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Photo-induced phase transitions in ferroelectrics

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Ferroic materials naturally exhibit a rich number of functionalities, which often arise from thermally, chemically or mechanically induced symmetry breakings or phase transitions. Based on Density Functional Calculations, we demonstrate here that light can drive phase transitions as well in ferroelectric materials such as the perovskite oxides lead titanate and barium titanate. Phonon analysis and total energy calculations reveal that the polarization tends to vanish under illumination, to favor the emergence of non-polar phases, potentially antiferroelectric, and exhibiting a tilt of the oxygen octahedra. Strategies to tailor photo-induced phases based on phonon instabilities in the electronic ground state are also discussed.

vert light into electric [5], mechanical [6–8] or cheminon-destructive reading [11]. Numerous works focused on the photo-induced deformation of the unit cell without apparent symmetry breaking [8]. However, a recent experimental work hinted at possible structural phase transitions occurring in ferroelectric BaTiO₃ nanowires under illumination [12]. Using first-principles calculations, we show that photo-induced phase transitions are indeed possible in ferroelectrics such as barium and lead titanates. In both cases, transitions to non-polar phases are predicted. Furthermore, material engineering strategies based on competing lattice instabilities are suggested to tailor the critical concentration of photo-induced carriers at which the transition occurs, thus opening the way to a rational design of materials exhibiting specific properties under illumination.

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ABO₃ perovskite oxides are versatile materials exhibiting a wide range of properties (magnetism, polar distortions, etc.). Their high symmetry reference structure (see Fig. 1) is cubic with the B cation enclosed in an octahedral cage of oxygens at the center, and the A cation at the ³⁹ edges of the cube. Simple atomic displacement patterns e.g. as the octahedra tilts or electric dipoles sketched 41 in Fig. 1 – make it possible to obtain various symme-42 try breakings and control the materials properties. For 43 convenience, here we denote the most relevant of such 44 distortions using a generalized and simplified Glazer no-45 tation [13] $\alpha \beta \gamma / u^{\alpha'} v^{\beta'} w^{\gamma'}$, which is introduced in the 80 tween two Self-Consistent Field (SCF) steps was smaller 46 caption of Fig. 1.

Ferroelectric materials possess a spontaneous electric 47 Among perovskite oxides, barium titanate (BTO) and polarization that is switchable by electric fields. Active 48 lead titanate (PTO) are prototypical ferroelectric materesearch in bandgap engineering [1] and unconventional 49 rials. They exhibit a number of phases [14, 15] associated photovoltaic mechanisms [2–4] have prompted perovskite 50 with lattice instabilities of the reference cubic phase [16]. oxide ferroelectrics as potential light absorbers to con- 51 The transitions between phases are commonly driven by 52 temperature [14, 15], pressure [17–19] or static electric cal energy [9], or for optical memory writing [10] and 53 fields [20]. Here, we demonstrate that light can also act 54 as a knob to control the crystal symmetry. Optical ma-55 nipulation of ferroic materials has mainly focused on the 56 transient manipulation of the ferroic order parameter, 57 relying on non-thermalized photoexcited carriers [21] or 58 strong anharmonic phonon couplings [22]. In contrast, we 59 presently describe the emergence of new phases caused by 60 modified lattice instabilities in the presence of thermal-61 ized photo-excited carriers, i.e. the electrons lying at the 62 bottom of the conduction band (CB) and the equal num-63 ber of holes sitting at the top of the valence band (VB) 64 of a semiconductor during illumination. To this end, we 65 used ab-initio calculations, detailed in the next section, 66 before examining the cases of barium and lead titanate 67 respectively. Finally, we discuss the origin of this phe-68 nomenon and test design strategies for the engineering of 69 photo-induced phase transitions in ferroelectrics.

> We performed Density Functional Theory (DFT) cal-71 culations using the Abinit package software [23], with 72 the Projector Augmented Wave (PAW) method [24, 25]. ⁷³ We employ a $2 \times 2 \times 2$ supercell (with respect to the 5-74 atom pseudo-cubic perovskite cell) containing 40 atoms. ⁷⁵ The k-point mesh used was a Γ -centered $8 \times 8 \times 8$ grid; 76 the plane-wave cut-off was 35 Ha. A Fermi-Dirac distri-₇₇ bution with 0.1 eV smearing was used to populate the 78 electronic states. The electronic density was considered 79 converged when the difference in the calculated forces be- $_{81}$ than 10^{-8} Ha.Bohr⁻¹. The structural relaxation was

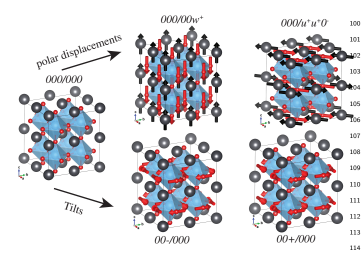


FIG. 1. Sketch of a $2 \times 2 \times 2$ supercell in the paraelectric cubic reference structure 000/000. A and B cations are in black and blue, oxygens in red. The symmetry of a phase is denoted by (i) its oxygen octahedra tilt pattern $\alpha\beta\gamma$, where α , β and γ are either 0 (no tilt), "+" (in-phase tilt) or "-" (antiphase tilt) and represent whether the oxygen octahedra tilts around the first, second and third pseudocubic axis, respectively. Similarly, we denote the electric dipole pattern using a Glazer-like notation $u^{\alpha'}v^{\beta'}w^{\gamma'}$, in which u, v, w represent the magnitude of the dipole inside the perovskite 5-atom cell, and α', β' and γ' describes whether dipole vectors [uvw] are aligned parallel ("+") or antiparallel ("-") when moving by one perovskite 5-atom cell along the first, second and third pseudocubic direction. Typical polar $(000/00w^+)$, antipolar $(000/u^+u^+0^-)$, in-phase $(00c^+/000)$ and antiphase tilt $(00c^-/000)$ displacement patterns are depicted using arrows.

82 stopped when all the forces on the atoms were smaller 83 than 5×10^{-7} Ha.Bohr⁻¹ and all components of the 84 stress tensor smaller than 0.1 MPa. BTO was described using the PBESol [26] exchange-correlation functional, while the Local Density Approximation [27, 28] was employed to treat PTO. Photo-excited thermalized carriers 88 are mimicked using Fermi-Dirac distribution with two 89 quasi-Fermi levels μ_e and μ_h , as commonly used to de-90 scribe photovoltaic effects in, for instance, p-n junctions [29]. As a result, during our DFT calculations, the 92 density is self-consistently converged under the constraint 93 of having $n_e = n_{ph}$ (and $n_h = n_{ph}$) electrons (holes) in 94 the CB (VB). In other words, at each Self-Consistent 95 Field (SCF) iteration, the following system of equations

$$n_h = \sum_{i \le N_v} \sum_{\mathbf{k}, \sigma} w_{\mathbf{k}} [1 - f(\varepsilon_{i\mathbf{k}, \sigma}, \mu_h)], \tag{1}$$

$$n_e = \sum_{i>N_e} \sum_{\mathbf{k},\sigma} w_{\mathbf{k}} f(\varepsilon_{i\mathbf{k},\sigma}, \mu_e), \qquad (2$$

₉₇ In the above equation, w_k is the weight of k-point k, ₁₅₃ tion [32]. Besides, the flattening of the energy landscape ₉₈ $f(\varepsilon_{ik},\mu)$ is the occupation number (Fermi-Dirac distri- ₁₅₄ around 0.03 e/f.u. ($\approx 5 \times 10^{20}$ e.cm⁻³) may be accompa-

 N_v is the index of the highest valence band. We assume 101 that there is no closing of the gap during our constrained DFT calculations, which is in practice observed for the concentrations of photo-excited carrier pairs n_{ph} considered in this work.

Note that all structures considered in this work were geometrically relaxed under the photo-excitation con-107 straint mentioned above. The criteria for convergence 108 of the structural degrees of freedom are set to be the same as in electronic ground state calculations.

Phonon calculations using the Phonopy package [30] were performed in the dark and under excitation on the 112 cubic perovskite oxides using a $2 \times 2 \times 2$ supercell after 113 the system was structurally relaxed. No analytic cor-114 rection, and thus no LO-TO splitting was considered, as 115 the required Born effective charges cannot be calculated 116 in photo-excited cases.

We first calculated the phonon band structure of cu-118 bic BTO in Figs. (2.a-d) for different concentrations of photoexcited carriers n_{ph} . In the dark $(n_{ph} = 0 \text{ e/f.u.})$ 120 Fig. (2.a)), the phonon dispersion exhibits imaginary 121 (negative in the figure) frequency modes characteris-122 tic of lattice instabilities. The main instability, at Γ , 123 is responsible for the emergence of the polar order in 124 BTO [16], which goes from a high temperature cubic $_{125}$ 000/000 structure to a ferroelectric tetragonal $000/00w^+$ phase below ~ 393 K, then to an orthorhombic structure $_{127}$ 000/0 v^+v^+ below \sim 280-270 K and finally to a rhombo- $_{128}$ hedral $000/u^{+}u^{+}u^{+}$ phase below ~ 183 K [14]. Upon 129 increasing the number of photo-excited carrier pairs to $n_{ph} = 0.05$ e/f.u. (in Fig. (2.b)), the frequencies of un-131 stable modes along the $X-M-\Gamma$ path get closer to real (positive) values, the ferroelectric soft optical mode at Γ becoming less unstable. Furthermore, for $n_{ph} \geq 0.1 \text{ e/f.u.}$ 134 (Figs. (2.c-d)) the cubic phase no longer exhibits unstable 135 phonon modes and is dynamically stable!

We calculated the energy (see Fig. (3.a)) of the four phases naturally occurring in bulk BTO, 000/000 (green 138 squares), $000/00w^{+}$ (blue diamonds), $000/0v^{+}v^{+}$ (vio-139 let triangles) and $000/u^+u^+u^+$ (ground state at 0 K in 140 dark conditions, black dashed baseline). The rhombohedral phase has the lowest energy for low concentrations of $_{142}$ photo-excited carriers (up to ≈ 0.0375 e/f.u.), while the tetragonal phase is slightly more stable at intermediate 144 concentrations $(n_{ph} \approx 0.0375 - 0.1 \text{ e/f.u.})$. When the cubic phase is dynamically stable $(n_{ph} > 0.1 \text{ e/f.u.})$, we find 146 accordingly that 000/000 is the lowest-energy structure, (1) 147 which is confirmed by the disappearance of polar atomic 148 displacements and the cell shape distortions of all con-149 sidered phases (see Supplementary Material [31]). The 150 predicted transition towards the 000/000 phase is con-151 sistent with the experimental observation, in BTO, of a is solved for μ_e and μ_h using a bisection algorithm. 152 downward shift of the Curie temperature under illumina-₉₉ bution) of state with eigenvalue ε_{ik} ; σ denotes the spin. ₁₅₅ nied by a monoclinic phase transition (see Supplemental

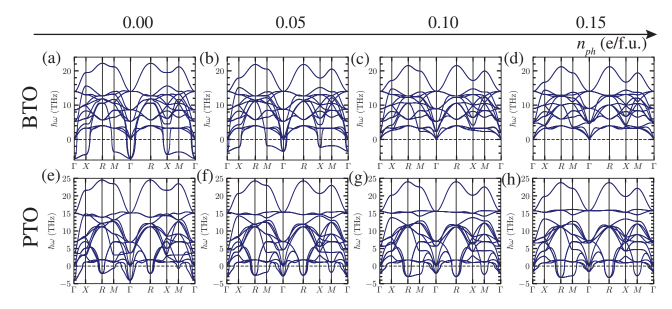


FIG. 2. (a-d) Phonon band structure in paraelectric cubic BTO and (e-h) PTO for different concentrations of photoexcited carrier pairs $n_{ph} = 0.00 \text{ e/f.u.}$ (a & e), $n_{ph} = 0.05 \text{ e/f.u.}$ (b & f), $n_{ph} = 0.10 \text{ e/f.u.}$ (c & g) and $n_{ph} = 0.15 \text{ e/f.u.}$ (d & h). The polar Γ dominant instability in BTO becomes stabilized (i.e. positive and real frequency) for $n_{ph} \geq 0.10 \text{ e/f.u.}$, as shown in (c) and (d). In PTO, the polar instability at Γ is replaced by a non-polar tilted instability at M and R.

in BTO nanowires under illumination [12, 33].

PTO exhibits a single transition from the non-polar cubic 000/000 phase to the ferroelectric tetragonal $000/00w^+$ structure below 765 K [15]. The phonon dispersion of the cubic phase for $n_{ph} = 0$ e/f.u. (Fig. (2.e)) exhibits one main polar instability at Γ , but also unstable modes at the R and M points associated with octahedra tilting 00- and 00+, respectively. The strength of the polar instability at Γ decreases with increasing the number of photo-excited carriers (see Figs. (2.e-h)) as in BTO. roelectricity in PTO is more robust against photocarrier generation. In contrast, the strengths of the tilting instabilities at R and M are slightly enhanced, and they become dominant for $n_{ph} \geq 0.1$ (Figs. (2.g-h)), suggestwith oxygen tilts under illumination.

Hence, we compute the energies of several prototypi-175 cal phases upon illumination, as shown in Fig. (3.b). The ferroelectric $000/00w^+$ phase, which is the ground state in dark conditions, is the black dashed baseline, while the cubic phase is plotted as green squares. We also plot a rhombohedral polar $---/u^+u^+u^+$ structure (navy circles) and its non-polar counterpart, a - - /000 (navy diamonds). We consider as well an orthorhombic phase 213 ₁₈₄ placement pattern [34]. A $00c^-/000$ non-polar tetrago-₂₁₅ outside the reach of experiments, and certainly not de-185 nal phase is also calculated (gold triangles). According to 216 sirable for applications. We can try to improve this

Material [31]). We note that such a critical concentration 187 stable up to $n_{ph} \approx 0.125$ e/f.u., the non-polar phase with would match recent observations of a monoclinic phase 188 tilts becoming dominant beyond that point. Overall, the antipolar $-0/u^-\bar{u}^-0^-$ phase is slightly more favor-₁₉₀ able (by 0.2-0.3 meV/f.u.) than the --/000 phase for $n_{ph} \geq 0.125$ e/f.u. Because of this small energy differ-192 ence, it is impossible to predict with certainty the sym-193 metry of PTO at large concentrations of photo-excited 194 carriers, and we can only reckon that this should be a 195 phase with oxygen tilts.

Unlike BTO, the polar atomic distortion in the $_{197} 000/00w^{+}$ phase remains rather large under photo-198 excitation (see Supplementary Material [31]). Moreover, However, it never totally disappears, indicating that fer- 199 both the antipolar displacement (for the $-0/u^-\bar{u}^-0^-$ 200 phase) and the tilt angle (for --/000 and -- $201 \ 0/u^-\bar{u}^-0^-$) increase with n_{ph} . In contrast, the polar dis-202 placements of the $---/u^+u^+u^+$ phase disappear with 203 increasing n_{ph} , and it thus collapses into the --/000ing a transition from a polar to a non-polar structure 204 phase, confirming that tilts and polarization are com-205 peting order parameters in PTO [35, 36], illumination 206 tipping the balance towards the former.

> Furthermore, calculations of minimum energy tran-208 sition paths between phases (see Supplementary Mate-209 rial [31]) show that $-0/u^{-}\bar{u}^{-}0^{-}$ and --/000 are 210 unstable in the dark; however, the $000/00w^+$ phase re-211 mains metastable under illumination, drawing the pic-212 ture of a first-order light-induced transition in PTO.

The large concentration of 0.125 e/f.u. needed to (violet triangles) with the $-0/u^-\bar{u}^-0^-$ antipolar dis- 214 destabilize the $000/00w^+$ phase in PTO is potentially Fig. (3.b), the ferroelectric $000/00w^+$ phase is the most 217 situation by mixing PTO with an antiferrodistortive

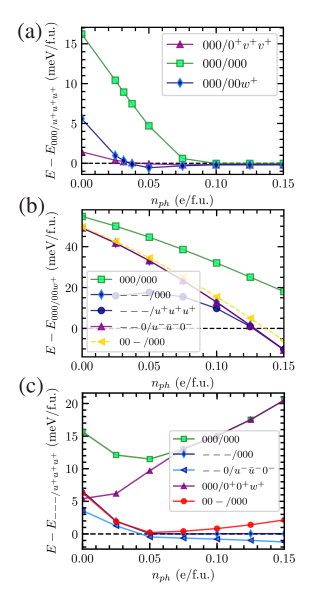


FIG. 3. (a) Energy of different phases in BTO with respect then the cubic 000 phase become more energetically favorable. (b) Energy of different phases in PTO with respect to the $000/00w^+$ structure (dashed dark line), which is the ground state in the dark, showing that non-polar antiferrodistortive phases become more stable at large concentration of photo-excited carriers. (c) Energy of different phases in PSTO 268 increasing concentration of excited carriers in both BTO with respect to the rhombohedral $--/u^+u^+u^+$ structure 269 and PTO, except that of the cubic 000/000 phase, which (dashed dark line), which is the lowest energy phase consid- 270 increases instead (see Supplementary Material [31]). Inered when no carriers are excited.

218 material such as SrTiO₃, and form the solid solution 274 drostatic compression; yet, the volume increase of the ²¹⁹ (Pb_{1/2}Sr_{1/2})TiO₃ (PSTO) to favor the tilt instability ²⁷⁵ cubic phase under illumination should in principle favor 221 several phases of PSTO with rocksalt order were cal- 277 behavior of the polar distortion upon illumination cannot 222 culated in Fig. (3.c). In particular, a polar rhombohe- 278 be understood solely in terms of a photo-induced volume $_{223}$ dral $--/u^+u^+u^+$ phase (black dashed line) is the $_{279}$ change [43], and that screening effects play a central role. $_{224}$ ground state in the dark. The $000/00w^+$ (purple tri- $_{280}$ There are, however, potential limitations of this work,

225 angles) and the high-symmetry cubic 000/000 phase (in green squares) have increasing energy with n_{ph} . On the contrary, the antipolar tilted phase $-0/u^-\bar{u}^-0^-$ (light blue triangles) becomes the most stable phase above $n_{ph} = 0.05$ e/f.u., thus supporting the proposed design strategy to reduce the critical concentration for the polarto-antipolar/antiferrodistortive transition.

From this work, it also becomes apparent that the ferroelectric instability can be hampered by the photogeneration of free carriers. Obviously, free carriers screen the long-range Coulombic interaction responsible for the polar order [37, 41], thus seemingly disfavoring polar phases with respect to non-polar ones. This is consistent with calculations [37, 41] and experiments [42] performed by doping BTO with free electrons. The former predicted the disappearance of the polar order in BTO at a concentration of extra-electrons of ≈ 0.1 e/f.u., which is exactly the concentration of photo-excited carriers that stabilizes the cubic phase in our calculations. We also note that the destruction of the polar order in BTO and the large stability (and metastability) window of the $000/00w^+$ phase in PTO are consistent with the doping calculations performed in Ref. [37]. In addition, photoexcitation tends to move electrons from O 2p states to Ti d states, thus destroying the covalent bonding that drives the ferroelectric order [40] and favoring non-polar orders. Looking at the 251 evolution of interatomic force constants (IFCs; see Sup-252 plementary Material [31]) under photoexcitation, shows that (i) almost all IFCs are reduced in magnitude under photoexcitation, and (ii) the most affected IFCs corresponds to 4th nearest neighbors interaction, i.e. similar atoms (Ti-Ti, O-O, etc.) interacting along the < 100 > cubic directions. This parallels the further stabilization of the paraelectric phase under photoexcitation in the 259 incipient ferroelectric PbTe, which results from screen-260 ing of the long-range Coulomb interactions by the redistributed photoexcited carriers along the < 100 > cubic to the $000/u^+u^+u^+$ structure (dashed dark line), which is 262 directions [38]. From the evidence presented here, as well the ground state in the dark. The tetragonal $000/00w^+$ and $_{263}$ as in other works involving doping and photoexcited cal-²⁶⁴ culations [37, 38], electrostatic screening by photoexcited 265 carriers appears as a universal mechanism for "dipoledipole driven" ferroelectric materials such as BTO [39].

In addition, the volume of most phases decreases with 271 terestingly, the decrease of volume of the polar phases 272 is consistent with the disappearance of the polar order, 273 and is reminiscent of the well-known [17–19] effect of hywith respect to the polar one. As a proof of concept, 276 the polar instabilities. Hence, our results suggest that the 281 since DFT is known to underestimate the quasi-particle 337 16-1-0227. 286 age - using DFT input from Quantum Espresso [46, 47] 342 DoD challenge grant for use of computing resources. We lence bands (see Supplementary Material [31]). The corrected band structure is overall not altered, except for an almost rigid upshift of the conduction bands. This shift slightly depends on the studied phase (with differences smaller than 100 meV between phases). The shift is smaller for non-polar phases, therefore facilitating further the light-induced transitions towards such latter states.

For BTO, we checked the robustness of our conclusions by performing calculations of zone center phonons 349 of the cubic phase with hybrid DFT. We compare calculations performed with the semi-local PBEsol functional and the hybrid HSEsol [48] functional. Both functionals 300 give similar lattice constants and similar frequency for 301 the imaginary frequency of the soft ferroelectric mode, even though the band gaps are quite different (1.8 eV for PBEsol and 3.4 eV, close to the GW value, for HSEsol). Despite the very different band gaps, the phonon frequencies as a function of the excited charge density evolve similarly: we observe a stabilization of the soft mode at an excited charge density of 0.09 electron per formula unit (see Supplementary Material [31]). Hence, although our DFT-based description may slightly alter the critical concentrations at which transitions occur, the present qualitative behavior should be correct, especially given our results in Fig. 2 for the dynamical lattice instabilities of the paraelectric cubic state.

At last, let us discuss the possibility of having photo-315 induced phase transitions to (small self-trapped, or ST) 316 polaronic states. Recent (doping) calculations have 372 shown that ST holes were unlikely in PTO [49, 50], but 373 could potentially be stabilized in cubic BTO [49]. In order to check the appearance of polaronic states, we ran calculations in BaTiO3 under photoexcitation with $_{\rm 321}$ $n_{ph}=0.075$ e/f.u. and $n_{ph}=0.150$ e/f.u (with no sym-322 metry). No trace of polaronic states was found.

In conclusion, we have shown that light absorption is 380 [15] viable route to switch on/off specific lattice instabilities in complex materials such as ferroelectric perovskite oxides. Via first-principles calculations we have demonstrated the microscopic mechanism leading to a lightcontrolled stabilization of the cubic phase in BTO but 329 not in PTO. The calculations are a promising tool to en-330 gineer and optimize such light-triggered transformations, 388 suggesting strategies for their practical realization in suitably chosen materials. The results presented in this work could offer a route to design light-controlled memory, by accessing light-induced metastable states (see Fig. 4 in the Supplemental Material [31] and related discussion).

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We also acknowledge support from the bandgap. Furthermore, excitonic effects are described 338 National Research Fund, Luxembourg through project within a mean-field approximation using constrained 339 INTER/ANR/13/20/NANOTMD (E. T. and L. W.) DFT to mimic excited states occupations. First, we per- 340 and the intermobility program (Grant No. 15/9890527 formed G_0W_0 [44] calculations with the Yambo [45] pack- 341 Greenox, J. I. and L. B.). C. P. thanks the AHPCC and a in order to accurately describe the conduction and va- 343 thank B. Dkhil and P. Ruello for interesting discussions.

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- See Supplementary Material at ... for more details which 412 includes Refs. [12, 14, 17-19, 37, 42, 43, 45-48, 51? 413 ? -58]. We detail the symmetry and atomic displace-414 ment evolution under excitation of carriers; Nudge Elas-415 tic Band calculations show that the transition is likely 416 to be first order and that the non-polar phase are not 417 metastable, but rather unstable; we discuss the occur-418 rence of a monoclinic phase in BaTiO₃; Volume changes 419 under illumination are compared with hydrostatic pressure results from the literature. The evolution of Inter-421 atomic Force Constants is also depicted. Finally, via the GW-approximation and hybrid DFT calculations, we discuss the relevance of many-body effects for the photo-424 iduced stabilization of the cubic phase. 425
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- Pump-probe experiments on BiFeO₃ in Ref. [7] estimate 428 $n_{nh} \sim 5 \times 10^{19} \text{e cm}^{-3}$ with maximum used fluences 429 of $\sim 50 \mu \text{J.cm}^{-2}$. In Ref. [12] (which observes a mono-430 clinic phase in BaTiO₃ nanowires) fluences ranging up to 431 $600\mu \text{J.cm}^{-2}$ are used, which would correspond to a car-432 rier concentration of $\sim \frac{600}{50}.5 \times 10^{19} \approx 6 \times 10^{20} \text{e cm}^{-3}$. 433 This order of magnitude matches the region where we 434 observed flattening of the energy landscape in Fig. 3a 435 (around the $000/u^+u^+u^+$ to $000/00w^+$ transition) which 436 could potentially harbor a monoclinic phase as discussed 437 in the Supplementary Material [31]. 438
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