

# Direct Observation of Intermediates in a Thermodynamically Controlled Solid-State Dynamic Covalent Reaction

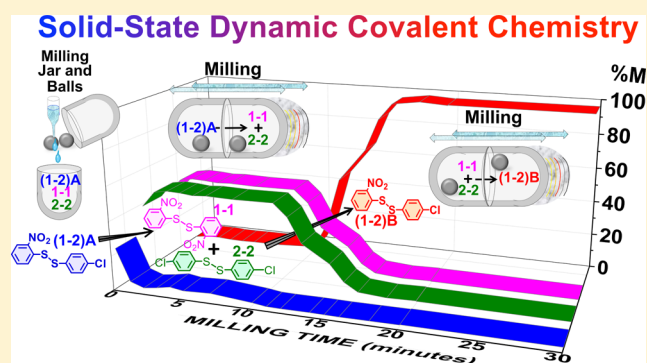
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**S** Supporting Information

**ABSTRACT:** We present the first polymorph interconversion study that uses solid-state dynamic covalent chemistry (DCC). This system exhibits unexpected and rich behavior, including the observation that under appropriate conditions the polymorph interconversion of a heterodimer proceeds through reversible covalent chemistry intermediates, and this route is facilitated by one of the two disulfide homodimers involved in the reaction. Furthermore, we demonstrate experimentally that in all cases a dynamic equilibrium is reached, meaning that changing the milling conditions affects the free energy difference between the two polymorphs and thus their relative stability. We suggest that this effect is due to the surface solvation energy combined with the high surface to volume ratio of the nanocrystalline powder.



## 1. INTRODUCTION

Using both ball mill neat grinding and ball mill liquid assisted grinding (LAG), we demonstrate here the rich and unexpected behavior of a simple dynamic covalent chemistry (DCC) system, focusing on three remarkable observations: (i) the system exhibits reversible interconversion of polymorphs of a disulfide heterodimer under thermodynamic control; (ii) in the presence of catalyst, polymorph interconversion involves sequential covalent reactions with observable covalent intermediates; (iii) one of the intermediates facilitates the covalent chemistry.

We conclude that the outcome of these ball mill grinding reactions is driven by thermodynamics associated with nanoparticulate surface solvation. More importantly, we believe this interpretation to be applicable to many other ball mill grinding reactions, not necessarily involving covalent chemistry. The system under study is illustrated in Figure 1.

Since this work is effectively the first detailed study of DCC in mechanochemical grinding (other than our preliminary report),<sup>2</sup> we first provide some background on these two previously separate fields.

**1.1. Dynamic Covalent Chemistry (DCC).** DCC provides an approach to the discovery of complex architectures and receptors that may be inaccessible or unimaginable by rational design.<sup>3</sup> It also has potential applications in a wide range of chemical and biological problems that involve binding equilibria in chemistry<sup>4</sup> and biology.<sup>3,5</sup> The members of a dynamic combinatorial library (DCL) are molecules that form in a combinatorial way by linking building blocks together through reversible covalent bonds that are dynamic, being continuously broken and re-formed. The concentration of each library member

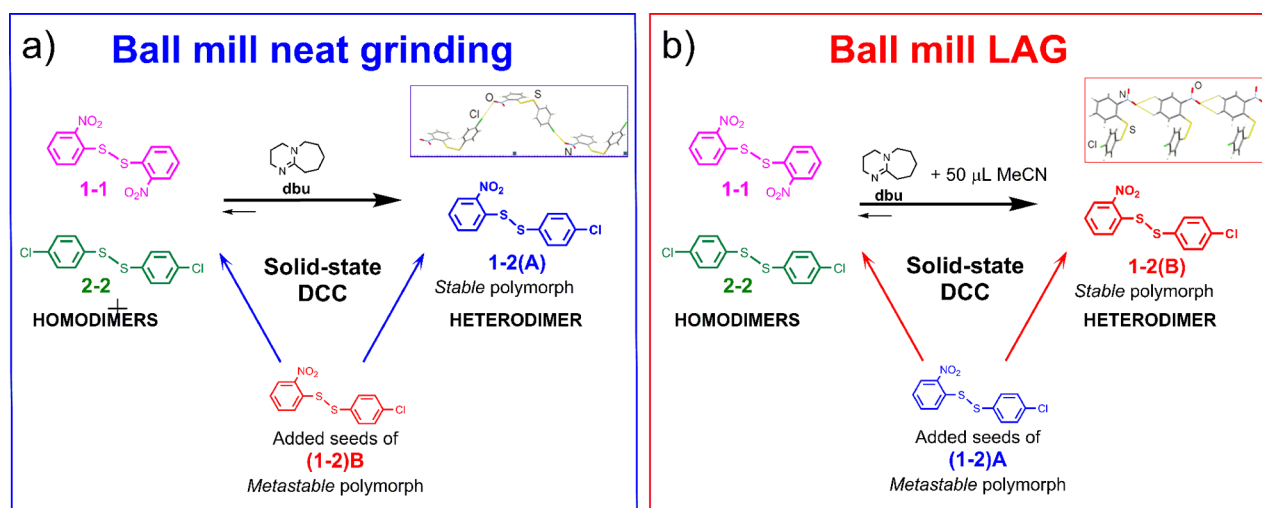
is dictated by its relative free energy; therefore, any phenomenon that can affect the stability of one or more DCL members will affect the concentration of all the library members in the DCL and therefore the overall DCL composition. For example, the addition of a template which selectively binds noncovalently to a specific library member amplifies the amount of this particular member at the expense of other less successful molecules.<sup>3c</sup> The dynamic equilibrium in DCL is easily demonstrated by the establishment of a new equilibrium in response to any perturbing stimulus. Many reversible reactions have been explored to date in DCC,<sup>3,6</sup> including the base-catalyzed disulfide exchange used in this work; adding acid quenches the exchange and freezes the composition for analysis.<sup>2</sup>

In this report, we extend the study of DCC from the traditional dilute solution to the solid state. Not only does this abolish the need for substantial quantities of solvents but also crystal stability<sup>2</sup> can act as a new driving force for establishing the position of the equilibrium.

**1.2. Mechanochemistry Induced by Ball Mill Grinding.** Mechanochemistry<sup>7</sup> is the field of chemical reactions induced by mechanical energy; this can involve reagents in any aggregate state (solid, liquid, or even gas), though typically it refers to fully solid state processes.<sup>8</sup> The term mechanochemistry embraces a broad range of areas, the mechanical energy required to activate the chemical reaction being provided by ultrasound,<sup>9</sup> an atomic force microscope (AFM),<sup>10</sup> manual and mechanical grinding,<sup>11</sup> vortex grinding,<sup>12</sup> or even simple mechanical forces, such as

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**Figure 1.** Solid-state DCC reaction through ball mill grinding: (a) under neat conditions; (b) under LAG conditions. From the solid-state DCC reaction of the homodimers (CCDC<sup>1</sup> codes ODNPS02 and DCPHDS for 1-1 and 2-2, respectively) only the relevant stable polymorph of the heterodimer crystallizes. When seeds of the metastable polymorph of the heterodimer are introduced in the grinding system, two simultaneous and competitive pathways are possible: a direct polymorph interconversion (route to the right) or a sequential transformation from (1-2)A to homodimers to (1-2)B (route to the left) under ball mill neat grinding conditions or vice versa under ball mill LAG conditions. The details of (1-2)A (CCDC code FUQLIM01) and (1-2)B (CCDC code FUQLIM) have been previously reported.<sup>2</sup>

gripping with tweezers or puncture by a needle.<sup>9b</sup> While the mechanochemistry of inorganic solids<sup>13</sup> and metal complexes<sup>14</sup> is a well-established field, developments have been taking place more recently in organic and organometallic<sup>15</sup> chemistry.

Organic reactions explored in manual or mechanical ball mill grinding<sup>16</sup> include the formation of disulfides from thiols,<sup>17</sup> disulfide exchange,<sup>2</sup> imine formation,<sup>8b,16c,18</sup> boronic ester formation,<sup>16c,18a</sup> hydrazone formation,<sup>8b,16c</sup> and a range of C–C bond forming reactions such as carbonyl condensations<sup>8b,16c,19</sup> and pericyclic reactions.<sup>8b,16a,c,20</sup> This has opened the field for the mechanochemical synthesis of complex chemical architectures,<sup>15b</sup> such as metal-coordinated cages,<sup>21</sup> fully organic cages,<sup>18a</sup> rotaxanes,<sup>22</sup> and capsules.<sup>19e</sup>

Much of the mechanochemistry literature has been generated using manual grinding with a pestle and mortar, where the reaction system is open: i.e., the solvent can escape from the system and the supplied mechanical energy is neither constant nor measurable. Here we perform ball mill grinding using a mechanical mixer mill (also called vibratory mill), which allows for reproducible and controlled milling frequencies and times.<sup>11a</sup> The milling jars are closed systems that allow the achievement of thermodynamic equilibrium, as has already been proposed in the field of metal alloys.<sup>23</sup> Some authors consider milling to be a nonequilibrium environment and refer to “pseudo-equilibria” or “equilibrium states of milling”.<sup>11a</sup>

Mechanical ball mill grinding can be performed neat (without added solvent) or solvent assisted (LAG): in the latter, very small amounts of added liquid can dramatically accelerate and even enable mechanochemical reactions between solids.<sup>11b,24</sup> As yet, little is known about the mechanisms and the driving forces involved in the chemical syntheses and supramolecular reactions induced by ball mill grinding. In two recent publications, the formation of organic and metal–organic supramolecular compounds by ball mill grinding under LAG and neat conditions was monitored in situ and in real time by X-ray diffractometry (XRD) at a synchrotron facility.<sup>25</sup> The resulting kinetic reaction curves clearly show a terminal plateau that may be indicative of a

thermodynamic equilibrium: the authors did not comment on the possible driving force of the reactions studied.

Crystallization from solution may be thermodynamically or kinetically driven or be a combination of both: the initial formation of nuclei promotes the exponential growth of a specific polymorph, as indicated by a sigmoidal kinetic curve.<sup>26</sup> In general, the kinetic curves for ball mill grinding reactions are observed to exhibit a sigmoidal shape, this being consistent with the initial formation of nuclei promoting the exponential growth of the product.<sup>25,27</sup>

Ball mill grinding is preceded or accompanied by a crystal size reduction of the starting materials, often down to an amorphous intermediate phase.<sup>24,25,28</sup> Whether this possible amorphous, and therefore nondiffracting, material includes more than one state or any crystal nuclei is not known.<sup>29</sup>

Whether the ball mill grinding process as a whole can be regarded as a purely solid state process is uncertain, even in the neat case, because of the difficulty in controlling or measuring the exact reaction conditions such as average and local pressure and temperature.<sup>13b</sup> Some authors propose that the heat generated in the course of a mechanochemical process can induce local melting of crystals,<sup>8a</sup> melting at the interface between crystals,<sup>30</sup> or formation of liquid eutectic intermediate phases,<sup>11b,13b,28b,31</sup> so that in such cases the reaction takes place in the liquid phase even though a solid product is ultimately produced. We previously investigated and excluded an eutectic-based mechanism for the current system,<sup>2</sup> but a discussion of these mechanistic aspects is beyond the scope of this paper. We will focus here on the relative energy minima of the ball mill grinding process as an approach to interpret our experimental results as a thermodynamic outcome. Indeed, we show below that, after the reaction reaches completion, equilibrium is achieved, with a stable phase composition.<sup>25,32</sup>

Unlike crystallization in solution or a slurry experiment, the ball mill grinding process leads to extremely small crystals whose size is hard to predict or even to estimate. One report of in situ and in real time monitoring by XRD of a ball mill grinding reaction describes the crystal size in a ball mill grinding reaction to

approach the order of tens of nanometers, giving a very different surface to volume ratio ( $S/V$ ) from microcrystals.<sup>25a</sup> Thus, while thermodynamics conventionally assumes surface effects to be negligible (i.e., infinite bulk structures as in the case of slurry experiments), this is not the case in continuously mechanically ground systems.<sup>8a</sup> These thermodynamic aspects are general and must apply to any milling system, independent of the mechanisms involved in the chemical reaction.

Small crystallites have higher enthalpies and free energies than large crystals because of a positive surface energy.<sup>33</sup> Reactions occur not in the whole bulk of the sample but at the interfaces between the phases.<sup>8a</sup> The role of mechanical action is usually to provide mixing, decreasing the particle size, and generating fresh surface for the contact.<sup>8a</sup> In contact with a solvent, as it happens in crystallization experiments in solution or in ball mill grinding under LAG conditions, the particle surfaces are solvated. Solvated surfaces are not simply a sharp boundary but have a finite depth that can extend up to 1 nm.<sup>34</sup> Thus, at high  $S/V$  ratio as in the case of nanocrystals obtained by ball mill grinding, the contribution of the solvated surface to the free energy minimum of a given polymorph is significant. In the case of ZnS nanocrystals (3 nm in diameter) absorption and desorption of methanol were found to reversibly change the atomic arrangement of the bulk structure.<sup>35</sup> Hence, the free energy minima of different polymorphs at the nanoscale will depend on the energetics of the bulk structures, the  $S/V$  ratio of the crystallites, and the extent of solvation of the surface of the crystallites. Therefore, the relative free energy of one polymorph versus another will depend on the solvent used in the case of ball mill LAG or in the lack of solvent in the case of ball mill neat grinding.<sup>36</sup>

**1.3. Solid-State DCC.** The solid-state DCC reaction shown in Figure 1 has been selected to utilize one of the simplest DCC systems available: the metathesis of two linear symmetric disulfide homodimers which can result only in the formation of a linear asymmetric disulfide heterodimer. What we learn from this simple solid-state DCC reaction should then be relevant to more complex oligomeric DCC systems, including other reversible chemistries. In solution, the system leads to a statistical mixture of the two homodimers (25% each) and the heterodimer (50%). Using ball mill grinding, the thermodynamic outcome is dramatically biased toward the heterodimer (97% yield) rather than homodimers (1.5% each).<sup>2</sup> We also demonstrated in earlier work that solid-state DCC can lead to two different polymorphs of the same product, depending on whether the ball mill grinding reaction is performed under neat or under LAG conditions.<sup>2</sup> This is a new aspect of DCC only possible in the solid state.

**1.4. Background to the Present Study.** In our previous paper we demonstrated that the solid-state DCC reaction in Figure 1 was under thermodynamic control. The two homodimers 2-nitrophenyl disulfide [(2-NO<sub>2</sub>PhS)<sub>2</sub>] and 4-chlorophenyl disulfide [(4-ClPhS)<sub>2</sub>] are here referred to as **1-1** and **2-2**, respectively, the corresponding heterodimer is referred to as **1-2**, and the base catalyst is 1,8-diazabicyclo[5.4.0]undec-7-ene (**dbu**).

Two separate tests proved that the constant composition of the product observed on reaching the plateau was thermodynamically determined: (a) the same final composition was obtained regardless of the composition of the starting state, provided that the equimolar stoichiometry of the added homodimers **1-1** and **2-2** was respected; (b) we proved that the plateau was not a kinetic sink, by demonstrating its dynamic nature.<sup>2</sup> The chemical composition of the reaction mixture was monitored using high-performance liquid chromatography (HPLC), and the poly-

morphic form of the quantitative product at equilibrium was determined by powder X-ray diffraction (PXRD). Brief investigations using solid-state NMR and solid-state FTIR did not reveal any useful quantitative information; therefore, we have not included them.

We also reported in our previous work that solid-state DCC can lead to different polymorphs of the heterodimer depending on whether the experiments correspond to ball mill neat grinding conditions, forming a polymorph of **1-2** referred to here as (**1-2**)**A**, or ball mill LAG conditions, forming a different polymorph of **1-2** referred to here as (**1-2**)**B**.<sup>2</sup> It was implied that both polymorphs (**1-2**)**A** and (**1-2**)**B** represent the final equilibrium product of the reaction under ball mill neat and ball mill LAG conditions, respectively, but no proof was provided. Indeed, the very idea that adding one drop of solvent to a milling jar could switch the thermodynamic equilibrium from one polymorph to the other is challenging. Thus, the thermodynamic aspects of these milling systems required the further investigations reported below.

In the present work we have added the solid-state composition obtained by Rietveld refinement of PXRD data collected throughout the grinding process, giving us the opportunity of identifying and quantifying which, and how much, of each polymorph is formed during the ball mill grinding reaction. All samples were also analyzed by HPLC. Performing kinetic studies of seeding experiments provides us with a powerful tool to elucidate the reaction pathway of solid-state DCC reactions by monitoring the solid-state composition as grinding proceeds.

Equipped with these tools, we now explore whether the heterodimer polymorph crystallization is determined by kinetics or thermodynamics, what are the intermediate products, if any, and how mechanochemical crystallization by ball mill grinding proceeds.

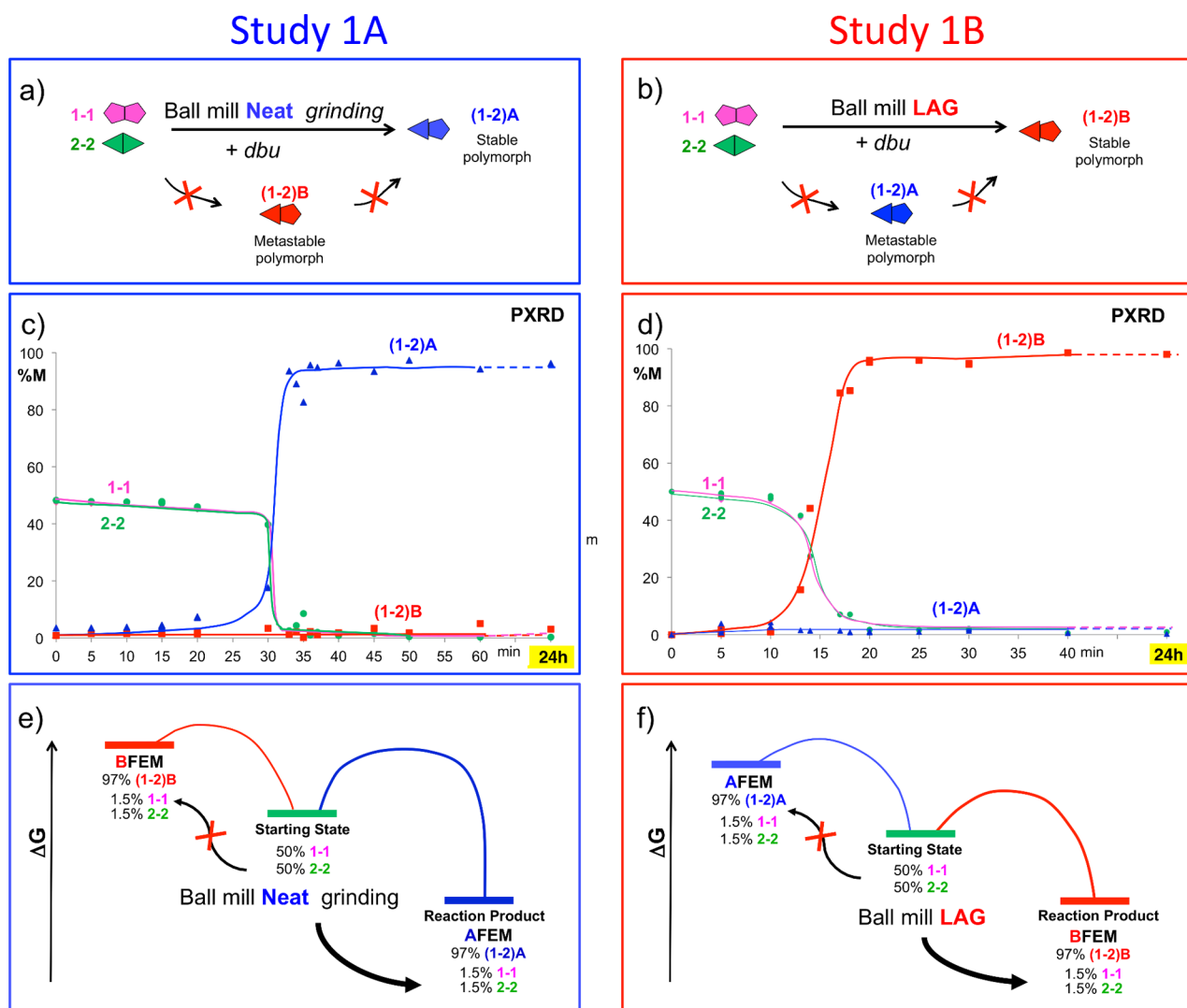
We show here that the experimental milling conditions of the DCC reaction, i.e. the presence or absence of a small quantity of solvent, which allows for solvated surfaces of the nanosized crystals, determine the polymorph selection at equilibrium (see section 3.1): under ball mill neat grinding conditions the DCC reaction exclusively yields polymorph (**1-2**)**A**, which is thus the stable polymorph under these conditions (Figure 1a); when the reaction is carried out under ball mill LAG conditions, (**1-2**)**B** is formed, which is the stable polymorph under ball mill LAG conditions (Figure 1b). If polymorph (**1-2**)**B** is added to the equimolar mixture of the two homodimers under ball mill neat grinding conditions (Figure 1a), then it is fully transformed to (**1-2**)**A**; (**1-2**)**B** is therefore the metastable polymorph under ball mill neat grinding conditions. Similarly, polymorph (**1-2**)**A** is metastable under ball mill LAG conditions and rapidly transforms to (**1-2**)**B** (Figure 1b). Polymorph transformations are crucial in the pharmaceutical industry and generally in any field where the final product is used and commercialized as a solid phase.<sup>37</sup>

## 2. EXPERIMENTAL DETAILS

We present here 14 kinetic studies, which were designed to explore the different kinetic and thermodynamic features of our solid-state DCC reaction, both under ball mill neat grinding conditions (i.e., in the absence of solvent) in studies A1–A7 and under the corresponding ball mill LAG conditions (i.e., with 50  $\mu$ L of acetonitrile added to 200 mg of powder in the grinding jar) in studies B1–B7.

The kinetic curves presented here were prepared from PXRD or HPLC data obtained from individual experiments, each experiment corresponding to a single grinding time. This approach avoids disrupting the delicate equilibrium achieved during grinding among the vapor, the





**Figure 2.** Solid-state DCC studies reacting 1-1 and 2-2 in an equimolar ratio (no added seeds of 1-2) in the presence of catalyst (**dbu**): (a, b) reaction scheme; (c, d) kinetic curves prepared as obtained from the Rietveld quantitative phase analysis; (e, f) free energy diagrams. The parts framed in blue on the left represent ball mill neat grinding conditions. The parts framed in red on the right represent ball mill LAG conditions using 50  $\mu\text{L}$  of MeCN.

liquid, and the solid components in the enclosed chamber of the snap-closed grinding jar.

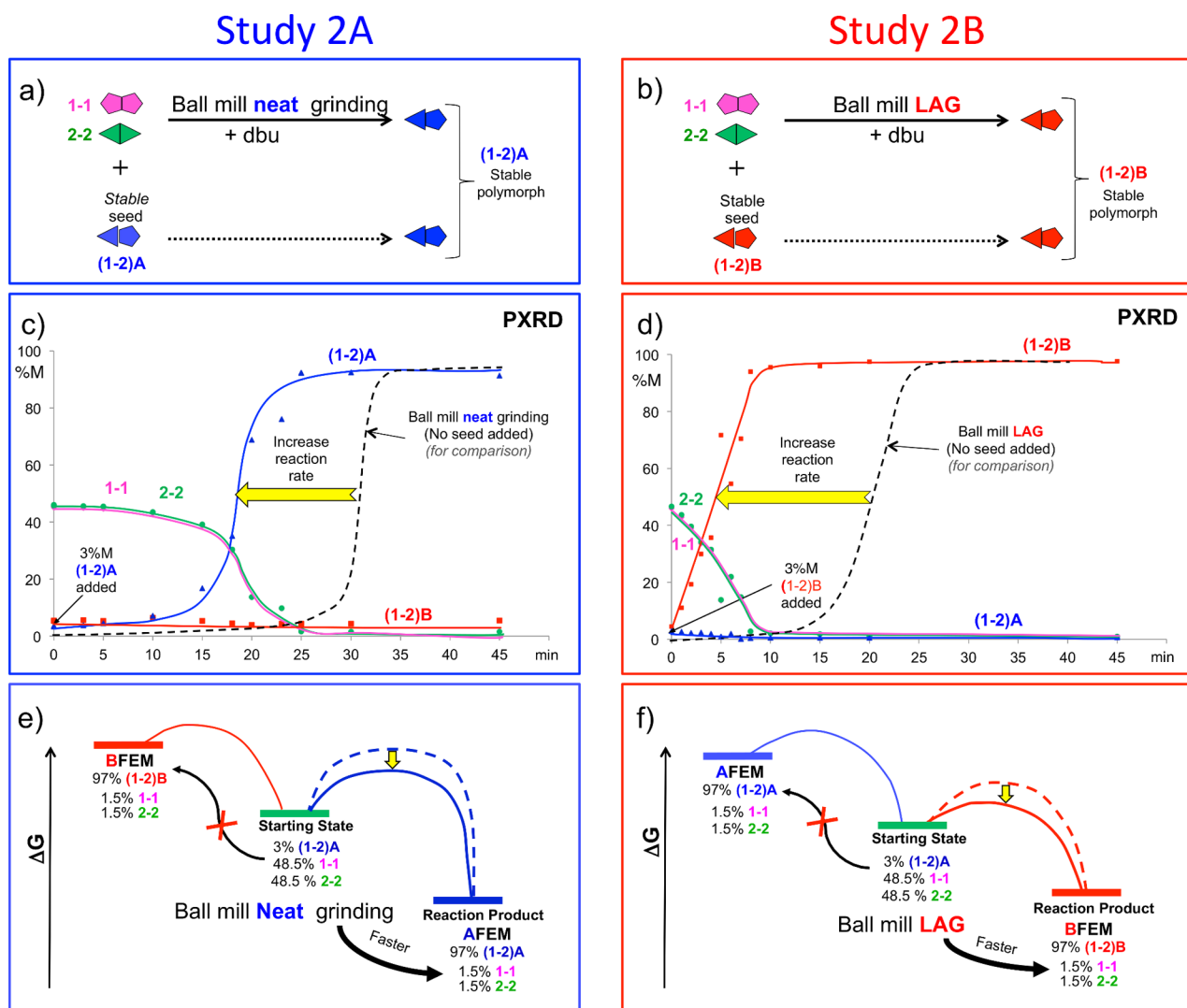
Samples of the components of the homodimers 1-1 and 2-2 and the added seeds of the polymorph of 1-2, as required, were accurately weighed, resulting in a typical loading of 200 mg. The solid powders were individually placed in a 14.5 mL stainless steel grinding jar together with two 7 mm i.d. stainless steel ball bearings. After the addition of 2  $\mu\text{L}$  (2% M) of catalyst (**dbu**) into the jar, nothing else was added for ball mill neat grinding, while 50  $\mu\text{L}$  of acetonitrile was added for ball mill LAG experiments. The jars were snap-closed, and grinding was performed at 30 Hz on a MM400 Retsch automated grinder (see Figures SI 39 and SI 40 in the Supporting Information) for the specified period of time. After completion of the grinding period, the jar was immediately opened and the contents were analyzed; the results obtained as % M concentration versus grinding time were used to construct the corresponding kinetic profiles.

The number of grinding experiments and grinding times for a given study depended on how many points were required to represent, with good definition, the sigmoidal segment of the kinetic curve and ascertain that the system had finally reached a plateau. Indeed, the rigorous experimental procedures detailed in the Supporting Information were found to be crucial for reproducibility. The solid-state composition of the samples, reported here as % M, was determined by Rietveld refinements from PXRD data. The chemical composition of the sample was obtained

by HPLC analysis. While the PXRD analysis is not as sensitive or accurate (estimated accuracy  $\pm 3\%$  M absolute and estimated sensitivity, limit of detection (LOD) as 3% M) as HPLC (estimated sensitivity 0.1% M relative to the main component), it supplies the phase composition. Therefore, all further discussion regarding kinetic profiles will be based on the Rietveld refinements of the PXRD data. The agreement between PXRD and HPLC analysis was found to be excellent (see the Supporting Information). More details about the analytical methods can be found in the Supporting Information.

### 3. RESULTS AND DISCUSSION

To help the reader, kinetic curves for ball mill neat grinding studies are shown framed in blue, while ball mill LAG experiments are shown framed in red for Figures 2–7. No fitting was performed, and the kinetic curves are only a guide to the eye. Each time point in these kinetic plots corresponds to a single grinding experiment. The findings of these studies are summarized at the top of Figures 2–7 in the form of reaction schemes for each kinetic study. To appreciate the thermodynamic aspects and to recognize the relative predominance of the competing kinetic pathways, we summarize these relative free energy minimum (FEM) states for the starting state, intermediate



**Figure 3.** Solid-state DCC studies reacting 1-1 and 2-2 in an equimolar ratio seeded with 3% M of the stable polymorph of 1-2 in the presence of catalyst (**dbu**): (a, b) reaction scheme; (c, d) kinetic curves prepared as obtained from the Rietveld quantitative phase analysis; (e, f) free energy diagrams. The parts framed in blue on the left represent ball mill neat grinding conditions. The parts framed in red on the right represent ball mill LAG conditions using 50  $\mu\text{L}$  of MeCN.

state, and reaction product at the bottom of each figure in the form of a free energy level diagram. These levels and the corresponding energy barriers between the starting state and intermediate state and reaction product have been inferred from our kinetic studies. Analysis of these kinetic data has allowed us to distinguish alternative reaction pathways, preferred as well as unlikely pathways, for each grinding study.

**3.1. Unseeded Solid-State DCC Experiments.** We will start this discussion with unseeded grinding studies. These solid-state DCC experiments were performed by grinding equimolar amounts of the homodimers, 1-1 and 2-2, in the presence of catalyst (2% M **dbu**). Figure 2 shows the findings of the ball mill neat grinding experiments (study 1A) on the left and ball mill LAG experiments with 50  $\mu\text{L}$  of MeCN (study 1B) on the right.

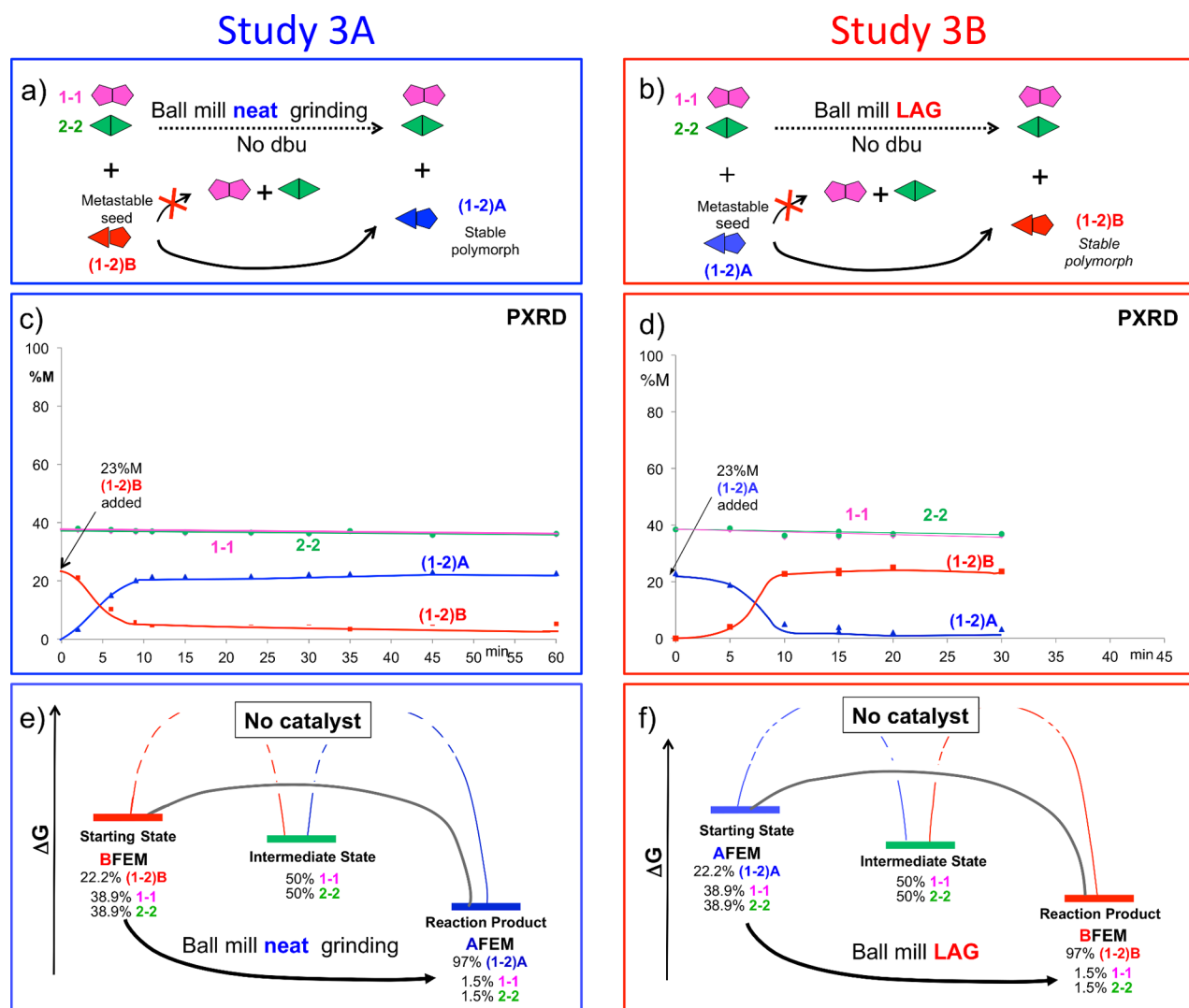
The solid-state DCC reaction reached equilibrium, always resulting in quantitative formation of the stable polymorph of 1-2, its composition being maintained for extended grinding periods (i.e., 24 h). The metastable polymorph of 1-2 was never observed at any time point of the grinding process. These unseeded reactions exhibited a long delay before a significant concentration (7% M) of the stable polymorph of 1-2 was observed: 25 min for

ball mill neat grinding studies (Figure 2c) and 10 min for ball mill LAG studies (Figure 2d), showing that the ball mill LAG process is faster than the ball mill neat process. It is known that very small amounts of added liquid can dramatically accelerate mechanochemical reactions between solids. In other words, given a mechanochemical reaction, ball mill LAG is generally faster than ball mill neat grinding.<sup>11b</sup> However, in our case ball mill LAG and ball mill neat grinding lead to two different polymorphs, (1-2)B and (1-2)A, respectively; therefore, their reaction rates cannot be directly compared.<sup>11b,38</sup>

Both ball mill neat grinding and ball mill LAG reactions exhibited a nucleation phase (around 5 min) and a sharp transition before reaching a constant plateau.

Any change in the phase composition affects the position of the free energy minima (FEM) levels and their corresponding energy barriers. For brevity, we designate **AFEM** as the FEM containing (1-2)A and **BFEM** as the FEM containing (1-2)B. The composition of the FEM level of the starting state for studies 1A and 1B was equimolar amounts of 1-1 and 2-2.

Under ball mill neat grinding conditions (Figure 2e) the equilibrium is represented by the **AFEM** containing (1-2)A,



**Figure 4.** Solid-state DCC studies reacting 1-1 and 2-2 in an equimolar ratio seeded with 23% M of the metastable polymorph of 1-2, in the absence of catalyst (dbu): (a, b) reaction scheme; (c, d) kinetic curves prepared as obtained from the Rietveld quantitative phase analysis; (e, f) free energy diagrams. The parts framed in blue on the left represent ball mill neat grinding conditions. The parts framed in red on the right represent ball mill LAG conditions using 50  $\mu$ L of MeCN.

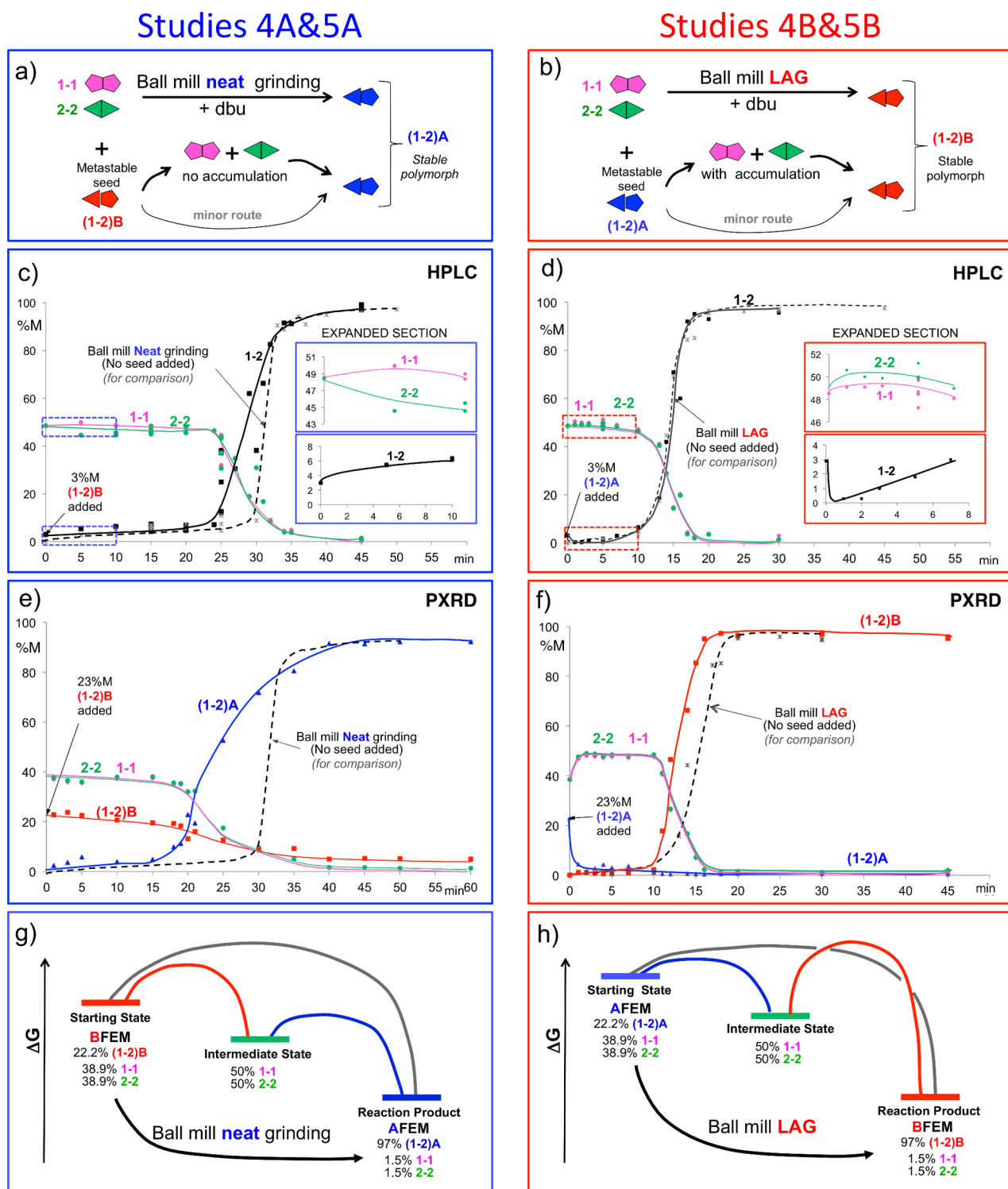
which is the stable polymorph and the only polymorph of 1-2 observed in the ball mill neat grinding reaction. Under ball mill LAG conditions (Figure 2f) the equilibrium corresponds to BFEM containing (1-2)B, which is the stable polymorph and the only polymorph of 1-2 observed in the ball mill LAG reaction. As the kinetic curves for ball mill neat grinding (Figure 2c) are slower than those for ball mill LAG (Figure 2d), the energy barrier between the starting state and the reaction product (AFEM) for ball mill neat grinding (Figure 2e) must be higher than that between the starting state and the reaction product (BFEM) for ball mill LAG (Figure 2f).

**3.2. Solid-State DCC Reaction Seeded with the Stable Polymorph of the Heterodimer.** Figure 3 displays the results from grinding experiments seeded with 3% M of the stable polymorph of 1-2 to an equimolar amount of 1-1 and 2-2 in the presence of catalyst (2% M dbu). The ball mill neat grinding experiments (study 2A) and the ball mill LAG experiments with 50  $\mu$ L of MeCN (study 2B) are on the left and right sides of the figure, respectively. These studies showed a decrease in the lag time by 10–12 min in comparison to the times for the unseeded studies, maintaining the shape of the sigmoidal segment, as

shown in Figure 3c for ball mill neat grinding studies and in Figure 3d for ball mill LAG studies.

The Rietveld refinement of PXRD data showed that only the stable polymorph was formed under these experimental conditions, but the reaction rate was greater than that in the corresponding unseeded studies. The interpretation of these data in the form of free energy level diagrams is shown in Figure 3e for ball mill neat grinding studies and Figure 3f for ball mill LAG studies: the increase in the reaction rate must be predominantly a consequence of a significant reduction of the energy barriers between the starting state and the reaction product.

**3.3. Direct Polymorph Interconversion in the Absence of Catalyst.** Direct polymorph interconversion from the metastable to the stable polymorph requires only a supramolecular rearrangement of the crystal lattices of (1-2)B  $\rightarrow$  (1-2)A for ball mill neat grinding studies and (1-2)A  $\rightarrow$  (1-2)B for ball mill LAG studies. Such direct transformation in milling experiments has been reported before in other systems.<sup>39</sup> We explored these direct polymorph interconversions in study 3-A under ball mill neat grinding conditions and in study 3-B under ball mill LAG conditions; their outcomes are shown on the left

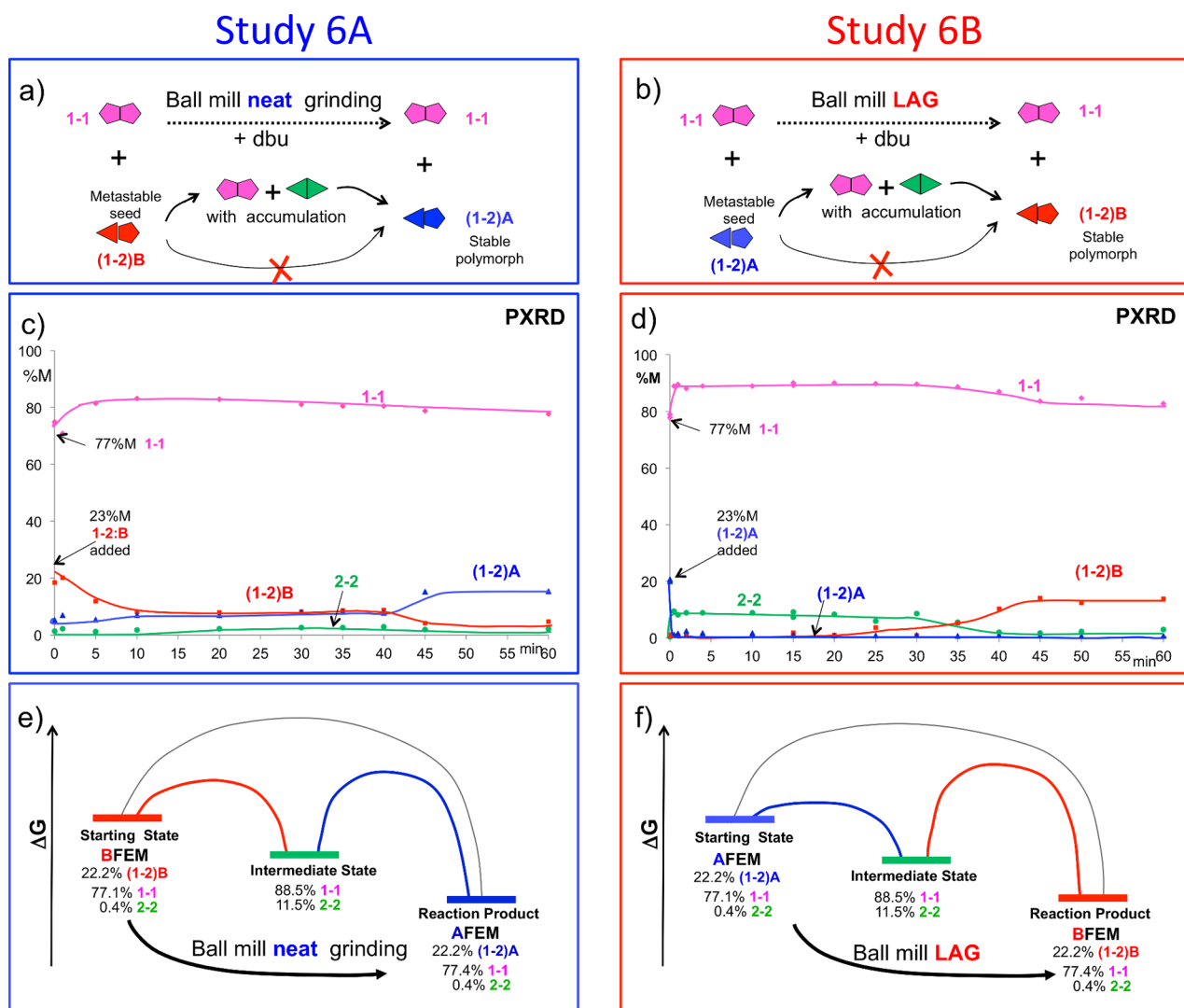


**Figure 5.** Solid-state DCC studies reacting 1-1 and 2-2 in an equimolar ratio seeded with the metastable polymorph of 1-2 in the presence of catalyst (dbu): (a, b) reaction scheme; (c–f) kinetic curves prepared as obtained from the Rietveld quantitative phase analysis; (g, h) free energy diagrams. The parts framed in blue on the left represent ball mill neat grinding conditions (3% M of seeds shown in (c)); 23% M of seeds shown in (e)). The parts framed in red on the right represent ball mill LAG conditions using 50  $\mu\text{L}$  of MeCN (3% M of seeds shown in (d); 23% M of seeds shown in (f)).

and right sides of Figure 4, respectively. In these experiments 23% M of the metastable polymorph of 1-2 was added to 77% M of equimolar amounts of 1-1 and 2-2 homodimers in the absence of dbu to prevent any DCC reaction from taking place.

As expected, the 77% M concentration of equimolar amounts of 1-1 and 2-2 remained unreacted. Under ball mill neat grinding conditions, the 23% M concentration of the metastable

polymorph (1-2)B was fully transformed into the stable polymorph (1-2)A in around 10 min, the kinetic curve presenting a shallow sigmoidal kinetic segment and the conversion starting soon after grinding was initiated (Figure 4c). Similar kinetic curves were observed under ball mill LAG conditions, where (1-2)A transformed into (1-2)B in around 10 min as well (Figure 4d). In summary, direct polymorph interconversion rapidly



**Figure 6.** Solvent-free DCC studies reacting 77% 1-1 with 23% M of the metastable polymorph of 1-2, in the presence of catalyst (*dbu*): (a, b) reaction scheme; (c, d) kinetic curves prepared as obtained from the Rietveld quantitative phase analysis; (e, f) free energy diagrams. The parts framed in blue on the left represent ball mill neat grinding conditions. The parts framed in red on the right represent ball mill LAG conditions using 50  $\mu\text{L}$  of MeCN.

transformed the metastable to the stable polymorph in very little time.

**3.4. Solid-State DCC Reaction Seeded with the Metastable Polymorph of the Heterodimer.** Unexpected phenomena were observed in studies 4 and 5, where seeds of the metastable polymorph of 1-2 were added to equimolar amounts of 1-1 and 2-2 in the presence of *dbu* catalyst (2% M). These experiments were initially performed at 3% M concentration of the added seeds (Figure 5c,d, study 4), the composition of which could only be analyzed by HPLC, 3% M being the limit of detection of the PXRD method. A higher concentration of seeds (23% M, Figure 5e,f, study 5) was required to monitor the solid-state composition by PXRD.

**3.4.1. Ball Mill LAG.** The kinetic curves for the ball mill LAG case in study SB illustrated in Figure 5f present an unexpected observation.

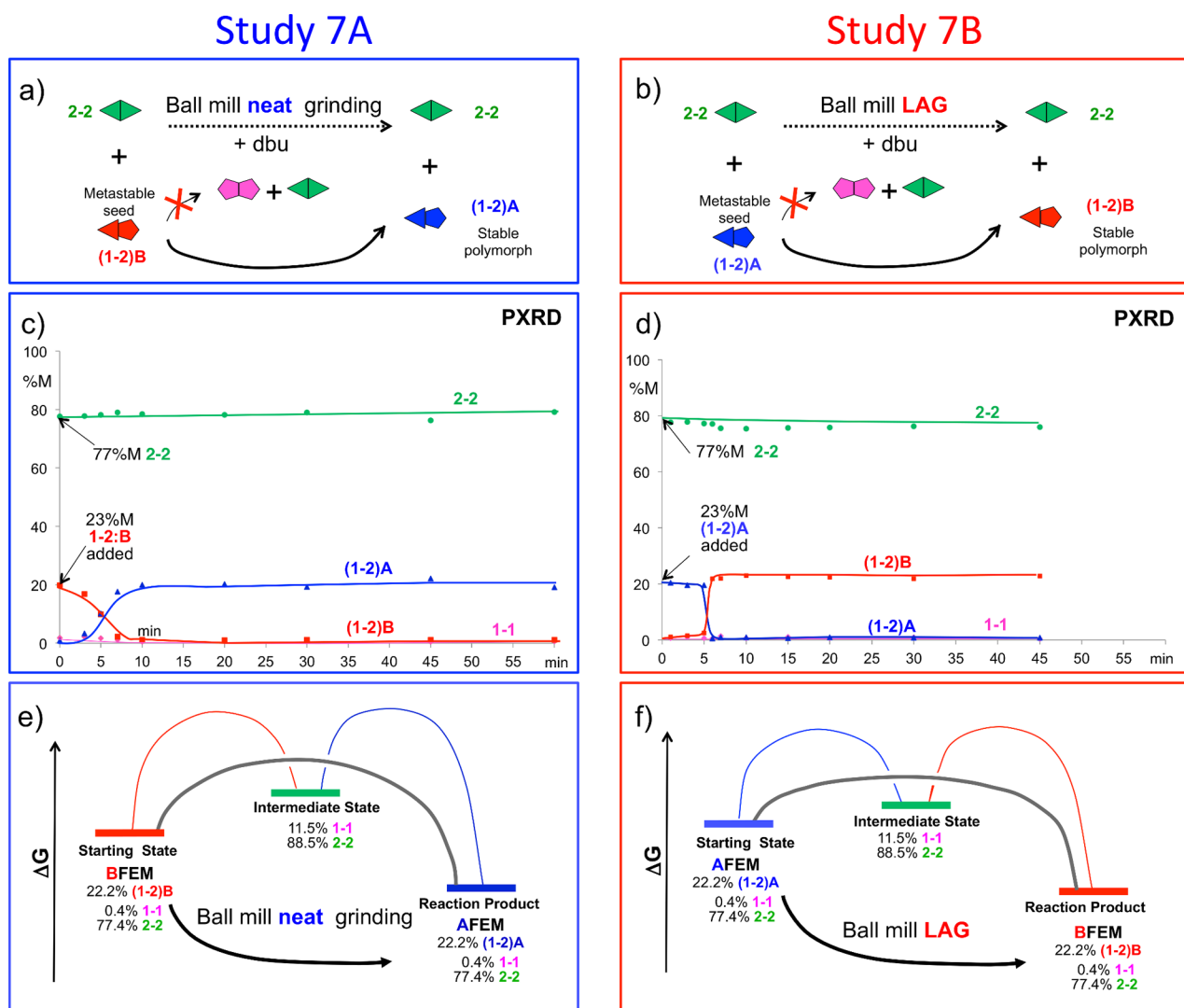
The rapid disappearance of (1-2)A from the initial concentration of 23% M down to 3–5% M within the first 1 min of ball mill grinding was accompanied by a simultaneous increase of the concentration of 1-1 and 2-2 from the initial concentration of 37.5% M up to 47–48% M. In other words, the reduced concentration of (1-2)A exactly matched the corre-

sponding increase in the concentration of 1-1 and 2-2. After the first 1 min and for the next 9 min, the concentration of 1-1 and 2-2 stayed constant at 48% M while the concentration of (1-2)A was around 3–5% M. The freshly formed 1-1 and 2-2 were therefore the covalent intermediates in the first step of the reaction ((1-2)A  $\rightarrow$  1-1 + 2-2) of a sequential pathway, where (1-2)A is transformed to (1-2)B. This process involves bond breaking and bond forming of the reversible disulfide bond. This is the first time we are aware of that a purely covalent reaction has been reported to be involved in a polymorph interconversion.<sup>40</sup>

The interpretation of this sequential mechanism in the form of free energy level diagrams is shown in Figure 5h. Since we observed an accumulation of the covalent intermediates for over 10 min, the second step of the sequential mechanism (1-1 + 2-2  $\rightarrow$  (1-2)B) must be the rate-determining step (RDS).

The kinetics in study SB were found to be significantly faster (3–4 min) in comparison to those for the corresponding unseeded experiments (study 1B). This result seems counterintuitive. If the two-step sequential transformation were to be the only operative pathway in study SB, it should have resulted in a delay of the kinetics. This slight increase in the reaction rate with respect to that of study 1B could indicate that an alternative minor





**Figure 7.** Solvent-free DCC studies reacting 77% M 2-2 with 23% M of the metastable polymorph of 1-2, in the presence of catalyst (*dbu*): (a, b) reaction scheme; (c, d) kinetic curves prepared as obtained from the Rietveld quantitative phase analysis; (e, f) free energy diagrams. The parts framed in blue on the left represent ball mill neat grinding conditions. The parts framed in red on the right represent ball mill LAG conditions using 50  $\mu$ L of MeCN.

but faster pathway (the direct polymorph interconversion) is acting competitively and simultaneously in study 5B to the sequential transformation as depicted in Figure 5h. We suggest that the residual fraction of (1-2)A not immediately consumed by the sequential pathway was transformed into (1-2)B within a few minutes by the direct interconversion pathway, these freshly created seeds of (1-2)B acting as a template and thus increasing the DCC reaction rate.

**3.4.2. Ball Mill Neat Grinding.** Unlike the case under ball mill LAG conditions in study 5B (Figure 5f), the kinetic curves of study 5A under ball mill neat grinding conditions (Figure 5e) showed no signs of accumulation of covalent intermediates. Despite this observation, we propose that the transformation of the metastable (1-2)B to the stable polymorph (1-2)A under ball mill neat grinding conditions also proceeds through two competitive and simultaneous pathways, the sequential mechanism being the predominant path (Figure 5g).

In the ball mill neat grinding case the sequential path has to be inferred from the reaction rate and the kinetic curve in Figure 5e, which shows a shallow sigmoidal shape. This shallow curve is different from the sharp sigmoidal segments for the corresponding unseeded experiments (Figure 2c, study 1A) or the

corresponding studies seeded with the stable polymorph (Figure 3c, study 2A). Had the direct polymorph interconversion been the predominant or exclusive pathway, we should have seen a sharp nucleation phase as in studies 1A and 2A.

The delay in the polymorph interconversion and the shallow kinetic curve can be explained by the transformation happening competitively and simultaneously through a direct polymorph interconversion and a sequential mechanism, the latter being the predominant one. Since there is no accumulation of the covalent intermediate, the first step of the sequential process ((1-2)B  $\rightarrow$  1-1 + 2-2) must be the RDS: as soon as the covalent intermediates 1-1 and 2-2 are formed in the RDS step 1, they are immediately consumed in the lower energy barrier step 2 to form (1-2)A.

**3.5. Effect of the Nature of the Homodimers on the Competition between the Sequential and the Direct Polymorph Transformation.** We were interested in understanding if the presence of the two homodimers, 1-1 and 2-2, had any effect on the polymorph interconversion reaction. We therefore designed kinetic studies so that 23% M seeds of the metastable polymorph of 1-2 were added to either 77% M of 1-1

only (Figure 6, study 6) or to 77% M of 2-2 only (Figure 7, study 7) in the presence of catalyst.

Adding metastable polymorph seeds just to 1-1 resulted in very similar kinetic curves for ball mill neat grinding (Figure 6c, study 6A) and ball mill LAG studies (Figure 6d, study 6B). Both graphs have the tell-tale signs of the sequential covalent route: immediate reduction of the metastable polymorph of 1-2 with simultaneous accumulation of 1-1 and 2-2 (Figure 6a,b). The accumulation time before conversion to the stable polymorph of 1-2 product molecule was now much longer in comparison to that of study 5B. This result indicates that the direct polymorph interconversion does not occur to any significant extent in study 6A-B, and the two-step reaction thus takes longer than an unseeded experiment. The free energy level interpretation of study 6 is consistent with a significant stabilization of energy level of this intermediate state with respect to the starting state, both under ball mill neat grinding (Figure 6e) and under ball mill LAG (Figure 6f); this resulted in a lowering of the energy barriers for the sequential mechanism, while the energy barrier for the direct conversion was increased. Under these experimental conditions, the sequential mechanism was predominant or exclusive, the second step of the sequential mechanism being the RDS, with relative accumulation of 1-1 and 2-2.

In contrast, when the same seeds of the metastable polymorph of 1-2 were added to 77% M of pure 2-2 (Figure 7), the direct transformation pathway was predominant or exclusive for both ball mill neat grinding (Figure 7c, study 7A) and ball mill LAG (Figure 7d, study 7B) and showed the same kinetic profile as those in study 3A (Figure 4c) and study 3B (Figure 4d) performed in the absence of *dbu*, where only the direct polymorph transformation could take place. The free energy level interpretation of study 7 is consistent with a significant destabilization of energy level of this intermediate state with respect to the starting state, both under ball mill neat grinding (Figure 7e) and under ball mill LAG conditions (Figure 7f); this resulted in an increase of the energy barriers for the sequential mechanism, while the energy barrier for the direct conversion was decreased. Under these experimental conditions, the direct mechanism was predominant or exclusive.

“Seeding assisted” polymorphic transformation was recently reported for the covalent synthesis of imines by manual grinding, where the addition of seeds of a specific polymorph directed the selection of the polymorph formed.<sup>41</sup> In contrast, for the solid-state DCC system studied here, we have demonstrated that while seeding with the stable polymorph leads to an acceleration of the reaction (Figure 3), “seeding assisted” polymorphic transformation did not take place: i.e., seeding with the metastable polymorph resulted in its complete transformation to the stable polymorph (Figures 4–7).

#### 4. CONCLUSIONS

We have provided extensive experimental evidence that only the more stable of the two polymorphs of the heterodimer is exclusively formed in almost quantitative yield in our solid-state dynamic covalent chemistry model reactions. No further change in the equilibrium composition was observed after up to 24 h of continuous grinding. The experimental conditions, especially the absence of solvent for ball mill neat grinding or the presence of a few drops of acetonitrile for ball mill LAG, determines which polymorph of the product is formed on grinding. Under ball mill neat grinding conditions (1-2)A is the stable polymorph and (1-2)B is the metastable polymorph. Under ball mill LAG conditions (1-2)B is the stable polymorph and (1-2)A is the

metastable polymorph. Therefore, we have experimentally demonstrated that the identity of the stable and metastable polymorphs is reversed under ball mill neat grinding and ball mill LAG conditions. Predictably, the rate of reaction was increased when the material was seeded with the stable polymorph. The addition of seeds of the metastable polymorph leads to a rich and unexpected reaction path affecting the kinetics without changing the final phase composition at equilibrium: the metastable polymorph is always converted into the more stable form. In the absence of catalyst the seeds of the metastable polymorph were transformed into the more stable polymorph exclusively by direct polymorph interconversion, as no DCC chemical reaction can take place. In the presence of catalyst, this polymorph interconversion occurred by two simultaneous and competitive routes: a direct transformation and a sequential route, which is predominant. In the latter route, the first step is the conversion of the metastable polymorph of the heterodimer into its corresponding homodimers, which thus are the covalent intermediates in this reversible reaction. This is the first time that a purely covalent bond breaking and re-forming has been reported in the literature to be involved in a polymorph interconversion. Under ball mill LAG conditions the RDS is the second step, while under ball mill neat grinding conditions the RDS is the first step of the sequential route. Furthermore, we have proved that the presence of 2-2 homodimer promotes direct polymorph interconversion of the heterodimer, while the presence of 1-1 homodimer promotes the two-step sequential polymorph transformation, which exploits the reversible covalent disulfide bond. This work may be seen as an important step toward an understanding of the mechanisms and potential of solid-state DCC.

The milling process reduces the crystallite size: we believe that the nanosized dimensions of the crystallites and/or the surface solvation free energy are key to understanding why the polymorph stability order is different in the presence or absence of the acetonitrile solvent. More importantly, this interpretation of the system may be general and may apply to many other grinding processes.

#### ■ ASSOCIATED CONTENT

##### Supporting Information

Text, tables, and figures giving detailed procedures for the solid-state DCC reaction, HPLC data, PXRD patterns, data, and their corresponding Rietveld refinement for each of the 14 studies, and a discussion of the methodology used for the Rietveld refinement. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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##### Notes

The authors declare no competing financial interest.

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