Atomic scale simulation of carbon nanotube nucleation from hydrocarbon precursors

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Atomic scale simulations of the nucleation and growth of carbon nanotubes is essential for understanding their growth mechanism. In spite of over twenty years of simulation efforts in this area, limited progress has so far been made on addressing the role of the hydrocarbon growth precursor. Here we report on atomic scale simulations of cap nucleation of single-walled carbon nanotubes from hydrocarbon precursors. The presented mechanism emphasizes the important role of hydrogen in the nucleation process, and is discussed in relation to previously presented mechanisms. In particular, the role of hydrogen in the appearance of unstable carbon structures during in situ experimental observations as well as the initial stage of multi-walled carbon nanotube growth is discussed. The results are in good agreement with available experimental and quantum-mechanical results, and provide a basic understanding of the incubation and nucleation stages of hydrocarbon-based CNT growth at the atomic level.

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Controlling and steering the growth of carbon nanotubes (CNTs) is often believed to require control over the nucleation stage. In spite of recent advances, gaining precise control over the resulting structure proves a daunting task. Current synthesis techniques include arc discharge, laser ablation, fullerene recrystallization, catalytic chemical vapour deposition and plasma-enhanced CVD, all of which are capable of producing nanotubes with well-defined structures to a certain extent. Catalytic chemical vapour deposition, in which a hydrocarbon gas is catalytically dissociated at the nanocatalyst surface at elevated temperature, is currently the most common experimental technique to grow CNTs due to its simplicity, high degree of control and scalability. However, despite numerous experimental, theoretical and simulation results to understand the CNT growth, the onset (that is, incubation and nucleation stages) of CNT formation using hydrocarbon \( C_nH_m \) molecules remains still unclear.

Although environmental transmission electron microscopy studies are capable of showing time-resolved details of metal-catalysed CNT nucleation and growth at the atomic level, current transmission electron microscopy (TEM) resolution is still insufficient to observe atomic scale dynamics. Consequently, a variety of computational techniques including quantum mechanical, tight-binding and classical approaches have been applied to simulate CNT growth on the atomic scale. A wide variety of effects, including the catalyst particle size, the importance of the carbon chemical potential, the necessity of a carbide phase, the interaction between metal and carbon network, the influence of growth precursor, using either single-wall or multi-wall carbon nanotubes and energetic C-ions as input gas, as well as the influence of ion bombardment, have been computationally studied.

Until now, however, none of these studies considered cap nucleation from a hydrocarbon growth precursor. Instead, pure carbon (either as C or C\(_2\)) is invariably used as carbon source. Hence, instantaneous hydrocarbon decomposition is implicitly assumed, whereas the role of hydrogen is never considered. Only very recently, the first dynamic simulation studies appeared on related processes using hydrocarbon molecules as carbon feedstock. Somers et al. and Liu et al. employed reactive molecular dynamics (MD) simulations to study the interaction of methane and methane-derived radicals with Ni-surfaces in the context of plasma catalysis. Shibuta et al. performed ab initio MD simulations to study the decomposition of \( CH_4 \) and \( C_2H_4 \) molecules on a Ni(111) surface and the reactivity of \( CH_4 \) on a Cu(111) surface in the context of metal-catalysed graphene growth. Recently, Wang et al. studied the interaction of acetylene with a Fe\(_{58}\) cluster by density-functional tight-binding simulations and concluded that cap formation is not a necessary condition for single-walled CNT (SWNT) nucleation and large hydrocarbon clusters can grow before hydrogen abstraction would allow the formation of a carbon-only tubular structure. In contrast, however, experimental in situ studies by Hofmann et al. and Yoshida et al. demonstrated cap lift-off preceding SWNT growth on a surface-bound Fe or Ni nanocatalyst using a \( C_2H_2 \) feedstock. In our previous work, we also demonstrated the formation of such vertical graphene nanowalls, which transform to a horizontally oriented carbon network, partially covering the catalyst surface. So far, however, there have been no reports on cap and nanotube nucleation from hydrocarbon species.

In this work, we report on CNT cap nucleation using different hydrocarbon species at different temperatures to unravel the initiation of CNT growth by combined reactive MD and time-stamped force-bias Monte Carlo (tfMC) simulations. We find that the competition between (re)hydrogenation and dehydrogenation processes during the incubation stage is critical for the dynamics of the cap formation and subsequent CNT growth process.

**Results**

**Dehydrogenation degree.** It is known that hydrogen may either enhance CNT growth or etch the growing CNTs, and thus has a dual role. Both effects depend on the H concentration at the surface. Therefore, steering the H concentration during the CNT growth is very important. Until now, however, the precise role of adsorbed H atoms in the onset of CNT growth has not yet been studied down to the atomic level. As demonstrated below, the growth from hydrocarbon gas precursors depends on the adsorption rate of \( C_nH_m \) precursors, and the desorption rate of \( H_2 \) molecules and \( C_{m-y}H_{m+y} \) species. The calculation of adsorption and desorption rates is thus elementary to assess the dehydrogenation degree of the adsorbed precursor and the catalyst cluster during the growth. The \( k \)-coefficient is introduced to estimate the dehydrogenation degree, to analyse the CNT growth stages:

\[
k = \frac{mN_C - nN_H}{mN_C + nN_H}
\]

where \( n \) and \( m \) are the number of carbon and hydrogen atoms, respectively, in the \( C_nH_m \) molecule and \( N_C \) and \( N_H \) are the total number of adsorbed carbon and hydrogen atoms in the cluster, respectively. The \( k \)-coefficient (\( k \in [0,1] \)) allows to easily differentiate between physisorption of the hydrocarbon molecule on the catalyst surface (\( k = 0 \)) and complete dehydrogenation of the adsorbed molecule and the catalyst (\( k = 1 \)). Note that the coefficient depends on both the \( H_2 \) desorption rate and the H-arrival rate, but it does not depend on how the precursor decomposes.

It is well known that different structures may be formed, depending on the adsorption (hydrogenation) and desorption (dehydrogenation) rates, temperature, partial pressure of the hydrocarbon molecule and the H-content in the molecule. Thus, methane (\( CH_4 \)), acetylene (\( C_2H_2 \)) and benzene (\( C_6H_6 \)) will typically yield different structures. Yet, we find that the overall chemical nature of the hydrogenation and dehydrogenation processes is very similar for these molecules, and we can therefore generalize all cases in a single mechanism, as depicted in Fig. 1.

The figure shows the evolution of the cap formation, the number of C and H atoms on and in the nanocluster, the formation of carbon hexagonal, pentagonal and heptagonal rings, as well as the evolution of the \( k \)-coefficient as a function of the simulation time. Based on this evolution, we distinguish three main stages in the onset of CNT growth: incubation (stage I), cap formation (stage II) and continued growth (stage III).

**Stage I.** The incubation stage is vital in understanding the onset of CNT growth. Although this stage has been studied before by many researchers, there is currently little knowledge on its role in relation to the subsequent cap formation stage, and in particular in the case of hydrocarbon-based growth (see Supplementary Table I for a comparison in this regard between earlier work and the current work). We therefore here first analyse stage I, which we divide in four sub-stages.

In the first substage (or supersaturation stage), all C–H and C–C bonds of the impinging hydrocarbon molecule are gradually broken and the C atoms eventually dissolve into the pure Ni\(_{55}\) nanocluster (Supplementary Fig. 1) in agreement with earlier studies. H adatoms freely diffuse over the nanocluster and recombine with other H atoms on the surface to...
desorb as H₂ from the nanoparticle. Also, at high-impingement flux and low temperature, some adsorbed molecules (for example, C₆H₆ or CH₄) do not completely dehydrogenate. The number of adsorbed/dissolved carbon atoms quickly increases due to the fast hydrogen desorption and therefore the k-coefficient quickly rises towards one (Fig. 1b, substage 1, see k). Generally, in this substage, this C₆H₆ adsorption/dissociation and H₂ desorption scenario continues until the cluster is (super)saturated. Owing to the dissolution of carbon in the particle, the wetting angle of the Ni₅₅ cluster is therefore the Ni₅₅-X adatom evolution of vertical (free-standing), horizontal (adsorbed on the nanocluster) and total number of carbon rings as a function of the simulation time.

Figure 1 | Analysis of the CNT growth stages. (a) Evolution of the CNT cap nucleation process up to 6.8 × 10⁷ (± 2 × 10⁶ MC) MD-steps; Ni, C and H atoms are coloured light green, blue and red, respectively; (b) evolution of the k-coefficient and the number of C atoms (including dissolved C_H and sp-hybridized C_sp carbon atoms) and H adatoms on and in the catalyst nanocluster as a function of the simulation time; (c) evolution of vertical (free-standing), horizontal (adsorbed on the nanocluster) and total number of carbon rings as a function of the simulation time.

The onset of the third substage (or formation of vertical nanowalls) of stage I is characterized by the formation of the first carbon ring on the surface. We find that the initial pentagon (Fig. 1a, 9.0 and Fig. 1c) or hexagon ring (Fig. 1c) can be formed in four distinct ways: (i) by incorporation of segregated or surface C atoms in an existing surface carbon structure; (ii) by folding of a polyyne (C₆H₆ or C₇H₇ or C₆H₅) chain into a ring structure (Supplementary Fig. 2); (iii) in the case of acetylene: by direct interaction of gas-phase C₂H₂ with surface C-structures such as a carbon trimer thereby forming a pentagon; or (iv) in the case of benzene: by incorporation of the impinging molecule in an existing surface carbon structure after partial dehydrogenation (see Supplementary Fig. 1, C₆H₆ case). Although the former two processes have been described earlier in terms of the cross-linking mechanism proposed by Eres et al. and the ‘pentagon-first’ mechanism presented by the group of Irle and Morokuma, the latter two processes have not yet been described before, as they are specific to hydrogen-containing growth precursors.

The appearing pentagons and hexagons are mobile and may either concatenate or connect with (sub)surface carbon atoms or other carbon structures to form a small graphitic nucleus. Meanwhile, diffusing H adatoms interact with and add to the various carbon surface structures including partially dissociated C₆H₆, polyyne chains, polyene chains and incipient pentagon–hexagon networks on the surface (‘rehydrogenation’), or they may recombine with other H atoms to desorb as H₂ from the surface (‘dehydrogenation’). Owing to the presence of the carbon surface structures, the H atoms cannot diffuse freely over the surface. As a result, the H₂ desorption rate decreases, and the number of surface H atoms significantly increases. This leads to a decrease of the k-coefficient (Fig. 1b, see H and k, substage 3). When rehydrogenation occurs, the surface carbon structures including polyyne chains and graphene-like networks become partially hydrogenated and vertically free-standing graphene sheets or carbon nanowalls are formed on the catalyst surface (Supplementary Fig. 3), corresponding to recent ab initio MD and density-functional tight-binding-based MD studies. Subsequently, these parallel graphene-like sheets may transform to hydrogen-terminated few-layer graphene and thereby form the initial structures leading to lift-off of a multi-walled CNT (MWNT) or a carbon nanofiber as proposed by Hofmann et al. Our results indicate that while a high temperature and low H-content yield the formation of SWNTs, a low temperature and high H-content favour MWNT nucleation (Supplementary Fig. 4), in qualitative agreement with the general experience of CVD experiments, that is, low-temperature CVD (600–900°C) yields MWNTs, whereas high-temperature (900–1200°C) conditions favour SWNT growth.

Also, we observe that smaller free-standing graphene patches as well as other initial carbon structures can be etched if the flux of H atoms is sufficiently high, as is the case for CH₄ as growth precursor in our simulations, which is in agreement with experimental reports using methane as growth precursor above 800°C (ref. 40). This indicates that the growth rate of CNTs from CH₄ can be much lower compared with growth from C₂H₂ and C₆H₆ due to the higher (re)hydrogenation rate compared with the dehydrogenation rate.

Thus, in this substage, both horizontally and vertically oriented carbon patches can exist (Fig. 1a, time 16.0 and time 26.0), due to competition between dehydrogenation and rehydrogenation, which in turn strongly depends on the growth temperature, type of hydrocarbon species and their gas-phase pressure (or flux) as well as the catalyst-substrate interaction. The competition between rehydrogenation, dehydrogenation and H etching thus explains how unstable carbon patches may appear in the incubation stage, as for example, observed during in situ TEM...
observations\(^8\). Such studies indeed found protrusions of various unstable carbon structures before eventual cap appearance and lift-off.

In the fourth and final substage of stage I, vertical nanowalls and horizontal nanosheets can be simultaneously found on the catalyst surface. However, the vertical free-standing networks invariably and irreversibly transform to horizontal networks covering the surface. Consequently, the graphic network starts to gradually cover the nanocatalyst surface.

In Fig. 2, potential energies per atom of five selected samples, which are found during the aforementioned stages, are shown (solid circles). From the figure, it is clear that horizontal graphene patches, covering the surface, are energetically favoured over their vertical free-standing counterparts. Accordingly, the interaction energy between a C\(_{40}\) graphene patch and the nanocluster significantly increases when the vertically oriented carbon nanowall (Ni\(_{55}\)C\(_{8}\) + C\(_{40}\)H\(_{12}\) and Ni\(_{55}\)C\(_{8}\) + C\(_{40}\)H\(_{8}\)) transforms to the horizontally oriented nanosheet (Ni\(_{55}\)C\(_{8}\) + C\(_{40}\)), after elimination of their edge-terminated H atoms (open circles in Fig. 2).

As a result of this transformation, the rate of hydrocarbon adsorption decreases due to the decreasing free-metal surface area. The increase in number of carbon atoms therefore drops relative to the previous stage, and remains constant throughout the rest of the simulation time (Fig. 1b, see C, stage 4). Likewise, the number of H adatoms decreases due to both a small adsorption rate of C\(_{2}\)H\(_{m}\), the high H\(_2\) desorption rate (Fig. 1b, see H) and thus the k-coefficient rises again. When H atoms connect to the edges of the horizontal graphene patches, the carbon sheets can only partially stand upright on the cluster (Fig. 1a, time 26.0), limiting the number of free-standing rings in this stage (Fig. 1c). Also, the number of polyyne chains with or without H-termination (that is, C\(_y\) or C\(_y\)H) becomes negligible for the rest of the simulation.

**Stage II.** This second main stage is characterized by the appearance of a carbon cap on the cluster, which is essential for the nucleation of a CNT (refs 7,8). In this stage, the adsorption rate of C atoms does not change, whereas the density of H atoms continues to decrease. The k-coefficient can therefore rise close to one (Fig. 1b, see k, stage II). The dehydrogenation rate is now somewhat slower compared with fourth substage of stage I. Also, the horizontally oriented graphene-like patches start to lose their H atoms from their edges. In agreement with first-principles calculations\(^39\), the cap on a curved Ni-cluster is more stable than a graphene sheet (Fig. 2), and at a sufficiently high temperature a cap hence develops and eventually lifts off (Fig. 1a, time 40.0).

In Fig. 3, a cap as formed during low-flux C\(_2\)H\(_2\) impacts at high temperature is shown. In the cap, the central pentagon ring is surrounded by five hexagon rings (Fig. 3b) indicating essentially defect-free CNT nucleation. The formation of this carbon dome, containing a hexagon-surrounded pentagon ring or a hemispherical fullerene structure, corresponds very well with both in situ experiments\(^7\) and ab initio MD simulations\(^35\).

After cap formation, a metal step appears between the tube and the substrate due to the smaller diameter of the cap than the diameter of the Ni cluster (Fig. 3). Although the formation of such steps is important for the growth of MWNTs (refs 7,54,56), we observe that the growing nanotube actually forces its shape onto the Ni cluster (Fig. 1a, time 68.0) and the metal steps disappear during the growth, corresponding to observations in in situ experiments for SWNT growth\(^7\).

Our simulations demonstrate that cap nucleation in CNT growth from hydrocarbons does not require the formation of long polyyne chains. This is in contrast to the many simulations predicting the formation of such chains on the surface when pure, hydrogen-free carbon is used as growth precursor\(^4\). Although the presence of such chains during the nucleation and growth has been suggested before\(^52\), no direct experimental evidence for their occurrence, however, has been produced yet. Indeed, our simulations demonstrate that cap nucleation does not require the formation of such long polyyne chains.

At low temperature, we do not observe the formation of a cap, corresponding to earlier MD simulations demonstrating how lift-off requires a weak work of adhesion, high growth temperature (or sufficient thermal kinetic energy) and/or a large catalyst or a low curvature of the catalyst surface\(^20,21\). Finally, note that in our simulations the substrate-bound semi-spherical nanocatalyst is not entirely encapsulated by the carbon network at low temperature and a significant part of the network edge is quickly saturated by H atoms.

**Stage III.** In the final stage, the CNT continues to grow on the substrate-bound catalyst surface (Fig. 1a, time 68.0) due to the incorporation of the segregated C atoms, diffusing C adatoms or through direct adsorption of partially dehydrogenated C\(_{n-x}\)H\(_{m-y}\) fragments near the metal/cap interface followed by dehydrogenation.

In Fig. 4, C diffusion and its incorporation mechanism during CNT growth is shown for the case of C\(_2\)H\(_2\) impacts. When the molecule impinges on the cluster, it quickly loses one of its H atoms (Fig. 4a). Consequently, the remaining ethynyl (C\(_2\)H) radical diffuses over the Ni surface before it incorporates into the nanotube base (Fig. 4b). After incorporation, one of the C atoms of the C\(_2\)H radical diffuses into the cluster (Fig. 4c) while the H atom diffuses over the surface (Fig. 4d). The C atom from the
remaining part of the ethynyl radical binds to a C atom in the tube edge and eventually a new hexagon ring appears at the tube-cluster interface (Fig. 4e). As a result, the second H atom is also eliminated from the ring and it can subsequently also diffuse over the surface (Fig. 4f). Finally, a C atom from the bulk may segregate at the surface (Fig. 4g) and subsequently incorporate into the CNT edge (Fig. 4h).

Our results indicate that C atoms initially diffuse into the catalyst cluster after C6H6 decomposition (Supplementary Fig. 1), according to the vapour–liquid–solid model46 and in agreement with various experimental observations8,54,57, two scenarios turn out to be possible after supersaturation: (a) either C atoms continue to diffuse into the cluster, provided they completely lose their hydrogens, or (b) they diffuse over the catalyst surface, in case they keep or only partially lose their hydrogen. In either case, the C atoms eventually incorporate into the growing network (Supplementary Figs 5–8).

Finally, note that in the case of high H-content (for example, CH4 and C6H6) at high flux conditions, the nanotube either grows very slowly or grows with many defects due to the presence of a significant number of H atoms41. These H atoms can react with defective sites of the CNT when hydrocarbon molecules dissociate on the nanocluster surface. In the last stage, the k-coefficient remains very close to one, indicating the presence of some H adatoms, signalling a suitable growth condition when using hydrocarbon feedstock (Fig. 1b, see H, stage III).

Discussion

The overall results demonstrate the important role of the dehydrogenation degree in determining the various growth stages. In the first two substages of the incubation stage (stage I), adsorbed C6H6 species quickly decompose and lose their H atoms. Consequently, C atoms dissolve into the cluster (Supplementary Fig. 1). In all hydrocarbon impact cases, the k-coefficient rises towards its maximum (that is, k tends to 1), as the H2 desorption rate is much higher than the H-arrival rate. Owing to the high degree of dehydrogenation, the system quickly saturates with C atoms and consequently initial C-structures appear on the cluster. Also in substage 4 (formation of horizontal graphene sheets on the surface) of stage I and in stage II (formation of a carbon cap), k is found to increase. When k \approx 1, the effect of hydrogen on the nucleation and growth process is negligible, and these stages are therefore very similar to the growth stages as found in growth simulations based on C-only precursors.

In the third substage of the incubation stage, k decreases significantly (that is, k tends to 0): the increase in the number of adsorbed H atoms exceeds the number of desorbed H2 molecules per unit of time: after supersaturation, surface C atoms increase fast and diffusing H atoms quickly connect to initial C-structures (rehydrogenation) rather than to find other H atoms to desorb from the surface as H2 (dehydrogenation). Such fast hydrogenation eventually leads to the formation of the free-standing graphitic patches (see Supplementary Figure 3). Therefore, a low dehydrogenation degree is characterized by the appearance of H-terminated free-standing graphene patches in the system. In this stage, the effect of the presence of hydrogen cannot be ignored, and simulations based on pure C-precursors cannot provide any information on this growth stage.

In the third stage of the nucleation process, the carbon cap covers a significant part of the catalyst nanoparticle and only a small area of the catalyst surface remains available for hydrocarbon species to impinge and dissociate before H2 desorption may occur (Supplementary Figs 6–8). The combination of a low growth rate, a high concentration of C atoms in the system, a low-sticking probability of gas-phase species and the H desorption and H arrival rates being equal to each other leads to the k-coefficient to be continuously high and close to one (that is, constant kmax), which is the essential condition for continued CNT growth from hydrocarbons.

In summary, using combined MD/Monte Carlo simulations, the nucleation of SWNT caps from hydrocarbon precursors (CH3, C2H2 and C6H6) is studied for the first time. The complete nucleation mechanism consists of three main consecutive stages: incubation (consisting of four substages), cap formation and continued growth.

In the incubation stage, impinging hydrocarbon molecules may contribute to initial ring formation after supersaturation, in addition to earlier suggested mechanisms. Free-standing graphene nanowalls as well as a horizontal carbon network are found on the catalyst cluster due to competition of (re)hydrogenation and dehydrogenation processes. We also find that dehydrogenation, rehydrogenation and H etching can explain a number of experimental observations: (1) the appearance of unstable carbon protrusions during the incubation stage in in situ transmission electron microscopy-observations; (2) the onset of MWNT nucleation through the formation of a multi-layer graphene structure in the case of low temperature and high H flux; and (3) the lower growth rate of CNTs from CH4 compared with growth from C2H2 and C6H6.

In the second stage, the cap formation mechanism proceeds from a horizontal carbon nanosheet through H-elimination. We find that cap nucleation in CNT growth from hydrocarbons does not require the formation of long polyyne chains. At low temperature, the substrate-bound semi-spherical nanocatalyst is not entirely encapsulated by the carbon network and a significant part of the network edge is quickly saturated by H atoms.

In the growth stage, C atoms may either diffuse into the cluster after full dehydrogenation, or diffuse over the surface in the case of partial dehydrogenation. In either case, they are found to eventually incorporate into the carbon network. During the growth, the growing nanotube is observed to force its shape onto the Ni cluster and consequently metal steps gradually disappear.

We thus conclude that the dehydrogenation degree can distinguish various growth (sub) stages and thus control over the k-coefficient is a highly important factor in controlling CNT nucleation and growth.

Methods

Hybrid MD/tfMC technique and ReaxFF. Combined reactive MD and tfMC simulations57,58,59 are used to simulate the catalysed CNT growth process. In tfMC calculations, all the atoms in the system are displaced at once in every time step with unit probability, and thus generate a system evolution in a MD-like fashion, instead of, for example, single-particle moves common in Metropolis MC (ref. 57). This technique can therefore very efficiently be coupled to canonical MD simulations: the MD module accounts for the impacts of the growth species on the
catalyst surface and for the initial chemical reactions taking place during the first few picoseconds, whereas the tMC module takes care of the longer timescale relaxation of the system in a single tMC simulation step, each atom is displaced in a direction that is dependent on the force (deterministic component), as well as on the temperature (stochastic component)\(^\text{59}\). The typical step size in tMC is about 0.1 \(\AA\), which is at least one order of magnitude longer than in MD (\(\sim 0.01 \text{ \AA}\)). Bond dissociation and other activated processes occur in MD only due to thermal fluctuations, rendering these events to be rare. In tMC, in contrast, the random component in the displacement algorithm allows atoms to move against the force, and thus allow the system to cross transition states and attain a more stable configuration much more quickly than is possible in MD. Although it is clear that the exact system dynamics are not reproduced by tMC, it has been demonstrated that tMC correctly accounts for detailed balance, provides a realistic (albeit not exact) dynamical path and reproduces end configurations in full agreement with (very long) MD simulations. Both during the MD and the tMC cycles, ReaxFF (refs 61,62) is applied to properly describe the bond dissociation and formation processes, that is, the C–C bond dissociation, dehydrogenation, rehydrogenation and H2 growth as well as reatom reactions. Currently, accelerated MD simulations are carried out to further extend the time scale of CNT growth simulations\(^\text{63}\).

### Simulation details

A Ni\textsubscript{100} nanocatalyst is equilibrated at the desired growth temperature using the canonical Busi thermostat\(^\text{64}\). The nanocatalyst isphisorbed on a virtual Al or Si substrate, employing a 2–3-integrated Lennard-Jones potential\(^\text{58}\). Subsequently, methane (CH\textsubscript{4}), acetylene (C\textsubscript{2}H\textsubscript{2}) or benzene (C\textsubscript{6}H\textsubscript{6}) molecules are allowed to impinge on the cluster. Throughout the simulation, the total number of gas-phase hydrocarbon species is kept constant. When a hydrocarbon molecule adsorbs on the Ni cluster, the resulting structure is allowed to relax by application of tMC (refs 37,60). During the relaxation, no new molecules are allowed to impinge on the cluster. Application of the combined tMC/MD technique was previously demonstrated to allow relaxation processes in general and the simulation of the CNT nucleation process in particular much more efficiently compared to pure MD simulations\(^\text{37,64}\). All simulations are performed at a temperature in the range of 1000–2000 K controlled by the canonical thermostat\(^\text{64}\).

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Author contributions
E.C.N. conceived the idea and U.K. performed all simulations. U.K. and E.C.N. carried out the data analysis. U.K., A.B. and E.C.N. contributed to the discussion and writing of the manuscript.

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