Dynamics of tungsten hexacarbonyl, dicobalt octacarbonyl and their fragments adsorbed on silica surfaces

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Tungsten and cobalt carboxyls adsorbed on a substrate are typical starting points for the electron beam induced deposition of tungsten or cobalt based metallic nanostructures. We employ first principles molecular dynamics simulations to investigate the dynamics and vibrational spectra of W(CO)$_6$ and W(CO)$_5$ as well as Co$_2$(CO)$_8$ and Co(CO)$_4$ precursor molecules on fully and partially hydroxylated silica surfaces. Such surfaces resemble the initial conditions of electron beam induced growth processes. We find that both W(CO)$_6$ and Co$_2$(CO)$_8$ are stable at room temperature and mobile on a silica surface saturated with hydroxyl groups (OH), moving up to half an Ångström per picosecond. In contrast, chemisorbed W(CO)$_5$ or Co(CO)$_4$ ions at room temperature do not change their binding site. These results contribute to gaining fundamental insight into how the molecules behave in the simulated time window of 20 ps and our determined vibrational spectra of all species provide signatures for experimentally distinguishing the form in which precursors cover a substrate.

I. INTRODUCTION

The study of the energetics and dynamics of individual molecules adsorbed on a substrate is relevant for several prominent fields of research such as molecular electronics, molecular magnetism and catalysis. The possibility to manipulate individual molecules adsorbed on surfaces using the tip of a scanning tunneling microscope (STM) leads to an increasing need for theoretical information of the geometries as well as adsorption and desorption mechanisms and reaction pathways of adsorbates. STM probes can not only be used to arrange complex molecules on surfaces but also to measure their vibrational spectra. Molecules adsorbed on insulating surfaces which due to insufficient conductivity are hard to study by STM techniques have been investigated successfully using non-contact atomic force microscopy.

Here, we will study molecules and substrates that are important in the context of electron beam induced deposition (EBID) of organometallic precursor molecules. This is a widely used method to grow size- and shape-controlled nanometer-sized structures. The obtained EBID deposits nevertheless possess a significant percentage of organic contaminants mainly from carbon and oxygen, lowering the conductivity of these deposits and thus limiting the possible applications of EBID. Several pre- and post-fabrication approaches have been employed to remove these contaminants, but reproducibility is still an issue. First principles calculations can provide a detailed description of the microscopic behavior of the deposits and are often used to improve the quality of the deposition processes. Important progress has been done, for instance, on atomic layer deposition and chemical vapor deposition processes.

Recently, in an attempt to understand the EBID growth process, we analyzed by means of density functional theory (DFT) calculations the interaction of precursors like W(CO)$_6$, Co$_2$(CO)$_8$ and CH$_3$(C$_5$H$_5$)Pt[CH$_3$]$_3$ on two different SiO$_2$ surfaces (fully and partially hydroxylated) as a representative for untreated and pretreated EBID surfaces. Hereafter, surfaces corresponds to SiO$_2$ surfaces unless otherwise mentioned. These studies illustrate the preferred orientation of the adsorbate and the nature of the interaction between the precursor molecules and the SiO$_2$ substrates. Further, interesting phenomena such as the spontaneous fragmentation of the carbonyl precursors (W(CO)$_6$ to W(CO)$_5$ and Co$_2$(CO)$_8$ to two Co(CO)$_4$ molecules) on the partially hydroxylated surfaces that correspond to pre-treated surfaces were observed.

It has recently been reported that the surface residence time of an organometallic precursor should be sufficiently long (lasting from microseconds to milliseconds) to have an efficient deposition yield. In order to understand these observations and to improve the conditions for the adhesion of precursor molecules to the substrate, it is essential to have knowledge on the behavior of the precursor molecules and their fragments on the surface SiO$_2$ substrates. Theoretical and experimental studies focussing on the adsorption process of free CO on various surfaces and several molecules on different SiO$_2$ substrates have been reported. However, little is known on the dynamics of W(CO)$_6$ and Co$_2$(CO)$_8$ precursors and their fragments on SiO$_2$ surfaces. Therefore, in this work we use first principles molecular dynamics simulations to investigate the nature of W(CO)$_6$ and Co$_2$(CO)$_8$ molecules adsorption on fully and partially hydroxylated SiO$_2$ surfaces and provide quantitative microscopic insight into the stability of these fragmented precursors on these surfaces and on their vibrational spectra.
II. COMPUTATIONAL DETAILS

Ab initio molecular dynamics (MD) simulations were performed in the framework of density functional theory (DFT). We employed the projector augmented wave (PAW) basis within the generalized gradient approximation in the parametrization of Perdew, Burke and Ernzerhof (PBE) as implemented in VASP. Spin polarized calculations including the corrections for long range van der Waals interactions were used in all calculations. All ions were relaxed using the conjugate gradient scheme until the forces were reduced by $\lesssim 0.01\text{ eV/Å}$ with a plane wave energy cut-off of 400 eV. The Brillouin zone for the substrate-precursor complex was sampled at the gamma point only.

The structures considered as initial configurations for the MD simulations are shown in Figs. 1 and 5 (see the $t = 0$ ps snapshot). MD simulations were performed for 20 ps on a canonical ensemble at a finite temperature of $T = 298\text{ K}$ using the Nose-Hoover thermostat. The temperature was chosen in accordance with the reported experimental results where the largest temperature rise during W(CO)$_6$ deposition was $1\text{ °C}$ when a 1.42 nA electron beam was used for depositing tungsten nanostructures. The Verlet algorithm in its velocity form with a time step of $\Delta t = 1\text{ fs}$ was used to integrate the equations of motion. We also performed MD simulations for the precursor molecules (W(CO)$_6$ and Co$_2$(CO)$_8$) in the gas phase by placing the molecules in a cubic box of $a = b = c = 30\text{ Å}$ using similar parameters used for the substrate-precursor complex. The length of the simulation for the molecule was limited to 10 ps.

The power spectrum $I(\omega)$ provides information about the distribution of the vibrational energy of the system. It can be computed by a Fourier transformation of the velocity autocorrelation function obtained from the velocities $\mathbf{v}_i(t)$ of all atoms in the course of the MD trajectory. The method we use is described in the Appendix. Initially, we have compared the vibrations of the gas phase precursor molecules obtained by the finite displacement method with the calculated power spectrum (Vibrations through the finite displacement method were computed using Turbomole 6.0). The geometries were optimized using triple $\zeta$ basis sets for all elements and we used an effective core potential (60 core electrons) for W in the case of W(CO)$_6$. Therefore, vibrations for the SiO$_2$ substrate and the complex precursor molecule-
FIG. 2. (Color online) Schematic illustration of the movement of W(CO)$_6$ on a fully hydroxylated surface. In the top panel, the polygon of the oxygen atoms of W(CO)$_6$ is shown, with one facet always shown in red in order to visualize the rotations of the molecule. In the lower panel the time evolution of the three center-of-mass (COM) coordinates of W(CO)$_6$ is shown. FOH in the figure corresponds to the fully hydroxylated surface.

FIG. 3. (Color online) DFT optimized structure of W(CO)$_5$. In the gas phase it possesses a C$_{4v}$ symmetric structure. The trigonal bipyramidal structure (middle panel) is 0.643 eV less stable than the square pyramidal structure (left and right panels) but it is stabilized by surface-molecule interaction on partially hydroxylated.

FIG. 4. (Color online) Variations in the structural features of W(CO)$_6$ on SiO$_2$ surfaces on fully hydroxylated (left panel) and partially hydroxylated (right panel). Bold (red) letters correspond to WC and normal letters (blue) for CO bond. The values shown in this figure are the standard deviation of the respective values in the initial structure.

III. RESULTS

A. Dynamics of W(CO)$_6$ on SiO$_2$ surfaces

Previous DFT calculations indicate that W(CO)$_6$ interacts through weak physisorption with surface hydroxyls on the fully hydroxylated surface and by strong chemisorption on the Si sites available on partially hydroxylated surfaces with substantial changes in the structure and electronic properties. The most stable configuration of W(CO)$_6$ on fully hydroxylated has an adsorption energy of -0.498 eV while the fragment W(CO)$_5$ together with a free CO ligand stabilize with an energy -1.262 eV on a partially hydroxylated surface. In Fig. 1 we show five snapshots in 5 ps intervals of the MD simulations of W(CO)$_6$ on a fully hydroxylated surface (up-
FIG. 5. (Color online) Adsorption of Co$_2$(CO)$_8$ on fully hydroxylated (top) and partially hydroxylated (POH) substrates. The illustrated snapshots are the configurations taken at every 5ps interval. S1 and S2 in partially hydroxylated case denotes the two bonding sites on the surface where Co(CO)$_4$ fragments are bonded. Color code: Blue - Co, throughout this manuscript. The orientation of the configuration is denoted by the coordinate system on the bottom left hand side of each panel.

FIG. 6. (Color online) Evolution of the three center-of-mass coordinates of Co$_2$(CO)$_8$ adsorbed on a fully hydroxylated surface during the 20 ps trajectory.

FIG. 7. (Color online) Calculated vibrational spectrum of the gas phase of W(CO)$_6$ molecules by Fourier transforming the velocity autocorrelation function obtained from the MD trajectory (upper panel) and by the finite displacement method (lower panel).

In a fully hydroxylated surface. In order to visualize this drifting, Fig. 2 shows a schematic depiction of the W(CO)$_6$ displacement together with the center-of-mass (COM) analysis. The calculated drifting distance on the surface after 20 ps is ca. 5 Å. These simulations illustrate that the undissociated W(CO)$_6$ molecule changes...
the orientation considerably but does not desorb away from the fully hydroxylated surface.

However, a chemisorbed W(CO)$_5$ molecule which is formed by the release of a CO ligand from W(CO)$_6$ (Fig. 1 lower panel) remains localized on its binding site on a partially hydroxylated surface. The dissociated CO ligand doesn’t recombine with the parent moiety and the vacant site on W remains empty and is not filled by surface hydroxyls as has been suggested as a possibility for Mo(CO)$_5$. It should be noted that W(CO)$_5$ in the gas phase is stable on a square pyramidal structure and different conformations are possible through the pathway shown in Fig. 3 involving a trigonal bipyramidal transition state. Analysis of the adsorbed W(CO)$_5$ structure (compare the transition state on Fig. 3 with the configuration at 0 ps in Fig. 1 lower panel) indicates that the W(CO)$_5$ molecule is stabilized in a trigonal bipyramidal structure on partially hydroxylated surfaces. The stabilization of such a transient intermediate has been proposed for Cr(CO)$_5$ and the present work supports such a proposal also for W(CO)$_5$. This stabilization, should have an impact over the kinetics of further release of intra molecular CO ligands.

Evaluation of the changes in the WC and CO bond lengths of the adsorbates W(CO)$_6$ and W(CO)$_5$ (see Fig. 4) during the first 20 ps of the trajectory show deviations from the initial configuration of the order of 1-2% and might be due to thermal fluctuations. We will analyze these deviations in more detail in Section III C.

The electronic structure of the final configurations of the complex molecule-substrate on both fully and partially hydroxylated cases shows only minor variations with respect to the initial configurations (results not shown).

B. Dynamics of Co$_2$(CO)$_8$ on SiO$_2$ surfaces

We now proceed with the adsorption scenario of Co$_2$(CO)$_8$ interacting with SiO$_2$ surfaces. In Ref. a weak bonding of Co$_2$(CO)$_8$ on fully hydroxylated surfaces was observed, similar to the case of W(CO)$_6$, with an adsorption energy of -0.76 eV. In contrast, Co$_2$(CO)$_8$ molecules on a partially hydroxylated surface fragment into two Co(CO)$_4$ moieties rather than eliminating a CO ligand as in the case of W(CO)$_6$. The fragmented
Co(CO)$_4$ moieties exhibit a strong chemisorption on partially hydroxylated surfaces with an adsorption energy of -1.77 eV. Considering the above configurations as the initial setting for our MD simulations, Fig. 5 shows snapshots in 5 ps intervals of Co$_2$(CO)$_8$ adsorbed on a fully hydroxylated surface (upper panel) and of Co(CO)$_4$ fragments adsorbed on a partially hydroxylated surface (lower panel).

We observe a significant drift of Co$_2$(CO)$_8$ on the fully hydroxylated surface but as this is a larger molecule than W(CO)$_6$, the displacement is less pronounced than in W(CO)$_6$. This can be seen by comparing Fig. 5 where the COM movement of Co$_2$(CO)$_8$ on a fully hydroxylated surface is depicted, with Fig. 2(b) which is analogous to W(CO)$_6$. The calculated drifting radius of Co$_2$(CO)$_8$ within 20 ps is about 4 Å.

We now investigate the adsorption of the Co(CO)$_4$ species on partially hydroxylated surfaces (Fig. 5 lower panel). We would like to note that, unlike in the case of W(CO)$_5$, the Co(CO)$_4$ species in the gas phase possesses a tetrahedral structure which remains stable with a slight distortion on the partially hydroxylated surfaces (see $t = 0$ ps in Fig. 5 lower panel). During the MD simulations, the fragmented Co(CO)$_4$ species are localized on the surface but we observe severe changes in the orientation of CO ligands. In particular, we find that the Co(CO)$_4$ fragment bonded to the S2 site on the partially hydroxylated surface as shown in Fig. 5 (lower panel) shows bonding of the Co atom to surface oxygen as time evolves (compare $t = 0$ ps and $t = 5$ ps snapshots). A similar situation was suggested by Rao et al. during the adsorption of Co$_2$(CO)$_8$ on dehydroxylated MgO and on SiO$_2$ surfaces. However, within our simulation window, we only observe this effect for the Co(CO)$_4$ fragment bonded to the S2 site but not for the fragment bonded to the S1 site (see Fig. 5 (lower panel)) of the partially hydroxylated SiO$_2$ surfaces. Such a metal-substrate bond might be due to activation of the substrate surface by dehydroxylation.

C. Vibrations of W(CO)$_6$ and Co$_2$(CO)$_8$ on a SiO$_2$ surface

1. W(CO)$_6$ adsorbed on SiO$_2$ surface

We first start with the analysis of the vibrational spectrum of W(CO)$_6$ in the gas phase and the modes arising due to SiO$_2$ surfaces. We observe W−C stretching modes and (OC)−W−(CO) bending modes at 300-600 cm$^{-1}$, and C=O stretching modes at 1900-2150 cm$^{-1}$ for the gas phase W(CO)$_6$. The bending vibrations that involve W−(CO) atoms are seen at 60-100 cm$^{-1}$. All these nor-
normal modes are seen in both upper (power spectrum) and lower panel (finite displacement method) of Fig. 7. Also, in Fig. 8 we compare the power spectrum calculated for W(CO)$_6$ adsorbed on a fully hydroxylated surface with experimental infrared and Raman spectra measured for SiO$_2$ substrates so as to analyze the vibrations of SiO$_2$. We observe a good overall agreement between the calculated SiO$_2$ modes (Infrared and Raman active peaks at 100-200, 400-450, 700-800 and 1000-1100 cm$^{-1}$) and the experimental spectrum. Apart from SiO$_2$ modes, the additional modes in Fig. 8 are due to the surface adsorbed W(CO)$_6$ molecule (compare with Fig. 7). This analysis indicates that the adopted methodology to calculate the power spectrum could be used to describe the vibrations of W(CO)$_6$ adsorbed on SiO$_2$ surfaces.

In Fig. 9 we present the total power spectrum of W(CO)$_6$ adsorbed on fully and partially hydroxylated surfaces (Fig. 9 (a)) and only the vibrations of physisorbed W(CO)$_6$ and chemisorbed W(CO)$_6$ extracted from it which are compared with the vibrations (obtained by finite displacement method, shown as vertical bars) of isolated gas phase structures (Fig. 9 (b) and (c)). From Fig. 9 (b), we observe only minor changes in the vibrational modes of W(CO)$_6$. In particular, the CO frequencies observed between 1900 and 2110 cm$^{-1}$ (two of which are Raman active) for the gas phase of W(CO)$_6$ have also been observed upon its bonding to fully hydroxylated surfaces without being perturbed indicating that the molecule retains its gas-phase characteristics. These features are indeed similar to W(CO)$_6$ adsorbed on hydroxylated alumina ($\gamma$-Al$_2$O$_3$) surfaces where all the molecular vibrations are retained indicating the weak interaction between W(CO)$_6$ and the substrate.

From the analysis of the chemisorbed W(CO)$_6$ fragments on partially hydroxylated surfaces (see Fig. 9 (c)) we find, in addition to the shifts of several peaks, two new modes occurring at 700-800 cm$^{-1}$ and 1400-1600 cm$^{-1}$. Peaks at 700-800 cm$^{-1}$ which correspond to the Raman vibrations of $\beta$-cristobalite are now seen, because of precursor bonding to the surface. Also, the peaks at 1400-1600 cm$^{-1}$ appear as a result of oxygen (of CO) interaction with substrate Si atoms. Due to this interaction the bonding between tungsten and the carbonyl in W(CO)$_6$ gets affected leading to a considerable split of peaks in the regions that correspond to the vibrations of C−O (1900-2100 cm$^{-1}$) and W−C (300-600 cm$^{-1}$) bonds of W(CO)$_6$ respectively.

2. Co$_2$(CO)$_8$ adsorbed on SiO$_2$ surface

There have been some studies aimed at understanding the vibrations of Co$_2$(CO)$_8$ adsorbed on SiO$_2$. However, they are either limited to the carbonyl region of the spectrum or they observe immediate conversion of Co$_2$(CO)$_8$ to subcarbonyl species. We first show in Fig. 10 the analysis of the vibrational spectrum of Co$_2$(CO)$_8$ in the gas phase from power spectrum and the finite displacement method and both methods agree with each other and the existing reports. The two sets of vibrations in the region 1850-1900 and 2100-2000 cm$^{-1}$ are assigned to the stretching modes of the bridging and terminal CO ligands and the Co-Co stretching mode in Co$_2$(CO)$_8$ is observed between 200-250 cm$^{-1}$.

We now investigate the vibrations of Co$_2$(CO)$_8$ adsorbed on SiO$_2$ by analyzing the power spectrum. In Fig. 11 (a) we show the power spectrum of Co$_2$(CO)$_8$ on a fully hydroxylated surface compared to the power spectrum of Co$_2$(CO)$_4$ fragments on a partially hydroxylated surface. The extracted vibrations of physisorbed Co$_2$(CO)$_8$ and chemisorbed Co$_2$(CO)$_4$ from the total power spectrum are compared with the vibrations of the isolated molecules and are shown in Fig. 11 (b) and (c) respectively. The results indicate that Co$_2$(CO)$_8$ on fully hydroxylated surface preserves the gas-phase molecule characteristics (cf. Fig. 10 and Fig. 11 (b)).

In the case of Co$_2$(CO)$_4$ on partially hydroxylated surfaces, the overall agreement of the power spectrum with that of the gas phase Co$_2$(CO)$_4$ moiety is relatively poor because the Co$_2$(CO)$_4$ moieties exhibit strong structural distortions upon bonding to the SiO$_2$ substrate which is absent in the gas-phase computations. In the power spectrum, we find that the modes in the region 220-250 cm$^{-1}$ (Co-Co stretching) and the mode at approx. 1850 cm$^{-1}$ (bridging carbonyls) disappear owing to the fact that the Co$_2$(CO)$_8$ molecule fragments on partially hydroxylated surfaces. Further, two new modes in the 600-800 cm$^{-1}$ and 1400-1600 cm$^{-1}$ regions appear in this case (cf. Fig. 11 (a) and Fig. 11 (c)) illustrating the interaction of surface Si atoms with the carbonyls through the oxygen atom of the CO. However, this situation is only observed for one of the Co$_2$(CO)$_4$ moieties indicating that the fragmentated species behave differently. This is a consequence of the interaction of surface oxygen atoms with the Co atom observed for the fragment bonded to the S2 site (see Fig. 5).

D. Discussion

Investigation of surface adsorption and residence-time of the precursors on the surface is necessary for improving the understanding of deposition processes. The present study elucidates the vibrational footprints of precursors which interact through weak physisorption with fully hydroxylated surfaces and by strong chemisorption with partially hydroxylated surfaces.

The authors of Ref found that physisorbed W(CO)$_6$ molecules on TiO$_2$ surfaces completely desorb when the system is cooled down to room temperature. Our study finds no desorption of W(CO)$_6$ molecules on fully hydroxylated surfaces in the considered length of simulation. Also, Co$_2$(CO)$_8$ molecules on fully hydroxylated surfaces were reported to fragment spontaneously but, in our calculations on fully hydroxylated surfaces we didn’t observe any tendency to fragmentation or an indication of
chemisorption. Other effects (probably extrinsic) may be responsible for the experimental observations.

The bond variations of $\text{W(CO)}_6$ and $\text{Co}_2(\text{CO})_8$ adsorbed on fully hydroxylated surfaces indicate uniform fluctuations ($W-C$ and $C-O$ bonds) on either side (i.e., oriented towards the surface and the vacuum) of the molecule. On the contrary, the relative bond values of $\text{W(CO)}_6$ on partially hydroxylated surfaces show non-uniform variations indicating that certain bonds (i.e., $W-C$ bonds oriented towards the vacuum and the CO bonds towards the substrate) experience larger changes than others. Thus, for favorable conditions, the bond between $C$ and $O$ for the CO bonded to the surface might cleave leaving the surface $\text{Si}$ atoms terminated with oxygens. This situation is not observed in the present MD simulations but the higher ratio of carbon contamination with respect to oxygen (before atmospheric air exposure) in the EBID-obtained samples might provide an evidence for such a fact.\textsuperscript{17–21} Similarly, we found in the case of $\text{Co}_2(\text{CO})_8$ on partially hydroxylated surfaces, that surface oxygen atoms are involved in bonding to the dissociated surface species. This suggests that the removal of oxygen components from the deposits might not be an easy task to achieve. Presence of oxygen contamination is also expected to occur as a result of exposure of EBID deposits to the air. Thus the composition of the EBID deposits are determined to a large extent by the number of available active $\text{Si}$ sites (alternatively the degree of hydroxylation), the frequency of these precursor molecules approaching such a site and the exposure time to the environments. Furthermore, the surface defects observed in $\text{SiO}_2$ during the interaction of $\text{Co(CO)}_4$ with partially hydroxylated surfaces might also act as an active site for activating the approaching precursor molecule that could account for the facile dissociation of $\text{Co}_2(\text{CO})_8$ on $\text{SiO}_2$.\textsuperscript{31,66}

On partially hydroxylated surfaces our results for both $\text{W(CO)}_6$ and $\text{Co}_2(\text{CO})_8$ show that the fragmented species remain localized, thus blocking active sites on the surface. Therefore, further deposition should occur on the deposited layers and this may be likely the reason for the increase in height of the deposits as the irradiation time increases during EBID experiments.\textsuperscript{67}

IV. CONCLUSIONS

The purpose of this work was to model by means of \textit{ab initio} molecular dynamics simulations the dynamics of two precursor molecules adsorbed on fully and partially hydroxylated $\text{SiO}_2$ surfaces in order to achieve a better understanding of the microscopics of electron-beam induced deposition of nanostructures. Our results reveal that $\text{W(CO)}_6$ and $\text{Co}_2(\text{CO})_8$ molecules preserve their gas-phase bonding characteristics on fully hydroxylated surfaces. Apart from a considerable drift, only minor variations in the structure and vibrations is observed. Therefore spontaneous dissociation of these precursor molecules will not be possible on fully hydroxylated surfaces, unless some surface active sites are created by external forces.

For the case of $\text{W(CO)}_6$ and $\text{Co}_2(\text{CO})_8$ on partially hydroxylated surfaces, the fragmented species retain the chemisorbed character on the surface and we do not observe any reformation of the parent precursor moiety, but instead a slight tendency towards fragmentation. We also observe a smaller weakening of the surface-oriented CO bonds compared to vacuum-oriented CO bonds in $\text{W(CO)}_6$ and $\text{Co}_2(\text{CO})_8$ on partially hydroxylated surfaces. Therefore, conditions that favor the formation of active sites (in this case surface $\text{Si}$ atoms) are needed in order to have high efficiency in fragmentation and improve the metal content of the deposit.

The calculated vibrational spectra of these carbonyl molecule/ $\text{SiO}_2$ substrate systems show clear fingerprints to be detected experimentally. We propose therefore the consideration of such simulations as a route to experimentally distinguish the form in which precursors cover a substrate.

Finally, while the present simulations provide insights on the surface-precursor interaction, the investigation of other processes like surface-electron, precursor-electron and deposit-molecule interaction remains a challenge for future work.

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Appendix A: Estimation of the power spectrum

We use the power spectrum for analysing the vibrational characteristics of precursor molecules in the gas phase and for the entire system of precursors chemi- or physisorbed on a silica substrate.\textsuperscript{68} The power spectrum is defined as \textsuperscript{69,70}

$$I(\omega) = \frac{1}{2\pi} \int_{-\infty}^{\infty} C(\tau) e^{-i\omega \tau} d\tau,$$

(A1)

where $C(\tau)$ is the velocity autocorrelation function

$$C(\tau) = \langle \vec{v}(0) \cdot \vec{v}(\tau) \rangle = \lim_{T \to \infty} \left[ \frac{1}{TN} \int_0^T \sum_{l=1}^N \vec{v}_l(t) \cdot \vec{v}_l(t + \tau) dt \right].$$

(A2)

$\vec{v}_l(t)$ represents the velocity of atom $l$ at time $t$ for all $N$ atoms of the system. As the calculation of the time averages $\langle \ldots \rangle$ of Eq. (A2) is quite inefficient computationally, we use the Wiener-Khinchin theorem which guarantees
that the power spectrum can also be calculated by individually Fourier-transforming the velocities $\mathbf{v}(t)$ and summing the squares of the result:

$$I(\omega) = \frac{1}{N} \sum_{l=1}^{N} \left[ \frac{1}{2\pi} \int_{-\infty}^{\infty} \mathbf{v}(t)e^{-i\omega t} dt \right]^2. \quad (A3)$$

Finite trajectories, calculated with a finite time step $\Delta t$, only yield estimates to the power spectrum. Instead of the Fourier integral in Eq. (A3), discrete Fourier sums have to be calculated according to

$$V_{\mu}^{l\mu} = \sum_{j=0}^{l-1} v_j^{\mu} e^{2\pi i j k}, \quad (A4)$$

where the $v_j^{\mu}$ stand for the $\mu$ component ($\mu \in \{x, y, z\}$) of the velocity of atom $l$ at time step $j$, and $J$ is the number of time steps of the trajectory. The so-called periodogram estimate for the power spectrum is then defined for $J/2 + 1$ frequencies:\n
$$P^{l\mu}(0) = P^{l\mu}(f_0) = \frac{1}{J} |V_0^{l\mu}|^2,$$

$$P^{l\mu}(f_k) = \frac{1}{J} \sum_{j=0}^{J/2} |V_{j}^{l\mu}|^2 + |V_{J-k}^{l\mu}|^2, \quad (A5)$$

$$P^{l\mu}(f_c) = P^{l\mu}(f_{J/2}) = \frac{1}{J} |V_{J/2}^{l\mu}|^2.$$

The highest frequency $f_c = \frac{1}{2\Delta t}$ is called Nyquist frequency and is determined by the time step $\Delta t$ of the MD calculation. Thus, the finite time estimate of the power spectrum can finally be written as

$$I^J(\omega) = \frac{1}{N} \sum_{l=1}^{N} \sum_{\mu=1}^{3} P^{l\mu}(\omega). \quad (A6)$$

The function $I^J(\omega)$ will approach the true power spectrum $I(\omega)$ of the system in the limit $J \to \infty$, i. e. in the limit of infinitely long trajectories. The power spectrum estimates now provide us with information about the distribution of the vibrational energy of a molecule or a solid over the frequencies. Power spectra at different temperatures may be used to investigate the differences in the population of vibrational modes. The integral over the power spectrum corresponds to the kinetic energy of the system.

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