

Cite this: *Chem. Sci.*, 2018, **9**, 8644

All publication charges for this article have been paid for by the Royal Society of Chemistry

Atomic structure of boron resolved using machine learning and global sampling†

Si-Da Huang, Cheng Shang, Pei-Lin Kang and Zhi-Pan Liu *

Boron crystals, despite their simple composition, must rank top for complexity: even the atomic structure of the ground state of β -B remains uncertain after 60 years' study. This makes it difficult to understand the many exotic photoelectric properties of boron. The presence of self-doping atoms in the crystal interstitial sites forms an astronomical configurational space, making the determination of the real configuration virtually impossible using current techniques. Here, by combining machine learning with the latest stochastic surface walking (SSW) global optimization, we explore for the first time the potential energy surface of β -B, revealing 15 293 distinct configurations out of the 2×10^5 minima visited, and reveal the key rules governing the filling of the interstitial sites. This advance is only allowed by the construction of an accurate and efficient neural network (NN) potential using a new series of structural descriptors that can sensitively discriminate the complex boron bonding environment. We show that, in contrast to the conventional views on the numerous energy-degenerate configurations, only 40 minima of β -B are identified to be within 7 meV per atom in energy above the global minimum of β -B, most of them having been discovered for the first time. These low energy structures are classified into three types of skeletons and six patterns of doping configurations, with a clear preference for a few characteristic interstitial sites. The observed β -B and its properties are influenced strongly by a particular doping site, the B19 site that neighbors the B18 site, which has an exceptionally large vibrational entropy. The configuration with this B19 occupancy, which ranks only 15th at 0 K, turns out to be dominant at high temperatures. Our results highlight the novel SSW-NN architecture as the leading problem solver for complex material phenomena, which would then expedite substantially the building of a material genome database.

Received 2nd August 2018
Accepted 11th September 2018DOI: 10.1039/c8sc03427c
rsc.li/chemical-science

1. Introduction

The mysteries of boron have persisted in chemistry since the discovery of the B_2H_6 molecule. To name a few, (i) unlike carbon with its typical bonding patterns (such as sp^2 and sp^3), the chemical bonding of boron is highly flexible and complex;¹ (ii) the most stable structures of boron clusters, B_{80} as a famous example, are generally unknown; (iii) the ground state structure of boron crystals has not been conclusively determined in the

sense that there are no conclusive experimental and theoretical results on the relative stability of α -B and β -B.^{2–4} The potential energy surface (PES) of boron is thus of general interest, but is highly challenging to rationalize by both experiments and theory. On the other hand, machine learning has emerged as a promising tool to solve complex physical problems that are “much too complicated to be soluble” using quantum mechanics. While state-of-the-art neural network (NN) techniques^{5–8} are now available to establish the link between molecular characteristics (e.g. geometry) and the observable properties, it remains an open question how far NN techniques can be applied to solve the top challenges in physical science, especially those involving the PES, for which a quantitative solution for energy is needed. The structure determination of β -B is a very hard problem.

Among the 16 known boron allotropes, the rhombohedral form (β -B) was recently proven to be the most stable phase.^{4,9–12} The basic framework of β -B is considered as the layer by layer packing of B_{12} icosahedral cages and B_{28} triple-fused icosahedral cages along the [111] axis in a rhombohedral lattice (105 atoms in total, i.e. β -B₁₀₅, #166, $\bar{R}\bar{3}m$), as shown in Fig. 1a. Recent work suggested that the stability of β -B might be related

Collaborative Innovation Center of Chemistry for Energy Materials, Key Laboratory of Computational Physical Science (Ministry of Education), Shanghai Key Laboratory of Molecular Catalysis and Innovative Materials, Department of Chemistry, Fudan University, Shanghai 200433, China. E-mail: zpliu@fudan.edu.cn

† Electronic supplementary information (ESI) available: SSW-NN methodology, including the HDNN architecture, the iterative dataset generation, statistical assessment of different structural descriptors, general guidelines for setting up the structural descriptors and network size; DFT calculation details; pair distribution function of α -B; parameters for the structural descriptors used in set-1 and final NN potential; phonon and free energy calculations; convergence of occupancy rates with respect to number of minima; electronic analyses of β -I-15; XYZ coordinates for low energy crystals of boron and the top 40 β -B structures; the 8000-dataset for benchmarking purposes. See DOI: [10.1039/c8sc03427c](https://doi.org/10.1039/c8sc03427c)



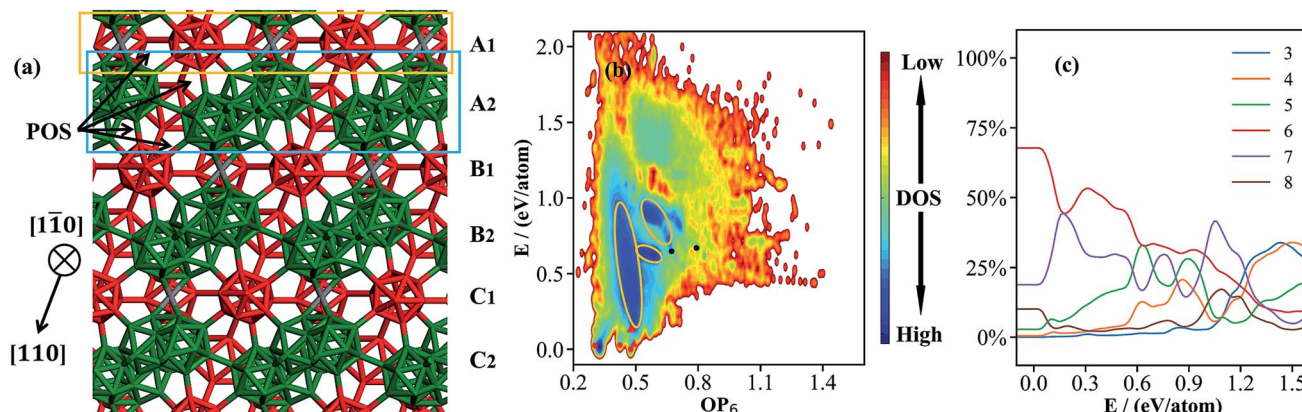


Fig. 1 (a) Atomic structure of the basic framework of β -B, β -B₁₀₅ (105 B atoms per rhombohedral lattice), highlighting the close packing of B₁₂ cages (red) and B₂₈ cages (green), and the connecting B15 sites (grey, detailed in Section 3). The cubic close packing pattern of β -B₁₀₅ (A₁A₂-B₁B₂C₁C₂A₁A₂...) together with the locations of possible partially occupied sites (POSSs); (b) the OP₆-E contour map of the global dataset from first principles. OP₆ is the distance-weighted Steinhardt order parameter in eqn (1) with $L = 6$, and the density of states (DOS) is indicated by color. The energy of α -B is set as zero. The red dots represent α -B, β -B₁₀₅ and γ -B, the red triangles represent #66 and honeycomb structures, and the black dots represent B₄₀-ball and B₄₀-flat. Their coordinates (E, OP₆) in the map are as follows: α -B: (0.00, 0.48); β -B₁₀₅: (0.03, 0.30); γ -B: (0.03, 0.42); #66: (0.04, 0.49); H-2: (0.14, 0.48); H-1: (0.14, 0.49); B₄₀-ball: (0.65, 0.67); B₄₀-flat: (0.67, 0.79); (c) the percentages of differently coordinated B for structures in the global dataset.

to the existence of numerous partially occupied sites (POSSs). These POSSs are located in or near the vacant spaces surrounded by B₁₂ and B₂₈ cages, and it is known from experiments that there are at least 10^7 likely arrangements for 42 POSSs in a rhombohedral cell, without even considering the other unknown POSSs. Traditionally, the simple “hand picking” strategy was utilized for searching for stable configurations of β -B based on the experimental XRD data.^{12,13} In 2009, an Ising model together with Monte Carlo sampling revealed that there were many energy-nearly-degenerate configurations for β -B.^{3,14,15} However, the Ising model fits the energetics of β -B minima structures containing the known dominant POSSs. Therefore, the Ising model cannot go further to predict new structures with unknown POSSs and importantly, cannot be used for geometry relaxation and global sampling. As an important consequence, current knowledge of the detailed occupancy of all possible POSSs in β -B is still far from satisfactory. How to search the complete configurational space of β -B by taking into account all likely geometry relaxation remains a great challenge and thus requires highly efficient PES exploration techniques.

The fast development of NN methods in recent years gives hope for the understanding of complex PES problems. The current NN methods generally involve the convolution of the 3-D atomic structure into numerical structure descriptors as input, and the subsequent learning against a big dataset, *i.e.* training the network parameters, for property prediction. Being the link between molecular/material structures and their properties, the structural descriptor plays key roles in the application type and predictive power of the NN. Apart from the straightforward Cartesian coordinates, many structural descriptors were proposed recently, *e.g.* extended-connectivity fingerprint,¹⁶ Coulomb matrix,¹⁷⁻¹⁹ graph convolution,^{20,21} SMILES strings^{22,23} and symmetry functions.^{24,25} For example, graph convolution has been utilized to encode organic

molecules for reaction prediction and drug toxicity prediction.^{20,21} Based on the idea of local internal coordinates, *e.g.* bond distances and bond angles, to construct a classical force field, Behler and Parinello proposed a high dimensional neural network (HDNN) approach for describing the PES of complex materials, where the atomic distances and angles are assembled to form atom-centered structural descriptors (also known as symmetry functions).^{24,25} In this approach, the total energy of the system is decomposed into the sum of individual atomic energies that can be obtained by NN training. We recently proposed a global-to-global scheme to generate a NN potential for describing the global PES of a material, which opens the possibility of material discovery from large-scale global PES scanning.²⁶

Here we aim to shed light on the global PES of boron by developing and applying machine learning methods. For this purpose, a large first principles calculation dataset is first constructed by using stochastic surface walking (SSW) global optimization^{27,28} to explore the global PES of boron. By developing new power-type structural descriptors, training the NN global potential and performing SSW global optimization on the boron NN potential, we are finally able to resolve the long-standing puzzles of the atomic structure of β -B. We reveal the general rules governing the stability of β -B configurations and resolve the physical origin of the strong temperature-dependence of the β -B structure, which has profound implications on the properties of boron.

This paper is organized as follows. In Section 2, we describe the boron global PES dataset generated by SSW PES exploration using first principles calculations, and develop a series of power-type structural descriptors for training such a complex dataset to obtain an accurate and transferable NN potential. In Section 3, we explore the PES of β -B using a SSW global search with NN potential, present all the low energy configurations of



β -B and determine the occupancy rate for the interstitial sites, which is compared with experimental data in detail.

2. Construction of boron global NN PES

a First principles dataset for boron global PES

A dataset for boron based on quantum mechanics calculations lays the foundation for machine learning of the boron PES. To reach high predictive power and transferability, this dataset needs to include as many different boron allotropes as possible, including solids and clusters. It therefore requires efficient PES sampling techniques,^{29–31} *e.g.* the SSW global optimization method^{27,28} utilized here, which can combine with quantum mechanics calculations to explore the PES. The SSW method has been applied for both structures and pathway searches of many complex systems, ranging from clusters (*e.g.* B₄₀,³² carbon fullerene³³), to surfaces,³⁴ to solids (*e.g.* TiO₂,³⁵ ZrO₂ (ref. 36)). Because the SSW simulation produces continuous trajectories when exploring the PES, it is an ideal tool to quickly generate a dataset with a variety of structural patterns, containing both minima and saddle points on the PES. More details on the SSW method and its recent combination with a HDNN for material discovery can be found in our previous work,²⁶ and also briefly in the ESI.†

In this work, the global dataset for boron is established *via* an iterative approach to incorporate as many boron bonding environments as possible. The first stage, being the most important and time-consuming step, was carried out using first-principles SSW global optimization in different systems with fewer than 40 atoms. Subsequently, a sample of structures from the first stage was taken as the training dataset for building a NN potential, which was then utilized to speed up the global optimization in large systems (up to 107 atoms). The HDNN architecture was utilized for the NN potential, and the network was trained by simultaneously matching the energy, force and stress (see ESI† and our previous work²⁶ for details). It should be mentioned that the NN potential trained for the purpose of expanding the global dataset does not need to be accurate and hence the structure descriptors and the network size utilized at this stage are generally small (these are discussed in depth in the ESI†).

To be more specific, the first stage of sampling involves up to 100 SSW simulations, each starting from a different initial structure, namely bulk, cluster and layer structures with different numbers of atoms (12, 14, 28, 40 per cell). These initial structures include the known solid allotropes (*e.g.* α -,² γ -B³⁷), the reported B₄₀ minima³² and randomly configured structures. All first-principles calculations were carried out using the plane-wave DFT code VASP³⁸ with the GGA-PBE functional,³⁹ and the details of the calculation setups are described in the ESI.†

Finally, we obtained a global dataset with 165 423 structures in total, containing 109 881 bulk, 4649 layer, and 50 893 cluster structures with different numbers of atoms, as detailed in Table S1.† In the dataset, the number of atoms per cell is usually 12 or 14, each with around 40 000–50 000 structures. These small

systems prevail in the dataset because they can represent major atomic environments and are computationally more efficient for both first-principles calculations and subsequent NN training. To provide an overview of this dataset, we constructed an energy *versus* geometry contour plot, as shown in Fig. 1b. The x-axis in Fig. 1b is the distance-weighted Steinhardt-type order parameter⁴⁰ (OP) defined by eqn (1) with the degree $L = 6$, as also utilized previously to distinguish the short–medium range ordering of solid structures.

$$OP_L = \left(\frac{4\pi}{2L+1} \sum_{m=-L}^L \left| \frac{1}{N_{\text{bonds}}} \sum_{i \neq j} e^{-\frac{1}{2} \frac{r_{ij}-r_c}{r_c}} Y_{Lm}(\mathbf{r}_{ij}) \right|^2 \right)^{\frac{1}{2}} \quad (1)$$

In eqn (1), Y_{Lm} is the spherical harmonic function, i and j are atoms in the lattice, \mathbf{r}_{ij} is the vector between atoms i and j , r_{ij} is the distance between them, r_c is set as 60% of the typical single bond length between i and j atoms (1.7 Å here for boron–boron single bonds), and N_{bonds} is the number of bonds in the first bonding shell (2.05 Å). The y-axis is the energy per atom (eV per atom), where zero energy is set as the energy of α -B hereafter, since its crystal structure is well defined.

Fig. 1b shows that at the bottom of the PES there are three major funnels belonging to three stable crystal phases: from left to right, they are β -B₁₀₅, γ -, and α -B (red dots in the figure) with OP₆ values of 0.30, 0.42 and 0.48, respectively. Many new boron crystal forms that are not reported in the literature can also be identified in these regions. For example, a high symmetry structure (Cccm, #66, red triangle) shares the same icosahedral packing skeleton as α -B but with 4 extra doping atoms at the interstitial sites per 52-atom unit cell. The honeycomb structures (H-1 and H-2, red triangles) are also typical in less stable B crystal forms,⁴¹ and form by packing two-dimensional boron sheets with different connections between neighboring layers (also see ESI† for these crystals). Above 0.15 eV per atom, there are dark blue zones with high density of states (circled by yellow lines), which correspond to amorphous solids and cluster structures. The lowest energy B₄₀ cluster (black dots) in the dataset is the B₄₀ fullerene (OP₆ = 0.67 in Fig. 1b), which is 0.65 eV per atom less stable than α -B.

It is of general interest to analyze the geometrical environment of boron in the global dataset. For this purpose, we have computed the B coordination number for all structures in the global set. The B–B distance of 2.05 Å is set as the criterion for B coordination, which takes into account the first nearest bonding neighboring atoms as indicated from the pair distribution function (see ESI†). In Fig. 1c, we plot the evolution of the B coordination number with increasing energy (x-axis). The y-axis is the percentage of different coordination numbers, which is calculated by counting and averaging the B coordination numbers for structures in the same energy interval, E to $E + dE$ ($dE = 1$ meV per atom). As shown, six and seven are the major coordination environments for the low energy crystal phases. Five coordination becomes popular in the amorphous solid region (0.15–0.6 eV per atom), and three and four coordination are the dominant coordination patterns only in the very high energy region (>1.0 eV per atom).



Fig. 1b and c demonstrate clearly that the bonding patterns of boron in the global dataset are highly complex, with coordination numbers ranging from 2 to 8 with few similarities to common polyhedral bonding. This feature must be attributed to the unique electronic structure of the B atom, which, to reach octet saturation, often adopts multi-center delocalized bonding.² This results in an extremely complex atomic environment for boron and thus leads to a great challenge in constructing either empirical or NN potentials for boron-containing materials.

b Structural descriptors with power radial function

Because the structure descriptor is the key to correlating a structure with its energetics in a HDNN, one would expect that a qualified structural descriptor needs to be sensitive enough to distinguish as many structures as possible on the PES. For the boron global dataset, which has great structural diversity, it is thus more difficult to identify the appropriate structural descriptors.

Let's first recall the structural descriptors originally proposed by Behler and Parrinello:^{24,25} the most used two-body G^2 and three-body G^4 functions are described in eqn (2)–(4):

$$f_c(r_{ij}) = \begin{cases} 0.5 \times \tan h^3 \left[1 - \frac{r_{ij}}{r_c} \right], & \text{for } r_{ij} \leq r_c \\ 0, & \text{for } r_{ij} > r_c \end{cases} \quad (2)$$

$$G_i^2 = \sum_{j \neq i} e^{-\eta(r - r_s)^2} \cdot f_c(r_{ij}), \quad (3)$$

$$G_i^4 = 2^{1-\zeta} \sum_{\substack{\text{all} \\ j, k \neq i}}^{} (1 + \lambda \cos \theta_{ijk})^\zeta \cdot e^{-\eta(r_{ij}^2 + r_{ik}^2 + r_{jk}^2)} \cdot f_c(r_{ij}) \cdot f_c(r_{ik}) \cdot f_c(r_{jk}), \quad (4)$$

where r_{ij} is the inter-nuclear distance between atoms i and j , and θ_{ijk} is the angle centered at the i atom with neighbors j and k (i, j, k are atom indices). The key ingredients in the Behler-type structural descriptors (BTSDs) are the cutoff function f_c which decays to zero beyond the r_c (eqn (2)), the Gaussian-type radial function and the trigonometric angular functions. By changing five parameters, r_c , r_s , η , ζ and λ , a set of two-body G^2 (eqn (3)) and three-body G^4 (eqn (4)) functions can then be generated, which serve to distinguish the atomic environment of the central atom i .

In fact, we initially tested the BTSDs for constructing the boron NN PES using the global dataset. However, it was unable to achieve a high accuracy (the root mean square (RMS) for energy was larger than 30 meV per atom). This implies that the global PES of boron, despite containing only a single element, is much too complex to describe using the BTSDs alone.

To solve this problem, we have designed a series of new structural descriptors, S^1 to S^6 (eqn (5)–(11)). Inspired by the Laguerre polynomials for atomic orbitals, all these structural descriptors utilize the power function as the radial function. We therefore named them power-type structural descriptors (PTSDs) to distinguish them from the BTSDs.

$$R^n(r_{ij}) = r_{ij}^n \cdot f_c(r_{ij}), \quad (5)$$

$$S_i^1 = \sum_{j \neq i} R^n(r_{ij}), \quad (6)$$

$$S_i^2 = \left[\sum_{m=-L}^L \left| \sum_{j \neq i} R^n(r_{ij}) Y_{Lm}(\mathbf{r}_{ij}) \right|^2 \right]^{\frac{1}{2}} \quad (7)$$

$$S_i^3 = 2^{1-\zeta} \sum_{j, k \neq i} (1 + \lambda \cos \theta_{ijk})^\zeta \cdot R^n(r_{ij}) \cdot R^m(r_{ik}) \cdot R^p(r_{jk}), \quad (8)$$

$$S_i^4 = 2^{1-\zeta} \sum_{j, k \neq i} (1 + \lambda \cos \theta_{ijk})^\zeta \cdot R^n(r_{ij}) \cdot R^m(r_{ik}), \quad (9)$$

$$S_i^5 = \left[\sum_{m=-L}^L \left| \sum_{j, k \neq i} R^n(r_{ij}) \cdot R^m(r_{ik}) \cdot R^p(r_{jk}) \cdot (Y_{Lm}(\mathbf{r}_{ij}) + Y_{Lm}(\mathbf{r}_{ik})) \right|^2 \right]^{\frac{1}{2}}, \quad (10)$$

$$S_i^6 = 2^{1-\zeta} \sum_{j, k, l \neq i} (1 + \lambda \cos \delta_{ijkl})^\zeta \cdot R^n(r_{ij}) R^m(r_{ik}) R^p(r_{il}). \quad (11)$$

In the PTSDs, S^1 and S^2 are two-body functions, S^3 , S^4 and S^5 are three-body functions, and S^6 is a four-body function. S^1 and S^3 mimic G^2 and G^4 , respectively, except for the change in the radial function. S^2 and S^5 have a common component, *i.e.* the spherical function, as seen in the Steinhardt-type order parameter (eqn (1)), which has been found to be sensitive for distinguishing the local coordination of an atom.^{40,42} S^6 is a four-body term, designed to describe the torsion angle. The torsion angle δ_{ijkl} in eqn (11) is the dihedral angle centered at the I and j atoms, with k and l being the neighboring atoms. In total, there are seven adjustable parameters, n, m, p, L, r_c, ζ and λ in the PTSDs.

The replacement of the Gaussian function in the BTSDs by the power function in the PTSDs has several advantages: (i) the computational cost of the numerical calculations is reduced; (ii) the adjustable parameters are reduced from two (r_s, η) to one (n) which simplifies the search for the optimal parameters for the two-body functions; (iii) the power function when combined with the decaying cutoff function can create radial distributions with flexible peak and shape, which fulfills the similar purpose of the Gaussian function; (iv) the introduction of different powers (n, m, p) in the three-body functions can conveniently couple different radial distributions. To illustrate point (iii), we plot in Fig. 2 the evolution of R^n versus r_{ij} for different n and the same cutoff radius $r_c = 3.2$ Å. As shown, the peak of the R^n function shifts to larger r_{ij} and becomes narrower with increasing n . This enables us to portray the neighboring atoms within any circular shell by adjusting r_c and n .

Overall, the new PTSDs incorporate flexible radial functions, spherical functions and up to four-body functions. To assess the structure discrimination ability of the different types of structural descriptors (detailed in ESI†), we have performed principle component analysis (PCA) as detailed in the ESI,† which demonstrates that the new PTSDs outperform the BTSDs in



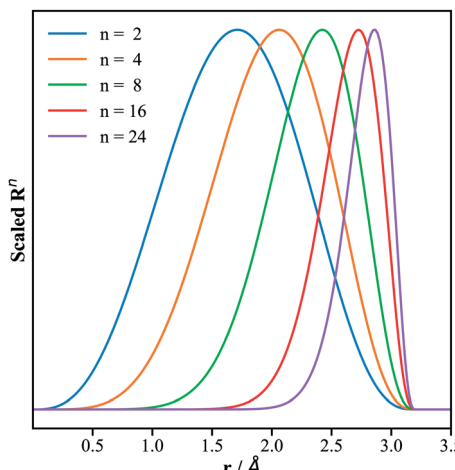


Fig. 2 Plots of the radial part of the PTSDs, eqn (5), for the same cutoff radius of 3.2 Å but different power n . The x-axis is the distance r , while the y-axis is the function value scaled to (0, 1).

describing the structures in the boron global dataset. In particular, it outlines the importance of the S^2 , S^4 and S^5 descriptors, which rank top out of the two-body and three-body functions. Apparently, the incorporation of the spherical harmonic function in the S^2 and S^5 PTSDs enhances substantially the structure discrimination ability.

c Boron NN potential

We are now ready to produce the boron NN potential using the global dataset. To achieve a high accuracy which is desirable for differentiating the energy-degenerate configuration isomers of β -B, we have adopted a large set of structural descriptors, which contains 173 PTSDs, *i.e.* 39 S^1 , 36 S^2 , 16 S^3 , 52 S^4 , 18 S^5 and 12 S^6 , and compatibly, the network utilized is also large, involving two hidden layers each with 110 neurons, equivalent to 31 461 network parameters in total. Our theoretical procedure to set up the structural descriptors and network size is discussed in detail in the ESI,† where we verify that the current structure descriptor set is complete and that the network size is the optimum to achieve a highly accurate NN potential. After training the network on the global dataset, we obtain the first boron global NN PES with RMS values for energy, force and stress of 12.4 meV per atom, 0.28 eV Å⁻¹, and 3.00 GPa, respectively. The overall accuracy is quite standard for a global NN PES considering that the energies of the structures in the dataset span a large window from 0 to 4 eV per atom.

We have examined the accuracy of the NN PES for the representative crystal/cluster boron structures and benchmarked it against DFT calculations, as listed in Table 1. These structures were fully optimized using the NN until the maximal force component was below 0.01 eV Å⁻¹ and the stress was below 0.01 GPa, and then refined using DFT for comparison. As shown, the energy RMS error is 2.73 meV per atom for these typical low energy minima, while the volume RMS error is ~0.45%. This accuracy is sufficient for a global structure and pathway search to identify the low energy candidates.

Table 1 Comparison between the NN and DFT for common boron crystal phases based on energy (E , meV per atom) and volume (V , Å³)

Name	Z^b	E^{DFT}	ΔE^{\ddagger}	V^{DFT}	ΔV^c
Important crystal structures					
α -B	12	0	2.77	7.25	0.66
β -B ₁₀₅	105	25.29	2.45	7.78	0.28
γ -B	28	27.31	0.88	6.99	-0.88
#66 ^a	52	39.52	2.74	7.62	-0.02
H-1 ^a	12	136.83	-4.67	6.88	0.34
H-2 ^a	19	143.01	4.25	6.88	-0.47
B ₄₀ -ball	40	649.40	0.29	—	—
B ₄₀ -flat	40	671.33	-3.74	—	—
RMS	—	—	2.73	—	0.45
Structures of β-B					
β -II-1	107	-0.75	0.81	7.64	0.03
β -II-2	107	-0.69	0.58	7.65	-0.06
β -II-3	107	-0.09	-0.93	7.65	-0.08
β -II-4	107	0.07	-0.47	7.65	-0.09
β -II-5	107	1.18	1.37	7.65	0.19
β -I-6	106	1.30	2.67	7.71	-0.1
β -I-7	106	1.42	2.86	7.72	-0.24
β -II-8	107	1.53	0.66	7.65	0.29
β -II-9	107	1.71	0.79	7.65	0.3
β -I-10	106	1.92	2.01	7.72	-0.28
β -II-11	107	2.51	0.41	7.65	0.05
β -II-12	107	2.68	0.51	7.65	0.07
β -II-13	107	2.89	-1.87	7.65	-0.04
β -II-14	107	3.4	-1.43	7.65	-0.05
β -I-15	106	4.03	0.85	7.72	-0.13
RMS	—	—	1.44	—	0.17

^a Crystal structures identified from global PES. ^b Z : number of atoms per unit cell. ^c $\Delta E = E^{\text{DFT}} - E^{\text{NN}}$ and ΔV is the percentage volume deviation between the NN and DFT.

3. Structure of β -B

a β -B PES from the SSW-NN

β -B was long believed to have numerous structural configurations as a “frustrated system”, similar to ice.¹⁴ Due to the electron deficiency of B₁₂ icosahedra and the intrinsic instability of B₂₈ triple-fused icosahedra,⁴³ the crystal is self-doped with the doping atoms appearing in many likely interstitial sites in the lattice. These sites, commonly known as partially occupied sites (POSSs), are only partly distinguishable from XRD experiments: the reported occupancy of the dominant POSSs varies from experiment to experiment, and about 3% of POSSs are even unassigned since their geometric positions are unknown.⁴⁴ Unlike previous work^{14,15} that utilized the Ising model to establish the static interaction between the known POSSs, we are now able to utilize the NN potential to explore all the likely structures on the boron global PES, where both the atom and the lattice can be fully optimized.

Starting from the known β -B rhombohedral lattice, a SSW global optimization was utilized to explore all the possible low energy structures of β -B. More than 20 SSW runs were carried out independently, with 10 000 minima to visit in each run. The structures contained either 105, 106 or 107 atoms per cell, since these are known as the most stable structures in the literature.

By removing duplicated minima, we finally obtained 15 293 distinct minima for β -B, and the OP_6 -energy PES contour plot for these minima is plotted in Fig. 3. The representative atomic structures of β -B are shown in Fig. 4.

It should be emphasized that the SSW sampling of large systems (>100 atoms) is extremely computationally demanding using first principles calculations. We estimated that the NN calculations are at least 3×10^3 times faster than DFT calculations,²⁶ and thus the SSW-NN allows more than 2×10^5 minima of β -B to be visited in a short time.

As shown in Fig. 3, there are two high density of states regions (deep blue zones) in the β -B PES, which are separated by a gap at ~ 0.15 eV per atom (red box). We found that all structures below the gap have intact B_{12} (red units in Fig. 4a) and B_{28} cages (dark green units in Fig. 4a) as the skeleton, the same as the known β -B, and that these cages start to melt (crack) for high energy structures above the gap (a typical melting B_{28} cage is shown in the insert of Fig. 3).

To determine an accurate energy sequence for the β -B isomers, we selected the 366 most stable minima (25 meV per atom above the most stable minimum) predicted by the NN PES and refined them using DFT. The energy root mean square error (RMSE) between the NN and DFT for these minima is 3.85 meV per atom, suggesting that these selected minima from the NN PES should cover the most stable minima of β -B. Table 1 provides a comparison between DFT and the NN for the 15 most stable structures, based on energy and volume. Clearly, for these most stable structures, the RMSEs in energy and volume for the NN prediction are rather low, being 1.44 meV per atom and 0.17%, respectively. We emphasize that all results below are referenced to the DFT calculations for the high accuracy setups (see ESI†).

Structural patterns

Now we are in a position to inspect closely the atomic structures of the most stable β -B isomers. From our results, there are three

different skeletons for β -B, namely **SK-I**, **SK-II** and **SK-III**, differing in the connecting blocks between neighboring B_{28} cages. **SK-I** and **SK-II** were known previously,³ and **SK-III** was newly found in this work. They are elaborated as follows.

(i) **SK-I**. The **SK-I** structure (Fig. 4b) features five B_{13} (orange balls) and one B_{15} (grey ball) in the connecting unit. The five B_{13} are all apex atoms of two B_{28} units, which are connected by the five-coordinated B_{15} . Since there is one missing apex site in one B_{28} unit, there are two least-coordinated B_3 (brown balls) that are only four-coordinated to the skeleton.

(ii) **SK-II**. The **SK-II** structure (Fig. 4c) features four B_{13} , one B_{15} , one B_{17} (purple ball) and one B_{18} (magenta ball) in the connecting unit. B_{17} and B_{18} can be considered as the relocation of two B_{13} into the vacant region between two B_{28} cages. B_{17} is four-coordinated to B_{15} , B_{18} and one B_{28} cage, while B_{18} links to B_{17} , one B_{28} cage and also the neighboring two B_{12} cages. Obviously, only one least-coordinated B_3 is present (brown ball) in **SK-II**.

(iii) **SK-III**. The **SK-III** structure (Fig. 4d) is similar to **SK-I** but with one B_{13} missing. Thus, only four B_{13} are left in **SK-III**. The connecting B_{15} has a planar four-coordination with the four B_{13} .

In these skeletons, every four neighboring cages, either B_{12} or B_{28} , form a tetrahedral void, which provides four possible doping sites in the hexagonal ring of the tetrahedral faces. B_{16} , B_{19} and B_{20} are such doping sites, dominant in the most stable β -B structures and accounting for $\sim 30\%$ of doping sites according to experiments (the other 70% are accounted for by the skeleton POSS B_{13} , B_{17} and B_{18}). B_{16} (light green ball) is the most frequently encountered, and resides in the ring connecting three B_{12} units. B_{19} (yellow ball in Fig. 4e) and B_{20} (light purple ball) are in the rings connecting two B_{12} units and one B_{28} unit. They link to B_{28} differently, *via* two B_3 (blue balls) for B_{19} , but *via* one B_3 and one B_8 (turquoise balls) for B_{20} . By taking B_{13} , B_{17} and B_{18} into account, there are 42 total POSS (six for B_{13} , B_{16} , B_{17} , B_{18} and B_{19} , and twelve for the B_{20} sites) in one rhombohedral unit cell and thus at least 10^7 possible configurations.

Interestingly, other unknown doping sites were also revealed by our global search. Two such sites, namely B_{21} and B_{22} , are illustrated in Fig. 4f and g and are described as follows. B_{21} (light purple ball in Fig. 4f) is located in the ring connecting two B_{12} units and one B_{28} unit *via* one B_8 and one B_{10} (green balls). B_{22} is located inside the tetrahedron surrounded by four B_{12} units, which can be occupied by a pair of atoms (light purple balls in Fig. 4g).

Energy spectrum

In Table 2, we list the key structural information for the important β -B isomers. From DFT, **SK-II** is the most stable skeleton, being present in the five most stable structures. The most stable β -B is β -II-1, which has two doping atoms in the B_{16} sites per rhombohedral cell. Hereafter the notation (β -II-1) follows this rule: the Roman number (II) represents the skeleton (**SK-II**) and the Arabic number (1) indicates the energy ranking. This global minimum (GM) from our SSW search is in

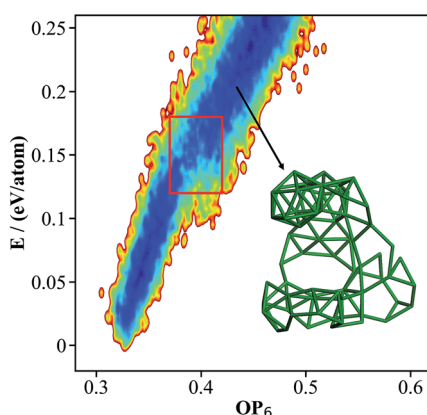


Fig. 3 The contour map (E against OP_6 , also see Fig. 1 caption for explanation) for the β -B minima from the SSW global optimization using the boron NN PES. The red box indicates the boundary above which the B_{28} cages start to melt. The inset shows a typical structure with broken B_{28} cages.



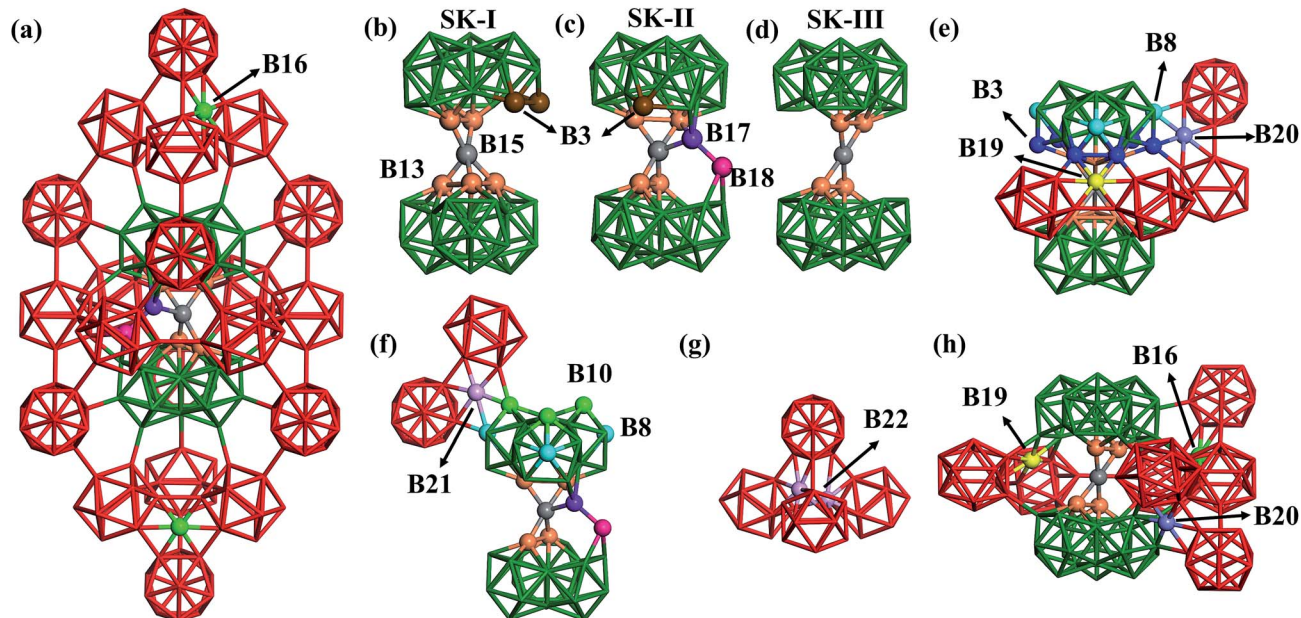


Fig. 4 Important structures of β -B from the SSW-NN search. (a) Atomic structure of the most stable configuration, β -II-1, highlighting the location of B16; (b) the key unit in the SK-I framework, highlighting the locations of B13 and the least-coordinated B3; (c) the key unit in the SK-II framework, showing the locations of B17, B18 and the least-coordinated B3; (d) the key unit in the SK-III framework; (e) the locations of B3, B8, B19 and B20; (f) the locations of B8, B10 and B21; (g) the location of B22; (h) the key unit in β -III-18.

accordance with the most stable structure reported in ref. 13. The second to the fourth most stable structures, *i.e.* β -II-2 to β -II-4 (<1 meV per atom), are structurally similar to the GM with the major difference being the B16 position. Due to symmetry

breaking in **SK-II**, there are six distinct B16 sites, as shown in Fig. 5a, numbered from 1 to 6. In the figure, *a*, *b*, and *c* represent the three lattice axes in the close packing (111) plane in the rhombohedral lattice.¹² Restricted by the three-fold symmetry,

Table 2 The key information for the important structures of β -B from the SSW-NN global search. The data listed include the DFT energy (E^{DFT} , meV per atom) of the structures, the number of filled POSS (B13 to B22), the structure pattern (P_1 to P_6) and the partition function contribution (C%) at different temperatures (K)

Name	E^{DFT}	C%											
		B13	B16	B17	B18	B19	B20	B21	B22	Pattern	1000	1500	2000
β -II-1	-0.75	4	2	1	1	0	0	0	0	P_1	18.2	8.0	4.2
β -II-2	-0.69	4	2	1	1	0	0	0	0	P_1	17.1	7.7	4.1
β -II-3	-0.09	4	2	1	1	0	0	0	0	P_1	8.1	4.7	2.8
β -II-4	0.07	4	2	1	1	0	0	0	0	P_1	6.6	4.1	2.5
β -II-5	1.18	4	1	1	1	0	1	0	0	P_2	11.0	9.8	7.4
β -I-6	1.29	5	2	0	0	0	0	0	0	P_3	4.6	4.5	3.5
β -I-7	1.42	5	2	0	0	0	0	0	0	P_3	3.9	4.0	3.2
β -II-8	1.53	4	1	1	1	0	1	0	0	P_2	7.1	7.3	5.9
β -II-9	1.71	4	1	1	1	0	1	0	0	P_2	5.8	6.4	5.3
β -I-10	1.92	5	2	0	0	0	0	0	0	P_3	2.1	2.7	2.4
β -II-11	2.51	4	1	1	1	0	1	0	0	P_2	2.1	3.3	3.2
β -II-12	2.68	4	1	1	1	0	1	0	0	P_2	1.7	2.9	2.9
β -II-13	2.89	4	2	1	1	0	0	0	0	P_1	0.2	0.4	0.4
β -II-14	3.4	4	2	1	1	0	0	0	0	P_1	0.1	0.3	0.3
β -I-15	4.03	5	1	0	0	1	0	0	0	P_4	7.6	18.3	23.2
β -II-17	4.32	4	1	1	1	0	0	1	0	P_5	0.2	0.8	1.1
β -III-18	4.42	4	1	0	0	1	1	0	0	P_5	0.1	0.3	0.5
β -I-21	4.65	5	1	0	0	0	0	0	2	P_5	0.0	0.1	0.2
β -II-26	4.93	4	1	1	1	1	0	0	0	P_6	0.0	0.1	0.1
β -II-27	4.98	4	1	1	1	0	1	0	0	P_2	0.1	0.4	0.7
β -I-29	5.15	5	1	0	0	1	0	0	0	P_4	1.9	7.3	11.7
β -I-34	5.64	5	1	0	0	0	1	0	0	P_6	0.0	0.1	0.2

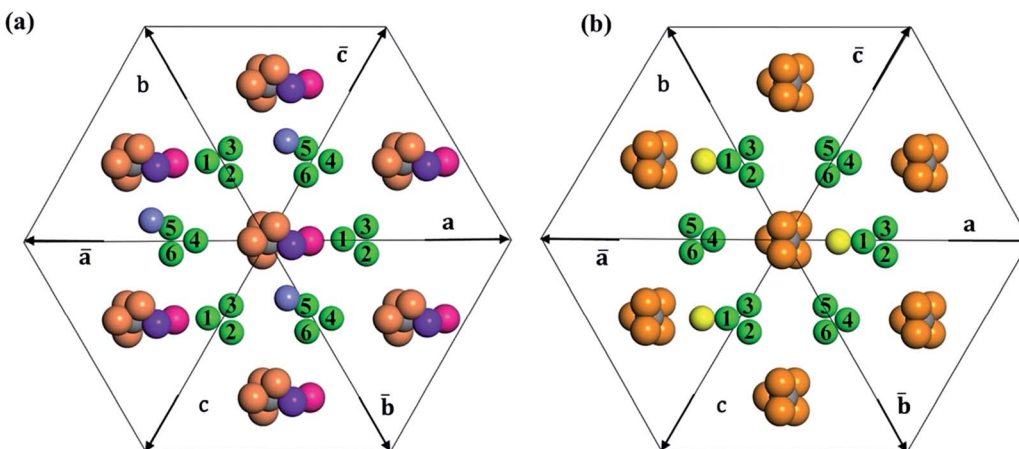


Fig. 5 (a) Positions of B16 (green balls) and B20 (light purple balls) with respect to B13 (orange balls), B15 (grey balls), B17 (purple balls) and B18 (magenta balls) in SK-II. All views are along the close packing [111] direction of the rhombohedral cell. The B20 location plotted is the most stable position of B20, appearing in β -II-5, β -II-8 and β -II-9; (b) positions of B16 and B19 (yellow balls) with respect to B13 and B15 in SK-I. The B19 location plotted is the most stable position of B19, appearing in β -I-15 and β -I-29.

the three sites (a_1, b_1, c_1) are equivalent, and so are (a_2, b_2, c_2) and (a_3, b_3, c_3). These four degenerate GM structures have the following B16 arrangements: β -II-1: $a_2\bar{a}_6$; β -II-2: $a_3\bar{a}_5$; β -II-3: $a_3\bar{a}_4$, β -II-4: $a_2\bar{a}_4$.

It is interesting to further understand why only four energy-nearly-degenerate isomers are present out of $C_6^2 = 30$ possible configurations. This is due to the fact that the following three scenarios are energetically not favored, *i.e.* exclusion rules: (i) two atoms in one tetrahedral void. The structural configurations with two atoms (*i.e.* B22 site, β -II-22) are at least 5.4 meV per atom less stable than the GM; (ii) the filling of the a_1 site in **SK-II**. This is due to the too short contact with the nearby B18 site. (Fig. 5a); and (iii) two occupied B16 in the same close packing plane, such as $a_2\bar{a}_5$. The structure with the $a_3\bar{a}_6$ B16 arrangement, β -II-13, is 3.64 meV per atom above the GM. The poorer stability is obviously due to the large strain induced by the two neighboring doping B16 atoms in the same close packing plane.

The fifth most stable structure, β -II-5, contains one B16 and one B20, and is 1.93 meV per atom above the GM. It also has two energy-nearly-degenerate isomers, β -II-8 and β -II-9. They share the same B20 site, which is 4.29 Å away from B17 as shown in Fig. 5a. This B20 site is coordinated to the least-coordinated B3 in the B₂₈ cage. The B16 sites for these three structures are: (i) \bar{a}_4 ; (ii) a_2 ; (iii) a_3 . The other possibilities, a_1 , \bar{a}_5 and \bar{a}_6 , are not favored due to the same exclusion rules as mentioned above for the four GM isomers.

SK-I appears in the sixth most stable structure, β -I-6, which contains two B16 sites (the same as reported in ref. 12). It has only two energy-nearly-degenerate structures, β -I-7 and β -I-10, apparently because only four distinct B16 sites are present in **SK-I** and the same exclusion rules must be obeyed. Their B16 arrangements are as follows: β -I-6: $a_1\bar{a}_6/a_3\bar{a}_4$; β -I-7: $a_2\bar{a}_6/a_2\bar{a}_4$; β -I-10: $a_1\bar{a}_5/a_3\bar{a}_5$. Next, the 11th to 14th most stable structures contain either B16 or B20 doping atoms in the less favored configurations.

Importantly, the B19 site only starts to be occupied in β -I-15, which is 4.78 meV per atom above the GM. This B19 site, being

3.8 Å away from B15, is connected to two least-coordinated B3 and thus can be considered as a relocation of B18 by shifting only 0.83 Å towards the nearby B₁₂ unit. The second stable structure with B19 is β -I-29, which is another 1.12 meV per atom above β -I-15. They contain the same B19 site, but with different B16 positions as shown in Fig. 5b: (i) \bar{a}_2/\bar{a}_3 ; (ii) \bar{a}_1 .

Overall, these low energy structures (top 40 structures, <7 meV per atom above the GM) can be summarized as six major patterns ordered by energy sequence:

- P₁: **SK-II** with 2 B16;
- P₂: **SK-II** with a B16 and a B20;
- P₃: **SK-I** with 2 B16;
- P₄: **SK-I** with a B16 and a B19;
- P₅: structures with B21/B22 doping sites or **SK-III**;
- P₆: **SK-II** with a B16 and a B19 or **SK-I** with a B16 and a B20.

With all these minima, most of them discovered for the first time, it is possible for us to assess current knowledge of the structure of β -B, which has been found to be either misleading or even incorrect.

(i) Only 91 structures are within 10 meV per atom and 40 are within 7 meV per atom above the GM, which is remarkably smaller than the $>10^7$ possible configurations from different arrangements of dominant POSS. Apparently, the energy degeneracy is much lower than expected. This is because (i) the exclusion rules are critical for pinning the position of B16; (ii) the filling of B19 strongly prefers only one position, *i.e.* that in β -I-15; (iii) the filling of B20 prefers five positions out of 12 possibilities in the top 40 structures. Ogitsu¹⁴ *et al.* stated that the filling of B20 must occur simultaneously with a vacant B13, which is indeed true for the most preferable B20 position (β -II-5). But, we also found that the filling of B20 near an occupied B13 site (β -II-11) is only 1.33 meV per atom less stable than β -II-5.

(ii) The filling of a particular B20 site can be energetically rather stable, which rationalizes the experimental observation of B20 filling. From our results, occupation at the B20 site (β -II-5) is slightly more preferable than that at the B19 site (β -II-26).



Apparently, occupation at the B20 site preferentially occurs in **SK-II**, while occupation at the B19 site preferentially occurs in **SK-I**. This corrects the view of Setten¹² *et al.*, who suggested that the filling of an arbitrary B19 is more stable than the filling of a B20. Moreover, we found that the simultaneous filling of one B19 and one B20 in one rhombohedral cell can also be stable. For example, β -III-18 (Fig. 4h) contains three doping atoms at B16, B19 and B20 sites, which is only 5.17 meV per atom above the GM.

(iii) Unknown POSSs are present and have low energies. This information was not available previously, but it is important for understanding the residual boron observed in experiments.⁴⁴ The main newly identified POSSs from our results are B21 and B22, occurring in β -II-17 (5.07 meV per atom above the GM, containing a B16 and a B21) and β -I-21 (5.4 meV per atom above the GM, containing a B–B pair in B22 sites).

b POS occupancy rates at different temperatures

With all these low energy minima, it is now possible to derive from theory the occupancy rates for POSSs at different temperatures, which can be compared with the fitted data from XRD experiments. The data available from experiment,⁴⁴ *i.e.* the three examples MG57, MG179 and EP, as listed in Table 3, exhibit a variation in POS occupancy rate of up to 6.5%. Since the MG57 and MG179 samples were cooled from the melt at rates of \sim 350 and $2.2\text{ }^{\circ}\text{C min}^{-1}$, respectively, the variation in the data suggests that the occupancy rates of the POSSs are sensitive to the sample preparation conditions, especially temperature. This suggests that one must take into account the free energy contribution in order to properly address the structure of β -B.

From theory, we can derive the occupancy rates of the POSSs using eqn (12) and (13), assuming thermodynamic equilibrium at a given temperature T :

$$P\% = \frac{\sum_i n_i^{\text{POS}} \cdot e^{-\frac{G_i - G^{\text{min}}}{RT}}}{N^{\text{POS}} \sum_i e^{-\frac{G_i - G^{\text{min}}}{RT}}} \times 100\%, \quad (12)$$

$$G_i = E_i + \text{ZPE}_i - TS_i. \quad (13)$$

Table 3 Occupancy rates for POSSs from DFT and the NN at different temperatures (1000, 1500 and 2000 K), and also from three experimental samples

	B13	B16	B17	B18	B19	B20	Res ^b
NN-1000	68.3	29.4	15.0	15.0	1.5	1.2	0.5
NN-1500	71.8	23.3	11.6	11.5	4.6	2.3	2.0
NN-2000	74.1	20.2	9.1	9.1	6.8	2.4	3.2
DFT-1000	70.1	26.8	13.2	13.2	1.6	2.4	0.6
DFT-1500	73.2	22.8	10.1	10.1	4.6	2.8	2.3
DFT-2000	75.0	20.7	8.1	8.1	6.7	2.7	3.7
MG179 ^a	73.0	28.4	9.7	7.4	7.0	2.5	3.2
EP ^a	74.5	27.2	8.5	6.6	6.8	3.2	2.1
MG57 ^a	77.7	25.8	3.2	5.8	7.2	0	3.9

^a Experimental data from ref. 44. ^b Residual atoms with unknown location.

At a given temperature T , the free energy G_i for the i -th configuration is first determined by correcting the zero-point energy (ZPE) and vibrational entropy (S). These thermodynamic properties are calculated from the full phonon dispersion by using the numerical finite difference approach to diagonalize the Hessian matrix (more details in the ESI†). The one with the lowest free energy G^{min} is then identified, and is utilized to establish the partition function and calculate the occupancy rate P . In the equation, N^{POS} is the total number of POSSs (six for B13, B16, B17, B18, and B19; twelve for B20), and n_i^{POS} is the number of occupied POSSs in the structure.

In Table 3, we listed the calculated POSS occupancies at three typical temperatures, 1000, 1500 and 2000 K, and compared them with the experimental data. It can be seen that the data predicted by the NN agrees well with that from DFT (within 3% deviation), and the theoretical occupancies are also consistent with the experimental observations, with the deviation generally within 5%. In particular, our results interestingly show that with an increase in temperature, the occupancy of B13 increases, while that of B16, B17 and B18 decreases monotonically. This temperature dependence of the POSS occupation concurs with the variation of the data from three different experimental samples. Table 2 also lists the detailed POSS contributions for selected important structures at different temperatures. These results are elaborated as follows.

In general, the B13 occupancy is around 75%, which is obviously an average value from the B13 occupancy in the two dominant skeletons, 83% in **SK-I** and 68% in **SK-II**. In addition, B17 and B18 have 17% occupancy in **SK-II** and zero occupancy in **SK-I**, which suggests that the occupancies of B17 and B18 are equivalent and around 8% on average. As for the B16 sites, one or two sites per rhombohedral cell are filled in the low energy structures (**P₁** to **P₄**), resulting in a B16 occupancy of around 25%. On the other hand, the filling of B19 and B20 sites generally occurs in combination with the filling of B16 sites, suggesting that their occupancies are below 8%.

Taking the POSS occupancy rates at 1500 K as an example, there are 73.2% B13, 22.8% B16, 10.1% B17 and B18, 4.6% B19 and 2.8% B20. At this high temperature, the top 20 structures with the lowest free energy, all belonging to **P₁** to **P₅**, have obvious contributions (95.4%) to the overall POSS occupancy rates. The most important configuration is β -I-15 (**P₄**), which contributes 18.3% to the final occupancy. The structure with the lowest contribution among them, β -II-27 (**P₂**), still contributes more than 0.4%. For the rest of the structures in the top 100 structures, their total contribution reaches up to \sim 4.5%, and the top 100 to 300 structures only have very low contributions of $<0.1\%$. Therefore, numerical convergence of the POSS occupancies calculated from the boron global PES is achieved with the 100–300 lowest energy structures (see ESI† for details).

To understand the origin of the free energy difference, we plotted the phonon spectra of four structures, *i.e.*, β -II-1, β -II-5, β -I-6 and β -I-15, corresponding to the lowest energy structures for **P₁** to **P₄**, in Fig. 6a. There are three major peaks at low frequencies (350 cm^{-1} , 580 cm^{-1} and 680 cm^{-1}). Obviously, compared to the other isomers, β -I-15 shows the highest phonon density of states (DOS) for these three peaks; β -II-5 has

a higher phonon DOS for the peaks at 580–700 cm^{−1} compared to β -II-1 and β -I-6; β -I-6 has a higher phonon DOS for the 350 cm^{−1} peak compared to β -II-1 and β -II-5.

By examining the phonon displacement vectors, we found that the 350 cm^{−1} peak is mainly due to the vibrational motion of B13 atoms, and that the 580–700 cm^{−1} peaks correspond to the soft translational/rotational motion of skeleton B₁₂ and B₂₈ cages. The more intense peak of β -I-15 at ~350 cm^{−1} can be attributed to the additional B19 vibrations due to the flat PES associated with the B19 atom (also see the next section for electronic structure analyses). For similar reasons, B17 and B18 doping (in β -II-1 and β -II-5) will restrict the motion of B13 atoms and reduce the phonon density at ~350 cm^{−1}. The more intense peaks of β -I-15 and β -II-5 at 580–700 cm^{−1} are related to their only singly-occupied B16 sites, the filling of which will hinder the collective motion of the B₁₂ cages (*c.f.* the two B16 present in β -II-1 and β -I-6).

By plotting the relative free energy ($G_i - G[\beta\text{-II-1}]$) for β -II-5 (green), β -I-6 (blue), and β -I-15 (red) with respect to β -II-1 (Fig. 6b), one can see that at elevated temperatures, β -I-15 has

the lowest free energy, followed by β -II-5 and then β -I-6. Above 1210 K and 1320 K respectively, the free energies of β -I-15 and β -II-5 are lower than that of the GM β -II-1. This provides clear evidence that at temperatures above ~1200 K, the GM β -II-1 is no longer the thermodynamically favored configuration.

This dramatic free energy contribution, up to 5.5 meV per atom, leads to a change in POS occupancy at 1000 to 2000 K. β -II-1 and β -II-2, belonging to \mathbf{P}_1 , contribute ~30% of the total POS occupancy at 1000 K, while β -I-15 and β -I-29, belonging to \mathbf{P}_4 , provide a slightly higher contribution (~34%) at 2000 K. The change in preference from SK-II to SK-I rationalizes the monotonic increase in B13 occupancy, and the decrease in B16, B17 and B18 occupancies, as also observed in experiments.

It is also interesting to note that the B16 occupancy observed experimentally is in general higher than our theoretical results by ~5%. This is equivalent to ~1 additional B16 atom per hexagonal lattice (~320 atoms per cell) in the experimental sample. A possible explanation is that our theoretical results are from bulk calculations and do not consider the surfaces and domain boundaries. Due to the electron deficient nature of surfaces, the filling of an extra B16 would provide additional electrons and thus stabilize the surface. Another difference between theory and experiment lies in the paired nature of B17 and B18: the B17 and B18 occupancies are the same from theory, but different by up to 3% from experiment. This peculiar phenomenon was also noticed previously and attributed to possible errors in the experimental measurements by Slack *et al.*⁴⁴ From our work, considering that the filled B19 is only 0.8 Å away from the nearby B18 site in low energy configurations, we suggest that the lower occupancy of B18 from experiment may be correlated with the higher occupancy of B19, due to the incorrect assignment of these two close-lying sites. Assuming that B17 and B18 must have the same occupancy rate, the B18 and B19 occupancies need to be adjusted to 9.7% and 4.7%, respectively for MG179, which then agrees nicely with the theoretical results at 1500 K (B18: 10.1% and B19: 4.6%).

Finally, we note that the occupancy of B20 in all experimental samples is below 3%, which is similar to that of the residual boron atoms (2–4%), whose locations are difficult to assign⁴⁴ (see Table 3). Indeed, our theoretical prediction for B20 is also below 3%, and the occupancies of the residual boron atoms, now assigned to the B21 and B22 positions, also range from 0–4% at different temperatures.

c Electronic structure of β -B

With the PES of β -B having been clarified, we are now in a position to discuss the electronic structure and bonding of β -B in the context of previous knowledge on boron chemistry. In fact, the instability of perfect (undoped) β -B₁₀₅ has been addressed in many previous papers,^{3,43,45–47} and is attributed to the electron deficiency of the B₁₂ cage^{45,46} and the electron abundance of the perfect B₂₈ cage.^{43,47} Each doping B atom inside the hexagonal B ring can transfer its three electrons into the neighbouring B₁₂ cage *via* multi-center bonding, which converts three two-center bonds into three three-center bonds^{14,48} (our wavefunction analyses of the doping B also

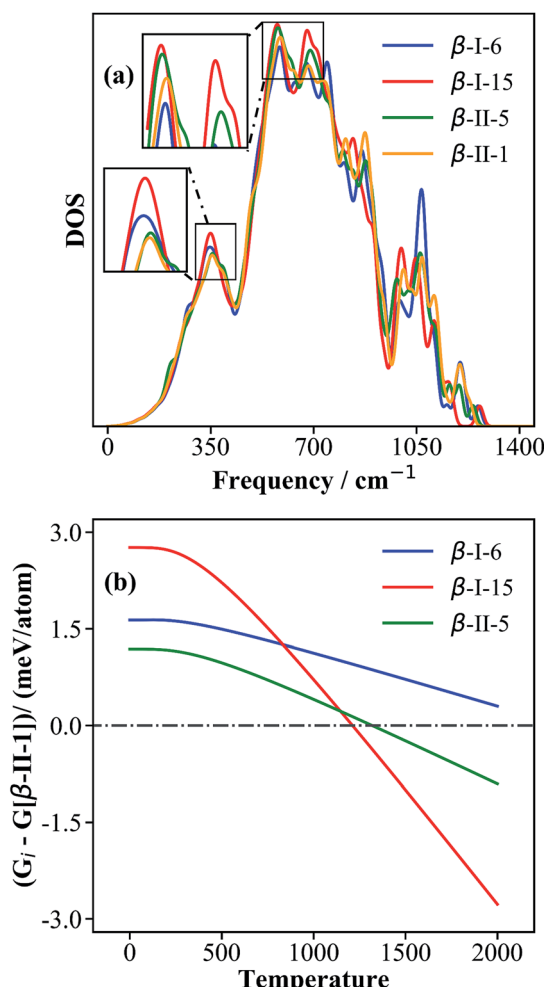


Fig. 6 The phonon density of states (a) and the relative free energies (b) for the key configurations, β -I-6, β -I-15, β -II-5 and β -II-1. The relative free energy (G_i in eqn (13)) is with respect to that of β -II-1, $G[\beta$ -II-1].



confirm this picture, see ESI Fig. S5†). Naturally, this doping atom may directly come from the B13 site of the B₂₈ cage. Both the B13 vacancy and the population of the B17 and B18 sites are beneficial for decreasing the electron abundance of the B₂₈ cage.^{14,43} Indeed, the low energy minima found in this work are consistent with previous understanding: (i) the doping atom is located either in a hexagonal ring (B16, B19, B20 and B21) or in a void (B22) surrounded by B₁₂ and B₂₈ cages, and generally achieves multi-center bonding with its neighbors; and (ii) all three skeletons (**SK-I** to **SK-III**) of β -B contain B13 vacancies: one in **SK-I**; two in **SK-II** and **SK-III**. Thus, the low energy β -B₁₀₆ structures prefer **SK-I**, while the low energy β -B₁₀₇ structures adopt **SK-II**.

Since the B19 and B16 doping sites play key roles in the temperature dependence of POS occupancy, we have further analyzed the electronic differences between them by taking β -I-15 as an example. For β -I-15, our Bader charge analyses show that the net charges of filled B16 and B19 are +0.35 and +0.26, respectively, which demonstrates that a filled B16 near three B₁₂ cages can donate more electrons than a filled B19 near two B₁₂ and one B₂₈ cages. Consistent with this, the center of the B16 2p band occurs at -6.35 eV below the valence band maximum (VBM), while that of the B19 2p band is at -6.02 eV below the VBM (see ESI Fig. S5†). All this information suggests that B16 interacts more strongly with the nearby B₁₂ cages and restricts the soft motion of the cages. This would in turn reduce the phonon density in the low frequency region (500 – 700 cm⁻¹) and lead to smaller entropy due to the B16 doping. On the other hand, B19 has a flatter PES with weaker bonding to nearby cages, which results in a higher entropy.

4. Conclusions

The ground state structure of boron crystals has long been a fascinating but challenging topic in fundamental science. Here, with newly developed machine learning methods to exhaustively explore the PES of boron with the SSW global optimization method, we are now able to establish the energy spectrum of all low energy configurations within the huge configuration space of β -B, which allows us to resolve in fine detail the β -B atomic structure at different temperatures. The structure determination paves the way towards a deep understanding and an accurate prediction of the physicochemical properties of boron crystals.

In total, 15 293 unique β -B configurations are found, but only 40 structures are within ~ 7 meV per atom above the global minimum. These low energy structures belong to three types of skeletons, namely **SK-I**, **SK-II** and **SK-III**, and can be further classified into six patterns, **P**₁ to **P**₆. The occupancy rates of different POSs are then derived and are found to be highly temperature dependent. It is the large vibrational entropy of the **P**₄ configurations that matters: **P**₄ becomes the dominant configuration at high temperatures. We demonstrate that, in contradiction to the long-held belief that β -B exhibits a huge energy degeneracy, only 20 configurations are of significance in the observed boron structures, and contribute the most ($>95\%$) to the overall POS occupancies.

In addition to new boron chemistry, this work also made great progress in methodology development for machine learning potentials. The great complexity of the boron global PES not only provides an excellent opportunity for developing sensitive structural descriptors, the key tool for correlating a structure with its quantitative properties (e.g. energy), but also enables us to identify the bottleneck and the key ingredients for generating high-dimensional NN potentials for complex materials. Major achievements are outlined below.

(i) This work reports a series of new structural descriptors, PTSDs, for describing the complex atomic bonding environment in boron. The PTSDs improve greatly the predictive power of the NN potential, which manages to reach an RMS accuracy of 12 meV per atom for energy and 0.28 eV Å⁻¹ for force for the global dataset (>160 000 structures spanning over 4 eV per atom in energy).

(ii) The first principles boron dataset established here, as also provided in the ESI,† can be utilized as a standard dataset for sharing, benchmarking and improving machine learning methods for building PESs in future.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by the National Key Research and Development Program of China (2018YFA0208600), the National Science Foundation of China (21533001, 91745201, 91645201 and 21603035), the Science and Technology Commission of Shanghai Municipality (08DZ2270500) and the Shanghai Pujiang Program (16PJ1401200).

Notes and references

- 1 W. N. Lipscomb, *Science*, 1977, **196**, 1047.
- 2 B. Albert and H. Hillebrecht, *Angew. Chem., Int. Ed.*, 2009, **48**, 8640–8668.
- 3 T. Ogitsu, E. Schwegler and G. Galli, *Chem. Rev.*, 2013, **113**, 3425–3449.
- 4 M. A. White, A. B. Cerqueira, C. A. Whitman, M. B. Johnson and T. Ogitsu, *Angew. Chem., Int. Ed.*, 2015, **54**, 3626–3629.
- 5 M. H. S. Segler, M. Preuss and M. P. Waller, *Nature*, 2018, **555**, 604.
- 6 M. K. Nielsen, D. T. Ahneman, O. Riera and A. G. Doyle, *J. Am. Chem. Soc.*, 2018, **140**, 5004–5008.
- 7 J. P. Janet and H. J. Kulik, *Chem. Sci.*, 2017, **8**, 5137–5152.
- 8 K. Yao, J. E. Herr, D. Toth, R. McKintyre and J. Parkhill, *Chem. Sci.*, 2018, **9**, 2261–2269.
- 9 D. E. Sands and J. L. Hoard, *J. Am. Chem. Soc.*, 1957, **79**, 5582–5583.
- 10 A. Masago, K. Shirai and H. Katayama-Yoshida, *Phys. Rev. B*, 2006, **73**, 104102.
- 11 S. Shang, Y. Wang, R. Arroyave and Z.-K. Liu, *Phys. Rev. B*, 2007, **75**, 092101.

12 M. J. van Setten, M. A. Uijttewaal, G. A. de Wijs and R. A. de Groot, *J. Am. Chem. Soc.*, 2007, **129**, 2458–2465.

13 M. Widom and M. Mihalkovič, *Phys. Rev. B*, 2008, **77**, 064113.

14 T. Ogitsu, F. Gygi, J. Reed, Y. Motome, E. Schwegler and G. Galli, *J. Am. Chem. Soc.*, 2009, **131**, 1903–1909.

15 T. Ogitsu, F. Gygi, J. Reed, M. Udagawa, Y. Motome, E. Schwegler and G. Galli, *Phys. Rev. B*, 2010, **81**, 020102.

16 D. Rogers and M. Hahn, *J. Chem. Inf. Model.*, 2010, **50**, 742–754.

17 M. Rupp, A. Tkatchenko, K.-R. Müller and O. A. von Lilienfeld, *Phys. Rev. Lett.*, 2012, **108**, 058301.

18 T. B. Hughes, N. L. Dang, G. P. Miller and S. J. Swamidass, *ACS Cent. Sci.*, 2016, **2**, 529–537.

19 K. T. Schütt, F. Arbabzadah, S. Chmiela, K. R. Müller and A. Tkatchenko, *Nat. Commun.*, 2017, **8**, 13890.

20 J. N. Wei, D. Duvenaud and A. Aspuru-Guzik, *ACS Cent. Sci.*, 2016, **2**, 725–732.

21 H. Altae-Tran, B. Ramsundar, A. S. Pappu and V. Pande, *ACS Cent. Sci.*, 2017, **3**, 283–293.

22 R. Gómez-Bombarelli, J. N. Wei, D. Duvenaud, J. M. Hernández-Lobato, B. Sánchez-Lengeling, D. Sheberla, J. Aguilera-Iparraguirre, T. D. Hirzel, R. P. Adams and A. Aspuru-Guzik, *ACS Cent. Sci.*, 2018, **4**, 268–276.

23 M. H. S. Segler, T. Kogej, C. Tyrchan and M. P. Waller, *ACS Cent. Sci.*, 2018, **4**, 120–131.

24 J. Behler and M. Parrinello, *Phys. Rev. Lett.*, 2007, **98**, 146401.

25 J. Behler, *J. Chem. Phys.*, 2011, **134**, 074106.

26 S.-D. Huang, C. Shang, X.-J. Zhang and Z.-P. Liu, *Chem. Sci.*, 2017, **8**, 6327–6337.

27 C. Shang, X.-J. Zhang and Z.-P. Liu, *Phys. Chem. Chem. Phys.*, 2014, **16**, 17845–17856.

28 C. Shang and Z.-P. Liu, *J. Chem. Theory Comput.*, 2013, **9**, 1838–1845.

29 S. Kirkpatrick, C. D. Gelatt and M. P. Vecchi, *Science*, 1983, **220**, 671.

30 D. J. Wales and H. A. Scheraga, *Science*, 1999, **285**, 1368.

31 D. M. Deaven and K. M. Ho, *Phys. Rev. Lett.*, 1995, **75**, 288–291.

32 H.-J. Zhai, Y.-F. Zhao, W.-L. Li, Q. Chen, H. Bai, H.-S. Hu, Z. A. Piazza, W.-J. Tian, H.-G. Lu, Y.-B. Wu, Y.-W. Mu, G.-F. Wei, Z.-P. Liu, J. Li, S.-D. Li and L.-S. Wang, *Nat. Chem.*, 2014, **6**, 727.

33 X.-J. Zhang, C. Shang and Z.-P. Liu, *J. Chem. Theory Comput.*, 2013, **9**, 3252–3260.

34 S.-C. Zhu, S.-H. Xie and Z.-P. Liu, *J. Am. Chem. Soc.*, 2015, **137**, 11532–11539.

35 W.-N. Zhao, S.-C. Zhu, Y.-F. Li and Z.-P. Liu, *Chem. Sci.*, 2015, **6**, 3483–3494.

36 S.-H. Guan, X.-J. Zhang and Z.-P. Liu, *J. Am. Chem. Soc.*, 2015, **137**, 8010–8013.

37 A. R. Oganov, J. Chen, C. Gatti, Y. Ma, Y. Ma, C. W. Glass, Z. Liu, T. Yu, O. O. Kurakevych and V. L. Solozhenko, *Nature*, 2009, **457**, 863.

38 G. Kresse and J. Furthmüller, *Comput. Mater. Sci.*, 1996, **6**, 15–50.

39 J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1997, **78**, 1396.

40 X.-J. Zhang, C. Shang and Z.-P. Liu, *Phys. Chem. Chem. Phys.*, 2017, **19**, 4725–4733.

41 W. H. Han, Y. J. Oh, D.-H. Choe, S. Kim, I.-H. Lee and K. J. Chang, *NPG Asia Mater.*, 2017, **9**, e400.

42 P. J. Steinhardt, D. R. Nelson and M. Ronchetti, *Phys. Rev. B*, 1983, **28**, 784–805.

43 E. D. Jemmis and M. M. Balakrishnan, *J. Am. Chem. Soc.*, 2001, **123**, 4324–4330.

44 G. A. Slack, C. I. Hejna, M. F. Garbauskas and J. S. Kasper, *J. Solid State Chem.*, 1988, **76**, 52–63.

45 W. H. Eberhardt, B. Crawford and W. N. Lipscomb, *J. Chem. Phys.*, 1954, **22**, 989–1001.

46 H. C. Longuet-Higgins and M. D. Roberts, *Proc. R. Soc. London, Ser. A*, 1955, **230**, 110.

47 M. Tillard-Charbonnel, A. Manteghetti and C. Belin, *Inorg. Chem.*, 2000, **39**, 1684–1696.

48 W. N. Lipscomb, *Boron hydrides*, Courier Corporation, 2012.

