A large energy-gap oxide topological insulator based on the superconductor BaBiO$_3$

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Mixed-valent perovskite oxides based on BaBiO$_3$ [1, 2] (BBO) are, like cuprates, well-known high-$T_c$ superconductors. Recent ab initio calculations [10] have assigned the high-$T_c$ superconductivity to a correlation-enhanced electron–phonon coupling mechanism, stimulating the prediction and synthesis of new superconductor candidates among mixed-valent thallium perovskites [4–6]. Existing superconductivity has meant that research has mainly focused on hole-doped compounds, leaving electron-doped compounds relatively unexplored. Here we demonstrate through ab initio calculations that BBO emerges as a topological insulator [7–10] (TI) in the electron-doped region, where the spin-orbit coupling (SOC) effect is significant. BBO exhibits the largest topological energy gap of 0.7 eV among currently known TI materials [11], inside which Dirac-type topological surface states (TSSs) exit. As the first oxide TI, BBO is naturally stable against surface oxidization and degrading, different from chalcoginide TIs. An extra advantage of BBO lies in its ability to serve an interface between the TSSs and the superconductor for the realization of Majorana Fermions [12].

The parent compound BBO crystallizes in a monoclinic lattice that is distorted from the perovskite structure, and this distortion is attributed to the coexistence of two valence states, Bi$^{3+}$ (6s$^2$) and Bi$^{5+}$ (6s$^0$), due to charge disproportion of the formal Bi$^{4+}$. Octahedral BiO$_6$ breathes out and in for Bi$^{3+}$ and Bi$^{5+}$, respectively [9]. Under hole-doping conditions, such as in Ba$_{1-x}$K$_x$BiO$_3$ (x $\sim$ 0.4) [2, 14] and BaBi$_{1-x}$Pb$_x$O$_3$ ($x \sim$ 0.3) [1, 13], the breathing distortion is suppressed, resulting in a simple perovskite lattice [10] in which superconductivity emerges. The O-breathing phonon mode of BiO$_6$ is believed to result in the pairing of superconducting electrons in the Bardeen–Cooper–Schrieffer (BCS) framework [10]. In addition to the breathing distortion, undoped BBO also presents extra O-tilting distortions [9], finally resulting in a monoclinic phase. However, in ref. 10 and earlier work [11, 13, 19], SOC was not taken into account in the theoretical study, since the electronic states in the superconducting (hole-doped) region mainly result from Bi-6s and O-2p orbitals whose SOC effect is usually negligible.

By including the SOC effect in density-functional theory calculations of the BBO band structure, we discovered a band inversion between the first (Bi-6s state) and second (Bi-6p state) conduction bands, which is stable against lattice distortions. This inversion indicates that BBO is a three-dimensional (3D) TI with a large indirect energy gap of 0.7 eV when doped by electrons instead of holes. The band structure of ideal cubic BBO (Fig. 1A) reveals that the conduction bands are modified dramatically when SOC is included due to the presence of the Bi-6p states. The first conduction band crossing the Fermi energy ($E_F$) has a considerable Bi-6s contribution over the whole Brillouin zone, except at the $R$ momentum point where the Bi-6p contribution is dominant. Without SOC there is a zero energy gap at $R$ because of the degeneracy of the $p$ states. When the SOC is included, the $|p, j = 3/2 >$ and $|p, j = 1/2 >$ states split, which results in the large indirect energy gap of 0.7 eV in the vicinity of the $R$ point. Here, the Bi-6s state lies above the Bi-6p states, causing band inversion around the energy gap (Fig. 1A). Unlike bulk HgTe [20, 21], a well-known TI, this inversion occurs between the $|s, j = 1/2 >$ state and the $|p, j = 1/2 >$ state, rather than $|p, j = 3/2 >$ state. Since the Bi atom is the inversion centre of the perovskite lattice, the $|s, j = 1/2 >$ and $|p, j = 1/2 >$ states have “+” and “−” parities, respectively. Thus, a TI state can be obtained if $E_F$ is shifted up into this energy gap. The parities of all the valence bands below this gap were also calculated at all time-reversal invariant momenta, $\Gamma$ (0 0 0), $X$ (0.5 0 0), $M$ (0.5 0.5 0), and $R$ (0.5 0.5 0.5), which yielded $Z_2$ topological invariants (1;111), confirming the topological nontrivial feature according to Fu and Kane’s [22] parity criteria. At a doping rate of one electron per formula unit, $E_F$ shifts inside the $s$–$p$ inversion gap, and all the Bi$^{5+}$ ions become Bi$^{3+}$. Consequently, a cubic phase appears when the BiO$_6$ breathing distortion is suppressed, similar to the hole-doping case [10]. The new cubic lattice is found to expand only slightly in comparison to the undoped lattice because the Bi$^{3+}$–O bond is longer than the Bi$^{5+}$–O bond due to the localized Bi-6s orbital, while the band inversion is unaffected.

To illustrate the TSSs, we calculated the surface band structure using a slab model. As an example, we take the surface to be oriented along the (001) direction on which the bulk $R$ point is projected onto the $M$ point of the surface Brillouin zone. The slab is thirty BBO units thick with the outermost atomic layers being Ba-O. The TSSs,
shown in Figs. 1C and D, exhibit a simple Dirac-cone-like energy dispersion. The Dirac cone starts warping at higher energies due to the cubic symmetry of the lattice. The Fermi surface below the Dirac point is estimated to be approximately 0.75 eV μs, and inside the large bulk energy gap, the TSSs are well localized on the surface atomic layers to about two BBO units or around 1 nm in thickness. On the other hand, the Fermi energy near the Dirac point is estimated to be approximately 0.5 eV μs, which is consistent with the \textit{ab initio} calculations. We can confirm that the TI phase is stable against lattice distortions. The monoclinic BBO band structure (Fig. 1B) shows that the R point of the cubic lattice is projected onto the Γ point due to band folding, but the s–p band inversion at this Γ point is still present and the indirect gap is unchanged (0.7 eV). Since the inversion strength (the energy difference between the |s; j = 1/2 > and |p; j = 1/2 > bands) is nearly 2 eV (1.2 eV for the electron-doped structure), the TI phase is not destroyed by the O-breathing or -tilting distortions. Furthermore, band structure calculations using the hybrid functional method [5, 6], which is known to treat the dynamical correlation effect well for BBO [15, 27], also validated the inversion for both the ideal cubic and distorted structures. (Details are described in the Supplementary Information.)

Experimentally, electron-doped BBO may be achieved in BaBi(O_{0.67}F_{0.33})_3 by substituting O with F atoms. The O and F atom have comparable atomic radii and electronegativities, which can keep the octahedral BiO\_6 stable. For example, F substitution for O was applied for (1), iron-based superconductor LaOFeAs to realize electron-doping [29]. It is also possible to employ a state-of-art electrolyte gating technique to BBO to induce heavy electron-doping, which was realized for several mixed-valent compounds [30, 32]. In addition, oxygen vacancies, which are commonly observed in experiments [33], are also natural electron donors in BaBiO\_3–x. On the other hand, although the TSSs are unoccupied in pristine BBO compounds, it may be possible to monitor these states directly via monochromatic two-photon photoemission, as was recently employed to monitor the empty TSSs of Bi chalcogenides [34].
Thus far we can state that BBO becomes a superconductor with hole doping and a potential TI with electron doping. If pn-junction-type devices are fabricated with BBO, an interface between the TSSs and the superconductor may be realized, which is necessary for the realization of Fu and Kane’s [12] proposal on Majorana Fermions for quantum computation. Here, we outline a double-gated thin-film configuration, as illustrated in Fig. 2. If the bottom and top regions of the film are doped as p and n type, respectively, the double-gated structure may feasibly induce a hole-rich bottom surface and an electron-rich top surface, resulting in TSSs and superconductivity states on the top and bottom surfaces, respectively. In the middle region of the slab, the TSSs overlap with the bulk bands and penetrate the bulk. These TSSs can then become superconducting as a result of the proximity effect with the bottom superconducting regime. Such a structure is likely to be attainable as high-quality BBO thin films, which have been successfully grown on SrTiO$_3$ [35,37] and MgO [37] substrates. And moreover, the O-tilting lattice distortion was recently found to be suppressed in a BBO(001) thin film on MgO [37], which is very close to our required cubic structure.

The band structure of BaBiO$_3$ can act as a prototype for designing new perovskite TIs. Ba can be substituted by Sc, Y, or La to obtain new compounds as analogues of an electron-doped BBO. We found in calculations that a similar band inversion exists in this case. However, these compounds are semimetals (the Sc/Y/La-3d orbitals are lower in energy than the Bi-p states) and induce topological semimetals. In contrast, CsTiCl$_3$-type halide perovskites, which are predicted to be superconductor candidates [4,5], have band structures that are similar to BBO. However, we did not observe s-p inversion for ATiX$_3$ (X = Cs, Rb, F, Cl, Br, or I), because the SOC of Ti is not strong enough. When we can substitute Ti with Sn or Pb, we find that heavier members of this family, such as CsSnI$_3$ and CsPbI$_3$, are near the boundary of a topological trivial–nontrivial phase transition. Compressive pressure is necessary to drive these boundary materials into the TI region, which is consistent with recent theoretical calculations of these halides [38].

We thank Prof. X.-L. Qi at Stanford University and Prof. S. S. P. Parkin at IBM Almaden Research Center San Jose for fruitful discussions. B.Y. acknowledges financial support from the ERC Advanced Grant (291472) and computing time at HLRN Berlin/Hannover (Germany).

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Supplementary Information

Method

In band structure calculations, we employed \textit{ab initio} density-functional theory (DFT) with the generalized gradient approximation (GGA) [1]. We employed the \textit{Vienna ab initio simulation package} with a plane wave basis [2]. The core electrons were represented by the projector-augmented-wave potential [3]. For hybrid-functional calculations, we adopted HSE06 [4–6] exchange-correlation functional and interpolated the band structures using Wannier functions [7], where the DFT wavefunctions were projected to Bi-$sp$, Ba-$d$ and O-$p$ orbitals. The cubic crystal structure of BaBiO$_3$(BBO) was taken from ref. [8] with lattice constant $a = 4.35$ Å and the monoclinic structure was from ref. [9].

Band structures of BBO with hybrid-functionals

We validated the band structures of both cubic and monoclinic lattices using HSE06 functionals. For the cubic lattice, spin-orbit coupling (SOC) was also included. Compared to GGA, HSE06 was found to shift both the conduction and valence bands (see Figs. S1A and B), consist with ref. [10]. One can see clearly that the Bi-$5s$ state is still above the energy gap, perserving the $s-p$ band inversion. From Fig.1 of the paper, SOC can dramatically modify the Bi-6$p$ bands, while it does not change the Bi-$s$ band. Therefore, the $s-p$ inversion will remain when SOC exists, if they are already inverted without SOC. Withoung SOC a clear feature of $s-p$ inversion is the p-state degeneracy at the $R$ point right above the Fermi energy (Fig. 1A of the paper). In many previous literatures that did not employ SOC, actually, this feature can bee clearly observed [10][11]. So we performed HSE06 calculations without SOC for the monoclinic lattice, in order to reduce the computational time of large supercells. Our result agrees with recent HSE06 calculations [15]. In Fig. S1C, one can also see the $s-p$ inversion. In all, we conclude that the band inversion is robust with hybrid-functionals for both ideal cubic and distorted lattices.

Band structures of ABiO$_3$(A=Sc, Y and La)

We calculated GGA band structures of ABiO$_3$(A=Sc, Y and La). The $s-p$ band inversion exists for all three compounds, in which the Fermi energy shifts into the
inversion gap (See the band structures of ScBiO$_3$ and YBiO$_3$ for examples in Fig. S2). Therefore, they have the same nontrivial topological $Z_2$ invariants (1;1,1,1). However, the Sc/Y-d states exhibit lower energy than Bi-p states, resulting in a zero indirect gap. So these compounds are topological semimetals. In addition, we optimized the lattice constants of the perovskite lattice as 4.35, 4.41, and 4.46 Å for Sc, Y and La compounds, respectively, before calculating band structures.

FIG. 4: Band structures of bulk ScBiO$_3$ and YBiO$_3$ calculated with GGA. Red balls present bands with Bi-s states. Parities are labeled.