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# LETTER Open Access



# Realization of photonic *p*-orbital higher-order topological insulators

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# **Abstract**

The orbital degrees of freedom play a pivotal role in understanding fundamental phenomena in solid-state materials as well as exotic quantum states of matter including orbital superfluidity and topological semimetals. Despite tremendous efforts in engineering synthetic cold-atom, as well as electronic and photonic lattices to explore orbital physics, thus far high orbitals in an important class of materials, namely, higher-order topological insulators (HOTIs), have not been realized. Here, we demonstrate *p*-orbital corner states in a photonic HOTI, unveiling their underlying topological invariant, symmetry protection, and nonlinearity-induced dynamical rotation. In a Kagome-type HOTI, we find that the topological protection of *p*-orbital corner states demands an orbital-hopping symmetry in addition to generalized chiral symmetry. Due to orbital hybridization, nontrivial topology of the *p*-orbital HOTI is "hidden" if bulk polarization is used as the topological invariant, but well manifested by the generalized winding number. Our work opens a pathway for the exploration of intriguing orbital phenomena mediated by higher-band topology applicable to a broad spectrum of systems.

**Keywords** Higher-band topology, Orbital degrees of freedom, Generalized chiral symmetry, Bulk polarization, Winding number, Breathing Kagome lattice

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In condensed matter systems, an important characteristic of electrons besides charge and spin is the orbital degree of freedom (ODoF), which plays a crucial role in understanding unconventional properties in solid-state materials as well as in "orbital physics" toward unveiling the science and technology of correlated electrons [1]. However, due to the complexity and various degrees of freedom simultaneously involved in real materials, it has always been a challenge to fully unravel the physics of strongly correlated electronic matter mediated by the ODoF via controlled experiments. On the other hand, the interest in synthetic orbital systems including trapped atoms in optical lattices surged rapidly over the past decade [2, 3], leading to "orbital-only" quantum emulators not only for fermions but also for bosons. For example, orbital lattices have been employed to demonstrate complex Bose-Einstein condensates and orbital superfluidity [4] as well as exotic topological semimetal phases [5]. Clearly, the capability of purposely preparing atoms to higher orbital



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bands in optical lattices has opened the door for understanding orbital physics in condensed matter systems, with the ultimate goal of exploring new quantum states of matter not found in natural solids.

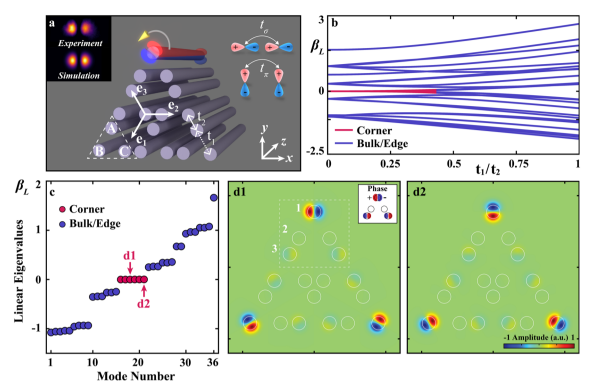
Apart from ultracold atoms in optical lattices, other synthetic platforms wherein the ODoF has emerged and caught increasing attention include semiconductor polaritonic lattices [6, 7], laser-written photonic lattices [8, 9], nanomechanical resonant structures [10], and atom-to-atom engineered electronic lattices [11, 12]. In particular, by employing the p-orbital bands of polariton micropillars arranged in a honeycomb lattice, both orbital edge states and exotic Dirac dispersions were observed [13, 14]. These artificial lattices can be used to flexibly manipulate orbital Dirac matter with transport properties and topological features difficult to achieve in conventional solid-state systems.

Using such synthetic lattice platforms, symmetry-protected topological (SPT) phases have been extensively explored due to their peculiar characteristics and unique potential applications associated with robust boundary states, especially in photonics [15–18]. Notable recent endeavors have focused on the realization of higher-order topological insulators (HOTIs) [19-25]—a new class of topological materials that do not obey the conventional bulk-edge correspondence principle [18, 26, 27]. While their underlying physics is still an ongoing subject of investigation, higher-order topological corner states have been extensively tested for various applications, including topological nanocavities and lasers [28-30]. However, to date, all experimental studies in HOTI systems relied on the s-orbital band, leaving higher-orbital HOTIs unexplored except for a recent proposal [31]. As a popular model of HOTIs, the breathing Kagome lattices (BKLs) exhibiting  $C_3$  rotational symmetry have been widely explored for demonstration of HOTIs in many different systems [32-41]. Although there is a continuing debate concerning the HOTI classification for the BKLs [42], experiments using metamaterials have unambiguously identified the higher-order topology for the s-orbital BKL-type topological crystalline insulators [39]. Furthermore, their robustness to symmetry-breaking perturbations has also been validated [41]. It is thus natural to ask: can we realize an HOTI manifesting higher orbitals using a synthetic BKL platform, and what are the distinctive topological features and new perspectives associated with the orbital HOTIs?

In this work, we experimentally demonstrate p-orbital HOTIs using photonic BKLs established by a continuous-wave (CW) laser-writing technique. In a triangle-shaped nontrivial BKL, both  $p_{x^-}$  and  $p_y$ -type orbital corner states are observed, with characteristic intensity and phase structures manifesting the zero-dimensional

"zero-energy" modes. Furthermore, corner excitation leads to dynamical rotation of a dipole beam due to the nonlinearity-induced lifting of  $p_x$  and  $p_y$  orbital mode degeneracy. We calculate the band structures and topological invariants of the orbital HOTIs and confirm their nontrivial topology from the winding number, albeit the topology is "hidden" in the conventional bulk polarization due to orbital coupling-induced band crossing. The SPT phase of orbital HOTIs is found to be inherited from the lower s-band Hamiltonian, being protected by the  $C_3$  rotational symmetry, the generalized chiral symmetry (GCS), as well as by a previously unknown orbital-hopping symmetry not applicable to the s-band HOTIs.

The photonic BKL platform used to demonstrate p -orbital HOTIs is illustrated in Fig. 1a, which has sixunit cells comprised of three sublattices (A, B, and C) with intracell and intercell hopping amplitudes  $t_1$  and  $t_2$ , respectively. When  $t_1 < t_2$ , the BKL exhibits a nontrivial topology, featuring SPT corner states as in previously studied s-orbital HOTIs [32-35]. In our laser-written BKL,  $t_1$  and  $t_2$  are tuned through the respective waveguide distances, while every waveguide supports two dipole modes (see left inset, Fig. 1a), allowing for both  $p_x$ and  $p_{\nu}$  orbitals. When the ODoF is taken into account, additional hopping amplitudes  $t_{\sigma}(t_{\pi})$  must be considered, being those responsible for the longitudinal (transverse) hopping when the p-orbitals are oriented parallel (perpendicular) to the bond direction [43] (see right inset, Fig. 1a). Under the tight-binding model, the full energy spectrum is calculated from the p-band Hamiltonian [31] (see "Methods") and plotted in Fig. 1b, where localized orbital corner states in the strongly topologically nontrivial regime are marked. Although the nontrivial phase occurs once  $t_1 < t_2$  in an infinite BKL, we found that localized corner states are present in a narrower window of parameters in an experimentally accessible BKL. Furthermore, the band structure for a representative nontrivial case is plotted in Fig. 1c, which displays six topological orbital corner modes in the bandgap with eigenvalues close to zero. Two typical corner modes exhibiting characteristic  $p_{x}$ - and  $p_{y}$ -type orbitals are shown in Fig. 1d1, d2: their amplitudes distribute equally among three corner sites, showing no leaking to the nearest-neighbor sites but only a weak amplitude in the next-nearest-neighbor (NNN) sites. If we consider, for example, the  $p_x$ -type orbital at the top vertex, the corner mode is exponentially localized at the A-sublattice sites with an opposite phase between the dipoles in the corner and the two NNN sites (Fig. 1d1)—a characteristic feature of the HOTI corner modes [34, 44]. Other orbital orientations with unequal amplitude distributions at three vertices of a finite-sized BKL are discussed in the Additional file 1. Here we point out the difference Zhang et al. eLight (2023) 3:5 Page 3 of 9



**Fig. 1** Illustration of "zero-energy" p-orbital topological corner states in a BKL. **a** Schematic diagram of the triangular BKL with six-unit cells, where the white dashed triangle marks one unit cell consisting of three sublattices (A, B, and C), and  $t_1$  and  $t_2$  indicate the intracell and intercell hopping amplitudes, respectively. Each lattice site corresponds to a laser-written waveguide, which supports p-orbital modes (see the left inset). Two types of orbital hopping amplitudes,  $t_{\sigma}$  and  $t_{\pi}$ , are illustrated in the right inset. In the top vertex of the lattice, we depict the rotation of an orbital corner state under nonlinear excitation. **b** Calculated eigenvalue spectrum  $\beta_L$  as a function of  $t_1/t_2$  at  $t_{\pi}/t_{\sigma} = -0.3$  for the finite-size BKL shown in (**a**), where distinct "zero-energy" corner states in the highly nontrivial regime are marked in red. **c** Calculated band structure for the BKL with  $t_1/t_2 = 0.1$ , showing six corner states (red dots) at  $\beta_L \approx 0$ , where two representatives ( $p_{x^-}$  and  $p_{y^-}$ type modes relative to the top corner) are plotted in (**d1**, **d2**). Note that a topological p-orbital corner mode, as the  $p_x$ -mode illustrated in (**d1**), is characteristically localized at the corner site 1, with a staggered phase distribution in the NNN site 3 but zero amplitude at the nearest-neighbor site 2

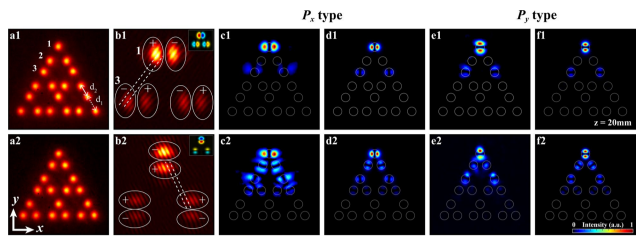
between the BKL employed in this work, where each constituting waveguide supports both fundamental and p-orbital modes, and the 2D Su–Schrieffer–Heeger (SSH) lattices used in previous work [44], where each waveguide supports a single mode and no p-orbitals were involved.

In our experiments, we write waveguides site-by-site in a nonlinear crystal [45], establishing both topologically nontrivial (Fig. 2a1) and trivial (Fig. 2a2) photonic BKLs. All waveguides remain intact during measurements, whereas probing of corner states is performed with an appropriately-shaped beam that can undergo linear or nonlinear propagation depending on whether a bias field is employed [44]. For a direct comparison of propagation through the 20-mm-long lattices, the amplitude and phase of the probe beam are pre-modulated to mimic the corner mode profiles in Fig. 1d (see Additional file 1). Measured results to illustrate orbital HOTI corner states are presented in Fig. 2b1-f2. Specifically, for the excitation of the  $p_x$ -type orbital corner states, the input probe beam is made of three x-oriented dipoles with a

nonuniform intensity distribution and staggered phase, i.e.,  $(0,\pi)$  for the top dipole vs.  $(\pi,0)$  for the other two (Fig. 2b1). The modulated probe beam is then launched into the BKLs at the top corner of the synthetic structure (the three uppermost A-sublattice sites). After linear propagation through the trivial and nontrivial BKLs, a significant difference is observed. In the nontrivial regime, the probe beam remains to be corner-localized in the A sublattice, with no intensity reaching the neighboring B and C sites (Fig. 2c1), thus confirming the formation of orbital HOTI corner states (see Fig. 1d1). On the contrary, the same probe beam becomes strongly distorted in the trivial regime, spreading into the edge and bulk sites with a tangible intensity distribution also in the B and C sites (Fig. 2c2). These results are corroborated by numerical simulations showing good agreement with the experiments (Fig. 2d1, d2).

In another set of measurements, the input probe beam is modulated so as to excite  $p_y$ -type orbital corner states, see Fig. 1d2. Similar results confirm the formation of a p

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**Fig. 2** Demonstration of p-orbital topological corner states in a photonic BKL. **a1**, **a2** Experimental BKLs in the nontrivial (**a1**) and trivial (**a2**) regimes. **b1**, **b2** Zoom-in interferograms of the probe beam with a reference plane wave, showing the out-of-phase relation between the corner site 1 and its NNN site 3 for both  $p_{X^-}$  (**b1**) and  $p_{Y^-}$  type (**b2**) orbital excitations. The white ellipses identify the three-dipole probe beams, where the plus/minus signs represent  $0/\pi$  phases for each dipole, and dashed straight lines are added to see the phase structure from the shifted (**b1**) and the aligned (**b2**) interference fringes, confirming the staggered phase structures (see also numerical results in right insets). **c1**, **c2** Observed output intensity patterns in nontrivial (**c1**) and trivial (**c2**) BKLs stemming from a  $p_{X^-}$  orbital corner excitation. **d1**, **d2** Numerical simulations corresponding to (**c1**, **c2**). Results from a  $p_{Y^-}$  orbital corner excitation are shown in (**e1**, **e2**) and (**f1**, **f2**). Grey circles in (**c1**–**f2**) mark the BKL sites. Lattice parameters are  $d_1 = 39 \, \mu \text{m}$  and  $d_2 = 31 \, \mu \text{m}$  for p-orbital excitation in the nontrivial BKL. Note the zero amplitude at the nearest-neighbor site 2 in (**c1**–**f1**)

-orbital mode in the orthogonal direction (see Fig. 2b2 and Fig. 2e1-f2). In addition, to validate the need for the staggered phase structure, a series of experiments and numerical simulations are performed by setting the input dipoles all in phase for direct comparison (see Additional file 1). Our results reveal that both  $p_x$ - and  $p_y$ -type orbital corner states can be realized under appropriate excitations.

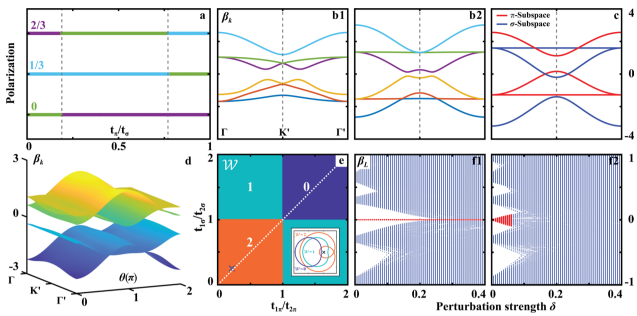
To demonstrate that the p-orbital BKL is indeed a symmetry-protected HOTI, we need to identify the topological invariant and prove the robustness of the corner states. The first candidate is the bulk polarization [46], or the  $Z_3$  Berry phase [31, 47] of the s-band BKL. In Fig. 3a, b1, b2, we plot the calculated polarization and 6-band structures as a function of the coupling ratio  $t_{\pi}/t_{\sigma}$ . Polarizations of the upper three bands are always 0, 1/3, and 2/3, however the order of these three values changes with  $t_{\pi}/t_{\sigma}$  as shown in Fig. 3a. The sum of the quantized polarizations of the upper three bands is always 1, yet this value must be regarded as mod 1, which implies that polarization vanishes. Although the vanishing of bulk polarization happens only in a parameter region where one needs to consider three bands instead of two or five [31], it suggests that polarization is not a good topological invariant at least for the *p*-orbital BKL.

To resolve the above issue, we introduce an auxiliary Hamiltonian  $H'(\mathbf{k}, \theta)$  obtained via a unitary transformation of the p-band Hamiltonian  $H(\mathbf{k})$  [31, 47], which employs a rotational operator  $R(\theta)$  (see "Methods" and

Additional file 1). The  $C_3$  symmetry and the GCS are both preserved as  $\theta$  is changed (see Additional file 1). At  $\theta = 4\pi/3$ , the auxiliary Hamiltonian is identical to the *p*-orbital BKL Hamiltonian, whereas at  $\theta = 0$  it is composed of two independent BKL Hamiltonians—one for the  $t_{\sigma}$  hopping and the other for the  $t_{\pi}$  hopping, just as if we had two decoupled s-band BKLs (Fig. 3c). For each of these Hamiltonian components, the topological invariant is given by the bulk polarization [46], which depends solely on  $t_1/t_2$ . In Fig. 3d we plot the band-gap structure of the auxiliary Hamiltonian as a function of  $\theta$ , showing that the gap does not close as  $\theta$  is varied from 0 to  $4\pi/3$ . Thus, the key point is that the orbital BKL described by the auxiliary Hamiltonian does not undergo a topological phase transition, so the p-orbital Hamiltonian at  $\theta = 4\pi/3$  is topologically non-trivial, just as for the s -band BKLs when  $t_1/t_2 < 1$  (see Additional file 1 for details).

Another interesting question that arises is whether there is a topological invariant that can manifest the nontrivial topology for the orbital HOTIs directly. To this end, we propose a generalized winding number  $\mathcal{W}$  as defined in "Methods" [14, 48]. Results are displayed in Fig. 3e:  $\mathcal{W}=2$  is found for the topologically nontrivial regime  $(t_1/t_2<1)$  and  $\mathcal{W}=0$  for the trivial regime  $t_1/t_2>1$ . We emphasize that  $\mathcal{W}$  is clearly a good topological invariant, independent of  $\theta$ . In fact, for BKL-like orbital HOTIs, the need of using the winding number  $\mathcal{W}$  is essential. This is because, for the Kagome geometry,

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**Fig. 3** Theoretical analysis of *p*-orbital HOTIs: topological invariant and corner-mode robustness. **a** Bulk polarization of the upper three bands (blue, green, and purple as plotted in (**b1**) and (**b2**) as a function of the orbital coupling ratio  $t_{\pi}/t_{\sigma}$ , exhibiting step jumps at the band-crossing points (marked by vertical dashed lines). The sum of the quantized polarizations of the upper three bands is (0 + 1/3 + 2/3)mod1 = 0. **b1-b2** Band structures plotted for (**b1**)  $t_{\pi}/t_{\sigma} = -0.19$  and (**b2**)  $t_{\pi}/t_{\sigma} = -0.77$  at  $t_1/t_2 = 0.6$  and  $\theta = 4\pi/3$ , showing the change in band-crossing for the upper three bands. **c** Band structure of the auxiliary Hamiltonian  $H'(\mathbf{k}, \theta)$  calculated for  $t_1/t_2 = 0.6$  and  $t_{\pi}/t_{\sigma} = -0.8$  at  $\theta = 0$ , which is equivalent to two sets of decoupled *s*-band BKLs: red for one set and dark blue for the other set. **d** 3D plot of the band structure of  $H'(\mathbf{k}, \theta)$  as a function of  $\theta$ , showing that the gap at zero energy remains open for any  $\theta$  (or for that matter, any orbital hybridization). **e** Calculated winding numbers, where the white dotted line marks the orbital-hopping-symmetry condition  $t_{1\sigma}/t_{2\sigma} = t_{1\pi}/t_{2\pi}$  required for topological protection of the orbital corner states. Here, circles in the lower inset illustrate distinct windings for each case, and the blue cross in the  $\mathcal{W} = 2$  region corresponds to the experimental parameters used to realize the *p*-orbital HOTI. **f1-f2** Robustness test of orbital corner states in a rhombic BKL obtained by applying random perturbations with increasing strengths  $\delta$  between site couplings that (**f1**) preserve the A-SubSy (i.e., without A-A coupling) and (**f2**) break the A-SubSy (i.e., with A-A coupling). The corner modes (red circles) remain at zero energy in (**f1**) but not in (**f2**). See Additional file 1 for details

one cannot physically decouple the system into two independent Hamiltonians—one for the  $t_{\sigma}$  and the other for the  $t_{\pi}$  hopping, akin to the chiral Hamiltonian for the quantum spin Hall effect [49, 50]. Orbital hybridization leads to band crossing and mixing from the  $\sigma$  and  $\pi$  subspaces as the parameter  $\theta$  is changed, so one cannot relate a single band to a particular subspace. The vanishing bulk polarization cannot unequivocally reveal the underlying nontrivial topology in the BKL. This is quite different from the 2D SSH-based HOTIs [19, 22-24, 44, 51], which can be decomposed into two independent Hamiltonians associated with orthogonal orbital hopping (see, e.g. Ref [51].). Given that BKL-based HOTIs are protected by the GCS, the use of the winding number can resolve this issue. If  $\theta$  changes progressively from 0 to  $4\pi/3$ , the band gap at zero-energy either remains open as shown in Fig. 3d or simply exhibits a "trivial" touching without band inversion—see Additional file 1. Consequently, the upper and lower three bands never topologically entangle. Since a continuous tuning of  $\theta$  does not induce a phase transition, the topological invariants are entirely inherited from the decoupled auxiliary Hamiltonian at  $\theta=0$ .

The case for W=1 in Fig. 3e merits further discussion. Since the BKLs possess a  $C_3$  symmetry, the winding numbers for each direction along the lattice unit vectors are identical (see "Methods"). W = 2 implies that both subspaces of the auxiliary Hamiltonian at  $\theta = 0$  are topologically nontrivial, thus there are two SPT states (corresponding to  $p_x$ - and  $p_y$ -orbitals) in each corner. In contrast, W = 1 indicates that only one subspace is topologically nontrivial while the other is indeed trivial. Intuitively, one would think that half of the corner states should persist as  $\theta$  is changed. However, as the trivial and nontrivial subspaces are coupled, the corner states of the whole system merge into the bulk. The absence of corner states for W = 1 is in agreement with the condition  $t_{1\sigma}/t_{2\sigma}=t_{1\pi}/t_{2\pi}$  that must be held in order to pin the orbital corner states at zero energy (see Additional file 1): the diagonal white line in Fig. 3e illustrating this condition does not cross the two W=1 quadrants. Thus, we only have topological corner states in the W = 2 region.

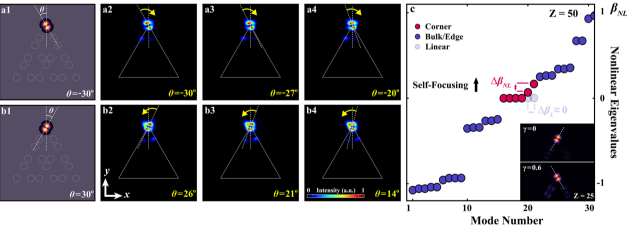
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To prove the immunity of an orbital corner state to perturbations, we investigate its robustness in a rhombic BKL, which hosts two orthogonal corner modes only at one corner, by following the approach recently developed for the study of sub-symmetry (SubSy)-protected topological states [41]. Here we assume that neglecting the long-range hopping is a reasonable approximation. Our results (Fig. 3f1, f2) show that orbital corner states residing in the A sublattice are protected under perturbations that respect the A-subsymmetry [41]. Moreover, we uncover that in addition to the  $C_3$  symmetry and the GCS, the topological protection of the p-orbital HOTI corner states requires a counterintuitive orbital-hopping symmetry (see "Methods" and Additional file 1). Such a symmetry demands  $t_{1\sigma}/t_{2\sigma}=t_{1\pi}/t_{2\pi}$  (marked by the white dashed line in Fig. 3e), which basically implies that the two orthogonal orbital hoppings should experience no difference when the lattice breathing characterized by  $t_1/t_2$  occurs. The lattice used in our experiments satisfies this condition with good approximation, as indicated by the blue cross in the region of W = 2 in Fig. 3e.

Finally, we observe an intriguing phenomenon—the dynamical rotation of orbital corner modes under the action of nonlinearity. As discussed in Additional file 1, the six p-orbital corner modes in a finite-size BKL are not all degenerate in their eigenvalues. Besides the two typical corner modes exhibiting characteristic— $p_x$  - and  $p_y$ -type orbitals shown in Fig. 1d1, d2, the other four orbital corner-mode eigenvalues are degenerate

in pairs. This leads to mode beating, and the amplitude distribution and orbital orientation in the three corners are susceptible to parameter variation such as in lattice size or potential. In our experiment, when a dipole-like beam is initially tilted away from the "equilibrium" position of the  $p_x$ - or  $p_y$ -orbital mode, excitation at one corner leads to a dipole rotation towards the preferred orbital orientation under a self-focusing nonlinearity. A typical example is shown in Fig. 4, where both clockwise (Fig. 4a1–a4) and counter-clockwise (Fig. 4b1–b4) rotations of about  $10^\circ$  are observed when the top corner is excited with a dipole beam tilted, respectively, at  $\theta=-30^\circ$  and  $\theta=30^\circ$  relative to the y-axis.

To better understand the underlying physical mechanism, we investigate numerically the observed phenomenon by use of the discrete NLSE for nonlinear beam propagation (see Additional file 1). We find that such rotations can be explained by the nonlinearityinduced lifting of the orbital mode degeneracy (Fig. 4c) since an oblique p-orbital can be regarded as the superposition of  $p_x$ - and  $p_y$ -type orbital components. Nonlinearity can introduce local variation of the lattice potential and break the lattice symmetry, and depending on the strength, it can also drive the corner modes away from zero-energy position and even form corner solitons [44]. Here we use a weak self-focusing nonlinearity as encountered in the experiment, and numerically show that the dynamical rotation can be explained by a dissimilar response of two orthogonal



**Fig. 4** Nonlinearity-induced dynamical rotation of *p*-orbital corner states. **a1**, **b1** Excitation of the top corner of a nontrivial BKL with a dipole-like beam initially tilted at (**a1**)  $\theta = -30^\circ$  and (**b1**)  $\theta = 30^\circ$  with respect to the *y*-axis. **a2**–**a4** Experimental output of the orbital corner state under an initial excitation corresponding to (**a1**), showing a clockwise rotation under the action of a self-focusing nonlinearity (with a bias field of 175 kV/m). **b2**–**b4** Opposite rotation observed under the same nonlinearity when the initial excitation corresponds to (**b1**). The grey triangle in each panel outlines the BKL boundaries. Note that no rotation occurs when the dipole is initially oriented along the *y*-axis. **c** Orbital band structure calculated from the discrete NLSE at a dimensionless distance Z = 50, for a probe beam incident at  $\theta = 30^\circ$  and propagating under both linear ( $\gamma = 0.6$ ) regimes. A tilted dipole-like beam excites two orthogonal orbital corner modes which are nearly degenerate in the linear regime ( $\Delta \beta_{L} \approx 0$ ). Here, the nonlinearity lifts the degeneracy and increases the difference  $\Delta \beta_{NL}$  between the nonlinear eigenvalues, in turn leading to a rotation of the orbital state. Insets illustrate the orientation of a probe beam at Z = 25 in the two regimes

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components to the action of nonlinearity, which lifts the degeneracy of the two orbital modes.

Before closing, we further emphasize the following differences between this work and that in Ref. [40], which also used a BKL-type lattice but focused on nonlinear s-orbital HOTI. (1) The ODoF requires each waveguide in the BKL can support at least the higher orbital p-mode, while the fs-laser-writing BKL waveguide as used for the s-orbital HOTI typically supports only a fundamental s-mode. (2) Different from the s-orbital, the coupling strength of the p-orbital is ori-

including hybrid quadrupole topological insulators [59] and orbital modes in non-Euclidean surfaces with disclinations [60]. Our results may prove relevant to those studies, as well as to the development of new photonic devices such as topological vortex waveguides and topological lasers.

#### 1 Methods

# 1.1 Orbital Hamiltonian and the winding number

Under the tight-binding approximation, the p-orbital corner states in a BKL can be found from the real-space Hamiltonian H expressed as [31]:

$$H = \sum_{m,n} t_{1\sigma} \left( a_{1m,n}^{\dagger} b_{1m,n} + b_{2m,n}^{\dagger} c_{2m,n} + c_{3m,n}^{\dagger} a_{3m,n} \right) + \sum_{m,n} t_{2\sigma} \left( a_{1m,n}^{\dagger} b_{1m+1,n} + b_{2m,n}^{\dagger} c_{2m-1,n+1} + c_{3m,n}^{\dagger} a_{3m,n+1} \right)$$

$$+ \sum_{m,n} t_{1\pi} \left( \widetilde{a}_{1m,n}^{\dagger} \widetilde{b}_{1m,n} + \widetilde{b}_{2m,n}^{\dagger} \widetilde{c}_{2m,n} + \widetilde{c}_{3m,n}^{\dagger} \widetilde{a}_{3m,n} \right) + \sum_{m,n} t_{2\pi} \left( \widetilde{a}_{1m,n}^{\dagger} \widetilde{b}_{1m+1,n} + \widetilde{b}_{2m,n}^{\dagger} \widetilde{c}_{2m-1,n+1} + \widetilde{c}_{3m,n}^{\dagger} \widetilde{a}_{3m,n+1} \right) + h.c.,$$

$$(1)$$

entation-dependent, so one needs two coupling  $t_{\pi}$  and  $t_{\sigma}$  to describe the orbital coupling dynamics. (3) For the s-orbital HOTI, the conventional bulk polarization can be used to characterize the topological properties of the corner states as in Ref. [40]. However, we have shown that the nontrivial topology of the p-orbital HOTI is "hidden" if the bulk polarization is used due to hybridization effects, but well manifested by the generalized winding number (see also discussion in Section IX of the Additional file 1). (4) We have found that the topological protection of p-orbital corner states demands an orbital-hopping symmetry, which has no counterpart whatsoever in the s-orbital HOTI. (5) Apart from different mechanisms for the employed nonlinearity itself, Ref [40] focused on nonlinear HOTI corner states and corner solitons, while the nonlinearity in our current work is used as "perturbation" to induce p-orbital corner mode rotation which is not applicable to the *s*-orbital HOTI.

We have thus shown that p-orbital HOTIs in photonic BKLs have a nontrivial band topology inherited from their s-orbital counterparts, which is characterized by the winding number rather than the quantized bulk polarization. Higher-order orbital corner states are topologically protected by the GCS along with an orbital-hopping symmetry not found in HOTIs where  $\sigma$ - and  $\pi$ -orbital couplings can be fully decoupled. These results have a direct impact on topological photonics involving the ODoF, especially given that different experimental platforms have enabled active and precise control of gain and loss, on-site energy, and coupling strength, in both real and synthetic dimensions [52–58]. For instance, using the ODoF, it is possible to construct a host of new topological phases

where  $a_{im,n}(\widetilde{a}_{im,n}), b_{im,n}(\widetilde{b}_{im,n})$  and  $c_{im,n}(\widetilde{c}_{im,n})$  (i=1,2,3) are the  $\sigma$ -type ( $\pi$ -type) annihilation operators at the A, B, and C sublattice sites in the (m,n) th unit cell, directed along the primitive lattice vectors  $\mathbf{e}_i$ . The coefficients  $t_{1\sigma}(t_{2\sigma})$  and  $t_{1\pi}(t_{2\pi})$  are the intracell (intercell) orbital coupling strengths (see Fig. 1a; Additional file 1). In Fig. 1, the linear band structure ( $\beta_L$ ) is obtained by diagonalizing H for different dimerization parameters  $t_1/t_2$  under an orbital hopping symmetry  $t_{1\sigma}/t_{2\sigma} = t_{1\pi}/t_{2\pi}$ . The p-orbital mode distribution is found by retrieving both  $p_x$ -and  $p_y$ -components from the calculated eigenvectors of H. In the momentum space, the Fourier transform of H for an orthogonal basis along x- and y-directions reads as

$$H(\mathbf{k}) = \begin{pmatrix} 0 & H_1^{\dagger} & H_3 \\ H_1 & 0 & H_2^{\dagger} \\ H_3^{\dagger} & H_2 & 0 \end{pmatrix}, \tag{2}$$

where the matrix entries  $H_i(i=1,2,3)$  are  $2\times 2$  matrices defined as  $H_i(\mathbf{k})=e_ie_i^\dagger(t_{1\sigma}+t_{2\sigma}e^{i\mathbf{k}\cdot e_i})+d_id_i^\dagger(t_{1\pi}+t_{2\pi}e^{i\mathbf{k}\cdot e_i})$ , with  $\mathbf{k}=(k_x,k_y)$  being the transverse wave vector, and  $\mathbf{d}_i$  the unit vector orthogonal to  $\mathbf{e}_i$ . The SPT phase of the p-band BKL model is identified from an auxiliary Hamiltonian  $H'(\mathbf{k},\theta)$ , which is achieved by applying a unitary transformation on  $H(\mathbf{k})$  (see Additional file 1 for details). The structure of  $H'(\mathbf{k},\theta)$  is identical to that in Eq. (2), with the matrix entries  $H_i(\mathbf{k})$  replaced by  $H_i'(\mathbf{k},\theta)=\begin{pmatrix} t_{1\sigma}+t_{2\sigma}e^{i\mathbf{k}\cdot e_i} & 0 \\ 0 & t_{1\pi}+t_{2\pi}e^{i\mathbf{k}\cdot e_i} \end{pmatrix} R(\theta)$ , where  $R(\theta)=\begin{pmatrix} \cos\theta-\sin\theta\\ \sin\theta&\cos\theta \end{pmatrix}$  is the rotation operator. The orbital Kagome Hamiltonian  $H(\mathbf{k})$  (Eq. 2) is fully equivalent to  $H'(\mathbf{k},\theta=4\pi/3)$ , i.e., at  $\theta=4\pi/3$ . In our analysis,  $R(\theta)$  is used to artificially tune the auxiliary Hamiltonian

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 $H'({\bf k},\theta)$  from the orbital-decoupled  $\theta=0$  to the physical  $\theta=4\pi/3$  Kagome model, to unequivocally identify the topological phase.

The winding number coefficients  $W_i$  are calculated from the matrix elements of  $H'(k,\theta)$  along each direction pointed by  $e_i$  in the first Brillouin zone through the following expression:

$$W_i = \frac{1}{2\pi} \int_0^{2\pi} dk_i \frac{d\Phi_i(k_i)}{dk_i}, \quad i = 1, 2, 3$$
 (3)

with  $\det[H_i'(\mathbf{k},\theta)] = |\det[H_i'(\mathbf{k},\theta)]|e^{i\Phi_i(\mathbf{k},\theta)}$ , where  $\Phi_i(\mathbf{k},\theta)$  is the argument of  $\det[H_i'(\mathbf{k},\theta)]$ , and  $k_i$  is the wave vector along the direction  $e_i$ . In general, for an arbitrary value of  $\theta$ , we have  $\mathcal{W} = \mathcal{W}_1 = \mathcal{W}_2 = \mathcal{W}_3$ , i.e., all  $\mathcal{W}_i$  are equal as long as the  $C_3$  symmetry is preserved in the system.

#### 1.2 Continuous model and numerical simulations

To provide a direct correspondence to the experimental results in our photonic platform based on a saturable photorefractive (PR) crystal (SBN:61), numerical simulations from the following continuous-model nonlinear Schrödinger-like equation (NLSE) are compared to the measurements [44]:

$$i\frac{\partial\Psi}{\partial z} = -\frac{1}{2k}\nabla_{\perp}^{2}\Psi - \frac{k\Delta n}{n_{0}}\frac{\Psi}{1 + I_{L}(x,y) + I_{P}(x,y)}.$$
 (4)

Here,  $\Psi(x,y,z)$  is the electric field envelope, where x and y are the transverse coordinates and z is the propagation distance.  $k=2\pi n_0/\lambda_0$  is the wavenumber in the medium, where  $n_0=2.35$  is the crystal refractive index and  $\lambda_0=532$  nm is the laser wavelength.  $\Delta n$  is the refractive index change determined by both the crystal electro-optic coefficient and the bias field.  $I_L(x,y)$  and  $I_P(x,y)$  are the intensity patterns of the lattice-writing and probing beams, respectively. The strength of the nonlinearity is controlled by the probe-beam intensity and the bias field [44]. Solutions to Eq. (4) for orbital corner states in both trivial and nontrivial BKL geometries are found via the split-step Fourier transform method.

# 2 Experimental method

Experimental BKLs in trivial and nontrivial geometries are established by a site-to-site writing process of each component waveguide using an ordinarily-polarized CW green laser beam [45]. To attain a variable planar distribution, a spatial light modulator (SLM) is employed to modulate the initial phase of the laser beam in the Fourier space. The effective PR "memory effect" guarantees that all waveguides remain intact during the measurement time window. The excitation of the orbital corner states

is carried out with a low-power extraordinarily-polarized probe beam, which undergoes linear propagation through the lattices for the results presented in Fig. 2. The probe beam is appropriately shaped via amplitude and phase modulation in order to excite the p-orbital corner modes (see Additional file 1). In the nonlinear regime, a tilted dipole-like probe beam is launched at the top corner of the nontrivial BKL, where the strength of the self-focusing nonlinearity can be controlled by the probe-beam intensity and the bias field.

# **Supplementary Information**

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**Additional file 1.** Supplementary Information including experimental details, calculation of topological invariants, and perturbation analysis.

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#### Author contributions

All authors contributed to different aspects of this work.

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#### Availability of data and materials

Data supporting this manuscript are available upon request.

## **Declarations**

# Ethics approval and consent to participate

Not applicable.

# Consent for publication

Not applicable.

#### **Competing interests**

There are no competing interests.

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