

Linear and Helical Cesium Iodide Atomic Chains in Ultranarrow Single-Walled Carbon Nanotubes: Impact on Optical Properties

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ABSTRACT: One-dimensional (1D) atomic chains of CsI were previously reported in double-walled carbon nanotubes with ~0.8 nm inner diameter. Here, we demonstrate that, while 1D CsI chains form within narrow ~0.73 nm diameter single-walled carbon nanotubes (SWCNTs), wider SWCNT tubules (~0.8–1.1 nm) promote the formation of helical chains of CsI 2 × 1 atoms in cross-section. These CsI helices create complementary oval distortions in encapsulating SWCNTs with highly strained helices formed from strained Cs₂I₂ parallelogram units in narrow tubes to lower strain Cs₂I₂ units in wider tubes. The observed structural changes and charge distribution were analyzed by density-functional theory and Bader analysis. CsI chains also produce conformation-selective changes to the electronic structure and optical properties of the encapsulating tubules. The observed defects are an interesting variation from defects commonly observed in alkali halides as these are normally associated with the Schottky and Frenkel type. The energetics of CsI 2 × 1 helix formation in SWCNTs suggests how these could be controllably formed.

KEYWORDS: atomic chains, carbon nanotubes, electron microscopy, density functional theory Raman spectroscopy, photoluminescence

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"True" one-dimensional (1D) atomic chains were first formed by relocating atoms with a scanning tunnelling microscope (STM) tip under cryogenic conditions,¹⁻⁴ demonstrating quantum scale device fabrication, albeit with a one-off approach that limited studies to the laboratory. More recently, $M_{1-x}M'_{x}X_{1-y}X'_{y}$ (M or M' = Mo or W and X' = S, Se, Te, or I) 1D nanowires and nanowire junctions were formed by electron beam lithography from the corresponding $M_{1-x}M'_{x}(X_{1-y}X'_{y})_{2}$ 2D dichalcogenides.⁵⁻⁸ Templating atomic chain formation in carbon nanotubes, however, offers by far the greatest compositional diversity and scalability for 1D materials fabrication observed so far. Nanotube insertion occurs from the liquid or vapor phase and its van Der Waals surface templates integral,⁹ helical^{10–12} or "zigzag"¹³ atomic columns in cross section, and in the limit, "true" 1D atomic chains in the narrowest tubules,^{12–26} although the characterization of single element materials including Se, Te, and S in porous materials in

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Figure 1. (a) Six HRTEM images and Wien-filtered details of linear single atomic width Cs–I chains in SWCNTs with measured diameters increasing from 0.73 to 0.81 nm. (b) Structure models of Cs–I linear chains inside SWCNTs with (n,m) conformations specifying diameters matched to the composites imaged in a. (c–h) HRTEM images, corresponding composite structure models and multislice image simulations (denoted SIM) of twisted Cs–I 2 × 1 atomic chains in SWCNTs spanning (c,d) average 0.88 nm diameter, (e,f) average 0.94 nm diameter and (g,h) average 0.95 nm diameter. All three nanotubes distort ~10% in the cross section in response to the twisting of the embedded 2 × 1 atomic chains (see also Figure 2a–f) but, as the diameter of the encapsulating SWCNT widens the internal I–Cs–I bond angles relax from ~72–83° (*i.e.*, c,d and e,f) to ~90° as indicated by the inset motifs.

aligned porous materials such as mordenite and AlPO₄ have also been extensively reported. 9,27,28

Several groups have demonstrated 1D and helical iodine chain formation in single-walled carbon nanotubes (SWCNTs)^{10,15} and double-walled carbon nanotubes (DWCNTs)^{16,17} and linear chains of S¹² carbyne¹⁷ Te¹⁸ P¹⁹ and As²⁰ in narrow SWCNTs. Functionality can be extended if the nanowires are formed from binary or ternary phases, and Senga et al. demonstrated linear chains (or "true" 1D crystals) of CsI¹³ and NaI, LiI, CsF, and Cs_{1-x}K_xI within DWCNTs.²³ Binary 1D chain filling in narrow SWCNTs was also demonstrated for chalcogenides SnTe²¹ and SnSe,²² for which computations with density-dunctional theory (DFT) indicated enhanced thermoelectric properties in isolated nanowires of the former. 1D properties have been investigated by Raman spectroscopy for I_3^- , I_5^{-24-26} KI,²⁹ and HgTe.³⁰ True 1D physical phenomena such as Peierls distortions (*e.g.*, Te_x)¹⁸ and Charge Density Waves (e.g., I_x^{-}) can potentially also be addressed.¹⁵ More extrinsic properties of 1D crystal encapsulated SWCNT (i.e., 1D@SWCNT) composites include the fine-tuning of nanotube properties by doping or charge transfer.^{24–26,30–33} Calculations from first-principles suggest that 1D CsI crystals perturb the band structure of the narrowest SWCNTs with fluctuations in the electron density in the CNT commensurate with the periodicity of the CsI chain and which diminish as the encapsulating diameter increases and/or the 1D structure deviates from linearity.¹

Our investigation establishes the precise experimental structural evolution of atomic chains of CsI formed inside narrow SWCNTs with diameters of 0.7-1.1 nm, producing structures that vary from "true" 1D CsI, to highly strained 2×1 CsI helices with diameter. We imaged these structures using a combination of high-resolution transmission electron microscopy (HRTEM) and annular dark field (ADF) in scanning transmission electron microscopy (STEM) imaging mode and also atomically resolved electron energy loss spectroscopic

(EELS) imaging. We reveal by Raman, optical absorption, and photoluminescence spectroscopy that the structural distortions of the SWCNTs induced by the CsI filling selectively modify the optoelectronic properties of tubules with specific (n,m) chiralities.

RESULTS AND DISCUSSION

Samples of CsI-filled SWCNTs were produced by filling 20 mg quantities of preoxidized SWeNT nanotubes by sublimation using an adapted procedure¹³ as described in the Methods. Preliminary imaging studies were performed at 80 kV in a double aberration corrected JEOL ARM200F instrument operated in phase contrast HRTEM and annular dark field (ADF) imaging modes. Further high angle annular dark field (HAADF) imaging studies were performed at 60 kV in a Nion UltraSTEM with EELS being recorded on a Gatan Enfina spectrometer, which also enabled the acquisition of atomically resolved Cs and I chemical maps with a 20 pA electron beam current and ~0.1 nm resolution. Further experimental details are recorded in the Methods.

HRTEM, ADF Imaging, and EELS Mapping. Figure 1 shows the evolution of the CsI microstructure with SWCNT cross-section for a SWCNTs ranging in diameter from 0.73 to 0.95 nm as imaged in plan view by aberration-corrected HRTEM. In the diameter range 0.73-0.81 nm (Figure 1a,b), linear -Cs-I-Cs-I- atomic chains are observed with an average interatomic separation of ~0.36 nm, comparable to the ~0.34 nm separation recorded within CsI-filled DWCNTs¹³ with innermost SWCNT diameters as small as ~0.80 nm. When the imaged SWCNT has a diameter just larger than ~0.87 nm (Figure 1c,d), a distortion of both the 1D crystal and the nanotube is observed in which the crystal forms a twisting 2×1 "zigzag" configuration with -I-Cs-I- internal angle of $\sim 71^{\circ}$ with a loss of linear symmetry. A $\sim 10\%$ oval distortion in the SWCNT cross section is observed coincident with the 2×1 nanostructure twist, comparable to distortions reported for



Figure 2. (a-c) HRTEM lattice image, multislice simulation and a model of a 2 × 1 helical chain of CsI formed inside a distorted ~0.90 nm average diameter SWCNT. The ~14 nm pitch and distortion can be seen by viewing (a-c) at a glancing angle in the direction indicated. The relative orientation of the two-atom CsI unit in ovally distorted SWCNTs relative to the electron beam (black arrow) is given by cross-sectional models at the bottom of c. (d-f) HRTEM lattice image, multislice simulation, and corresponding models of a longer ~36 nm pitch 2 × 1 helical chain imaged inside a ~0.94 nm diameter SWCNT, assignations as for (a-c). (g) Four ADF STEM images (I–IV) of five CsI 2 × 1 helical chains inside SWCNTs showing the variability in pitch, even within a single SWCNT. Cs–I atomic distances $(Cs-I_{av})$, measured longitudinally for all micrographs (*i.e.*, (a), (d), (g(I-IV))), average in the range 0.36–0.38 nm. Note that for all images, whenever the electron beam passes through a two-atom column, we see a doubling of the observed atom contrast.

SWCNT encapsulating quasi-1D chains of iodine, ^{10,15} HgI₂, ¹¹ and CoI₂.³⁴ The SWCNT in Figure 1c,d has an average diameter of 0.88 nm with narrower ~0.81 and wider ~0.91 nm sections visible. In a second example (Figure 1e,f), a SWCNT is filled with 2×1 CsI in which the average diameter increases to 0.94 nm with an increase in the I–Cs–I angle to ~75°. In a third example (Figure 1g,h), a 2×1 CsI chain with undistorted 90° internal I–Cs–I angle is observed, but it exhibits a short 180° helical twist defect with a corresponding distortion in the SWCNT (average diameter ~0.95 nm). In the above, local variations in the local imaged plan view diameter are sometimes seen as a result of debris on the SWCNTs, knock-on damage, or radiolysis, but the overall fluctuations in diameter are smooth.

In Figure 2a–g, seven 2 × 1 CsI chains imaged by HRTEM (Wien-filtered) and ADF STEM with evident helical symmetry are presented. Viewing each imaged object at glancing angles in the indicating arrowed directions enables the helicity to be more easily visualized. The multislice simulations in Figure 2b,e are based on the corresponding models in Figure 2c,f, respectively, with the orientation of a two-atom CsI unit with respect to the electron beam at each horizontal position indicated below. The pitch of the helices imaged in Figure 2 varies from ~5 nm (Figure 2gII) to ~36 nm (Figure 2d). The Cs–I distances, averaged longitudinally along each helical chain, vary from 0.36 nm up to 0.38 nm, comparable to those recorded by Senga *et al.*¹³ but significantly longer than the I–I distances reported for charged iodine chains (see also Figure 5d–f).^{10,14–16}

Confirmation of the chemistry and local stoichiometry of both linear and helical chain CsI filling is given in Figure 3a–i. Aberration-corrected HAADF images and atomically resolved elemental EELS maps and matching models are presented for a linear –I–Cs–I–Cs–I– chain (Figure 3a–c) and for a helical 2 \times 1 Cs–I chain (Figure 3e–g). EEL spectra (Figure 3d) obtained from the boxed regions in Figure 3b show M_{4.5} edges from discrete Cs and I atoms corresponding to the -I-Cs-I-Cs–I– model (Figure 3c). An EEL spectrum, obtained from an imaged Cs–I atom pair with doubled Z contrast (Figure 3f), shows both Cs and I M_{4.5} edges (Figure 3h), whereas discrete atoms indicated by the boxes in the lower part of Figure 3f produce separate Cs and I M_{4.5} edges (Figure 3i) in good agreement with the perspective helical model in Figure 3g. Average diameters of the SWCNTs in Figure 3a,e are 0.78 and 0.91 nm, respectively. Average Cs-I (i.e., Cs-I_{av}) spacings for linear or helical 2×1 chains along SWCNTs were measured by direct imaging, by either fast Fourier transforms (FFTs)¹³ calculated from HRTEM images or direct electron diffraction in both cases obtained from aligned filled SWCNT bundles.35 Direct imaging measurements averaged over 20 10-20 nm long CsI linear chains reveal a Cs-I_{av} of ~0.36 nm. Cs-I_{av} lengthens to ~0.37 nm for a similar number of measured 2×1 helical Cs–I chains, reflecting the somewhat higher strain in the latter. FFTs obtained from bundles of SWCNTs filled with CsI (Figure S2a,b) reveal some variability in $Cs-I_{av}$ from bundle to bundle, which indicates population differences in linear 1D CsI chains versus 2×1 Cs–I chains in these composites.

DFT Modeling and Bader Analysis. The structures of atomic chains of CsI in varying degrees of confinement can be explored by DFT^{36,37} (Supporting Information) combined with charge distribution analyzed by Bader analysis.³⁸ We modeled the isolated CsI 1D wires initially from relaxed and stretched ion approaches, applying chain perturbations and examining the efficiency of the calculations using finite chains. A basic force-



Figure 3. (a) HAADF image of a discrete CsI atomic chain imaged in a ~0.78 nm diameter SWCNT. (b) HAADF detail and corresponding Cs and I EELS chemical maps showing alternating Cs and I. (c) Representative model of a 1D CsI atomic chain in an undistorted (6,6) SWCNT. (d) Cs and I $M_{4,5}$ EEL spectra obtained from the 2nd and 3rd atom, respectively, according to the indicating boxes in (b). (e) HAADF image of a rotating 2 × 1 helical chain imaged in a ~10% sympathetically distorting ~0.91 nm (average diameter) SWCNT. (f) HAADF detail and corresponding Cs and I $M_{4,5}$ EELS chemical maps showing coincident Cs and I atom pairs (top) and discrete CsI 1D chains (bottom) and Cs and I EELS chemical maps. (g) Perspective structure model of rotating 2 × 1 helical chain corresponding to the HAADF detail in (f). (h) EEL spectra obtained from a single 2 × 1 atom column obtained from the yellow boxed region in (f) in which both the Cs and I $M_{4,5}$ edges can clearly be seen. (i) CsI $M_{4,5}$ spectra obtained from the two discrete Cs and I atoms and indicated by the orange and blue boxed regions in (f).



Figure 4. (a) Energy change in a twisting 2×1 chain of alternating dipoles into a helix of varying pitch. Results for both PBE-DFT and the fitted potential (parameter set 2, see the SI). Inset: dipole helix with 12 ion pairs. (b) Formation energy (E_f) of encapsulated CsI in SWCNTs calculated per atom energy difference between a formed complex structure (E_c) , a pristine SWCNT (E_{CNT}) , and the sum of the chemical potentials of Cs and I atoms in a mixture of noninteracting atomic gases, E_{at} (*i.e.*, $E_f = E_c - E_{CNT} - E_{at}$) and plotted versus effective SWCNT diameter and averaged for an ovally distorted SWCNT. (c) Plot showing the difference in energy of a "Tilted" 2×1 CsI chain with -I-Cs-I- bond angles <90° versus a "Ladder" 2×1 CsI chain with -I-Cs-I- bond angles <90° versus stress relative to the bulk.

field approach was utilized to examine different twisting conformations, summarized in Figure 4a, with which the fitted potential of a twisted helical 2×1 Cs–I chain is compared with the same result established by DFT incorporating the Perdew–Burke–Ernzerhof (PBE) functional.³⁷ At the DFT level of accuracy, we then established the dependence of the formation energy of atomic chains of CsI as a function of the average diameter of undistorted and ovally distorted SWCNTs, respectively (Figure 4b, see also Figure S13b). A comparison of the energy trends for the experimentally observed linear –I–

Cs–I–Cs–I– and 2 × 1 CsI chains reveals a minimum ~0.82 nm SWCNT average diameter (note: in the case of a 2 × 1 CsI chain, the numerical "average" diameter of the encapsulating oval SWCNT corresponds to the numerical mean of its narrowest and widest measured diameter (*i.e.*, Figure 1c–h and Figure 2a–f) in which the formation of both is energetically possible but the linear -I-Cs-I-Cs-I- is energetically preferred (Figure 4b). We next identify a SWCNT diameter of ~0.92 nm in which linear CsI chains coexist with highly strained zigzag 2 × 1 CsI chains with reduced I–Cs–I angles

(green and red curves, Figure 4b). Above ~0.92 nm diameter, relaxed 2×1 CsI chains with ~90° internal I–Cs–I angles (red curve, Figure 4b) in SWCNTs, in good agreement with experimental results (Figure 1c–h). In Figure S13b, the formation energy study is extended to the next structure in the series, 2×2 CsI, for which the diameter crossover is ~1.05 nm.

The expected C, Cs, and I charge redistributions for linear (1D) Cs–I chains and 2×1 CsI as a function of encapsulation in narrow to wide armchair (*n*,*n*) SWCNTs predicted by Bader analysis are presented in Table 1. Linear -I-Cs-I-Cs-I-

Table 1. Bader Charge Analysis in Terms of Outermost Valence Electrons and Formation energies (E_f) of Linear Chain 1D and 2 × 1 Chain 2D CsI@SWCNT

		average population of electronic orbitals			
	SWCNT diameter (nm)	Cs, [] 5s5p6s	I, [] 5s5p	С, []2р	formation energy $(E_{t^{p}} eV)$
1D@(4,4) SWCNT	0.525	7.196	8.270	4.011	3.261
1D@(5,5) SWCNT	0.678	7.662	8.200	3.994	-1.517
1D@(6,6) SWCNT	0.814	7.752	8.157	4.002	-2.135
2D@(6,6) SWCNT	0.814	7.655	8.257	4.002	-1.332
2D@(7,7) SWCNT	0.945	7.717	8.201	3.993	-2.264

chains encapsulated in very narrow SWCNTs (*i.e.*, (4,4) SWCNTs) are calculated to have a larger charge redistribution and higher formation energy than linear chains formed in (5,5), (6,6), and (7,7) SWCNTs, suggesting mostly steric confinement of CsI within SWCNTs. In Figure 4c, using the electrostatic potential as steric confinement for an "implicit" 0.814 nm diameter (6,6) SWCNT nanotube,¹⁸ we have studied that different degrees of stress may cause 2×1 CsI to vary in structure from a highly strained "tilted" 2×1 parallelogram form to a more relaxed or square-planar or "ladder" form as indicated by the inset structures. "Tilted" and "relaxed" 2×1 CsI ladder structures may coexist within (6,6) SWCNTs with the latter being ~3.8 meV/atom lower in energy at 0 K.

Dynamic Imaging, Comparison with Iodine. Figure 5 reveals CsI dynamical imaging behavior in SWCNTs and images of the same tubules filled with iodine for comparative spectroscopic measurements. In Figures 5a,b we see the motion of short 1D CsI fragments under the electron beam in straight and curved SWCNTs using $0.88-1.44 \text{ pA/cm}^2$ e-beam dose. In both cases, short CsI chains move in a nonsmooth way along SWCNTs (Figure 5a,b; Movies S1 and S2). If a SWCNT is continuously filled with long CsI chains, little motion is observed (Figure 5c and Movie S3) and local deviations from the average Cs–I separation (CsI_{av}) can be detected. In the straightened detail in Figure 5c, the Cs–I distance varies from ~0.33 to ~0.37 nm. For bundles of SWCNTs filled with iodine, we obtain similar average I–I spacings (*i.e.*, $I-I_{av}$) to those reported within DWCNTs¹⁶ or narrow SWCNTs.^{10,15,16,25,26} Using the FFT protocol for imaged filled SWCNTs bundles (Figure 5d, inset),^{13,35} the $I-I_{av}$ is ~0.30 nm. Iodine chains in individual SWCNTs show significant local differences, with some continuous I_x chains and ~0.55–0.58 nm long I_3^- trimers (Figure 5e). The extent to which trimer formation occurs in DWCNTs was reported to increase as the inner SWCNT

diameter increased from 0.73 to 0.89 nm,¹⁵ consistent with our results. A schematic comparison of the relative microstructure of I_x chains versus CsI linear chains is shown in parts f and g, respectively, of Figure 5.

Raman, Optical Absorption, and Photoluminescence Measurements. Pristine, CsI-filled, and I-filled SWCNT samples were characterized by Raman, optical-absorption spectroscopy (OAS), and photoluminescence (PL) spectroscopies, allowing (n,m) chiral distributions and changes in electronic structure to be evaluated for filled (6,4) to (10,6) SWCNTs, with outlier contributions from wider SWNTs. Resonant Raman spectroscopy was performed at 532 and 660 nm (Figures 6a, S14c, and S14a,b) for pristine, CsI-filled, and iodine-filled SWCNTs.⁵ The 266 and 275 cm⁻¹ radial breathing mode (RBM) peaks in unfilled, CsI-filled, and iodine-filled SWCNTs can be assigned to (7,6) and (8,4) tubes, respectively.³⁹⁻⁴¹ More striking are three additional peaks at 108 and 153 cm⁻¹ for iodine-filled SWCNTs and a 192 cm⁻¹ enhanced resonance, denoted $(12,5)^{E}$, observed exclusively for CsI-filled SWCNTs. The 108 and 153 cm⁻¹ resonances correspond to iodine trimers, I_5^- fragments, or I_r polymeric chains²⁵ and are also observed at 660 nm (Figure S14a). Similar weak resonances are also seen in SWCNTs filled with CsI, indicating that a small amount of elemental iodine may have separated to create iodine filled SWCNTs, also observed in NiI₂ filled SWCNTs.¹⁶ The 192 cm⁻¹ resonance is exclusive to the CsI filled SWCNTs and conforms to an outlier semiconducting (12,5) nanotube⁴⁰ with a diameter of \sim 1.20 nm, which would accommodate $2 \times 2 \text{ CsI}^{11}$ (see also Figure S13b). This indicates "tuning into resonance" similar to reported behavior for 2×2 KI crystals in ~1.14 nm diameter SWCNTs.⁴¹ That this occurs in larger SWCNT for CsI may be a consequence of the larger diameter Cs⁺ ion (*i.e.*, 181 pm versus 152 pm for K^+)⁴² which would incur a $\sim 8\%$ expansion in the 2 \times 2 CsI cross section, commensurate with an increase in the observed diameter. The results in Figures 6a and S14a-c can be taken as an indication that the CNTs are filled, but a more detailed study of the influence of the filling on the RBM modes requires either micro-Raman on individual filled CNTs or a multiwavelength study on an ensemble of filled CNTs.

In Figure 6b, the OAS of unfilled SWCNTs show strong bands at 590–860 and 860–1300 nm corresponding to E_{22}^{S} and E_{11}^{S} excitonic transitions from numerous overlapping (n,m) chiralities.^{39,40} Following filling with CsI, most E_{11}^{S} and E_{22}^{S} absorbance features are unchanged apart from those for (6,5)SWCNTs, which can only fill with linear -I-Cs-I-Cs-Ichains which show enhanced absorbance at 660 and 976 nm, respectively. In (7,5) and (7,6) nanotubes, which fill with 2×1 CsI, E_{11}^{s} and E_{22}^{s} absorbances at 660 and 1147 nm are still present. In nanotubes with diameters >1.12 nm, filling with 2×2 CsI is expected which theory predicts will *n*-dope the nanotubes.⁴³ Literature values of the work function of each constituent can also be examined to assess the potential for charge transfer, *i.e.*, ~5 eV for SWCNTs,⁴⁴ ~5.5 eV for iodine (bulk),⁴⁵ and 3.4 eV for CsI (bulk)⁴⁶ with the latter two reducing with lowered dimensionality.^{47,48} Iodine therefore acts as an electron acceptor, receiving electrons from the SWCNT and making the nanotube p-type,⁴¹ whereas CsI should act as an electron donor and create *n*-type nanotubes. The degree of charge transfer was assessed from the absorption strength in the E_{11}^{s} range^{49,50} while the strong reduction in absorbance for iodine filling (Figure 6b) indicates a large shift in chemical potential. CsI filling does not noticeably alter the absorption



Figure 5. (a) Two images from an 18-image sequence CsI atoms in a partly filled straight SWCNT (0.74 diameter) showing disordered dynamical motion (yellow arrows, see also Movie S1). (b) Pair of images from a 20-image sequence in which canonical motion is observed in a partially filled curved 0.75 nm diameter SWCNT (yellow arrows, see also Movie S2) regulated by a defect near the center of the SWCNT. (c) Pair of images obtained 18s apart in 30-image sequence of continuously CsI-filled 0.74 nm diameter SWCNTs. In the animation, it is clear that the Cs and I atoms are rigidly fixed (orange arrows, see also Movie S3). A filtered, straightened region (bottom) shows a 14-atom CsI sequence with an average Cs–I separations of 0.35 nm but with variations from 0.33 to 0.38 nm. (d) ADF-STEM image of an aligned bundle of SWCNTs containing exclusively 1D iodine chains. Inset FFT from boxed region I indicates an average I–I separation of 0.301 nm.¹³ (e) ADF-STEM image of a discrete iodine chain forming trimers as reported in ref 15 with lengths indicated. The "stretched" trimer at right is an outlier. (f–g) Models of I and CsI chains embedded within appropriate diameter SWCNTs.

strength, suggesting small charge transfer, or a degree of charge compensation by a small parasitic I phase in the CsI samples (apparent in Figure 6a). A clearer view of the influence of CsI filling upon the optoelectronic properties of the SWCNTs was obtained from PL spectra (Figure 6c) of dispersed tubules in solution. Under optical excitation at 650 nm, emission from (8,3), (7,5), and (7,6) chiralities was visible while 575 nm excitation probed (6,5) and (8,4) tubules.^{39,40} Vertical solid lines indicate demarcation between "linear" -I-Cs-I-Cs-Iand 2×1 CsI filling and dashed lines indicate resonance wavelengths of unfilled SWCNTs. For (6,5) SWCNTs filled with I-Cs-I-Cs-I-- chains the E^S₁₁ emission was red-shifted with respect to the unfilled reference. The (7,5), (7,6), and (8,4)chiralities instead display blue-shifts when filled by 2×1 helical CsI. E^s₁₁ emission was efficiently suppressed for all the iodinefilled (n,m) tubules at both excitation wavelengths as a result of their *p*-type doping blocking radiative emission by phase space filling.

Figure 6d shows the full 2D photo-luminescence (PL) excitation for empty SWCNTs and CsI-filled SWCNTs, with the

dashed lines indicating the profiles used in Figure 6c (these plots are reproduced in Figure S15 with the 2D plot for I@SWCNT added for comparison). Solid curves in the first two plots indicate onsets for linear -I-Cs-I-Cs-, 2 × 1 and 2 × 2 CsI formation with widening SWCNT diameter. The PL plots for pristine SWCNTs indicate the predominating (9,4) and (7,6) semiconducting SWCNTs. After CsI filling, the distribution was dominated instead by -I-Cs-I-Cs- filled (6,5) SWCNTs, with relatively weaker emission from (7,5), (9,4), (7,6), and (8,4) 2 × 1-filled SWCNTs. Relatively weaker emission from the larger diameter 2 × 1-filled SWCNTs may result partially from small *n*-type doping. However, the oval distortion created by the helical 2×1 CsI filling could also disrupt the radiative emission from excitons, either by changing the intrinsic properties of the SWCNTs (e.g., electronic band structure, exciton binding energy) or by introducing disorder (e.g., fluctuations in tube diameters) on length scales shorter than the exciton diffusion length (typically >100 nm).



Figure 6. (a) Resonant RBMs of pristine SWCNTs, iodine, and CsI-filled SWCNTs at 532 nm laser excitation (660 nm excitation, Supporting Figure S14(a)). Raman shifts at 250, 266, and 275 cm⁻¹ correspond to (9,4), (7,6), and (8,4) SWCNTs, respectively. Resonances due to polymeric iodine chains $(I_3^-, I_5^-, or \text{ continuous } I_x)$ in the iodine and CsI-filled SWCNTs and an enhanced SWCNT resonance (denoted $(12,5)^E$) for the CsI-filled sample are visible between 100 and 200 cm⁻¹. (b) Optical absorption spectra, normalized to the π -plasmon of empty SWCNTs, of empty SWCNTs, I_x and CsI filled SWCNTS with (n,m) peak assignments.⁴⁰ Vertical lines indicate shifts in the peaks due to filling. (c), 900 to 1300 nm region from (b) showing E_{11}^s shifts with (n,m) peaks indicated. Nearly all are suppressed for iodine-filled SWCNTs with only (6,5) nanotubes having a significant absorbance and blue shift. CsI-filled (7,5) and (7,6) SWCNTs show prominent blue shifts, whereas CsI-filled (6,5) SWCNTs are unshifted. (d) 2D PL emission contour plots of empty, CsI and iodine-filled SWCNTs (left to right). The CsI-filled SWCNTs retain strong (6,5) and blue-shifted (7,5) and (7,6) emissions, respectively, with nearly all E_{11}^s emissions for iodine-filled SWCNTs being quenched (see also Supporting Figure S15).

CONCLUSIONS

In this work, we reveal in detail the structural transformation of "True 1D" CsI atomic chains in narrow SWCNT with diameters from 0.73 to 0.81 nm to 2×1 CsI atomic chains in ovally distorting ~0.84-0.95 nm diameter SWCNTs. For encapsulated 2×1 CsI helical chains, electron imaging reveals regulation of I-Cs-I angles by the SWCNT internal cross section and spectroscopic imaging confirms the chemical identify of linear -I-Cs-I-Cs- chains versus 2 \times 1 CsI helices, the latter forming in oval cross section SWCNTs. Raman, OAS, and PL optical measurements reveal the highly selective nature of doping that these ionic CsI chains produce in different narrow SWCNTs with (6,5), (7,5), and (7,6) all exhibit markedly different behavior. Our computational studies reveal the energetics of 2×1 CsI formation, and the resulting helical structure is found to be stable in the range ~0.88 to 1.08 nm but with I-Cs-I strain increasing as the SWCNT cross section reduces. These will be proportionally different for comparable alkali halide and other binary ionic systems but should

demonstrate generality for linear or helical crystal growth inside smooth narrow tubules such as carbon nanotubes. Annealing helical structures with a specific (n,m) conformation should also generate helices of constant pitch as their formation energy will reach a minimum for a specific diameter. These extended defects differ from the Schottky and Frenkel type normally found in alkali halides⁵¹ with longer range defects normally being stochastically induced by radiation or electron beam damage in thin halide films.⁵² By encapsulating, annealing and refining the diameter distribution of SWCNTs, we now have a genuine prospect of controlling the formation of helical chains in CsI and related materials with interesting consequences for encapsulate and composite physical properties as a result.

METHODS

Preparation of CsI- and I_x-Filled SWCNTs for AC-TEM and AC-ADF-STEM. Bulk CsI (Sigma-Aldrich, 99.999%) was vapor phase transported and deposited inside CoMoCAT SWeNT SWCNTs, originally supplied by SouthWest NanoTechnologies, with a specified diameter distribution of 0.7–1.1 nm (Product No. 404121). Filling of

these nanotubes was effected *via* the sublimation method using similar conditions to those described elsewhere.^{13,18,21} Twenty milligrams of SWCNTs were first oxidized in open air at 750 K before being placed in one side of a 20 cm long two compartment silica quartz ampule with 50 mg CsI being placed at the end in the center of the hot zone of a Carbolite tube furnace. The hot end was heated to 825 K and the CsI was allowed to sublime along the natural temperature gradient created by the single hot zone of the furnace for 7 days into the preopened CoMoCAT SWCNTs maintained in the second compartment.

Electron Microscopy Imaging and Simulation. A doublecorrected JEM-ARM 200F microscope operating at 80 kV equipped with CEOS imaging aberration and probe correction and a Gatan SC1000 ORIUS camera with a 4008 \times 2672 pixel CCD was used for preliminary HRTEM and ADF STEM investigations. Electron beam densities were adjusted to between 0.8 to 1.88 pÅ/cm². High-resolution high angle annular dark field (HAADF) imaging and electron energy loss spectroscopy (EELS) were carried out in a Nion UltraSTEM100 dedicated scanning transmission electron microscope (STEM), equipped with a Gatan Enfina spectrometer. The instrument was operated at 60 kV acceleration voltage to minimize knock-on damage to the SWCNTs, and the STEM probe forming optics were adjusted to provide a 20 pA beam with 33 mrad convergence angle-resulting in an estimated probe size of approximately 1 Å. The EEL spectrometer entrance aperture half an gle acceptance was 36 mrad, while the HAADF detector inner and outer angles were 85 and 190 mrad, respectively. Chemical maps were produced from acquired EELS data by integrating the intensity of the relevant ionization edges (Cs and I M_{45}) over a 75 eV window above their respective onsets, after subtraction of the decaying background using a standard power law model. A spectral dispersion of 0.5 eV/channel was used to capture both edges simultaneously (resulting in a point-spread limited effective energy resolution of $\sim 1 \text{ eV}$; to further boost signal-to-noise the data was binned (\times 2) spectrally prior to generating the EELS maps, but no statistical denoising was applied.

DFT and Bader Analysis Technical Details. All of the calculations are conducted with CASTEP-16.0 computational suite³⁶ exploiting DFT with a plane-wave basis approach. Exchange and correlation was considered within a generalized gradient approximation in the form of the Perdew-Burke-Ernzerhof (PBE) functional.³⁷ The core electrons were replaced by ultrasoft pseudopotentials while the electronic wave functions were expanded with a converged value for cutoff energy at 600 eV. All our models contain at least 12 Å vacuum layers between periodic images that exclude spurious interactions. The Brillouin zone was integrated on a $1 \times 1 \times 11$ points of the Monkhorst and Pack grid. Geometry was optimized until forces on the atoms were smaller than 0.001 eV/Å. The calculated charge was then processed with a grid-based Bader analysis.³⁸ To account for possible effects of stress arising from incommensurate periodic models of CsI atomic chains and host CNTs, we include a stress compensation term in assessment of formation energy, which we find via calculations within "implicit nanotubes" where the repulsive electrostatic potential mimics steric confinement with CNTs.¹

Raman, Optical Absorption, and Photoluminescence Spectroscopies. Optical absorbance spectroscopy was performed using a Lambda 1050 UV/vis/IR spectrophotometer with a spectral range of 250–1500 nm. Raman spectra were obtained using confocal microspectrometers (Labram, Jobin–Yvon Horiba, and Renishaw inVia Reflex) with 660 and 532 nm excitation, a spectral resolution of 0.1 and 2 cm⁻¹, and a spatial resolution of 1.5 μ m. Photoluminescence spectra were measured by a Fluorolog UV/vis/IR fluorescence spectrophotometer (Horiba Jobin Yvon).

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsnano.1c03705.

Information about image straightening procedures to used facilitate helical chain measurements; additional images and FFTs of CsI filling in SWCNT bundles; *ab* *initio* modeling of CsI wires and SWCNT interactions including by DFT (CASTEP), force-field fitting by PotFit, PBE-DFT, SWCNT-ion interactions, LAMMPS simulations, Bader analysis; additional Raman measurements and additional PL data (PDF)

Movie of unsmooth motion of linear CsI chain in straight SWCNT (AVI)

Movie of unsmooth motion of linear CsI chain in curved SWCNT (AVI)

Movie of static linear CsI chain in continuosly filled SWCNT (AVI)

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Author Contributions

J.S., R.J.K., M.B., A.V., P.V. C.M., A.J.M., D.Q., Q.R., and J.L.-H. conceived the project. R.J.K. performed the sample preparation and aberration corrected transmission electron microscopy characterization of filled nanotube samples with further imaging and analysis being carried out by J.S. and HAADF imaging and EELS being performed by Q.R. in Daresbury. Optical characterization of the samples was performed by M.B. and J.L.-H. Theoretical studies were performed by A.V., P.V.C.M., A.J.M., and D.Q. in Warwick and Birmingham. J.S., R.J.K., M.B., A.V., D.Q., and J.L.-H. wrote the manuscript and J.S. prepared the figures, including additional image processing, image simulation,

and structure modeling with contributions from A.V., D.Q., R.J.K, Q.R., MB., and J.L-H. All authors discussed the results and commented on the manuscript.

Notes

The authors declare no competing financial interest. Data used in this work are available via the Warwick data repository at wrap.warwick.ac.uk/156210.

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REFERENCES

(1) Eigler, D. M.; Schweizer, E. K. Positioning Single Atoms with a Scanning Tunneling Microscope. *Nature* **1990**, *344*, 524–526.

(2) Crommie, M. F.; Lutz, C. P.; Eigler, D. M. Confinement of Electrons to Quantum Corrals on a Metal Surface. *Science* **1993**, *262*, 218–220.

(3) Yeom, H. W.; Takeda, S.; Rotenberg, E.; Matsuda, I.; Horikoshi, K.; Schaefer, J.; Lee, C. M.; Kevan, S. D.; Ohta, T.; Nagao, T.; Hasegawa, S. Instability and Charge Density Wave of Metallic Quantum Chains on a Silicon Surface. *Phys. Rev. Lett.* **1999**, *82*, 4898–4901.

(4) Nadj-Perge, S.; Drozdov, I. K.; Li, J.; Chen, H.; Jeon, J.; MacDonald, A. H.; Bernevig, B. A. Observation of Majorana Fermions in Ferromagnetic Atomic Chains on a Superconductor. *Science* **2014**, *346*, 602–607.

(5) Lin, J.; Cretu, O.; Zhou, W.; Suenaga, K.; Prasai, D.; Bolotin, K. I.; Cuong, N. T.; Otani, Minoru; Okada, S.; Lupini, A. R.; Idrobo, J.-C.; Caudel, D.; Burger, A.; Ghimire, N. J.; Yan, J.; Mandrus, D. G.; Pennycook, S. J.; Pantelides, S. T. Metallic Nanowires with Self-Adaptive Contacts to Semiconducting Transition-Metal Dichalcogenide Monolayers. *Nat. Nanotechnol.* **2014**, *9*, 436–442.

(6) Lin, J.; Zhang, Y.; Zhou, W.; Pantelides, T. Structural Flexibility and Alloying in Ultrathin Transition-Metal Chalcogen Nanowires. *ACS Nano* **2016**, *10*, 2782–2790.

(7) Koh, A. L.; Wang, S.; Ataca, C.; Grossman, J. C.; Sinclair, R.; Warner, J. H. Torsional Deformations in Subnanometer MoS Interconnecting Wires. *Nano Lett.* **2016**, *16*, 1210–1217.

(8) Nagata, M.; Shukla, S.; Nakanishi, Y.; Liu, Z.; Lin, Y.-C.; Shiga, T.; Nakamura, Y.; Koyama, T.; Kishida, H.; Inoue, T.; Kanda, N.; Ohno, S.; Sakagawa, Y.; Suenaga, K.; Shinohara, H. Isolation of Single-Wired Transition-Metal Monochalcogenides by Carbon Nanotubes. *Nano Lett.* **2019**, *19*, 4845–4851.

(9) Sloan, J.; Kirkland, A. I.; Hutchison, J. L.; Green, M. L. H. Integral Atomic Architectures of 1D Crystals Inserted into Single Walled Carbon Nanotubes. *Chem. Commun.* **2002**, 1319–1332.

(10) Fan, X.; Dickey, E. C.; Eklund, P. C.; Williams, K. A.; Grigorian, L.; Buczko, R.; Pantelides, S. T.; Pennycook, S. J. Atomic Arrangement of Iodine Atoms inside Single-Walled Carbon Nanotubes. *Phys. Rev. Lett.* **2000**, *84*, 4621–4624.

(11) Sloan, J.; Kirkland, A. I.; Hutchison, J. L.; Green, M. L. H. Aspects of Crystal Growth within Carbon Nanotubes. *C. R. Phys.* 2003, *4*, 1063–1074.

(12) Fujimori, T.; Morelos-Gómez, A.; Zhu, Z.; Muramatsu, H.;
Futamura, R.; Urita, K.; Terrones, M.; Hayashi, T.; Endo, M.; Hong, S.
Y.; Choi, Y. C.; Tománek, D.; Kaneko, K. Conducting Linear Chains of Sulphur inside Carbon Nanotubes. *Nat. Commun.* 2013, *4*, 2162/1–8.
(13) Senga, R.; Komsa, H.-P.; Liu, Z.; Hirose-Takai, K.;
Krasheninnikov, A. V.; Suenaga, K. Atomic Structure and Dynamic Behaviour of Truly One-Dimensional Ionic Chains inside Carbon Nanotubes. *Nat. Mater.* 2014, *13*, 1050–1054.

(14) Guan, L.; Suenaga, K.; Shi, Z.; Iijima, S. Polymorphic Structures of Iodine and Their Phase Transformation in Confined Nanospace. *Nano Lett.* **2007**, *7*, 1532–1535.

(15) Komsa, H.-P.; Senga, R.; Suenaga, K.; Krasheninnikov, A. V. Structural Distortions and Charge Density Waves in Iodine Chains Encapsulated inside Carbon Nanotubes. *Nano Lett.* **2017**, *17*, 3694–3700.

(16) Nie, C. Y.; Gabilert, A.-M.; Soula, B.; Datas, L.; Sloan, J.; Flahaut, E.; Monthioux, M. The Unexpected Complexity of Filling Double-Wall Carbon Nanotubes with Nickel (and Iodine) 1-D Nanocrystals. *IEEE Trans. Nanotechnol.* **2017**, *16*, 759–766.

(17) Shi, L.; Rohringer, P.; Suenaga, K.; Niimi, Y.; Kotakoski, J.; Meyer, J. C.; Peterlik, H.; Wanko, M.; Cahangirov, S.; Rubio, A.; Lapin, Z. J.; Novotny, L.; Ayala, P.; Pichler, T. Confined Linear Carbon Chains as a Route to Bulk Carbyne. *Nat. Mater.* **2016**, *15*, 634–639.

(18) Medeiros, P. V. C.; Marks, S.; Wynn, J. M.; Vasylenko, A.; Ramasse, Q. M.; Quigley, D.; Sloan, J.; Morris, A. J. Single-Atom Scale Structural Selectivity in Te Nanowires Encapsulated inside Ultranarrow Single-Walled Carbon Nanotubes. *ACS Nano* **2017**, *11*, 6178–6185.

(19) Hart, M.; White, E. R.; Chen, J.; McGilvery, C. M.; Pickard, C. J.; Michaelides, A.; Sella, A.; Shaffer, M. S. P.; Salzmann, C. G. Enapsulation and Polymerization of White Phosphorous inside Single-Walled Carbon Nanotubes. *Angew. Chem., Int. Ed.* **201**7, *56*, 8144–8148.

(20) Hart, M.; Chen, J.; Michaelides, A.; Sella, A.; Shaffer, M. S. P.; Salzmann, C. G. One-Dimensional Arsenic Allotropes: Polymerization of Yellow Arsenic inside Single-Wall Carbon Nanotubes. *Angew. Chem., Int. Ed.* **2018**, *57*, 11649–11653.

(21) Vasylenko, A.; Marks, S.; Wynn, J. M.; Medeiros, P. V. C.; Ramasse, Q. M.; Morris, A. J.; Sloan, J.; Quigley, D. Electronic Structure Design of Sub-Nanometre 1D SnTe *via* Nanostructuring within Single-Walled Carbon Nanotubes. *ACS Nano* **2018**, *12*, 6023–6031.

(22) Slade, C. A.; Sanchez, A. M.; Sloan, J. Unprecedented New Crystalline Forms of SnSe in Narrow to Medium Diameter Carbon Nanotubes. *Nano Lett.* **2019**, *19*, 2979–2984.

(23) Senga, R.; Suenaga, K. Single-Atom Detection of Light Elements: Imaging or Spectroscopy ? *Ultramicroscopy* **2017**, *180*, 150–155.

(24) Grigorian, L.; Williams, K. A.; Fang, S.; Sumanasekera, G. U.; Loper, A. L.; Dickey, E. C.; Pennycook, S. J.; Eklund, P. C. Reversible Intercalation of Charged Iodine Chains into Carbon Nanotube Ropes. *Phys. Rev. Lett.* **1998**, *80*, 5560–5563.

(25) Chorro, M.; Kané, G.; Alvarez, L.; Cambedouzou, J.; Paineaud, E.; Rossberge, A.; Kociak, M.; Aznar, R.; Pascarellia, S.; Launois, P.; Bantignies, J. L. 1D Confinements of Polyiodides inside Single-Walled Carbon Nanotubes. *Carbon* **2013**, *52*, 100–108.

(26) Tonkikh, A. A.; Obraztsova, E. D.; Obraztsova, E. A.; Belkin, A. V.; Pozharov, A. S. Optical Spectroscopy of Iodine-Doped Single-Wall Carbon Nanotubes of Different Diameter. *Phys. Status Solidi B* **2012**, *B249*, 2454–2459.

(27) Poborchii, V. V. Polarized Raman and Optical Absorption Spectra of the Mordenite Single Crystals Containing Sulfur, Selenium or Tellurium in the One-Dimensional Nanochannels. *Chem. Phys. Lett.* **1996**, *251*, 230–234.

(28) Poborchii, V. V.; Kolobov, A. V.; Caro, J.; Zhuravlev, V. V.; Tanaka, K. Dynamics of Single Selenium Chains Confined in One-Dimensional Nanochannels of AlPO₄-5: Temperature Dependencies of the First- and Second-Order Raman Spectra. *Phys. Rev. Lett.* **1999**, *82*, 1955–1958.

(29) Ivanov, V. G.; Kalashnyk, N.; Sloan, J.; Faulques, E. Vibrational Dynamics of Extreme 2×2 and 3×3 Potassium Iodide Nanowires Encapsulated in Single-Walled Carbon Nanotubes. *Phys. Rev. B: Condens. Matter Mater. Phys.* **2018**, *98*, 125429.

(30) Spencer, J. H.; Nesbitt, J. M.; Trewhitt, H.; Kashtiban, R. J.; Bell, G.; Ivanov, V. G.; Faulques, E.; Sloan, J.; Smith, D. C. Raman Spectroscopy of Optical Transitions and Vibrational Energies of ~ 1 nm HgTe Extreme Nanowires within Single Walled Carbon Nanotubes. *ACS Nano* **2014**, *8*, 9044–9052.

(31) Ilie, A.; Bendall, J. S.; Roy, D.; Philp, E.; Green, M. L. H. Effects of KI Encapsulation in Single-Walled Carbon Nanotubes by Raman and

Optical Absorption Spectroscopy. J. Phys. Chem. B 2006, 110, 13848–13857.

(32) Vasylenko, A.; Wynn, J.; Medeiros, P. V. C.; Morris, A. J.; Sloan, J.; Quigley, D. Encapsulated Nanowires: Boosting Electron Transport in Carbon Nanotubes. *Phys. Rev. B: Condens. Matter Mater. Phys.* **2017**, *95*, 121408.

(33) Kharlamova, M. V.; Kramberger, C.; Rudatis, P.; Pichler, T.; Eder, D. Revealing the Doping Effect of Encapsulated Lead Halogenides on Single-Walled Carbon Nanotubes. *Appl. Phys. A: Mater. Sci. Process.* **2019**, *125*, 320.

(34) Philp, E.; Sloan, J.; Kirkland, A. I.; Meyer, R. R.; Friedrichs, S.; Hutchison, J. L.; Green, M. L. H. An Encapsulated Helical 1D Cobalt Iodide Crystal. *Nat. Mater.* **2003**, *2*, 788–791.

(35) Hirahara, K.; Suenaga, K.; Bandow, S.; Kato, H.; Okazaki, T.; Shinohara, H.; Iijima, S. One-Dimensional Metallofullerene Crystal Generated inside Single-Walled Carbon Nanotubes. *Phys. Rev. Lett.* **2000**, *85*, 5384–5387.

(36) Clark, S. J.; Segall, M. D.; Pickard, C. J.; Hasnip, P. J.; Probert, M. J.; Refson, K.; Payne, M. C. First Principles Methods Using CASTEP. *Z. Kristallogr. - Cryst. Mater.* **2005**, *220*, 567–570.

(37) Perdew, J. P.; Burke, K.; Ernzerhof, M. Generalized Gradient Approximation Made Simple. *Phys. Rev. Lett.* **1996**, *77*, 3865–3868.

(38) Tang, W.; Sanville, W. E.; Henkelman, G. A Grid-Based Bader Analysis Algorithm without Lattice Bias. J. Phys.: Condens. Matter 2009, 21, 084204.

(39) Christian, T.; Reich, S. In Raman Scattering in Carbon Nanotubes" in *Light Scattering in Solids IX*; Cardona, M., Merlin, R., Eds.; Springer: Berlin, Heidelberg, 2007; pp115–234.

(40) Weisman, R. B.; Bachilo, S. M. Dependence of Optical Transition Energies on Structure for Single-Walled Carbon Nanotubes in Aqueous Suspension: An Empirical Kataura Plot. *Nano Lett.* **2003**, *3*, 1235– 1238.

(41) Ilie, A.; Bendall, J. S.; Roy, D.; Philp, E.; Green, M. L. H. Effects of KI Encapsulation in Single-Walled Carbon Nanotubes by Raman and Optical Absorption Spectroscopy. *J. Phys. Chem. B* **2006**, *110*, 13848–13857.

(42) Mann, J. B. Atomic Structure Calculations II. Hartree-Fock Wave Functions and Radial Expectation Values: Hydrogen to Lawrencium, LA-3691; Los Alamos Scientific Laboratory, USA, 1968.

(43) Bichoutskaia, E.; Pyper, C. Theoretical Study of the Structures and Electronic Properties of All-Surface KI and CsI Nanocrystals Encapsulated in Single Walled Carbon Nanotubes. *J. Chem. Phys.* **2008**, *129*, 154701.

(44) Shiraishi, M.; Ata, M. Work Function of Carbon Nanotubes. *Carbon* **2001**, *39*, 1913–1917.

(45) Yamamoto, H. Ultraviolet Photoelectron Spectroscopy of Solid Iodine. J. Chem. Phys. **1987**, 86, 1775–1779.

(46) Verma, R. L. Potential and Limitations of Cesium Iodide as a Dynode Material for Use in Electron Multipliers. *J. Phys. D: Appl. Phys.* **1978**, *11*, 63–71.

(47) Avery, N. R. LEED, Auger and Work Function Study of Iodine Adsorbed on W (100). *Surf. Sci.* **1974**, *43*, 101–122.

(48) Fairchild, S. B.; Back, T. C.; Murray, P. T.; Cahay, M. M.; Shiffler, D. A. Low Work Function CsI Coatings for Enhanced Field Emission Properties. *J. Vac. Sci. Technol., A* **2011**, *29*, 031402.

(49) Kazaoui, S.; Minami, N.; Jacquemin, R.; Kataura, H.; Achiba, Y. Amphoteric Doping of Single-Wall Carbon-Nanotube Thin Films as Probed by Optical Absorption Spectroscopy. *Phys. Rev. B: Condens. Matter Mater. Phys.* **1999**, *60*, 13339–13342.

(50) Burdanova, M. G.; Tsapenko, A. P.; Satco, D. A.; Kashtiban, R.; Mosley, C. R.; Monti, M.; Staniforth, M.; Sloan, J.; Gladush, Y. G.; Nasibulin, A. G.; Lloyd-Hughes, J. Giant Negative Terahertz Photoconductivity in Controllably Doped Carbon Nanotube Networks. *ACS Photonics* **2019**, *6*, 1058–1066.

(51) Royce, B. S. H. The Creation of Point Defects in Alkali Halides. *Prog. Solid State Chem.* **1967**, *4*, 213–243.

(52) Catlow, C. R. A.; Diller, K. M.; Hobbs, L. W. Irradiation-Induced Defects in Alkali Halide Crystals. *Philos. Mag. A* **1980**, *42*, 123–150.