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### **PERSPECTIVE**

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# Chalcogen bonding in synthesis, catalysis and design of materials

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Chalcogen bonding is a type of noncovalent interaction in which a covalently bonded chalcogen atom (O, S, Se or Te) acts as an electrophilic species towards a nucleophilic (negative) region(s) in another or in the same molecule. In general, this interaction is strengthened by the presence of an electron-withdrawing group on the electron-acceptor chalcogen atom and upon moving down in the periodic table of elements, from O to Te. Following a short discussion of the phenomenon of chalcogen bonding, this Perspective presents some demonstrative experimental observations in which this bonding is crucial for synthetic transformations, crystal engineering, catalysis and design of materials as synthons/tectons.

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#### Introduction 1.

Noncovalent interactions were firstly taken into consideration by van der Waals in 1873, helping to revise the equation of state for real gases. In comparison to covalent bonds, intraand intermolecular noncovalent interactions are in general weak and exhibit much lower energy and directionality, as reflected by the term "noncovalent". Nevertheless, in many cases these interactions can collectively play a dominant role in the synthesis, catalysis and design of materials.



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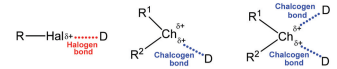
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Currently, based on the nature of the particular elements or synthons involved in the interactions, noncovalent bonds are classified into hydrogen, agnostic, anagostic,  $\pi-\pi$  stacking,  $n-\pi^*$ ,  $\pi$ -cation,  $\pi$ -anion and others. These usually lead to the formation of molecular associates or clusters with directionalities and distances specific for a particular type of interactions.  $^{2-30}$  Recently, due to the application of modern techniques, such as NMR or X-ray diffraction analysis, new types of noncovalent interactions, such as chalcogen, pnicogen, tetrel and icosagen bonds, have been identified. Although they are usually considered as weak interactions, their strength ranges from a few to several hundreds of kJ mol $^{-1}$  depending on their type.

In view of the amphoteric (electrophilic and nucleophilic) behaviour of chalcogens, the chalcogen-donor interactions have long been known as "secondary bonding interactions",  $^{31-34}$  however being recognized as a particular chalcogen (Ch) bonding in recent years. Its general structure (together with the halogen bond, for comparison) is depicted in Scheme 1, in which Ch is an electrophilic chalcogen atom (O, S, Se or Te) and D is a donor centre (electron donor atom, anion,  $\pi$ -system, radical, *etc.*). Surprisingly, the strength of chalcogen bonding is sometimes comparable with that of hydrogen bonds and other types of noncovalent interactions.  $^{35}$ 



**Scheme 1** Schematic representation of halogen and chalcogen bonds  $(R^1 \text{ is a stronger electron-withdrawing group than } R^2)$ .

Generally the formation of a chalcogen bond is explained in terms of the electrostatic potential present on the outermost portion of the chalcogen's surface, opposite to the R–Ch bond (s) (Scheme 1). This region of positive electrostatic potential is called the " $\sigma$ -hole" and refers to the electron deficient outer lobe of a p orbital involved in a covalent bond, especially when one of the atoms is highly electronegative. The cases when a region of positive electrostatic potential is perpendicular to a portion of a molecular framework, it is named the " $\pi$ -hole". Thus, due to the unequal occupation of the valence orbitals, the electrostatic potential around the chalcogen atom is anisotropic. The size of the  $\sigma$ -hole increases from the lighter to the heavier chalcogen atoms as polarizability increases and electronegativity decreases (O < S < Se < Te), and the remainder of the molecule becomes more electron-



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withdrawing. In the multivalent chalcogen atoms the respective  $\sigma$ -holes are localized on the axes of the covalent bonds (*e.g.* opposite to R<sup>1</sup> and R<sup>2</sup>, Scheme 1), <sup>39,40</sup> while the R–Ch covalent bonds elongate upon the formation of the chalcogen bond.

Since the mode of association is intimately related to the steric and electronic nature of the R substituent, modification of the R group can be used to direct the chalcogen bond assisted synthesis, catalysis and design of materials. Thus, a divalent chalcogen may be involved in multiple chalcogen bonding, as in 1,4-bis(methyltellanyl)buta-1,3-diyne, where each tellurium atom behaves as a two-electron donor and two-electron acceptor centre (Scheme 2). Each Te atom, placed at the edge of a stack, is in close contact with four Te atoms of neighbouring molecules. As a result, each Te forms *zigzag* like contacts (3.74 and 3.82 Å) to two neighbouring atoms (Scheme 2), with less than the van der Waals distances (Te···Te 4.12 Å). Leach Te distances (Te···Te 4.12 Å).

Due to the presence of the lone pairs of the chalcogen atom, a  $\sigma$ -hole is usually surrounded by a region of negative molecular electrostatic potential, and this determines a high directionality of the chalcogen bonds. This linear geometry of the R-Ch···D fragment provides a maximal electrostatic attraction between a  $\sigma$ -hole and an area of negative molecular electrostatic potential around lone pairs of the heteroatom or  $\pi$ -system and a minimal repulsion between areas of negative molecular electrostatic potential. For instance, the highly directional interactions between sulphur atoms and aromatic rings in the side chains of amino acids are crucial for the formation, stability and unique properties of  $\alpha$ -helices in proteins.  $^{43,44}$ 

Another interesting property of chalcogen bonds is their tunability, *i.e.* when their strength and geometric properties can be relatively easy modified or "tuned". <sup>45–47</sup> It should be noted that the recognized chalcogen bonds mainly involve divalent chalcogens, although they are commonly engaged in other valencies, <sup>48</sup> namely with four or six separate covalent bonds. However, a few recent works address tetravalent chalcogens. <sup>49,50</sup>

Similar to the hydrogen bond, the chalcogen bonds are classified into several fundamental types: negative charge assisted chalcogen bond (Ch···D<sup>-</sup>), positive charge assisted chalcogen bond (Ch···D), conventional (or "neutral") chalcogen bond (Ch···D) and resonance assisted chalcogen bond (RACB).  $^{40,51-56}$  Similar to the hydrogen and halogen bonds,  $^{29,57-59}$  chalcogen interactions assisted by a charge or a conjugated  $\pi$ -system (in the case of RACB) can be characterized

$$Te^{\delta^{+}} \xrightarrow{3.74 \text{ Å}} Fe^{\delta^{+}} \xrightarrow{\delta^{-}} Fe^{\delta^{+}} \xrightarrow{\delta^{-}} Te$$

$$= \frac{\delta^{+}}{\delta^{+}} e^{\delta^{+}} \xrightarrow{3.74 \text{ Å}} Te$$

$$\frac{3.82 \text{ Å}}{\delta^{-}} \xrightarrow{3.82 \text{ Å}} Fe^{\delta^{+}} \xrightarrow{\delta^{-}} Te$$

Scheme 2 Multiple chalcogen bonding in 1,4-bis(methyltellanyl)buta-1,3-diyne (one fragment is shown).  $^{41}$ 

by the enhanced strength of the chalcogen bond in comparison with the conventional chalcogen bonding. 40,51–56

There are several specific features of chalcogen bonds; 3,5,6,8,24,60,61 from those we can list the following ones. (i) Directionality: Due to the localization of a region of positive electrostatic potential on the outermost, opposite to R, part of the chalcogen atom, the R-Ch fragment elongates and "linearly" approaches D, with the typical R-Ch···D bond angle close to 180°. The stronger the chalcogen bond is, the shorter is the Ch···D distance; (ii) Tunability: The strength of the chalcogen R-Ch···D bonding can be tuned by the variation of R or D and by the replacement of the Ch atom by another one (O, S, Se or Te); (iii) Hydrophobicity: The chalcogen-bond donors are hydrophobic in comparison to conventional hydrogen bonding. In fact, positive electrostatic potential(s) on the outermost surface of the chalcogen atom is (are) hidden by the next covalent bond(s). In other words, the attached groups on the chalcogen atom prevent the solubility of organochalcogens in aqueous or polar solvents; (iv) Donor atom size: Due to the short van der Waals radii of hydrogen atoms (1.20 Å) compared to the chalcogen atoms (1.52, 1.80, 1.90, and 2.06 Å for O, S, Se, and Te, respectively),42 the hydrogen bond is less sensitive to steric hindrance than the chalcogen bond, and this causes differences in some physical and chemical properties; (v) Multiplicity: Unlike the hydrogen atom, the chalcogen atom shows variable valences, e.g., di-, tetra- and hexa-valence, which provide multiple chalcogen bonding. This allows the creation of new types of materials with unique properties, and may be used in synthetic transformations, catalysis, crystal engineering, etc.

As a result of the mentioned distinctive features, application of the noncovalent interactions became a growing area of modern chemistry, while chalcogen bonding started to be used as a driving force toward desired selectivity in synthesis. For example, the reaction of NaSPh with *syn* or *anti* hydroxy epoxide in THF afforded a mixture of the *syn-syn* and *syn-anti* lactones I and II in a ~30:70 ratio, while other two possible isomeric lactones (*anti-syn* or *anti-anti*) were not observed (Scheme 3).<sup>62</sup> The crystal structures of *syn-syn* lactones revealed two attractive 1,4-intramolecular interactions (chalcogen bonds) of divalent sulfur with both hydroxyl and carbonyl oxygens. These stabilizing noncovalent S···O interactions were

**Scheme 3** Chalcogen bond assisted stereoselective synthesis of

considered to be a driving force for this reaction and an important factor in favouring the observed stereochemical outcome.  $^{62}$ 

Furthermore, various modifications of chalcogenated ligands have been performed to drive molecular self-assembly in the synthesis of coordination compounds and to modulate their properties. <sup>29,63–68</sup> Thus, the reaction of *cis*- $[PdCl_2(CNXyl)_2]$  (Xyl = 2,6-Me<sub>2</sub>C<sub>6</sub>H<sub>3</sub>) with heterocyclic 1,3-

thiazol-2-amines or 1,3,4-thiadiazol-2-amines yields a mixture of kinetically (I with S···Cl chalcogen bond) and thermodynamically (II with S···N chalcogen bond) controlled regio-isomers of binuclear diaminocarbene complexes at room temperature and on heating, respectively (Scheme 4).<sup>67</sup> The ratio of isomers (I and II) depends on the solvent polarity and temperature (Scheme 4). According to DFT calculations, the chalcogen bond energy of the thermodynamic isomers II

Scheme 4 Difference in energy between S···Cl and S···N types of chalcogen bonds drives regioisomerization of binuclear (diaminocarbene)Pd<sup>II</sup> complexes.<sup>67</sup>

Scheme 5 Proposed mechanisms of asymmetric dihydropyranone formation via chalcogen bond involved intermediates.<sup>72</sup>

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(4.6-5.3 kcal mol<sup>-1</sup>) is higher than that of the kinetic ones I  $(2.8-3.0 \text{ kcal mol}^{-1})$  (Scheme 4).

As hydrogen and halogen noncovalent interactions, the chalcogen bond can also be used in catalysis providing an additional stabilization of intermediate(s). 65,69-72 For example, the chalcogen bond donor catalyst (2S,3R)-3-isopropyl-2-phenyl-3,4-dihydro-2*H*-benzo[4,5]thiazolo[3,2-*a*]pyrimidine (Scheme 5, black) promotes the asymmetric annulation of a range of nucleophiles such as 1,3-diketones, β-ketoesters and azaarylketones to (E,E)- $\alpha,\beta$ -unsaturated anhydrides, yielding either functionalized esters (upon ring opening), dihydropyranones or dihydropyridones with high enantioselectivity (up to 97% ee) and yields (up to 86%) via a chalcogen bond assisted intermediate.72 The interaction of the catalyst with an (E,E)-α,β-unsaturated anhydride leads to an intermediate **I**, with the carbonyl oxygen adopting a syn-conformation with respect to the isothiourea sulfur atom due to a stabilizing chalcogen bonding in the formed 5-membered ring (Route I in Scheme 5). Michael addition of diketone enolate to the Re face of I gives intermediate III, which can also be obtained by a [3,3]-rearrangement of **II**. A subsequent proton transfer/lactonisation process yields an enantioenriched dihydropyranone IV, which, upon ring opening with MeOH, produces the ester product V in high (up to 97%) ee (Routes II-V in Scheme 5).

Chalcogen bonding can play a crucial role not only in synthesis and catalysis, but also in the design of materials. For instance, chalcogen S...S bonding has been used to construct supramolecular nanotubes which are able to host other molecules (Scheme 6).41 In the case of 1,4,9,12,17,20-hexathiacyclotetracosa-2,10,18-triyne, the pre-made rings are stacked together by chalcogen bonding (Scheme 6).41 The diameter of the thus-built tubes is approximately 6 Å, the size appropriate to constrain n-hexane. The channel size is controllable by a number of -CH<sub>2</sub>- or -C=C- units (Scheme 6), as well as by replacement of S by Se or Te atoms. Accordingly, those materials can host various guests such as chlorobenzene, p-xylene, nitrobenzene, etc. 41

Apart from the above representative cases, new prospective approaches to the application of chalcogen bonding appear every day, and below we highlight some of them. In some cases the crucial role of chalcogen bonds was not recognized by authors, but we spotlight them based on the reported interatomic contacts (which should be less than the sum of the van der Waals radii) and the specific bond directionality.

### **Synthesis**

The inter- or intramolecular chalcogen bonding plays a significant role in many cases of chemical synthesis, drug design, enzyme mimetics, vapor deposition, etc.73-78 The intramolecular chalcogen bonding is classified into 1-3, 1-4, 1-5 and 1-6 types (Scheme 7) with the corresponding type of the quasi-ring involving Ch···D synthons (D = O, S, Se, Te, N, P, halogen, anion,  $\pi$ -system, radical, etc.). As expected, in most cases the Ch···D distance decreases from 1-3 to 1-6 rings, in parallel with the increase of thermodynamic stability of the corresponding chalcogen bond. Moreover, the steric energy of a molecule, which accounts for its geometry at the energy minimum, is expressed in terms of a number of contributions including chalcogen bonds. 78,79 In this section, the chalcogen bond assisted synthesis of organochalcogens and coordination compounds will be demonstrated by several examples where the Ch···D interactions play a role as a driving force and/or selectivity synthon in the course of the reaction.

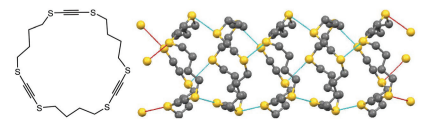
#### 2.1. Organic synthesis

Functionalized chalcophenes are key structural units in conducting polymers, microelectronic actuators, chemical sensors, electrochromic devices, organic light emitting diodes, nonlinear optics, bioactive pharmacophores, natural products, and the inter- and intramolecular chalcogen bonding can contribute to their synthetic transformations. For example, the ring-closure addition-elimination reaction of 2,3-dimethoxybuta-1,3-diene with TeCl<sub>4</sub> in the presence of CH<sub>3</sub>SiSiCH<sub>3</sub> and CH<sub>3</sub>COONa produces 3,4-methoxytellurophene stabilized by strong intermolecular chalcogen bonds (Scheme 8).80 The formation of the Te···Te bond assisted 6-membered ring in 3,4methoxytellurophene was accounted for by the metallic character of tellurium, what can be used to create new organic electronic materials.80

Another interesting motif that relies upon a 1,5 N···Se interaction was encountered in the single-crystal X-ray structure of bis{[2-(N,N-dimethylamino)methyl]phenyl}tetraselenide,



Scheme 7 Quasi-ring synthons in organochalcogens. 78,79



Scheme 6 Supramolecular nanotubes constructed from 1,4,9,12,17,20-hexathiacyclotetracosa-2,10,18-triyne.<sup>41</sup>

Scheme 8 Synthesis of 3,4-dimethoxytellurophene.80

obtained by the treatment of *N*,*N*-dimethylbenzylamine with *n*-butyllithium and elemental selenium, followed by oxidation under aqueous conditions (Scheme 9).<sup>81</sup> The formation of the two five-membered cycles with the assistance of intramolecular chalcogen bonding results in the production of the corresponding organotetraselenide compound (Scheme 9).

A 1,4 N···S chalcogen bonding appears to influence the biological activity and conformation of heterobiaryls and can also drive the reaction towards the target product. For example, the Suzuki–Miyaura coupling of 6-chloro-2-(4-fluorophenyl)imidazo[1,2-b]pyridazine with thiophen-2-ylboronic acid leads to 2-(4-fluorophenyl)-6-(thiophen-2-yl)imidazo[1,2-b] pyridazine (Scheme 10).

Other 1,4-(S···O) and 1,4-(Se···Cl) chalcogen bonding synthons were found in *N*,*N*-diisopropyl-4-methoxy-3-(methylthio) picolinamide and 3-(benzylselanyl)-4-chloro-*N*,*N*-diiso-propylpicolinamide, respectively (Scheme 11). The formation of a chalcogen bond assists the C-H bond activation in 4-substituted-*N*,*N*-diisopropylpicolinamide, while no interaction is observed between S (or Se) and the carbonyl oxygen as the oxygen atom points away from the S (or Se) centre.

$$H_3C$$
 $H_3C$ 
 $H_3C$ 

 $\begin{array}{lll} \textbf{Scheme 9} & \textbf{Synthesis} & \textbf{of} & \textbf{bis}\{[2\text{-}(\textit{N},\textit{N}\text{-}dimethylamino})\text{methyl}] \textbf{phenyl} \\ \textbf{tetraselenide.}^{81} & \\ \end{array}$ 

Scheme 10 Synthesis of 2-(4-fluorophenyl)-6-(thiophen-2-yl)imidazo [1,2-b]pyridazine.<sup>84</sup>

**Scheme 11** Synthesis of chalcogen (S and Se) derivatives of 4-chloro-and 4-methoxy-*N*,*N*-diisopropylpyridine-2-carboxamide. 85

Chalcogen bonding assisted 1,3-dipolar cycloaddition of  $N^3$ ,4-bis(4-bromophenyl)-1,2,4-thiadiazolidine-3,5-diimine with N-(4-bromophenyl)cyanamide produces (E)-1,2-bis(4-bromophenyl)-1-(3-((4-bromophenyl)amino)-1,2,4-thiadiazol-5-yl) guanidine, which contains a 1,5-(S···N) synthon with a distance of 2.538 Å (Scheme 12), suggesting that there is a strong noncovalent interaction between the electron-donating guanidine N atom and the acceptor S atom of the thiadiazole ring.  $^{86}$ 

Chalcogen bonding between oxygen atoms O···O has been observed in a large number of organic compounds, contributing to their synthesis. From those, one can consider a reaction of (2-methoxyphenyl)-bis(methylperoxy)- $\lambda^3$ -iodane with dimethyl malonate to give dimethyl 2-((2-methoxyphenyl)- $\lambda^3$ -iodanylidene)malonate stabilized by chalcogen bonding with a short O···O distance (Scheme 13). From those of the specific properties of the specific properti

Apart from the stabilization of a particular product, chalcogen bonding can increase the stability of organic radicals. <sup>89–92</sup> Thus, the reduction of 1,2,3,5-dithiadiazolyl salt [NC–(CF<sub>2</sub>)<sub>4</sub>–CNSNS]SbF<sub>6</sub> with ferrocene (FeCp<sub>2</sub>) led to the 1,2,3,5-dithiadiazolyl radical NC–(CF<sub>2</sub>)<sub>4</sub>–CNSSN which is stabilized by multiple chalcogen bonding (Scheme 14). <sup>91</sup> In fact, this compound is dimerized in the *trans*-cofacial fashion with intra-dimeric

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$$\begin{array}{c} Br \\ HN \\ S-NH \end{array}$$

Scheme 12 Synthesis of (E)-1,2-bis(4-bromophenyl)-1-(3-((4-bromophenyl)amino)-1,2,4-thiadiazol-5-yl)guanidine. 86

Synthesis of dimethyl 2-((2-methoxyphenyl)- $\lambda^3$ -iodanylidene) Scheme 13 malonate.87

chalcogen bonding S...N, and the dimeric units are also linked by S...N<sub>cvano</sub> interactions forming an infinite chain. Both S atoms in the dithiadiazolyl ring behave as a two-centred chalcogen bond donor towards the N' radical and the nitrogen atom of the cyano group (Scheme 14). When the temperature is increased from 305 to 335 K, NC-(CF<sub>2</sub>)<sub>4</sub>-CNSSN\* shows magnetic bistability with colour change from dark green (dimeric solid) to dark brown (paramagnetic liquid).

Another example is related to ortho-chalcogen substituted phenoxyl radicals stabilized by strong chalcogen bonding (Scheme 15).92 A homolytic abstraction of the hydrogen atom from the -OH group of 4-methoxy-6-methylbenzo[b]thiophen-7-ol by a peroxyl radical yields the chalcogen bond protected 4-methoxy-6-methylbenzo[b]thio-phenoxyl-7 radical (Scheme 15). 92 From this example, one can conclude that the antioxidant properties of phenoxyl radicals can be controlled with the assistance of chalcogen bonding.

Scheme 15 Stabilization of the 1,2,3,5-dithiadiazolyl radical. 92

#### 2.2. Inorganic synthesis

Exploiting the analogy with hydrogen and halogen bonds, the chalcogen bond is comparatively a new type of weak noncovalent interaction, which can be used by organic tectons to control the secondary coordination sphere of metal ions. 29,63-68 As for other types of noncovalent interactions, chalcogen bonding can be intra- and intermolecular, in coordination compounds. 29,63-68 In intramolecular chalcogenbonded coordination compounds, the coordination entities can be employed with two different roles: as donors and as acceptors.29 Thus, metal coordination may increase the electrophilicity of the chalcogen atom of a ligand which can act as the chalcogen bond donor towards other coordination entities or species.<sup>29</sup> Chalcogen bond containing coordination entities can also act as the chalcogen bond acceptor nodes through their ligands. 29,63-68 The potential of chalcogen bonding in the synthesis of coordination compounds has recently been pointed out by us (we refer to the readers),<sup>29</sup> but, in this subsection, we shall discuss more recent examples on this topic. An interesting case has already been presented above, in the

Scheme 14 Stabilization of the 1,2,3,5-dithiadiazolyl radical NC-(CF<sub>2</sub>)<sub>4</sub>-CNSSN<sup>\*</sup>.91

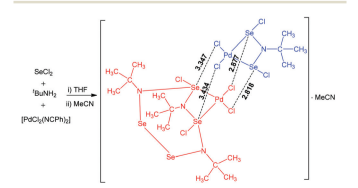
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Scheme 16 Synthesis of a macrocyclic Pd complex with intramolecular chalcogen bonds.66

Introduction section, concerning the regioisomerization of binuclear (diaminocarbene)Pd<sup>II</sup> complexes (Scheme 4).<sup>67</sup> Other examples of inorganic synthesis assisted by chalcogen bonding are as follows.

Chalcogen bonding may be involved in self-assembly of a variety of supramolecular aggregates that include cyclic tetramers and hexamers, which can act as receptors for host species such as tetrahydrofurane, Pd<sup>2+</sup>, etc. <sup>66</sup> For instance, the auto-association of [Pd(NCCH<sub>3</sub>)<sub>4</sub>](BF<sub>4</sub>)<sub>2</sub> with the negative charge assisted chalcogen bond supported tetramer of isotellurazole N-oxides in a CH2Cl2/acetonitrile mixture affords a macrocyclic Pd complex (Scheme 16).66 In this compound the metal centre displays a square planar coordination geometry with Pd-Te distances of 2.5804(4) Å, while the Te···O chalcogen bonding is 2.174 Å, being significantly shorter than the sum of the van der Waals radii of the interacting atoms, of 3.58 Å (Te + O = 2.06 + 1.52 = 3.58 Å). Thus, each of this strong intramolecular negative charge assisted chalcogen bonding closes a five-membered metallacycle leading to the Pd<sup>2+</sup> hosted macrocycle, which can contribute thermodynamically to the complexation reaction.

Multiple chalcogen bonding participates in the interaction of [PdCl2(NCPh)2] with the reaction product of SeCl2 with <sup>t</sup>BuNH<sub>2</sub> leading to a dinuclear palladium(II) adduct (Scheme 17).68 This product consists of two different palladium(II) complexes in the same crystalline lattice, linked



Scheme 17 Multiple chalcogen bonding in the synthesis of a dinuclear palladium(II) adduct.68

together by multiple Se...Cl chalcogen bonding. The coordination environment of both palladium centres in this chalcogen bond supported adduct is square-planar. The Cl-Se-N and Cl-Se-Cl angles in the range of 145.62-166.69° are in agreement with chalcogen bond formation with a moderate directionality.

#### 3. **Catalysis**

Noncovalent interactions play an important role in many catalytic cycles, e.g., offering additional stabilization to particular intermediate(s) and thus providing high selectivity in the case of asymmetric catalysis. 65,69-72,93-95 The stabilization of an intermediate depends on the types of noncovalent interactions and reaction conditions. Although individually these interactions are relatively weak, their orchestrated action can provide a powerful tool for reaching high yields and controlling selectivity.

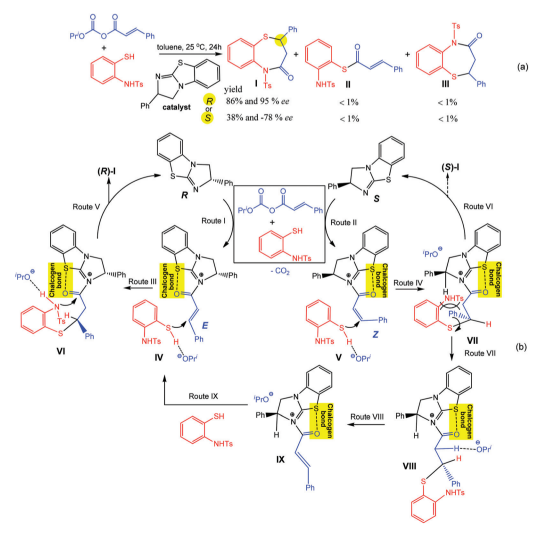
The role of noncovalent interactions, such as  $\pi$ -stacking interactions, 70 hydrogen 69 and halogen 24 bonding, in catalysis has been well recognized. Taking into account the similarities between those weak forces and chalcogen bonding, the latter should also be explored as an important tool in catalysis. Moreover, the higher directionality of chalcogen bonds may allow achieving a higher selectivity. In particular, interesting catalytic results can be expected if catalysis is performed in nonpolar media when the hydrophobic character of the chalcogen bond plays an additional role.<sup>71</sup> Although data on the use of chalcogen bond donor molecules in catalysis still remain rather scarce, 71,72,93-95 we found some interesting examples, and one of them has been discussed in the Introduction section (Scheme 5),<sup>72</sup> while others follow below.

Thus, the chalcogen bond accelerates the hydrogenation of quinolones and imine when dithieno[3,2-b;2',3'-d]thiophenes or their diimides are used as chalcogen bond donor catalysts (Scheme 18).<sup>71</sup> Cooperation of two chalcogen bonds in a specific binding motif (as "tongs") facilitates the expected substrate-catalyst interactions. According to the proposed mechanism, an intermediate is stabilized by chalcogen bonds leading to a 5-membered ring, which initiates the hydrogenation of the C=N bond in quinolines and imines (Scheme 18).

Similar to the above example of Scheme 5, a chiral isothiourea catalyst, (R)- or (S)-2-phenyl-2,3-dihydrobenzo[d]imidazo[2,1-b]thiazole, is effectively converted into a chalcogen bond supported α,β-unsaturated acylammonium intermediate by its reaction with an  $\alpha,\beta$ -unsaturated anhydride, which undergoes two sequential chemoselective nucleophilic attacks by 2-aminothiophenols producing 1,5-benzothiazepines I and a negligible amount (<1%) of by-products II and III (Scheme 19a).<sup>93</sup>

According to the proposed mechanism (Scheme 19b), the reaction of (R)- or (S)-2-phenyl-2,3-dihydrobenzo [d]imidazo [2,1-b]thiazole with a mixture of E- or Z-isomers of α,β-unsaturated anhydrides and 2-aminothiophenols goes through (i) the formation of acylammonium intermediates IV

Scheme 18 Chalcogen bond assisted hydrogenation of quinolines (Route I) and imines (Route II).<sup>71</sup>



 $\hbox{\bf Scheme 19} \quad \hbox{\bf Synthesis of 1,5-benzothiazepines (a) and proposed mechanism (b).} ^{93}$ 

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and V, stabilized by intramolecular S...O chalcogen bonding, with the liberation of <sup>1</sup>PrO<sup>-</sup> and CO<sub>2</sub> where the former behaves as a hydrogen bond acceptor towards the S-H moiety of a 2-aminothiophenol (Routes I and II in Scheme 19); (ii) a nucleophilic attack of a sulfur atom of 2-aminothiophenol to an olefin carbon atom leading to the intermediates VI and VII (Routes III and IV in Scheme 19), followed by cyclization with the formation of I (in comparison to Route V, Route VI may be inhibited by a favourable conformational change driven by the release of repulsion energy); (iii) due to unstable conformation, an intermediate VII is converted into VIII (Route VII), bearing a more stable conformation, and 2-aminothiophenol eliminates to generate the (E)-intermediate IX (Route VIII), which again affords R-1,5-benzothiazepine via intermediates IV and VI (Routes IX, III and V). In all the proposed intermediates, the chalcogen bond increases the electrophilic character of the olefinic carbon atom towards nucleophilic attack by the sulfur atom of 2-aminothiophenol, the enantioselectivity of this reaction being directed by the difference of cyclization rates between the intermediates VI and VII.

As has been mentioned above and illustrated by the above examples, the directionality and versatility of chalcogen bonding offers several possibilities for the creation and stabilization of different types of intermediates in catalytic cycles. The cooperation of chalcogen bonding with other type(s) of noncovalent interactions can further facilitate a desired reaction, while a particular orientation could enhance the selectivity. For example, cooperating chalcogen and halogen bonds are observed in the peri-substituted naphthalene (Scheme 20a) catalyzed regioselective deiodination of thyroxine (T4) to 3,5,3'-

triiodothyronine (T3) and 3,3',5'-triiodothyronine (rT3) by outer- and inner-ring deiodination pathways, respectively (Scheme 20b). 94,96 Although the formation of any I...Se halogen bond elongates (weakens) the C-I bond, a regioselective and effective bond activation begins only when the two chalcogen atoms are in the peri position in the catalysts (Scheme 20a). Thus they interact with each other leading to chalcogen bonding that facilitates the transfer of a higher electron density to the C-I σ\* orbitals through halogen bonds (Scheme 20c). As a selenol provides stronger chalcogen bonding than a thiol, replacement by selenium of either one or both sulfur atoms of the peri-naphthalene-dithiol considerably increases the yield (75-fold more) of the deiodination.

#### Design of materials 4.

Materials containing organochalcogens or coordination compounds with chalcogen bond(s) have been extensively investigated due to their unique properties, such as electric and thermal conductivities, molecular self-assemblies, nanotube formation, ion exchange capacity, molecular recognition, nonlinear optics, photoelectrochemical behaviour, drug delivery, etc. 78,97-119 All these properties are dependent on the nature of the constituent elements, directionality and tunability of chalcogen bonding, as well as the corresponding size of the quasiring involving the Ch···D synthon and planarity of this ring. All the Ch···D containing quasi-rings are almost planar and can be involved in other types of noncovalent interactions leading to better materials with enhanced device performance. 97,98

Scheme 20 Synthetic transformations of thyroxines (a) and peri-substituted naphthalenes (b). Proposed actions of halogen and chalcogen bonds in the activation of the C-I bond (c).94

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Accordingly, chalcogen bonding in organic/inorganic materials can not only provide a fundamental understanding of structure–property relationships<sup>120</sup> but also has great potential for many technological applications.<sup>27,41,117–124</sup> In this respect, switching to heavier chalcogen atoms with a larger polarizability, such as selenium or tellurium, is of particular interest since the larger atomic size may strengthen the inter- or intramolecular chalcogen bond interactions in solution and in the solid state.<sup>125</sup> For instance, many polymers reported to date are based on thiophene or its fused ring analogues, while a conjugated polymer with selenophene may exhibit a better performance in organic field-effect transistors, retaining the identical polymer backbone.<sup>126–129</sup>

The variation of substituent(s) and chalcogen bond acceptor side(s) in the monomeric unit can also significantly affect the charge transport properties of these materials. Thus, the use of selenophene spacers in the polymer poly(2,7-carbozole-*alt*-diselenienylbenzothiadiazole) (Scheme 21a) led to increased crystallinity and higher power conversion efficiency (4.1% *vs.* 3.6%) in comparison to the analogous thiophene device due to Se···Se chalcogen bonding. <sup>130</sup> Moreover, the replacement of sulfur or selenium with tellurium in conjugated polymers leads to the narrower HOMO–LUMO gap, redshifted optical absorption, higher polarizability and dielectric constant. <sup>97,131</sup>

The variation of comonomers (thiophene, selenophene and tellurophene) in cyclopentadithiophene polymer (Scheme 21b)<sup>132</sup> offers a red-shifted absorption with narrowing the band gap. At the same time, the polymer tends to aggregate in solution due to the strong Te···Te chalcogen bonding. The intensity of the absorption peak depends on the strength of the chalcogen–chalcogen interactions, *i.e.*, a polymer with a tellurophene comonomer demonstrates the most intense aggregation-induced absorption. <sup>132</sup>

As can be seen from the above examples, chalcogen bonding can control the arrangement of molecules in the solid state, and this is very important for the successful design of materials with desired magnetic, conducting, non-linear optic, *etc.* properties. Although considerable progress towards the application of hydrogen and halogen bonding for the design of new materials has been achieved, only limited examples are available regarding the related role of chalcogen bonding.<sup>62,67,113–116</sup> Hence, in this section we highlight the role of chalcogen bonding in crystal engineering and for molecular recognition.

#### 4.1. Crystal engineering

Along with the well-explored hydrogen- and halogen-bonded motifs, in recent years chalcogen bonding has also attracted significant interest in crystal engineering, drug design,<sup>78</sup> supramolecular synthons, *etc.*, as in most cases it combines two essential properties: strength and directionality.<sup>51,90,101,102,108,109,133–171</sup> The manipulation of dimensionality can be achieved by modifying the type and/or strength of chalcogen bonds present in the crystal.

A variety of chalcogen bond involved synthons with electron donor or acceptor character were employed for controlling the assembly of molecular building blocks into designed supramolecular architectures. Both types of synthons are of great importance owing to the fact that by changing the molecular geometries of organochalcogens one can manipulate the structure of crystals. Hence, an understanding of the nature of the chalcogen bonding towards its control of the molecular arrangement in the solid state can be a fundamental issue for obtaining the desired organic semiconducting or switching materials.

One of the examples demonstrating these possibilities relates to (E/Z)-2,2'-bibenzo[d][1,3]oxaselenolylidene where the inter- and intra-molecular chalcogen bonds contribute to the  $E \leftrightarrow Z$  switching in solution and to the isolation of E- or Z-isomers in the solid state (Scheme 22). The E-isomer is stabilized by multiple inter- and intramolecular noncovalent interactions, such as chalcogen (Se···O in a 1,4-membered quasi-ring) and hydrogen (C-H····Se) bonds, and two types of  $\pi$ - $\pi$  interactions ( $\pi$ phenyl- $\pi$ oxaselenole and Se- $\pi$ se····Oring). Not only intramolecular homonuclear chalcogen bonds, but also

Scheme 22 Chalcogen bonding in (E/Z)-2,2'-bibenzo[d][1,3] oxaselenolylidene. <sup>140</sup>

Scheme 21 Poly(2,7-carbozole-*alt*-diselenienylbenzothiadiazole) (a);<sup>130</sup> cyclopentadithiophene polymer with different comonomers (thiophene, selenophene and tellurophene) (b).<sup>132</sup>

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intermolecular short  $Se(2)\cdots Se(2)$  (3.449 Å,  $C_{oxaselenole}$ – $Se(2)\cdots Se(2)$  167.81°) interactions have an impact on the stabilization of the *Z*-isomer in the solid state. Thus, the chalcogen bond involved synthon can play a donor or an acceptor role in the construction of supramolecular architectures.

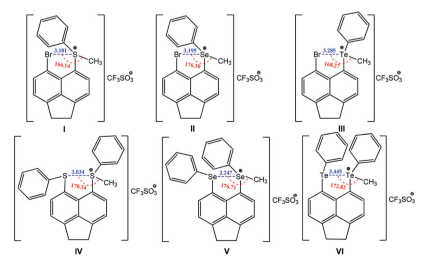
Other types of chalcogen bonding have also been identified and applied in crystal engineering. For example, a positive charge assisted chalcogen bonding synthon is found in a series of peri-substituted chalconium acenaphthene salts I-VI (Scheme 23). 154 The same equatorial-axial arrangement is observed in I-III, with the methyl group positioning along the acenaphthene plane with axial occupation of a phenyl group. The planar location of the methyl group with the acenaphthene moiety provides the "correct" geometry for the existence of a positively charged assisted chalcogen bonding with moderate directionality, 166-176° (Scheme 23). Upon switching from S to Te, the Ch+...Br distances increase in the following order: 3.181(S) < 3.195(Se) < 3.285(Te) Å (Scheme 23), which are 13-16% shorter than the sum of van der Waals radii for the two interacting atoms. A similar tendency occurs in the homonuclear positively charge assisted chalcogen bonding synthons Ch<sup>+</sup>···Ch IV-VI: 3.134(S) < 3.247(Se) < 3.445(Te)Å (16-19% shorter than the sum of van der Waals radii) and Ch<sup>+</sup>···Ch−CH<sub>3</sub> angles approaching 180° (170–177°) (Scheme 23), which may promote the delocalization of a chalcogen lone pair in a 5-membered quasi-ring. In each case the two phenyl rings are located axially to the acenaphthene plane but with either a trans (IV) or a cis (V and VI) orientation (Scheme 23).

The strength of the chalcogen...nucleophile interactions is related to the degree of charge transfer between the chalcogen bond donor and the acceptor, illustrating the importance of electrostatic interactions in the formation of a negative charge assisted chalcogen bond. Thus, a strong and directional chalcogen bonding is expected when an anion is used as a chalcogen bond acceptor. Furthermore, the negative charge assisted chalcogen bond may cooperate with other types of noncovalent

Scheme 24 Negative charge assisted chalcogen bonding in 2-(2,6-diisopropylphenyl)-4-((2,6-diisopropylphenyl)amino)-3-methylisothiazol-2-ium chloride. 148

interactions, such as hydrogen bonding or ionic interactions, 2-(2,6-diisopropylphenyl)-4-((2,6-diisopropylphenyl) amino)-3-methyliso-thiazol-2-ium chloride (Scheme 24), where cooperation of a negative charge assisted chalcogen bonding with ionic interactions and hydrogen bonding was found. 148 In this compound, the S···Cl<sup>-</sup> distance (2.848 Å) is significantly lower than the sum of van der Waals radii (S + Cl = 3.55 Å), and the N<sup>+</sup>-S···Cl<sup>-</sup> angle approaches 180° (174.82°). In the intermolecular charge assisted hydrogen bonding system, the N-H···Cl angle and the N···Cl distance are 146.05° and 3.267 Å, respectively, whereas the intramolecular distance between the Cl<sup>-</sup> anion and the N<sup>+</sup> of the thiazol ring is 4.553 Å, which falls in the recognized range of ionic interaction distances. 172 Thus, cooperation of all the above mentioned noncovalent interactions contributes to the stabilization of this organic salt in the solid state.

The chalcogen bond contributes to the stabilization of one tautomeric form of the pharmaceutical ingredient sulfamethizole (Scheme 25). <sup>159</sup> Due to the  $\pi$ -conjugation in the 5-membered quasi-ring with the formation of RACB, the tautomer III is stable in the solid state (Scheme 25). <sup>159</sup> The strengthening and directionality of RACB in III can be manipulated by cocrystalization (IV), protonation (V) or deprotonation (VI) (Scheme 24). In all cases, the O···S distance increases



Scheme 23 Positive charge assisted chalcogen bonding in *peri*-substituted chalconium acenaphthene salts. 154

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Scheme 25 RACB in molecular (IV), cationic (V) and anionic (VI) forms of sulfamethizole. 159

(2.799-2.938 Å) in comparison to the starting material II (2.762 Å), whereas the O···S-C<sub>thizole</sub> angle can decrease (in **IV-VI**) or increase (in VI) depending on the exhibited noncovalent interactions. Apparently, the participation of the RACB moiety of VI as a hydrogen bond acceptor synthon towards protonated 4-aminopyridine does not affect the O···S-C<sub>thizole</sub> angle (Scheme 25).

#### 4.2. Molecular recognition and anion transport

It is well established that noncovalent interactions are responsible for the detection of many organic, inorganic and bio-relevant molecules. In order to develop an artificial receptor (carrier or ionophore) selective towards a specific analyte, the receptor should contain a variety of synthons, organized to complement the size and shape of the analyte. Similar to the halogen bonding, 21,173-177 the relatively strong and directional chalcogen bond can also be used for selective recognition; however, quantitative data regarding the chalcogen bonding in solution are sparse. 112-115,178-183

Due to the hydrophobic character of the chalcogen bond, a desired (high) analytical signal can be expected when the molecular recognition is carried out in apolar media. Moreover, there are three possible ways for the application of chalcogen bonding in molecular recognition: (i) the receptor may have a chalcogen bonding system as a synthon, and it can interact with the analyte, or (ii) the analyte may have a chalcogen bonding system as a synthon, and it can interact with the receptor; or (iii) a chalcogen bond synthon can be formed during the analytic reaction (detection process).

Accordingly, the multiple chalcogen bond containing 2,2'-(benzo[1,2-c:4,5-c']bis([1,2,5]chalcadiazole)-4,8-diylidene) dimalononitrile receptors (I and II) (Scheme 26a) have been employed in the selective separation of p-xylene or 2,6-dimethylnaphthalene from the corresponding isomeric mix-

tures. 179 Both receptors are planar molecules and chalcogen atoms act as bifurcated chalcogen bond donor centres towards the cyano group nitrogens of neighbouring molecules with moderate directionality (167.87° N-S···N<sub>cyano</sub> and 169.49° N-Se···N<sub>cvano</sub> angles). Short chalcogen bonds connect the molecule with four neighbours (Scheme 26b), thus leading to nearly coplanar "sheetlike" networks, which form infinite layers with face-to-face overlapping between heterocycles. 179 Thus, in order to maintain the chalcogen bond assisted supramolecular architecture (Scheme 26b), these receptors selectively capture the symmetric para-disubstituted benzene rather than the less favoured ortho- or meta-disubstituted benzenes. 179 In fact, the packing diagram of I-p-xylene (analytical form) shows the existence of a similar "sheetlike" network with short S···N<sub>cyano</sub> contacts (3.22 and 3.24 Å), followed by decreasing the directionality of chalcogen bonding (137.24 and 142.82° N-S···N<sub>cyano</sub> angles). 179

A chalcogen bond assisted synthon can also be formed during the detection process. For instance, the reaction of (R)-(8-(dimesitylboryl)naphthalen-1-yl)(methyl)(phenyl) chalconium trifluoromethane-sulfonate with a fluoride anion (analyte) leads to a positive charge assisted chalcogen bond containing (R)-(8-(dimesitylboryl)naphthalen-1-yl)(methyl) (phenyl)chalconium fluoride (Scheme 27). The replacement of S by Te in the receptor increases its fluoride affinity because the larger size of Te allows for a reduction of intraligand repulsions, which generates a more favourable chalcogen bonding with the F<sup>-</sup> analyte. In fact, the Te<sup>+</sup>···F (2.505 Å) distance is shorter than the  $S^+ \cdots F$  (2.550 Å), whereas the latter is more directional (C<sub>Ph</sub>-S<sup>+</sup>···F 176.61°) in comparison to the former one ( $C_{Ph}$ - $Te^+$ ···F 174.02°).

The electron-deficient perfluoroaryl-substituted tellurophenes (I-IV) act as anion receptors through negative charge Perspective Dalton Transactions

Scheme 26 Chalcogen bonding in 2,2'-(benzo[1,2-c:4,5-c']bis([1,2,5]chalcadiazole)-4,8-diylidene) dimalononitrile receptors.<sup>179</sup>

$$\begin{array}{c} \text{Mes} \\ \text{Ce}_{\text{H}_{3}} \\ \text{Ce}_{\text{H}_{5}} \\ \text{Ch} \\ \text{S} \\ \text{CF}_{3} \\ \text{SO}_{3} \\ \text{CF}_{3} \\ \text{SO}_{3} \\ \text{CF}_{3} \\ \text{SO}_{3} \\ \text{Ch} \\ \text{S} \\ \text{CH}_{3} \\ \text{Ch} \\ \text{S} \\ \text{C} \\ \text{H}_{3} \\ \text{C} \\ \text{C} \\ \text{C} \\ \text{F}_{3} \\ \text{C} \\ \text{C$$

Scheme 27 Positive charge assisted chalcogen bonding in the recognition of  $F^-$  by (R)-(8-(dimesitylboryl)naphthalen-1-yl)(methyl)(phenyl) chalconium trifluoromethanesulfonates.  $^{181}$ 

assisted chalcogen bonding in THF (Scheme 28). <sup>113</sup> For all the anions, the bidentate chalcogen bonding in IV–X<sup>-</sup> leads to a significant increase of the association constant ( $K_a$ ) in comparison to the monodentate ones III–X<sup>-</sup> (Scheme 28), in view, in the former case, of the effect of an ethynylene linker of chalcogen-based heterocycles. In fact, both associates III–X<sup>-</sup> and

Scheme 28 Negative charge assisted chalcogen bonding in the recognition of  $\rm X^-$  by perfluoroaryl-substituted tellurophenes.  $^{113}$ 

Scheme 29 Chalcogen bond assisted anion transport. 114

IV-X<sup>-</sup> were optimized using DFT calculations, the minimumenergy geometry being stabilized by chalcogen bonding interactions having the Te····Cl<sup>-</sup> distances of 3.07 and 3.23 Å and the 168 and 170° C-Te····Cl<sup>-</sup> angles, respectively (Scheme 28).

The transmembrane transport of anions is another important direction due to various analytical and medical applications, and noncovalent interactions have successfully been employed in this field. Thus, an application of chalcogen bonding in anion transport with electron-deficient dithieno [3,2-b:2',3'-d] thiophenes (Scheme 29) has been reported. The calculated anion binding energy in 1:1 complexes increases with the enhancement of the electrophilic character of a sulfur atom in a neighbouring ring, as observed by the oxidation of the bridging S atom in I to S=O (II) and S(=O)<sub>2</sub> (III) (Scheme 29). Thus, the electrophilic character of the chalcogen bond donor centre(s) can be regulated by the addition of an electron-withdrawing group(s) to the anionophore moiety.

### 5. Conclusions

In comparison with the well-established hydrogen and halogen bonding, the chalcogen bond (a family member of Dalton Transactions Perspective

 $\sigma$ -hole interactions) is much less explored. The properties of chalcogen bonding (directionality, tunability, hydrophobicity and donor atom size) can be exploited in synthetic operations similar to those of halogen bonding, but the multiplicity of chalcogen bonding provides additional manoeuvering in synthesis and design. For instance, multiple chalcogen bonds between sulfur, selenium and tellurium centres favour the formation of nanotube-like structures in the solid state that are able to host other molecules.  $^{41}$ 

Furthermore, the chalcogen bond can direct asymmetric organic reactions, 62 stabilize organic radicals 91,92 or support thermodynamic isomers. 67 In catalysis, 5-membered chalcogen bond closed intermediates can determine the destination of the reaction. 71,72,93-95 Cooperation of the chalcogen bond with other types of noncovalent interactions, e.g. halogen bond, is particularly useful.<sup>94</sup> In materials chemistry and crystal engineering, all types of chalcogen bonding, viz. negative charge assisted, positive charge assisted, conventional (or "neutral") resonance assisted, have been employed. 140,148,154,159 Chalcogen bonds have been also used in the recognition of neutral molecules and anions, 113,179,181 as well as in anion transport. 114

Chalcogen bonding has mainly been observed in the solid state, but several studies have shown its role in solution, and this area of research is of particular importance due to potential application in analytical chemistry and medicine. Moreover, the role of chalcogen bonding (mainly S···O interactions) in biological systems is undeniable and the strategic inclusion of chalcogens in artificial drug design and in the development of synthetic transmembrane anion transporters can be provisioned.

We hope the considered examples and their discussion will attract additional attention to this exciting field of chalcogen bond assisted applications. Moreover, as for halogen bonding, <sup>61</sup> we may expect the recognition of chalcogen bonding by the International Union of Pure and Applied Chemistry (IUPAC) in the near future.

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