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# Interplay between Electrostatic Properties of Molecular Adducts and Their Positions at Carbon Nanotubes

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dipole moments and the local charges induced by various molecular adducts on the energy and localization properties of the optically active defect-based exciton in a (10,5) SWCNT. The interplay of two effects plays a role in the localization of the exciton and, therefore, its red shift, but at different scales: The source of the leading order contribution is the defect conformation, resulting in the red shift of the defect-associated exciton with respect to the  $E_{11}$  band of the pristine SWCNT on the order of ~100 meV, while the individual dipoles and polarization properties of molecular



adducts lead to significantly smaller red shifts on the order of ~10 meV. While the species-dependent trends in defect-induced charges do not directly correlate to the exciton red shift, the charge at the sp<sup>3</sup>-defect exhibits a distinct behavior between *ortho-* and *para*-defect configurations and is relevant to the chemical reactivity of the defect position depending on the adduct type. Overall, our computational results may be helpful for diverse synthetic strategies to fine-tune emission of SWCNTs toward desired applications.

# 1. INTRODUCTION

Single-walled carbon nanotubes (SWCNTs) exhibit electronic and optical properties intrinsic to their structural features, including their diameter and chiral angles, which make them promising materials for applications in electronics, sensors, and photovoltaics.<sup>1-8</sup> In addition, the photophysical properties of SWCNTs can be modified by covalently binding small organic molecules to the SWCNT surface. Such functionalization changes the sp<sup>2</sup>-hybridized carbon atom of the nanotube lattice to sp<sup>3</sup>-hybridized by bonding with a pair of functional groups to a single carbon ring of the SWCNT.<sup>9,10</sup> Formation of the sp<sup>3</sup>-defect results in new optically active transitions,  $E_{11}^*$ , which are red-shifted from the main excitonic band, E<sub>11</sub>.<sup>3,11</sup> In contrast to the lowest energy transitions of the pristine SWCNTs, the lowest energy defect-associated transitions are optically active, enhancing photoluminescence with quantum yields up to 30%.<sup>12-14</sup> The precise energy of the defectassociated transitions for a given chirality of a SWCNT has been found to be dependent on the topological characteristics of the defect on the nanotube surface.<sup>9,13</sup> The inducting ability of the molecular adduct has been also shown to impact the emission energy.<sup>15</sup> These dependences provide chemical tools for fine-tuning emission features targeting a specific application. For instance, aryl-functionalized SWCNTs have demonstrated unique room-temperature single-photon emission extending to telecom wavelengths suitable for quantum communication applications.<sup>16-1</sup>

While tuning of emission energies and their quantum yields is made possible through functionalization of SWCNTs with different chemical adducts,<sup>13,20-24</sup> the chemical effects responsible for the red shift and their dependence on the type of the functional group are not completely understood. The magnitude of red shifts has been attributed to inductive effects in the functional group, with an increasing red shift of  $E_{11}^*$  with a stronger electron-withdrawing group.<sup>15,25</sup> However, calculations based on Density Functional theory (DFT) demonstrate that one or two halide substitutions in the aryl adduct has negligible effect on the lowest transition  $E_{11}^*$ , despite increased dipole moment of the defect group.<sup>3,10,12</sup> On the other hand, it was shown that the dipole moment of the molecular adduct modifies the spatially dependent dielectric function of the functionalized SWCNT, which should affect the lowest energy exciton.<sup>10,26</sup>

Here, we have performed an extensive computational study aiming to clarify how the difference in the dipole moments of functional groups affects the defect-associated optical transition

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in SWCNTs. Applying time-dependent DFT (TD-DFT), we have found that the localization of exciton density near and on the chemical defect is greatly affected by two identical highly polar species bound to the same carbon ring at the surface of a (10,5) SWCNT, which results in the largest red shift of the  $E_{11}^*$  exciton. In contrast, the defect comprising either polar or nonpolar species and a hydrogen atom exhibits much less exciton density at the molecular adduct, leading to smaller red shifts of  $E_{11}^*$  transitions with respect to the parent  $E_{11}$  band. Nonetheless, neither the electrostatic dipole of the system nor the local charge induced by a molecular adduct at the nanotube atom is directly correlated with the red shift of the defectassociated exciton E<sub>11</sub>\* across all considered functional groups. Instead, the defect position at the nanotube surface plays the leading role in the localization and red shift of the trapped exciton.

# 2. METHODOLOGY AND COMPUTATIONAL DETAILS

**2.1. Description of Considered Structures.** A SWCNT of (10,5) chirality and ~10 nm (3 primitive unit cells) in length was prepared using the Visual Molecular Dynamics (VMD) package.<sup>27</sup> Two dangling bonds on each end of the SWCNT were passivated with methylene groups, and the remaining dangling bonds were passivated with hydrogen atoms. This scheme has been shown to eliminate finite size effects and recover the intrinsic electronic structure of the infinite length system.<sup>9,28</sup> A functional group, R<sub>1</sub>, is chosen from a class of alkyls, aryls, or non-carbon linkers (Figure 1a).



**Figure 1.** Structure of (10,5) SWCNT of 10.278 nm in length (3 primitive unit cells) with two functional groups  $R_1$  and  $R_2$  covalently attached to the same carbon ring of the SWCNT (a). The functional group  $R_2$  can be placed in six unique positions with respect to  $R_1$  (red circle): Three *ortho* (dashed circles) and three *para* (solid circles) (b). The functional groups are chosen from three classes of radicals: fully fluorinated and non-fluorinated alkyls with chain lengths  $n \in \{1, 2, 6\}$  (Class I), phenyl, *p*-bromophenyl, and pentafluorophenyl (Class II), and fluorine and difluorophosphine (Class III) (c). The second functional group,  $R_2$ , is either the same as the first group (2 $R_1$  structures) or hydrogen ( $R_1$ /H structures).

For alkyl groups with various chain lengths and dipole moments, we consider  $C_nH_{2n+1}$  and  $C_nF_{2n+1}$  (Class I), where n = 1, 2, or 6. Aryl groups are represented by  $C_6H_5$ ,  $C_6H_4Br$ , and  $C_6F_5$  (Class II). For the non-carbon groups, we explore F and PF<sub>2</sub> groups (Class III).

Placing a single molecular adduct on the surface of the nanotube creates an open-shell system that is highly reactive.

As such, a second species,  $R_2$ , (an auxiliary group) is added in the vicinity of the first addition, so that both R<sub>1</sub> and R<sub>2</sub> adducts form a single sp<sup>3</sup>-defect at the tube surface.<sup>3,29</sup> Two schemes with different auxiliary groups are studied here: one where both substituents are the same (called " $R_1/R_1$ " or " $2R_1$ ") and another with a hydrogen atom as the auxiliary group (called " $R_1/H$ "). In previous studies, it has been shown that six nearest neighbor positions (ortho and para) are available to place this auxiliary group to produce emissive (bright) defect-associated transitions.<sup>3,9,13,29</sup> This allows for three distinct ortho- and three distinct para-functionalization schemes. We utilize the notation previously established for defect positions, as illustrated in Figure 1b, where each ortho and para position is denoted by their approximate angle between  $R_1$ ,  $R_2$ , and the nanotube axis.<sup>30</sup> Note, the exact values of these angles for the (10,5) SWCNT are 19.1°, 79.1°, and -40.9°, which are rounded to  $20^{\circ}$ ,  $80^{\circ}$ , and  $-40^{\circ}$ , respectively. Due to the absence of a resonance structure with charge in the meta position, placing an auxiliary group there generates a synthetically nonrelevant structure and is therefore not considered in this study. In light of recent work investigating densely packed, interacting defects on the SWCNT surface, we emphasize that we are working in the noninteracting limit (i.e., sparsely functionalized at low concentrations of reagent), where the nearest defect may be more than  $\sim 100$  nm away, on average, much larger than the native exciton size of  $\sim 10$ nm.<sup>31,32</sup>

2.2. Ground and Excited State Calculations. All calculations were performed using the Gaussian 16 rev. B.01 package.<sup>33</sup> Geometries were preoptimized using the semi-empirical Austin Model 1 (AM1).<sup>34</sup> DFT calculations were then performed using the long-range corrected hybrid functional CAM-B3LYP.<sup>35</sup> This functional has been shown to properly capture the localization of the electron density and the exciton to the region of the defect.<sup>10</sup> Basis set dependence on these systems was explored as an initial step: we examine three unique combinations of basis sets to yield computationally inexpensive, yet accurate, results.<sup>36-42</sup> One of three schemes was used for the basis set of the defect atoms: (i) all atoms calculated under STO-3G, (ii) only the halides calculated with 6-31G\* with all other atoms calculated using STO-3G, and (iii) all atoms in the functional group plus tube carbon atoms adjacent to the defect calculated with 6-31G\* with all other atoms calculated under STO-3G. For functional groups containing only carbon and hydrogen atoms (aryl and alkyl groups), an increase in the basis set from STO-3G to 6-31G\* has a negligible effect on the electronic structure of the functionalized SWCNT (Figure 2a).

However, this is not the case for fluorinated alkyls. Using 6-31G\* for fluorine atoms instead of STO-3G results in the larger difference (by ~50 meV) in the HOMO-LUMO gap between the functionalized species and the pristine nanotube ( $\Delta E_{\rm HL}$ ), compared to those calculated by STO-3G only, which is further increased when 6-31G\* is used for the entire functional group according to the scheme (iii) (Figure 2a). Furthermore, the values of  $\Delta E_{\rm HL}$  calculated utilizing different basis sets are somewhat constant, regardless of the functional group or its position at the tube surface. Therefore, the energy gaps calculated by the reduced STO-3G basis set can be corrected as a constant energy shift, as has been described in our previous studies.<sup>18</sup> However, due to its substantially decreased HOMO-LUMO gaps for functionalization with



**Figure 2.** Difference in the HOMO-LUMO gap ( $\Delta E_{\rm HI}$ ) between the pristine and functionalized (10.5) SWCNT as a function of the R<sub>2</sub> adduct position on the nanotube surface. (a) Comparison of  $\Delta E_{\rm HL}$  of structures with two fluorinated alkyl groups attached to the same carbon ring of the nanotube,  $2C_6F_{13}$  (black), with those of nonfluorinated alkyls, 2C<sub>6</sub>H<sub>13</sub> (red), calculated by STO-3G (filled stars), and STO-3G/6-31G\*, where the 6-31G\* basis is used either for F atoms (slashed stars) or for the whole functional group, including the two carbons of the tube at the defect sites (empty stars). Empty and filled red stars overlap for the non-fluorinated alkyls pointing to the negligible effect of the basis set. This is not the case for fluorinated alkyls. (b)  $\Delta E_{\rm HL}$  comparison between structures with one alkyl and hydrogen, R<sub>1</sub>/H (filled symbols and solid lines), and two identical alkyl groups,  $2R_1 = R_1/R_1$  (empty symbols and dashed lines). The different number of carbons in alkyls (red) and fluorinated alkyls (black) corresponds to different symbols shown in the legend of the (c) panel. (c) Comparison of  $\Delta E_{HL}$  for the fluorinated alkyls of various lengths and (d) for non-carbon groups of fluorine (empty and filled black diamonds) and difluorophosphine (empty and filled cyan triangles). In all cases, the O<sub>20</sub> defect position results in the smallest  $\Delta E_{\rm HL}$ , whereas the P<sub>20</sub> position leads to the largest  $\Delta E_{\rm HL}$ .

strongly polar groups, the scheme (iii) for the basis set has been chosen for all ground and excited state calculations.

The TD-DFT methodology was then used to obtain excited states<sup>43–49</sup> by applying the same density functional and basis sets as for the ground state calculations. The oscillator strengths were taken from the TD-DFT calculations, and, along with the transition energies, a continuous spectrum was constructed using the finite-width Gaussian curve, where the empirical width corresponds to a thermal broadening of 0.05 eV. To visualize the localization of the excitons in space, the Natural Transition Orbitals (NTOs) and the spatially resolved transition density were calculated.<sup>50–52</sup> To quantitatively analyze the localization of the transition density, we employed a participation ratio, PR<sub>k</sub>, which is a heavily used technique for solid state physics applications in the analysis of defects and impurities.<sup>53,54</sup> The PR can be defined as

$$PR_{k} = \frac{\sum_{i}^{N} \rho(\mathbf{r}_{i})^{2}}{\left[\sum_{i}^{N} \rho(\mathbf{r}_{i})\right]^{2}} = \frac{\sum_{i}^{N} \psi(\mathbf{r}_{i})^{4}}{\left[\sum_{i}^{N} \psi(\mathbf{r}_{i})^{2}\right]^{2}}$$
(1)

where  $\rho$  is the electron density,  $\psi$  is a wave function (electron, hole, or quasi-particle), *i* is some spatial coordinate that uniquely defines each point in space with a total of N points, and *k* identifies a specific electronic state (e.g., Molecular Orbitals, MOs—for the ground state—or NTOs—for excitons). The PR<sub>k</sub> value varies from 0 to 1. When PR<sub>k</sub> approaches one, a small number of spatial points contribute more significantly than the other points, evidencing a localized nature of an exciton. Conversely, if  $PR_k$  is small, every point contributes nearly the same to the overall density, indicating in a completely delocalized exciton.

# 3. RESULTS AND DISCUSSION

The calculated HOMO-LUMO gap is 2.25 eV for the pristine (10,5) SWCNT, which is significantly blue-shifted compared to experimental values due to the lack of electron correlations for the ground state calculations and confinement effect of the finite-size (10 nm) nanotube model we use. However, this discrepancy does not affect the qualitative trends for comparing the electronic structure of pristine and functionalized SWCNTs, as investigated in our previous work.55 Consistent with previous studies, functionalization results in destabilization of the HOMO and stabilization of the LUMO-both associated with the sp<sup>3</sup>-defect site-leading to a decreased energy gap.<sup>3,10,13</sup> While the magnitude of this reduction is dependent on both the conformation and the chemical composition of the functional group, the topological position of the molecular adduct dominates (Figure 2b-d). In fact, changing the geometry of the defect results in a noticeable deviation between the energy gap of the pristine and functionalized SWCNTs ( $\Delta E_{\rm HL}$ ) up to 350 meV. In contrast, the  $\Delta E_{\rm HL}$  value reaches no more than 100 meV, even for functional groups with highly electronegative fluorines (Figure 2b).

A steric effect between two long alkyl groups attached to the same carbon ring of the nanotube, 2C<sub>6</sub>H<sub>13</sub>, negligibly impacts the electronic structure of the functionalized SWCNTs. As a result, the 2C<sub>6</sub>H<sub>13</sub> system exhibits similar values of  $\Delta E_{\rm HI}$  to the cases with a single alkyl and a hydrogen, C<sub>6</sub>H<sub>13</sub>/H, only demonstrating the dependence on the defect position (Figure 2b). This indicates that the impact of the second group on the electronic structure of the SWCNT is marginal for groups with weak electron-withdrawing ability, thereby validating the use of a hydrogen atom as the auxiliary group for modeling such systems. This is not the case for their fluorinated counterparts, where functionalization with two fluorinated chains,  $2C_nF_{2n+1}$ , results in a noticeable deviation of  $\Delta E_{\rm HL}$  from those of  $C_n F_{2n+1}/H$  structures (Figure 2c). A similar behavior is observed for two non-carbon adducts with a strong electronwithdrawing (F) or electron-donating  $(PF_2)$  character (Figure 2d). For all these cases, a strong electrostatic repulsionstrengthened by the inductive effect from an increased number of fluorine atoms in the alkyl chain-is expected to significantly affect the electron density of the SWCNTs, contributing to an observed reduction in the HOMO-LUMO gaps of the functionalized SWCNT, compared to its pristine counterpart. This gap energy reduction is the most pronounced for the structures with defects at the O<sub>-40</sub> and  $\mathbf{\hat{P}}_{20}$  positions. In contrast, a dipole moment increase of a single functional group  $(C_nF_{2n+1}/H)$  results in much smaller (a few meV) deviations of  $\Delta E_{\rm HL}$  values between functional species (Figure 2c,d). Aryl groups act very similarly to  $C_n H_{2n+1}/H$  and  $C_n F_{2n+1}/H$  alkyls, showing an insignificant effect of the halide substituents on the values of  $\Delta E_{\rm HL}$ , Figure S1a in the Supporting Information (SI).

Having analyzed trends of the ground state energies, we next inspect the behavior of the excited state energies calculated with TD-DFT. The calculated  $E_{11}$  optical transition is at 1.52 eV for the pristine (10,5) SWCNT being closer to experiment owing to electronic correlations effects in TD-

DFT. Similar trends for two functional groups  $(2R_1)$  versus one group  $(R_1/H)$  are observed for the excited states. Figure 3



**Figure 3.** Red shift of the first absorption peak  $E_{11}^*$  of the functionalized SWCNT from  $E_{11}$  absorption band of the pristine SWCNT ( $\Delta E_{11} = E_{11} - E_{11}^*$ ), as a function of the defect position. (a) Comparison between structures with two identical alkyl groups ( $2R_1 = R1/R_1$ , empty triangles and solid lines) and structures with one alkyl and hydrogen ( $R_1/H$ , filled triangles and dashed lines). (b) The effect of increasing the fluorinated alkyl chain length on the  $\Delta E_{11}$  for  $2R_1$  structures (empty triangles and solid lines) and  $R_1/H$  structures (filled triangles and dashed lines). (c) Comparison between non-carbon groups of fluorine (empty and filled black diamonds) and difluorophosphine (empty and filled cyan triangles) with the  $2C_6F_{13}$  (empty black triangles).

shows the red shift of the defect-associated lowest optical transition,  $E_{11}^*$ , from the main absorption band,  $E_{11}$ , of the pristine nanotube:  $\Delta E_{11} = E_{11} - E_{11}^*$ . Similarities in the behavior of the ground and excited state electronic structures are rationalized by a predominant contribution of the HOMO-LUMO pair to the lowest energy excitonic wave function of SWCNT.<sup>3,9,13</sup> Therefore, the  $\Delta E_{HL}$  at the ground state mainly follows the trends and characteristics of the low-lying excitons,  $\Delta E_{11}$ , as evidenced by comparison of Figures 2 and 3. Overall, using two polar groups as adducts produces larger red shifts of the lowest excitons than substituting one with a hydrogen atom, which is the most pronounced for  $2C_6F_{13}$  (Figure 3a,b) and 2F (Figure 3c) cases. These two structures provide the strongest red-shifted E<sub>11</sub>\* transitions, well-distinct from E<sub>11</sub>\* of other functional groups by about 60–100 meV at  $O_{-40}$  and  $P_{20}$ conformations (Figure S2 in SI), evidencing a dominant effect of the electrostatic interactions between neighboring functional groups on the lowest energy exciton.

Recent experimental data using high-resolution spectroscopy on a functionalized (6,5) SWCNT has detected a 19 meV difference between  $E_{11}^*$  excitons associated with defects due to functionalization by the 4-nitroaryl (strong electron-withdrawing group) versus 4-methoxyaryl (weaker electron-withdrawing group) and much smaller splitting of 4–6 meV between  $E_{11}^*$  of mixed defects when they are in very close proximity to each other.<sup>56</sup> Our calculations may explain these experimental trends. Assuming that similar types of molecular adducts can be attached to the same carbon ring of the nanotube, the species having strong electron-withdrawing abilities are expected to behave similarly to our 2F or  $2C_6F_{13}$ model, resulting in a stronger red shift of the  $E_{11}^*$  exciton, compared to those of structures with weak electron-withdrawing characters (like our  $2C_6H_{13}$  and aryl models). However, the mixture of two adducts with different electronwithdrawing abilities attached to the same carbon ring of the nanotube is expected to behave similarly to our F/H or  $C_6F_{13}/$ H models, resulting in insignificant  $E_{11}^*$  splitting between different species. Note that our calculations do not consider several sp<sup>3</sup>-defects and interactions between them. Instead, two adduct molecules contribute to a single defect, while the difference in their interactions—due to variations in electrostatic properties of adducts—uniquely affects the exciton red shifts.

A recent experimental study reported an inductive effect of functional groups in connection to the defect-associated spectral features of SWCNTs.<sup>25</sup> A linear correlation between red shifts of emission energies and electron inductive ability of the functional group has been suggested. To get atomistic insights into this question, we provide the dependence of  $\Delta E_{\rm HL}$  and  $\Delta E_{11}$  on the dipole moments of the SWCNT with various adducts to gauge the relationship between the electrostatic dipole and the defect-originated excitons in SWCNTs (Figure 4).

For a given type of the defect geometry, both  $\Delta E_{\rm HL}$  and  $\Delta E_{11}$  tend to increase with the dipole moment of the functionalized (10,5). This trend is better pronounced for the most red-shifted geometries of O<sub>-40</sub> and P<sub>20</sub> (filled triangles and empty squares in Figure 4) and is very minor for O<sub>20</sub> and P<sub>-40</sub> defect positions (empty triangles and filled squares in Figure 4) having the least distorted electronic structure compared to the pristine SWCNT. We also note the lack of obvious correlations between the optical intensity of the lowest exciton and the system dipole moment (Figure S3).

For all defect positions, the increase in  $\Delta E_{\mathrm{HL}}$  and  $\Delta E_{\mathrm{11}}$  with the dipole moment of the system is very minor until the dipole reaches the value of 6 D. The trend is better pronounced after an inflection point at around 6 D, which mainly corresponds to structures having two adducts with strong electrostatic interactions, such as  $2C_6F_{13}$ , 2F, and  $PF_2/F$  (Figure 4). This trend is more conspicuous for the ground state  $\Delta E_{\rm HL}$  than the excited state transitions  $\Delta E_{11}$  (compare Figure 4a to Figure 4b). For example, the exciton red shift of the 2F system (purple symbols in Figure 4b) is noticeably larger than that for the  $2C_6F_{13}$  system (brown symbols in Figure 4b) that has a higher dipole moment compared to the 2F structure. The deviations in the trend point to the importance of direct interactions of the highly electronegative fluorines with the SWCNT rivaling the inductive capacity of the fluorinated alkyl group. This is expected to directly modify charge distributions in the region of the defect beyond dipole influence. As such, the dipole moment of the adduct alone does not seem to be a good indicator of the defect-originated exciton red shifts for most species studied here.

To investigate charge redistribution and charge-charge interactions between different functional groups, we examine the charge of the tube atom connected to the molecular adducts  $(Q_T)$ , which is modified by hybridization, dipole, and inductive effects of the covalent attachment. Inductivity has been previously suggested to play a major role in resolving species-dependent shifts.<sup>25,29</sup> Here, we further explore this hypothesis, extending the studies to multiple binding configurations, including 2R<sub>1</sub> groups. Figure 5 demonstrates



**Figure 4.** Dependence of the (a) ground state energy gap difference,  $\Delta E_{\rm HL}$ , and (b) excited state red shift,  $\Delta E_{11}$ , on the electrostatic dipole moment of the functionalized SWCNT. Different colors of symbols represent different functional groups named at the top X-axis. The shape of symbols corresponds to the position of the defects at the SWCNT lattice. The size of the symbols corresponds to the oscillator strengths of  $E_{11}^*$  transitions: the larger the symbol, the larger the oscillator strength. Both the ground state  $\Delta E_{\rm HL}$  and the excited state  $\Delta E_{11}$  nearly monotonically increase with the dipole moment almost for all defect types and positions. This trend is less pronounced for  $\Delta E_{11}$  of structures at  $O_{20}$  (filled squares) and  $P_{-40}$  (empty triangles) positions, which correspond to the brightest, delocalized, and least red-shifted  $E_{11}^*$  excitons.

the correlations between the lowest energy exciton red shift  $\Delta E_{11}$  and the local charges of the defect-connecting tube atom,  $Q_{\rm T}$ , for each carbon-linking functional species and configuration we studied. Results for all functional groups, including 2F, F/H, PF<sub>2</sub>/H, and PF<sub>2</sub>/F, are shown in Figure S4. For the  $C_nH_{2n+1}$ , aryl, and F species, we observe that the charge induced on the nanotube atom has a positive sign, indicating that these molecular adducts work as electron-withdrawing groups (EWG) upon functionalization, with fluorines being the strongest EWG. In contrast, the fluorinated alkyl and PF<sub>2</sub> species have the opposite effect on  $Q_{\rm T}$  as electron-donating groups (EDG), with PF<sub>2</sub> acting as the strongest EDG.

Near the zero-charge values, we note an interesting difference between the *ortho* and *para* positions (Figure 5). The alkyl/aryl EWG species with a positive  $Q_T$  exhibit the value of the charge 3–4 times smaller in the *para* than the *ortho* configurations. The reverse trend, although less pronounced, is true for the fluorinated alkyl EDG species inducing negative charge  $Q_T$ . For EWG alkyl/aryl species, this trend can be rationalized by a larger number of sp<sup>2</sup>-hybridized nearest neighbor carbons in *para* (3), compared to *ortho* (2) configurations, which effectively minimizes the value of the positive  $Q_T$  in the *para*-defect. Significantly reduced  $Q_T$  in *para* positions of EWG can be also associated with a less reactive

ability of para versus ortho positions. It was previously shown that there is a predominant selectivity for ortho binding configurations of aryl groups on the nanotube lattice, due to  $\pi$ -orbital misalignment enforced at the *ortho*-defect, making it highly reactive.  $\overset{3,29,30}{}$  However, for the negative  $Q_T$  case in fluorinated alkyls, the reverse trend does not follow this argument, showing less distinction in the  $Q_{\rm T}$  charge between the para- and ortho-defects. Overall, our data suggest that EWG functionalization leads to more localized positive charge at the defect in ortho attachment, whereas EDG functionalization more strongly localizes the negative charge in the para-defect. However, either the largest or the smallest positive or negative charges on the defect atom does not necessarily provide the largest red shift of the defect-associated excitation. For example, the red shift of the longest fluorinated alkyl chains,  $2C_6F_{13}$ , is the greatest in all defect positions but does not induce the largest negative charge in any configuration. This implies that there are additional factors determining the exciton red shift.

Consistent with previous reports,<sup>3</sup> there is a straightforward correlation between the exciton localization around the defect and its red shift: The higher the degree of exciton localization, the more the defect-associated exciton is red-shifted with respect to the  $E_{11}$  band (Table S1). Thus, regardless of the adduct species, a higher degree of localization of the transition charge density is seen near the defect in the  $O_{-40}$  and  $P_{20}$ positions, leading to the largest exciton red shift in comparison to the highly delocalized exciton of the  $O_{20}$  and  $P_{-40}$  positions, resulting in insignificant splitting between the defect-associated exciton and the  $E_{11}$  band (Table S1 and Figure S2). In our previous studies, we have provided physical interpretations of the difference in red shifts of optical features caused by the adduct positions: Similar to our finding here, strong red shifts in emission energies of certain defect geometries have been correlated to exciton localization, which is attributed to geometry-dependent variation in the nodes of the frontier orbitals.<sup>37</sup> In particular, when ortho-functionalization is performed along the less reactive axial bonds in the mod2 SWCNTs, such as  $O_{20}$  in (10,5) considered here, the resulting bond lies parallel to the nodes of the frontier orbitals and perturbations to the electronic structure are minimal.<sup>57</sup> Thus, the electronic structure and optical properties of the SWCNT functionalized along the axial O<sub>20</sub> direction only marginally deviate from the pristine nanotube, resulting in the smallest red shifts. In contrast, the strongest red shift in mod2 nanotubes is the result of the defect along the highly reactive off-axial bonds  $(O_{-40}$  in (10,5) SWCNT), where the nodes of the frontier orbitals are bisected by the defect, resulting in strong exciton localization due to a significant disturbance to the orbital nodal structure in the vicinity of the defect.<sup>57</sup> Similar bisected node structures of the frontier orbitals in the  $\mathrm{P}_{20}$  position also result in strong localization of the orbitals near the defect site, leading to the largest red shift of the lowest exciton.

This localization effect is analogous to exciton trapping by a quasi-one-dimensional potential energy well formed by the lattice defect.<sup>12–14</sup> The well depth and width are defined mostly by the geometric configuration of the defect associated with geometry-dependent variations in the nodes of the frontier orbitals<sup>57</sup> and to a weaker extent by the chemical species themselves. The latest is rationalized by the localization of the electron-hole density mainly on the defect, rather than on the functional group. However, the P<sub>20</sub> configuration allows for some density to be drawn onto the anchoring atoms of the

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**Figure 5.** Charge on the nanotube carbon connected to the molecular adduct  $(Q_T)$  plotted as a function of the exciton red shift,  $\Delta E_{11} = E_{11} - E_{11}^*$ , for each configuration (a-f) and only carbon-linking functional groups. Increasing size of the symbol is correlated with increasing dipole moment and the chain length of the system.

functional group regardless of the species (Table S1). Similar behavior is observed for  $O_{-40}$ . This occurs even for a weakly EWG species such as non-fluorinated alkyls. This is due to the reduced width of the trapping well, which implies highly increased exciton localization in  $P_{20}$  and  $O_{-40}$  configurations, with some extension to the functional groups. Therefore, the exciton red shifts are more sensitive to the adduct species in  $P_{20}$  and  $O_{-40}$  configurations, compared to other defect positions (Figure 3).

We use PRs (as defined in eq 1) to provide a quantitative description of exciton localization in the considered systems, where larger PR values indicate increased localization. Two contrasting examples of such dependence can be seen in the top panel of Figure 6, where the NTOs corresponding to delocalized (left) and localized (right) excitons differ in PR by almost a factor of 10. Figure 6a shows the dependence of the averaged  $Q_{\rm T}$  charge on the PR, where the charge values are averaged over all carbon-containing functional groups, separately presented for the EWG groups with the positive charge and EDG with the negative charge. For both EWG and EDG, the para-defects exhibit negligible changes in  $Q_T$  values on the exciton localization, with nearly neutral charge for EWG and the maximum negative values for EDG. In contrast, orthodefects show a slight, but well-resolved, decrease in the positive charge with the exciton localization for EWG groups and increase in the negative charge for EDG groups with even stronger dependence (Figure 6a). These trends have potential importance for determining the reactive defect configuration implying that ortho-defects are likely more selective toward specific exciton energy, as it was previously suggested in literature reports.<sup>3,29,30</sup>

The dependence of the red shift of the defect-originated exciton on its localization for each defect configuration indicates a nearly ideal linear trend, when the values are averaged over all  $R_1/H$  species: The red shift of the defectassociated exciton strongly increases with its wave function localization as indicated by the PR (Figure 6b). More thorough analysis of this dependence focusing on individual contribution of each species is shown in Figure 6d and Figure S5.

Within each defect type, there is a weak increase of the exciton red shift with its PR, exhibiting more deviations for adducts at the defect positions with  $\Delta E_{11} > 100$  meV. Comparison between the averaged data and individual ones shown in Figure 6b,d suggests the interplay of two effects playing a role in the localization of the excitonic state and, therefore, its red shift, but at different scales: defect configuration – ca. 100 meV, adduct species – ca. 10 meV.

As we have shown in Figure 4, most functional groups exhibit weak dependence of the exciton red shift on the dipole moment of the system, where only adducts with the largest dipoles cause significant effects. Figures 6c and S5b also demonstrate the minor dependence of the exciton localization on the dipole moment by showing the PR as a function of the dipole moment for each system. Within each defect configuration, on average, nearly linear dependence between the PR and dipole moments can be observed, although the values for each species are significantly scattered from the averaged line. While the exciton localization insignificantly affects the  $Q_{\rm T}$  charge for *para*-defects (Figure 6a), the dipole moment of species at the para positions demonstrate much stronger dependence, compared to *ortho* positions (Figure 6c). The exception is O<sub>80</sub> and P<sub>80</sub> defects, which we single out as having the largest standard deviations from the linear trend for these cases, compared to other defects. In general, significant scattering of data in Figure 6c points to a stronger dependence of the dipole moment on the chemical composition of adducts,



Figure 6. Correlation between individual electrostatic properties of molecular adducts and the participation ratio (PR) of functionalized (10,5) SWCNT. Two natural transition orbitals (NTOs) for the 2F O<sub>20</sub> and P<sub>20</sub> geometries corresponding to the hole with their associated PR value, depicting how exciton localization is quantified using the PR (top panel). (a) Charge on the nanotube atom connected to the chemical adduct  $(Q_T)$  as a function of the participation ratio. The positive-(negative-)charge panel is the average charge of the alkyl and aryl (fluorinated alkyl) species for all carbonbased adducts in each configuration, including both R<sub>1</sub>/H and 2R<sub>1</sub> systems. The points are connected with dashed lines for better visualization of the trend. (b) Red shift of the defect-originated exciton from the main  $E_{11}$  band  $(\Delta E_{11} = E_{11} - E_{11}^*)$  as a function of its participation ratio. The symbols represent an average over all R<sub>1</sub>/H molecular adducts in a given defect configuration. (c) The dependence of the participation ratio on the dipole moment of the system for all considered functional groups. (d) The dependence of the red shift  $\Delta E_{11}$  on the participation ratio for R<sub>1</sub>/H molecular adducts in a given defect configuration. A linear fit is shown by a solid (ortho-defects) and dashed (para-defects) lines.

rather than the ability of the sp<sup>3</sup>-defect to localize the exciton. This difference can be rationalized by polarization properties of the excited state due to the charge density redistribution upon excitation that compensates the effect of electrostatic charge distribution associated with the electrostatic dipole moment of the ground state.

# 4. CONCLUSIONS

Using DFT-based methodologies, we have investigated the effect of the intrinsic molecular dipole moments and the local charges induced by various molecular adducts on the energy and wave function localization properties of the optically active defect-associated excitons in a (10,5) functionalized SWCNT. Our results evidence that neither the dipole moment nor inductive features—including local charge on the SWCNT-defect interface—can serve as reliable descriptors of the energy splitting between the defect-associated exciton and the main  $E_{11}$  band of the SWCNTs. When considering the electrostatic dipole moment of the system, the overall trends are consistent with the expectation that the red shift of the defect-associated exciton alized

SWCNT, but exhibiting very weak dependence, except for significantly large values of the dipole moments (>6 D). Notably, such a large dipole moment of the system can be introduced only by two strong EWG or EDG, such as 2F or  $2C_6F_{13}$  and  $PF_2/F$  attached to the same carbon ring of the nanotube. However, the effect of a single group—F/H,  $C_nF_{2n+1}/H$ , and  $PF_2/H$ —is nearly the same as for weak EWG or EDG groups, such as aryls and non-fluorinated alkyls. In addition, the dipole moment of the species poorly correlates with the exciton localization, while the latter exhibits a strong linear dependence on the red shifts of the defect-associated excitons. Our calculations demonstrate that the degree of localization of the defect-associated exciton, which can be measured using the participation ratio, is a suitable empirical parameter to determine the energy splitting between the defect-originated exciton and the  $E_{11}$  band in the covalently functionalized SWCNTs. However, no adduct-dependent parameter was found to strongly correlate with this variable.

The charge at the sp<sup>3</sup>-defect shows a distinct behavior between ortho- and para-defect configurations, regardless of molecular adducts, and likely relevant to the chemical reactivity of the defect position. However, the species-dependent trends in charges induced by molecular adducts do not directly correlate to the exciton red shift across all considered functional groups. We explain such a weak dependence by the redistribution of the exciton density of the excited state that weakens the polarization effect of the electrostatic dipole moment of the ground state. This leads to the localization of the lowest energy exciton mainly around the sp<sup>3</sup>-defect of the nanotube with a negligible extent of its density distributed to the anchoring atoms of the functional group. Therefore, the defect configuration plays the leading role in the red shift of the defect-associated exciton with respect to the  $E_{11}$  band of the pristine SWCNTs on the order of ca. 100 meV, while the polarization properties of molecular adducts lead to significantly smaller red shifts on the order of ca. 10 meV. In ref 57, it was shown that all investigated trends in electronic and optical properties relating to the defect configuration are similar for SWCNTs of the same mod(n-m,3). As such, we also can assume that all trends obtained for (10,5) can be generalized to other SWCNTs of mod2. These computational results may aid in finding improvements of synthetic protocols to achieve desired emission of functionalized SWCNTs.

## ASSOCIATED CONTENT

#### **Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpcc.0c10157.

Various additional plots to supplement most main figures, simulated absorption plots for comparing the effects of fluorination and Natural Transition Orbitals (NTOs) to showcase the size of the exciton and its locality to and on the defect, and tables of data for most of the necessary observables in this work, as there are many overlapping data points in the figures (PDF)

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#### Notes

The authors declare no competing financial interest.

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