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Spin–orbit coupling and electron spin resonance for interacting electrons in carbon nanotubes

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Abstract

We review the theoretical description of spin–orbit (SO) scattering and electron spin resonance (ESR) in carbon nanotubes. Particular emphasis is laid on the effects of electron–electron interactions. The SO coupling is derived and the resulting ESR spectrum is analysed using both the effective low-energy field theory and numerical studies of finite-size Hubbard chains and two-leg Hubbard ladders. For single-wall tubes, the field theoretical description predicts a double-peak spectrum linked to the existence of spin–charge separation. The numerical analysis basically confirms this picture, but also predicts additional features in finite-size samples.

1. Introduction

Nanotubes constitute a new class of mesoscopic quantum wires characterized by the interplay of strong electron–electron interactions, low dimensionality, disorder and unconventional spin dynamics [1–8]. In a sense, they represent an ideal model for strongly correlated mesoscopic systems, where in fact basically all known effects in mesoscopic physics have been experimentally observed. Two main classes of nanotubes can be distinguished, namely single-wall nanotubes (SWNTs), which consist of just one wrapped-up graphene sheet with radius *R* in the nanometre regime, and multiwall nanotubes, which contain additional inner shells [5]. Here we will focus on the conceptually simplest case of *metallic SWNTs*, where interactions should completely destroy the Fermi liquid picture and imply a so-called Luttinger liquid (LL) state of matter [6, 7]. The LL is the generic low-energy description of metallic 1D (single-channel) quantum wires [9].

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Evidence for the LL behaviour of interacting 1D electrons has been reported for charge transport in SWNTs [2–4]. However, in such materials one also expects to find more dramatic consequences of the breakdown of Fermi liquid theory, most notably the phenomenon of spincharge separation. This many-body effect asserts that electrons brought into a LL effectively break up into a charge and a spin part that travel with different velocities and hence will be spatially separated after some time. A recent proposal to detect evidence for spin-charge separation in SWNTs has been based on spin transport [8]. A different (and perhaps easier to realize) proposal based on electron spin resonance (ESR) is reviewed in this paper, expanding on our short paper [10]. ESR is a valuable experimental tool for probing the intrinsic spin dynamics of many systems. In ESR experiments one applies a static magnetic field and measures the absorption of radiation polarized perpendicular to the field direction. In the absence of SU(2) spin symmetry breaking terms in the Hamiltonian, the absorption intensity is then simply a δ peak at the Zeeman energy [11, 12].

Since spin–orbit (SO) interactions are generally the leading terms breaking the SU(2) invariance, deviations in the ESR intensity from the δ peak, e.g. shifts or broadenings, are directly connected to these couplings. Below we theoretically address the SO interaction and the resulting ESR spectrum for interacting SWNTs, using both a continuum field theory and a Hubbard model description. Within the effective field theory, the single δ peak is split into *two* narrow peaks in SWNTs if spin–charge separation is realized. Otherwise the ESR spectrum would form a broad band with thresholds at the lower and upper edges [13]. This qualitative difference is caused by the fact that the SO interaction in SWNTs does not spoil spin–charge separation to leading order. Experimental observation of the peak splitting could therefore provide evidence for the elusive phenomenon of spin–charge separation [9]. To experimentally check the predictions made below, samples free of magnetic impurities have to be used. Such impurities have probably spoiled previous ESR measurements for nanotubes [5].

The outline of this paper is as follows. In section 2 the LL theory of carbon nanotubes is reviewed in detail and we give an introduction to ESR theory relevant for our purposes. The SO interaction is derived in section 3, followed by a detailed discussion of the lowenergy theory predictions for the ESR spectrum in section 4. An alternative approach is to use numerical methods to compute the ESR spectrum for microscopic lattice fermion models. We shall use both a Hubbard chain and the more realistic two-leg Hubbard ladder formulation of interacting SWNTs [14]. While charge transport does not allow for such a description due to the importance of long-range interactions, it turns out that, for ESR spectra, only shortrange interactions are important. These are correctly captured by Hubbard-type models, and therefore such models are expected to be appropriate for the quantitative description of ESR spectra in SWNTs. We discuss this approach in section 5 and compare the numerical results with the field-theoretical predictions. Finally, in section 6 conclusions and a brief outlook are provided. In most of this paper, we use the conventions $\hbar = c = 1$ to simplify notation.

2. Basics

2.1. Luttinger liquid theory

Starting from a microscopic lattice description of the SWNT, inclusion of the interactions among electrons leads to rather complicated models. In the case of short-ranged interactions, one can study Hubbard-type models and we will do so in section 5. For low-energy phenomena such as ESR, however, only bands close to the Fermi energy matter. The relevant electronic properties of (not too thin) SWNTs are then caused only by the π electrons of the wrapped graphene sheet. On energy scales $|E| < D = \hbar v_F/R \approx 1$ eV around the Fermi energy E_F

(here v_F is the Fermi velocity), the graphene band structure takes a simple form allowing us to develop a powerful field-theoretic framework for SWNTs [6] reviewed below. We mention in passing that it can be explicitly demonstrated that bands sufficiently far away from the Fermi surface will not change the results obtained from field theory; for an explicit discussion, see [9].

Simple tight-binding band structure calculations for graphene reveal that there are only two linearly independent Fermi points ('flavours') with coordinates $\alpha \vec{K}$ in the first Brillouin zone ($\alpha = \pm$), instead of a continuous Fermi surface [1]. For |E| < D, the dispersion relation around the Fermi points is highly linear (two-dimensional light cone). Since the basis of the graphene honeycomb lattice contains two atoms, there are two sublattices $p = \pm$, and hence two degenerate Bloch states

$$\varphi_{p\alpha}(\vec{r}) = (2\pi R)^{-1/2} \exp(-i\alpha \vec{K} \cdot \vec{r}) \tag{1}$$

at each Fermi point $\alpha = \pm$. Here $\vec{r} = (x, y)$ lives on the sublattice *p* under consideration and the correct normalization for nanotubes has been anticipated. The Bloch functions are defined separately on each sublattice such that they vanish on the other. One can then expand the electron operator in terms of these Bloch functions. The resulting effective low-energy theory of graphene is the 2D massless Dirac Hamiltonian, as follows also from standard $\vec{k} \cdot \vec{p}$ theory.

Wrapping the graphene sheet into a cylinder then leads to transverse momentum quantization, and hence to the effectively 1D band structure of a metallic SWNT. Taking the *x* axis along the tube direction and the circumferential variable as $0 < y < 2\pi R$, quantization of transverse motion now allows for a contribution $\propto \exp(imy/R)$ to the wavefunction. However, excitation of angular momentum states other than m = 0 costs a huge amount of energy, of the order of *D*. Assuming that the SWNT is not excessively doped, following our above remarks, in field theory we may then omit all transport bands except m = 0. The theory will then apply on energy scales |E| < D and length scales larger than the graphene lattice spacing $a \approx 0.246$ nm.

Evidently, a SWNT then forms a truly 1D quantum wire with only two spin-degenerate bands intersecting the Fermi energy. To take this into account, the electron operator for spin $\sigma = \pm$ is written as

$$\Psi_{\sigma}(x, y) = \sum_{p\alpha} \varphi_{p\alpha}(x, y) \psi_{p\alpha\sigma}(x), \qquad (2)$$

which introduces 1D fermion operators $\psi_{p\alpha\sigma}(x)$. Neglecting interactions for the moment, the Hamiltonian is

$$H_0 = -v_{\rm F} \sum_{\rho\alpha\sigma} p \int \mathrm{d}x \,\psi^{\dagger}_{\rho\alpha\sigma} \partial_x \psi_{-\rho\alpha\sigma},\tag{3}$$

which is equivalent to a massless 1D Dirac Hamiltonian, where $v_{\rm F} = 8 \times 10^5 \text{ m s}^{-1}$.

Next we discuss interactions mediated by the (possibly externally screened) Coulomb potential $U(\vec{r} - \vec{r}')$. The precise form of this potential will, of course, depend on details of the set-up. In the simplest case, bound electrons and the effects of an insulating substrate are described by a dielectric constant κ , and for the long-range Coulomb interaction

$$U(\vec{r} - \vec{r}') = \frac{e^2/\kappa}{\sqrt{(x - x')^2 + 4R^2 \sin^2[(y - y')/2R] + a_z^2}},$$
(4)

where $a_z \approx a$ denotes the average distance between a $2p_z$ electron and the nucleus, i.e. the 'thickness' of the graphene sheet. Electron–electron interactions are then described by

$$H_{\rm I} = \frac{1}{2} \sum_{\sigma\sigma'} \int d\vec{r} \int d\vec{r}' \, \Psi_{\sigma}^{\dagger}(\vec{r}) \Psi_{\sigma'}^{\dagger}(\vec{r}') U(\vec{r} - \vec{r}') \Psi_{\sigma'}(\vec{r}') \Psi_{\sigma}(\vec{r}) \tag{5}$$

which is brought into a 1D form by inserting equation (2) for the electron field operator, allowing us to employ the large arsenal of theoretical methods available only in 1D [9]. The result is

$$H_{\rm I} = \frac{1}{2} \sum_{pp'\sigma\sigma'} \sum_{\{\alpha_i\}} \int dx \, dx' \, V^{pp'}_{\{\alpha_i\}}(x - x') \psi^{\dagger}_{p\alpha_1\sigma}(x) \psi^{\dagger}_{p'\alpha_2\sigma'}(x') \psi_{p'\alpha_3\sigma'}(x') \psi_{p\alpha_4\sigma}(x), \tag{6}$$

with 1D interaction potentials

$$V_{\{\alpha_i\}}^{pp'}(x-x') = \int dy \, dy' \varphi_{p\alpha_1}^*(\vec{r}) \varphi_{p'\alpha_2}^*(\vec{r}') U(\vec{r}-\vec{r}'+p\vec{d}\delta_{p,-p'}) \varphi_{p'\alpha_3}(\vec{r}') \varphi_{p\alpha_4}(\vec{r}). \tag{7}$$

These potentials only depend on x - x' and on the 1D fermion quantum numbers. For interactions involving different sublattices $p \neq p'$ for \vec{r} and $\vec{r'}$ in equation (5), a sublattice shift vector \vec{d} arises [7]. To simplify the resulting 1D interaction (6), one can exploit momentum conservation. Provided we stay away from the charge neutrality point $E_{\rm F} = 0$, umklapp electron–electron scattering processes can be ignored, and the situation simplifies considerably. We then have only 'forward scattering' processes [7], where $\alpha_1 = \alpha_4$ and $\alpha_2 = \alpha_3$, and 'backscattering' processes, with $\alpha_1 = -\alpha_2 = \alpha_3 = -\alpha_4$.

Next we introduce the potential

$$V_0(x - x') = \int_0^{2\pi R} \frac{\mathrm{d}y}{2\pi R} \int_0^{2\pi R} \frac{\mathrm{d}y'}{2\pi R} U(\vec{r} - \vec{r}').$$
(8)

For the unscreened Coulomb interaction (4), this can be explicitly evaluated [7]. From equations (7) and (1), the forward scattering interaction potential is $V_0(x) + \delta_{p,-p'} \delta V_p(x)$, with

$$\delta V_p(x) = \int_0^{2\pi R} \frac{\mathrm{d}y \,\mathrm{d}y'}{(2\pi R)^2} [U(x + pd_x, y - y' + pd_y) - U(x, y - y')], \qquad (9)$$

which is only present if \vec{r} and $\vec{r'}$ are located on different sublattices. Thereby information about the discrete nature of the graphene network has been kept despite the low-energy continuum approximation. Since $V_0(x)$ treats both sublattices on an equal footing, the resulting forward scattering interaction part couples only the total 1D electron densities:

$$H_{\rm I}^{(0)} = \frac{1}{2} \int \mathrm{d}x \, \mathrm{d}x' \,\rho(x) V_0(x-x')\rho(x'),\tag{10}$$

where $\rho(x) = \sum_{p\alpha\sigma} \psi^{\dagger}_{p\alpha\sigma} \psi_{p\alpha\sigma}$. This part of the electron–electron interaction is the most important one and is responsible for the LL behaviour. Note that it is due to the *long-ranged* tail of the Coulomb interaction. The remaining interactions originate from short-ranged interaction processes, and since these are effectively averaged over the tube circumference, their amplitude is quite small, scaling as 1/R. Such couplings are seen below to cause only exponentially small gaps.

For $|x| \gg a$, the term $\delta V_p(x)$ is extremely small. However, for $|x| \leqslant a$, an additional term beyond equation (10) arises due to the hard core of the Coulomb interaction. At such small length scales, the difference between inter- and intra-sublattice interactions matters and $\delta V_p(0)$ must be computed microscopically, leading to a coupling constant f characterizing the additional forward scattering contribution:

$$H_{\rm I}^{(1)} = -f \int \mathrm{d}x \sum_{p\alpha\alpha'\sigma\sigma'} \psi^{\dagger}_{p\alpha\sigma} \psi^{\dagger}_{-p\alpha'\sigma'} \psi_{-p\alpha'\sigma'} \psi_{p\alpha\sigma}, \qquad (11)$$

where $f/a = \gamma_f e^2/R$ with a dimensionless constant γ_f depending on the tube chirality. An estimate for armchair SWNTs yields $\gamma_f \approx 0.05$, implying that f is very small. A similar

reasoning applies to the backscattering contributions $\alpha_1 = -\alpha_2 = \alpha_3 = -\alpha_4$ in equation (6). Because of a rapidly oscillating phase factor, again the only contribution comes from $|x - x'| \leq a$. Furthermore, only the part of the interaction which does not distinguish between the sublattices is relevant, leading to

$$H_{\rm I}^{(2)} = b \int \mathrm{d}x \sum_{pp'\alpha\sigma\sigma'} \psi^{\dagger}_{p\alpha\sigma} \psi^{\dagger}_{p'-\alpha\sigma'} \psi_{p'\alpha\sigma'} \psi_{p-\alpha\sigma}.$$
(12)

For the unscreened interaction (4), $b/a = \gamma_b e^2/R$ with $\gamma_b \approx \gamma_f$. For externally screened Coulomb interaction, however, one may have $b \gg f$.

Further progress can be made by using the Abelian bosonization approach [9]. For that purpose, one brings the noninteracting Hamiltonian (3) into the conventional form of the 1D Dirac model by switching to right-and left-movers $(r = \pm)$, which are linear combinations of the sublattice states $p = \pm$. In this representation, an Abelian bosonization formula [6, 7, 9] applies with four bosonic phase fields $\theta_a(x)$ and their canonical momenta $\Pi_a(x)$. The four channels are obtained from combining charge and spin degrees of freedom as well as symmetric and antisymmetric linear combinations of the two Fermi points, a = c+, c-, s+, s-. The bosonized expressions for H_0 and $H_1^{(0)}$ are

$$H_{0} = \sum_{a} \frac{v_{\rm F}}{2} \int dx \left[\Pi_{a}^{2} + K_{a}^{-2} (\partial_{x} \theta_{a})^{2} \right]$$
(13)

$$H_{\rm I}^{(0)} = \frac{2}{\pi} \int dx \, dx' \, \partial_x \theta_{\rm c+}(x) V_0(x - x') \partial_{x'} \theta_{\rm c+}(x'). \tag{14}$$

The bosonized form of $H_{\rm I}^{(1,2)}$ [6] leads to nonlinearities in the θ_a fields for $a \neq c+$. Although bosonization of equation (3) gives $K_a = 1$ in equation (13), interactions will renormalize these parameters. In particular, in the long-wavelength limit, $H_{\rm I}^{(0)}$ can be incorporated into H_0 by putting

$$K_{c+} = \{1 + 4\tilde{V}_0(k \simeq 0)/\pi \hbar v_{\rm F}\}^{-1/2} \leqslant 1,$$
(15)

while, for all other channels, the coupling constant f gives rise to a tiny renormalization, $K_{a\neq c+} = 1 + f/\pi \hbar v_F \simeq 1$. The plasmon velocities of the four modes are $v_a = v_F/K_a$ and hence the charged (c+) mode propagates with higher velocity than the three neutral modes. The dimensionless Luttinger parameters measure the correlation strength in the system, with the noninteracting point at $K_a = 1$ and repulsive interactions leading to $K_{c+} < 1$. For the long-ranged interaction (4), the logarithmic singularity in $\tilde{V}_0(k)$ requires the infrared cut-off $k = 2\pi/L$ due to the finite length L of the SWNT, resulting in

$$K_{c+} = \left\{ 1 + \frac{8e^2}{\pi\kappa\hbar v_{\rm F}} \ln(L/2\pi R) \right\}^{-1/2}.$$
 (16)

Since $\hbar c/e^2 \simeq 137$, we estimate $e^2/\hbar v_F = (e^2/\hbar c)(c/v_F) \approx 2.7$, and therefore K_{c+} is typically in the range 0.2–0.3. This estimate does only logarithmically depend on L and R and therefore is expected to be almost independent of the sample under study. The Luttinger parameter (16) can also be written in the form

$$K_{\rm c+} = \left(1 + \frac{2E_{\rm c}}{\Delta}\right)^{-1/2},$$

where E_c is the charging energy and Δ the single-particle level spacing. The small value predicted here implies that a metallic SWNT should be a strongly correlated system displaying pronounced non-Fermi liquid effects.

It is clear from equations (13) and (14) that, for f = b = 0, a SWNT constitutes a realization of the LL. We therefore have to address the effect of the nonlinear terms associated

with the coupling constants f and b. This can be done by means of the renormalization group approach. Together with a solution via Majorana refermionization, this procedure allows for the complete characterization of the non-Fermi-liquid ground state of a clean nanotube [6, 7]. From this analysis, one finds that, for temperatures above the exponentially small energy gap

$$k_{\rm B}T_b = D \exp[-\pi\hbar v_{\rm F}/\sqrt{2}b] \tag{17}$$

induced by backscattering processes, the SWNT is adequately described by the LL model and $H_{\rm I}^{(1,2)}$ can effectively be neglected. A rough order-of-magnitude estimate is $T_b \approx 0.1$ mK. In the remainder, we focus on temperatures well above T_b , where the nonlinearities can be neglected and the Luttinger picture applies.

2.2. Sugawara formulation

In the following discussion of ESR theory, it is mandatory to keep the SU(2) spin symmetry explicit at all stages. To do so it is advantageous to avoid the Abelian bosonization used above, which breaks the spin symmetry by hand, but rather employ the Sugawara formulation [9] which manifestly respects SU(2) spin invariance. This formulation is, in fact, fully equivalent to a Wess–Zumino–Witten theory, even with the flavour degeneracy due to the two Fermi points [9]. Since our main interest is in spin properties, we shall, however, suppress the flavour index for most of what follows but return to the complexities added by it later, see section 5. Instead of four channels, we then have a two-channel LL, with interaction constants for charge, $K_c = (K_{c-}+K_{c+})/2 \simeq (1+K_{c+})/2$, and spin, $K_s = (K_{s-}+K_{s+})/2 \simeq 1$. In Abelian bosonization, in order to impose the correct spin symmetry, one then fixes $K_s = 1$ [9]. This procedure is not necessary in the Sugawara treatment below. Inclusion of magnetic Zeeman fields *B* (orbital effects play no role here) only affects the spin sector and, in general, could renormalize K_s . However, this renormalization is irrelevant to ESR, which probes the finite energy scale $\approx B$ [11]. In addition, the velocities $v_{c/s} = v_F/K_{c/s}$ are used in the following.

The Sugawara formulation uses the charge $(J_{L/R})$ and spin currents $\vec{J}_{R/L}$ for right- and left- $(r = \pm = R/L)$ moving electrons described by chiral 1D fermion operators $\psi_{r=R/L}(x, t)$, where we suppress the additional spin index σ . In terms of these field operators, chiral SU(2) spin current operators are given by

$$J_{R,L}(x) = \frac{1}{2} : \psi_{R/L}^{\dagger}(x) \vec{\sigma} \psi_{R/L}(x) :,$$
(18)

where the colons denote normal ordering and Pauli matrices $\vec{\sigma}$ act in spin space. They obey Kac–Moody commutation relations (μ , $\nu = x, y, z$):

$$[J_{L/R}^{\mu}(x), J_{L/R}^{\nu}(x')] = \pm i\delta'(x - x')\delta^{\mu\nu}/4\pi + i\epsilon^{\mu\nu\lambda}J_{L/R}^{\lambda}(x)\delta(x - x').$$
(19)

Likewise, charge current operators are defined as

$$J_{\rm R,L} = :\psi_{\rm R/L}^{\dagger}(x)\psi_{\rm R/L}(x):, \tag{20}$$

where spin indices are summed over. Using these current operators, the LL in the $U(1) \times SU(2)$ -invariant Sugawara formulation is [9]

$$H_0 = H_c + H_s, \tag{21}$$

with decoupled charge and spin parts. The charge sector is described by the U(1)-invariant Hamiltonian:

$$H_{\rm c} = \frac{\pi v_{\rm c}}{2} \int \mathrm{d}x \, (:J_{\rm R} J_{\rm R} + J_{\rm L} J_{\rm L}: + g_{\rm c}: J_{\rm L} J_{\rm R}:).$$
(22)

The coupling g_c is determined by the Luttinger parameter K_c . Explicit expressions can be found in [9], but they are not required below. In particular, in the noninteracting limit, $g_c = 0$ and $v_c = v_F$. The SU(2)-invariant spin Hamiltonian H_s commuting with H_c is

$$H_{\rm s} = \frac{2\pi v_{\rm s}}{3} \int dx : \vec{J}_{\rm R} \cdot \vec{J}_{\rm R} + \vec{J}_{\rm L} \cdot \vec{J}_{\rm L} : + g_{\rm s} \int dx : \vec{J}_{\rm R} \cdot \vec{J}_{\rm L} :, \qquad (23)$$

where $v_s \approx v_F$ is the spin velocity and $g_s = bv_s$ is the coupling constant for electron–electron backscattering processes. In a dynamical spin-sensitive ESR measurement, one should not simply discard this coupling despite the above thermodynamic argument invoking the smallness of gaps [8]. The LL Hamiltonian (21) completely decouples when expressed in terms of spin and charge currents (or the spin/charge bosons of Abelian bosonization). This remarkable fact leads to the phenomenon of spin–charge separation, and unless the SO interaction couples spin and charge sectors, ESR will only probe the spin sector.

2.3. Electron spin resonance

Adopting the conventional Faraday configuration, the ESR intensity at frequency ω is proportional to the Fourier transform of the transverse spin–spin correlation function [11]:

$$I(\omega) = \int dt \, e^{i\omega t} \langle S^+(t)S^-(0) \rangle, \qquad (24)$$

where the static magnetic field points along the *z* axis, $\vec{S} = \sum_i \vec{S}_i$ is the total spin operator and $S^{\pm} = S^x \pm iS^y$. The Hamiltonian can be written as $H = H_0 + H_Z + H'$, where H_0 represents the SU(2)-invariant nanotube model (21) including electron–electron interactions, $H_Z = -g_e \mu_B \vec{B} \cdot \vec{S}$ is the Zeeman term (below, often $g_e \mu_B = 1$) and H' represents SU(2)spin-symmetry breaking terms, in particular the SO coupling. Inserting a complete set of eigenstates $|a\rangle$ of H in equation (24), the ESR intensity follows as

$$I(\omega) = \frac{1}{Z} \sum_{a,b} e^{-E_b/k_{\rm B}T} \delta(\omega - (E_a - E_b)) |\langle a|S^-|b\rangle|^2.$$
⁽²⁵⁾

In the absence of H', there are only contributions from matrix elements between eigenstates with equal total spin $S_a = S_b$. Then all states with $S_a^z = S_b^z - 1$ will lead to a δ peak at frequency $\omega = B$. For instance, at zero temperature, the application of a magnetic field B, taken to be large enough to overcome a spin gap possibly present at B = 0, leads to a ground state with finite magnetization, $S_0 \neq 0$ and the states with $S_a^z = S_0^z - 1$ again yield the δ peak. This can be made explicit as follows. Since $[H_0 + H_Z, S^-] = BS^-$, one has $(H_0 + H_Z)S^-|0\rangle = S^-(H_0 + H_Z)|0\rangle + BS^-|0\rangle$ and identifying $S^-|0\rangle = |a\rangle$ one gets $E_a = E_0 + B$ and thus $I(\omega) = I_0\delta(\omega - B)$. Any perturbation preserving SU(2) invariance will neither shift nor broaden this peak, even at finite temperature [12]. To get nontrivial ESR spectra, one has to identify the leading perturbation breaking SU(2) invariance, which will cause a finite linewidth and shift of the ESR peak. In metallic systems like SWNTs, one has to consider the SO interaction. This is done in the next section.

3. Spin-orbit coupling

3.1. Microscopic derivation

In our derivation of the SO term, we shall neglect electron–electron interactions. Local electric fields exerted by other electrons on a given electron are typically weak compared to the ionic fields [15] and will generally only weakly renormalize the SO couplings from their

noninteracting values. In a single-particle picture, the SO interaction then appears because an electron moving in the electrostatic potential $\Phi(\vec{r})$ experiences an effective magnetic field $\vec{v} \times \nabla \Phi$ in its rest frame. With $\vec{p} = m\vec{v}$ the SO interaction is, in second-quantized form,

$$H' = -\frac{g_{\rm e}\mu_{\rm B}}{4m} \int d\vec{r} \,\Psi^{\dagger}[(\vec{p} \times \nabla\Phi) \cdot \vec{\sigma}]\Psi.$$
⁽²⁶⁾

This represents the starting point for our discussion of the SO coupling.

For a microscopic lattice description, the electron field operator $\Psi_{\sigma}(\vec{r})$ (whose low-energy expansion is given above in equation (2)) can now be expressed in terms of the electron operators c_i for a honeycomb lattice site i at \vec{r}_i :

$$\Psi_{\sigma}(\vec{r}) = \sum_{i} \chi(\vec{r} - \vec{r}_{i})c_{i\sigma}, \qquad (27)$$

where $\chi(\vec{r} - \vec{r}_i)$ is the corresponding Wannier wavefunction centred at lattice site \vec{r}_i . These localized Wannier orbitals can be chosen as real-valued functions even when hybridization with s-orbitals is important. For the simplest case, $2p_z$ orbital wavefunctions could be used, with *z* perpendicular to the graphene plane:

$$\chi(\vec{r}) = (z/4a_0\sqrt{2\pi})\exp(-r/2a_0)$$

where $a_0 = \hbar^2/6me^2$ is the effective Bohr radius.

Inserting the expansion (27) into (26), the SO interaction is [10, 16, 17]

$$H' = \sum_{\langle jk \rangle} \mathrm{i}c_j^{\dagger}(\vec{u}_{jk} \cdot \vec{\sigma})c_k + \mathrm{h.c.}$$
⁽²⁸⁾

which explicitly breaks SU(2) symmetry. The SO vector $\vec{u}_{jk} = -\vec{u}_{kj}$ has real-valued entries and can be written as

$$\vec{u}_{jk} = \frac{g_e \mu_B}{4m} \int d\vec{r} \, \Phi(\vec{r}) [\nabla \chi(\vec{r} - \vec{r}_j) \times \nabla \chi(\vec{r} - \vec{r}_k)]. \tag{29}$$

The on-site term (j = k) is identically zero, and since the overlap decreases exponentially with $|\vec{r}_j - \vec{r}_k|$, we keep only nearest-neighbour terms in equation (28). Equation (28) also allows us to incorporate electric fields due to impurities or close-by gate electrodes. To connect the SO vector to the experimentally measurable SO relaxation rate, τ_{SO}^{-1} , we estimate this rate using Fermi's golden rule. Assuming a constant \vec{u} , the probability for a transition from the initial state $|i\rangle = |k, \uparrow\rangle$ to the final state $|f\rangle = |k', \downarrow\rangle$ is given by $\Gamma_{i\to f} = 2\pi |\langle f|H'|i\rangle|^2 \delta(E_i - E_f)$ and summing over all final states:

$$\tau_{\rm SO}^{-1} = 8(u_x^2 + u_y^2)/3v_{\rm s}.\tag{30}$$

For SWNTs, SO couplings are generally expected to be small. This is in accordance with our approach since the SO vector (29) vanishes by symmetry for an ideal 2D honeycomb lattice. A finite nearest-neighbour SO coupling can only arise due to the finite curvature, stray fields from nearby gates or due to impurities, which break the high symmetry. Focusing for clarity on the curvature-induced SO coupling, the SO vector for achiral tubes can be seen to depend only on bond direction:

$$\vec{u}_{\vec{r}_i,\vec{r}_i+\vec{\delta}_n} = \vec{u}_n,\tag{31}$$

where the nearest-neighbour bonds δ_n (n = 1, 2, 3) connect the two sublattices of the graphene sheet in real space. For explicit representations of the vectors δ_n , see, for example, [17].

3.2. Low-energy form

From now on we then restrict our analysis to achiral tubes and construct the low-energy field theory Hamiltonian H' describing the SO interaction (28), including the flavour indices for the moment. Using equation (31), the microscopic form is

$$H' = \sum_{\vec{r}} \sum_{n=1}^{5} (ic_{\vec{r},p=+}^{\dagger} \vec{u}_n \cdot \vec{\sigma} c_{\vec{r}+\delta_n,p=-} + \text{h.c.}),$$

2

where \vec{r} runs over all lattice sites of the sublattice p = + and we have made the sublattice dependence of the lattice electron operators explicit (again, spin indices are implicit). The expansion (2) implies

$$c_{\vec{r},p,\sigma} = (2\pi R)^{-1/2} \sum_{\alpha} \mathrm{e}^{-\mathrm{i}\alpha \vec{K} \cdot \vec{r}} \psi_{p\alpha\sigma}(x),$$

which yields, after y integration and Taylor expansion of $\psi_{p=-,\alpha\sigma}(x+\hat{e}_x\cdot\vec{\delta}_n)$, the result

$$H' = \int \mathrm{d}x \, \sum_{\alpha,n} \mathrm{e}^{-\mathrm{i}\alpha\vec{K}\cdot\vec{\delta}_n} \psi_{p=+,\alpha}^{\dagger}(x) (\mathrm{i}\vec{u}_n\cdot\vec{\sigma}) [\psi_{p=-,\alpha}(x) + (\hat{e}_x\cdot\vec{\delta}_n)\partial_x\psi_{p=-,\alpha}(x)] + \mathrm{h.c.}$$
(32)

where we use the unit vector \hat{e}_x along the tube axis. Terms involving higher-order derivatives of the field operators are highly irrelevant and can safely be ignored. In equation (32), we have also neglected oscillatory terms that vanish in doped SWNTs because of axial momentum conservation. These oscillations are governed by the wavevector $2|\mu|/v_F$ corresponding to the doping level μ . In practice, intrinsic doping is unavoidable, but for a commensurate situation ($\mu = 0$), additional terms do arise that are ignored in this section.

Next we define vectors given in terms of the vectors specified in (31):

$$\vec{u}_{1\alpha} = \sum_{n=1}^{3} \exp(-i\alpha \vec{K} \cdot \vec{\delta}_n) \vec{u}_n$$
$$\vec{u}_{2\alpha} = \sum_{n=1}^{3} \exp(-i\alpha \vec{K} \cdot \vec{\delta}_n) \vec{u}_n (\hat{e}_x \cdot \vec{\delta}_n)$$

We then obtain from equation (32) the form $H' = H_1 + H_2$ with

$$H_1 = \int \mathrm{d}x \, \sum_{\alpha} \psi_{p=+,\alpha}^{\dagger}(x) (\mathrm{i}\vec{u}_{1,\alpha} \cdot \vec{\sigma}) \psi_{p=-,\alpha}(x) + \mathrm{h.c.}$$
(33)

and

$$H_2 = \int \mathrm{d}x \, \sum_{\alpha} \psi^{\dagger}_{p=+,\alpha}(x) (\mathrm{i}\vec{u}_{2,\alpha} \cdot \vec{\sigma}) \partial_x \psi_{p=-,\alpha}(x) + \mathrm{h.c.}$$
(34)

It is then apparent that H_1 will give the dominant contribution (relevant in the renormalization group sense, scaling dimension 1), while H_2 is marginal (scaling dimension 2). Since SO interactions are always small, the renormalization group arguments around the Luttinger fixed point with no SO couplings can be safely used here. Note that all contributions neglected here either violate momentum conservation or have scaling dimensions ≥ 3 , i.e. are highly irrelevant.

The final step is to move from the sublattice description to the r = R/L basis using a unitary transformation. Noting that the term involving Re $\vec{u}_{1\alpha}$ involves oscillatory contributions that can be neglected by virtue of momentum conservation, the dominant low-energy SO contribution takes the form

$$H_1 = \int \mathrm{d}x \, \sum_{\alpha} \vec{\lambda}_{\alpha} \cdot (\vec{J}_{\mathrm{L},\alpha} - \vec{J}_{\mathrm{R},\alpha}), \tag{35}$$

with effective SO vectors

$$\vec{\lambda}_{\alpha} = 2 \operatorname{Im} \vec{u}_{1,\alpha} = 2 \operatorname{Im} \sum_{n} e^{-i\alpha \vec{K} \cdot \vec{\delta}_{n}} \vec{u}_{n}.$$
(36)

In the analysis of section 4, for clarity, we shall ignore the flavour index, but it is clear from the above how to add it to the theory. The subleading term H_2 is (ignoring the flavour index and again omitting oscillatory terms)

$$H_2 = \int \mathrm{d}x \, \sum_{r=\mathrm{R/L}} \psi_r^{\dagger} \vec{\lambda}' \cdot \vec{\sigma} \mathrm{i} \partial_x \psi_r + \mathrm{h.c.}$$
(37)

where

$$\vec{\lambda}' = \sum_n \mathrm{e}^{-\mathrm{i}\vec{K}\cdot\vec{\delta}_n} (\hat{e}_x\cdot\vec{\delta}_n)\vec{u}_n.$$

4. ESR spectrum from field theory

In the following, we will keep only the leading contribution (35) and neglect the marginal term H_2 . In the numerical analysis of section 5, the full SO interaction will be considered to assess the accuracy of these approximations. Since SO couplings are typically very small, however, the above perturbative reasoning, suggesting we drop H_2 and the oscillatory terms, is expected to work. Remarkably, the SO interaction then acts exclusively in the spin sector and hence does not spoil spin–charge separation. Therefore the ESR intensity can be computed from the spin sector alone.

For convenience, we define

$$\vec{\lambda}_{r=R/L=\pm} = \vec{B} \pm \vec{\lambda},\tag{38}$$

which represents an effective magnetic field acting separately on the chiral (right/left-moving) spin currents. The presence of two pseudo-fields already hints at the outcome of our calculation below, namely the emergence of a double peak spectrum. Their position will then be given by the absolute values $\lambda_r = |\vec{\lambda}_r|$. The spin Hamiltonian governing the ESR spectrum $I(\omega)$ is then

$$H = H_{\rm s} - \sum_{r} \int \mathrm{d}x \,\vec{\lambda}_r \cdot \vec{J}_r,\tag{39}$$

with H_s given in equation (23). The Heisenberg equation of motion for the chiral spin current operators is

$$(\partial_t \pm v_s/3\partial_x)\vec{J}_r = \mp (g_s/4\pi)\partial_x\vec{J}_{-r} + g_s\vec{J}_R \times \vec{J}_L - \vec{\lambda}_r \times \vec{J}_r.$$
(40)

Taking $\vec{B} = B\hat{e}_z$, and using $\vec{S} = \vec{J}_R + \vec{J}_L$, the ESR spectral density is

$$I(\omega) = \int dt \, dx \, e^{i\omega t} \sum_{rr'} \langle J_r^+(x,t) J_{r'}^-(0,0) \rangle.$$
(41)

We then compute the ESR spectrum using the Sugawara spin Hamiltonian (39). An independent route would proceed via fermionization [10]. For simplicity, we discuss the case $g_s = 0$ (no backscattering), but the method is flexible enough to allow treatment of the general case as well. However, the numerical analysis of section 5 indicates that, indeed, the backscattering interaction does not affect the results in a significant way.

Since the spin Hamiltonian decouples in the chiral spin currents, we just need to add the contributions due to the two chiralities. In equilibrium, using the spin susceptibility $1/4\pi v_s$ of a chiral fermion, we have

$$\langle \vec{J}_{\rm R} \rangle = \frac{1}{4\pi v_{\rm s}} \vec{\lambda}_{\rm R}. \tag{42}$$

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Using the fluctuation-dissipation theorem, we can now express $I(\omega)$ in (41) in terms of the imaginary part of the susceptibility tensor $\chi_r(q, \omega)$:

$$I(\omega) = \sum_{r=\pm} \sum_{\mu,\nu=x,y} \frac{2}{1 - \exp(-\omega/k_{\rm B}T)} \,{\rm Im}\,\chi_r^{\mu\nu}(q\to 0,\omega).$$
(43)

The susceptibility can in turn directly be obtained from the equation of motion (40). Formally defining the matrix $(\Lambda_r)^{\mu\nu} = \epsilon^{\mu\nu\alpha}\lambda_r^{\alpha}$, we find

$$\chi_r(q \to 0, \omega) = \frac{1}{4\pi v_{\rm s}} [i\omega + \Lambda_r]^{-1} \Lambda_r,$$

and straightforward algebra then yields the ESR spectrum:

$$I(\omega) = \sum_{r=\pm} [1 + \lambda_r^z / \lambda_r]^2 \frac{\lambda_r}{4v_s (1 - e^{-\lambda_r / k_B T})} \delta(\omega - \lambda_r).$$
(44)

As expected, the single δ peak splits into two peaks but there is no broadening. The peak separation is $|\lambda_+ - \lambda_-|$, see (38), and the peak heights are generally different. To lowest order in λ/B , the two peaks are located symmetrically around $\omega = B$. Notice that, for \vec{B} perpendicular to the effective SO vector $\vec{\lambda}$, the splitting is zero.

It should be stressed that these results hold both for the noninteracting and the interacting cases. However, for the interacting case realized in SWNTs the double-peak structure is only possible if spin-charge separation is present. Otherwise the charge sector will mix in, leading to broad bands with threshold behaviours [13]. Closer inspection shows that inclusion of the subleading term (37) preserves the splitting into two peaks but the peaks now acquire a finite width $\sim |\vec{\lambda}'|$. Similarly, the effects of backscattering are expected to be small. It is important to point out that the double-peak spectrum would also show up in the noninteracting case, and hence its experimental observation does not represent a true proof for spin-charge separation. However, it would certainly give strong evidence, as in the interacting situation the double peak survives only when spin-charge separation is realized.

In practice, to get measurable intensities, one may have to work with an ensemble of SWNTs. The proposed experiment may be possible using electric-field-aligned SWNTs, or by employing arrays of identical SWNTS. In more conventional samples with many SWNTs, however, the SO vector $\vec{\lambda}$ can take a random direction. For results on the averaged spectrum, see [10].

5. ESR spectra for armchair SWNTs: two-leg Hubbard ladders

In this section we discuss an alternative description of the ESR spectrum via a lattice fermion description. We shall numerically compute ESR spectra for a Hubbard chain and a two-leg Hubbard ladder [14]. These models ignore the long-range part of the Coulomb interactions, which makes them less accurate for the description of charge transport. Since the long-range tails of the interaction do not affect the spin dynamics, however, this does not create a problem for the spin problem at hand.

Modelling the SWNT by a two-leg Hubbard ladder is attractive for the following reasons:

- Backscattering interactions as well as commensurability effects (umklapp scattering) are included.
- The two flavour degrees of freedom are taken care of.
- Band curvature effects are taken into account.
- The marginal SO term (37) as well as all irrelevant contributions can be kept.

- For the Hubbard chain, it is known that any magnetic field spoils spin-charge separation [18]. The importance of this effect for SWNTs can be assessed.
- Accurate numerical techniques are available for such models.

To gain a broader perspective and also to make contact with other 1D conductors, before dealing with the more complex two-leg ladder system, we first consider a 1D Hubbard chain [9] with uniform SO coupling,

5.1. Hubbard chain

The 1D Hubbard model with a uniform SO coupling vector $\hat{\lambda}$ is defined by

$$H = \sum_{n} [c_n^{\dagger}(-t + i\vec{\lambda} \cdot \vec{\sigma})c_{n+1} + h.c.] + U \sum_{n} n_n \uparrow n_n \downarrow - B \sum_{n} S_n^z.$$
(45)

For $\lambda = 0$, equation (45) leads to a LL phase at low energies [9]. It is worth noting that then the exact solution of (45) shows that, with a magnetic field, the charge and spin sectors get mixed [18]. To make progress, we gauge away the SO term by the unitary transformation

$$c_n \to d_n = \begin{cases} c_n & (n \text{ odd}),\\ \exp(i\alpha\hat{\lambda} \cdot \vec{\sigma})c_n & (n \text{ even}), \end{cases}$$

where $\alpha = \arctan(\lambda/t)$, $\hat{\lambda} = \vec{\lambda}/\lambda$ with $\lambda = |\vec{\lambda}|$. The hopping amplitude then changes as $t \to \tilde{t} = t + \lambda \sin \alpha$ and one can easily see that $n_{n\uparrow}n_{n\downarrow} = \tilde{n}_{n\uparrow}\tilde{n}_{n\downarrow}$, where $\tilde{n}_{n\sigma} = d_{n\sigma}^{\dagger}d_{n\sigma}$. Then the Hamiltonian (45) can be rewritten as

$$H = -\tilde{t}\sum_{n} (d_{n}^{\dagger}d_{n+1} + \text{h.c.}) + U\sum_{n} \tilde{n}_{n\uparrow}\tilde{n}_{n\downarrow} - B\sum_{n} \tilde{S}_{n}^{z}$$

and the spin operator as

$$\vec{S}_n = \begin{cases} \frac{1}{2} d_n^{\dagger} \vec{\sigma} d_n & (n \text{ odd}), \\ \frac{1}{2} d_n^{\dagger} (\mathrm{e}^{\mathrm{i}\alpha\hat{\lambda}\cdot\vec{\sigma}} \vec{\sigma} \mathrm{e}^{-\mathrm{i}\alpha\hat{\lambda}\cdot\vec{\sigma}}) d_n & (n \text{ even}). \end{cases}$$

The transformed picture is easier to use in practical calculations. We have calculated the ESR spectrum for (45) using exact diagonalization for small lattices.

The main features emerging from exact diagonalization can be seen in figure 1. For weak interactions (see figure 1(a)) we recover the two-peak structure predicted by field theory, see (44), with the peaks symmetrically arranged around $\omega = B$ but with different peak heights. In addition, no splitting is seen for \vec{B} perpendicular to the SO vector $\vec{\lambda}$. The interaction dependence of the ESR spectrum is shown in figure 1(b). Strong interactions suppress one of the peaks and enhance the other and also give rise to more structure in the spectrum. This behaviour can be traced back to the effects of band curvature. Note that, for increasing U, the distance between both peaks decreases. This feature can be qualitatively explained by a decrease of the spin velocity v_s with U, i.e. for a flatter band the SO splitting is less effective.

5.2. Two-leg Hubbard ladder: armchair SWNTs

We are now ready to generalize the above treatment to armchair SWNTs. Mapping the interacting honeycomb lattice Hamiltonian for armchair SWNTs onto the two-leg Hubbard model [14], one finds

$$H_0 = \sum_{ni} (-tc_{ni}^{\dagger} c_{(n+1)i} - t_{\perp} c_{n1}^{\dagger} c_{n2} + \text{h.c.}) + U \sum_{n,i} n_{n,i,\uparrow} n_{n,i,\downarrow}.$$
 (46)

While under the mapping, $t_{\perp} = t$ up to 1/R corrections [14], we shall allow for $t_{\perp} \neq t$ to have better numerical accuracy. Here we also want to include the SO interaction (28) in this



Figure 1. Exact diagonalization results for the T = 0 ESR spectra of a Hubbard chain with $\lambda = 0.1$, B/t = 0.2, and artificial broadening 0.05t. In (a) ESR spectra for U/t = 1 are shown for different angles γ between $\vec{\lambda}$ and \vec{B} . In (b) the U dependence is shown for $\gamma = 0$. All results are for 12 lattice sites and 4 electrons.

mapping. We use the SO vectors (31), and for simplicity set $\vec{u}_{n=1} = 0$, since this SO vector does not appear in the leading low-energy SO vector $\vec{\lambda}$ in (36) for armchair SWNTs. This simplification also allows for numerically feasible calculations. Writing $\vec{w}_n = (-1)^n \vec{W} + \vec{w}$, where $\vec{w} = (\vec{u}_2 - \vec{u}_3)/2$ and $\vec{W} = (\vec{u}_2 + \vec{u}_3)/2$, the SO term can then be written as

$$H' = \sum_{n} i(c_{n,1}^{\dagger} \vec{w}_n \cdot \vec{\sigma} c_{n+1,1} + c_{n,2}^{\dagger} \vec{w}_{n+1} \cdot \vec{\sigma} c_{n+1,2}) + \text{h.c.},$$
(47)

consisting of a uniform and an alternating contribution:

$$c_{n,1/2}^{\dagger}\vec{w}_{n}\cdot\vec{\sigma}c_{n+1,1/2} = c_{n,1/2}^{\dagger}\vec{w}\cdot\vec{\sigma}c_{n+1,1/2} \pm (-1)^{n}c_{n,1/2}^{\dagger}\vec{W}\cdot\vec{\sigma}c_{n+1,1/2}$$

The resulting model is studied in the remainder of this section for the special case that both \vec{w} and \vec{W} are parallel to the uniform magnetic field. Note that \vec{w} corresponds to the leading SO vector $\vec{\lambda}$ in the low-energy theory, while \vec{W} is related to the subleading vector $\vec{\lambda}'$, see equation (36). Moreover, away from half-filling, following standard reasoning, one may expect that the alternating terms average out. Nevertheless, they will be kept below, but we indeed confirm that in large systems they lead only to small effects. When the above gauge transformation is applied again, we arrive at a two-leg Hubbard ladder (46), but with the on-chain hoppings *t* carrying both a uniform phase $\exp(i\phi_u)$ and an alternating phase $\exp(i\phi_a)$ determined by \vec{w} and \vec{W} , respectively, see figure 2.

Let us first discuss the noninteracting case. For U = 0, the solution of equations (46) and (47) is straightforward. The eigenenergies are given by (*a* is the lattice constant)

$$E_{1\pm}(k_x) = -\sqrt{(2t\cos(k_xa + \sigma\phi_u))^2 \pm 4tt_\perp\cos(k_xa + \sigma\phi_u)\cos(\phi_a) + t_\perp^2}$$
$$E_{2\pm}(k_x) = \sqrt{(2t\cos(k_xa + \phi_u))^2 \pm 4tt_\perp\cos(k_xa + \phi_u)\cos(\phi_a) + t_\perp^2}$$



Figure 2. Sketch of the two-leg Hubbard ladder including SO couplings of the type considered here. The arrows indicate the phase for spins up. Spins down have the opposite phase.



Figure 3. (a) ESR spectrum and (b) dispersion relation of the noninteracting model for $t = t_{\perp} = 1$ and different phase configurations. The dotted (full) curves in (b) correspond to spins down (up).

where $\sigma = \pm 1$ for up or down electrons, respectively. In figure 3 we show the noninteracting T = 0 ESR spectrum in the presence of both the uniform and the alternating phase. The main excitations contributing to each peak are sketched in figure 3(b). The uniform phase case (see the dotted curve) produces a splitting of the principal Zeeman peak. Due to the different parity of the k = 0 and π bands, for the pure uniform case there is no ESR interband transition. When the alternating phase is turned on, the folding of the Brillouin zone produces peaks at



Figure 4. ESR spectra for periodic quarter-filled ladders with a uniform phase $\phi_u = 0.1, t = 1$ and $t_{\perp} = 1.2$.

higher energies. As the parity symmetry is broken, interband transitions are now possible. The opening of the small gap due to the alternating phase is manifested in the splitting seen in peaks A, B and C. The broad structure (F and D) between the highest peaks corresponds to transitions between branches with opposite curvature.

Next we discuss results for U > 0 using the density matrix renormalization group (DMRG) technique [19, 20]. We have used periodic boundary conditions, keeping 256 states and the finite-system algorithm. The ESR spectrum (24) at T = 0 has been calculated using the dynamical DMRG technique [21, 22] for a two-leg ladder with uniform phase ϕ_u only. Additional small alternating phases ϕ_a did not change the spectrum significantly. In figure 4 we present these results for a quarter-filled chain and $t_{\perp} = 1.2$ and t = 1. These parameters are used in order to clearly see the effect of the SO coupling, since for this filling the excitations occur near the linear term of the cosine bands for system sizes that are multiples of 4. The effect of the correlations mainly consists of a small shift of the peaks, but the double-peak spectrum is preserved. Therefore the numerical results lend support to the basic prediction of the analytical low-energy theory and show that the expected double-peak ESR spectrum is stable with respect to the above-mentioned perturbations.

6. Conclusions

We have reviewed the analysis of the ESR spectrum produced by the SO coupling in SWNTs. The effective field theory analysis shows that, at low energy, the SO interaction only acts in the spin sector and the single Zeeman peak, characteristic of a system with SU(2) spin symmetry, splits into two peaks with no broadening. This result relies in an essential way on the property of spin–charge separation characteristic of the LL state realized in the SWNT. Thus the observation of such a splitting would point to this elusive feature.

The field theory analysis has been complemented by the exact numerical calculation of the ESR spectrum for Hubbard models on a single chain and on a two-leg ladder. While these models are not able to describe reliably the charge transport properties of SWNTs due to neglect of the long-range tails of the Coulomb interaction, they are appropriate descriptions of this spin problem. The numerical analysis confirms the field theory predictions, including the validity of the approximations involved in their derivation. In addition, it reveals additional structure in the spectrum due to band curvature and higher energy processes, which are not captured by the field theory approach.

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