Rapid and direct synthesis of complex perovskite oxides through a highly energetic planetary milling

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The search for a new and facile synthetic route that is simple, economical and environmentally safe is one of the most challenging issues related to the synthesis of functional complex oxides. Herein, we report the expeditious synthesis of single-phase perovskite oxides by a high-rate mechanochemical reaction, which is generally difficult through conventional milling methods. With the help of a highly energetic planetary ball mill, lead-free piezoelectric perovskite oxides of (Bi, Na)TiO₃, (K, Na)NbO₃ and their modified complex compositions were directly synthesized with low contamination. The reaction time necessary to fully convert the micron-sized reactant powder mixture into a single-phase perovskite structure was markedly short at only 30–40 min regardless of the chemical composition. The cumulative kinetic energy required to overtake the activation period necessary for predominant formation of perovskite products was ca. 387 kJ/g for (Bi, Na)TiO₃ and ca. 580 kJ/g for (K, Na)NbO₃. The mechanochemically derived powders, when sintered, showed piezoelectric performance capabilities comparable to those of powders obtained by conventional solid-state reaction processes. The observed mechanochemical synthetic route may lead to the realization of a rapid, one-step preparation method by which to create other promising functional oxides without time-consuming homogenization and high-temperature calcination powder procedures.

Complex perovskite oxides containing suitable chemical elements are highly versatile and present a broad range of useful functionalities with unique physico-chemical properties. Traditionally, the majority of these materials have been produced by a solid-state reaction process, in which several intricate processing steps, including a long period of mixing or grinding of the reactant powders, multiple and prolonged calcination steps at high temperatures, and intermediate wet milling processes, are essential to create the proper chemical homogeneity and the desired structures¹². Recently, the field of mechanochemistry has seen a considerable upsurge of interest, mainly driven by the quest for a solvent-free, direct reaction and the low-temperature synthesis of various functional oxides³. The mechanochemical synthetic route is simple to implement, sustainable, and highly energy-efficient when typically carried out using a milling technique. The greatest potential lies in the possibility of promoting solid-state chemical reactions by which any reactant changes into any product at room temperature, as this is possible because the reaction is triggered by means of mechanical stress instead of heat to overcome all thermodynamic reaction barriers⁴–⁶. For the past two decades, intense efforts have been devoted to preparing various functional oxides, including PbTiO₃⁷, PbZrO₃⁸, Pb(Zr, Ti)O₃⁹–¹¹, (Pb, La)(Zr, Ti)O₃¹², Pb(Mg, Nb)O₃¹³–¹⁷, Pb(Mg, Nb)O₃-PbTiO₃¹⁸,¹⁹, BaTiO₃²⁰,²¹, NaNbO₃²²,²³, KNbO₃²⁴, (K, Na)NbO₃²²,²⁵, and BiFeO₃²⁶, mechanochemically using different types of a mill. Among them, a planetary ball mill has been the most popular type of mill adopted in mechanochemical studies of oxides, as is characterized by higher milling energy and thus more efficient in reducing the particle size of initially larger particles²⁷–⁵, compared to vibratory and attrition mills.

Nevertheless, it has been difficult to stabilize a single-phase perovskite in a direct and rapid manner from the initial reactant powder mixture usually consisting of micron-sized particles, which are cheap, easily affordable and typically used in conventional solid-state synthesis. Extending the milling time (typically, from tens to hundreds of hours) has usually been ineffective when attempting fundamental chemical changes. In many cases, a nano-crystalline or amorphous structure of the reactant powders resulted, often with contamination from the

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milling medium. Several studies have shown that the mechanical stress generated by ball impacts can modify the reactant powders to lower the calcination temperature and processing time\(^{25,28,30}\). As the milling time required for nano-crystallization, which is a prerequisite for desired mechanochemical reaction, significantly depends on the initial particle size\(^{31}\). Ohara \textit{et al.} have shown that the total duration to achieve the mechanochemical transformation can be greatly reduced by using the reactant powders at the nanoscale\(^{31}\).

The lack of the rapid and direct synthesis of a perovskite product by a planetary ball mill is attributed to a kinetic limitation associated with difficulties in developing a mill capable of providing highly energetic impacts that are critical to manage the mechanochemical reaction and related yield. Theoretical modeling of kinematic equations describing the energy transfer of the ball to the powder in a planetary mill was well established by Burgio \textit{et al.}\(^{32}\). Kinetic considerations of the mechanochemical behavior of alkali niobate oxide by correlating the milling parameters were also formulated by Rojac \textit{et al.}\(^{33}\). Although the experimental details from different studies were in some cases incomplete, thus thwarting deeper insights into this kinetic issue, our extensive literature survey and the calculation based on Burgio’s model indicated that the existing total energy transferred to the powder or cumulative kinetic energy (as determined by the ball-impact energy, ball-impact frequency and milling time) afforded by planetary mills was typically less than ca. 35 kJ/g for 1 h of milling, depending on design of the mill, the milling medium, the milling speed, and other related factors (Table 1).

Given that the level of energy and number of impacts transferred to the reactant powders may be correlated with an intrinsic activation energy barrier of the reaction system, at a higher activation energy level of a given reaction, a more energetic milling condition is needed to cause a reaction between starting materials mechanochemically. This also helps to facilitate chemical reactions at a faster rate and minimize chemical losses and/or contamination. From this perspective, we show here that single-phase perovskite oxides can be expeditiously synthesized from micron-sized reactant powders via a mechanochemical reaction by a highly energetic ball milling process, especially with significantly higher ball-impact energy and frequency levels. Experimental results are presented for two widely studied piezoelectric oxides, (Bi\(_{0.5}\)Na\(_{0.5}\)\()\)TiO\(_3\) (BNT) and (K\(_{0.5}\)Na\(_{0.5}\))NbO\(_3\) (KNN), as well as their modified complex compositions, 0.76(Bi\(_{0.5}\)Na\(_{0.5}\))TiO\(_3\) - 0.04(Bi\(_{0.5}\)Li\(_{0.5}\))TiO\(_3\) - 0.2(Bi\(_{0.5}\)K\(_{0.5}\))TiO\(_3\) (BNT-BLT-BKT) and 0.955 (K\(_{0.48}\)Na\(_{0.52}\))NbO\(_3\) - 0.03Bi\(_{0.5}\)(Na\(_{0.3}\)K\(_{0.7}\))ZrO\(_2\) - 0.01(Bi\(_{0.5}\)Na\(_{0.5}\))TiO\(_3\) (KNN-BNKLZ-BNT), as the solid-state syntheses of BNT- and KNN-based oxides and their chemical modifications are currently major challenges facing research on lead-free piezoelectrics\(^{34}\). Using the mechanochemically derived powders, the sintering characteristics and bulk piezoelectric properties are demonstrated and compared to those obtained from a conventional solid-state process.

**Methods**

**Materials.** The reactants used in this study were K\(_2\)CO\(_3\) (\(>99.0\%\), \(-150\mu\)m, Sigma-Aldrich), Na\(_2\)CO\(_3\) (\(>99.5\%\), \(-10\mu\)m, Sigma-Aldrich), Li\(_2\)CO\(_3\) (99.997\%, \(-20\mu\)m, Sigma-Aldrich), Nb\(_2\)O\(_5\) (99.9\%, \(-2\mu\)m, Sigma-Aldrich), Bi\(_2\)O\(_3\) (99.9\%, \(-10\mu\)m, Sigma-Aldrich), ZrO\(_2\) (99.0\%, \(-5\mu\)m, Sigma-Aldrich), and TiO\(_2\) (\(>99.9\%\), \(-1\mu\)m, Sigma-Aldrich), which are typically used in a conventional synthetic process. The carbonate powders of K\(_2\)CO\(_3\), Na\(_2\)CO\(_3\), and Li\(_2\)CO\(_3\) were dried at 120°C for 24 h before use owing to their hygroscopic characteristics. Also, all treatments of these powders, including weighing and drying, were carefully performed within a glove box filled with an Ar gas atmosphere. Stoichiometric Bi\(_2\)O\(_3\)-Na\(_2\)CO\(_3\)-TiO\(_2\) and K\(_2\)CO\(_3\)-Na\(_2\)CO\(_3\)-Nb\(_2\)O\(_5\) powder mixtures were used for the syntheses of pure BNT and KNN, respectively. The general chemical reactions of the reactants for the formation of BNT and KNN are as follows:

\[
0.25\text{Bi}_2\text{O}_3 + 0.25\text{Na}_2\text{CO}_3 + \text{TiO}_2 \rightarrow (\text{Bi}_{0.5}\text{Na}_{0.5})\text{TiO}_3 + 0.25\text{CO}_2 \uparrow
\]

\[
0.25\text{K}_2\text{CO}_3 + 0.25\text{Na}_2\text{CO}_3 + 0.5\text{Nb}_2\text{O}_5 \rightarrow (\text{K}_{0.5}\text{Na}_{0.5})\text{NbO}_3 + 0.5\text{CO}_2 \uparrow
\]

Additionally, their respective modified compositions were used as templates to assess the feasibility of preparing more complex compositions, specifically, 0.76BNT-0.04BLT-0.2BKT and 0.955KNN-0.03BNKLZ-0.015BNT, which were recently discovered to possess excellent piezoelectric properties\(^{35,36}\). These modified oxides were prepared using the stoichiometric Bi\(_2\)O\(_3\)-Na\(_2\)CO\(_3\)-TiO\(_2\) and K\(_2\)CO\(_3\)-Na\(_2\)CO\(_3\)-Nb\(_2\)O\(_5\) powder mixtures, respectively.

**Highly energetic planetary milling and mechanochemical synthesis.** Mechanochemical treatments were conducted using a planetary ball mill that was specially developed to provide the much higher milling speed required to increase both the impact energy and frequency. Several design concepts were used to maximize the rotational speed, including the anti-directional rotation mode of the disk and vial, the frictional contact between the disk and vial, the dimensions (diameter, height) of the disk and vial, the mechanical balance of the vial-embedded supporting disk at higher rotational speeds and efficient cooling with water to realize the high-speed milling operation at room temperature inside the vial (Figure S1)\(^{11}\). The vial was 5.2 cm in diameter \(D_v\) and 100 cm\(^2\) in volume. The distance between the rotational axes of the supporting disk and vial \(R_v\) was 5.3 cm. The ratio of the rotational angular speeds of the vial \((\omega_v)\) and supporting disk \((\omega_p)\) was 2.4 \((\omega_v = \frac{2.4 \cdot \omega_p}{\omega_p})\). The attainable values of rotational speeds of the disk and vial were ca. 1100 and ca. 2640 rpm at a maximum, respectively. In all cases, ca. 5 g of a stoichiometric powder mixture was placed in a vial with balls \((5, 8, 10 \text{ and } 12 \text{ mm in diameter } d_b)\), both made of tungsten carbide \((15.1 \text{ g/cm}^3 \text{ in density } \rho_b)\). The ball-to-powder weight ratio was kept at 36:1. Hence, the numbers of balls \((N_b)\) incorporated were 185, 45, 23 and 13 when the ball diameters were 5, 8, 10 and 12 mm, respectively. The rotational speeds of the disk and vial were set to 1000 and 2400 rpm, respectively, in the present experiments. The milling experiments were performed at different times in an argon atmosphere \((ca. 0.3 \text{ MPa})\).
Table 1. Kinetic parameters of the high-energy planetary ball milling process for the mechanochemical synthesis of various perovskite oxides. BNT: (Bi, Na)TiO₃, KNN: (K, Na)NbO₃, PT: PbTiO₃, PZ: PbZrO₃, PZT: Pb(Zr, Ti)O₃, PLZT: (Pb, La)(Zr, Ti)O₃, PMN: Pb(Mg, Nb)O₃, PMN-PT: Pb(Mg, Nb)O₃-PbTiO₃, BT: BaTiO₃, NN: NaNbO₃, KN: KNbO₃. All mills including that used in this work were of the planetary ball-mill type: P4 (Fritsch Pulverisette 4), P5 (Fritsch Pulverisette 5), and PM400 (Retsch PM400). ▼ Ball-impact energy[^32]:

\[
\Delta E_b = \frac{1}{2} \rho_b \pi d_b^3 \left( \frac{3}{6} \right) \omega_p^2 \left( \frac{W_p}{W_b} \right)^2 \left( \frac{D_b - d_b}{2} \right)^2 \left( 1 - 2 \frac{W_p}{W_b} \right) - 2 R_t \left( \frac{W_p}{W_b} \right)^2 \left( \frac{D_b - d_b}{2} \right)
\]

[^32]: Ball-impact frequency[^32]:

\[
\nu_i = N_b K \left( W_p - W_v \right)
\]

[^32]: Cumulative kinetic energy[^32]:

\[
E_{\text{cum}} = \Delta E_b \nu t / m_p
\]

Here, \( \rho_b \) is the density of the balls, \( d_b \) is the diameter of the balls, \( W_p \) is the rotational angular speed of the supporting disk, \( W_b \) is the rotational angular speed of the vial, \( D_b \) is the diameter of the vial, \( R_t \) is the distance between the rotational axes, \( N_b \) is the number of balls, \( K \) is a constant, \( t \) is the milling time and \( m_p \) is the powder weight. The constant \( K \) was 1.5 cited from ref. 46. For other studies, the values of \( E_{\text{cum}} \) calculated assuming 1 h of milling are given in the parentheses for comparison with those of this work. The mechanochemical formation of the perovskite product was assessed based on the reported XRD patterns: ● (phase-pure), ○ (dominant but with contamination or an unreacted phase), △ (partly formed), × (not formed).

Four different kinetic conditions, referred to as condition numbers 1, 2, 3 and 4, were adopted for the mechanochemical treatments. The related kinetic parameters are presented in Table 1. The key parameters of the ball-impact energy \( \Delta E_b \) and the ball-impact frequency \( \nu_i \) were controlled such that balls with a larger diameter resulted in higher \( \Delta E_b \) and smaller \( \nu_i \) values at a constant ball-to-powder weight ratio. According to the model derived by Burgio et al.\[^{32}\], \( \Delta E_b \) increases from 115 to 1209 mJ/hit when the ball diameter increases from 5 to 12...

| Material | Mill | Ball/\( d_b \) (mm) | Disk speed (rpm) | \( \Delta E_b \) (mJ/hit) | \( \nu_i \) (s^{-1}) | \( t \) (h) | \( E_{\text{cum}} \) (kJ/g) | Product | ref |
|----------|------|---------------------|-----------------|------------------------|-------------------|----------------|----------------|----------|-----|
| BNT/KNN No. 1 | WC/5 | 1000 | 115 | 15725 | 2.5 | 3257(1303) | ● ○ | This work |
| BNT/KNN No. 2 | WC/8 | 1000 | 421 | 3825 | 2 | 2520(1160) | ● ●● | This work |
| BNT/KNN No. 3 | WC/10 | 1000 | 760 | 1955 | 2 | 2140(1070) | ●●● | This work |
| BNT/KNN No. 4 | WC/12 | 1000 | 1209 | 1105 | 2 | 1924(962) | ○ ●● | This work |
| BNT/KNN No. 5 | YSZ/3 | 1000 | 10.8 | 72250 | 5 | 2800(560) | × × | This work |
| BNT/KNN No. 6 | YSZ/5 | 1000 | 46.4 | 15725 | 2.5 | 1312.5(525) | ○ ○ | This work |
| PT | P5 | WC/20 | 250 | 671 | 200 | 20 | 322(16.1) | ○ | [7] |
| PZ | P5 | WC/20 | 200 | 429 | 160 | 24 | 198(8.2) | △ | [8] |
| PZT | P5 | WC/20 | 200 | 429 | 160 | 80 | 659(8.2) | ○ | [9] |
| PLZT | P5 | WC/20 | 200 | 429 | 160 | 24 | 198(8.2) | ○ | [10] |
| PLZT | PM400 | WC/10 | 200 | 43 | 1500 | 10 | 77(7.7) | △ | [11] |
| PMN | P5 | WC/20 | 200 | 429 | 160 | 40 | 329(8.2) | ○ | [14] |
| PMN | PM400 | WC/20 | 300 | 560 | 340 | 60 | 206(3.4) | ○ | [15] |
| PMN | PM400 | WC/20 | 200 | 250 | 230 | 94 | 97(1.0) | ○ | [16] |
| PMN-PT | P5 | WC/20 | 200 | 429 | 160 | 20 | 165(8.2) | ○ | [18] |
| PMN-PT | PM400 | WC/20 | 300 | 560 | 340 | 64 | 219(3.4) | ○ | [19] |
| BT | PM400 | WC/20 | 200 | 43 | 1500 | 10 | 77(7.7) | × | [20] |
| NN | PM400 | YSZ/10 | 300 | 35 | 900 | 40 | 90(22.7) | ○ | [22] |
| NN | PM400 | YSZ/10 | 200 | 15 | 600 | 400 | 2592(6.5) | ● | [23] |
| NN | P4 | WC/10 | 300 | 370 | 360 | 96 | 1534(16.0) | ● | [23] |
| KN | PM400 | YSZ/10 | 300 | 35 | 900 | 40 | 90(22.7) | × | [22] |
| KN | P4 | WC/20 | 270 | 300 | 320 | 350 | 12096(34.6) | △ | [24] |
| KNN | PM400 | YSZ/10 | 300 | 35 | 900 | 10 | 227(22.7) | × | [22] |
mm. In contrast, the corresponding value of $v_2$ decreased greatly from 15725 to 1105 s$^{-1}$ owing to the strong dependence of this value on the number of balls ($N_v$) incorporated. This counterbalance yielded the cumulative kinetic energy $E_{\text{cum}}$, which is directly proportional to $\Delta E_v$ and $v_2$, between 962–1303 kJ/g for 1 h.

In Table 1, scattered kinetic data available from different studies are also presented for comparison. They clearly indicate that our milling conditions were much more energetic, as characterized by the higher values of $\Delta E_v$, $v_2$, and $E_{\text{cum}}$. Note the remarkably higher $E_{\text{cum}}$ values calculated based on an identical milling time of 1 h in comparison with those from other studies. For comparison, we also prepared powders using a conventional powder process. The identical stoichiometric powder mixtures were homogenized in ethanol for 24 h using a roller mill machine operating at 70 rpm and were dried at 120 °C for 24 h. Calcination was then carried out at 850 °C for 2 h for the BNT-based powder mixtures and at 850 °C for 6 h for the KNN-based powder mixtures.

**Sintering.** To investigate the sintering characteristics, the powders prepared by both methods were mixed with poly-vinyl alcohol (PVA) as a binder and axially compacted into disks 10 mm in diameter and 1 mm thick under shaping pressures of ca. 62 MPa for the BNT-based and ca. 187 MPa for the KNN-based powders. After burning off the PVA at 650 °C, traditional pressureless sintering was carried out in air. The sintering conditions under shaping pressures of

**Characterization.** The structural phases were characterized by an X-ray diffractometer (XRD; D/Max-2500; Rigaku, Tokyo, Japan) using Cu Kα radiation at a power of 40 kV and 15 mA and at a scan speed of 1°/min. A transmission electron microscope (TEM; TES-2100F; JEOL, Tokyo, Japan) was also used. After milling, the degree of contamination from the milling medium was examined using an inductively coupled plasma method (ICP; Optima 8300DV; Perkin Elmer, Waltham, MA, USA). A thermogravimetric analysis (TGA; Setsys Ev 1750; SETARAM, Caluire-et-Cuire, France) was conducted to examine the mechaenochemical reaction of the milled powders. Approximately 50 mg of the powder was placed in a Pt/Rh crucible and heated to 1000 °C at a rate of 10 °C/min in a flowing air atmosphere. The morphology of the sintered samples was studied using a field-emission scanning electron microscope (FE SEM; Sirion; FEI, Eindhoven, Netherlands) with an operating voltage of 20 kV equipped with an energy-dispersive X-ray spectrometer (EDXS). For piezoelectric measurements of the sintered samples, both sides of the disks were polished and painted with a silver paste, after which they were fired at 650 °C for 10 min. The samples were then poled in silicone oil at room temperature. The poling field and poling time were set to 60 kV/cm and 10 min, respectively, after optimization through pre-tests of poling over time and in an electric field. The piezoelectric coefficient ($d_{33}$) was measured using a piezo-$d_{33}$ meter (ZJ-6B; IACAS; Beijing, China).

**Results and Discussion**

Figure 1a shows XRD patterns of the stoichiometric Bi$_5$O$_7$-Na$_2$CO$_3$-TiO$_2$ powder mixture mechaenochemically treated as a function of the milling time under condition number 2 ($\Delta E_v = 421$ mJ/hit, $v_2 = 3825$ s$^{-1}$). With only 5–10 min of milling, most of the crystalline reactant peaks vanished due to nano-crystallization or amorphization and the characteristic diffraction peaks relevant to a perovskite structure began to appear. Meanwhile, an intermediate reaction product identified as Bi$_5$Ti$_4$O$_{12}$ was detected, likely resulting from a mechaenochemical reaction between Bi$_2$O$_3$ and TiO$_2$. This phase completely disappeared after 20 min. Surprisingly, a predominant perovskite structure was stabilized within 20 min, after which only a pure perovskite structure was evident, without any trace of a second phase. The diffraction peaks were indexed as BNT with monoclinic symmetry (JCPDS 01-080-4260, space group: Cc) (3). After 90 min, the Bi phase was detected, indicating decomposition of the perovskite BNT. The finely crystalline characteristic of the perovskite BNT phase was estimated from the broadening of the XRD peaks; according to Scherrer's formula, the crystallite size was ca. 17.0–18.5 nm. Such a structural characteristic is well supported by the TEM results shown in Fig. 1b.

Similar outcomes, such as reductions of the crystallite size, amorphization and the formation of Bi$_5$Ti$_4$O$_{12}$ and perovskite BNT, were noted for the powders prepared under condition numbers 1, 3 and 4 (Figure S2). For condition number 1 ($\Delta E_v = 115$ mJ/hit, $v_2 = 15725$ s$^{-1}$), however, a trace of the perovskite phase began to appear after 30 min of milling. Moreover, considerable reactant peaks such as stable TiO$_2$ were observed up to 60 min, indicating an incomplete reaction during the milling process (Figure S2a). For condition numbers 3 and 4, no appreciable changes were visible compared to condition number 2, but the partly reacted TiO$_2$ remained for prolonged periods of time and the formation of the further promoted Bi phase via PVA became severe (Figure S2b and c).

Figure 2a shows XRD patterns of the stoichiometric K$_2$CO$_3$-Na$_2$CO$_3$-Nb$_2$O$_5$ powder mixtures mechaenochemically treated as a function of the milling time under condition number 2. The intensity of the Nb$_2$O$_5$ peaks rapidly decreased owing to the reduced crystallite size. The peaks belonging to the alkali metal carbonates of K$_2$CO$_3$ and Na$_2$CO$_3$ disappeared after a very short period of milling (5–10 min), accompanied by a slight increase of the background near their reflections, which indicated amorphization. The oxide-assisted amorphization of carbonates during milling is well known in A$_2$CO$_3$-Nb$_2$O$_5$ (A = K or/and Na) systems, for which the reconstruction of CO$_2$$^-$ ions into a carbonato complex results in a loss of long-range periodicity of the carbonates (4). After 30 min, the Nb$_2$O$_5$ peaks disappeared and the only phase detected was an orthorhombic KNN phase (JCPDS 01-071-0946, space group: Amnm2) with a crystallite size of ca. 16.5–16.7 nm based on Scherrer’s formula. The formation of a nano-crystalline perovskite structure was further confirmed by TEM (Fig. 2b).

In contrast, the reactant powders, when subjected to condition number 1, showed continuous peak broadening and a decreased intensity accompanied by a gradual increase in the background with an increase in the milling time. The perovskite phase began to appear after 60 min and then became dominant (Figure S3a). This was notable because the reactant powders experienced much more vigorous collisions ($v_2 = 15725$ s$^{-1}$) as compared to those in condition number 2 ($v_2 = 3825$ s$^{-1}$), clearly indicating that the rate of the mechaenochemical reaction...
was decreased appreciably by the lower $\Delta E_b$ ($=115 \text{ mJ/hit}$). For condition number 3, the perovskite KNN dominated after 40 min, but with the presence of unreacted Nb$_2$O$_5$ (Figure S3b), similar to the BNT case. Despite the application of a higher value of $\Delta E_b$ compared to that used in condition number 2, the observed unaccomplished reaction may be accounted for by the effect of the reduced number of ball collisions due to the lower value of $v_t$ ($=1955 \text{ s}^{-1}$), which was close to half that of condition number 2. Such a sluggish reaction was more evident for condition number 4 with a higher value of $\Delta E_b$ ($=1209 \text{ mJ/hit}$) and lower $v_t$ values ($=1105 \text{ s}^{-1}$) (Figure S3c).

Figure 1. (a) XRD patterns of the mechanochemically treated stoichiometric Bi$_2$O$_3$-Na$_2$CO$_3$-TiO$_2$ powder mixture as a function of the milling time under condition number 2 ($\Delta E_b = 421 \text{ mJ/hit}$, $v_t = 3825 \text{ s}^{-1}$). (b) TEM bright-field image and selected-area diffraction pattern of the powders mechanochemically treated for 40 min.

Figure 2. (a) XRD patterns of the mechanochemically treated stoichiometric K$_2$CO$_3$-Na$_2$CO$_3$-Nb$_2$O$_5$ powder mixtures as a function of the milling time under condition number 2 ($\Delta E_b = 421 \text{ mJ/hit}$, $v_t = 3825 \text{ s}^{-1}$). (b) TEM bright-field image and selected-area diffraction pattern of the powders mechanochemically treated for 40 min.
Additionally, we investigated the possibility of synthesizing more complex compositions, specifically, 0.76BNT-0.04BLT-0.2BKT and 0.955KNN-0.03BNKLZ-0.015BNT, under an identical milling condition (40 min under condition number 2). Figure 3a and b show that the relatively well-defined XRD peaks precisely correspond to a perovskite structure without any second phases in both compositions. The crystallite sizes of the perovskite phase were ca. 16.4 nm for the BNT-BLT-BKT and ca. 17.6 nm for the KNN-BNKLZ-BNT, similar to those of the unmodified powders. These results confirmed that the rapid formation of nano-crystalline perovskite oxides by a high-rate mechanochemical reaction was highly reproducible, even if many reactants were involved. Promisingly, the minor influence of the number of starting reactants and the induced complexity of the composition on the formation of the perovskite structure offer more freedom in the choice of compositions in a multi-component oxide study.

Regarding contamination from the milling medium during milling, no trace of WC was visible in the XRD patterns of any of the powders prepared in this work (condition numbers 1, 2, and 3). An additional ICP analysis revealed that the amounts of tungsten (W) and cobalt (Co) as possible contaminants were at an acceptable level of less than 100 ppm and 20 ppm, respectively. However, prolonged milling with an excessive value of ΔEb as in the case of condition number 4 induced contamination from the milling medium (Figure S3c).

The observed drastic structural change, leading to the rapid formation of single-phase perovskite oxides, was obviously a result of a high mechanochemical reaction rate among the reactants of a type not normally observed in previous mechanochemical studies or under conventional milling conditions. Regardless of the number and nature of the reactant powders, such a high-rate solid-solution reaction process was realized by enhancing the key kinetic parameters of ΔEb and v, The underlying mechanisms could be understood on the basis of defect-enhanced solid-state reaction processes in which the reactant particles undergo a significant size reduction via the fracturing and generation of sub-grain structures, typically leading to a nano-crystalline or amorphous structure and hence the enhanced degree of solid-state diffusivity and chemical reactivity.

The XRD results for the different milling conditions always indicated the presence of a nano-crystalline or amorphous state, generally referred to as “mechanochemically activated” in other studies. This state usually appeared before the first appearance of the crystalline perovskite product. This trend was sensitive to the applied ΔEb, determining the mechanochemical reaction pathway. No chemical reaction for the formation of the perovskite product could commence below a certain minimum value of ΔEb, as shown by the syntheses of BNT and KNN under condition numbers 5 and 6 using low-density YSZ balls (Figure S4a and b). Moreover, at these energy levels, the perovskite product could be governed neither by v nor by Ecum, the values of v and Ecum utilized for condition number 5 showing no formation of the perovskite BNT and KNN were sufficiently high at v = 72250 s⁻¹ and Ecum = 2800 kJ/g. It is therefore reasonable to assume that a critical ball-impact energy Ec, above or below which either a nano-crystalline/amorphous or a perovskite product is formed, exists for reactions (1) and (2), and this critical energy is correlated with an intrinsic activation energy barrier possessed by each reaction. Although the precise determination of Ec was challenging due to the limited data available from the present sets of experiments, the minimum energy requirement (i.e., Ec) to trigger the reactions was clearly material-dependent, that is, higher for KNN (46.4–115 mJ/hit) than for BNT (10.8–46.4 mJ/hit) based on the XRD results. We also note that the observed energy levels were within a similar range of existing ball-impact energy levels applied in previous mechanochemical studies of several oxide systems. On the other hand, once the applied ΔEb exceeds Ec and hence the balls collide with sufficiently high kinetic energy for the reaction to be successful, the mechanochemical reaction can then be promoted by enhancing v, This situation was evidenced by comparing the case of condition number 2 with those of condition numbers 3 and 4 with higher ΔEc and lower v values, demonstrating the presence of certain unreacted oxide reactants (e.g., TiO2, Nb2O5). By clarifying the individual roles of ΔEb and ΔEc, the reaction pathway can be better understood.
During the mechanochemical reaction, our findings also contradict earlier reports by Abdellaoui et al.\textsuperscript{27,43}, who reported that the cumulative kinetic energy $E_{cum}$ or shock power, defined as the product of $\Delta E_b$ and $v_t$, is the only factor responsible for the formation of the end product.

In Fig. 4a, the energy map, i.e., $\Delta E_b$ (equation 3) vs. $E_{cum}$ (equation 5), was constructed by assessing the presence of perovskite BNT and KNN phases based on the XRD patterns with different milling conditions. The resulting maps fully defined the energy regions required for both the initiation and, more importantly, completion of the perovskite product of each oxide system: none of the studies showed such complete maps, only providing evidence of the initiation of the perovskite product for several oxide systems owing to limited kinetic energy levels\textsuperscript{33}. From these maps, the milling times required for the initiation ($t_i$) and completion ($t_f$) of each perovskite product were determined with different values of $\Delta E_b$ (Fig. 4b). Both $t_i$ and $t_f$ varied significantly with $\Delta E_b$, revealing different reaction thresholds and ends for each oxide system. The shortest $t_i$ (ca. 5 min for BNT and ca. 20 min for KNN) and $t_f$ (ca. 20 min for BNT and ca. 30 min for KNN) observed for condition number 2 suggested that the reaction rate was the highest. However, the reaction rate decreased for conditions 1, 3 and 4 with increases in $t_i$ and $t_f$. These results clearly suggest that the interplay between $\Delta E_b$ and $v_t$ strongly affects the reaction rate, demanding the proper conditions between them.

The milling-time-dependent TGA scans provided important information about the mechanochemical reaction of the powder mixtures. Because the evolution of CO$_2$ gas with a loss of weight is always indicative of the occurrence of chemical reactions (1) and (2), we were able to evaluate the degree of the completion of the reactions from the amount of CO$_2$ gas released from the thermal decomposition of the carbonates remaining in the milled powder. The TGA curves (Figs 5 and S5) show a decrease in the weight loss of both powder mixtures with a longer milling time. This trend implies the occurrence of reactions (1) and (2) for the formation of the perovskite BNT and KNN. The overall weight losses of the non-milled Bi$_2$O$_3$-Na$_2$CO$_3$-TiO$_2$ and K$_2$CO$_3$-Na$_2$CO$_3$-Nb$_2$O$_5$ mixtures amounted to ca. 5.09\% and ca. 11.58\%, respectively, slightly higher than those calculated when assuming complete decomposition of the alkali metal carbonates, i.e., ca. 4.94\% for Bi$_2$O$_3$-Na$_2$CO$_3$-TiO$_2$ and ca. 11.35\% for K$_2$CO$_3$-Na$_2$CO$_3$-Nb$_2$O$_5$. The small discrepancy between the theoretical and experimental values may have originated from the evaporation of small amounts of H$_2$O adsorbed by the hygroscopic reactants during the preparation processes. Notably, their respective weight losses greatly decreased to ca. 2.07\% and ca. 3.07\% after 40 min of milling under condition number 2.
For both of the powder mixtures (Fig. 5a and b), there was a common trend in that the carbon content gradually decreased at a certain milling time, after which there was a sudden decrease followed by no significant change, suggesting that the reaction approached a steady state. The observed sudden change was clearly caused by the intense formation of perovskite BNT and KNN products from the accelerated mechanochemical reaction. Furthermore, the existence of a certain milling duration when this sudden drop occurred implied that the mechanochemical reaction began with an activation period, during which the size reduction and crystal deformation processes continually took place to create chemically active sites, eventually facilitating the chemical reactions.

According to Fig. 5, the activation period of reaction (2) for the predominant formation of KNN under condition number 2 was ca. 30 min, which was longer than that of reaction (1) for BNT (ca. 20 min); for condition numbers 1 and 3 with lower and higher values of $\Delta E_b$, respectively, the activation periods became longer. All of these results were consistent with the XRD behavior (Figs 1a and 2a, S2a and S3a). Consequently, the cumulative kinetic energy $E_{cum}$ necessary to overcome such activation periods, characterized by a nano-crystalline or an amorphous state based on the XRD results, was calculated and found to be ca. 387 kJ/g for the BNT and ca. 580 kJ/g for the KNN for condition number 2. The degree of the completion of the reaction, i.e., the reaction yield, was determined from the perovskite periods (steady levels) for each condition (insets of Fig. 5). For condition number 2, these values were ca. 63.3 ± 11.7% for BNT and ca. 78.9 ± 2.7% for KNN. The lower reaction yield observed for BNT, despite the fact that its formation was easier than that for KNN (Fig. 4), may have been related to the formation of a further reacted Bi phase. Although the Bi phase was invisible within the perovskite periods in the XRD patterns, we cannot exclude the possibility of the coexistence of a Bi phase with the perovskite BNT before its crystallization. In contrast, the KNN showed a predominant formation of perovskite products. Nevertheless, some carbonates, although undetectable by our XRD system due to their amorphous characteristics, likely did not decompose. Considering the required milling time for different conditions, the reaction efficiency (i.e., the ratio of the reaction yield to the milling time) was also highest for condition number 2.

Conclusively, we ascribe the present findings of a much shorter reaction time (ca. 30–40 min under condition number 2) necessary to stabilize a single-phase perovskite structure to the highly energetic milling conditions using the greater rotational speed of the disk (1000 rpm), which allows an enhanced reaction rate induced by much more vigorous ball collisions at more than a certain minimum level of kinetic energy. Given equations 3–5, the most critical milling parameter to increase both $\Delta E_b$ and $v_i$ is known to be $W_p$. For condition number 2, equation 5 approaches this, as shown below.

$$E_{cum} = 3.22 \times 10^{-10} \times W_p^{3.5} \times t$$  (6)
This relationship implies that $E_{\text{cum}}$ is highly sensitive to $W_p$ with linear $W_p^3$ dependence; see the fitted curves of $E_{\text{cum}}$ vs $W_p$ in Fig. 6 when $t = 1, 2$ and $3$ h. In this figure, the literature data pertaining to $E_{\text{cum}}$ and $t$ required for the dominant formation of various perovskite oxides are also presented and compared to those used in this work. The prolonged period of milling between 20–400 h needed in many mechanochemical studies is attributed to the lower level of $W_p$ (200–300 rpm). For instance, the milling time of 400 h required to reach an $E_{\text{cum}}$ value of 2592 kJ/g at a disk speed of 200 rpm (used in ref. 23) can be dramatically reduced to less than 3 h when adopting condition number 2 as defined in this work.

Using mechanochemically derived powders, the sintering characteristics were investigated and compared to those obtained from a conventional solid-state reaction process. All of the ceramics including the BNT, BNT-BLT-BKT, KNN, and KNN-BNKLZ-BNT types, had pure perovskite structures without any second phases (Figure S7). The absence of changes in the peak positions or intensities for the ceramics prepared by both methods indicated nearly identical structural characteristics (BNT, monoclinic; BNT-BLT-BKT, pseudo-cubic\textsuperscript{44}; KNN, orthorhombic; KNN-BNKLZ-BNT, diphasic rhombohedral-tetragonal\textsuperscript{36}). According to SEM backscattered electron images (BEIs) of the ceramics prepared by mechanochemical synthesis, the homogeneously distributed main elements confirmed nearly complete perovskite solid solutions, together with the formation of well-developed perovskite grains with clear edges, similar to those prepared by conventional solid-state synthesis (Figure S8). The values of the piezoelectric coefficient $d_{33}$ obtained for the ceramics prepared by mechanochemical synthesis were similar to or slightly higher than those for the ceramics prepared by the conventional powder process (Fig. 7). Particularly, the modified BNT-BLT-BKT and KNN-BNKLZ-BNT ceramics prepared by mechanochemical synthesis showed excellent $d_{33}$ values of ca. 211 ± 1.2 pC/N and ca. 267 ± 3.5 pC/N, respectively. These values are comparable to the best properties reported for identical material systems prepared by the conventional solid-state synthesis method (ca. 190 pC/N by Hiruma et al.\textsuperscript{35} and ca. 147–231 pC/N by Lin et al.\textsuperscript{45} for BNT-BLT-BKT and ca. 285 pC/N by Cheng et al.\textsuperscript{36} for KNN-BNKLZ-BNT).

In summary, we have demonstrated that complex perovskite oxides can be expeditiously synthesized from micron-sized reactant powders via a mechanochemical reaction using a highly energetic ball milling process. Only a very short milling time of ca. 30–40 min was required to yield nano-crystalline, single-phase perovskite BNT and KNN oxides reliably, even given their more complex compositions. This was enabled by the high mechanochemical reaction rate induced by enhancements of key kinetic parameters, in this case the ball-impact energy ($\Delta E_b = 421$ mJ/hit) and frequency ($v_t = 3825$ s$^{-1}$). From the mechanochemical reaction outcomes with different values of $\Delta E_b$ and $v_t$, it was demonstrated that $\Delta E_b$ should be higher than a critical kinetic energy $E_c$. 

**Figure 6.** Comparison of the cumulative kinetic energy $E_{\text{cum}}$ and the milling time $t$ taken from the literature (Table 1) with those used in this work. The literature data were limited to those corresponding to the dominant formation of perovskite oxides.

**Figure 7.** Piezoelectric coefficient $d_{33}$ of ceramics sintered using powders prepared both by mechanochemical synthesis for 40 min (denoted by “M”) and conventional solid-state synthesis (denoted by “C”).
(10.8–46.4 mJ/hit for BNT, 46.4–115 mJ/hit for KNN) to initiate the mechanochemical formation of the perovskite product. Additionally, the mechanochemical reaction could be promoted by enhancing $v_t$, but only if the balls had sufficiently high kinetic energy for the reaction to occur. According to the TGA results, the formation of perovskite BNT and KNN required an activation period, during which the solid-state chemical reactivity of the reactant powders continued to increase, ultimately inducing the intense formation of the perovskite product. Furthermore, the formation of BNT was easier than that of KNN in terms of both the energy requirement and the activation period. The values of $E_{\text{act}}$ necessary to overcome the activation period for the formation of a dominant perovskite structure were $ca.$ 387 kJ/g for BNT and $ca.$ 580 kJ/g for KNN. Finally, the mechanochemically derived perovskite powders, when consolidated into bulk ceramics, showed encouraging piezoelectric properties that were not inferior to those prepared by conventional solid-state synthesis. Benefitting from the ability to skip the conventional high-temperature and time-consuming powder procedures, the high-rate mechanochemical route used here is not limited to the presently investigated oxides. It can be readily extended to other technologically important multi-component oxides, possibly irrespective of the choice of composition or the nature of the reactant (e.g., chemical stability).

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**Author Contributions**

G.L. contributed to the mechanochemical reaction discussion and wrote the manuscript. J.P. and M.L. conceived the main idea and supervised the manuscript preparation. E.P. performed the ball milling experiments and structural analysis. S.Y. and S.B. performed the bulk ceramic preparations and piezoelectric measurements. The manuscript was written through contributions of all authors.

**Additional Information**

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