IV. For the S(0, 2) states, we get four terms. The cross terms vanish, because they cannot have $\tau = \tau'$ and s = s' at the same time. We therefore conclude that there are only diagonal, direct Coulomb interaction terms, i.e., no exchange corrections. Assuming

the charge distribution of both Kramers pairs to be approximately the same, we thus have

$$\langle \psi | V^{\text{LR}} | \psi \rangle = E_{\text{C}}, \text{ where } \psi \in S(0,2).$$
 (B40)

Consider now the symmetric states containing different longitudinal modes S'(0, 2). There are now eight terms in the interaction integral. The nonzero terms have the same valley and spin for the same electron label. If $\tau = \tau'$ and $s \neq s'$, there are two positive cross terms. The same is true if $\tau \neq \tau$ and s = s' [because of the fast oscillations that lead to Eq. (B39)]. We therefore conclude

$$\langle S'(0,2)|V^{LR}|S'(0,2)\rangle = E_{C} + 2\mathcal{C},$$
 (B41)

where

$$\mathcal{C} = \frac{1}{4} \langle 1Ks, 2K's | V^{\text{LR}} | 1K's, 2Ks \rangle \tag{B42}$$

is an exchangelike Coulomb integral.

The situation is similar for the antisymmetric states AS(0, 2), except now the sign is opposite, i.e., there is an energy reduction due to exchange integrals. These energy shifts can be absorbed into the definition of $\Delta_{S,AS}$ and $\Delta_{AS,S'}$, which is what is done in Table IV.

9. Exchange integrals due to short-range Coulomb interaction

For the short-range Coulomb interaction the valley selection rules in Eq. (B39) do not apply. Within the S(0, 2) multiplet we find that states with antiparallel spin are raised in energy, assuming a repulsive short-range interaction U_0 . This is because in such a state, two electrons can occupy the same atomic site, in contrast to the spin-polarized states where spatial antisymmetry forbids this.

The antisymmetric states AS(0, 2) are almost unaltered by the short-range interaction (Secchi and Rontani, 2013), because their longitudinal symmetries differ from those in S(0, 2). This can be intuitively understood by considering the limit $\Delta_{SO} \rightarrow 0$, which allows us to separate each twoelectron wave function into a longitudinal part and a spinvalley part. In this limit, AS states have zero amplitude to occupy the same cross section of nanotube and therefore the same atomic site. As an example, consider the valleypolarized state $K \downarrow, K \uparrow$. For $\Delta_{SO} \rightarrow 0$ it can be rewritten as $(|\nu, \nu'\rangle - |\nu', \nu\rangle)|K, K\rangle(|\downarrow, \uparrow\rangle + |\uparrow, \downarrow\rangle)$, with an asymmetric longitudinal part as expected. In contrast, the analogous state in S(0, 2) is longitudinally symmetric: $K \downarrow, K \uparrow \rightarrow |\nu, \nu\rangle|K, K\rangle(|\downarrow, \uparrow\rangle - |\uparrow, \downarrow\rangle)$.

10. Two-electron states and Pauli blockade

In Sec. IV we discussed Pauli blockade of two-electron states at zero magnetic field. We noted that for some of the blocked states Pauli blockade can be lifted by a dephasing mechanism only, whereas other blocked states require that the spin or valley quantum number within one of the dots be flipped. Here we give some examples.

As an example for dephasing, consider the blocked state $AS(1, 1)K\downarrow, K'\uparrow$. From Table IV we can write out the wave

function by identifying $\nu = 1$ (2) with the lowest longitudinal shell in the right (left) dot. After rearranging some terms this state has the form

$$AS(1,1)K\downarrow, K'\uparrow = |1K\downarrow\rangle_1 |2K'\uparrow\rangle_2 - |2K'\uparrow\rangle_1 |1K\downarrow\rangle_2 + |1K'\uparrow\rangle_1 |2K\downarrow\rangle_2 - |2K\downarrow\rangle_1 |1K'\uparrow\rangle_2.$$
(B43)

We are interested in the time evolution of this state, and model dephasing by assuming an effective Kramers splitting in the left dot that is slightly larger than that in the right dot: $(E_{1K'\uparrow} - E_{1K\downarrow}) = (E_{2K'\uparrow} - E_{2K\downarrow}) + \delta$. Up to a trivial dynamical phase, the state evolves with time *t* to

$$\begin{aligned} |1K\downarrow\rangle_1|2K'\uparrow\rangle_2 - |2K'\uparrow\rangle_1|1K\downarrow\rangle_2 \\ + e^{-2i\delta t}(|1K'\uparrow\rangle_1|2K\downarrow\rangle_2 - |2K\downarrow\rangle_1|1K'\uparrow\rangle_2). \quad (B44) \end{aligned}$$

When $e^{-2i\delta t} = -1$ the blocked state $AS(1,1)K\downarrow, K'\uparrow$ has dephased into the unblocked state $S'(1,1)K\downarrow, K'\uparrow$. This is similar to the rapid dephasing from T_0 to S due to an Overhauser field difference in a conventional double dot.

Next, consider the blocked state $AS(1, 1)K\downarrow, K\downarrow$. We show that a valley flip in either dot will lift Pauli blockade. We can write out the state as

$$AS(1,1)K\downarrow, K\downarrow \propto |1K\downarrow\rangle_1 |2K\downarrow\rangle_2 - |2K\downarrow\rangle_1 |1K\downarrow\rangle_2.$$
(B45)

A valley flip in, e.g., the left dot $(2K \downarrow \rightarrow 2K' \downarrow)$ leads to the state

$$|1K\downarrow\rangle_1|2K'\downarrow\rangle_2 - |2K'\downarrow\rangle_1|1K\downarrow\rangle_2, \tag{B46}$$

which is a superposition of $AS(1,1)K\downarrow, K'\downarrow$ and the unblocked state $S'(1,1)K\downarrow, K'\downarrow$. Alternatively, a spin flip in the left dot results in a superposition of $AS(1,1)K\downarrow, K\uparrow$ and $S'(1,1)K\downarrow, K\uparrow$, whereas a combined spin-valley flip results in a superposition of $AS(1,1)K\downarrow, K'\uparrow$ and $S(1,1)K\downarrow, K'\uparrow$. In all cases, Pauli blockade is circumvented by the admixture of unblocked states (i.e., longitudinal symmetric states). However, because of spin-orbit coupling, a single flip in one of the dots requires emission or absorption of energy, whereas a spin and valley flip may be allowed by energy conservation. Therefore, "spin-valley" blockade can occur not primarily due to spin and valley conservation, but due to energy conservation. Thus energy conservation plays a central role in establishing Pauli blockade, just as in conventional quantum dots (cf. Fig. 27). In nanotubes the relevant energy scale is the spin-orbit energy rather than the orbital level spacing. Therefore, Pauli blockade in nanotubes is generically weaker than a singlet-triplet Pauli blockade in conventional semiconductors.

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