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Structural and chemical mechanisms governing stability of inorganic Janus nanotubes

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One-dimensional inorganic nanotubes hold promise for technological applications due to their distinct physical/chemical properties, but so far advancements have been hampered by difficulties in producing single-wall nanotubes with a well-defined radius. In this work we investigate, based on Density Functional Theory (DFT), the formation mechanism of 135 different inorganic nanotubes formed by the intrinsic self-rolling driving force found in asymmetric 2D Janus sheets. We show that for isovalent Janus sheets, the lattice mismatch between inner and outer atomic layers is the driving force behind the nanotube formation, while in the non-isovalent case it is governed by the difference in chemical bond strength of the inner and outer layer leading to steric effects. From our pool of candidate structures we have identified more than 100 tubes with a preferred radius below 35 Å, which we hypothesize can display distinctive properties compared to their parent 2D monolayers. Simple descriptors have been identified to accelerate the discovery of small-radius tubes and a Bayesian regression approach has been implemented to assess the uncertainty in our predictions on the radius.

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INTRODUCTION

In the last decades miniaturization of devices has been a main trend driving the electronics industry. In addition to reducing the usage of raw materials, nanomaterials often show improved properties compared to their larger counterparts. Among these nanomaterials are two-dimensional (2D) sheets, one-dimensional (1D) structures such as nanotubes and nanoribbons, and zero-dimensional (0D) nanoparticles.

Since their discovery, nanotubes have shown promise for a wide range of applications including gas separation and capture, catalysis, solid lubrication and controlled drug delivery. In addition to the well-known carbon nanotubes numerous inorganic nanotubes have been synthesized. Although the first successful synthesis of single-wall MoS₂ nanotubes has been reported, such structures usually appear together with numerous multi-wall tubes showing a distribution of radii and wall thicknesses. These multi-wall structures alleviate the built-in strain energy through van der Waals interactions in between the layers leading to an increase in stability. Overall, this has made it difficult to establish an experimental synthesis pathway to produce single-wall tubes with a specific radius and controllable physio-chemical properties.

A possible solution to this problem is the approach of considering asymmetric sheets, which can naturally wrap and form nanotubes with a well-defined size. Due to the asymmetry, the unsupported sheet is expected to be unstable compared to other curled shapes, such as tubes or scrolls. Pauling, already in the 1930s, mentioned that the driving force of sheets to curve is related to the lattice-mismatch between the two inner and outer atomic layers. Single-wall inorganic nanotubes with well-defined diameters hold promise for technological applications, not only because of their reduced dimensionality, but also for their distinctive properties, often inherently different from the ones of the corresponding asymmetric sheets. An example of a small-radius, single-wall nanotube formed from an asymmetric sheet is imogolite (Al₂SiO₄(OH)₄), which was first discovered in volcanic ash soil and later synthesized. Other tubular minerals include chrysotile (Mg₃Si₂O₅(OH)₄) and halloysite (Al₂Si₂O₅(OH)₄) that however occur as multi-wall tubes. Besides naturally occurring nanotubes, misfit layered compounds, composed of two separate sheets, make use of the lattice mismatch between the two sheets to induce a natural driving force to form a tube.

One of the possible classes of materials forming asymmetric 2D monolayers are Janus sheets, like MoSSe or BiTeI, which can be wrapped to form 1D tubes. A recent work has shown that radii well below 35 Å are needed to create single-wall Janus transition metal dichalcogenide (TMD) tubes, which have significantly different (electronic) properties from the corresponding asymmetric sheet. Although facile synthesis routes for the production of single-wall inorganic nanotubes has long been actively researched, not much attention has been paid to the question of which materials would be able to make such a structure avoiding the creation of multi-wall tubes. Consequently, a high throughput study on the stability of a wide range of Janus-based nanotubes, would provide valuable information for guiding future synthesis of small-radius single-wall nanotubes.

In this work, we present a comprehensive screening study in the framework of Density Functional Theory (DFT) on the stability of 135 different inorganic nanotubes generated from the rolling of asymmetric 2D Janus sheets along both the armchair and zigzag directions. The calculations focus on the stability and strain energy of the chosen nanotubes. The total number of DFT relaxations performed in this work amounts to ~4500. We show that for pure chalcogen or halogen tubes (isovalent anions), the wrapping mechanism is mostly governed by the lattice-mismatch between the two inner and outer atomic layers, while for mixing anions (non-isovalent anions) this is dominated by the difference in valency between the X/Y elements. These findings provide a physical foundation for designing Janus nanotubes with optimal (small) radii.

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RESULTS

Initial pool of materials

The nanotubes considered in this study consist of three layers (MXY, as illustrated in Fig. 1) composed of different mid-layer elements (M = (Ti, Zr, Hf, V, Nb, Ta, Cr, Mo, W, Fe, Ge, Sn, As, Sb, Bi)) decorated with inner X and outer Y chalcogen and halogen atoms (X,Y = (O, S, Se, Te, Cl, Br, I)). Here we denote all group 16 elements including oxygen as chalcogens. For the three pnictogens (As, Sb, and Bi) having 3+ as one of their possible oxidation states, we mix chalcogens and halogens in the structures. For the remaining 12 elements, the inner X and outer Y elements are either chalcogens or halogens. The idea of mixing chalcogens and halogens to form 2D MXY Janus sheets was recently explored in ref.17, but has so far not been pursued in the context of nanotubes. We construct the tubes by rolling up 2D layers in both the T- and H-phase crystal structures, corresponding to the crystal structures found for the experimentally synthesized MoSSe13 and BiTeI14 2D sheets, along both the armchair and zigzag directions.

Nanotube strain energy

Two main quantities that are needed to characterize an asymmetric nanotube are the optimal radius, which defines the most stable nanotube size, and the strain energy, which defines the energy associated with the wrapping of a 2D sheet into a nanotube (negative strain energies indicate a spontaneous wrapping).

The strain energy is defined as the difference between the energy of the nanotube and the corresponding 2D sheet18. In formula:

\[ E_{\text{strain}}(R) = \frac{E_{\text{tube}}}{N_{\text{tube}}} - \frac{E_{\text{MXY}}}{N_{\text{MXY}}}, \]

where \( E_{\text{tube}} \) is the energy of a nanotube with \( N_{\text{tube}} \) atoms and \( E_{\text{MXY}} \) is the energy of the corresponding 2D Janus sheet with \( N_{\text{MXY}} \) atoms in the unit cell. In the infinite limit \( R \to \infty \), the strain energy is zero, since the energy per atom of a tube is equal to the energy per atom of an infinite 2D Janus sheet.

It has been shown that for symmetric tubes (carbon, for example) the nanotube strain energy follows a \( 1/R^2 \) dependence19,20. This relationship does not hold for asymmetric tubes in which the strain energy curve exhibits a minimum. Instead, it can be more accurately described using the equation21:

\[ E_{\text{strain}}(R) = \frac{a}{R^2} + \frac{b}{R}. \]

Extrapolating the function using the obtained DFT data and evaluating the function at the minimum strain energy \( E_{\text{strain-min}} \) leads to the optimal tube radius \( R_{\text{opt}} \) (see also Fig. 2). We note that, although Eq. (2) fits well in the region around the optimal tube radius, large strain energies can lead to a deviation25. We take this into consideration during the screening using Bayesian statistics (details in the Supplementary Note 3) which helps to identify cases where the function chosen does not capture the observed data points well across all tube radii.

As an example, Fig. 2 shows the strain energy as a function of the radius for the symmetric MoS2 nanotube and the two asymmetric NbSSe and BiSI nanotubes. The label in brackets indicates the prototype (H-/T-phase) and the wrapping direction (a/z indicating armchair/zigzag).

![Fig. 1 Basic principle of the computational workflow. The symmetric MY2 and MX2 are the parent sheets for the asymmetric 2D Janus MXY sheet, which can be wrapped up to form a 1D Janus nanotube. In this case M = Bi (light purple), X = Te (dark yellow), and Y = I (dark purple).](image)

![Fig. 2 Strain energy as a function of the nanotube radius. Strain energy as a function of the radius for the symmetric MoS2 nanotube and the two asymmetric NbSSe and BiSI nanotubes. The label in brackets indicates the prototype (H-/T-phase) and the wrapping direction (a/z indicating armchair/zigzag).](image)
Bayesian regression. In general, the values for the optimal amount of data points lead to an extended uncertainty interval. Additional data points are not necessary. However, the limited sufficient to identify the mid-point as the minimum and adding uncertainty on the radius is estimated to be larger than 30 Å. It is ensured that at least three data points after filtering according to the criteria described in the Supplementary Note 1 exist (otherwise marked with a hatched box). The asterisk indicates that the difference in energy between the H-/T-phase is <10 meV/atom and that the smaller optimal tube radius is chosen given the extrapolated optimal radius from both prototypes. The lowercase letters a/z indicate which wrapping direction (armchair/zigzag) is preferred.

Stability and optimal radius

Figure 3 shows the optimal tube radii for all studied materials and their extrapolated optimal radius according to Eq. (2). It is ensured that at least three data points after filtering according to the criteria described in the Supplementary Note 1 exist (otherwise marked with a hatched box). The asterisk indicates that the difference in energy between the H-/T-phase is <10 meV/atom and that the smaller optimal tube radius is chosen given the extrapolated optimal radius from both prototypes. The lowercase letters a/z indicate which wrapping direction (armchair/zigzag) is preferred.

The calculated values for the convex hull energy of the tube at the optimal tube radius. Blue and bright colors indicate good stability, while a red color indicates poor stability. Values exceeding the metastability criterion of 0.2 eV/atom are shown in black.

Giving the mid-layer element is in its preferred oxidation state, the convex hull used here can be found in the literature. To compare the energy of the tube to its most stable 3D bulk structures, we calculate the convex hull energy of the tube for at least one of the calculated combinations (here we define a combination stable when the energy of the candidate compound is within 0.2 eV/atom above the convex hull to account for a possible metastability). Given the mid-layer element is in its preferred oxidation state, the resulting tube shows higher stability compared to the case of an
unfavored oxidation state as expected. Ge, for example, generates more stable tubes when combined with two halogens (Ge_2^{2+}) than with two chalcogens (which would need a Ge_4^{4+}, instead). We do not find stable nanotubes for 23 combinations (hatched boxes in Fig. 4). 18 of these materials can be attributed to the transition metal in its unfavored 2^{2+} oxidation state, when paired with two halogens. The remaining 5 materials contain either Fe, Sn or Ge which form in general less stable nanotubes for most of the studied combinations (i.e., more than 0.1 eV/atom above the convex hull). Because of their stability in the 3D form, oxygen-containing tubes are in general more prone to decompose compared to the pure chalcogenide ones.

For almost 90% of the materials, the energy difference between the armchair and zigzag wrapping direction is below 10 meV/atom, which indicates that there is only a weak driving force causing wrapping up around a specific direction. Although the armchair and zigzag wrapping directions only have a minor impact on the stability of the tube, it can be expected that the wrapping direction has a larger impact on the electronic properties due to the difference in bond distances in these tube configurations. We have recently demonstrated this for MoSTe, but a more comprehensive study focusing specifically on the electronic properties would be needed to establish general design rules.

The combination of the three metals As, Sb, and Bi mixed together with a chalcogen sitting inside and a halogen element sitting outside of the structure generates stable and small-radius nanotubes with a rather small strain energy associated with its optimal radius. For instance, AsSI has a minimum strain energy of −80 meV/atom which is ~40 meV smaller than that of the experimentally observed imogolite nanotube when compared to computational reference data found in literature. The MXY nanotubes share with imogolite, which is known to exist as a single-wall nanotube, the shape of the strain energy curve (see BiSI in Supplementary Fig. 2).

Figure 5 reports the correlation between various quantities, namely the extrapolated optimal radius $R_{\text{opt}}$ versus the lattice-mismatch of the corresponding parent sheets (lattice mismatch calculated as $a_{MX_2}/a_{MY_2}$ with $a$ being the lattice constant) (a), the minimum strain energy at the optimal radius (b) and the Mulliken ratio of the corresponding Mulliken electronegativities of outer and inner elements (Mulliken ratio X/Y) (c) are shown. The ratio in layer thickesses in the 2D Janus sheet versus the lattice-mismatch. The different colors represent the three classes of pairing the mid-layer with either two chalcogens (Janus-Cal-Cal, {O, S, Se, Te}, green), two halogens (Janus-Hal-Hal, {Br, I, Cl}, red) or a mix of chalcogens inside and halogens outside (Janus-Cal-Hal, yellow).

**Fig. 5** Property correlations derived from nanotubes and sheets. $R_{\text{opt}}$ versus the lattice mismatch of the corresponding parent sheets (lattice mismatch calculated as $a_{MX_2}/a_{MY_2}$ with $a$ being the lattice constant) (a), the minimum strain energy at the optimal radius (b) and the Mulliken ratio of the corresponding Mulliken electronegativities of outer and inner elements (Mulliken ratio X/Y) (c) are shown. The ratio in layer thickesses in the 2D Janus sheet versus the lattice-mismatch. The different colors represent the three classes of pairing the mid-layer with either two chalcogens (Janus-Cal-Cal, {O, S, Se, Te}, green), two halogens (Janus-Hal-Hal, {Br, I, Cl}, red) or a mix of chalcogens inside and halogens outside (Janus-Cal-Hal, yellow).
chalcogens (Janus-Cal-Cal class following trend OTe > OSe > OS > Ste > SeTe > SeS) or two halogens (Janus-Hal-Hal class following ClI > BrI > CIBr) with each other.

The lattice-mismatch between the two parent sheets can give an estimate on the optimal tube radius in the case of isovalent anions, while it fails for mixing halogens with chalcogens (Janus Cal-Hal class). Here, it would predict the halogen to sit inside of the tube, instead of on the outside, as several of the materials show a lattice mismatch $d_{MX}/d_{MY}$ larger than 1 (Fig. 5a, yellow dashed line). For instance, the MoO2 parent sheets (H-phase) have a lattice mismatch of 0.79 ($d_{MoO2} = 2.82 \text{ Å}$, $d_{MoTe2} = 3.55 \text{ Å}$) leading to the small radius of 7.2 Å. Conversely, the BiTeCl parent sheets (T-phase) have a lattice mismatch of 1.06 since the parent structure BiCl2 (a = 3.68 Å) has a smaller lattice constant than BiTe2 (a = 3.9 Å). Based on the lattice mismatch, the tube should wrap in a way that the Chlorine atoms are inside of the tube. The three different classes appear well-separated when the radius is plotted versus the ratio of the Mulliken electronegativities of the X and Y element (Fig. 5c). Having more electronegative elements sitting on the inside of the tube does not seem to be a necessary criterion for forming small-radius nanotubes (Janus-Cal-Hal, yellow).

A possible explanation on the reason why the chalcogen is sitting inside of the non-isovalent tube can be made by investigating the bond lengths in the non-isovalent structures closer. Shevelkov et al.33 studied the experimental 3D-bulk crystal structure of layered BiTeI and found the Bi–X bond distance in these structures to be significantly longer (ionic) than the ones found in bismuth trihalides. In addition, the geometry of the (BiTe) layer is shown to be comparable to the one found in Bi-bulk metal and bismuth trihalides. In these cases, the element forming stronger bonds sitting inside the non-isovalent tube can be made by.

![Fig. 6 Investigating inverse nanotube wrapping. Side (or cross-section) view for the energetically favored armchair BiSI nanotube, the infinite 2D Janus layer and the unstable inverse wrapped BiIS nanotube. Measures are given in Å.](image)

Optimal radius descriptors
As we have reported above, to estimate the optimal radius, we need to calculate the tubes at different sizes. These calculations are approximately three orders of magnitude more computationally-demanding than to simulate the corresponding infinite Janus sheets. Identifying descriptors to predict the optimal radius based exclusively on the 2D Janus sheet is a valuable pathway to faster navigate through the studied phase space and discover small-radius nanotubes.

All studied tubes are more stable if the chalcogen element is placed inside. In fact, for some of the studied non-isovalent nanotubes the optimal tube radius is independent of the halogen sitting on the outside of the tube (e.g., SbScI, SbSBr, and SbSi). Thus, only considering MX2 and MY2 parent structures provides an inaccurate picture when studying non-isovalent nanotubes. In addition, the parent structures of non-isovalent tubes will each have a M-element oxidation state different from the 2D Janus sheet (e.g., BiI2, BiS2, and BiI2). Two models from literature use the parent structure lattice constants as descriptors for predicting the optimal tube radius of isovalent nanotubes. The first approach is based on the plate theory as described by Timoshenko,34,35 while the second approach makes use of the Poison ratios of the parent sheets (Poisson model)15. Both models require the calculation of the stiffness tensors of the parent structures. Our approach, which we refer to as inner-bond model, is instead.
based on solely geometrical parameters, which can be readily obtained from existing 2D databases (e.g.36). By circumventing the time-consuming calculation of the stiffness tensor, we are able to calculate these descriptors at least one order of magnitude faster. The Inner-bond model uses the lattice constants of the 2D MXY Janus and MX2 parent sheets as well as the 2D Janus tM-X layer thickness, as discussed above. It is initially assumed that the optimal tube radius is determined by the lattice mismatch between the MXY Janus sheet and the corresponding MX2 parent sheet. The derivation of the Inner-bond model and the used formulas for the two reference models are provided in Supplementary Note 2.

All three models capture the rolling mechanism for isovalent anions for a selected set of materials (with discrepancies up to 30% compared to the calculated DFT radius—Fig. 7a), while only the Inner-bond model predicts the optimal radii of non-isovalent nanotubes to be small. Nevertheless, it fails to capture that the optimal DFT radius for some tubes, in which a chalcogen sits inside the tube, is almost independent of the chosen halogen element sitting outside of the tube (see orange bars for SbSCl, SbSBr and SbSI in Fig.7a). Here, the layer thickness tM-X is so thin, that the steric effects of the halogen atoms at small radii do not decisively impact the stability of the tube. In this picture, the wrapping mechanism is solely governed by the steric effects of the chalcogen elements. Controversely, the 2D Janus sheet lattice constants increases for SbSCl < SbSBr < SbSI due to the size difference of the halogen atoms, leading to a mismatch in predicted values for the Inner-bond model.

One feature all the studied materials share is that the lattice mismatch between the parent MX2 and the Janus MXY sheets is always smaller than 1 (Fig.7b). This is different from the idea of comparing MX2 and MY2 lattice constants, for which, in the case of mixing chalcogen and halogen X/Y elements, the lattice mismatch exceeds 1 (Fig.5a). Although only giving quantitative predictions, the Inner-bond model shows that the bond distances of the elements in the inner MX2 sheet together with the MXY lattice constant are good descriptors for predicting the optimal radius within the studied compound space. In the case of isovalent anions the lattice-mismatch of the parent structures can also be used as a descriptor.

Strain energy descriptors

The Janus sheets show an asymmetry between the two sides of the layer. We can make a sheet where the mirror symmetry along the mid-layer (M-layer) is restored through rearranging the X/Y elements into alternating X/Y rows (as shown in Fig. 8). The graph in Fig. 8 shows the energy difference between the alternating and the Janus sheet (E_{2D-alt} - E_{2D-Janus}) as a descriptor for the strain energy minimum (E_{strain-min}). Few outliers (>50 meV/atom error in prediction) correspond to structures with a low convex hull stability, i.e., E_{CH} > 0.2 eV/atom.

For instance, this descriptor, which does not require the calculation of the tubes, could be used to indicate if a sheet exfoliated onto a host structure might undergo spontaneous curling. The strain energy would need to be larger than the adsorption energy of the sheet on the host structure, which is for
example not the case for the experimentally exfoliated BiTeI sheet\textsuperscript{14}. Whether a large strain energy is needed to form single-wall nanotubes (in favor of multi-wall nanotubes or other forms of rolled structures) requires a deeper experimental investigation, or, at least, to consider growing and other experimental conditions which might impact the formation mechanism\textsuperscript{15}.

Different synthesis procedures can be imagined leading to the synthesis of the Janus nanotubes suggested here. Besides the exfoliation of a monolayer onto a host structure and its subsequent spontaneous wrapping, which might need lithography techniques to cut the monolayer into the required dimensions, the intercalation of ions into Janus multilayered materials, combined with ball-milling\textsuperscript{16}, can be a viable experimental synthesis path allowing to produce larger quantities. A common issue with these methods would be preventing unwanted reactions occurring at unsaturated bonds. A possible alternative synthesis pathway could be to use Atomic Layer Deposition combined with lithography to cut the 2D layer into a desired shape before it spontaneously wraps. Regardless the synthesis procedure, the intrinsic driving force of the 2D Janus sheets to form 1D tubes (or alternatively other curled shapes) should be large enough to be observed.

A promising starting material to experimentally verify our predictions is the layered BiTeI 3D bulk crystal structure for which experimental synthesis routes have been established\textsuperscript{17}. Other interesting candidates include tubes based on Vanadium or Titanium, which, by displaying a variety of oxidation states, are suitable for electrochemical applications\textsuperscript{18,19}. Curled multilayer VO\textsubscript{x} structures with much larger inner radii than the tubes predicted here have already been reported\textsuperscript{20}. Other non-isovalent anion based tubes, such as BiSL, might be feasible. However, the weaker ionic halogen bonds might drive reconstructions, in contrast to less reactive covalent bonds.

**DISCUSSION**

In this work, we investigated the formation mechanism of nanotubes by studying their stability and optimal nanotube radius. 135 different Janus nanotubes have been calculated using a structure prototype approach (T/H-phase). Each three-layered material consisted of one of 15 different cation mid-layer elements in combination with inner and outer atomic layers being occupied by either chalcogens, halogens or their mixture as anions. For isovalent anions, the wrapping mechanism could be explained by the lattice-mismatch between the two inner and outer atomic layers, while for non-isovalent anions steric effects caused through short pnictogen-chalcogen inside and longer pnictogen-halogen bonds outside of the material drive the stability. These effects are beneficial to the formation of some of the smallest identified nanotubes showing optimal radii below 10 Å. We noted that in general a large minimum strain energy is not needed to find tubes with an optimal radius smaller than 35 Å. In addition, the minimum strain energy was reasonably well estimated using the energy difference between a 2D Janus and alternating sheet as a descriptor. We employed Bayesian statistics to assess the quality of our fitting in order to identify uncertainties in our predictions of the optimal radius due, for example, to a misfit between the obtained data and the underlying strain energy curve function.

Nanotubes based on BiTeI and related compositions appear to be a particular interesting starting point for experimental verification due to their synthesizability in a 3D structure and exfoliability into 2D layers, as well as their predicted stability in 1D form. Following the interest in Vanadium- and Titanium-based nanotubes for electrochemical applications, we suggest that the metastable combinations of these metals paired with either OTe, OSe, or OS can be additional interesting candidates.

The findings reported here, shed light on the mechanism behind the curling of 2D Janus sheets and define a possible path for the synthesis of nanotubes with small radii, for which the lattice mismatch and the bonding character of the anions play a fundamental role.

**METHODS**

**Computational workflow**

The first step to create our library of nanotubes is to relax the 2D Janus sheets, taken from the Computational 2D Materials Database (c2db)\textsuperscript{36}. If a 2D Janus sheet is not present in the database, the 2D Janus sheet is created by averaging the lattice constants from its constituent MX\textsubscript{x}, and MX\textsubscript{y} parent sheets. Subsequently, tubes are generated by repeating and wrapping the 2D sheets both along the armchair and zigzag wrapping directions, thereby obtaining tubes with various radii (similar to what is shown in Fig. 1). In details, the initial number of unit cell repetitions is $n = (6, 8, 10, 12, 14)$ for the armchair and $n = (10, 13, 16, 19, 22)$ for the zigzag wrapping direction, which correspond to tubes with a radius smaller than 20 Å. We apply a set of filters to decide whether or not the relaxed structure is accepted for further investigation. These filters include assuming that a tube retained its circular shape and that no unwanted changes into different prototypes occurred during the relaxation. The filters discard ~40% of the data generated. A detailed discussion on how the data is filtered prior to visualization can be found in Supplementary Note 1. For consistent and reproducible calculations, we implement a workflow combining the Atomic Simulation Environment (ASE)\textsuperscript{28} and the workflow scheduling system MyQueue\textsuperscript{41}. Inspired by the CUSTODIAN package\textsuperscript{30}, we establish an "ASE error handler" to handle common DFT errors thus limiting the need for user intervention. A similar approach has been recently implemented to autonomously discover battery electrodes\textsuperscript{32}.

**DFT calculations**

All calculations are carried out with the Vienna ab initio Simulation Package (VASP) using a plane-wave basis set with an energy cutoff of 550 eV\textsuperscript{22,43}. In order to approximate the exchange-correlation effects the Perdew-Burke-Ernzerhof (PBE) form generalized gradient approximation (GGA) is used\textsuperscript{44}. A k-point density $\sim 4.7 / \text{Å}^{-1}$ is used to sample the Brillouin zone. All forces are converged to <0.01 eV/Å. The structures are relaxed in a non-magnetic configuration, i.e., without applying initial magnetic moments on the elements. A minimum vacuum in between repeating images of 16 Å is assured. Dipole corrections are applied along the non-periodic direction for materials with an out-of-plane dipole moment. To assess the stability versus 3D phases, we use a convex hull analysis, where the reference structures are the ones defining the convex hull in the Materials Project database\textsuperscript{37}. These structures are then relaxed with the matching input parameters used in this work\textsuperscript{45}, while the reference energy of oxygen is obtained by calculating the difference in energy between water and hydrogen in the gas phase, including the zero-point energy (ZPE) corrections\textsuperscript{46}. For the calculation of the stiffness tensor we use the workflow described by Haastrop et al.\textsuperscript{38}. Here, the k-point density applied for calculating elastic constants is >120 /Å\textsuperscript{3} together with an applied strain of ±1%.

**DATA AVAILABILITY**

The structures presented in this work are available on DTU DATA with the identifier \texttt{https://doi.org/10.11583/DTU.13153355}.

**CODE AVAILABILITY**

The workflow is continuously developed and may be accessed at https://gitlab.com/asc-dtu/workflows/nanotubes/.

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AUTHOR CONTRIBUTIONS

F.T.B. wrote all scripts for the analysis of the data and the draft of the manuscript. F.T.B. and A.E.G.M. implemented the workflow and performed the calculations. T.V. and I.E.C. designed the project. All authors contributed to discussing the results as well as writing and revising of the manuscript.

COMPETING INTERESTS

The authors declare no competing interests.

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