Nucleating a Different Coordination in a Crystal under Pressure:  
a Study of the B1-B2 Transition in NaCl by Metadynamics

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Here we propose an NPT metadynamics simulation scheme for pressure-induced structural phase transitions, using coordination number and volume as collective variables, and apply it to the reconstructive structural transformation B1/B2 in NaCl. By studying systems with size up to 64,000 atoms we reach a regime beyond collective mechanism and observe transformations proceeding via nucleation and growth. We also reveal the crossover of the transition mechanism from Buerger-like for smaller systems to Watanabe/Tolédano for larger ones. The scheme is likely to be applicable to a broader class of pressure-induced structural transitions, allowing study complex nucleation effects and bringing simulations closer to realistic conditions.

Structural phase transitions in crystals induced by pressure or volume are complex phenomena of great fundamental and practical importance. Most of them are reconstructive, thermodynamically first order, and involve crossing of free-energy barriers via a non-trivial concerted motion of atoms, representing a rare event. These transitions give rise to a number of important phases with unique properties such as, e.g., diamond created from graphite at high-pressure conditions. In the process of synthesis of such phases, kinetics plays a key role in determining the outcome of the transition, which might not necessarily be the thermodynamically most stable form, but rather a metastable one (e.g., after compression of silicon in the cubic-diamond structure to 11 GPa and decompression to ambient pressure, the BC8 phase is found) [1].

In the past two decades spectacular progress has been made in the prediction of crystalline phases, due to the advent of methods such as evolutionary search [2], random search [3, 4], particle swarm optimisation [5], minima hopping [6], etc. These approaches very effectively address the thermodynamics of the problem, identifying stable and metastable structures as global or local minima of the enthalpy surface. However, an understanding the mechanisms of the transitions, the pertinent barriers in the free-energy surface (FES) and the resulting kinetics still lags behind and more detailed information about the FES is needed to make progress. Commonly used theoretical approaches to uncover possible mechanisms and estimate energetic barrier per unit cell are based on geometric modelling [7–10], group-theory [11–14], phenomenological Landau theory [15], or, more recently, exploration of the FES [16–18]. However, by assuming collective transformation throughout the crystal, they cannot by construction assess the size of the nucleation region and determine the true nucleation barrier. A realistic simulation must therefore reach beyond collective behaviour and include nucleation. We note that one of the methods allowing mapping of FES, metadynamics (MetaD) [19] (for recent review see Ref. [20]) was successfully applied to the problem of crystallisation from liquid [21–26] which has a number of similar features to the problem of solid-solid transitions.

The application of MetaD to structural transitions in crystals started in Refs. [27, 28], using the h-matrix of the supercell vectors (similarly to the Parrinello-Rahman variable-cell MD [29, 30]) as the generic 6D collective variable (CV). This approach is efficient in inducing structural transitions in a number of systems [31–43], however, the use of a 6D CV essentially limits the use of MetaD to escaping FES minima and precludes the FES reconstruction. For an efficient reconstruction of the FES [44] CVs with dimensionality up to 3 are usually chosen. Moreover, the supercell-based CV by construction works well only for relatively small systems where transitions proceed via collective mechanisms but is unlikely to allow the study of nucleation in a large system. Several approaches addressing an autonomous construction or a choice of CVs have been proposed recently [45–54]. Applications of MetaD to structural transitions not based on the supercell CV are presented in Refs. [55–60].

We present in this Letter a simple and general scheme based on physically motivated CVs such as coordination number (CN) and volume (V) that should be applicable to the important class of pressure-induced structural transitions. This choice is primarily motivated by one of generic rules of high-pressure chemistry formulated by Prewitt and Downs [61–63] that states that pressure-induced transitions are typically accompanied by an increase of CN in the 1st coordination sphere. In a more general context, CN was proposed as a reaction coordinate in constrained MD in Ref. [64]. It was also employed in an early MetaD study of a structural transition in carbon [55] and in a MetaD study of the B1-B2 transition in colloidal clusters [65]. Thermodynamically, in first-order transitions an abrupt densification of the system takes place, with a jump in volume from a few % up to 10-20%. We show here that the combination of CN & V appears to provide an effective 2D CV able to drive pressure-induced structural transition.

We demonstrate the applicability of this scheme on the pressure-induced B1-B2 transition in NaCl, which represents a paradigmatic but also very complex example of a reconstructive transition [15]. It occurs at room temperature at $p = 26.6$ GPa and involves a volume drop of 5% [66]. Several theoretical collective mechanisms were proposed for this transi-
tion, falling essentially into two groups. The ones by Shoji [7], Buerger [8] and Stokes & Hatch [12] are mainly driven by lattice strain while the other class by Hyde & O’Keeffe [9], Watanabe et al. (WTM) [10] and Toledáno et al. [15] involves more shuffling of atoms [13]. Computational studies include overpressurized variable-cell MD [67, 68] and transition path sampling by Zahn & Leoni [69].

We describe NaCl by the well-known and computationally simple Born-Mayer-Huggins-Fumi-Tosi (BMHFT) potential [70, 71], which yields for the equilibrium transition pressure at \( T = 0 \) a value of \( p_{eq} = 19.25 \) GPa. Details of the simulation are provided in the Supplemental Material [72].

Structurally, the \( B1-B2 \) transition in NaCl is accompanied by the transfer of two ions with opposite charges from the \( 2^{nd} \) to the \( 1^{st} \) coordination shell, increasing the CN from 6 in \( B1 \) to 8 in \( B2 \). The average CN between the \( Na^+ \) and \( Cl^- \) ions can be calculated by means of a switching function as

\[
CN = \frac{2}{N} \sum_{i \in Na^+} \left( 1 + \left( \frac{r_{ij} - d_0}{r_0} \right)^6 \right)^{-1},
\]

where \( r_{ij} \) is the distance between the \( i^{th} \) cation and \( j^{th} \) anion and \( N \) is the total number of atoms. The choice of the parameters \( d_0 \) and \( r_0 \) requires some attention. The switching function should allow to clearly differentiate between the initial state (e.g., \( B1 \)), the transition state, and the final state (e.g., \( B2 \)) [20]. Moreover, its slope should be sufficiently high at the positions of the radial distribution function (RDF) peaks of the \( B1 \) phase corresponding to the first and the second coordination sphere in order to drive an easy exchange of ions between the two spheres. A suitable switching function meeting both requirements is shown in Fig. 1 [79].

For a system in the \( B1 \) phase with 512 atoms, we performed both MetaD with only CN as well as one with CN & V as CVs. In both versions, both forward and reverse transitions can be seen; see Supplemental Material [72], Figs. S4-S7. However, the character of the CN evolution in the two cases is different. When only the CN is used as CV, even after the first forward and reverse transitions, the system continues to jump between the two phases indicating that the CN does not have full control over the system. On the other hand, when V is added, the evolution of CN and V after the 1\( ^{st} \) transitions becomes much more diffusive. This can be seen in the cross-correlation between CN and V; see Supplemental Material [72], Figs. S8 and S9. We conclude that CN and V thus represent a good choice of CVs. However, the reconstructed FES in Fig. 2 shows that the structural phases are represented as rather long and narrow valleys. The soft direction (SD) represents “breathing” of the crystal preserving the structure, while the perpendicular - hard one (HD) represents a direction of structural change. To improve sampling of such a shaped FES, we introduce a rotation of CVs with origin at the equilibrium point \([CN, V](p, T)\) of \( B1 \). Deposited Gaussians thus respect the shape of the valleys, being wide in the SD and narrow in the HD. We first rescale CN and V with respect to \( B1 \) and then rotate them by an orthogonal transformation, whose components are orthonormal eigenvectors of the covariance matrix of the rescaled coordinates. The covariance matrix was obtained from a short 200 ps unbiased NPT MD simulation at given pressure \( p \) and temperature \( T \) in the \( B1 \) phase. A detailed description of the approach is provided in the Supplemental Material [72].

For illustration, the evolution of the structure of the system is added, the evolution of CN and...
across the transition at 40 GPa is shown in the Supplemental Material [72], Fig. S21. The intermediate state (d) is similar to the B33 structure that appears in some theoretically proposed mechanisms (see later). It is seen that the whole system first transforms to this transient short-living state which quickly converts to B2, pointing to a collective mechanism of the transition. For some alkali-halides, a two-step transition mechanism through the intermediate bulk B16 or B33 phases was proposed by Toledano et al. [15].

The presented scheme can be readily applied to larger systems allowing to study precursor effects, nucleation and growth and access to information about free energy, size, shape, and structure of the critical nucleus. We performed the simulations at T = 300 K for various system sizes, N = 512, 4096, 13824, 32768 and 64000 atoms. Because the size of the critical nucleus at 20 GPa, close to equilibrium, is expected to be very large, we chose to work at pressures of 30 and 40 GPa. We note that non-classical nucleation theories (see later) [80] predict also divergence of the size of the critical nucleus upon approaching the point of dynamical instability for our system we found this at 60 GPa.

In Fig. 3 we show the evolution of the transition barrier as a function of system size for two values of pressure [81]. For p = 30 GPa the curve appears to grow in a nearly linear manner up to N = 13824, indicating that even at this moderate overpressurization very large system sizes are necessary to properly accommodate the large critical nucleus. For systems smaller than 4096 atoms, the barrier per atom agrees well with the estimate based on the static Buerger mechanism (see Supplemental Material [72], Fig. S3), showing that the transition proceeds via a collective mechanism. The barrier height in the thermodynamic limit must be larger than 10^2 eV, revealing that homogeneous nucleation in such a regime is physically impossible. At the higher pressure of 40 GPa the curve appears to eventually converge to a value above 90 eV, still too high for a physical transition. Since experimentally the transition at 300 K occurs at p = 26.6 GPa [66], it must be assisted by extrinsic factors such as lattice defects [83–87], dislocations [88–95], grain boundaries [96], surfaces [83–85], or non-hydrostatic pressure [97, 98]. This observation is similar to the one found for nucleation of melting [94], crystallisation of ice [99] and transformation of graphite to diamond [87].

The slow convergence of the barriers can be explained by the presence of long-range (~ 1/ρ^3) elastic strain fields [83–85, 100–102]. We note that the elastic energy of the nucleus and surrounding lattice [83–85, 102] is taken into account in non-classical nucleation theory [80, 102–115] but is missing in standard static approaches [7–15] which assume a strictly collective character of the transformation with no interface between the parent and the new phase. It would be fully taken into account in simulation provided the system is sufficiently large.

We now focus on the structural aspects of the transition. In Fig. 4 we see the critical nucleus (determined as the first timestep from which an unbiased MD proceeds towards the B2 basin) in the system of 64000 atoms at 40 GPa. Even at this system size, the critical nucleus represents a cylinder extending across the periodic boundary conditions (PBC) along one dimension. In all simulations, the nucleus formation starts by creation of strain in a large region of the lattice that extends across the PBC. In this region the primary nucleus is eventually formed, followed by the creation of a secondary nucleus. At all system sizes and pressures presented, the size of the critical nucleus is not small compared to the system size and PBC artefacts are present. The dependence of shape and size of the critical nucleus on system size and pressure can be found in the Supplemental Material [72].

We further analyzed the detailed transformation mechanism and its dependence on pressure and system size. For convenience, we provide in the Supplement Material [72] a review of previous results found in the literature. The four idealised collective mechanisms proposed in Ref. [13] can be characterised based on the transformation of the local environment of each atom. B2 is formed from B1 after adding two second neighbours of the opposite type to the first coordination shell. All eight of these second neighbours are corners of the conventional fcc cell with a given atom in the centre. If both these additional second neighbours join at the same time and originally form an edge of the conventional fcc cell, one finds the WTM mechanism [10]. On the contrary, if they are located at opposite corners, one finds the Buerger mechanism [8].
FIG. 4. (Color online) Critical nucleus of the \(B_2\) phase (with the shape of a cylinder) in the 64000 atoms system at 40 GPa. (a) Critical nucleus (axis of the cylinder perpendicular to the plane). (b) Perpendicular view of the critical nucleus where only atoms with coordination \(\geq 6.5\) are shown for clarity. (c) Shear around the critical nucleus in the plane perpendicular to the axis of the cylinder. (d) A localized nucleus with the shape of an ellipsoid (red), 41.7 ps prior to the critical nucleus frame (a). The ellipsoid grows into the cylinder along the axis in which the strain field extends across the PBC (distortion along horizontal direction). View (d) is perpendicular to both (a) and (b). The pictures were produced using OVITO [116].

FIG. 5. (Color online) WTM-like mechanism (Toledano) observed during the formation and growth of the critical nucleus in the 64000 atoms system at 40 GPa. The central atom and its 1st and 2nd coordination shell are shown without distinguishing Na and Cl atoms while atoms forming the conventional unit cell of the emerging \(B_2\) phase are shown in red. All atoms are within the critical nucleus and the cutoff for bonds is set to 3 Å. The position of the central atom and the point of view are fixed. (a) Atoms in the \(B_1\) lattice. (b)-(e) Atomic configuration at specified times where (d) corresponds to the critical nucleus. The pictures were produced using OVITO [116].

Adding the two atoms independently in two steps instead results in the Toledano [15] (modified WTM) and Stokes and Hatch (modified Buerger) mechanisms, both of which create an intermediate \(B_{33}\)-like structure [117]. This analysis allows the possibility of different parts of the system transforming at different times via distinct mechanisms. It was performed for all systems considered in the Supplemental Material [72]. The WTM and Toledano mechanisms are facilitated by the local lattice shear strain which amounts to a compression along a (110) direction. This breaks the cubic symmetry and brings four out of eight second neighbours closer to the central atom [see Fig. 5(b)]. It is likely that an application of such uniaxial stress in experiments would reduce hysteresis and facilitate the observation of the transition closer to the thermodynamic transition pressure.

In our simulations, for systems up to 4096 atoms, the dominant mechanism is related to the creation of the intermediate (bulk) \(B_{33}\) structure that subsequently transforms to \(B_2\) via the Stokes and Hatch mechanism. For a system size of 4096 atoms, only parts of the system locally transform through the Stokes and Hatch mechanism, see Supplemental Material [72], Fig. S28(e). Finally, for systems larger than 4096 atoms, all atoms within the critical nucleus transform via the Toledano mechanism. This involves the displacement of planes, as can be seen in Fig. 5(c). In larger systems the Stokes and Hatch mechanism would cost too much energy and therefore nucleation via a zig-zag pattern (WTM or Toledano mechanism), which causes less strain, appears to be preferable. Nuclei are surrounded by 7-coordinated atoms, but this layer does not resemble \(B_{33}\)-like structures in large systems.

Our approach is likely to work for a broader class of pressure-induced structural transitions and uncover their microscopic mechanisms in the regime of nucleation and growth, including calculation of free-energy barriers. It might represent a bridge between atomistic modelling of structural phase transitions and effective phase-field theories [80, 102–115, 118–127]. The use of simple and physically naturally motivated CVs allows a MetaD simulation without prior knowledge of the transition and the final states. This would enhance the predictive value, in particular in cases in which the final state might be either a metastable one or one which is stabilized by entropy and therefore falls beyond the reach of standard \(T=0\) structural prediction methods. It is essential to access long time (\(\sim 10\) ns) as well as length scales...
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