A combinatory ferroelectric compound bridging simple ABO$_3$ and A-site-ordered quadruple perovskite

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The simple ABO$_3$ and A-site-ordered AA$'_{3}$B$_4$O$_{12}$ perovskites represent two types of classical perovskite functional materials. There are well-known simple perovskites with ferroelectric properties, while there is still no report of ferroelectricity due to symmetry breaking transition in A-site-ordered quadruple perovskites. Here we report the high pressure synthesis of an A-site-ordered perovskite PbHg$_3$Ti$_4$O$_{12}$, the only known quadruple perovskite that transforms from high-temperature centrosymmetric paraelectric phase to low-temperature non-centrosymmetric ferroelectric phase. The coordination chemistry of Hg$^{2+}$ is changed from square planar as in typical A-site-ordered quadruple perovskite to a rare stereo type with 8 ligands in PbHg$_3$Ti$_4$O$_{12}$. Thus PbHg$_3$Ti$_4$O$_{12}$ appears to be a combinatory link from simple ABO$_3$ perovskites to A-site-ordered AA$'_{3}$Ti$_4$O$_{12}$ perovskites, sharing both displacive ferroelectricity with former and structure coordination with latter. This is the only example so far showing ferroelectricity due to symmetry breaking phase transition in AA$'_{3}$B$_4$O$_{12}$-type A-site-ordered perovskites, and opens a direction to search for ferroelectric materials.
Ferroelectrics and their derivatives show many interesting physical, chemical and mineral properties such as ferromagnetism, ferroelectricity, piezoelectricity, ion conductivity, photocatalysis and superconductivity11-18 that can be modified dramatically by the coordination chemistry for a given composition. There is a special class of perovskite-type materials with the general chemical formula $\text{AA}'_3\text{B}_2\text{O}_{12}$, named A-site-ordered, or quadruple perovskites, which received much attention owing to their fascinating structural and wide varieties of physical properties10, including colossal magnetoresistance under weak field11, charge disproportionation12 and giant dielectric constant over a wide temperature range13,14. For the simple $\text{ABO}_3$ perovskite, the 12-fold coordinated A-site is often occupied by large size ions such as alkali metal, alkaline earth or lanthanide cations and the 6-fold coordinated B-site is often occupied by transition metals (TMs) to satisfy the so called tolerance factor $t = (r_A + r_b)/(\sqrt{2}(r_A + r_b))$ ($r_A$, $r_b$ and $r_O$ represent for the ion radius of A, B, and O, respectively) with $t$ usually in the range of 0.75 to 1.05 for stable perovskite compounds1,18. Generally, TM ions with strong Jahn-Teller distortions like Mn$^{3+}$ and Cu$^{2+}$ preferentially occupy the A′-site with square-planar coordination20-24. The small TM ion at the A′-site causes the distortion of BO$_6$ octahedron in $\text{AA}'_3\text{B}_2\text{O}_{12}$ perovskite, usually resulting in a cubic crystal structure with space group Im-3.25 From a structural point of view, the flexibility for tilting the BO$_6$ octahedra in $\text{AA}'_3\text{B}_2\text{O}_{12}$ perovskite is limited in order to maintain the square-planar coordination at the originally 12-fold coordinated A-site in the simple perovskite structure.21

Ferroelectricity is one of the most important properties for application in actuators, sensors, and memory storage devices, etc. Large numbers of ferroelectric materials possess simple perovskite structures, such as $\text{BaTiO}_3$, $\text{PbTiO}_3$, $\text{Pb(Ti, Zr)}_2\text{O}_3$, $\text{BiFeO}_3$.26-30 The origin of ferroelectricity is due to either the lone pair 6$d$ effect of cations (Pb$^{2+}$, Bi$^{3+}$), or the second-order Jahn-Teller active cations with electron configuration of $d^0$ (Ti$^{4+}$, Zr$^{4+}$); these compounds are usually referred to as displacive-type ferroelectrics. However, ferroelectricity is seldom observed in $\text{AA}'_3\text{B}_2\text{O}_{12}$ type A-site-ordered compounds, even when they contain second-order Jahn-Teller active cations or lone pair effect cations.28 One reason for the absence of ferroelectricity in this series of compounds is the unusual presence of a centrosymmetric structure, and the variation of temperature generally causes isostructural or centrosymmetric -to- centrosymmetric phase transition12,21,25, hence there cannot be spontaneous polarization. Currently the rare examples in this series that show ferroelectricity are $\text{AMn}_3\text{O}_{12}$ ($\text{A} = \text{Ca}$, Bi or Pb)31-33 and $\text{AMn}_3\text{Cr}_3\text{O}_{12}$ ($\text{A} = \text{La}$ or Bi)20,22. Although these compounds have centrosymmetric structures, all are spin-driven multiferroic systems, due to strong magnetoelectric coupling effects.20,22,32. Up to now, there is still no report of ferroelectricity from symmetry breaking transition in A-site-ordered perovskite compounds.

In this work, an $\text{AA}'_3\text{B}_2\text{O}_{12}$ type A-site-ordered perovskite oxide, $\text{PbHg}_3\text{Ti}_4\text{O}_{12}$ (henceforth PHTO) was designed and synthesized at high pressure and high temperature conditions. It is found that the Hg$^{2+}$ ion occupies the A′-site in $\text{AA}'_3\text{B}_2\text{O}_{12}$ type perovskite in sharp contrast to previous reports where 3$d$ TM ions ($\text{Cu}^{2+}$, Mn$^{3+}$) usually locate at the A′-site. Moreover, ferroelectricity was observed in PHTO, when its ambient centrosymmetric phase transformed to a non-centrosymmetric phase at 250 K. The origins of ferroelectricity are discussed based on the comprehensive characterizations of crystal structure using both neutron and synchrotron x-ray diffractions and density functional theory (DFT) calculation.

Results and Discussion

Crystal structure. The NPD refinements are shown in Fig. 1a, which confirm that PHTO crystallizes in the $\text{AA}'_3\text{B}_2\text{O}_{12}$ type A-site-ordered perovskite. The refined structure parameters of PHTO based on NPD data collected at 295 K are listed in Table 1. No anomaly is observed in the occupation parameters at any site, including full occupancy for the oxygen site as well. Thus, PHTO should have stoichiometric composition. Bond valence sum (BVS) calculations based on the refined structure from NPD data give valences of +2.26 for Pb ions, +1.99 for Hg ions and +3.88 for Ti ions (see Table 1), which are consistent with x-ray absorption spectroscopy (XAS) results discussed later. Hence the obtained sample is an A-site-ordered perovskite with stoichiometric $\text{PbHg}_3\text{Ti}_4\text{O}_{12}$ formula and space group Im-3 as shown in Fig. 1b. The structure model is consistent with the refinements of SXRD data as shown in Supplementary Fig. S1 and Table S1. Moreover, for Hg ions, if

![Fig. 1 Neutron diffraction and crystal structure of PHTO.](https://example.com/figure1.jpg)
perovskites (7.3–7.5 Å), while the $\angle$Ti–O–Ti angle (155.8°) of PHTO is less distorted from the ideal 180° than those for other AA$'$3B4O12 type perovskites ($\angle$B–O–B = 140°). The larger $\angle$Ti–O–Ti angle and correspondingly smaller $\angle$A$'$–O–Ti angle in PHTO imply the less tilting of TiO6 octahedron, which is attributed to the large ionic size of Hg2+$^{+}$ about 0.96 Å. Figure 2 shows the tolerance factor $t$ versus average ionic radius of A-site ions for simple perovskites ATiO3 and A-site-ordered perovskites AA$'$3Ti4O12 where the $t$ factor is calculated assuming a simple ABO3 form with the average ionic size at A site. It is found that from CaTiO3 to BaTiO3, the ionic radius of A-site increases from 1.34 Å to 1.61 Å while the tolerance factor $t$ increases from 0.97 to 1.07. The simple perovskites ATiO3 are located at the upper right panel of Fig. 2. For most of the AA$'$3Ti4O12 compounds, the average ionic radius of A-site is about 0.8 Å and the tolerance factor $t$ is around 0.77. Therefore, AA$'$3Ti4O12 perovskites are located at the lower left panel of the Fig. 2. However, different from these ATiO3 and AA$'$3Ti4O12 compounds, the average ionic radius of A-site ion for PHTO is large, about 1.1 Å and the tolerance factor $t$ is about 0.88. Moreover as aforementioned, the coordination number for Hg ions at the A-site in PHTO is eight, not square planar as in A$'$-TM quadruple perovskites AA$'$3Ti4O12, and it is located in the middle area of Fig. 2, between those of simple ATiO3 (coordination number 12) and AA$'$3Ti4O12 quadruple perovskites (coordination number 4). Thus, PHTO appears to be a link from simple ATiO3 perovskite to AA$'$3Ti4O12 type A-site-ordered perovskite (Fig. 2). The A-site coordination number for ABO3 perovskite is 12 while the A-site and A$'$-site coordination numbers in the A-site-ordered quadruple perovskite AA$'$3B4O12 compound are 12 and 4 respectively. However in PHTO, the coordination number of A$'$ (Hg2+$^{+}$) is 8, which is just in between that for simple ATiO3 and AA$'$3Ti4O12. This result provides further evidence supporting the scenario aforementioned. Considering that the atomic size of Sr2+$^{+}$ (1.45 Å) is similar to that of Pb2+$^{+}$ (1.49 Å), we also fabricated the A-site-ordered perovskite compound SrHg3Ti4O12, which is isostructural with PHTO, but without the observation of ferroelectricity. The result about SrHg3Ti4O12 will be published elsewhere. According to the previous report37, the low energy level of $d^{10}$ orbital prefers a relatively high coordination number owing to the weak d-s and d-p hybridization in perovskite compounds. As to PHTO system, the first principles calculation result indicates that the energy level

| Compounds         | $a$/Å | $r_a$/Å | $\Delta r_a$/Å | $\Delta A'$–O – Ti/deg | $\Delta Ti$–O – Ti/deg |
|-------------------|-------|---------|----------------|------------------------|------------------------|
| PbHg3Ti4O12       | 7.7234| 1.49    | 0.96           | 0.53                   | 102.60                  |
| CaPd3Ti4O12       | 7.4977| 1.34    | 0.64           | 0.70                   | 107.17                 |
| CaFe3Ti4O12       | 7.4672| 1.34    | 0.64           | 0.70                   | 107.82                 |
| CaCo3Ti4O12       | 7.3730| 1.34    | 0.57           | 0.77                   | 108.98                 |
| SrCu3Ti4O12       | 7.4275| 1.44    | 0.57           | 0.87                   | 109.18                 |

Table 2 Lattice constant $a$, ionic radius of A-site $r_a$, ionic radius of A-site $r_a$, difference between ionic radii of A- and A$'$-site $\Delta r_a$, metal–oxygen bond angles $\angle A'$–O–Ti and $\angle Ti$–O–Ti for some AA$'$3Ti4O12 type perovskite compounds. In PHTO, the $\Delta$Ti–O–Ti angle is large, about 155.8°. Therefore, PHTO is different from simple perovskites ATiO3 and A-site-ordered perovskites AA$'$3Ti4O12, where the $t$ factor is calculated assuming a simple ABO3 form with the average ionic size at A site. It is found that from CaTiO3 to BaTiO3, the ionic radius of A-site increases from 1.34 Å to 1.61 Å while the tolerance factor $t$ increases from 0.97 to 1.07. The simple perovskites ATiO3 are located at the upper right panel of Fig. 2. For most of the AA$'$3Ti4O12 compounds, the average ionic radius of A-site is about 0.8 Å and the tolerance factor $t$ is around 0.77. Therefore, AA$'$3Ti4O12 perovskites are located at the lower left panel of the Fig. 2. However, different from these ATiO3 and AA$'$3Ti4O12 compounds, the average ionic radius of A-site ion for PHTO is large, about 1.1 Å and the tolerance factor $t$ is about 0.88. Moreover as aforementioned, the coordination number for Hg ions at the A-site in PHTO is eight, not square planar as in A$'$-TM quadruple perovskites AA$'$3Ti4O12, and it is located in the middle area of Fig. 2, between those of simple ATiO3 (coordination number 12) and AA$'$3Ti4O12 quadruple perovskites (coordination number 4). Thus, PHTO appears to be a link from simple ATiO3 perovskite to AA$'$3Ti4O12 type A-site-ordered perovskite (Fig. 2). The A-site coordination number for ABO3 perovskite is 12 while the A-site and A$'$-site coordination numbers in the A-site-ordered quadruple perovskite AA$'$3B4O12 compound are 12 and 4 respectively. However in PHTO, the coordination number of A$'$ (Hg2+$^{+}$) is 8, which is just in between that for simple ATiO3 and AA$'$3Ti4O12. This result provides further evidence supporting the scenario aforementioned. Considering that the atomic size of Sr2+$^{+}$ (1.45 Å) is similar to that of Pb2+$^{+}$ (1.49 Å), we also fabricated the A-site-ordered perovskite compound SrHg3Ti4O12, which is isostructural with PHTO, but without the observation of ferroelectricity. The result about SrHg3Ti4O12 will be published elsewhere. According to the previous report37, the low energy level of $d^{10}$ orbital prefers a relatively high coordination number owing to the weak d-s and d-p hybridization in perovskite compounds. As to PHTO system, the first principles calculation result indicates that the energy level

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determine the valence state, local environment, and of Hg2+ shows a low total energy in the state of 8 coordination for 4 coordination (See Supplementary Fig. S2a). Moreover, Pb2 and PbNiO3 as Pb4+ reference. The weak pre-edge feature in PbNiO3 due to excitation from a 2p core electron to the 6s orbital. It has been established that this weak pre-edge feature is a sensitive fingerprint of the Pb valence state in solid state materials. The reason is that the Pb 5d orbitals are fully occupied, the valence state is then reflected by the 6s occupation. However, as shown in the inset, PHTO has the Pb-L3 XAS profile similar to the Pb2+ reference PbTiO3, detail, no 2p-6s related excitation is observed in PHTO, which indicates the valence state of Pb2+ with fully occupied 6s2 state in this compound. Figure 3c shows the Hg-L edge XAS spectra of PHTO as well as HgO. The same energy position and very similar spectral profile of both PHTO and HgO indicate the Hg2+ valence state. The XAS results show that the electronic configuration of PHTO is Pb2+Hg2+Ti4+O12, which is fully consistent with the BVS results based on the crystal structure.

### Magnetic and transport characteristics

The magnetic susceptibility $\chi$ is almost constant and negative near zero from 300 K to 30 K, while $\chi$ increases rapidly on further decrease of temperature as shown in Supplementary Fig. S3. These results revealed that PHTO is nonmagnetic due to the 3d$^9$, 5d$^{10}$ and 6s$^2$ electron configuration for Ti$^{4+}$, Hg$^{2+}$ and Pb$^{2+}$, respectively, and consistent with the BVS calculations and XAS results. The resistance of PHTO at room temperature is too large to be measured, indicating the high electrical insulating property and few vacancies in the sample. Supplementary Fig. S4a shows the Ultraviolet-Visible-Near Infrared (UV-Vis-NIR) absorption spectrum of PHTO. The curve exhibits a steep decrease from 420 to 640 nm, which is a typical optical response of semiconductors. The optical band gap of the PHTO was estimated to be 2.12 eV.

### Electric and ferroelectricity

The permittivity and dielectric loss measured with different frequencies from 2 K to 300 K are shown in Fig. 4a, b, respectively. The relative dielectric constant, $\varepsilon_r$ of PHTO is over 220 in the measured temperature range. The temperature dependence of $\varepsilon_r$ shows a clear peak around 250 K at all the frequencies of measurements. The relative temperature-dependent $\varepsilon_r$ strongly indicates a low-temperature ferroelectric phase transition to a high-temperature paraelectric phase at $T_{FE} \approx 250$ K. Furthermore, the transition temperature is independent of measurement frequency, indicating that PHTO is not a relaxor-type ferroelectric material. The relative increase of the dielectric loss around 250 K is consistent with the paraelectric-ferroelectric transition. Moreover the dielectric loss keeps at a very low value (<0.03) with all of frequency in the whole measurement range, which indicate the intrinsic dielectric property of PHTO sample is stable at ambient up to 973 K as revealed by thermogravimetry analysis measurements shown in Supplementary Fig. S5.

### Valence state analysis

Soft X-ray absorption spectroscopy at the L$_{2,3}$ edge of TM 3d ions is a highly sensitive method to determine the valence state, local environment, and orbital occupation of the ions. Figure 3a shows the Ti-L$_{2,3}$ XAS spectra of PHTO together with SrTiO$_3$ as a pure Ti$^{4+}$ reference. The very similar multiplet spectral features in PHTO and SrTiO$_3$ clearly correspond to the Ti 2p$^5$3d$^0$ $\rightarrow$ 2p$^5$3d$^1$ transition, demonstrating the Ti$^{4+}$ valence state in PHTO. Note that the PHTO spectrum is shifted by about 0.25 eV to lower energy with respect to the SrTiO$_3$ spectrum, reflecting the different Ti local environments. Actually the Ti-O distance is 1.974(7) Å in PHTO and 1.951(7) Å in SrTiO$_3$, which suggests a weaker crystal field effect in PHTO. Figure 3b shows the Pb-L$_3$ XAS spectra of PHTO together with PbTiO$_3$ as a Pb$^{2+}$ and PbNiO$_3$ as Pb$^{4+}$ reference. The optical band gap of the PHTO was estimated to be 2.12 eV.
PTHO. In order to further identify the ferroelectricity of PHTO, the isothermal polarizations were measured by the PUND method\textsuperscript{20}. No loops of polarization versus electric field (P–E) were observed in polycrystalline PHTO at room temperature. However, canonical P–E hysteresis loops were observed below 250 K as shown in Fig. 4c, d. It is clear that polarization is enhanced at low temperature. At fixed temperature, the P–E loops significantly expand with increasing electric field. For example, the P value increases from 0.028 to 0.11 µC cm\(^{-2}\) as the maximum electric field changes from 17 to 32 kV cm\(^{-1}\) at 10 K. As known, PbTiO\(_3\) is a classic and extensively studied ferroelectric material with Curie temperature about 763 K, which excludes the possibility that the ferroelectricity of PHTO originated from PbTiO\(_3\). Thus it can be concluded that the ferroelectricity is due to PHTO.

**Temperature dependent crystal structure transition.** To further understand the mechanism of ferroelectricity from the structure point of view, we performed the temperature dependent SXRD, as shown in Supplementary Fig. S6. The characteristic (400) diffraction peak at different temperatures is collected in Fig. 5a. It is obvious that the diffraction peaks split into two sub-peaks between 240 K and 260 K, which indicates the occurrence of temperature induced phase transition at 250 K is due to symmetry reduction from a cubic - to - orthorhombic phase. Checking the SXRD data measured at 300 K and 90 K, all the high symmetry diffraction peaks, such as (200), (220), (222), (400) and (402) of the high temperature phase, are split into two sub-peaks on cooling to low temperature (see Supplementary Fig. S7). There is a clear long-range structural transition with decreasing temperature. The SXRD data collected at 90 K is presented in Fig. 5b. The crystal structure of PHTO can be fitted very well by an orthorhombic phase of non-centrosymmetric space group Imm\(_2\) (No.44). The inset shows the crystal structure of the low temperature phase. The Rietveld refinement results are listed in Supplementary Table S2. Figure 5c shows variation of the lattice parameters based on refinements of SXRD data. The lattice parameter \(a\) decrease linearly from 300 K to 250 K. Below the temperature of the phase transition, the lattice parameter \(b\) and \(c\) continue to decrease while parameter \(a\) increases. Figure 5d shows the evolution of TiO\(_6\) octahedron before and after phase transition. In the high temperature phase, TiO\(_6\) forms a regular octahedron with \(\angle\text{Ti-O-Ti}\) angle of 155.8° and Ti-O bond length of 1.974 Å respectively. When this cubic phase transformed to an orthorhombic non-centrosymmetric phase below 250 K, both \(\angle\text{Ti-O-Ti}\) angles and Ti-O bond length divergences occurred in TiO\(_6\) octahedron. These distortions originate from the relative displacements of Ti and O around all the axis directions. The polarization in the \(ab\)-plane was canceled due to the random relative displacements of Ti and O, leaving the polarization vector only along the [001] direction. As to HgO\(_8\) polyhedron, the single type HgO\(_8\) polyhedron with only two-types of Hg-O bond length in the high temperature phase evolves into three different type of HgO\(_8\) polyhedron with different bond length in the low temperature phase, but without contribution to the polarization. Supplementary Fig. S8 illustrates the evolution of HgO\(_8\) during the paraelectric-ferroelectric transition. Therefore, PHTO is a displacive-type ferroelectric. The ionic spontaneous polarization can be calculated by multiplying the effective charges and the distance between the negative and positive valence weighted mean center along the c-axis then divided by the volume of the unit cell\textsuperscript{17}. Based on the refined results of NPD data collected at 5 K in Table 3, the calculated polarization value is about 13.65 µC cm\(^{-2}\). Although temperature-dependent phase transition is typical in A-site-ordered perovskites, it is usually isostructural or centrosymmetric - to - centrosymmetric phase transition.

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**Fig. 4 Dielectricity and ferroelectricity of PHTO.** Temperature dependence of a relative dielectric constant \(\varepsilon_r\) and b dielectric loss at selected frequencies. The P-E hysteresis loops measured at c 10 K and d 200 K under selected electric fields.
Fig. 5 Temperature induced structural transition in PHTO. a The characteristic diffraction peaks (400) collected at different temperatures. b Rietveld refinements based on SXRD data at 90 K. The arrow denotes the polarization direction. c Temperature dependence of the lattice constant and d schematic illustration for the changing of TiO$_6$ octahedron along different axis.

Table 3 Refined structure parameters of PHTO based on NPD data collected at 5 K.

| Crystallographic data for PbHg$_3$Ti$_4$O$_{12}$ based on NPD at 5 K |
|-------------------------------------------------|-----------------------------|-----------------------------|
| **Atom**                                       | **Wyck**    | **x** | **y** | **z** | **U$_{iso}$ (Å$^2$)** |
| Pb                                            | 2a           | 0.5   | 0.5   | 0.5   | 0.008(8)                |
| Hg1                                           | 2b           | 0.5   | 0.5   | 0.5   | 0.007(1)                |
| Hg2                                           | 2a           | 0.5   | 0.5   | 0.5   | 0.005(1)                |
| Hg3                                           | 2b           | 0.5   | 0.5   | 0.5   | 0.004(4)                |
| Ti                                            | 8e           | 0.2353(7) | 0.2396(3) | 0.2455(8) | 0.013(9)               |
| O1                                            | 4c           | 0.1966(0) | 0      | 0.2905(6) | 0.007(0)               |
| O2                                            | 8e           | 0.2786(9) | 0.2103(7) | 0.0006(9) | 0.013(4)               |
| O3                                            | 4d           | 0     | 0.2802(5) | 0.2527(9) | 0.027(0)               |
| O4                                            | 4c           | −0.2008(6) | 0      | −0.3022(4) | 0.025(1)               |
| O5                                            | 4d           | 0     | −0.2696(8) | −0.1963(7) | 0.022(2)               |

| **Bond length (Å)**                           | **Bond angle(°)** |
|------------------------------------------------|-------------------|
| Pb-O1: 2.708(6)                                | Hg1-O1: 2.853(4)   |
| Pb-O2: 2.701(6)                                | Hg1-O3: 2.580(1)   |
| Pb-O3: 2.907(2)                                | Hg1-O5: 2.331(4)   |
| Pb-O4: 2.801(7)                                | Hg2-O1: 2.220(1)   |
| Pb-O5: 2.569(6)                                | Hg2-O2: 2.814(7)   |
| Ti-O1: 1.993(9)                                | Hg2-O4: 2.179(1)   |
| Ti-O2: 2.004(2)                                | Hg3-O2: 2.360(5)   |
| Ti-O3: 1.981(7)                                | Hg3-O3: 2.547(8)   |
| Ti-O4: 1.967(7)                                | Hg3-O4: 2.179(1)   |
| Ti-O5: 1.967(7)                                | Hg3-O5: 2.360(5)   |

Space group: Immm (No. 44) a = 7.7568(9) Å, b = 7.7040(1) Å, c = 7.709(8) Å, α = β = γ = 90°; V = 460.26 Å$^3$; Z = 2; R$_p = 4.08%$, R$_wp = 3.39%$. 
from $Imm\text{'}-3$ to $Pn\text{'}-3$. Most of those phase transitions are mainly induced by charge transfer between the A$'$-site and B-site ions or charge disproportionation at B-site ions\textsuperscript{12,21,25}. PHTO is an example of AA$'$\textsubscript{3}B\textsubscript{4}O\textsubscript{12} type A-site-ordered perovskite that exhibits a centrosymmetric to non-centrosymmetric phase transition.

**First principles calculations.** In order to get a deeper insight into the ferroelectricity of PHTO at low temperature, first principles calculations have been performed. Supplementary Fig. S9 shows the phonon spectrum for the cubic structure of PHTO: three imaginary frequency modes with irreducible representation $T_d$ degenerate at $I$ point. By moving atoms along these soft phonon modes, the non-centrosymmetric crystal structure with space group $Imm\text{'}2$ can be obtained. Free energy per primitive cell versus moving amplitude of atoms along one of the soft modes is plotted in Fig. 6. The energy well suggests a spontaneous structure phase transition with the biggest contribution coming from the Ti-O mode. The imaginary modes at $H$ point only consist of the Ti-O phonon modes, which are consistent with the experimental conclusion. In addition, the $6s^2$ lone pair electrons of Pb$^{2+}$ also contribute to the site instability according to the theory analysis. However, the contribution from the lone pair mechanism does not play a dominant role from the energy perspective (see Fig. 6). Using Berry phase method, the polarization of ferroelectric structure and Born effective charge have been calculated. The calculated polarization value is about 16.87 $\mu$C cm$^{-2}$, which can also be obtained by multiplying the Born effective charge tensor with the corresponding atom displacements. This result is roughly in agreement with that of ionic spontaneous polarization calculation result of 13.65 $\mu$C cm$^{-2}$ based on neutron diffraction measurements. Both values are larger than the experimental result. This is because the calculated value is based on single crystal data, while the experimental measured value is from a polycrystalline sample of randomly distributed grains. For example, in ceramic samples of BiFeO$_3$, much lower values of about 8.9 $\mu$C cm$^{-2}$ of polarization have been measured at room temperature while in single crystals the polarization is up to 60 $\mu$C cm$^{-2}$ along [012] axis and 100 $\mu$C cm$^{-2}$ along [001] axis\textsuperscript{48,49}. Another reason for the low experimental value of polarization may be due to the grain boundary effect and/or leaking current issue preventing the application of maximum electric field in the experiment\textsuperscript{48,50}. Supplementary Fig. S10 shows the SEM image of the polycrystalline sample of PHTO. It can be seen that the grain size distributes around several microns to nearly 20 microns and generates a large amount of grain boundaries. Those grains are disorderly distributed, which will severely affect the polarization.

The A-site-ordered perovskite PbHg$_3$Ti$_4$O$_{12}$ was synthesized at high pressure and high temperature. PbHg$_3$Ti$_4$O$_{12}$ is a nonmagnetic insulator with a direct energy gap of $\sim$1.2 eV and relative dielectric constant larger than 220. PbHg$_3$Ti$_4$O$_{12}$ undergoes a transition from a high temperature centrosymmetric, cubic and paraelectric phase to a low temperature non-centrosymmetric, orthorhombic and ferroelectric phase with a record high Curie temperature at 250 K for an A-site-ordered quadruple perovskite. The ferroelectric distortion is dominated by Ti-O phonon mode anomaly. Because of the large average size of the A-site cation $\sim$1.1 Å in PHTO, the coordination of Hg$^{2+}$ is not square planar as in typical A-site-ordered quadruple perovskites, but closer to 8. Therefore PHTO can be considered to be structurally linking ABO$_3$ simple and AA$'$\textsubscript{3}B\textsubscript{4}O\textsubscript{12} quadruple perovskites. The established A-site-ordered perovskites AHg$_3$B$_4$O$_{12}$ may provide a potential pathway to find a class of ferroelectric materials of high Curie temperature. The work provides one of unique examples that high pressure synthesis can lead to the compounds with unique properties that are otherwise hard to get access at ambient. Moreover perovskites are the most abundant minerals in the broad lower Mantle of the Earth where high pressure is prevailing. Hence it might shed lights to further understand the coordination chemistry of perovskites like compounds at high pressures.

**Methods**

**Sample fabrication.** The polycrystalline sample of PHTO was synthesized by a solid state reaction under high pressure of 6 GPa and high temperature of 1273 K. The starting materials HgO (Aldrich, 99.0% pure), PbO (Alfa, 99.995% pure) and TiO$_2$ (Alfa, 99.995% pure) were mixed homogeneously in a molar ratio 1:3:4 and pressed into a pellet with diameter of 6 mm in an argon gas protected glove box with oxygen and H$_2$O level of less than 1 ppm. The pre-pressed pellet was sealed in a gold capsule. High pressure experiments were performed with the cubic anvil type high pressure apparatus\textsuperscript{31,52}. After the pressure was gradually increased to 6 GPa, the sample was heated to 1273 K and maintained for 30 min. Then the temperature was quenched to ambient before the release of pressure. In this process, yellow polycrystalline PHTO was obtained.

**Structure characterization.** The crystal structure was characterized by neutron powder diffraction (NPD) with the BT-1 high-resolution neutron powder diffractometer with a Ge (311) monochromator at the Center for Neutron Research (NCNR) of the National Institute of Standard & Technology (NIST). The neutron wavelength was 2.0774 Å. The intensities were measured with steps of 0.05° in the 2θ range of 10°–160°. The sample was also measured by synchrotron x-ray diffraction (SXRD) at the 11-BM-B at the Advanced Photon Source (APS) in Argonne National Laboratory. The X-ray wavelength was 0.412726 Å. Diffraction data were collected in the angle (2θ) range from 0.5° to 50° with steps of 0.002°. The obtained NPD and SXRD data were analyzed by the Rietveld method with the GSAS program\textsuperscript{53}.

**Valences determination.** The valence states of the Ti, Pb and Hg ions were determined by x-ray absorption spectroscopy (XAS). The soft XAS at the Ti-$L_3$ edge was measured with total electron yield mode at the Dragon beamline while the hard XAS at the Pb-$L_3$ and Hg-$L_3$ edge were measured with transmission geometry at the BL07A beamline at National Synchrotron Radiation Research Center (NSRRC) of Taiwan. SrTiO$_3$, PbTiO$_3$, PbNiO$_3$ and HgO were also measured at the respective Ti-$L_3$, Pb-$L_3$ and Hg-$L_3$ edges as reference materials.

**Permittivity and ferroelectricity characterization.** The permittivity was measured with different frequency by an Agilent-4980A LCR meter on a solid pellet with 4.0 mm in diameter and 230 μm in thickness. The ferroelectric hysteresis loops were measured at 20 Hz with a Radiant Precision Premier-II Ferroelectric Test System at different temperatures based on the proposed positive-up negative-down (PUND) method. Detailed experimental descriptions can be found in Ref. 20.

**Data availability**

The data that support the findings of this study are available from the corresponding authors upon reasonable request.
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Author contributions

C.Q.J. conceived and supervised the project. C.Q.J., R.Z.Y. and H.M.W. coordinated the research. J.F.Z., X.W., W.M.L., J.Z., Z.D., L.D., C.W.R., Y.W.L., J.L.Z., R.Z.Y. and X.C.W. performed the X-ray diffraction and crystal structure analysis. Q.Z.H. and F.J.Z. collected the powder neutron diffraction data and carried out the data analysis with the help of others. X.S. and R.C.Y. carried the TEM measurements. J.F.Z., Z.W.H., L.P.C., W.M.L., S.C.W., H.J.L., C.T.C. and L.H.T. carried out the XAS measurements. J.C.G., Y.T.Q., Z.L. and H.M.W. did the theoretical calculations. J.F.Z., R. Y., C.Q.J. and discussed with M.G. the experimental results in many details. J.F.Z., R.Z.Y. and C.Q.J. wrote the manuscript with comments from H.M.W., M.G. & Z.W.H. All authors contributed to the version.

Competing interests

The authors declare no competing interests.
Additional information

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