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Theoretical trend of ion exchange ability with divalent cations in hydroxyapatite

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First-principles calculations are performed for hydroxyapatite (Hap) in order to investigate the relative ion exchange ability with divalent cations such as Mg^{2+} , Ni^{2+} , Cu^{2+} , Zn^{2+} , Cd^{2+} , Ba^{2+} , and Pb^{2+} . Their ionic substitutional energies are calculated from total energies of supercells and chemical potentials for Ca^{2+} and the foreign cations determined under chemical equilibrium between HAp and its saturated solution. It is found that, in most cases, the ion exchange ability is basically dependent on the ionic sizes and the larger or smaller-sized cations than Ca^{2+} tend to exhibit more difficulty of substitution for Ca^{2+} in HAp. However, Pb^{2+} exhibits the extremely small substitutional energy, which originates from covalent bonds with the adjacent oxygen atoms. In particular, Pb^{2+} at the Ca-1 site of HAp has covalency with the second-nearest-neighboring oxygen atoms as well as the first nearest-neighboring ones. The covalent bond formation plays an important role for the distinct ion exchange ability for Pb^{2+} by HAp observed experimentally.

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I. INTRODUCTION

Hydroxyapatite (HAp) $[Ca_{10}(PO_4)_6(OH)_2]$ is now attracting much attention in materials science due to its practical application to biomaterials. In addition, since HAp has versatile physical and chemical properties, it is expected to be utilized as catalysts, chromatographic adsorbents for protein separation, scaffold materials in tissue engineering, and so on. One of the interesting properties of HAp is an ion exchange ability for various foreign cations and anions and would be useful to remove harmful ions in waste water.^{1,2} It is known that HAp minerals in human bones also contain a variety of foreign ions.³ The incorporated impurities and dopants in HAp are successively released into or uptake from the surrounding body fluids during the bone remodeling processes. It can be said, therefore, that the ion exchange of HAp is also a key factor to determine the biological properties of HAp.

There were a number of experimental reports on ion exchange properties of synthetic HAp materials.^{1–7} Suzuki *et al.*¹ reported the ranking of ion exchange ability of Cd^{2+} , $Zn^{2+} > Ni^{2+} > Ba^{2+}$, Mg^{2+} and showed the ranking of $Pb^{2+} > Cu^{2+} > Mn^{2+} \cong Co^{2+}$ in their separate paper.² Takeuchi and Arai⁴ also investigated the removal of Pb^{2+} , Cu^{2+} , and Cd^{2+} by synthetic HAp and found that Pb^{2+} ions are more readily incorporated into HAp by exchange with Ca^{2+} , as compared to Cu^{2+} , and Cd^{2+} . As a general trend, it seems that cations having the larger atomic numbers tend to be more easily exchanged with Ca^{2+} in HAp.

The difference in ion exchange ability of divalent cations was previously interpreted by ionic sizes and electronegativity of the exchanged cations. From simple geometrical considerations, it is easily expected that cations having similar ionic radii with Ca^{2+} (0.10 nm in the sixfold coordination⁸) can be favorably exchanged with Ca^{2+} in HAp. Suzuki *et al.*² argued that cations easily uptake from an aqueous solution into HAp tend to have ionic radii of around 0.09–0.13 nm.

Moreover, they also mentioned the tendency of the good ion exchange ability for cations having much higher electronegativity than Ca^{2+} , which may suggest the importance of the detailed chemical bonding states around the exchanged cations. Recently, Zhu *et al.*⁹ studied the atomic structures of HAp solid solutions with Pb²⁺, Sr²⁺, and Cd²⁺ and argued the site preference in the HAp lattice based on the ionic sizes and electronegativity. Although the ionic sizes and electronegativity are convenient measures to qualitatively interpret the difference in ion exchange ability, however, it is desirable to reveal a physical and chemical origin of the ion exchange ability in more detail. In this regard, first-principles calculations have been proven to be suitable for studying electronic structures and energetics of substitutional defects in materials.

In this study, various kinds of substitutional divalent cations in HAp are calculated in a first-principles manner, and the defect formation energies by ion exchange with Ca²⁺ in HAp are evaluated from total energies of supercells and chemical potentials determined under solid-liquid equilibrium. It is noted that this type of calculations corresponds to bulk substitutions of foreign cations in HAp. Since actual ion exchange takes place through interfaces between HAp and aqueous solution, structural characteristics of HAp surfaces in contact with solution may affect absorption and incorporation of foreign cations into HAp.¹⁰ Nevertheless, it is beneficial, as a first step, to reveal essential acceptability of foreign cations by bulk HAp via ion exchange from calculations of the bulk substitutions.

A number of first-principles studies on substitutional defects in HAp have been reported so far.^{11–17} However, most of the previous studies investigated atomic and electronic structures around substitutional defects and did not systematically reveal the trend in stability of foreign cations substituting for Ca^{2+} in HAp by ion exchange. The results obtained by thermodynamic treatments of first-principles total energies are compared with available experimental data, and the

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origin of the ion exchange ability of HAp will be discussed in terms of atomic and electronic structures localized around the exchanged divalent cations.

II. COMPUTATIONAL METHODS

The projector-augmented-wave (PAW) method is used for electronic structure calculations of substitutional defects in HAp using the VASP code.^{18–20} In the cases of substitutional transition-metal ions having unfilled *d* orbitals (Ni²⁺ and Cu²⁺), spin polarization is also taken into account. The generalized gradient approximation (GGA) for the exchange-correlation potential is employed.²¹ Electronic wave functions are expanded by plane waves up to a cutoff energy of 500 eV. Brillouin-zone integration is performed on *k*-point meshes by the Monkhorst-Pack scheme.²² All atoms in supercells for defect calculations are relaxed, and the structural optimization is truncated when their forces converge to less than 0.05 eV/Å.

The present study deals with eight kinds of substitutional divalent cations in HAp such as Mg²⁺, Ni²⁺, Cu²⁺, Zn²⁺, Sr²⁺, Cd²⁺, Ba²⁺, and Pb²⁺. This is mainly because the ion exchange behavior for these ions by HAp was experimentally examined¹⁻⁷ and also the inclusion of many of these ions in HAp are known to play an important biological role. It is noted that the substitutional cases of Mg²⁺ and Zn²⁺ ions in HAp were already reported in our previous paper¹⁷ and the results are also partly used here for comparison. For the defect calculations, 352-atom supercells generated from the hexagonal crystal structure of HAp are used,^{3,23} and one of the Ca ions in the perfect supercell is replaced by a particular foreign cation to generate the defective supercell. More details of the supercell construction were described elsewhere.^{17,24} In the supercell calculations, only the Γ point is used for the Brillouin-zone sampling.

Two inequivalent Ca sites, Ca-1 and Ca-2, in HAp are replaced by the substitutional cations. Local atomic configurations of Ca-1 and Ca-2 are illustrated in Fig. 1. The Ca-1 site is surrounded by six PO_4^{3-} tehrahedra, where six oxygen ions at the vertices of PO_4^{3-} are located at the first nearestneighboring (NN) sites and additional three oxygen ions are present at the second NN sites. The Ca-2 site is also coordinated by six oxygen atoms at the first NN sites, and yet one of the first NN oxygen atoms belongs to the OH group. In addition, one oxygen atom is located at the second NN site. It is noted, therefore, that the atomic coordination difference between the two Ca sites is sometimes described by the oxygen coordination numbers within the second NN atomic coordination shell, the ninefold coordination for Ca-1 and the sevenfold for Ca-2.

From total energies $E_{\rm T}$ of the HAp supercells with or without the substitutional defects, the defect formation energies ($\Delta H_{\rm f}$) are evaluated. When a substitutional ionic species M^{2+} is exchanged by a Ca²⁺ in HAp, $\Delta H_{\rm f}$ can be given by

$$\Delta H_{\rm f} = E_{\rm T}(\text{defective}) - E_{\rm T}(\text{perfect}) + \mu_{\rm Ca^{2+}} - \mu_{\rm M^{2+}}.$$
 (1)

It is noted here that the substitutional cations are isovalent with Ca^{2+} , and thus additional charge compensating defects do not need to be considered for the ion exchange.



FIG. 1. (Color online) (a) Crystal structure of HAp viewed along the c axis, (b) local atomic structures of Ca-1, and (c) Ca-2. In the illustrations of (b) and (c), the first NN oxygen ions are indicated by connecting with the Ca ions with the solid lines, while the second NN oxygen ions with the broken lines.

 $\mu_{Ca^{2+}}$ and $\mu_{M^{2+}}$ in Eq. (1) indicate chemical potentials of Ca^{2+} and M^{2+} , which are determined from a particular chemical equilibrium. In this study, it is assumed that HAp crystals equilibrate with an aqueous solution saturated with respect to HAp and containing a particular amount of M^{2+} ions. According to the thermodynamic treatment for ions in solution,^{25,26} the chemical-potential difference in Eq. (1) can be expressed as



FIG. 2. Equilibrium Ca^{2+} concentration in the saturated aqueous solution as a function of solution pH.

$$\mu_{\text{Ca}^{2+}} - \mu_{\text{M}^{2+}} = \mu_{\text{Ca}^{2+},\text{aq}}^{\circ} - \mu_{\text{M}^{2+},\text{aq}}^{\circ} + k_{\text{B}}T \ln(a_{\text{Ca}^{2+}}/a_{\text{M}^{2+}}),$$
(2)

where $\mu_{M^{2+},aq}$ and $a_{M^{2+}}$ are the standard chemical potential and the activity of a cation in solution, respectively. k_B is the Boltzmann constant and a temperature of *T* is set at 298 K throughout the present study. The $\mu_{Ca^{2+},aq}^{\circ} - \mu_{M^{2+},aq}^{\circ}$ term is further written using the standard Gibbs formation energies for ions in aqueous solution $\Delta G_{f}^{\circ}(M^{2+},aq)$ as¹⁷

$$\mu^{\circ}_{Ca^{2+},aq} - \mu^{\circ}_{M^{2+},aq} = \Delta G^{\circ}_{f}(Ca^{2+},aq) - \Delta G^{\circ}_{f}(M^{2+},aq) + \mu^{\circ}_{Ca,s} - \mu^{\circ}_{M,s}.$$
(3)

The experimental $\Delta G_{\rm f}^{\circ}$ values are tabulated in the thermodynamic data, which are used in this study.²⁷ On the other hand, $\mu_{\rm Ca,s}^{\circ}$ and $\mu_{\rm M,s}^{\circ}$ indicate chemical potentials of solid Ca and metal M. In this case, these quantities are considered to be the total energies per atom for the pure metals having stable crystal structures in the standard state, which are also calculated in a first-principles manner.

Regarding the ionic activity term of Eq. (2), the activity coefficients are assumed to be unity in this study. This is because HAp is sparing soluble and its saturated solution can be regarded as a dilute solution. The ionic concentrations of impurity ions $[M^{2+}]$ are assumed to be a particular value, and then the calcium concentration in solution $[Ca^{2+}]$ is determined from the solubility of product for HAp, the ionic product of water, three different acid dissociation constants for phosphate ions $(HPO_4^{2-}=PO_4^{3-}+H^+, H_2PO_4^{-}=HPO_4^{2-}+H^+,$ and $H_3PO_4 = H_2PO_4^- + H^+$, and charge-neutrality requirement in solution. More details of the computational procedure to evaluate concentrations of ionic species in the saturated solution can be found elsewhere.^{17,24} In Fig. 2, only the calcium concentration in solution $[Ca^{2+}]$ as a function of pH is depicted. Since [Ca2+] depends on the solution pH value,17,24,28 the defect formation energies calculated from Eq. (1) also vary against the pH condition of the aqueous solution.



FIG. 3. Interatomic distances from the Ca ions to the neighboring oxygen ions against the ionic radii of the substituted cations. For the ionic radii, the experimental values in the sixfold coordination are used (Ref. 8). The distances to the first NN oxygen ions are plotted by open circles, while those to the second NN oxygen by open squares.

III. RESULTS AND DISCUSSION

A. Structural characteristics around exchanged cations

When foreign divalent ions M²⁺ are substituted for Ca²⁺ of HAp, the ionic sizes are generally different from that of Ca^{2+} , and thus the surrounding ions may undergo atomic relaxation. In this section, therefore, characteristics of the atomic structures around the substitutional cations in HAp are investigated. Figure 3 displays the calculated interatomic distances between M²⁺ and its surrounding oxygen ions as a function of the ionic radii. In this case, the Ca-1 and Ca-2 sites in the perfect HAp lattice are basically sixfold coordinated by oxygen ions at the first NN sites, and thus the experimental ionic radii in the sixfold coordination by Shannon are used.⁸ In perfect HAp, the Ca-1 ion is coordinated by three of the first NN oxygen ions with a bond length of 0.244 nm, and the other three are located at 0.248 nm in distance from Ca-1. At the second NN sites, three oxygen ions are also present with a bond length of 0.280 nm. On the other hand, the local atomic configuration of Ca-2 is more complicated than that of Ca-1. The Ca-2 ion is also surrounded by six oxygen ions at the first NN sites, and yet their distances from Ca-2 vary from 0.236 to 0.252 nm. An oxygen ion at the second NN site is situated at a distance of 0.281 nm from Ca-2.

When the divalent foreign ions replace Ca-1, the interatomic distances from Ca-1 strongly depend on their ionic sizes [see Fig. 3(a)]. For the smaller-sized cations of Ni²⁺, Mg^{2+} , Cu^{2+} , and Zn^{2+} , the first NN oxygen ions tend to considerably move toward Ca-1 (more than 7% relaxation on average), whereas the second NN oxygen ions undergo outward relaxation by more than 10% in distance. This can be explained from the local atomic configuration of Ca-1, as shown in Fig. 1(b). The first and second NN oxygen ions are situated at the vertices of the adjacent PO_4^{3-} tetrahedra, and substitution of the smaller-sized cations brings about inward displacements of the PO_4^{3-} groups toward the Ca-1 site. However, the inward relaxation of the surrounding PO_4^{3-} groups simultaneously accompanies close proximity of the oxygen ions, and then the PO_4^{3-} groups rotate so as to minimize electrostatic repulsions between the oxygen ions at their vertices. As a result, the second NN oxygen ions tend to move away from the Ca-1 site, as shown in Fig. 3(a). Such a behavior of the surrounding ions can be found also in the Cd²⁺ case, although the atomic relaxations around Cd²⁺ are much smaller due to its similar ionic radius with Ca^{2+} .

In the cases of the larger-sized cations $(Sr^{2+}, Pb^{2+}, Ba^{2+})$ at the Ca-1 site, the first NN oxygen ions displace their positions away from the Ca-1 site. For instance, the Ba-O distances at the first NN sites are by about 9% larger than those of Ca-O. It is noted, however, that the distances of the second NN oxygen ions from the larger cations at the Ca-1 site are almost the same with that from Ca-1 in perfect HAp.

Interatomic distances of the foreign cations at Ca-2 vary in the similar way to the Ca-1 substitutional case, depending on the ionic radii [Fig. 3(b)]. As stated in Sec. II, however, the atomic configuration at Ca-2 in the original HAp lattice exhibits lower symmetry than at Ca-1, and thus the substituted cations exhibit somewhat wide variations in the interatomic distances with the first NN oxygen ions. As a result, some of the smaller-sized cations tend to have smaller coordination numbers to oxygen at the first NN sites, as compared to the case in perfect HAp (sixfold coordination). In the cases of Cu^{2+} and Zn^{2+} , their coordination numbers to the first NN oxygen ions are four with bond lengths of 0.207-0.213 nm, and the other oxygen ions at the sites beyond the second NN coordination shell are located at a distance of more than 0.24 nm. The decreased coordination numbers of Cu^{2+} and Zn^{2+} can be understood from the fact that these ions have the fourfold coordination with oxygen in bulk CuO and ZnO. Likewise, the coordination number of Ni²⁺ to the first NN oxygen is found to be 5, and their bond lengths range from 0.210 to 0.221 nm. Ni2+ ions favor the sixfold coordination to oxygen in bulk NiO, but the coordination number decreases from 6 to 5 in HAp, because of its extremely small ionic size and the resultant electrostatic repulsions between the surrounding oxygen ions.

B. Energetics of ionic substitution

Substitutional energies of different divalent cations M^{2+} by exchange with Ca^{2+} in HAp are evaluated. As a typical



FIG. 4. Ionic substitutional energies at the Ca-1 site of HAp against the solution pH.

example, the ionic substitutional energies at the Ca-1 site are plotted against pH in Fig. 4. These results are obtained under assumption of the chemical equilibrium of HAp with the HAp-saturated aqueous solution having the impurity concentration $[M^{2+}]$ of 1.0×10^{-3} mol/1 at 298 K. It is noted here that the substitutional energies also depend on $[M^{2+}]$ in the solution. It is confirmed that the substitutional energies for all impurities systematically decrease by about 0.12 eV in the case of the larger concentration $[M^{2+}]=1.0 \times 10^{-1}$ mol/1 whereas increase by around 0.18 eV for the smaller concentration $[M^{2+}]=1.0 \times 10^{-6}$ mol/1.

It can be seen that the ionic substitutional energies tend to increase with decreasing pH. This is because the Ca^{2+} concentration in the surrounding saturated solution ($[Ca^{2+}]$) becomes larger with decreasing solution pH, as shown in Fig. 2. Owing to such pH dependence of $[Ca^{2+}]$ on the solution, Ca^{2+} ions are more difficult to be released from HAp into the solution by the ion exchange in the lower pH condition so that the ionic substitutional energies become larger in more acidic conditions.^{17,24} Among the foreign cations, Pb²⁺, Cd²⁺, and Sr²⁺ exhibit the much smaller substitutional energies and, in particular, the substitutional energies of Pb²⁺ and Cd²⁺ are negative in the almost entire pH range.

As a matter of course, the ionic substitutional energies at the Ca-2 site exhibit the same pH dependence with those at Ca-1, but the absolute energy values are slightly different depending on the substitutional sites. In order to also show the Ca-site dependence of the substitutional energies, the calculated values at pH=7 (neutral condition) at the two independent Ca sites are plotted against the ionic radii of the foreign cations in Fig. 5, where the ionic radii in the sixfold coordination are used.⁸

It is found that the variations in the substitutional energies against the ionic radii for the Ca-1 and Ca-2 substitutions are quite similar to each other. It is obvious that the smaller- or larger-sized cations than Ca^{2+} exhibit larger substitutional energies while substitution of Cd^{2+} , Sr^{2+} , and Pb^{2+} ions relatively similar in size to Ca^{2+} is energetically more favored. This can easily be imagined from simple consideration of ionic size mismatches between the foreign ions and Ca^{2+} . As



FIG. 5. Ionic substitutional energies at the Ca-1 and Ca-2 sites of HAp against the ionic radii. The energy values correspond to the ones at pH=7 (see also Fig. 4).

displayed in Fig. 3, the smaller- and larger-sized foreign cations induce larger lattice relaxation of their surrounding ions by substitution and then suffer more significant elastic energy expenses.

It should be mentioned here, however, that the Pb²⁺ substitution exhibits the exceptionally small substitutional energy, even compared to the Sr^{2+} substitution with the quite similar ionic size. In such a case, the simple consideration of the ionic sizes is not available, and thus further detailed analyses of the electronic structures would be required. For this purpose, the partial densities of states (PDOS) of Pb²⁺ at Ca-1 and its surrounding first NN oxygen ions are displayed in Fig. 6. For comparison, the PDOS curves for Ca^{2+} , Sr^{2+} , and Cd²⁺ at the Ca-1 site are also shown. As stated in our previous studies, the upper valence band (VB) of perfect HAp is composed mainly of O 2p orbitals, and the lower VB comprises O 2s and Ca 3p orbitals. For chemical bonding states in HAp, orbital overlap in the upper VB would be important, and the originally unoccupied Ca 3d4s orbitals only slightly contribute to the upper VB [see Fig. 6(a)], which indicates a typical ionic nature of bonding between Ca-O. This is also the case for Sr^{2+} and Cd^{2+} in Figs. 6(b) and 6(c). It is noted that, in the case of Cd^{2+} , the Cd 4dorbitals considerably overlap with the O 2p orbitals in the lower part of the upper VB, but the Cd-4d orbitals are originally fully occupied by electrons because the formal electronic configuration of Cd^{2+} is described as $[Kr]4d^{10}5s^0$. Therefore, the orbital overlap between Cd 4d and O 2p does not directly contribute covalent interactions between Cd and О.

In the case of Pb²⁺, the original electronic configuration can be described as $[Hg]6s^26p^0$, and the electron-occupied 6s orbitals appear in the lower part of the upper VB. It is worth mentioning here that the unoccupied 6p orbitals are more admixed with O 2p in the upper part of the upper VB, as compared to the Ca²⁺, Sr²⁺, and Cd²⁺ cases. It can be expected, therefore, that covalent interactions with the surrounding oxygen ions may be present in the Pb²⁺ substitution.

The role of Pb 6sp orbitals for covalent bond formation has been qualitatively discussed.^{29,30} In an isolated state of



FIG. 6. PDOS curves of the cations and their first NN oxygen ions in HAp. In these plots, the valence-band maxima are set at 0 eV.

Pb²⁺, two 6*s* electrons can simply be described as an inactive lone pair. However, depending on the atomic coordination environment, the unoccupied Pb 6*p* orbitals are admixed with the 6*s* orbitals. This results in formation of a "stere-ochemically active" lone pair.^{29,30} The active lone pair can strongly interact with O 2*p* valence electrons in a bonding manner so that significant covalent bond formation around Pb²⁺ can be realized. This situation actually corresponds to the partial DOS curves in Fig. 6(d), where the Pb 6*sp* components are well admixed with the O 2*p* components in the upper VB.

Figure 7 depicts contour maps of electron densities around Pb^{2+} and Sr^{2+} at the Ca-1 site on different cross sectional planes [(1100) for (a) and (c) and (0001) for (b) and (d)]. It is noted that, in Fig. 7(a), Ca²⁺ and Pb²⁺ ions at the Ca-1 sites and their first NN oxygen ions are located close to



FIG. 7. Contour maps of electron densities on the [(a) and (c)] $(1\overline{100})$ and [(a) and (c)] (0001) planes for HAp with Pb²⁺ and Sr²⁺ at the Ca-1 sites. The contour lines are drawn from 0.00 to 0.50 with an interval of 0.05e/ Å (Ref. 3).

the cross sectional plane while the cations and their second NN oxygen ions are present close to the cross section in Fig. 7(b). This is also the case for Sr^{2+} in Figs. 7(c) and 7(d).

It can be seen in Fig. 7(a) that more electrons tend to be accumulated between Pb-O, as compared to those in the interatomic region of Ca-O [see Fig. 7(a)]. Moreover, the Ca-1 ions do not exhibit apparent electron densities in the interatomic region with the second NN oxygen ions, but it is obvious that the doped Pb²⁺ ion also shows more electrondensity overlap with the second NN oxygen [see Fig. 7(b)]. These results are an indication of the strong tendency of covalent bond formation of Pb²⁺ with the surrounding oxygen atoms. In this regard, Ellis et al.³⁰ also investigated Pb substitution in HAp by the similar first-principles method and analyzed the local atomic coordination and chemical bonding state. They showed that Pb substitution in HAp cannot simply be described by substitution of a rigid Pb^{2+} ion but accompanies strong covalent interactions with the surrounding oxygen atoms, which is consistent with the present result. It is noted that relativistic effects on Pb valence orbitals were also examined by the above authors but were found not to significantly alter the covalent bonding character between Pb and O.³⁰

In the case of the Sr^{2+} substitution at the Ca-1 site [Fig. 7(c) and 7(d)], the ionic size of Sr^{2+} is quite similar to that of Pb²⁺ (see Fig. 3), but the substitutional energy is considerably larger than that of Pb²⁺ (see Fig. 5). In this case, the larger electron-density overlap between Sr and its surrounding oxygen can also be observed (especially with the second NN oxygen), as compared to that of Ca-O. However, the electron densities of Sr-O are still smaller than those of Pb-O. Therefore, the extremely stable substitution of Pb²⁺ can be attributed to the formation of strong covalent bonds with the adjacent oxygen atoms.

Based on the above results, the ion exchange ability by HAp is predicted to be in the order of $Pb^{2+}>Cd^{2+}>Sr^{2+}$

 $>Zn^{2+}>Cu^{2+}>Mg^{2+}>Ba^{2+}>Ni^{2+}$. The previous ion exchange experiments by Suzuki *et al.*^{1,2} also showed that Pb²⁺ and Cd²⁺ ions are more favorably uptaken by HAp from the solution. Yasukawa *et al.*³¹ discussed the ranking of preferential incorporation of divalent foreign cations into HAp based on their experimental data and indicated the order of Ca²⁺>Mg²⁺ and Pb²⁺>Ca²⁺>Sr²⁺>Ba²⁺. They also mentioned that Pb²⁺ and Cd²⁺ are, in particular, easy to be incorporated by exchange with Ca²⁺ in HAp. There were also a number of experimental studies synthesizing Pb₁₀(PO₄)₆(OH)₂, Sr₁₀(PO₄)₆(OH)₂, and Cd₁₀(PO₄)₆(OH)₂ by solution-precipitation methods.^{9,31-35} In this regard, the present results show that the substitutional energies of Pb²⁺, Cd²⁺, and Sr²⁺ are much smaller than those of the other ions (Figs. 4 and 5), whose tendency is consistent with the previous experimental results.

It is noted that the calculated ranking of the ion exchange ability except for Pb²⁺, Cd²⁺, and Sr²⁺ seems different from the experimental result of Cd^{2+} , $Zn^{2+} > Ni^{2+} > Ba^{2+}$, Mg^{2+} by Suzuki *et al.*^{1,2} It is plausible that the discrepancy may be related to the presence of a transition layer between HAp and the surrounding solution. As discussed by Tung *et al.*,³⁶ it is thought that surface structures of HAp in contact with an aqueous solution are different from the case in contact with vacuum and a disordered and/or hydrated transition layer may be present at the interface. Such a transition layer is expected to strongly affect uptake and exchange processes for foreign ions by HAp so that the transition layer may affect the apparent solubility of the cations. This would be especially true for the ions having the relatively large substitutional energies into HAp. There is a possibility that the cations cannot be incorporated into the HAp lattice but can be favorably included in the transition layer. In fact, Mg²⁺ and Zn²⁺ incorporations in the transition layer were investigated in our previous study, where octacalcuim phosphate (OCP) was used as a prototype for the transition layer.¹⁷ It was found that the substitutional energies in the transition layer were much smaller than in HAp, depending on the substitutional sites. Therefore, further studies will be required to make more quantitative discussion on the ranking of the cations having the high substitutional energies.

Regarding the site preference of the substituted cations in HAp, the present results in Fig. 5 indicate that the smallersized cations prefer the Ca-2 substitution while the larger cations favor the Ca-1 substitution except for Pb²⁺. Table I shows the available experimental data of the preferential occupation sites of divalent foreign cations.^{9,37-40} For Cd²⁺ and Pb^{2+} , an agreement between the present calculations and the experiment can be achieved, but there are the discrepancies for Sr²⁺ between theory and experiment. Kikuchi et al.³⁸ showed that Sr²⁺ prefers the Ca-1 site substitution in HAp, which is consistent with the present result. However, Hughes et al.³⁹ experimentally analyzed naturally occurring Srsubstituted apatite samples with more than about 3 mol % of Sr and showed that the Sr substitution is favored to the Ca-2 site. In this case, it is noted that the samples also contained other trace elements of sodium, silicon, and so on, which may affect the Sr substitutional behavior. More recent Rietveld analyses of solid solutions of Sr-Ca HAp by Zhu et al.⁹ also indicated that Sr²⁺ favors occupation of the Ca-2 site

TABLE I. Comparison of preferential substitutional site for divalent cations between experiment and the present study.

Cation species	Experiment	The present work
Cd ²⁺	Ca-2 ^a , ^b	Ca-2
Sr ²⁺	Ca-1, ^c Ca-2 ^a , ^d	Ca-1
Pb ²⁺	Ca-2 ^a , ^e	Ca-2

^aReference 9.

^bReference 37. ^cReference 38.

^dReference 39.

^eReference 40.

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in the range of higher Sr content, whereas Sr^{2+} does not show significant site preference in the lower Sr content. Strictly speaking, the present calculations correspond to the dilute substitution of Sr^{2+} in HAp, and thus it is not reasonable to make direct comparison of the present result of the site preference with the above-mentioned experiment with much higher Sr^{2+} content. As shown in Fig. 5, however, the difference in substitutional energies between the two Ca sites for Sr^{2+} is somewhat small (about 0.07 eV) and therefore Sr^{2+} may not exhibit pronounced site preference in the low Sr^{2+} content range, as shown in Ref. 9.

It is finally noted that the Sr content in HAp can be roughly calculated to be around 1 mol % in the range of pH=7-8 based on the result of Fig. 4.¹⁷ Although the calculated Sr content in HAp is much smaller than those by the previous experiments,⁹ it is worth mentioning that the equilibrium concentration may strongly depend on experimental conditions to produce the HAp materials. Even in the present case, as shown in Fig. 4, the Sr-substituted energy tends to decrease with increasing pH and $[Sr^{2+}]$ in solution. Therefore, more intense Sr substitution may be possible, depending on experimental conditions of the ion exchange process (more alkaline and higher $[Sr^{2+}]$ conditions). In the system with more Sr substitutions, however, Sr-Sr interactions and the distribution and configuration of substitutional Sr ions would become important to calculate the equilibrium Sr concentration, which needs to be correctly taken into account in future, from a theoretical viewpoint.

IV. CONCLUSIONS

First-principles calculations were performed for HAp, to study ion exchange ability of the divalent cations. Chemical equilibrium between HAp and its saturated solution was assumed, and the chemical potentials for the foreign cations then obtained were used to evaluate the ionic substitutional energies. It was found that the ion exchange ability is basically dependent on the ionic sizes. The larger or smallersized cations than Ca^{2+} tend to exhibit more difficulty of substitution for Ca in HAp. However, it is worth mentioning that Pb^{2+} exhibits the extremely small substitutional energy, which originates from covalent bond formation with the adjacent oxygen ions. Therefore, the detailed chemical bonding state plays an important role for the good ion exchange ability for Pb^{2+} by HAp.

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