

MOLECULAR DYNAMICS SIMULATIONS OF DISCRETE BREATHERS IN THE NICKEL HYDRIDE

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Computational assessment of the discrete breathers, i.e. spatially localized and periodic in time nonlinear excitations in crystals, is performed in nickel hydride with an even stoichiometry by means of molecular dynamics simulations. The breathers consisting of hydrogen and metallic atoms were excited following the experience obtained earlier by modelling the breathers in pure metallic systems. Stable breathers were found in the nickel hydride system for the hydrogen atoms oscillating along $\langle 100 \rangle$ and $\langle 111 \rangle$ polarization axes. Two types of the stable breathers involving single oscillating hydrogen and a pair of hydrogen atoms beating in antiphase mode were discovered. Analysis of the breather characteristics reveals that its frequency is in the phonon gap or lying in the optical phonon band of phonon spectrum near the upper boundary. Analysis of the movement of atoms constituting the breather was performed to understand the mechanism that enables the breather stabilization and long-term oscillation without dissipation its energy to the surrounding atoms. Thus, our analysis provides an explanation for the existence of the long-living stable breathers in nickel hydride.

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INTRODUCTION

Discrete breathers (DBs) are the spatially localized and periodic in time nonlinear excitations of the lattices [1]. The two critical components such as nonlinearity and discreteness must combine to make a DBs to exist. Nonlinearity allows the strongly excited localized modes to have a fundamental frequency outside the spectrum of small amplitude oscillations (phonons). In crystalline solids, DBs may result in a *catalytic acceleration* of chemical and nuclear reactions in their vicinity [2–4]. The triggering mechanism of DBs is based on *irradiating* a crystal with fast particles or electromagnetic waves that could excite DBs with a minimum production of phonons, i.e., without substantial heating of the crystal lattice [5].

The majority of pioneering theoretical studies on DBs dealt with the idealized one- or two dimensional nonlinear lattices of coupled oscillators interacting via simplified pairwise potentials with polynomial nonlinearity. It has been shown that integrable nonlinear lattices, such as Ablowitz-Ladik lattice [6, 7], support exact solutions in the form of spatially localized excitations. For instance, Bogdan has obtained the exact DB solution for the Hirota lattice equation [8]. In further publications the exact Hamiltonian dynamics [9] and pair interactions processes [10] of these excitations were investigated analytically. However a number of exactly solvable and physically important nonlinear discrete lattices is rather small. That is why it was important to prove rigorously the existence and stability of DB in non-integrable nonlinear chains [11–13].

The role of DBs has been extensively discussed in the fields of nonlinear optics [14, 15], superconducting Josephson junctions [16, 17], electric circuit arrays [18, 19], micromechanical and macromechanical cantilever arrays [20, 21], mechanical systems [22, 23], granular crystals [24], etc.

Discrete breathers in crystalline solids have been actively studied in recent years in relation with their physical and mechanical properties [25, 26]. The atomic-scale nature of DBs, quantum effects behind and scarcity of events make the observation and experimental registration of these excitations challenging. Nevertheless, there are a number of experiments demonstrating the existence of DBs in different crystals. In the experiments with the nonlinear halide-bridged mixed-valence quasi-one-dimensional transition metal complex PtCl [27] strong evidence for the existence of DBs has been found by measuring resonance Raman spectra. The redshift in the resonances indicated the localized excitation of Pt-Cl bonds. Manley et al. [28, 29] have reported an experimental observation of gap DBs (i.e. DBs with the frequency laying in the gap of the phonon spectrum of bi-atomic system) in thermodynamic equilibrium in NaI crystal. In a series of experimental works, evidence supporting existence of DBs in alpha-uranium has been obtained [30–32]. Recent reviews [33–35] summarize the results of the theoretical and experimental investigations and the molecular-dynamics simulations of DBs in two- and three-dimensional crystals.

Particular importance attributed to DBs is defined by their potential role in determining rates of non-equilibrium processes in real material (e.g. chemical diffusion, catalysis, defect migration and annealing, activation of dislocations and other atomic-scale mechanisms) [36–38]. DBs can enhance heat capacity by storing some energy [30]. They can also reduce thermal conductivity by scattering phonons [39]. On the other hand, mobile DBs can enhance heat conductivity by transporting energy [31].

In the recent years, DBs-mediated effects in solid state physics and materials science have been the focus of a rapidly growing number of studies based on more

realistic atomistic models of crystals. With the appearance of powerful computational methods, the research focus has shifted to simulations in 3D lattices using realistic many-body potentials and investigating properties of breathers at finite temperature [40–42] or in non-ideal crystals containing lattice defects [43]. Molecular dynamics (MD) was applied to study breathers not only in single-atomic systems like covalent silicon and diamond [44, 45], pure metals [43, 46–50] but also in the ordered alloys [51–53], as well as in ionic crystals with NaCl structure [54–57]. Not only MD [58–64] but also *ab initio* simulations support existence of DBs in 2D crystalline structures such as graphene and graphane [65, 66].

The existence of breathers in NiH systems may have a number of important applications for the hydrogen technology. Hydrogenation is a chemical reaction between molecular hydrogen and another compound or element, usually in the presence of a *catalyst* such as nickel or palladium. There is no single theory of catalysis, but only a series of principles to interpret the underlying processes. An important parameter of the reaction kinetics is the activation energy, i.e. the energy required to overcome the reaction barrier. The lower is the activation energy, the faster the reaction rate, and so a catalyst may be thought to reduce somehow the activation energy. Dubinko et al [36–38] have suggested that DBs arising in the vicinity of a reaction cite result in a time-periodic modulation of the potential landscape, which can be described by a combination of time-periodic modulation of the activation barrier height and shape. The height modulation effect was shown to result in a drastic acceleration of escape from the potential well driven by thermally-activated ‘jumps’ over the reaction barrier [36, 37]. The shape modulation effect was shown to result in a strong acceleration of tunnelling through the potential barrier at sufficiently low temperatures, where the reaction rates are controlled by quantum rather than by thermal fluctuations [38]. These results support the concept of DB mediated catalysis at both high and low temperatures, which shows the need for more detailed atomistic modeling of DBs in various physical systems.

Here, we investigate the existence of DBs in the nickel hydrides by means of modern atomistic computational tools involving usage of *ab initio*-derived many body interatomic potentials, which accurately reproduce a number of features of the studied hydrides. By performing MD simulations at zero and finite temperature we investigate the existence of breathers and their characteristics regarding the location of their frequencies in the phonon spectrum. As this is the first step, we take an idealized structure of the perfectly stoichiometric hydrides with chemical formulas NiH having the lattice structure equivalent to NaCl ionic crystal.

COMPUTATIONAL DETAILS

The assessment of DB’s characteristics in NiH has been performed using classical MD simulations in 3D periodic crystals. The NiH structure was generated using Latgen code supplied with LAMMPS package [67]. We have modelled the stoichiometric hydrides

where fcc hydrogen sub-lattice occupies octahedral positions in the fcc metallic (palladium or nickel) sub-lattice.

MD simulations were carried out by employing the many-body type interatomic potentials derived on the basis of available experimental data and *ab initio* results. The NiH potential was taken from [68]. These potential was developed to account for specific properties of H in Ni crystal, overall showing a good agreement with available experimental and *ab initio* data as what concerns basic properties. In this work we did not involve density functional theory calculations as yet, because it imposes high computational demand, given that a high number of various configurations must be studied. MD simulations with realistic many-body potentials are known to be a good compromise to encompass the research, while *ab initio* methods can be applied in future to validate the MD results.

MD simulations were done in the virtual 3D-periodic crystals with three principal axes oriented along the $\langle 100 \rangle$, $\langle 010 \rangle$, and $\langle 001 \rangle$ directions, respectively. The size of the MD supercell was chosen to be 10, 10, and 10 atomic planes along x, y, and z. Such MD box is sufficiently large to avoid the self-interaction of a highly localized DB via periodic boundaries.

A DB was created in the crystal by setting an initial displacement along a selected close packed direction, where the breather is mostly expected to be stable, following the example of earlier studies in pure metals [46–50]. The key feature of the excitation of the breather is the choice of the initial displacement direction and selection of configuration involving one, two or a higher number of initially excited atoms. The central offset displacement will be referred in the following as d_0 . Its variation determines the DB amplitude and frequency and, ultimately, its oscillating lifetime. In practice, such an initial condition of the DB excitation corresponds to the recoil processes which can be the result of exposure to high-energy particle beam or a shock wave.

Prior to initiating a DB a full relaxation to zero pressure and zero temperature was achieved by the conjugate gradient method. All initial velocities of atoms were set to zero. Two types of breathers were probed in this study. The first type implies the excitation of a single atom, such that the oscillation of a hydrogen atom between two metallic atoms realizes. Thus, such configuration is referred to as “M-H-M” breather, where M stands for Metal (in our case Ni). A schematic representation of its initial configuration is shown in Fig. 1. One hydrogen atom, originally located in the centre of the picture, is displaced to the left. In what follows, as we shall discuss characteristics of DBs, the analysis of the distance between the H atom constituting the DB and hydrogen/metallic nearest neighbour atom will be performed. These two distances are shown in Fig.1 as $r(\text{H-H})$ and $r(\text{H-M})$. Fig. 2 shows the initial configuration of the H-M-H breather which consists of H1 and H2 atoms beating in the antiphase and being separated by M2 atom.

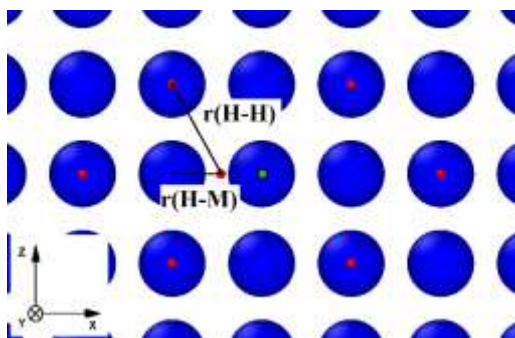


Fig. 1. Schematic representation of the initial configuration of the M-H-M type breather. The length of the H-M and H-H bond, shown respectively as $r(H-M)$ and $r(H-H)$, is analysed during the simulation. Large blue balls showing metallic atoms and small red balls showing hydrogen atoms are drawn at a scale of ionic ratio for nickel (Ni) and hydrogen (H). Green small ball in the center represents the equilibrium position of the H atom

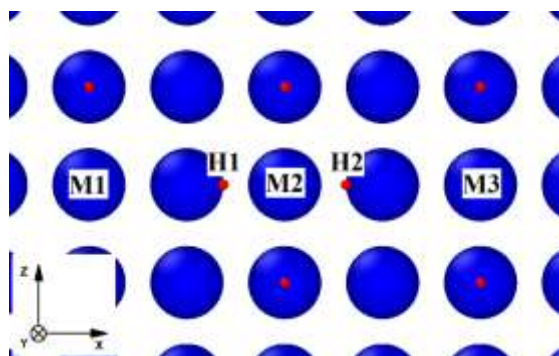


Fig. 2. Schematic representation of the initial configuration of the H-M-H type breather. Blue large balls showing metallic atoms and red small balls showing hydrogen atoms are drawn at a scale of ionic ratio for nickel (Ni) and hydrogen (H)

Each type of breather was excited by gradually increasing d_0 up to the situation when the point defect was generated as a result of the evolution of atomic motion. In order to quantify characteristics of stable DBs, the profiles of atomic positions and potential energy function of atoms constituting the breather were analyzed. The time evolution of the displacement of H and M atoms (constituting the breather) from their equilibrium positions was used to define the eigen frequencies of the DBs. To complete the analysis of the breather characteristics, the phonon density of states for the NiH systems were derived using the procedure proposed in [69]. In [69], the methodology to calculate the phonon dispersion of a crystal using molecular dynamics simulations is proposed and implemented as an open source routine to be plugged to LAMMPS code. The dynamical matrix is used to compute the phonon spectra by evaluating its eigenvalues. The dynamical matrix is constructed by recording the displacements of atoms during a representative MD run, making the use of the fluctuation–dissipation theory.

The finite temperature simulations were also performed to investigate potential existence of breathers at elevated temperature, following the methodology proposed in Refs. [40, 41]. Those works exploited an

idea of spatial and temporal temperature (i.e. atomic velocity) analysis. From such a combined analysis, the breathers were deduced on the basis of a specific numerical criterion which reflects local temperature within a certain timespan. In this case, the MD simulations of NiH at finite temperature using NVE micro-canonical ensemble were carried at the temperature of 300 K. From the finite temperature simulations, the velocity profile of all the atoms in the system was analysed and post-processed to deduce the presence of the breather according to the definition introduced in [41].

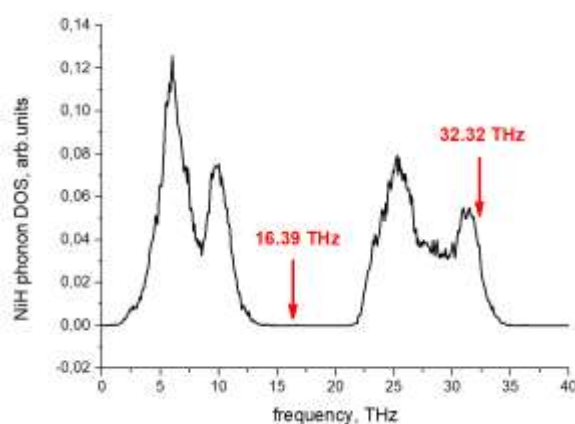


Fig. 3. Phonon density of states for the NiH lattice. The frequency of stable (i.e. long-living) breathers found in this work are indicated on the figure by red arrows

RESULTS

MODELING DISCRETE BREATHERS AT ZERO TEMPERATURE

The phonon density of states for NiH, obtained at low temperatures (1 K), are presented in Fig. 3. It can be seen that for the NiH lattice there is a gap in the 13.5...21.5 THz frequency interval and the upper edge of the phonon spectrum is 34.5 THz. Thus, we expect to observe breathers in NiH lattice with frequencies either lying in the gap or exceeding the upper limits of the phonon spectra.

As was explained earlier, the breathers with different polarization axes were excited by gradually increasing the initial displacement d_0 (see Figs. 1 and 2). The general trend for the stability of the breather with increasing d_0 , that was observed for all polarization axes, is presented in Fig. 4 by the example of the $\langle 100 \rangle$ H-M-H breather in Ni-H system. If the excitation amplitude is too small (Fig. 4,a), the H oscillation quickly decays and the DB energy is quickly dissipated to the surrounding atoms, i.e. to the lattice. Given a sufficient offset displacement distance, the oscillation of H atom is stabilized after a short transient period. An example of the evolution of the atomic coordinates of five atoms forming the breather is shown in Fig. 4,b. It is seen that shortly after the breather is launched, the M1 and M3 metallic atoms bounding the oscillating H1 and H2 atoms slightly displace to extend a volume for the oscillating H atoms. Centers of vibration of the two H atoms vibrating out-of-phase also shift from the equilibrium positions. Then, the system comes to quasi-

stationary motion and such a breather performs its stable oscillation up to 400 cycles of full vibration periods at nearly constant amplitude, without losing its energy. Analysis of the atomic motion shows that such a breather represents fully symmetric system where M2 (see Fig. 2) is the center of the symmetry. This long-living breather was analyzed in details to calculate the kinetic energy stored in this defect. The kinetic energy of the breather was calculated as the average kinetic energy of five atoms (two hydrogen and three metallic atoms) forming the breather. It was found that the kinetic energy of the breather shown in Fig. 4,b is 1.23 eV.

Increasing the initial amplitude further leads to the displacement of H atom(s) from their original sites and subsequent creation of point defect(s). Such example is shown in Fig. 4,c, where one of H atoms is eventually displaced out of its original site. Thus, the excitation of the stable breather requires careful choice of d_0 as was originally anticipated.

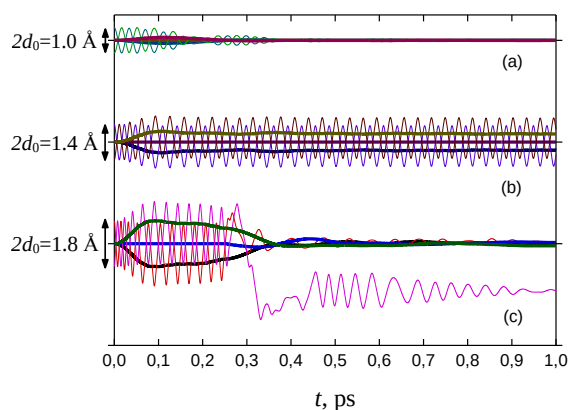


Fig. 4. Evolution of the displacement of atoms from their equilibrium positions for the case of H-M-H $\langle 100 \rangle$ breather in NiH system: small ($d_0 = 0.5 \text{ \AA}$) (a), middle ($d_0 = 0.7 \text{ \AA}$) (b), and large ($d_0 = 0.9 \text{ \AA}$) (c) initial displacement, see text for details. Thick/thin lines correspond to the Ni/H atoms displacements from the equilibrium positions. The arrows next to the ordinate show the absolute value of the atomic oscillation relative to the equilibrium position\

The same type of calculations were repeated in the NiH for $\langle 110 \rangle$ and $\langle 111 \rangle$ polarization axes. It was found that depending on specific polarization axis, optimal d_0 and therefore resulting DB frequency differ. Moreover, for some of the polarization directions the excitation of both types of the breathers was not successful. Hence, depending on the polarization axis, different types of breathers may exist. Analysis of the oscillations of the long-living breathers in NiH system revealed that 16.39 and 32.32 THz are the frequencies of the most stable gap-type and high frequency type M-H-M breathers, respectively.

Several examples of the atomic oscillations, which were found to be the most stable, are provided for the $\langle 100 \rangle$ (Fig. 5), $\langle 110 \rangle$ (Fig. 6), and $\langle 111 \rangle$ (Fig. 7) polarizations of the M-H-M breathers, respectively. The animation of the atomic motion near the location of each of the breather, shown on those figures, is included in the supplementary material. An example of the high

frequency M-H-M breather with $\langle 100 \rangle$ polarization axis is provided in Fig. 5, where the displacement of hydrogen from its initial position is shown as the function of time.

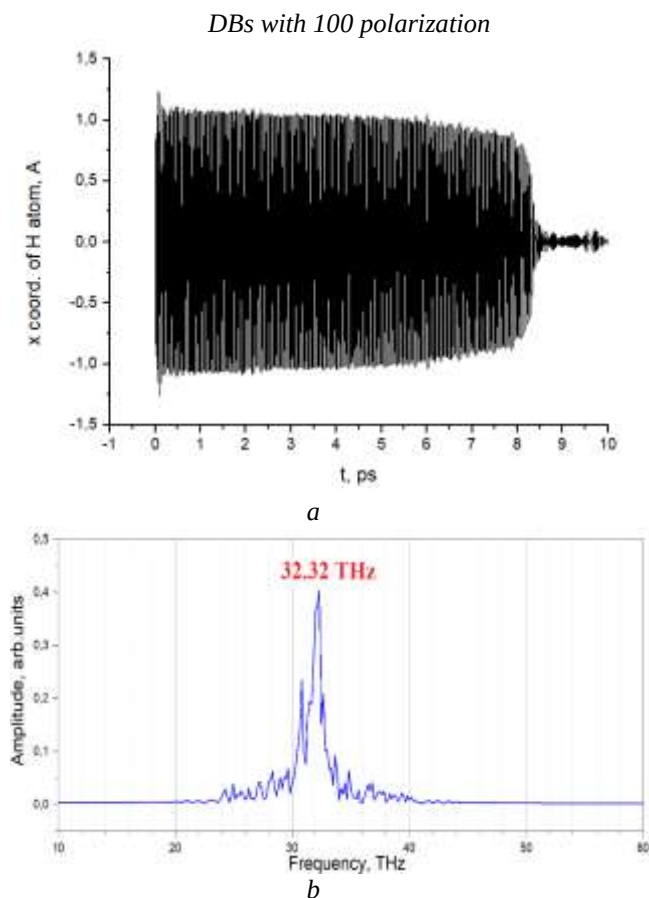


Fig. 5. Oscillation of a hydrogen atom forming the M-H-M $\langle 100 \rangle$ breather in NiH lattice initiated with $d_0 = 0.84 \text{ \AA}$. The resulting oscillation amplitude is $1.05 \text{ \AA}$, oscillation frequency is 32.32 THz, oscillation period is 0.03 ps and lifetime is 8.3 ps, which is equivalent to 276 oscillation periods (a). Fourier spectrum constructed using the trajectory presented in Fig. a. The position of the peak corresponds to the oscillation frequency of 32.32 THz (b)

By analyzing the atomic trajectory over the stable oscillation time interval, the following breather parameters were deduced: the oscillation amplitude is $1.05 \text{ \AA}$, oscillation frequency is 32.32 THz, oscillation period 0.03 ps and lifetime is 8.3 ps, which is equivalent to 276 oscillation periods. Another example of the M-H-M breather, which was excited in $\langle 101 \rangle$ direction, is presented in Fig. 7. This breather decayed rather quickly within less than 1 ps, however, its oscillation frequency is found to be 18.88 THz, thus it is the gap-type breather. The variation of d_0 did not result in the formation of any other stable breather configurations for this excitation direction. Changing the excitation direction to $\langle 111 \rangle$, we found a very stable M-H-M gap-type breather. The oscillation profile is given in Fig. 8, where we present the zoomed area for 1 ps simulation time span and 25 ps segment. This breather was seen to be stable for more than 100 ps. The resulting oscillation

frequency is 16.39 THz and period 0.061 ps, proving that this is gap-type breather.

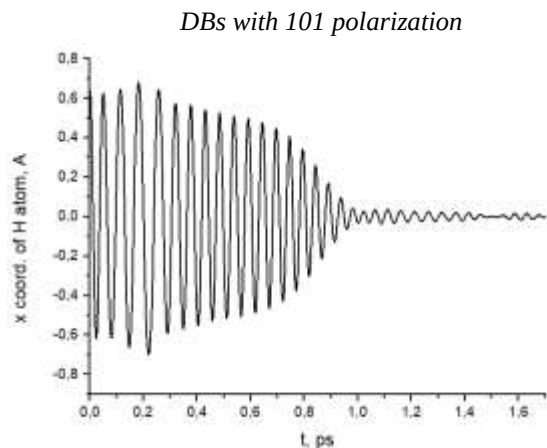


Fig. 6. Oscillation of a hydrogen atom forming the M-H-M $\langle 101 \rangle$ breather in NiH lattice with $d_0=0.9 \text{ \AA}$. The resulting oscillation amplitude is $0.5 \text{ \AA}$, oscillation frequency is 18.88 THz, oscillation period is 0.053 ps and lifetime is 0.86 ps, which is equivalent to 15 oscillation periods

To further rationalize the reasons for the difference in the oscillation behavior of atoms in nickel and palladium hydrides we calculated the effective potential pair profile for hydrogen-hydrogen and metal-hydrogen and calculated the interaction force for the M-H-M type DB polarized along $\langle 100 \rangle$ direction.

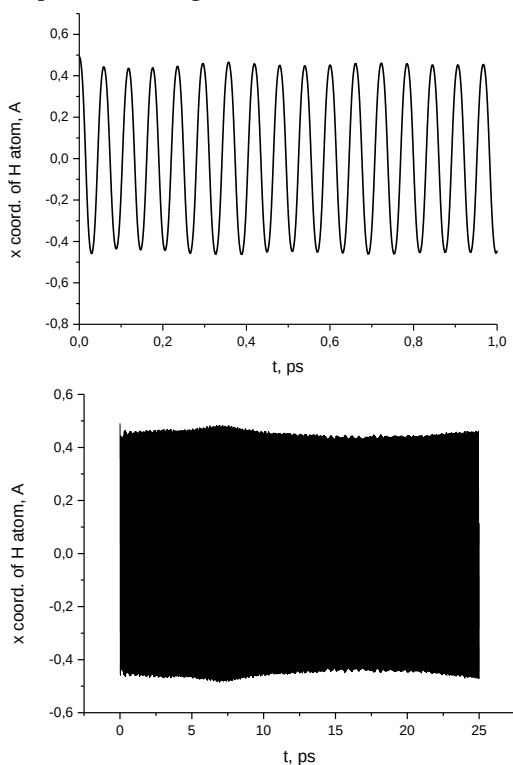


Fig. 7. Oscillation of a hydrogen atom forming the M-H-M $\langle 111 \rangle$ breather in NiH lattice with $d_0=0.85 \text{ \AA}$. The resulting oscillation amplitude is $0.79 \text{ \AA}$, oscillation frequency is 16.39 THz, oscillation period is 0.061 ps and lifetime is higher than 100 ps ~ 1640 oscillation periods (the simulation run was stopped before the breather oscillation decayed)

This is shown in Fig. 5 for the NiH. Each figure contains the profile of the effective pair interaction force i.e. negative derivative of the effective pair potential. The effective pair potential is calculated using molecular static approach by moving an atom in the lattice, while keeping the surrounding neighbors constrained in their positions. The vertical lines labeled as $r_{\min/\max}^{M-H}$ and $r_{\min/\max}^{H-H}$ denote the minimum M-H/H-H distance which realizes as the breather oscillates in the steady-state mode, where M denotes nickel (Ni). Given the spatial boundaries, specified by vertical lines on the figures, one can understand the evolution of the force acting on the oscillating H atom during the whole oscillation cycle of a DB. Positive (negative) force in Fig. 8 is repulsive (attractive).

Considering the case of the DB in NiH lattice (see Fig. 8). Note that within the span of H atom oscillation, the H-H interaction is negligibly weak as compared to the Ni-H interaction.

Thus, the stability of a DBs oscillation process in NiH can be qualitatively explained by the short-range H-H interaction in this lattice.

SUMMARY AND CONCLUSIONS

In this work, we have analysed the existence of discrete breathers in NiH. The assessment was performed by employing 3D atomistic simulations with realistic many-body interatomic potentials, developed to account for specific properties of hydrogen in these metallic lattices. The simulations were performed in the fully relaxed lattice at initial temperature of zero Kelvin, and at elevated temperature using conventional MD simulations. The simulations performed at zero temperature were used to explore typical characteristics of the hydrogen breathers by varying the polarization axis, intrinsic oscillation energy and hydrogen-metal configuration defining the breather structure (i.e. H-M-H or M-H-M type of oscillation).

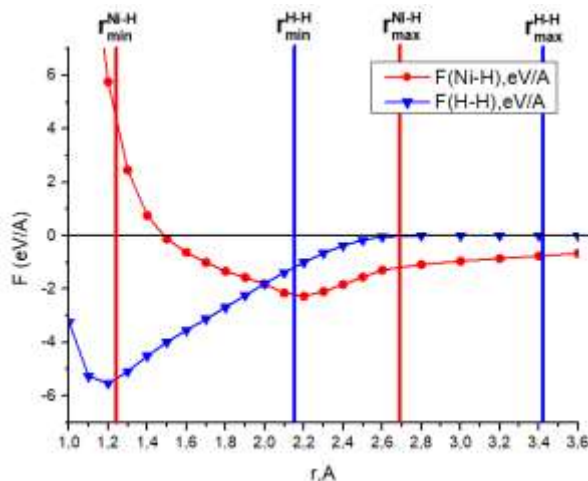


Fig. 8. Interaction force of the Ni-H and H-H atomic pairs as a functions of the distance between the atoms embedded in NiH lattice for the case of M-H-M DB polarized along $\langle 100 \rangle$ direction. Negative values correspond to the attractive interaction, while the positive values to the repulsive interaction

Parametric study was applied to scan a large range of the excitation amplitudes (related to the intrinsic oscillation energy) to deduce the most stable breathers, where stability refers to the number of oscillations prior the breather is absorbed by the lattice. To classify the type of the breather, the phonon density of states for both hydride systems were calculated using the selected set of potentials. By comparing the frequency of the breathers as measured in zero temperature simulations with the density of states, we conclude that both gap-type and high frequency breathers exist in NiH system.

On the basis of the results and discussion provided above it is possible to state the following:

(i) Both, gap-type and high frequency breathers are discovered in the nickel hydride system using classical molecular dynamics simulations with reliable many-body interatomic potentials. Depending on the polarization axis the breather has the frequency that falls in the gap of the phonon spectrum, or its frequency is near the upper phonon band edge.

(iv) With respect to the potential technological applications these results are important because breathers in NiH systems can act as catalysts of various reactions of technological importance [36–38, 70–72].

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МОЛЕКУЛЯРНО-ДИНАМІЧНЕ МОДЕЛЮВАННЯ ДИСКРЕТНИХ БРИЗЕРІВ У ГІДРИДІ НІКЕЛЮ

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Обчислювальна оцінка дискретних бризерів (також відомих як власні локалізовані моди) виконувалась у гідриді нікелю з рівною стехіометрією за допомогою моделювання молекулярної динаміки. Бризери, що складаються з атомів водню та металу, були збуджені після досвіду, отриманого раніше шляхом моделювання бризерів у чистих металевих системах. У системі гідриду нікелю виявлено стійкі бризери для атомів водню, що коливаються вздовж осей $\langle 100 \rangle$ і $\langle 111 \rangle$. Виявлено два типи стабільних бризерів, що включають одиничний коливальний водень і пару атомів водню, що б'ються у протифазному режимі. Аналіз характеристик бризера показує, що його частота знаходиться у фононному проміжку або лежить в оптичній фононній смузі фононного спектра поблизу верхньої межі. Аналіз руху атомів, що утворюють бризер, був проведений для зрозуміння механізму, який забезпечує стабілізацію бризера та довгострокові коливання без розсіювання його енергії на навколишні атоми. Отже, наш аналіз дає пояснення існування довгоживучих стабільних бризерів у металогідридних системах.