

Efficient Racemization of the Pharmaceutical Compound Levetiracetam using Solvent Free Mechanochemistry

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Abstract

We present the racemization of an active pharmaceutical ingredient Levetiracetam using a novel approach. We demonstrate the design of a 100% solvent-free process that proceeds by high energy milling inside a regular mixer mill. The kinetics of the racemization process is drastically improved compared to the solution-based approach and illustrate the tremendous potential of mechanochemistry. In this study, we highlight the importance of mixing efficiency regarding data reproducibility and show, in particular, that water contamination has a negative impact on the reaction rate. Moreover, *in-situ* X-ray diffraction gives us first insight of the mechanisms involved at the solid state during the mechanochemical racemization process.

Introduction

More than 50% of commercialized active pharmaceutical ingredients (APIs) are chiral substances.¹ The two enantiomers of a given API have identical physico-chemical properties but can act differently in asymmetric media such as living organisms: if one enantiomer of the API owns the desired therapeutic effect, the counter-enantiomer could exhibit undesired effects.² Access to enantiopure materials is therefore paramount and the development of new methods for the production of pure enantiomers is a central concern in chemical research. Most often, the synthetic routes of the APIs are non-stereoselective and racemic mixtures (*i.e.*, a mixture made of both enantiomers in equal proportions) are finally obtained. The counter-enantiomer is thus considered as an impurity and must be removed, for instance by using chiral HPLC.^{3,4} In this context, methods based on crystallization principles can also be used.^{5–7} Yet, whatever the separation route, the maximum theoretical yield is only 50%. It implies a loss of half of the product and does not fit with the 1st and 2nd principles of Anastas and Warner's green chemistry principles, both reporting the necessity to avoid waste.⁸

Increasing the yield of the process to 100% is possible by converting the undesired enantiomer into a racemic mixture, which can be resolved once again. This transformation, called racemization, can be performed if the chiral entity can reach an achiral state either by using a racemizing agent or by using external parameters (heating, UV irradiation, dissolution etc.).^{9–11} This reaction is generally done in solution as high molecular mobility is required for reactivity purposes, which is lacking in the solid state. Some processes take advantage of the racemization reaction to turn a racemic mixture into an enantiopure end state. Deracemization processes such as Viedma ripening (VR) or Temperature Cycle Induced Deracemization (TCID), convert a racemic conglomerate crystal suspension in equilibrium with a saturated solution into an enantiopure material.^{12,13} In the case of VR, this is permitted by constant attrition of the suspension using glass beads that allows to amplify population imbalances between enantiomeric crystals.¹⁴ Also, productivity and yield of the well-known Preferential Crystallization process^{15,16} can be improved by solution-racemization in a process known as Second Order Asymmetric Transformation.^{17,18} In this process, the constant interconversion of the enantiomers at the liquid state provides a steady mass input to feed the enantiopure seeds and offers control of the supersaturation of the counter enantiomer. Nevertheless, the enantiopurification methods described above all suffer from the same drawback: they require the use of substantial amounts of solvent(s) which can lead to a costly and toxic process. Furthermore, solvents are often hard to remove, leading to high carbon footprint processes. Adding a solvent to facilitate a reaction is not in agreement with the 5th of Anastas and Warner's green chemistry principles that aims at reducing the amount of added auxiliary to a reaction.⁸

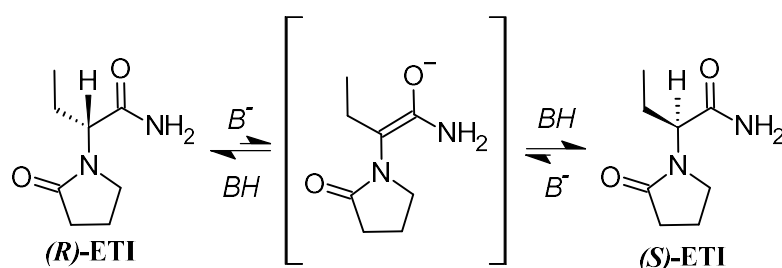
Mechanochemistry is a promising approach to considerably reduce, or suppress, the amount of liquid substances used in chemical processes and has been applied for solvent-free organic synthesis and the preparation or screening of new solid phases of multicomponent materials.^{19,20} It is commonly accepted that transformations under high mechanical stress occur due to energy inputs that may decrease transition energies, alter transition states, create local and transient intense heating, etc.^{20,21} Reaction pathways under high mechanical stress may also differ from the solution pathways, yielding processes with different regioselectivities.^{22,23} Mechanochemical processes are usually performed via Neat-Grinding (NG, *i.e.* in dry conditions) using vibratory or planetary ball mills. The use of a minimal amount of solvent, a.k.a. Liquid-Assisted Grinding (LAG),²⁴ is also common and often improves reactivity, or can be used for the formation of specific solvated solid phases.²⁵ Friščić *et al.* introduced the η parameter to describe the ratio between the volume of solvent and the

mass of solid (expressed in $\mu\text{L.mg}^{-1}$). These authors propose to use this parameter to differentiate mechanochemical ($<1\text{-}2\ \mu\text{L.mg}^{-1}$) from solution-based processes ($>1\text{-}2\ \mu\text{L.mg}^{-1}$).²⁶

Beside numerous advantages mechanochemistry is inherently confronted with mixing and homogenization issues. Ball milling often results in solid materials that exhibit gummy-like texture, leading to sticky materials characterized by poor flowability. The materials are difficult to handle, and often stick to the wall of the grinding jar. The occurrence of such behavior during NG is erratic and is likely due to a number of factors. Solvent formation during the chemical reaction during the process likely has a negative effect on mixing. It is also the case when the generated product has a rubber-like aspect. The so-called “*snow-ball effect*” has been coined to describe the typical coating of the grinding beads by a pasty and sticky materials upon milling.^{27–29} The snow-ball effect may have a positive impact on reaction rates for a part of the powder but most often reduces mixing efficiency.³⁰ Using solid lubricants such as NaCl, SiO₂ or even talc, mitigates this issue.^{29,31}

The use of mechanochemistry in the field of enantiomer conversion and separation remains largely unexplored. Ikekawa *et al.*³² reported the racemization of the natural amino acid L-leucine by NG, but chemical degradation occurred and the yields were poor. Recently, we have published the first example of deracemization using LAG.³³ The process may be considered as VR transferred in a high energy milling apparatus with much lower solvent loads and with much lower process duration ($\eta \sim 0.05\ \mu\text{L.mg}^{-1}$ with a process time of less than 2 hours in our work, while $\eta \sim 5\ \mu\text{L.mg}^{-1}$ and lasts more than 24h in the case of VR).

In the present study, we focus on the chiral API 2-(2-oxopyrrolidin-1-yl)butanamide, (**Scheme 1**) which is used to treat epilepsy.³⁴ The racemate is referred as Etiracetam ((*RS*)-ETI) with the pharmaceutically active *S*-enantiomer referred to as Levetiracetam ((*S*)-ETI). Concerning the solid landscape of Levetiracetam, no hydrate nor polymorphic forms are known. In contrast, Etiracetam, crystallizes as a stable racemic compound and exhibits two enantiotropically related polymorphic forms, with a transition temperature equals to 30.5 °C. A racemic dihydrated solid form has also been reported.^{35,36} (*S*)-ETI being the active enantiomer and (*R*)-ETI having no pharmaceutical activity, (*R*)-ETI is typically transformed into (*RS*)-ETI by racemization with further (*S*)-ETI recovered via chiral resolution.



Scheme 1 : Racemization reaction of (*S*) or (*R*)-ETI using a base as a racemizing agent

A solvent-based racemization route of Levetiracetam which requires the use of sodium methanolate as a racemizing agent and refluxing the mixture for 12 hours has been reported.^{37,38} The racemization mechanism has also been investigated *in-silico* by Li *et al.*³⁹ who concluded that the process requires proton abstraction using hydroxide ions, giving rise to an achiral intermediate (Scheme 1), before proton re-addition. In this paper, our aim is to evaluate the possibility to perform racemization of Levetiracetam without the use of a solvent through a mechanochemical NG process. This work focuses on the efficiency of mixing and process

reproducibility to demonstrate the potential of mechanochemistry in the field of enantiomer conversion.

Materials and methods

Chemicals

(*S*)-2-(2-oxopyrrolidin-1-yl)butanamide ((*S*)-ETI or Levetiracetam) was purchased from Xiamen Top Health Biochem. Tech. Co., Ltd. Sodium chloride (NaCl, 99.5% of purity), was purchased from VWR whereas sodium hydroxide (NaOH, 97% of purity) and sodium methoxide (25/30%wt in methanol) were purchased from Alfa Aesar. A list gathering the purity and supplier of all used compounds is given in the supporting information (Supporting Information, Table S1).

Milling procedure

All milling experiments were performed with a Mixer Mill 400 from Retsch in a 10mL jar containing one 12 mm diameter bead both composed of ZrO₂. Milling was performed with a frequency of 30 Hz for variable duration expressed as t_{milling} . For experiments with controlled humidity, jar preparation was entirely performed within a glove bag in which relative humidity (RH) was controlled with N₂ gas below 20%. The materials to be milled (*i.e.*, (*S*)-ETI, NaOH and NaCl) were all stored for a minimum of one week inside a desiccator under P₂O₅. (Figure S8 in Supporting Information). The jar was sealed with parafilm inside the glove bag prior to milling (performed under ambient conditions). Once the milling stops, the powder was immediately analyzed by cHPLC.

cHPLC

cHPLC analyses were performed on a Ultimate 3000 system, with a UV detection at 210 nm. Milled material was dissolved in a mixture of Heptane/IPA/TFA (80/20/0.1, v/v/v). The mobile phase was a heptane/IPA (90/10 v/v) mixture with a 1mL/min flow rate. The stationary phase was a Chiralcel OD-H 5 μ m 250x4.6 mm column from Daicel. The retention times were 16.3 and 20.1 min for (*R*)- and (*S*)-enantiomers, respectively.

In situ Powder X-ray Diffraction measurements

Levetiracetam (100 mg) and NaCl (50 mg) with NaOH (7 mg) were placed in a customized Perspex jar⁴⁰ of 12 mm diameter, and were then neat ground for 4 hours with a 10 mm zirconium ball at 50 Hz using a Fritsch Vibration Ball Mill (Pulverisette 23, Fritsch, Germany).

In situ X-ray diffraction measurements were performed at 30 seconds intervals at the μ Spot beamline (BESSY II, Helmholtz Centre Berlin for Materials and Energy).⁴¹ The experiments were conducted with a wavelength of 0.7314 Å using a double crystal monochromator (Si 111), and the resulting scattering images were integrated with the Dpdak-software.⁴² The data obtained were plotted simultaneously with the software OriginPro 2023 to obtain a temporal evolution of the system mechanically activated.

Results & Discussion

Enabling mechanochemical racemization of levetiracetam

A solution-based reference racemization protocol was reported by Leyssens *et al.*^{37,38} and repeated here both at reflux and at ambient temperature. For that purpose, 200 mg of (*S*)-ETI were dissolved in 10 mL of methanol (MeOH) containing 0.05 eq of sodium methoxide (MeONa). The quantity of MeOH has been chosen to afford a comparable volume to the one of the jars used in milling experiments. **Figure 1** shows that the racemization experiment performed at reflux yield to an enantiomeric excess (*e.e.*) of 88% after 240 minutes with no racemization occurring at ambient temperature after 24h.

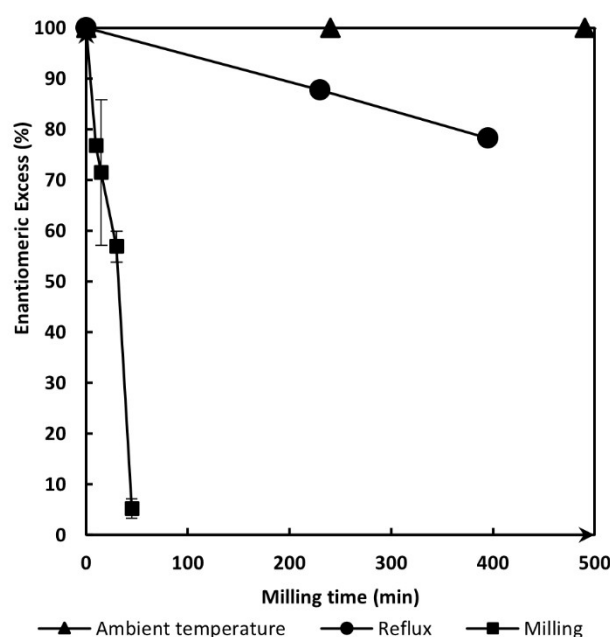


Figure 1 : Evolution of the enantiomeric excess of ETI as a function of time: under milling (black squares), in solution at reflux (black circles), in solution at ambient temperature (black triangles).

When performing the mechanochemical process, through liquid assisted grinding (LAG) using 200mg of (*S*)-ETI and 0.1 equivalent of MeONa added as a MeONa-methanol solution ($\eta = 0.1 \mu\text{L}.\text{mg}^{-1}$) a much faster process is observed. For each milling time (t_{milling}), experiments were repeated at least 4 times and the associated standard deviation is represented by the error bar. **Figure 1** (black squares) highlights that almost complete racemization (*e.e.* of 5%) was reached in a mere 60 minutes. These results show that the conversion rate of the racemization reaction can be considerably increased using a mechanochemical approach.

Albeit successful, the use of MeONa nevertheless comes with some drawbacks: (i) it is usually formulated as a methanol solution which limits the approach to LAG only (no Neat-Grinding), (ii) the base solution is not stable under ambient conditions as reaction with atmospheric water occurs, (iii) the base shows health hazards and its use in conjunction to pulverulent materials should be avoided.

Screening of alternative bases suitable for neat grinding racemization

We therefore decided to look for a base allowing NG racemization. Initially, attempts were performed in the absence of a racemizing agent. This did not lead to any enantio-conversion, hereby confirming the result of the DFT study of Li *et al.* and underlining the central role of proton carriers in the racemization mechanism.³⁹ A screening was then performed to identify a racemizing agent suitable for NG. All the compounds tested are gathered in **Table 1** alongside their pKa values. Milling experiments were systematically performed with $t_{\text{milling}} = 30$ min at 30 Hz using 1 eq. of base.

Table 1 : Summary of the bases tested for milling racemization by NG and their associated pKa. Experiments have been performed at 30Hz with a t_{milling} of 30min.

Base	pKa (measured in water at 25°C) ⁴³	Final % <i>e.e.</i> for $t_{\text{milling}} = 30\text{min}$
1,8-Diazabicyclo(5.4.0)undec-7-ene (DBU)	13.0 ^a	>99%
CaCO ₃	10.3 and 6.4 ^b	>99%
MgCO ₃		>99%
K ₂ CO ₃		>99%
Mg(OH) ₂	- ^c	>99%
Ca(OH) ₂	12.7	>99%
LiOH	13.8	>99%
NaOH	14.8	70%

a, estimated pKa; b, for the first and second acidity respectively; c, As there is no dissociation for magnesium dihydroxide, no pKa value can be measured or estimated

Concerning trials with the notably strong base DBU, no racemization occurred despite the rather high pKa value of 13. It is possible that the large molecular volume of this organic base hinders the proton abstraction of (*S*)-ETI. All the NG experiments performed with carbonate bases did not lead to racemization of (*S*)-ETI, with the *e.e.* upon milling remaining >99%. These bases have lower pKa values (10.3 and 6.4 for the first and second acidity respectively), which could explain the absence of racemization. Lithium, magnesium and calcium hydroxides failed to trigger the racemization reaction. Both the addition of solvent (water or methanol) or the use of a planetary, instead of vibration mill did not impact the outcome. In contrast, NG experiments performed with the strong base NaOH successfully triggered the racemization reaction (Table 1). Based on these results, it appears that the key parameter to trigger proton abstraction in mechanochemical conditions remains the strength of the base. Our results suggest that a pKa threshold corresponding to that of NaOH must be achieved to trigger racemization (*i.e.*, pKa ≥ 14.8). As a comparison, the pKa of MeONa is estimated at 15.5.

Racemization Efficiency

All experiments reported in this section were performed using 200 mg of (*S*)-ETI, 0.1 eq. of NaOH with a grinding duration of $t_{\text{milling}} = 60$ min. Preliminary NG experiments (i.e., $\eta = 0\mu\text{L.mg}^{-1}$) performed were found to be non-reproducible in terms of racemization efficiency: for a total of 8 experiments performed in the same conditions, the final *e.e.* varied from 0.4 to 96.7% with a standard deviation of 42 as shown by the error bars. (**Figure 3**, Exp#1).

This lack of reproducibility likely finds its origin in the high hygroscopic character of solid NaOH,⁴⁴ leading to uncontrolled amounts of water resulting amongst others in poor quality of mixing. Indeed, larger amounts of water can generate a paste-like material due to solubilization of the starting compounds.³⁰ The subsequent low homogeneity during milling can be a source of lacking reproducibility.

To confirm the influence of water on the mixing quality, two sets of milling experiments were performed. A first set (Exp #2, 8 repetitions) is performed by adding 10 μL of water to the grinding jar. This water aliquot does not come from the racemization reaction, therefore the experiments should be considered as LAG ($\eta(\text{H}_2\text{O}) = 0.05\mu\text{L.mg}^{-1}$). The second set (Exp #3, 20 repetitions) is performed by filling the jar under controlled humidity using a glove bag (GB, hereafter) and with prior storage of all compounds under dry conditions (in presence of a desiccant).

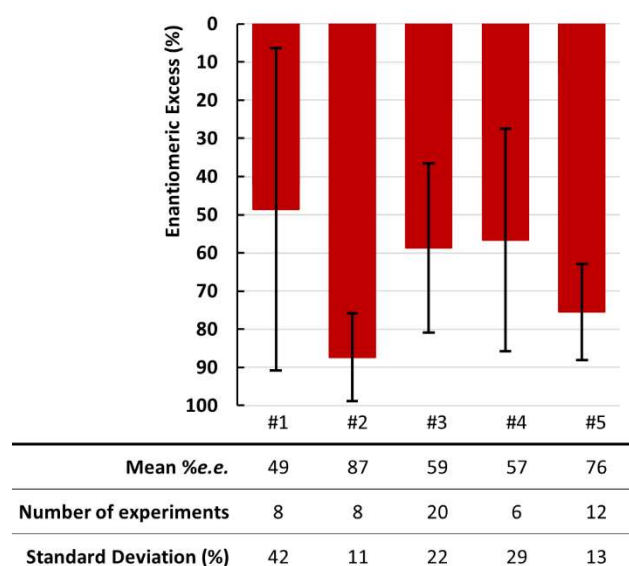


Figure 2 : Graphical representation of the mean value and standard deviation of different sets of milling experiments all performed for 60min at 30Hz : #1 Classic NG; #2 LAG with 10 μL of H_2O ; #3 NG + GB; #4 NG + NaCl; #5 NG + NaCl + GB

The impact of these parameters on the racemization efficiency is examined by comparing the results of Exp#1 to 3 (**Figure 2**). In the case of Exp #2, the results show that the racemization kinetics is drastically reduced to a mean value of 87% *e.e.* when adding only 10 μL of water. As a consequence, the standard deviation is also reduced to 11% (compared to 42.2% for Exp #1). In parallel, these experiments were repeated using 50 μL of water ($\eta(\text{H}_2\text{O}) = 0.25\mu\text{L.mg}^{-1}$) resulting in the same outcome in terms of data reproducibility and racemization rate (Table S8, in the supporting information). In contrast, filling the jar using a GB (Exp #3) had almost no impact on the mean *e.e.* value compared to Exp #1, but the standard deviation decreased to 22%, showing improved reproducibility. Further experiments were performed by filling the jar

under controlled humidity and by adding water with racemization clearly slowing down. These experiments confirm that water contamination has a negative impact on both racemization kinetics as well as reproducibility.

Literature shows several studies highlighting the advantage of adding an inert solid lubricant (such as NaCl, SiO₂, sand, talc, etc...) to the milling media ^{22,29} to provide a better homogeneity inside the jar. ETI generates H₂O when racemizing (**Scheme 1**) thus decreasing the flowability of the powder, we therefore decided to perform NG experiments adding 100 mg of NaCl to investigate the influence of a solid lubricant on racemization kinetics. The 6 repetitions of Exp #4 did not involve the use of a GB to fill the jar. The results show a mean value of 57 % and a standard deviation of 30 % (*i.e.*, between 13 and 87 %*e.e.*). These data are similar to those obtained for Exp #3. Even if the use of NaCl is expected to improve mixing, the rather poor data reproducibility of Exp #4 can still be associated to water trapping by this salt. Therefore, we combined the use of NaCl and water control, filling the jars in the GB in a final series of experiments (Exp #5). This successfully renders the process more reproducible with a standard deviation of only 12 %. In contrast, however, the mean value of the *e.e.* remained at 75.5 %. showing lower racemization kinetics compared to all other experiments (but Exp #2). This can be explained by the dilution effect, leading to a reduced number of contacts between (*S*)-ETI and the racemizing agent NaOH (*i.e.*, due to the increase of (*S*)-ETI/NaCl and NaOH/NaCl contacts).

The results shown in Figure 2 were correlated to a visual observation of the ground material just after grinding. For Exp #1 to 4, the materials were systematically found glued un-evenly on the walls of the jar, especially on its extrema and a remnant of a snow-balling was observed (Figure S3, in the supporting information). Besides, NG experiments performed with a higher amount of NaOH (0.5 eq.) resulted in the material being entirely stuck to the grinding ball, hereby confirming the occurrence of a snow-balling effect (Figure S7). In contrast, conditions of Exp #5 lead to a uniform distribution of the powder and the presence of lumps and sticky material was strongly reduced. This confirms that data reproducibility is correlated to the mixing efficiency, which in turn is favored by a synergetic combination of low water content and the use of a solid lubricant.

Using the optimal racemization and mixing conditions found in Exp #5, a kinetic follow-up of the racemization reaction was performed. For this, a series of 24 experiments has been performed and stopped after different t_{milling} (**Figure 3**). For each milling time, statistics were made over 6 experiments (except for 60min for which 12 experiments were performed). The optimized reproducibility of the results allows to evaluate the $t_{1/2}$ of the racemization reaction at around 80 minutes which shows much faster kinetics compared to solution-based deracemization for which only 80%*e.e.* are reached after 400min (**Figure 1**). Due to the absence of exact temperature control, the data cannot be used to determine an accurate activation energy of the mechanochemical racemization process.

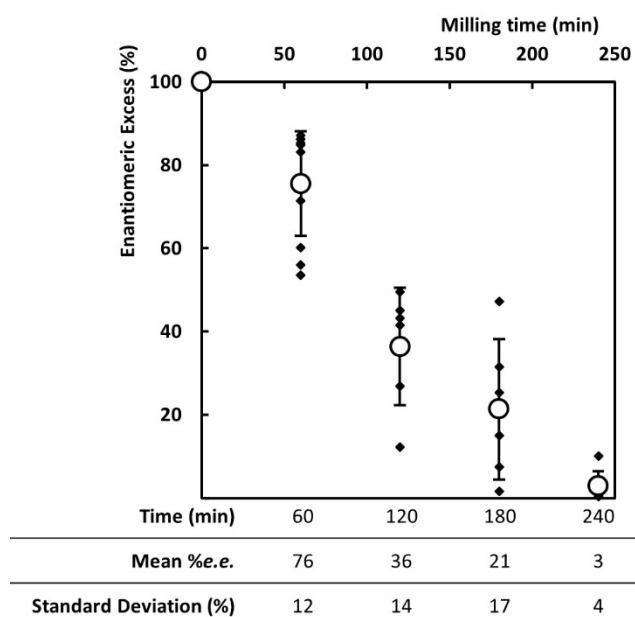


Figure 3 : Graphical representation of the kinetic follow-up of (S)-ETI racemization using 0.1 eq. of NaOH and $m_{\text{NaCl}}/m_{\text{S-ETI}} = 0.5$. The milling frequency is fixed at 30Hz. Black diamonds stand for the raw data whereas empty circles stand for the mean value. The error bars are the standard deviation.

First insights into the racemization mechanism using in-situ PXRD monitoring

To gain insight into the racemization mechanism, we performed *in-situ* X-ray diffraction monitoring of the mechanochemical reaction using the conditions of Exp #4 (*i.e.*, use of NaCl but no control of the humidity during both jar filling and milling) since it was easier to replicate these experimental conditions at the synchrotron facilities.

As shown in **Figure 4**, the initial system (*(S)*-ETI) goes through different steps before ultimately leading to form I of (*RS*)-ETI. First, only the diffractograms of both (*S*)-ETI and NaCl are detected. Their diffracted intensity progressively decreases within the first 10 minutes and become almost undetectable at 17 minutes. The disappearance of the diffraction peaks cannot be merely attributed to amorphization of (*S*)-ETI since the peaks of NaCl also decrease. Instead, the absence of a PXRD signal is more likely due to the occurrence of a snow-ball effect. After 17 minutes, the peaks of Form I of (*RS*)-ETI and NaCl are suddenly detected and remain until the end of the experiment. At the same time, it is also possible to observe the signal of Form II of (*RS*)-ETI (peak in the red box) but the latter is weak and disappears progressively. When this experiment was reproduced (see Supporting Information), the same trend was observed but the time after which the racemic phases appear varied from one experiment to another, consistently with what observed when sampling (**Figure 3**).

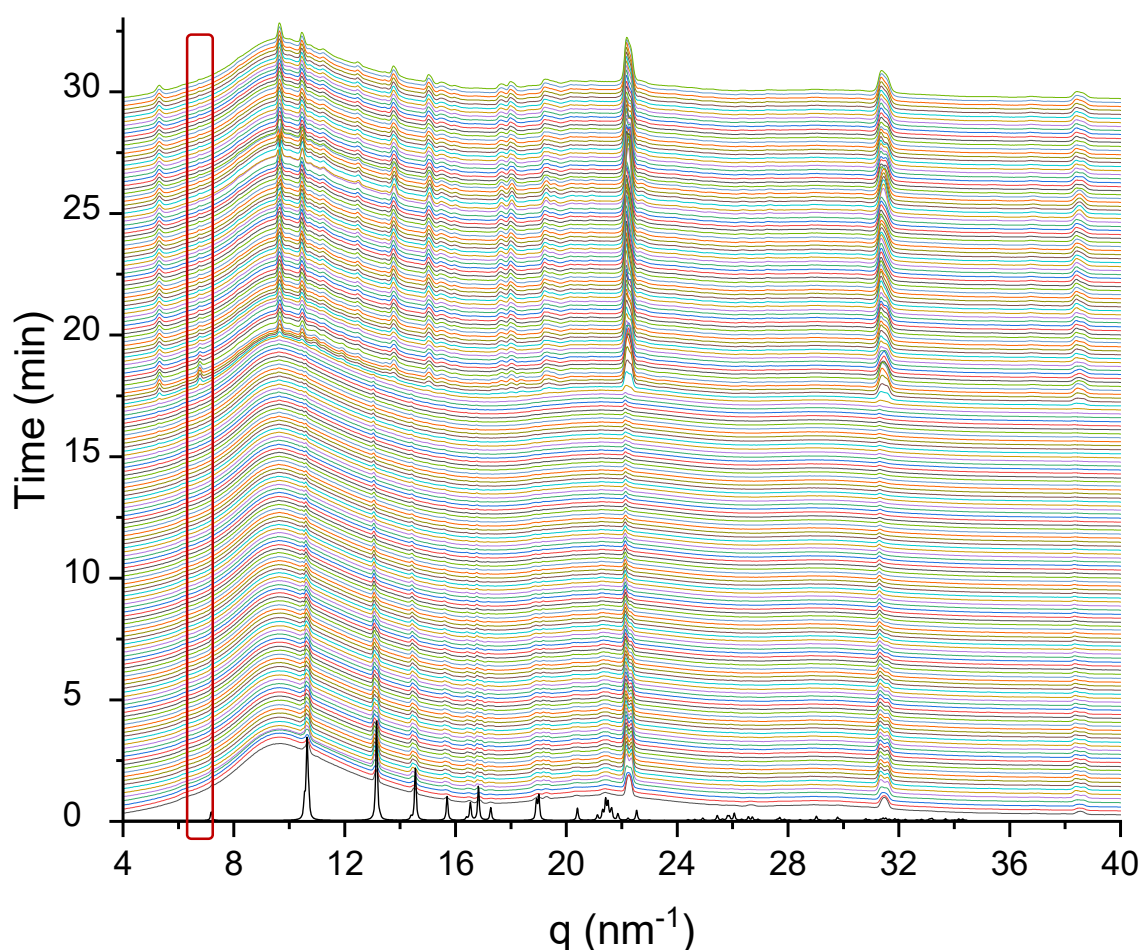


Figure 4. Temporal evolution of PXRD patterns collected at BESSY every 30 seconds vs the diffractogram of (*S*)-ETI (black line). No background subtraction was applied: the big slope at low angle is due to the jar contribution. Peaks around 22.2 and 31.5 nm⁻¹ belong to NaCl

Although further investigation would be needed to fully understand the kinetics, we can already extract unprecedented information on the mechanism of the mechanochemical racemization. From the *ex-situ* sampling data (**Figure 3**) a continuous evolution of the *e.e.* as a function of grinding time from 100 % to 0 % is to be expected. Instead, both *in-situ* dataset using conditions of Exp #4 show a sudden appearance of (*RS*)-ETI crystals after the occurring snowball effect. The combination of both findings confirms that the racemization reaction should not occur at the crystalline state but likely requires the formation of a disordered phase that is not detected by X-ray diffraction (nor by thermal analysis after sampling). Such a transient phase could contain a large amount of amorphous parts as a result of complex interactions between the ground material and trapped water.

Conclusions

In this paper, we show that Levetiracetam can easily be racemized using LAG and MeONa (in MeOH) as a racemizing agent. The mechanochemical racemization occurs much more rapidly than the solution-based process reaching a racemic mixture in less than 100min instead of 80%*e.e.* after a mere 400min. Moreover, we show that neat grinding can also efficiently lead to Levetiracetam racemization in only 240 min using NaOH as a base. However, the racemization kinetics drastically slow down when water is present, even in very small amounts. Water either comes from the racemization reaction or from atmospheric contamination, creating a paste-like material giving rise to a snowball effect. This leads to mixing issues and inhomogeneity created during neat grinding. To improve the reproducibility of the results, the humidity inside the milling jars was reduced and a solid auxiliary (NaCl) added (to prevent the formation of the pasty material and to support homogeneous mixing). The combined effect of these two adjustments enables satisfactory levels of reproducibility. Thus, we achieve a fast, reproducible and sustainable solvent-free route to racemization through high energy milling. Further to this, unprecedented information regarding the mechanism of mechanochemical racemization could be gained by following the reaction *in-situ* using X-ray diffraction. Our data shows that the process requires the formation of a disordered phase in which racemization of the substance takes place before a sudden recrystallization of the racemic phase.

Conflicts of interest

There are no conflicts to declare

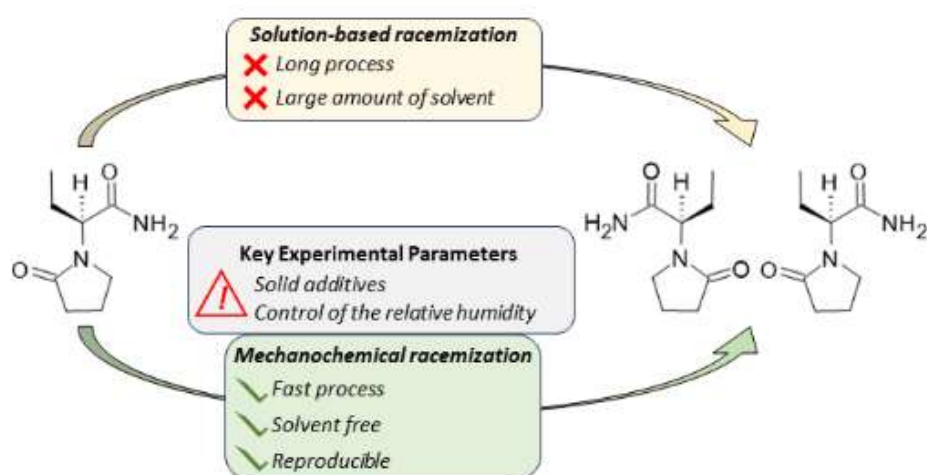
Data availability

The data supporting this article have been included as part of the Supplementary Information.

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