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Christian Wagner, Jörg Schuster, and André Schleife Phys. Rev. B **99**, 075140 — Published 20 February 2019 DOI: 10.1103/PhysRevB.99.075140

Strain and screening: Optical properties of a small-diameter carbon nanotube from first principles

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18	(Dated: January 25, 2019)
19	Carbon nanotubes (CNTs) are a one-dimensional material system with intriguing physical prop-
20	erties that lead to emerging applications. While CNTs are unusually strain resistant compared to
21	bulk materials, their optical-absorption spectrum is highly strain dependent. It is an open question,
22	as to what extent this is attributed to strain-dependent (i) electronic single-particle transitions,
23	(ii) dielectric screening, or (iii) atomic geometries including CNT radii. We use cutting-edge the-
24	oretical spectroscopy to explain strain-dependent electronic structure and optical properties of an
25	(8,0) CNT. Quasiparticle effects are taken into account using Hedin's GW approximation and exci-
26	tonic effects are described by solving a Bethe-Salpeter-equation for the optical polarization function.
27	This accurate first-principles approach allows us to identify an influence of strain on screening of
28	the Coulomb electron-electron interaction and to quantify the impact on electronic structure and
29	optical absorption of one-dimensional systems. We interpret our thoroughly converged results using
30	an existing scaling relation and extend the use of this relation to strained CNTs: We show that it
31	captures optical absorption with satisfactory accuracy, as long as screening, quasiparticle gap, and
32	effective electron and hole masses of the strained CNT are known.
33	Keywords: carbon nanotubes, optical properties, excitons, strain, first-principles calculations, density func-
34	tional theory, many-body perturbation theory, screening

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I. INTRODUCTION

Carbon nanotubes (CNTs) possess interesting mate-⁵⁷ 36 rial properties: Their mechanical behavior is dominated ⁵⁸ 37 by high stiffness and large rupture strain,¹⁻⁵ they are ⁵⁹ 38 chemically very stable,^{6,7} and show a sizable shift of ⁶⁰ 39 electronic energy levels as a function of axial strain.^{8–12} ⁶¹ 40 This shift renders *optical* transitions sensitive to strain, ⁶² 41 as has been observed experimentally $^{13-16}$ and explained 63 42 theoretically.¹⁷ For this reason, CNTs are excellent can-⁶⁴ 43 didates for electronic and optical strain sensing and op-⁶⁵ 44 tical strain characterization, which is a promising tech-⁶⁶ 45 nique due to the practical ease of optical readout and ⁶⁷ 46 the higher precision compared to alternative approaches ⁶⁸ 47 such as indirect electronic characterization. In particular, 69 48 optical strain sensors with extremely high, mechanically 70 49 tunable sensitivity can be built in combined CNT/micro-71 50 opto-electro-mechanical systems (MOEMS).^{14,18-20} 72 51

Unfortunately, there is no simple, quantitative picture 73 52 of the explicit strain behavior of optical transitions, since 74 53 their dependence on the single-particle band gap of the 75 54

CNT is not straightforward.¹⁷ This can partly be attributed to strong many-body effects: In low-dimensional systems such as quasi-one dimensional (1D) CNTs, there is less surrounding material than in bulk systems, leading to weak dielectric screening of the electron-electron and electron-hole interaction. As a consequence, quasiparticle (QP) shifts can be as large as 1.2 eV and excitonic effects can be equally strong.^{17,21–25} However, in order to achieve precise strain sensing based on CNTs, a thorough understanding of electronic and optical properties, as well as their strain dependence, needs to be developed. Quantitative insight is essential for the development of MOEMS, such as strain-tunable emitters based on CNTs or tunable optical sensors.

On a more fundamental level, CNTs are a well-suited test bed for obtaining deeper insight into the physics of the strain dependence of screening and, hence, the screened Coulomb electron-electron interaction W. Understanding this is important for modern many-body perturbation theory, since in GW approximation (GW) and Bethe-Salpeter equation (BSE) calculations, W plays a

⁷⁶ crucial role for the renormalization of electronic QP ener-134

⁷⁷ gies and optical transition energies. Large deformations¹³⁵
⁷⁸ are possible in CNTs before rupture, which allows explor-¹³⁶
⁷⁹ ing a much larger strain range than in bulk materials. ¹³⁷

This understanding is also needed since difficulties of-138 80 ten arise during the interpretation of experiments, e.g.¹³⁹ 81 for exciton binding energies: While in (homogeneous)¹⁴⁰ 82 bulk material, a spatial average is a good approximation¹⁴¹ 83 that describes dielectric screening using a dielectric con-142 84 stant ϵ , this cannot a priori be assumed for CNTs. The¹⁴³ 85 spatially resolved dielectric function $\epsilon(\mathbf{r}, \mathbf{r}')$ is needed be-144 86 cause the material response, i.e. screening, is restricted145 87 to the actual electron density of the CNT^{24} and is, thus,¹⁴⁶ 88 strongly direction dependent. In reciprocal space, spa-147 89 tial resolution corresponds to a dependence on q=k-k',¹⁴⁸ 90 which means that $\epsilon(\mathbf{q}=\mathbf{k}-\mathbf{k}')$ must be considered instead¹⁴⁹ 91 of a constant ϵ . Dynamical screening is captured by the 92 frequency-dependent dielectric function $\epsilon(\mathbf{q}, \omega)$, which is 93 required when energy-dependent integrals occur. 150 94

Furthermore, as 1D materials, CNTs show negligible¹⁵¹ 95 optical response perpendicular to the CNT axis, i.e. along 96 the z direction. Hence, screening $\epsilon(q_z = k - k')$ with¹⁵² 97 $\mathbf{q} = q_z \mathbf{e}_z$ and Brillouin zone (BZ) sampling are effectively 98 1D. This needs to be taken into account when using an-99 alytical model functions to describe dielectric screening,154 100 since usually their q-dependence is fitted to three dimen-₁₅₅ 101 sional (3D) semiconductors with a dielectric constant as_{156} 102 low-q limit.²⁶ However, in CNTs the low-q (large dis- $\frac{1}{157}$ 103 tance) limit is vacuum screening.^{24,27} Therefore, the q_{-158} 104 dependence of ϵ and the emerging local-field effects must₁₅₉ 105 be calculated accordingly. Nevertheless, in many stud-160 106 ies only the dielectric constant ϵ is used as a screening₁₆₁ 107 model for the description of excitons in CNTs, since it_{162} 108 is a much simpler quantity.^{22,28,29} This neglect of local₁₆₃ 109 fields for the description of screening and the scaling of_{164} 110 excitons in CNTs with respect to their radius, as pro-165 111 posed by Perebeinos *et al.*,²⁹ is an approximation that $\frac{100}{100}$ 112 requires careful revision. 113 167

In this work we use first-principles electronic-structure₁₆₈ 114 calculations to provide a deeper understanding of these $_{169}$ 115 questions. We use $DFT^{30,31}$ to compute ground-state₁₇₀ 116 geometries and total energies of a small-diameter (8,0)-171 117 CNT in equilibrium and under axial strain. Hedin's GW_{172} 118 approximation³² is used to account for QP effects on elec-173 119 tronic energy levels. Using the G_0W_0 approximation, we₁₇₄ 120 derive strain-induced shifts of valence- and conduction-175 121 band energies and compare to results from a computa-176 122 tionally cheaper hybrid exchange-correlation functional.₁₇₇ 123 Finally, by solving the BSE for the optical polarization₁₇₈ 124 function³³ we account for excitonic effects in optical- $_{179}$ 125 absorption spectra. We study the influence of $Coulomb_{180}$ 126 truncation, a scheme used to mitigate finite-size effects₁₈₁ 127 in supercell calculations for low-dimensional systems, on₁₈₂ 128 resulting optical spectra of the CNT under axial strain. 183 129

These detailed calculations of optical transitions al-184 low us to disentangle the influence of strain on QP ener-185 gies and on excitonic effects. Using our data we explore 186 whether the scaling relation by Perebeinos *et al.*²⁹ for 187 the exciton binding energy in different CNTs also holds for strain in a CNT. Finally, the relation between exciton binding energy, reduced effective mass, and dielectric constant is explored. The resulting strain dependencies of exciton binding energies and optical transitions are essential ingredients for design and layout of MOEMS.

The remainder of this work is structured as follows: Section II summarizes technical aspects of DFT, GW, and BSE calculations. In Sec. III we use these techniques to discuss the strain-dependent electronic structure based on G_0W_0 and hybrid-functional calculations. The solution of the BSE is shown and exciton binding energies are analyzed. We then revisit the scaling relation of Ref. 29 and explore its applicability for the Coulomb-truncated case of a strained CNT. Finally, Sec. IV summarizes and concludes our work.

II. THEORETICAL APPROACH AND COMPUTATIONAL DETAILS

A. Ground-state properties

We use $DFT^{30,31}$ to compute total energies and, via minimization of Hellman-Feynman forces, optimized ground-state geometries of a (8,0)-CNT in equilibrium as well as under axial strain. For these calculations the local-density approximation (LDA) is used to describe exchange and correlation³⁴ and the electron-ion interaction is described using norm-conserving pseudopotentials based on the parametrization by von Barth and Car.³⁵ Wave functions are expanded into a plane-wave basis up to a cutoff energy of 550 eV (40 Ry). To ensure accuracy, we also tested a plane-wave cutoff of 1100 eV (80 Ry), for which total energies are converged up to 9 meV/electron (36 meV/atom). In both cases, the resulting DFT as well as QP gaps agree within 20 meV, which we include in our error bars for QP energy calculations (see below). All DFT calculations are carried out using the Quantum Espresso code.³⁶

We construct a simulation cell that contains the (8,0)-CNT, oriented along the z axis and surrounded by vacuum in the other two directions. We choose a supercell size of $19.5 \times 19.5 \times 4.26$ Å³, such that two periodic images of CNTs are separated by 13.2 Å. This is by far enough vacuum to obtain converged results and to suppress finite-size effects in DFT calculations for the neutral CNT. The geometry optimization is performed using a $1 \times 1 \times 20$ Monkhorst-Pack (MP)³⁷ k-point grid and all atoms are relaxed until the remaining forces are smaller than 0.01 eV/Å. All our results for relaxed atomic geometries can be found in the supplemental material at [URL will be inserted by publisher].

We then compare to calculations within the Vienna *Ab-initio* Simulation Package.^{38,39} For these we use the generalized-gradient approximation by Perdew, Burke, and Ernzerhof⁴⁰ and the projector-augmented wave method.⁴¹ The calculations are carried out using ¹⁸⁸ a plane-wave cutoff energy of $400 \,\mathrm{eV}$ and the same \mathbf{k} -²⁴⁰

189 point grid discussed above. Relaxed atomic geometries

 $_{190}$ $\,$ from both approaches differ only very slightly (GGA in- $_{241}$

¹⁹¹ troduces about 0.2% strain, see supplemental material at ²⁴² [URL will be inserted by publisher]), which is reassuring ²⁴³

¹⁹³ for the comparison of excited-state properties below. 243

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B. *GW* calculations

In order to describe QP effects on electronic single-252 195 particle energies, we use Hedin's GW approximation for₂₅₃ 196 the electronic self energy.³² We use the Yambo package⁴²₂₅₄ 197 to compute QP energies within one step of perturbation₂₅₅ 198 theory, i.e. without updating G or W, which is $known_{256}$ 199 as $G_0 W_0$ approach. The fully frequency-dependent di-200 electric response function $\epsilon(\mathbf{q}, \omega)$, that enters W, is com-₂₅₈ 201 puted within random phase approximation (RPA) $using_{259}$ 202 real-axis integration. Local-field effects play an impor-260 203 tant role and are converged for a G-vector cutoff of $35 \,\mathrm{eV}_{_{261}}$ 204 $(\approx 2.4 \text{ Ry})$, which results in less than 0.5% change of₂₆₂ 205 the dielectric function at several q-points with respect₂₆₃ 206 to the extrapolated value or less than 20 meV change₂₆₄ 207 in the G_0W_0 gap (see Fig. 2 in the supplemental ma-₂₆₅ 208 terial at [URL will be inserted by publisher]). Con-266 209 verged calculations require a $1 \times 1 \times 40$ MP **k**-point grid₂₆₇ 210 $(1 \times 1 \times 60 \text{ MP } \mathbf{k} \text{ points in case of Coulomb truncation}_{,268})$ 211 due to the sharper profile of $\epsilon(q_z)$, see Ref. 21) and at_{269} 212 least 256 bands, which is four times the number of occu- $_{270}$ 213 pied states. Thus, the default parameter set is 256 bands_{271} 214 and 60 k-points for GW calculations, unless other val-272 215 ues are explicitly given. In addition, the singularity of_{273} 216 the Coulomb integral in \mathbf{k} space has to be circumvented, 274 217 which is achieved using the random-integration method₂₇₅ 218 (RIM) described by Marini *et al.*⁴² for GW calculations.₂₇₆ 219

While the vacuum size in our supercell calculations is²⁷⁷ 220 sufficiently large to achieve convergence in DFT, a thor-221 ough unit cell convergence is not feasible for the screened 222 Coulomb interaction W, due to its long-range character.²⁷⁸ 223 The slow decay of the Coulomb interaction with distance 224 renders it impossible to eliminate artificial interactions²⁷⁹ 225 between periodic images.^{43–46} This can be compensated,²⁸⁰ 226 e.g. when calculating defect-formation energies, by sub-227 tracting the electrostatic contribution of all repeating₂₈₁ 228 cells.⁴⁴ For converged calculations of W, Coulomb trun-282 229 cation schemes were developed.^{21,43} the Yambo imple-283 230 mentation of which is used in this work and described₂₈₄ 231 in Ref. 43. Using this scheme renders a lateral unit cell₂₈₅ 232 size of 19.5 Å and a truncation cylinder for the Coulomb₂₈₆ 233 interaction (radius 9.75 Å) sufficient. All details on con-287 234 vergence tests for \mathbf{k} points, number of bands, cell size, 288 235 and Coulomb truncation, including the non-trivial con-289 236 vergence studies with Yambo, are described in detail in₂₉₀ 237 the supplemental material at [URL will be inserted by₂₉₁ 238 publisher]. 292 239

C. Bethe-Salpeter calculations

Excitonic effects are taken into account in the description of optical absorption by solving a BSE for the optical polarization function.³³ For BSE calculations, the screened electron-hole interaction W is computed using the static limit of the response function and the same local-field effects as for GW calculations.^{42,47} Ten valence and ten conduction bands are included for the solution of the BSE. Convergence with respect to **k** points is achieved using a $2 \times 2 \times 80$ MP grid and the same lateral unit cell size of 19.5 Å is found to be sufficient. These convergence tests are discussed in detail in the supplemental material at [URL will be inserted by publisher]. The Yambo package (version 3.4.2) is used for all GW and BSE calculations.

In order to better understand the influence of dielectric screening, we compare Yambo results to BSE calculations from a recent VASP-based implementation.^{48,49} These are carried out using the same parameters as for Yambo: $2 \times 2 \times 80$ MP k points, ten valence and ten conduction bands, as well as the same simulation cell size. Local-field effects are included up to 35 eV G-vector cutoff, as discussed above for GW calculations. This allows calculating exciton binding energies that are converged to within about 1% with respect to the dielectric **G**-vector cutoff. This error estimate stems from the extrapolation of the estimated error of the dielectric function (see Fig. 2) in the supplemental material at [URL will be inserted by publisher]). An accurate extrapolation scheme is used to circumvent the Coulomb singularity.⁴⁹ This BSE implementation currently does not support Coulomb truncation to remove the interaction between super cells, which is discussed in Sec. III E. In order to study the influence of the screened interaction W, we compare the results using a dielectric constant to an analytical model dielectric function⁵⁰ for screening. This comparison allows us to quantitatively discuss the interplay between Coulomb truncation, screening, and strain effects.

III. RESULTS AND DISCUSSION

A. Electronic structure of the unstrained (8,0)-CNT

In Fig. 1 band structures computed using DFT (scissor-shifted to $1.84 \,\mathrm{eV}$) and $G_0 W_0$ are compared. Our $G_0 W_0$ gap of $1.84 \pm 0.02 \,\mathrm{eV}$ ($1.81 \pm 0.02 \,\mathrm{eV}$ when extrapolated to an infinite number of bands and **k**-points, see supplemental information [URL will be inserted by publisher]) agrees very well with early work by Spataru *et al.*, who reported $1.75 \,\mathrm{eV}$,²¹ and later work by Lanzillo *et al.*, who reported $1.81 \,\mathrm{eV}$.⁵¹ The small difference to Spataru *et al.*²¹ can be explained by slightly different computational parameters: They used a plasmon-pole approximation to describe the ω -dependence of the dielectric function and a slightly smaller 16 Å unit cell with



FIG. 1. (Color online.) G_0W_0 band structure (red dashed) is compared to scissor-shifted DFT results (black solid). Besides the scissor shift, the inset shows a linear dependence of QP shifts on KS eigenvalues. The fit shows that the GW valence band is stretched by about 1.15 and the GW conduction band by about 1.06 with respect to corresponding KS bands. The Fermi level of the GW band structure is chosen to be at zero energy.

Coulomb truncation beyond 7 Å cylinder radius. 293

327 The inset of Fig. 1 shows that G_0W_0 QP shifts de-328 294 pend approximately linearly on DFT Kohn-Sham (KS) 295 eigenvalues: In addition to the scissor shift that opens₃₃₀ 296 up the gap, band stretching parameters β describe the₃₃₁ 297 We find that valence band (VB) and_{332} linear slope. 298 conduction band (CB) are stretched by $\beta_{\rm vb}=1.15$ and $_{333}$ 299 $\beta_{\rm cb}=1.06$. This implies a small correction of effective₃₃₄ masses, $m^{\rm GW}=\sqrt{\beta} m^{\rm DFT}$, and needs to be taken into ac-₃₃₅ 300 301 count when solving the BSE with much finer k-point sam-₃₃₆ 302 pling. 303

In order to calculate effective masses of the π bands, ³³⁷₃₃₈ 304 we use a hyperbolic fit that resembles the G_0W_0 bands₃₃₉ as closely as possible.¹² The expression stems from the₃₄₀ 305 306 tight binding (TB) zone folding approach together with₃₄₁ 307 the Dirac cone approximation⁵³ for describing CNT band_{342} 308 structures. The fit to DFT data yields effective masses of_{343} 309 $m_{\rm cb}^{\rm DFT}=0.422\,m_0$ for the conduction and $m_{\rm vb}^{\rm DFT}=0.310\,m_{0_{344}}^{-3.3}$ 310 for the valence band. The effective masses of the respec- $_{345}$ 311 tive $G_0 W_0$ bands are $m_{\rm cb} = 0.418 \, m_0$ and $m_{\rm vb} = 0.278 \, m_{0,346}$ 312 in quantitative agreement with band stretching. 313 347

Hybrid functional for approximate QP energies в. 314

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The QP correction of the DFT gap within the $G_0 W_0$ 315 approach is sizable: The extrapolated shift is $1.21 \pm 0.02_{350}$ 316 eV, compared to a DFT gap of 0.60 eV. This large shift₃₅₁ 317 is attributed to weak dielectric screening in the 1D CNT, 352 318 clearly indicating the need for using a QP correction₃₅₃ 319 scheme. Unfortunately, the G_0W_0 approach is computa-354 320 tionally expensive and becomes unaffordable, e.g. when 355 321 a large number of CNTs or many different strained con-356 322 figurations are studied. For these cases, an approximate³⁵⁷ 323 description of QP corrections is beneficial and using a hy-358 324



FIG. 2. (Color online.) Left: Band gaps of DFT, G_0W_0 , and modified HSE06 calculations (66% HF exact exchange) for the (8,0)-CNT under strain ε . Right: Band stretching parameters $\beta_{\rm CB,VB}$ of $G_0 W_0$ and unmodified HSE06 calculations under strain (256 bands and $1 \times 1 \times 60$ k points). The blue dashed line indicates the DFT-LDA reference.

brid exchange-correlation functional, such as the one by Hevd, Scuseria, and Ernzerhof (HSE06)^{54–56} has proven successful. It comes at much reduced computational cost, since no Coulomb truncation or expensive convergence with respect to empty states is needed. For the (8,0)-CNT, the reduction of cost is about a factor of 6.

The HSE06 functional contains 25% of Hartree-Fock (HF) exact exchange and leads to a band gap of 1.06 eV for the (8,0)-CNT. Increasing the fraction of HF exchange to 66% reproduces the G_0W_0 band gap (see details in the supplemental material at [URL will be inserted by publisher]). Such a large fraction of HF exact exchange is not unusual for low-dimensional systems since screening is much weaker than in bulk materials, revealing almost bare electron-electron interaction. Clearly, using a hybrid exchange-correlation functional *without* adjusting the mixing parameter does not give correct band gaps for CNTs. As an example, the work of Matsuda *et al.* publishes a band gap of about 1.28 eV for the (8.0)-CNT, using the B3LYP functional without adapting the mixing parameter.⁵⁷ Next, we investigate whether the same fixed fraction of HF exchange results in sufficiently precise strain-dependent band gaps for the (8,0)-CNT, compared to $G_0 W_0$ results.

С. Electronic structure of the strained (8,0)-CNT

To investigate the strain dependence of the electronic structure, Fig. 2 shows fundamental gaps computed using DFT, modified HSE06, and G_0W_0 , and band stretching parameters for several relative axial strains up to 6%. This illustrates the strong dependence of the fundamental gap on strain, which is significantly enhanced when QP effects are included, as seen from the different slopes of blue and green curves in the left panel of Fig. 2.

This effect can be understood by invoking strain-

dependent dielectric screening, in addition to strain-359 dependent shifts of KS eigenvalues computed in DFT 360 (blue curve in Fig. 2): The smaller the band gap of the 361 strained CNT, the stronger the dielectric screening, and, 362 thus, the weaker is the electron-electron repulsion. Since 363 QP shifts are small in a material with strong dielectric 364 screening, the $G_0 W_0$ gap of the CNT with the largest 365 axial strain (smallest gap) is closer to the DFT gap than 366 for less strained CNTs. 367

Figure 2 also illustrates that the band gap computed using the modified HSE06 functional with 66 % exact exchange is very similar to the one computed using the G_0W_0 approach for all strains investigated here. The remaining difference is less than 0.1 eV, showing that axial strains up to ≈ 6 % have no influence on the required amount of HF exchange.

We also note that while band stretching β differs be-375 tween CB and VB, it only slightly changes with strain: 376 $\beta_{\rm cb}$ is reduced from 1.06 to 1.00 at 6 % strain and $\beta_{\rm cb}$ re-377 mains at a constant value of 1.15. The strain dependence 378 of QP corrections modifies the strain-dependent effective 379 mass of the CB by less than 3%. The VB is stretched 380 by 15% (7% change of the effective mass), independent 381 of the strain value. Overall, this means that the ratio 382 of DFT and GW corrected effective mass is close to 1.0 383 and, thus, barely strain dependent. However, the abso-384 lute value of the effective mass (either from GW or from 385 DFT) is strongly strain dependent as discussed in Sec. 386 IIIF (see Fig. 6). 387

³⁸⁸ D. Optical properties of the strained (8,0)-CNT

We now discuss the strain dependence of the optical 389 spectrum via strain-induced shifts of the transitions E_{11} , 390 E_{22}, E_{33} , and E'_{11} as depicted in Fig. 3, where the index nn indicates allowed transition from the $n^{\text{th}} \pi$ -VB to 391 392 the $n^{\text{th}} \pi$ -CB.⁵⁸ The unprimed transitions denote first-393 order excitons, whereas the primed transition E'_{11} is a 394 second-order exciton (n=2) that originates from the same 395 electronic bands as E_{11} . This assignment relies on the 396 numerical diagonalization of the exciton Hamiltonian, 397 whose eigenstates are superpositions of non-interacting 398 KS states. We analyzed these contributions for the dif-399 ferent strained cases (see details in the supplemental⁴¹⁴ 400 material at [URL will be inserted by publisher]) and⁴¹⁵ 401 our assignment agrees with Spataru et al. for the un-416 402 strained case.²¹ In the following, results with and with-417 403 out Coulomb truncation are discussed and the origin of₄₁₈ 404 the exciton binding energy is investigated. The visual-419 405 ization of strain-dependent optical transitions in Fig. 4420 406 shows that the first and second optical transition shift⁴²¹ 407 in opposite directions under strain. This is consistent⁴²² 408 with the most simple TB calculation of the CNT elec-423 409 tronic bands with the zone-folding method applied to₄₂₄ 410 (strained) graphene,⁸ which predicts a downshift of CNT₄₂₅ 411 bands with strain for odd transitions (n=1,3,...) and $_{426}$ 412 upshifts of even CNT bands (n=2, 4, ...). 427 413



FIG. 3. (Color online.) Strain-dependent optical spectra of the (8,0)-CNT computed using the BSE approach with Coulomb truncation. The E_{11} and E'_{11} transitions shift to lower energies, whereas E_{22} and E_{33} shift to higher energies. E'_{11} denotes a higher-order exciton (n=2). Black lines are guides to the eye to highlight the shift of transitions.



FIG. 4. (Color online.) Optical transition energies for the strained (8,0)-CNT with (BSE, blue solid) and without (GW, black dashed) excitonic effects, that shift strongly with applied axial strain, covering the visible spectral range.

In contrast to this TB picture, we observe an upshift for the third optical transition that we attribute to σ - π hybridization. Since the curvature of the (8,0)-CNT is large, σ - and π -bands hybridize and the respective band energies are lowered. This effect becomes stronger for higher bands and leads to reordering of the n=3 and n=4states. As a consequence, the third optical transition shifts in the direction opposite to what is predicted by the zone-folding model, which does not include an effect of a curved CNT surface.

For the (8,0)-CNT, the first optical transition E_{11} , which is often observed in photo- or electroluminescence, appears in the infrared and shifts towards lower energies. For the unstrained CNT we observe E_{11} at 1.51 ± 0.03 eV,



FIG. 5. (Color online.) The first three optical transitions computed with Yambo using RIM and homogeneous screening ("Yambo RIM") for the 19.5 Å unit cell. For comparison, the result with Coulomb truncation is included ("Yambo trunc"). $^{479}_{480}$

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which is nearly identical to 1.55 eV reported by Spataru₄₈₃ 428 et al.²¹ We explain the small difference with the slightly $_{434}$ 429 different gaps, the use of RPA instead of plasmon-pole 430 approximation (PPA), and the slightly smaller unit cell. 431 Under strain, the GW+BSE result for the E_{11} transi-432 tion shows a downshift to 1.02 eV at 6% tensile strain. 433 Qualitatively, this trend follows the GW results, but the 434 exciton binding energy $E_{\rm B}$, defined as difference between⁴⁸⁶ 435 GW (dashed black line in Fig. 4) and GW+BSE transi-436 tion (blue line with markers in Fig. 4), significantly re-487 437 duces with strain. We explain this via the strain-induced $_{488}$ 438 increase of the dielectric constant, i.e. screening (see Fig. 489 439 6 and discussion in Sec. IIIF), that leads to a reduction₄₉₀ 440 of the exciton-binding energy. 441 491

The E_{22} and higher transitions are observed₄₉₂ 442 in optical absorption, photoluminescence (PL), $^{15,58,59}_{493}$ Rayleigh scattering, 60,61 and via photocurrents due to₄₉₄ 443 444 absorption.^{62–64} Figure 3 illustrates that E_{22} and $E_{33_{495}}$ 445 each consist of a series of peaks. Their intensity-weighted $\frac{1}{496}$ 446 average, depicted in Fig. 4, shows that E_{22} and E_{33} shift₄₉₇ 447 approximately linearly in energy with strain by a $large_{498}$ 448 value of about 200 meV/%. For larger strained armchair $_{400}$ 449 CNTs, such as (11,0) and (17,0), that show a reduced $\sigma_{-\pi}^{-500}$ π -hybridization, this value is only about 150 meV/%.¹⁷₅₀₁ 450 451 501 Since these CNTs should possess about the same strain₅₀₂ 452 dependence within the TB model with the zone fold- $_{\scriptscriptstyle 503}$ 453 ing scheme, it appears that the σ - π -hybridization itself₅₀₄ 454 is strain dependent. This leads to an enhanced strain- $_{\scriptscriptstyle 505}$ 455 sensitivity of electronic bands and corresponding optical $_{\tt 506}$ 456 transitions of the (8,0)-CNT. 457 507

458 E. Exciton binding energies and long-range 459 Coulomb interaction

In order to understand the physics of screening in
strained CNTs, we compare BSE results with and without the Coulomb truncation scheme used for eliminating

artificial Coulomb interactions of CNTs in adjacent super cells. We use Yambo and the RIM to solve the BSE for the untruncated case and compare to the truncated case in Fig. 5. We also compare to the VASP-BSE implementation, which uses a different solver⁴⁹ and find that both codes agree almost perfectly, as documented in the supplemental material [URL will be inserted by the publisher].

Figure 5 shows that optical transitions appear at lower energies when the Coulomb interaction is truncated, which means that corresponding exciton binding energies are larger. While in the untruncated case the electronhole interaction is (artificially) affected by periodic images over long distances, in the truncated case, no periodic images are present and only the much smaller vacuum screening contributes. Thus, the truncation affects the low- q_z behavior of $\epsilon(q_z)$, which determines the screening of the electron-hole interaction in the longrange limit. The reduction of screening for low q_z due to Coulomb truncation explains the *enhancement* of exciton binding energies. Next, we establish detailed, quantitative insight into the scaling of exciton-binding energies with strain.

F. Scaling of the exciton binding energy with strain

Figure 5 also illustrates that the energies of optical transitions for untruncated and truncated cases depend on strain and approach each other for large strain. The reason is that the band gap is reduced with increasing strain, leading to increased screening that even becomes metallic for about 9% strain. In the metallic case, the truncation has almost no effect on the, then very large, screening.²¹

This effect of strain-dependent screening on exciton binding energy and GW gap does not just occur in 1D materials such as CNTs: The exciton binding energy in bulk ZnO decreases from approximately 70 meV to 55 meV between +2% and -2% strain due to different screening, see Ref. 65. The effect is smaller in bulk, compared to low-dimensional systems, since screening is much stronger in 3D. For various 2D materials with band gaps less than about 2 eV, where screening effects are almost as strong as in CNTs, Zhang *et al.* showed that there is a simple, almost linear dependence of the exciton binding energy on the fundamental band gap.⁶⁶ They also showed that the absolute exciton binding energy is about 50 % of the band gap and reported that it changes as the band gap changes, e.g. due to strain.

Next, we interpret the strain dependence of the exciton-binding energy via a scaling relation: Perebeinos *et al.* derived this for CNTs, using a TB Hamiltonian together with an Ohno potential in order to solve the BSE.²⁹ By introducing a single parameter α , they extended the well-known exciton scaling relation in homo-



FIG. 6. (Color online.) Strain-dependent reduced effective mass, dielectric constant (for the 19.5 Å unit cell), and CNT radius. These parameters enter the scaling relation for the exciton binding energy, Eq. (1).

geneous, isotropic materials,⁶⁷ $E_{\rm B} \sim \mu_{\rm eff} \epsilon^{-2}$, to

$$E_{\rm B} \approx A_{\rm B} \mu^{\alpha - 1} \epsilon^{-\alpha} r_{\rm CNT}^{\alpha - 2},\tag{1}$$

where $A_{\rm B}$ is the exciton-binding energy in a reference 510 state, $r_{\rm CNT}$ is the CNT radius, μ the reduced mass of elec-545 511 tron and hole, and ϵ is the dielectric constant. Perebeinos⁵⁴⁶ 512 et al. found a value of $\alpha = 1.40$ for $\epsilon > 4$ for CNTs. An in-547 513 dependent confirmation of the parameter is given by Ped-548 ersen, who predicted a scaling of $E_{\rm B} \sim r_{\rm CNT}^{-0.6}$ using a vari-514 515 ational approach for wave functions on a cylinder surface 516 and homogeneous, background dielectric screening.^{22,28550} 517 This result corresponds to the same value of α =1.4 and ⁵⁵¹ $\sim r_{\rm CNT}^{\alpha-2}$. While the above relations were developed for a background dielectric screening, we now show that this screening (i.e. no Coulomb truncation) and local fields ⁵⁵⁵ (i.e. with Coulomb truncation) are related 518 519 520 521 (i.e. with Coulomb truncation) are related. 522

To analyze the validity of this scaling relation for $\rm CNTs^{556}$ 523 under strain, we depict our first-principles results for the 524 three materials parameters that enter Eq. (1) in Fig. 6. 525 The dielectric constant is obtained from RPA calcula-⁵⁵⁹ tions using Yambo and the reduced effective mass results 526 527 from our $G_0 W_0$ data. This figure shows that the CNT⁵ 528 radius depends only weakly on strain; the Poisson ratio₅₆₂ 529 of about 0.2 leads to a shift in the exciton binding energy₅₆₃ 530 of about 0.7 % at 6 % tensile strain. Conversely, the elec-564 531 tronic structure is much more sensitive, leading to signifi-565 532 cant changes of effective masses and, via the fundamental $_{\rm 566}$ 533 gap, of the dielectric constant.^{8,9,11,12} The two parame-567 534 ters μ and ϵ , thus, determine the influence of strain on 568 535 the exciton binding energy via Eq. (1). 536 569

In order to compare this to our BSE results, we depict⁵⁷⁰ 537 the strain-dependent exciton-binding energy of the $E_{11^{571}}$ 538 transition in Fig. 7. This data is computed using the572 539 strain-dependent dielectric function $\epsilon(q_z)$ for screening⁵⁷³ 540 of the electron-hole interaction and we compare results⁵⁷⁴ 541 based on Coulomb truncation (see Fig. 8) to those com-575 542 puted without the truncation scheme. As expected, the 576 543 resulting exciton binding energies differ in magnitude,577 544



FIG. 7. (Color online.) Scaling of the strain-dependent exciton binding energy $E_{\rm B}$ of the E_{11} transition with ("Yambo trunc") and without Coulomb truncation ("Yambo RIM"). Symbols represent BSE results and dashed lines represent the scaling relation, Eq. (1), with different values of α . The value of 1.40 given by Perebeinos *et al.*²⁹ is compared to a fit to BSE data. Inset: Data for the Coulomb-truncated case as a function of the scaling parameter $\mu^{\alpha-1} \epsilon^{-\alpha} r^{\alpha-2}$.

since the underlying screening models deviate between truncated and non-truncated case, especially for low q_z (see supplemental material [URL will be inserted by publisher]).

More importantly, Fig. 7 illustrates that the scaling relation, Eq. (1), holds: Fitting to results without Coulomb truncation yields a value of $\alpha \approx 1.29 \pm 0.03$ and shows almost perfect agreement with our data, despite the fact that BSE calculations take local-field effects into account, whereas Eq. (1) was derived under the assumption of a constant, homogeneous dielectric screening. Since there may be a significant influence from strong σ - π hybridization due to CNT curvature, it is not surprising that the value of α slightly differs from 1.40 given by Perebeinos.²⁹ We note that the data in Fig. 7 was computed using Yambo; VASP data is shown in the supplemental material at [URL will be inserted by publisher].

Fitting to data with Coulomb truncation, yields a slightly different value of $\alpha \approx 1.21 \pm 0.03$, since local-field effects with Coulomb truncation are not captured by the static, homogeneous screening entering the TB models used by Pedersen^{22,28} or Perebeinos.²⁹ While this difference in α is, thus, not a surprise, it is remarkable that the scaling relation also holds in the Coulomb-truncated case, and we explore this in more detail in the next section. We point out that for this fit, we used the dielectric constant from the untruncated case (see Fig. 6) to mimic background screening, since Coulomb truncation would imply $\epsilon = 1.0$. This is also addressed in the next section, where we introduce a geometry-dependent parameter C_1 to substitute ϵ in the scaling relation. It characterizes the inhomogeneity and describes screening for confined carriers in the truncated geometry.



FIG. 8. (Color online.) In (a) the real part of the dielectrics⁹⁹ loss function $\epsilon^{-1}(q)$ is shown versus strain. Squares represent₆₀₀ first-principles data with Coulomb truncation and lines rep-₆₀₁ resent the model fit using Eq. (2). The figures (b) and (c)₆₀₂ show the fit coefficients C_1 and C'_2 as a function of ϵ and $E_{11,_{603}}$ respectively.

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578 G. Inhomogeneous dielectric screening and scaling 606 579 relation 607

For a single CNT, as a localized, spatially inhomo-⁶⁰⁹ geneous system, the wave-vector dependence of $\epsilon(q_z)$ is crucial when describing screening.^{21,24,45,68} In order to⁶¹¹ incorporate this into the scaling relation, we use the analytic expression for the dielectric function of an infinite⁶¹³ 1D cylinder, derived by Deslippe *et al.*²⁴ using the Penn⁶¹⁴ model:⁶¹⁹

$$\begin{aligned} \epsilon_{1\mathrm{D}}^{-1}(q_z) &= 1 + \chi(q_z) \, v_{\mathrm{trunc}}(q_z) & {}^{617} \\ &\approx 1 - C_2 \frac{R}{E_{11}} \frac{C_1 q_z^2}{1 + C_1 q_z^2} \left[2 \, I_0(q_z R) \, K_0(q_z R) \right] & {}^{618} \\ & {}^{619} \\ & {}^{620} \end{aligned}$$

$$= 1 - C_2' R \frac{C_1 q_z^2}{1 + C_1 q_z^2} \left[2 I_0(qR) K_0(q_z R) \right] \qquad (2)_{622}^{621}$$

Here, C_1 , C_2 , and $C'_2 = C_2/E_{11}$ are constants and R is₆₂₄ 580 the CNT radius. I_0 and K_0 are modified Bessel func-625 581 tions of the first and second kind, respectively. We fit₆₂₆ 582 this expression to our first-principles data for $\epsilon(q_z)$ in 627 583 Fig. 8 and observe very good agreement. This means₆₂₈ 584 that the model of the 1D cylinder mimics screening in a₆₂₉ 585 CNT, once the influence of the supercell is removed via630 586 Coulomb truncation. 587 631

As shown in Fig. 8, the resulting fit parameter $C_1 \lim_{\epsilon_{32}} \epsilon_{33}$ early depends on the strain-dependent dielectric constant₆₃₃ ϵ ; C'_2 is strain independent. Therefore, C_1 carries the₆₃₄ strain dependence of the screening function that was de-₆₃₅ scribed by ϵ before Coulomb truncation was applied. The₆₃₆ relation between C_1 and ϵ is almost linear, which explains₆₃₇ why the scaling relation, Eq. (1), also holds in the case of Coulomb truncation. We can, therefore, rewrite Eq. (1) using C_1 instead of ϵ :

$$E_{\rm B} = A_{\rm B} r_{\rm CNT}^{\alpha-2} \mu^{\alpha-1} C_1^{-\alpha} \tag{3}$$

Hybrid DFT calculations can then yield effective masses and, after adjusting the fraction of exact exchange, strain-dependent corrected gaps. In combination with the RPA, these calculations also yield the inhomogeneous screening as a function of strain and, thus, the parameter C_1 . This shows that for an isolated, strained CNT, the exciton-binding energy can be related to that of the unstrained state by means of a scaling relation, Eq. (3).

IV. CONCLUSIONS

We use first-principles electronic-structure calculations, based on the GW+BSE approach, to compute strong, strain-related shifts of peaks E_{nn} in the opticalabsorption spectrum of an (8,0)-CNT, consistent with earlier literature. We find that the exciton binding energy in strained CNTs is a function of the band gap and our work leads to the important conclusion that this arises directly from strain-dependent inhomogeneous dielectric screening. This shows that deformation potentials of electronic eigenvalues and exciton binding energies need to be considered explicitly, in order to predict strain-dependent optical spectra of CNTs.

While this implies that the effect of many-body physics on optical spectra in strained CNTs is crucial, we then show that a more simple scaling relation for the exciton binding energy is applicable also to strained CNTs. This scaling relation allows us to extrapolate the shift of optical transitions from the unstrained state to the strained state, based on the strain-induced shift of electronic energy levels and the strain dependence of $\epsilon(q)$ and μ . We then showed that the modified HSE06 hybrid functional, with a fraction of 66 % exact exchange, mimics QP corrections for the unstrained CNT quite well, allowing us to avoid expensive *GW* calculations of strained CNTs to determine these parameters.

Finally, we provide detailed understanding of why the scaling relation works for strained CNTs, even though it relies on the dielectric constant as a parameter and neglects the influence of local-field effects. To this end, we demonstrate that in low-dimensional materials, a wavevector dependent screening function $\epsilon(q_z)$ must be used. In addition, in first-principles excited-state calculations the Coulomb interaction must be truncated in order to obtain supercell convergence, which influences the longrange, low- q_z part of the screening function. We show that a suitable screening function $\epsilon(q_z)$ for CNTs can be obtained from a 1D Penn model of a charge on an infinitely long, hollow cylinder and connect the parameters of this model to our first-principles data, leading to an excellent fit. We envision that this significantly advances the study of optical transitions in strained CNTs and

enables broader applications of this interesting material⁶⁴³
 system.

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ACKNOWLEDGMENTS

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- Part of this work was supported by the National Sci ence Foundation under Grant No. DMR-1555153. Sup-
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port by the Deutsche Forschungsgemeinschaft DFG research unit 1713 and funding of the German academic exchange service (DAAD) is gratefully acknowledged. We thank the Yambo developers, who gave important feedback concerning technical issues during some calculations.

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