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 Cite this: *CrystEngComm*, 2016, 18, 2408

 Received 12th February 2016,
Accepted 9th March 2016

DOI: 10.1039/c6ce00366d

www.rsc.org/crystengcomm

Exploiting polymorphism in second sphere coordination: thermal transformation, NLO properties and selective mechanochemical synthesis†

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The racemic organic building block L acting as a first sphere ligand yields a new second sphere adduct $[\text{LH}]^+ \cdot [\text{FeCl}_4]^-$ crystallizing as racemic polymorphs α and β . Solid-state DFT calculations show that polymorph β is the kinetic adduct as observed experimentally. The β -phase crystallizes in the polar space group $Pna2_1$ and displays NLO properties. Mechanochemical synthesis yields only the β -phase.

The combination of organic cations and metal anions to synthesize hybrid metal–organic structures *via* second sphere coordination was used in the early 80s.¹ The importance of second sphere coordination has been demonstrated in supramolecular chemistry² recognition phenomena in biological systems,³ metal-based anion receptors^{4–6} and catalysis.⁷ In the solid state, second sphere coordination has been studied much less, but recently, gas adsorption,⁸ metal-ion separation,⁹ structural rearrangements,¹⁰ and chiral systems¹¹ have been reported.

As defined by Gibb's phase rule, polymorphism refers to the situation in which two or more crystalline phases have the same chemical composition but different structures. Polymorphism,¹² usually confined to pharmaceutical organic materials,¹³ is of fundamental importance in academy and industry due to its effect on the solid-state properties of a solid material. Interestingly, polymorphic examples of hybrid metal–organic materials self-assembled *via* outer sphere coordination

are rare despite the long history of such interactions. Self-assembling organic molecules and metals in a second sphere adduct in which the organic part is racemic can indeed increase the options of discovering a variety of structures such as racemic second sphere adducts, crystallization of pure *R* or *S* enantiomers (*i.e.*, conglomerate), or even polymorphism in the second sphere adduct. However, to the best of our knowledge, racemic organic crystals as first sphere ligands have not been applied to explore polymorphism in second sphere adducts.

Herein, we report a racemic crystalline form of the organic *N,N,N',N'*-tetrabenzyl-*trans*-1,2-diaminocyclohexane (L), and how racemic polymorphism is also observed in the second sphere adduct $[\text{LH}]^+ \cdot [\text{FeCl}_4]^-$. Upon the self-assembly of L with $\text{FeCl}_3 \cdot 6\text{H}_2\text{O}$ and HCl, $[\text{LH}]^+ \cdot [\text{FeCl}_4]^-$ crystallizes as two polymorphs: α -phase ($P2_1/c$) and β -phase ($Pna2_1$). Both polymorphs contain *R* and *S* enantiomers in the structure, and so are racemic. Weak, attractive C–H \cdots Cl and repulsive C–H \cdots H–C supramolecular interactions play important roles in driving the molecules into non-polar and polar packing. Solid-state DFT calculations showed that the most stable structure is the thermodynamic α -phase. The structural transformation from the acentric and polar $Pna2_1$ to the centric $P2_1/c$ structure is described upon thermal treatment in the solid state. The selective formation of the β -phase can be obtained by means of liquid-assisted grinding (LAG). Moreover, the single crystal second harmonic generation (SHG) behavior for the polar β -phase is reported, which shows significant efficiency compared to that of urea.

Ligand L was synthesized by reacting *trans*-1,2-diaminocyclohexane and benzyl bromide in the presence of $\text{LiOH}/\text{CH}_2\text{Cl}_2$ (Scheme 1).¹⁴ Crystals of L suitable for single crystal XRD (SC-XRD) were obtained upon evaporation from ethanol. In the crystalline state, L packs in the $C2/c$ space group with half a molecule in the asymmetric unit (see the ESI†). The crystal structure of L is racemic¹⁴ while the acentric (*R,R,N,N'*-tetrabenzyl-1,2-diaminocyclohexane) crystallizing in the

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† Electronic supplementary information (ESI) available: Experimental details, crystal structure description, additional figures. CCDC 1452830–1452832 (α -phase, β -phase, L) contain the supplementary crystallographic data for this paper. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c6ce00366d



monoclinic I_2 space group is the enantiopure form.¹⁵ The bulk powder of the as synthesized L belongs to the phase crystallizing in the $C2/c$ space group.¹⁴ In L, the bond angles around N1 were $112.9(1)^\circ$, $112.4(1)^\circ$, and $110.0(1)^\circ$ for C1–N1–C2, C1–N1–C5 and C2–N1–C5, respectively. This geometric information indicates that the conformation around each nitrogen atom is near sp^3 hybridised.

The lone pairs of each nitrogen atom are oriented towards the diamine pocket formed by the two N,N' -benzyl substituents on each side.

The ability of L to form second sphere adducts was explored in the solid state. The second sphere supramolecular structure was prepared by mixing a $CH_2Cl_2/MeOH$ solution of $FeCl_3 \cdot 6H_2O$ with L in the presence of HCl. After three days, crystals with two different habits were obtained: a large amount of yellow hexagonal crystals (α) and a small amount of yellow needle crystals (β).¹⁴

The structural characterization of α - and β -phases was carried out by SC-XRD. X-ray crystallography studies reveal that α - and β -phases are racemic polymorphs that crystallize in the monoclinic space group $P2_1/c$ and orthorhombic space group $Pna2_1$, respectively. The different phases have been confirmed by X-ray powder diffraction (XRPD) experiments (Fig. 4). The β -phase belongs to a noncentrosymmetric and polar space group. It is worth noting that polar crystals are highly demanded in materials science because they can display piezoelectric, pyroelectric, ferroelectric, and SHG properties (*vide ante*).

Both second sphere adducts contain an independent $[FeCl_4]^-$ anion and a $[HL]^+$ cation in the asymmetric unit. Each $[FeCl_4]^-$ anion in α - and β -phases has almost the same coordination environment and adopts a distorted tetrahedral geometry coordinated by four Cl atoms.¹⁴

It is worth mentioning that although the monocation is protonated and the N–H group in the diamine pocket is a po-

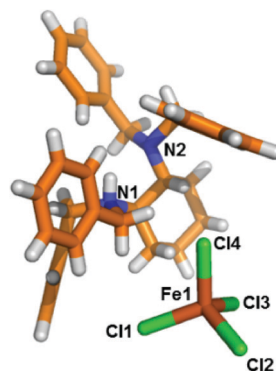
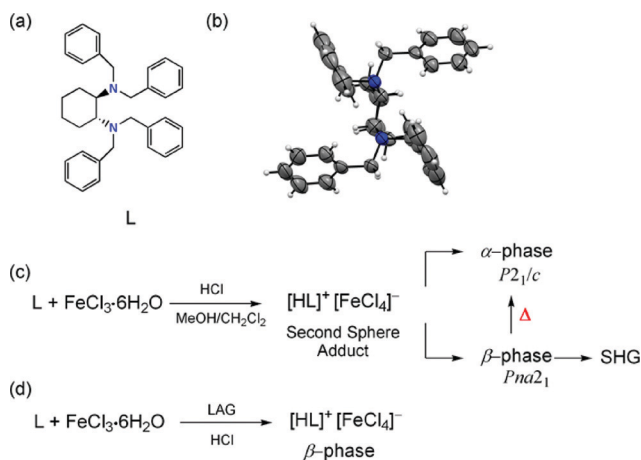


Fig. 1 Stick model of the asymmetric unit of $[LH]^+ \cdot [FeCl_4]^-$ (α -phase). Color code: carbon: orange; nitrogen: blue; chloride: green; iron: brown; hydrogen: white.

tential hydrogen bonding donor, the structural analysis indicated that it is “locked” by the four extended benzyl rings. There are no charge-assisted hydrogen bonds among $[HL]^+$ and the $[FeCl_4]^-$ anion participating in the structural formation. Thus, no *quasi*-chelating motif is observed in the supramolecular adduct.^{19c} Instead, weak interactions such as C–H \cdots Cl contacts play a critical role in the self-assembly of the second sphere adduct.

For the α -phase, as seen in Fig. 2, each $[FeCl_4]^-$ anion interacts with three neighbouring $[HL]^+$ cations through five C–H \cdots Cl interactions, with the methylene group (i. C11–H11A \cdots Cl3: 3.798(6) Å, $149.1(4)^\circ$; ii. C11–H11B \cdots Cl1: 3.486(6) Å, $128.8(4)^\circ$), and the other two are from $[HL]^+$ at (x, y, z) with the benzene ring and the methylene group, respectively (iii. C15–H15 \cdots Cl2: 3.794(8) Å, $151.8(4)^\circ$; iv. C13–H13A \cdots Cl2: 3.771(6) Å, $145.0(4)^\circ$). The structure contains both enantiomers. The 1D chains are further connected with each other through extensive C–H \cdots Cl interaction (v. C24–H24A \cdots Cl1: 3.83(1) Å, $160.6(7)^\circ$), forming a helical layer along the b -axis, as seen in Fig. S10.†

A rather diverse set of interactions are observed in the β -phase. In this structure, the $[FeCl_4]^-$ anion interacts with two $[HL]^+$ cations (Fig. 3). The first anion–cation interaction



Scheme 1 Chemical structure (a) and a view of the molecular structure. Displacement ellipsoids are drawn at the 50% probability level and H atoms are shown as small spheres of arbitrary radius (b). Synthetic route for the second sphere adduct $[LH]^+ \cdot [FeCl_4]^-$ giving rise to α - and β -phases in solution (c). LAG synthesis of the β -phase (d).

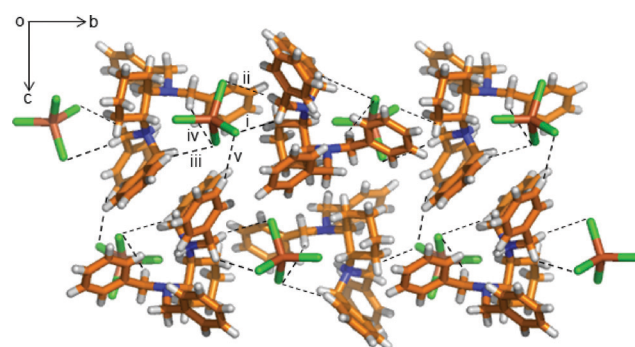


Fig. 2 Crystal structure of the α -phase viewed along the a -axis showing the various C–H \cdots Cl interactions among $[HL]^+$ cations and $[FeCl_4]^-$ anions. The hydrogen bonding interactions are shown with a dashed black line. Color code as in Fig. 1.



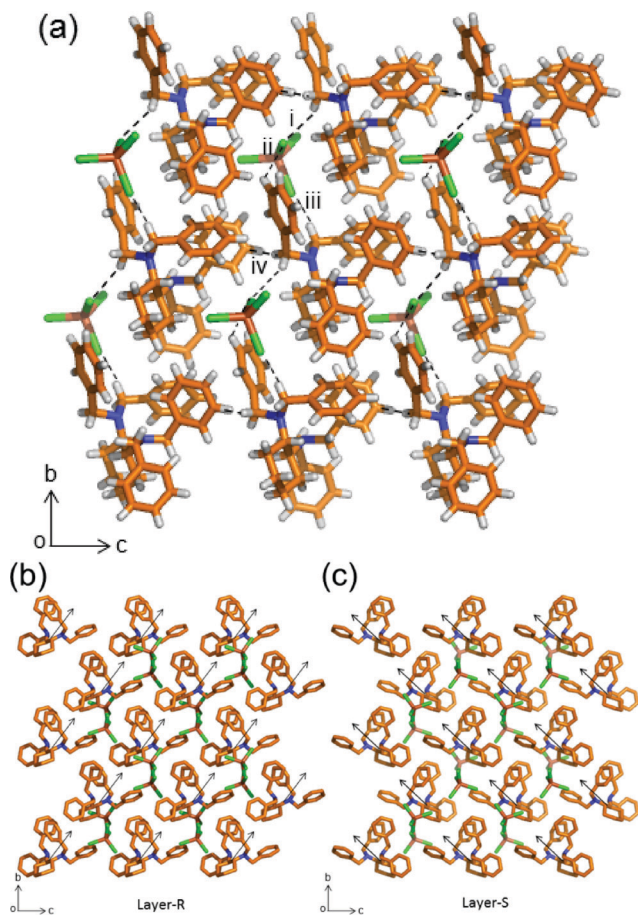


Fig. 3 Crystal structure of the β -phase viewed along a -axis formed by C–H \cdots Cl interactions among [HL] $^+$ cations and [FeCl $_4$] $^-$ anions and repulsive C–H \cdots H–C intermolecular interactions. The hydrogen bonding interactions are shown with a dashed black line (a). Packing of half a unit cell (a -axis) showing the non-centrosymmetric and polar orientation (black arrows) of layers of the enantiomers R (b) and S (c). Color code as in Fig. 1.

is through one C–H \cdots Cl contact with the methylene group (i. C14–H14B \cdots Cl1: 3.78(2) Å, 150.4(9) $^\circ$).

Then the second ion pair interaction involves contacts ii. C21–H21B \cdots Cl3: 3.62(2) Å, 129.1(9) $^\circ$ and iii. C19–H19 \cdots Cl1: 3.61(2) Å, 129(1) $^\circ$. As in the α -phase, the structure is composed of both enantiomers of L but arranged in a non-centrosymmetric manner (Fig. S12 †).¹⁴ There are also C–H \cdots H–C repulsive interactions between the aryl and methylene groups (iv. C14–H14A \cdots H33: 2.4881(3) Å). Further proof of the different crystal packings for both polymorphs is confirmed by IR experiments which show that the C–N and C–H stretching bands of $-\text{CH}_2$ are different for polymorphs α and β (Fig. S14 and S15 †).

Therefore, weak attractive C–H \cdots Cl and repulsive C–H \cdots H–C intermolecular interactions have played important roles in driving the molecules into different packing assemblies. Such non-attractive contacts are also relevant in maintaining the stability of the supramolecular adduct. The polar nature in the β -phase can be observed in the orientation of the [HL] $^+$ cations “towards” the b -axis as shown by the

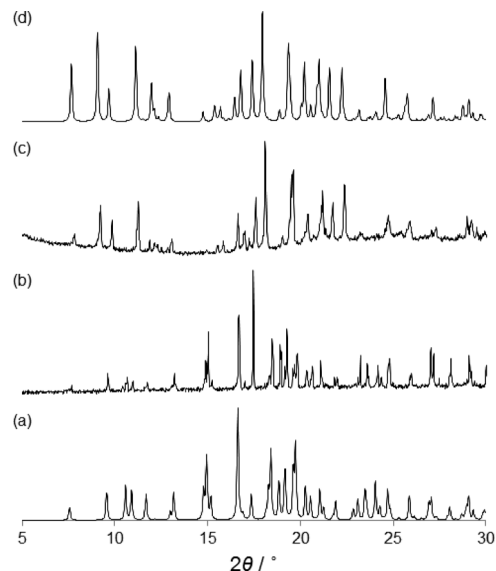


Fig. 4 The simulated isotropic XRPD patterns of β (a) and α (d) phases calculated from single crystal structures. Experimental XRPD pattern corresponding to the β phase at room temperature (b) and that of the β phase at 110 $^\circ\text{C}$ measured *ex situ* at room temperature (c).

black arrows in Fig. 3b and c. The overall crystal packing is non-centrosymmetric (Fig. S12 †). Racemates tend to crystallize in centrosymmetric crystals; however, a racemate can crystallize in centro- and non-centrosymmetric space groups as polymorphic forms as in this case.

Given that the calculated densities for polymorphs α and β are very similar (1.300 and 1.334 g cm $^{-3}$, respectively), it may be assumed that the free energy of these two polymorphic forms are quite similar.¹⁶ Thus, in order to investigate the stability of the two phases, solid-state Density Functional Theory (DFT) calculations were performed.¹⁴

The calculations were accomplished by fixing cell parameters and by optimising atomic coordinates within the experimental lattices. Explicit van der Waals corrections were included in the calculations¹⁷ as these terms can be important for subtle inter- and intra-particle interactions (*i.e.*, attractive and repulsive).¹⁸ This approach has been proved to be adequate in a number of cases, such as supramolecular complexes,¹⁹ crystalline systems containing charged particles²⁰ and crystalline phases of thiophene-based oligomers and polymers.²¹

The calculations showed that the α -phase (the lower density system) possesses stronger cation–anion interactions, within the crystalline phase, with respect to those corresponding to the β -phase (the higher density system) by about 16 or 14 kcal mol $^{-1}$ (*i.e.*, higher sublimation energy) obtained by DND (double-numeric quality basis set plus polarisation functions on heavy atoms, *i.e.* comparable to Gaussian 6-31G *) and DNP (double-numeric quality basis set plus polarisation functions on all atoms, comparable to Gaussian 6-31G **) sets, respectively. Therefore, according to the DFT results, the α -phase is the thermodynamic structure (lower energy) and the β -phase is the kinetic structure (higher



energy). Thus, the β -phase could be transformed into the α -phase in the solid state, for instance by heating.

To further investigate the above mentioned phase transition of the β to the α form in the solid state, we have carried out *ex situ* XRPD experiments on heated samples of the β form (Fig. 4). The microcrystalline β -phase was heated from room temperature to 140 °C. When the β -phase reached 80 °C the XRPD pattern shows that there was only one phase (β), but reaching 100 °C a mixture of β and α -phases is observed (see the ESI†). However, when 110 °C was achieved a complete phase transformation occurs and only the α -phase is present (Fig. 4c). Further heating shows that the crystallite melts at *ca.* 140 °C. The phase transition and the melting of the second sphere adducts have been corroborated by DSC experiments.¹⁴ Clearly, the polymorphic phase transition is carried out in the solid state.

Therefore, in light of the observed results, we can describe the α -phase as the thermodynamic phase and the β phase as the kinetic product. The formation of the β -phase can be described as the structure pertaining to a local minima in the energy landscape during the self-assembly of the metal anions and organic cations (*i.e.*, small activation energy). The formation of the thermodynamic α -phase corresponds to the global minimum, which in this case is obtained by applying temperature to the system, the β -phase.

In general, controlling the formation of a desired polymorphic form is difficult. Here we have monitored various crystallization methods using different solvents such as methanol, ethanol, THF, propanol, acetone, acetonitrile, ethyl acetate, and cyclohexane, but the outcomes resulted in mixtures of the two phases, an indication of their comparable energies. As an alternative crystallization route, solid-state grinding has become an efficient method for co-crystal grinding,²² affording good opportunities for the selective preparation of a given polymorph.²³ Herein we found that using LAG it is possible to control the formation of only the β polymorph. In the presence of various solvents, the grinding ligand L and $\text{FeCl}_3 \cdot 6\text{H}_2\text{O}$ with one drop of HCl gave the β -phase but not the α -phase (Fig. S16†).²⁴ Therefore, the selective formation of the kinetic and noncentrosymmetric phase is possible *via* LAG but not from solution crystallization. Kinetic products have been obtained by mechanochemical methods in other systems.^{23c}

Crucially, the properties of materials in the solid state depend not only on their building blocks, but also on how these subcomponents are arranged within the crystalline lattice. In the β -phase the organic cations are oriented in one particular way (Fig. 3), and for this reason the space group where crystallizes is non-centrosymmetric and polar ($Pna2_1$). This is relevant because the β -phase can show nonlinear optical properties such as SHG as well as ferroelectricity and piezoelectricity.²⁵ To test this phenomenon, SHG experiments were carried out using single crystals of the β -phase.^{14,26}

From the experimental results shown in Fig. 5, the β -phase displays a single crystal SHG efficiency 0.18 times that of urea.²⁷ We note that the single crystal after the SHG

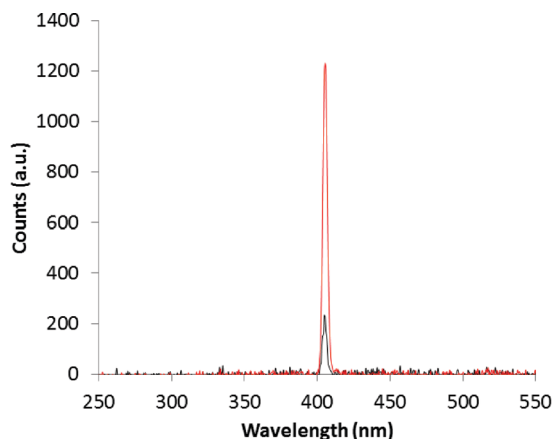


Fig. 5 Single crystal SHG signals for urea (red) and the β -phase (black) of the super-complex $[\text{LH}]^+ \cdot [\text{FeCl}_4]^-$.

experiment has the ability to withstand laser irradiation as it was not burned out and survived at least three measurements. This is important for potential applications in areas of photonics such as electro optics and frequency conversion. To the best of our knowledge, this is one of the few reports of single crystal SHG measured in a second sphere adduct on polar crystals.

In summary, the potential of using racemic organic molecules as templates in the self-assembly with $[\text{FeCl}_4]^-$ has led to the discovery of a new racemic polymorphic system in a second sphere adduct.²⁸ X-ray crystallography reveals that in the β -phase the $[\text{HL}]^+$ cations are oriented within the crystal-line lattice giving its polar nature despite being a racemic crystal, with good SHG efficiency. Mechanochemical LAG has demonstrated that only the β -phase is obtained in the solid state. DFT solid phase calculations in combination with X-ray experiments showed that the thermodynamic product is the α -phase.

The stability and non-linear optical (NLO) behavior of the β -phase could render this material suitable for functional applications in areas such as electro-optics and frequency conversion. Ferroelectricity and piezoelectricity phenomena can also be exploited in the second sphere adduct reported due to the polar nature of the β -phase. Second sphere adducts self-assembled with racemic or enantiopure organic building blocks certainly are good candidates to synthesize new hybrid metal-organic structures with potential in many technological applications. We are currently designing new organic building blocks to be used as first sphere coordination in outer sphere adducts.

Acknowledgements

This research was supported by NSFC (no. 21571090, 20772054), the program for Liaoning Excellent Talents in University (LJQ 2011003), and Liaohe Scholar support from Liaoning University. A. F. acknowledges the support of CINECA and PRIN (project 2010XLLNM3_007).



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