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Stepwise dissolution and composition determination of samples of multiple crystals using a dissolution medium containing aqueous alcohol and fluorocarbon phases.

Humphrey A. Moynihan and Declan Armstrong

SUPPLEMENTARY INFORMATION

Contents

Synthetic Procedures

Synthesis of 2-nitro-4-tert-butylacetanilide (3): page 3

Fig. S1. Chemical structure of 2-nitro-4-t-butylacetanilide (3): page 3

NMR Spectra

Fig. S2. ¹H NMR spectrum of 3: page 4

Fig. S3. ¹³C NMR spectrum of 3: page 4

Fig. S4. ¹⁹F NMR spectrum of 4: page 5

Fig. S5. ¹H NMR spectrum of 4: page 5

Fig. S6. ¹⁹F NMR spectra of 4: page 6

HPLC Calibration Data: page 7

Particle Sizing: page 11

Fig. S7. An image highlighting the typical perimeter established for area measurements and the typical dimension established for length measurements.

Impurity incorporation: page 12

Fig. S8. Graph displaying incorporation level versus solution impurity level of 4-methyl-2nitroacetanilide (2) in "host" crystals of 4-trifluoromethyl-2-nitroacetanilide.

Table S1. Overall incorporation (% composition by HPLC) of 4-methyl-2-nitroacetanilide (2) and 4-tertbutyl-2-nitroacetanilide (3) in crystals of 4-chloro-2-nitroacetanilide (1) obtained by crystallisation from solutions in toluene containing various quantities of 2 and 3 at a σ value of 1.5. Data is shown for two crystallisation batches.

Powder X-ray Diffraction: page 14

Fig. S9. Powder X-ray diffraction patterns obtained for crystals of 1 grown from toluene, 1 grown from toluene with concentrations of 5 mol% 2, and 2 grown from toluene: page 14

Fig. S10. Powder X-ray diffraction patterns obtained for crystals of 1 grown from toluene, 1 grown from toluene with concentrations of 4.5 mol % 3, and 3 grown from toluene: page 15

Differential Scanning Calorimetry: page 16

Fig. S11. DSC curves obtained from crystals of 1 containing 5 mol % 2; compared to DSC curves of component compound 1 and compound 2: page 16

Fig. S12. DSC curves obtained from crystals of 1 containing 4.5 mol % 3; compared to DSC curves of component compound 1 and compound 3: page 16

Partial Dissolution Graphs: page 17

- **Fig. S13.** Chart comparing particle area versus the ranking of each particle in a partial dissolution series of **1** doped with 1.5 mol % of **2**: page 17
- **Fig. S14.** Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 1.5 mol % of **2**: page 17
- **Fig. S15.** Plot of percentage by HPLC of added impurity in crystals of compound **1** *vs.* the dissolution mid-point for the sample of crystals grown from solutions containing 1.5 mol % of additive **2**: page 17 **Fig. S16.** Chart comparing particle area versus the ranking of each particle in a partial dissolution series
- of 1 doped with 2.0 mol % of 2: page 18
- **Fig. S17.** Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 2.0 mol % of **2**: page 19
- **Fig. S18.** Plot of percentage by HPLC of added impurity in crystals of compound **1** *vs.* the dissolution mid-point for the sample of crystals grown from solutions containing 2.0 mol % of additive **2**: page 19 **Fig. S19.** Chart comparing particle area versus the ranking of each particle in a partial dissolution series of **1** doped with 2.5 mol % of **2**: page 20
- **Fig. S20.** Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 2.5 mol % of **2**: page 20
- **Fig. S21.** Plot of percentage by HPLC of added impurity in crystals of compound **1** vs. the dissolution mid-point for the sample of crystals grown from solutions containing 2.5 mol % of additive **2**: page 21 **Fig. S22.** Chart comparing particle area versus the ranking of each particle in a partial dissolution series
- of **1** doped with 3.0 mol % of **2**: page 21
- **Fig. S23.** Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 3.0 mol % of **2**: page 22
- **Fig. S24.** Plot of percentage by HPLC of added impurity in crystals of compound **1** vs. the dissolution mid-point for the sample of crystals grown from solutions containing 3.0 mol % of additive **2**: page 22
- **Fig. S25.** Chart comparing particle area versus the ranking of each particle in a partial dissolution series of **1** doped with 3.5 mol % of **2**: page 23
- **Fig. S26.** Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 3.5 mol % of **2**: page 23
- **Fig. S27.** Plot of percentage by HPLC of added impurity in crystals of compound **1** vs. the dissolution mid-point for the sample of crystals grown from solutions containing 3.5 mol % of additive **2**: page 24
- **Fig. S28.** Chart comparing particle area versus the ranking of each particle in a partial dissolution series of 1 doped with 4.0 mol % of 2: page 24
- **Fig. S29.** Chart comparing particle length versus the ranking of each particle in a partial dissolution series of 1 doped with 4.0 mol % of 2: page 25
- **Fig. S30.** Plot of percentage by HPLC of added impurity in crystals of compound **1** *vs.* the dissolution mid-point for the sample of crystals grown from solutions containing 4.0 mol % of additive **2**: page 25
- **Fig. S21.** Chart comparing particle area versus the ranking of each particle in a partial dissolution series of **1** doped with 4.5 mol % of **2**: page 26
- **Fig. S32.** Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 4.5 mol % of **2**: page 26
- **Fig. S33.** Plot of percentage by HPLC of added impurity in crystals of compound **1** vs. the dissolution mid-point for the sample of crystals grown from solutions containing 4.5 mol % of additive **2**: page 27 **Fig. S34.** Chart comparing particle area versus the ranking of each particle in a partial dissolution series
- of **1** doped with 5.0 mol % of **2**: page 27
- **Fig. S35.** Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 5.0 mol % of **2**: page 28
- **Fig. S36.** Plot of percentage by HPLC of added impurity in crystals of compound **1** *vs.* the dissolution mid-point for the sample of crystals grown from solutions containing 5.0 mol % of additive **2**: page 28
- **Table S2.** Comparisons of parent batch average and weighted averages from stepwise dissolutions: page 29.

Fig. S37 Comparisons of parent batch average and weighted averages from stepwise dissolutions: page 29.

Table S3. Dissolution mid-points calculated for analysed 4.0 mol % **2**-doped **1** crystals from both observed measurements and 'theoretical' values: page 29.

References: page 30

Synthetic Procedures

Synthesis of 2-nitro-4-tert-butylacetanilide (3)

Fig. S1 Chemical structure of 2-Nitro-4-t-butylacetanilide (3).

2-Nitro-4-*tert*-butylaniline (3.825 g, 19.693 mmol, 1 eq.), acetic anhydride (3.03 g, 2.8 mL, 29.676 mmol, 1.5 eq.) and two drops of sulphuric acid were stirred together at room temperature to form an orange suspension. The mixture was heated, at approximately 55 °C the mixture solidified but eventually formed an orange coloured solution on continued heating towards 90 °C, at which point it was heated at 90 °C for two hours. After two hours, the reaction was allowed to cool to room temperature and then 30 mL of water was added to precipitate the product. The precipitate was isolated by filtration and washed with two 50 mL portions of water and air dried. The crude product was then recrystallized from 45 mL of hot ethanol, isolated by filtration and washed with two 20 mL portions of ice-cold ethanol. Yield: 2.913 g, 63.14 %. M.P. (DSC) 107.68 °C.

¹H NMR (300 MHz, CDCl₃): δ 10.18 (s, 1H, NH), 8.63 (d, ${}^{3}J_{HH}$ = 8.9 Hz, 1H, H-6), 8.17 (d, ${}^{4}J_{HH}$ = 2.4 Hz, 1H, 3-H), 7.67 (dd, ${}^{3}J_{HH}$ = 8.9 Hz, ${}^{4}J_{HH}$ = 2.4 Hz, 1H, 5-H), 2.27 (s, 3H, H₃CC(O)), 1.33 (s, 9H, C(CH₃)₃).

¹³C NMR (75 MHz, CDCl₃) δ 169.07 (s, C(O)), 147.09 (s, 2-C), 136.40 (s, 4-C), 133.52 (s, 5-C), 132.43 (s, 1-C), 122.26 (s, 6-C), 122.13 (s, 3-C), 34.70 (s, $\underline{C}(CH_3)_3$), 31.09 (s, $\underline{C}(\underline{C}H_3)_3$), 25.63 (s, $\underline{C}(CH_3)_3$).

Elemental analysis: found (calculated) for $C_{12}H_{16}N_2O_3$ (236.12 g mol⁻¹): C, 60.99 (61.00); H, 6.73 (6.83); N, 10.83 (11.86).

NMR Spectra

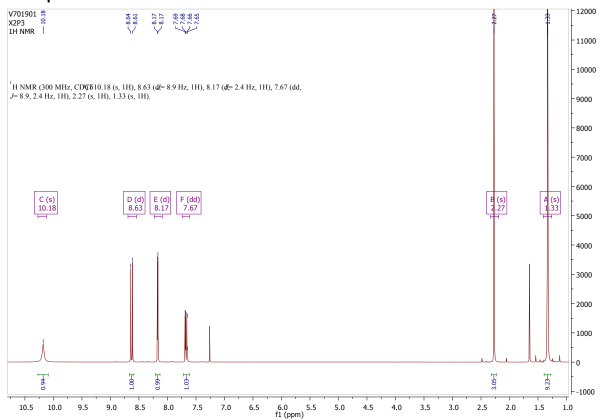


Fig. S2 ¹H NMR spectrum of 3 in CDCl₃.

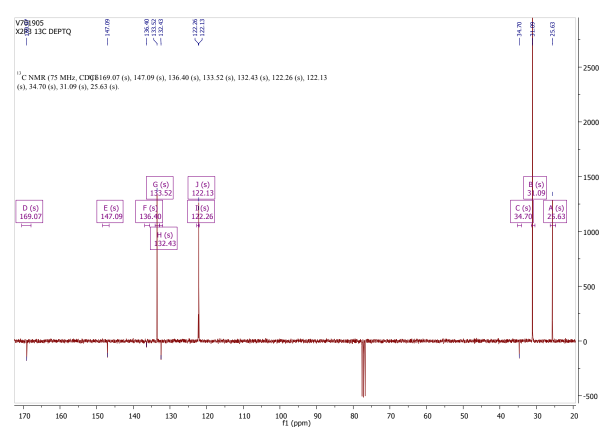


Fig. S3 ¹³C NMR {¹H} (DEPTQ-135) spectrum of **3** in CDCl₃.

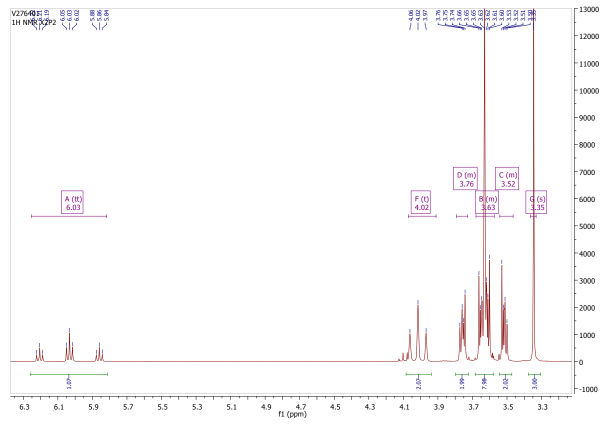


Fig. S4 ¹H NMR spectrum of 4 in CDCl₃.

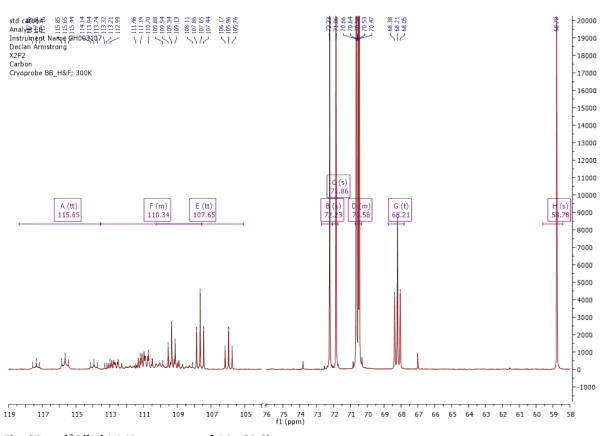


Fig. S5 ${}^{13}C{}^{1}H$ NMR spectrum of 4 in CDCl₃.

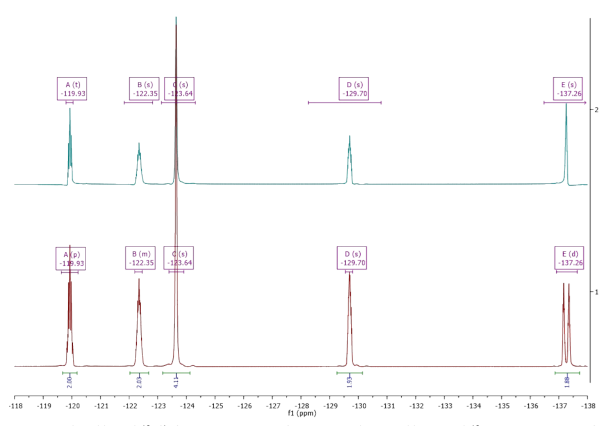


Fig. S6 (Top)(cyan) $^{19}F\{^1H\}$ NMR spectrum of **4** in CDCl₃. (Bottom)(maroon) ^{19}F NMR spectrum of **4** in CDCl₃.

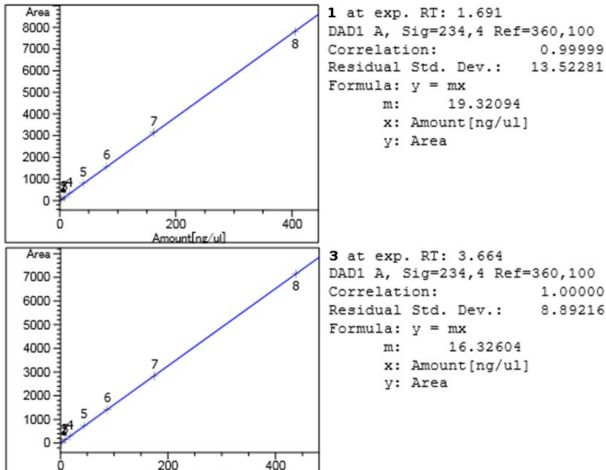
HPLC Calibration Data

```
HPLC calibration data for method analysing mixtures of compounds {\bf 1} and {\bf 3}
              General Calibration Setting
______
Calib. Data Modified : Monday, October 01, 2018 9:49:07 AM
Signals calculated separately:
                                No
Rel. Reference Window: 5.000 %
Abs. Reference Window: 0.000 min
Rel. Non-ref. Window: 5.000 %
Abs. Non-ref. Window: 0.000 min
Uncalibrated Peaks: not reported
Partial Calibration: Yes, identified peaks are recalibrated
Correct All Ret. Times: No, only for identified peaks
Curve Type: Linear
Origin
                         Forced
Weight
                         Equal
Recalibration Settings:
Average Response :
                         Average all calibrations
                      Floating Average New 75%
Average Retention Time:
                       Signal Details
______
Signal 1: DAD1 A, Sig=234,4 Ref=360,100
_____
                      Overview Table
______
                        Area Rsp.Factor Ref ISTD # Compound
  RT Sig Lvl Amount
             [ng/ul]
1.691 1 1 1.62000 32.07008 5.05144e-2 No No 1 2 4.05000 76.83419 5.27109e-2
             8.10000 155.67540 5.20313e-2
          3
            16.20000 321.02591 5.04632e-2
40.50000 808.80182 5.00741e-2
          5
            81.00000 1569.91272 5.15952e-2
          7 162.00000 3149.05396 5.14440e-2
          8 405.00000 7813.45020 5.18337e-2
```

| RT Sig Lvl | Amount [ng/ul] | Area I | Rsp.Factor | Ref : | ISTD | # | Compound |
|------------|-------------------|------------|------------|-------|------|---|----------|
| | | | | - | | - | |
| 3.664 1 1 | 1.75000 | 28.87975 | 6.05961e-2 | No. | No | | 3 |
| 2 | 4.37500 | 68.86250 | 6.35324e-2 |) | | | |
| 3 | 8.75000 | 139.87663 | 6.25551e-2 | | | | |
| 4 | 17.50000 | 289.56149 | 6.04362e-2 | | | | |
| 5 | 43.75000 | 729.74640 | 5.99523e-2 | | | | |
| 6 | 87.50000 | 1419.08252 | 6.16596e-2 | | | | |
| 7 | 175.00000 | 2844.21362 | 6.15284e-2 |) | | | |
| 8 | 437.50000 | 7148.04834 | 6.12055e-2 | 2 | | | |
| | | | | | | | |

8

Calibration Curves



HPLC calibration data for method analysing mixtures of compounds 1 and 2

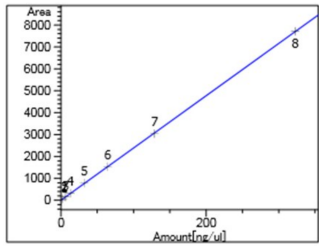
General Calibration Setting

Calib. Data Modified : Monday, October 01, 2018 10:15:31 AM Signals calculated separately: No Rel. Reference Window: 5.000 % Abs. Reference Window: 0.000 min Rel. Non-ref. Window : 5.000 % Abs. Non-ref. Window : 0.000 min : not reported Uncalibrated Peaks Yes, identified peaks are recalibrated Partial Calibration Correct All Ret. Times: No, only for identified peaks Curve Type Linear Origin Forced Weight Equal Recalibration Settings:

Amount[ng/ul]

```
Signal Details
Signal 1: DAD1 A, Sig=234,4 Ref=360,100
______
                        Overview Table
  RT Sig Lvl Amount
                        Area Rsp.Factor Ref ISTD # Compound
             [ng/ul]
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 3.648 1 1 1.33000 30.08573 4.42070e-2 No No 2 2 3.32500 74.27542 4.47658e-2
            6.65000 146.36064 4.54357e-2
13.30000 318.12717 4.18072e-2
33.25000 776.79382 4.28042e-2
          3
             66.50000 1501.62561 4.42853e-2
          7 133.00000 2968.40796 4.48052e-2
8 332.50000 7522.88916 4.41984e-2
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                        Area Rsp.Factor Ref ISTD # Compound
             [ng/ul]
5.121 1 1 1.29000 29.90119 4.31421e-2 No No 1 2 3.22500 75.31383 4.28208e-2
          3
             6.45000 148.62427 4.33980e-2
          4 12.90000 325.70456 3.96064e-2
          5
            32.25000 793.28381 4.06538e-2
            64.50000 1534.63477 4.20295e-2
          6
          7 129.00000 3038.24951 4.24587e-2
          8 322.50000 7712.12939 4.18172e-2
```

Calibration Curves

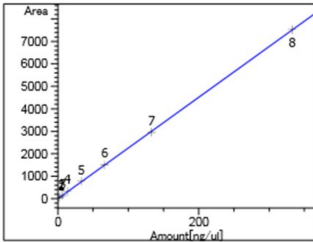


1 at exp. RT: 5.121 DAD1 A, Sig=234,4 Ref=360,100 Correlation: 0.99998 Residual Std. Dev.: 20.00348

Formula: y = mx

m: 23.86897 x: Amount[ng/ul]

y: Area



2 at exp. RT: 3.648

DAD1 A, Sig=234,4 Ref=360,100

Correlation: 0.99998

Residual Std. Dev.: 18.61429

Formula: y = mx

m: 22.59089 x: Amount[ng/ul]

y: Area

Particle Sizing

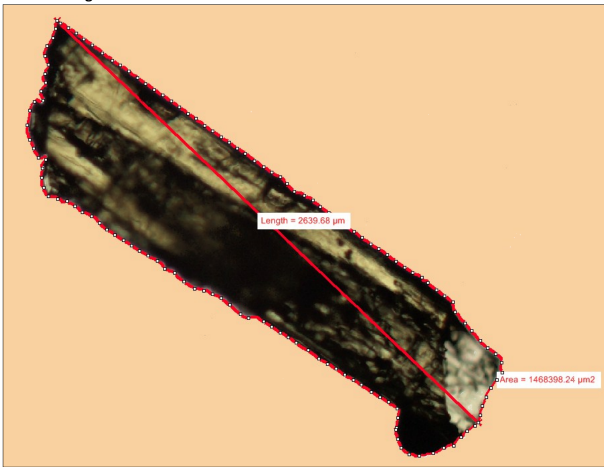


Fig. S7 Image displaying an isolated crystal of compound **1** from a solution doped with 4.0 mol % of compound **2**. This image displays how size measurements were taken; the area in μ m² was recorded from the area within the external perimeter (highlighted as a red line with white circles); the length in μ m was recorded from the longest dimension of the particle (highlighted as a red line bisecting the particle).

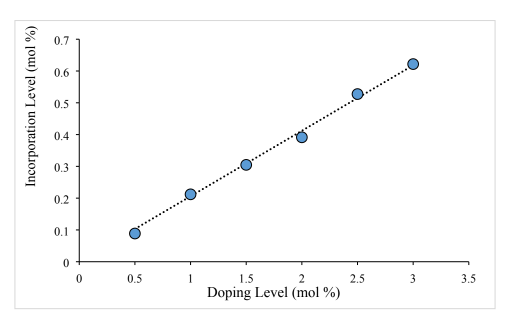


Fig. S8 Graph displaying incorporation level versus solution impurity level of 4-methyl-2-nitroacetanilide (**2**) in "host" 4-trifluoromethyl-2-nitroacetanilide.

Table S1. Overall incorporation (% composition by HPLC) of 4-methyl-2-nitroacetanilide (**2**) and 4-*tert*-butyl-2-nitroacetanilide (**3**) in crystals of 4-chloro-2-nitroacetanilide (**1**) obtained by crystallisation from solutions in toluene containing various quantities of **2** and **3** at a σ value of 1.5. Data is shown for two crystallisation batches.

| Level (mol %) of | % 2 incorporated into | % 3 incorporated | % 3 incorporated into |
|----------------------|-----------------------|---------------------|-------------------------|
| impurity (2 or 3) in | crystals | into first batch of | second batch of crytals |
| solution | | crystals | |
| 5.0 | 0.980 (0.03) | 0.0992 (0.0002) | 0.0599 (0.0002) |
| 4.5 | 0.917 (0.012) | 0.1867 (0.0005) | |
| 4.0 | 0.838 (0.011) | 0.01302 (0.00014) | 0.0934 (0.0003) |
| 3.5 | 0.690 (0.015) | 0.0852 (0.0003) | 0.0304 (0.0002) |
| 3.0 | 0.630 (0.05) | 0.0564 (0.0002) | 0.0717 (0.0004) |
| 2.5 | 0.497 (0.017) | 0.0520 (0.0003) | 0.02615 (0.00007) |
| 2.0 | 0.442 (0.003) | 0.0705 (0.0003) | |
| 1.5 | 0.340 (0.06) | 0.0212 (0.0002) | |
| 1.0 | 0.1980 (0.0014) | 0.03897 (0.00009) | |
| 0.5 | 0.101 (0.003) | 0.0172 (0.0003) | |

PXRD

Fig. S9 shows the PXRD pattern for $\bf 1$ grown in toluene at $\sigma=1.5$ against the PXRD patterns for $\bf 1$ grown in toluene at $\sigma=1.5$ with 5 mol % additions $\bf 2$. The PXRD pattern for $\bf 2$ grown under the same conditions is also added for reference. PXRD patterns were obtained for crystals of $\bf 1$ with additive concentrations as low as 0.5% grown under the aforementioned conditions, however, for none of the samples, as can be seen at the highest additive concentration of $\bf 2$, there is no discernible formation of additional diffraction peaks. All obtained patterns displayed peaks corresponding to that of $\bf 1$ with only minor differences in the intensities of some of the peaks peaks.

A similar image comparing the PXRD patterns based around compound **3** can be seen in Fig. S10. This figure compares the PXRD patterns for crystals related to pure **1**, 4.5 mol % **3**-doped **1**, and pure **3**, all grown in toluene at $\sigma = 1.5$. Again, the sample doped with impurity displays no diffraction peaks other than from the pure **1** sample with minor differences in intensity.

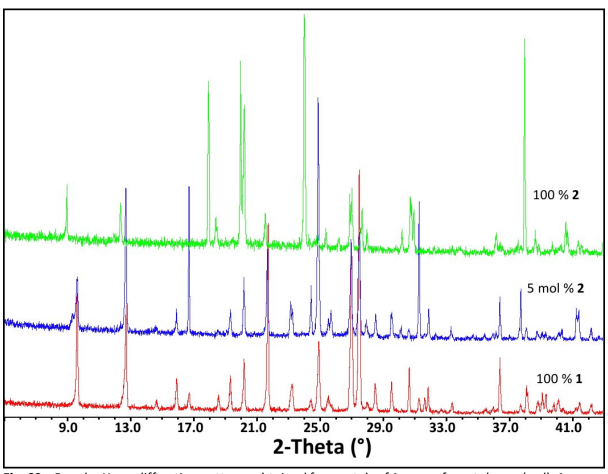


Fig. S9 Powder X-ray diffraction patterns obtained for crystals of **1** grown from toluene (red), **1** grown from toluene with concentrations of 5 mol% **2** (blue), and **2** grown from toluene (green).

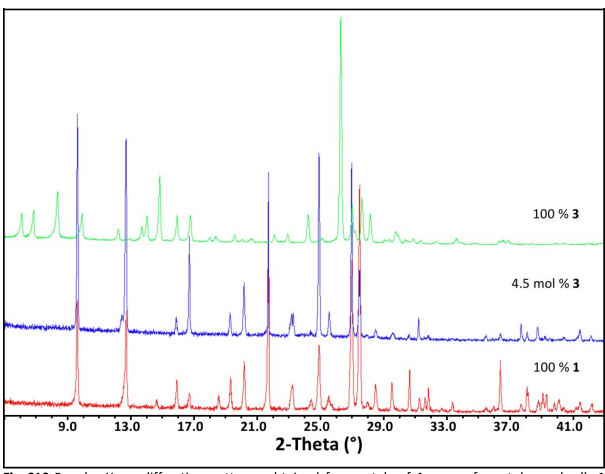


Fig. S10 Powder X-ray diffraction patterns obtained for crystals of 1 grown from toluene (red), 1 grown from toluene with concentrations of 4.5 mol % 3 (blue), and 3 grown from toluene (green).

Differential Scanning Calorimetry

The DSC curves for pure compounds **1** and **2**, and 5 mol % **2**-doped **1** are displayed in Fig. S11. The inclusion of additive at the highest level in this study did not alter the melting point of the crystals in any significant way, as the melting of 99.51 °C is for the impurity-doped sample is marginally higher than the melting point of 99.44 °C for pure compound **1**.

Fig. S12 shows the DSC curves for pure compounds **1** and **3**, and 4.5 mol % **3**-doped **1**. Again the impurity-doped sample shows a minimal difference in melting point, rising to 100.11 °C from 99.44 °C for pure compound **1**.

None of the samples displayed in Fig. S11 and Fig. S12 showed any secondary events such as minor melting points or polymorph changes under the tested conditions.

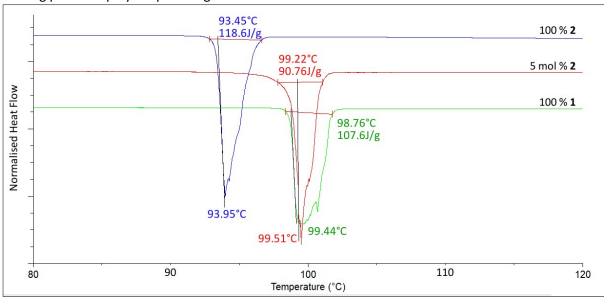


Fig. S11 DSC curves obtained from crystals of **1** containing 5 mol % **2** (red); compared to DSC curves of component compounds **1** (green) and **2** (blue).

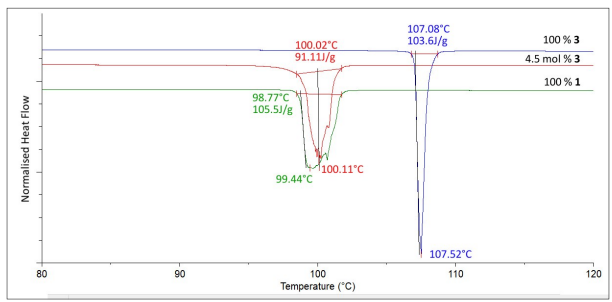


Fig. S12 DSC curves obtained from crystals of **1** containing 4.5 mol % **3** (red); compared to DSC curves of component compounds **1** (green) and **3** (blue).

Partial Dissolution Graphs

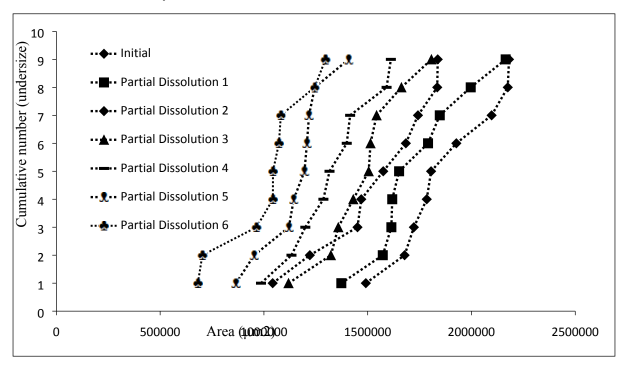


Fig. S13 Chart comparing particle area versus the ranking of each particle in a partial dissolution series of **1** doped with 1.5 mol % of **2**.

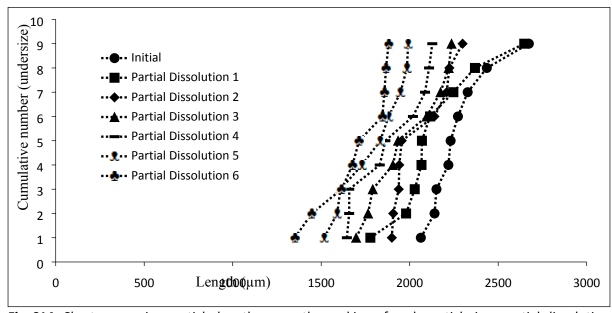


Fig. S14 Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 1.5 mol % of **2**.

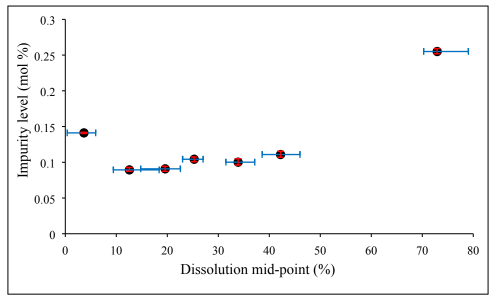


Fig. S15 Plot of percentage by HPLC of added impurity in crystals of compound **1** *vs.* the dissolution mid-point for the sample of crystals grown from solutions containing 1.5 mol % of additive **2**.

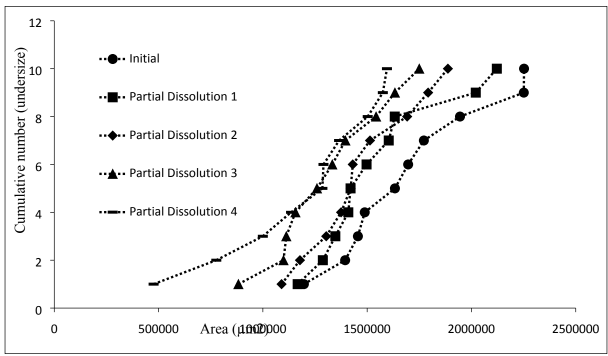


Fig. S16 Chart comparing particle area versus the ranking of each particle in a partial dissolution series of **1** doped with 2.0 mol % of **2**.

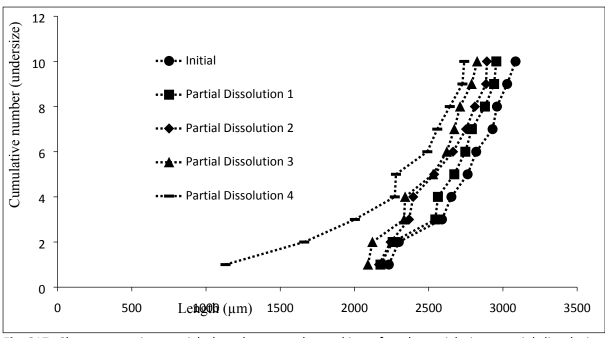


Fig. S17 Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 2.0 mol % of **2**.

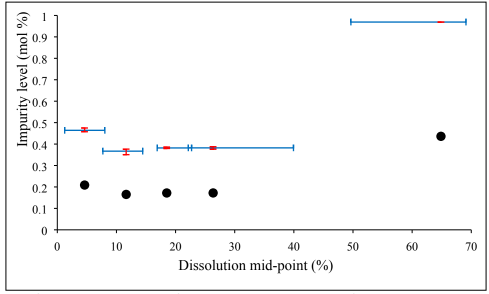


Fig. S18 Plot of percentage by HPLC of added impurity in crystals of compound **1** *vs.* the dissolution mid-point for the sample of crystals grown from solutions containing 2.0 mol % of additive **2**.

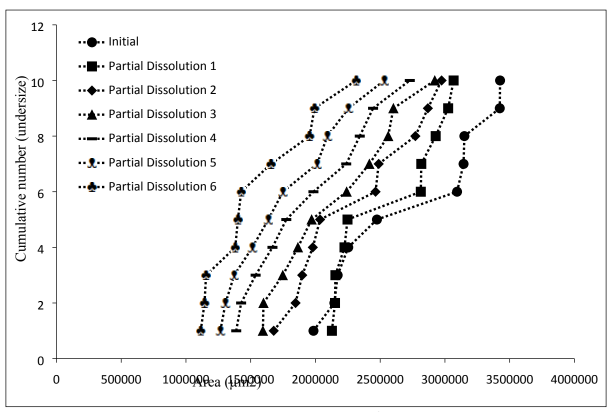


Fig. S19 Chart comparing particle area versus the ranking of each particle in a partial dissolution series of **1** doped with 2.5 mol % of **2**.

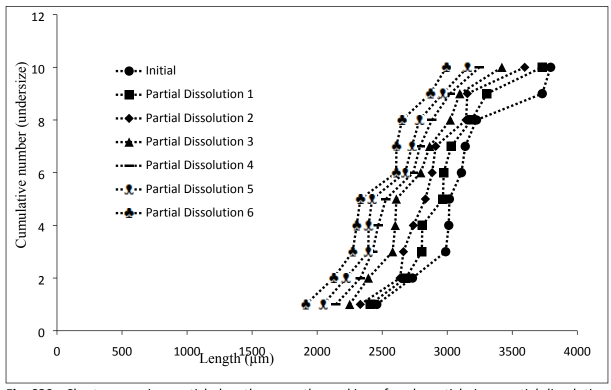


Fig. S20 Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 2.5 mol % of **2**.

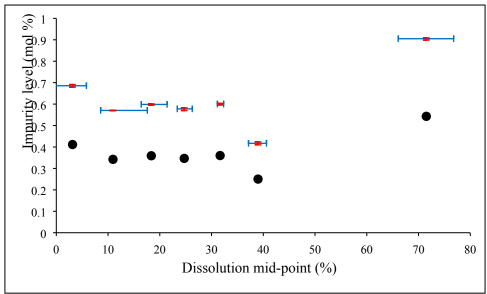


Fig. S21 Plot of percentage by HPLC of added impurity in crystals of compound **1** *vs.* the dissolution mid-point for the sample of crystals grown from solutions containing 2.5 mol % of additive **2**.

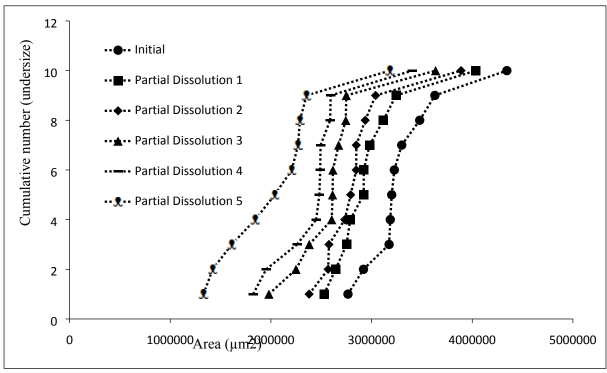


Fig. S22 Chart comparing particle area versus the ranking of each particle in a partial dissolution series of $\bf 1$ doped with 3.0 mol % of $\bf 2$.

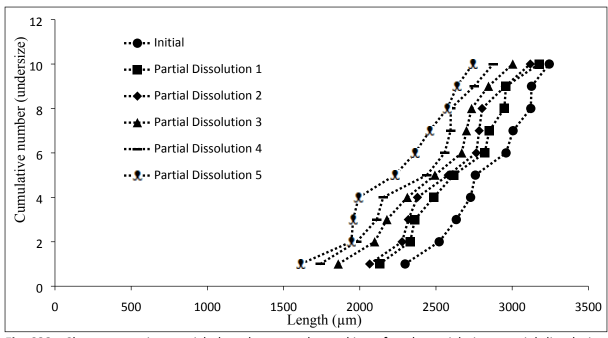


Fig. S23 Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 3.0 mol % of **2**.

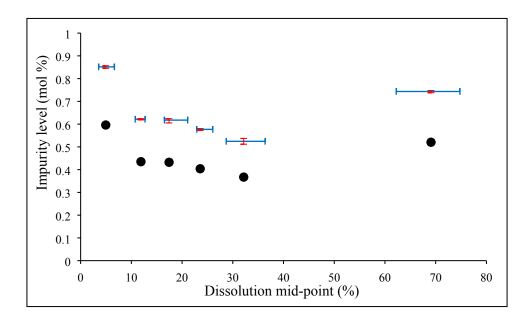


Fig. S24 Plot of percentage by HPLC of added impurity in crystals of compound **1** *vs.* the dissolution mid-point for the sample of crystals grown from solutions containing 3.0 mol % of additive **2**.

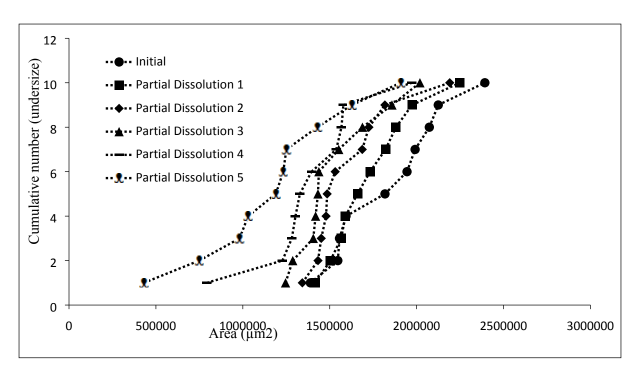


Fig. S25 Chart comparing particle area versus the ranking of each particle in a partial dissolution series of **1** doped with 3.5 mol % of **2**.

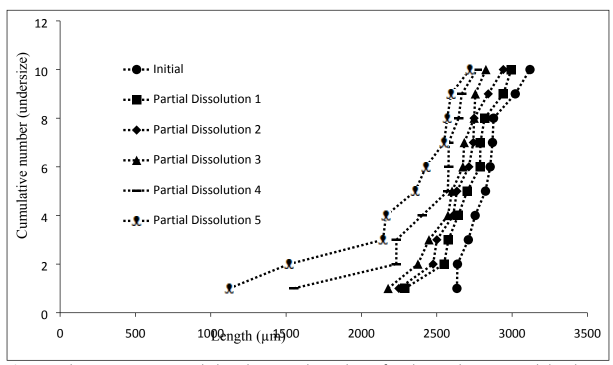


Fig. S26 Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 3.5 mol % of **2**.

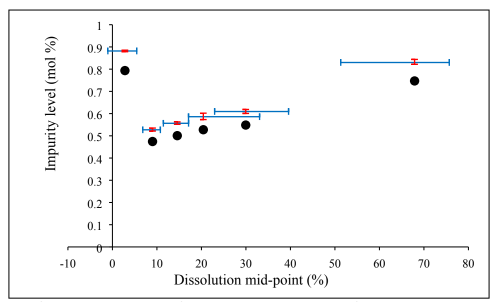


Fig. S27 Plot of percentage by HPLC of added impurity in crystals of compound **1** *vs.* the dissolution mid-point for the sample of crystals grown from solutions containing 3.5 mol % of additive **2**.

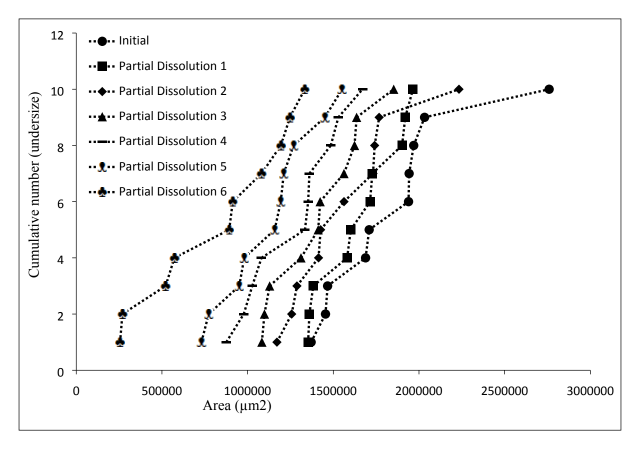


Fig. S28 Chart comparing particle area versus the ranking of each particle in a partial dissolution series of **1** doped with 4.0 mol % of **2**.

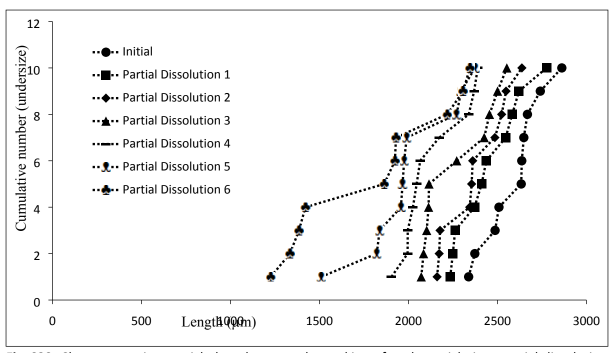


Fig. S29 Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 4.0 mol % of **2**.

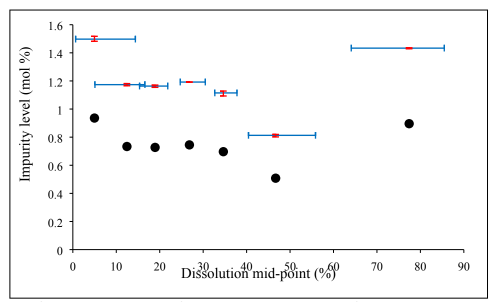


Fig. S30 Plot of percentage by HPLC of added impurity in crystals of compound **1** *vs.* the dissolution mid-point for the sample of crystals grown from solutions containing 4.0 mol % of additive **2**.

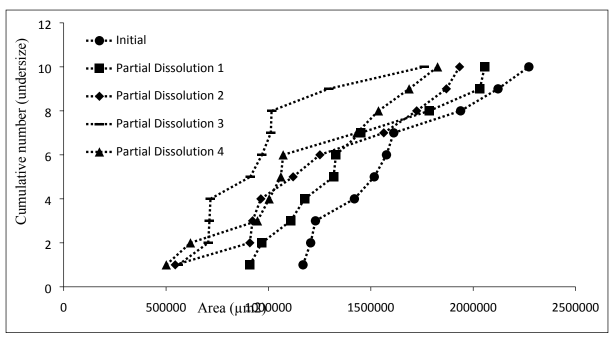


Fig. S31 Chart comparing particle area versus the ranking of each particle in a partial dissolution series of **1** doped with 4.5 mol % of **2**.

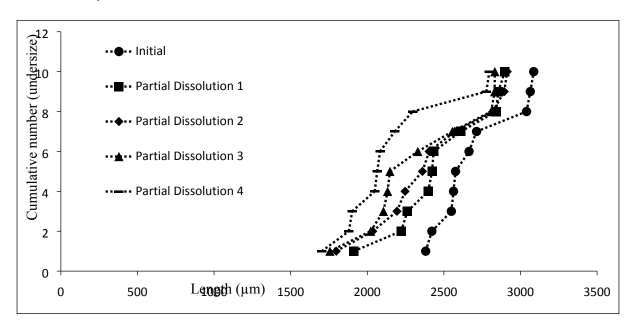


Fig. S32 Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 4.5 mol % of **2**.

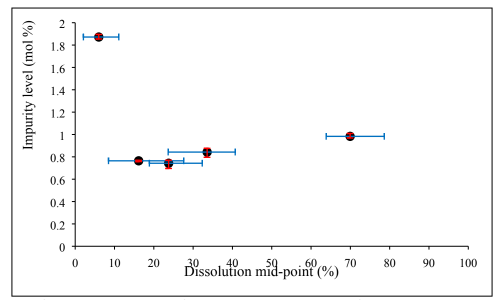


Fig. S33 Plot of percentage by HPLC of added impurity in crystals of compound **1** *vs.* the dissolution mid-point for the sample of crystals grown from solutions containing 4.5 mol % of additive **2**.

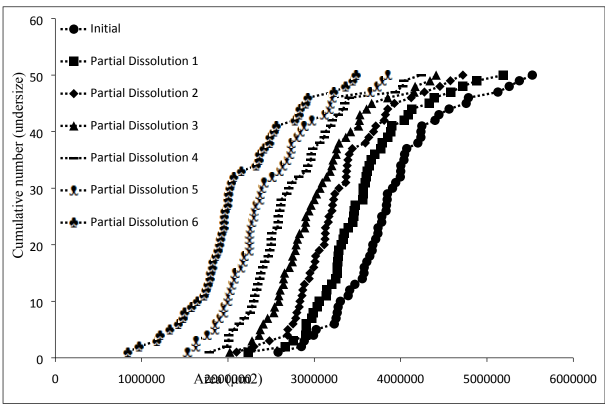


Fig. S34 Chart comparing particle area versus the ranking of each particle in a partial dissolution series of **1** doped with 5.0 mol % of **2**.

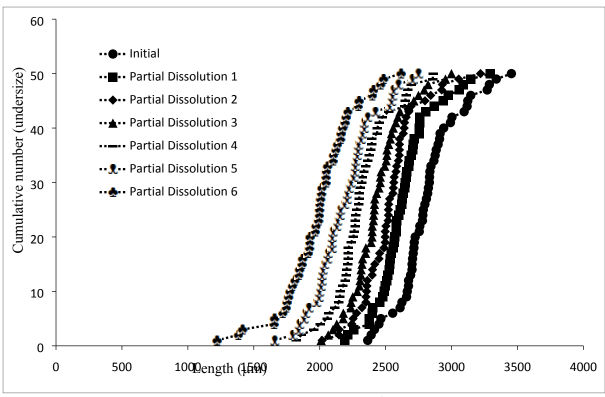


Fig. S35 Chart comparing particle length versus the ranking of each particle in a partial dissolution series of **1** doped with 5.0 mol % of **2**.

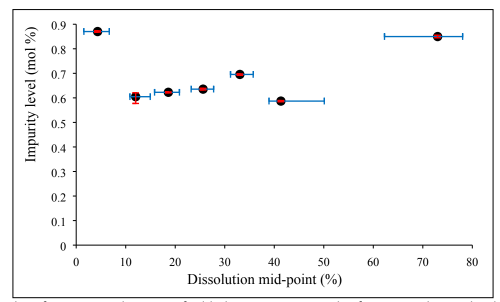


Fig. S36 Plot of percentage by HPLC of added impurity in crystals of compound **1** *vs.* the dissolution mid-point for the sample of crystals grown from solutions containing 5.0 mol % of additive **2**.

Table S2. Comparisons of parent batch average and weighted averages from stepwise dissolutions.

| Entry | Additive | %additive in soln. | %additive in parent sample | Weighted% additive in dissolution sample* | Difference (%) |
|-------|----------|--------------------|----------------------------|---|---------------------|
| 1 | 2 | 1.5 | 0.340 (0.06) | 0.1664 (0.0014) | +0.18 (0.06) |
| 2 | 2 | 2.0 | 0.442 (0.003) | 0.3573 (0.0005) | +0.085 (0.003) |
| 3 | 2 | 2.5 | 0.497 (0.017) | 0.4054 (0.0011) | +0.091 (0.017) |
| 4 | 2 | 3.0 | 0.630 (0.05) | 0.4961 (0.0015) | +0.13 (0.05) |
| 5 | 2 | 3.5 | 0.690 (0.015) | 0.665 (0.005) | +0.025 (0.016) |
| 6 | 2 | 4.0 | 0.838 (0.011) | 0.7866 (0.0018) | +0.051 (0.011) |
| 7 | 2 | 4.5 | 0.917 (0.012) | 1.014 (0.012) | -0.097 (0.017) |
| 8 | 2 | 5.0 | 0.980 (0.03) | 0.739 (0.003) | +0.24 (0.03) |
| 9 | 3 | 4.5 | 0.1867 (0.0005) | 0.03995 (0.00015) | +0.1468 (0.0005) |

^{*}The weighted average was determined from the HPLC data using the following formula:

$$\sum_{i=1}^{n} \left(\frac{D_i}{\sum_{i=1}^{n} D_i} \right) \cdot R_i$$

where n is the number of partial dissolutions in a dissolution series, D_i is the number of moles of analytes dissolved in a partial dissolution step i, and R_i is the percentage of additive in the solution obtained from partial dissolution step i.

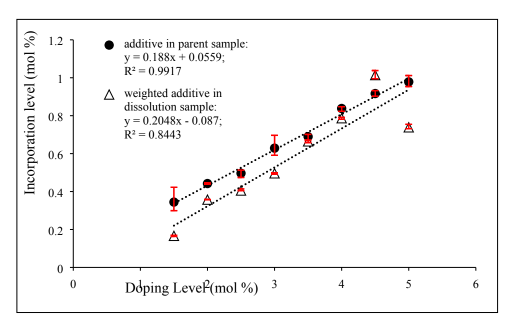


Fig. S37 Comparisons of parent batch average and weighted averages from stepwise dissolutions.

Table S3. Dissolution mid-points calculated for analysed 4.0 mol % **2**-doped **1** crystals from both observed measurements and 'theoretical' values.

| Sample | Obs. Dissolution Area MdPt [%] | 'Theor.' Dissolution Area MdPt [%] | Area MdPt Difference [%] [Obs Theor.] | Obs. Dissolution Length MdPt [%] | 'Theor.' Dissolution Length MdPt [%] | Length MdPt Difference [%] [Obs Theor.] |
|---------|--------------------------------|---|--|----------------------------------|---|---|
| Initial | 5.0 (3.9) | 3.4 | 1.6 (3.9) | 2.71 (1.08) | 1.73 | 0.98 (1.08) |
| PD1 | 12.5 (8.4) | 10.3 | 2.2 (8.4) | 6.8 (2.2) | 5.3 | 1.5 (2.2) |
| PD2 | 18.9 (3.5) | 17.5 | 1.4 (3.5) | 10.3 (2.6) | 9.2 | 1.1 (2.6) |
| PD3 | 26.84 (5.08) | 25.01 | 1.83 (5.08) | 15.05 (3.92) | 13.43 | 1.62 (3.92) |
| PD4 | 34.6 (5.9) | 32.9 | 1.7 (5.9) | 20.2 (4.0) | 18.1 | 2.1 (4.0) |
| PD5 | 46.7 (8.4) | 41.4 | 5.3 (8.4) | 26.7 (7.5) | 23.5 | 3.2 (7.5) |
| PD6 | 77.4 (17.7) | 72.9 | 4.5 (17.7) | 65.3 (13.9) | 63.2 | 2.1 (13.9) |

References

S. Krakert, N. Ballav, M. Zharnikov and A. Terfort, *Phys. Chem. Chem. Phys.*, 2010, **12**, 507–515.