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Effects of Charge and Substituent on the S···N Chalcogen Bond

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ABSTRACT
Neutral complexes containing a S···N chalcogen bond are compared with similar systems in which a positive charge has been added to the S-containing electron acceptor, using high level ab initio calculations. The effects on both XS···N and XS+···N bonds are evaluated for a range of different substituents X = CH₃, CF₃, NH₂, NO₂, OH, Cl, and F, using NH₃ as common electron donor. The binding energy of XMeS···NH₃ varies between 2.3 and 4.3 kcal/mol, with the strongest interaction occurring for X=F. The binding is strengthened by a factor of 2-10 in charged XH₂S+···NH₃ complexes, reaching a maximum of 37 kcal/mol for X=F. The binding is weakened to some degree when the H atoms are replaced by methyl groups in XMe₂S+···NH₃. The source of the interaction in the charged systems, like their neutral counterparts, derives from a charge transfer from the N lone pair into the σ*(SX) antibonding orbital, supplemented by a strong electrostatic, and smaller dispersion component. The binding also derives from small contributions from a CH···N H-bond involving the methyl groups, which is most notable in the weaker complexes.

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INTRODUCTION

A chalcogen bond is formed when a member of this family of atoms (e.g. S or Se) engages in an attractive and direct noncovalent interaction with an electronegative atom like N or O \(^1\text{-}^{15}\). As such, it is closely related to the halogen \(^{16}\text{-}^{25}\) and pnictogen \(^{26}\text{-}^{35}\) bonds which have received a great deal of recent attention. There are also parallels with H-bonds (HBs), in that the interaction is at least partly due to a charge transfer from the electron donor lone pair to a \(\sigma^*\) antibonding orbital of the acceptor; the increased occupation of the latter leads to a lengthening of the corresponding internal bond within the electron acceptor unit. The importance of chalcogen bonds has been underscored by their strength, which is comparable to, and sometimes exceeds that of HBs. For instance, there is a direct interaction of S of FHS with N of NH\(_3\), forming a strong S···N noncovalent bond \(^{36}\) with a binding energy of 8 kcal/mol.

Just as the electron acceptor can be any of several chalcogen atoms, the electron donor does not have to be N, but also P, S or O \(^{37}\text{-}^{39}\) serve as other examples. Nor must the transferred charge originate in a lone pair, but it can also be extracted from various \(\pi\)-systems including simple double and triple bonds, as well as conjugated and aromatic systems. In such cases, the bonding can be reinforced \(^{37}\) by back donation in the reverse direction, from the chalcogen lone pairs into \(\pi^*\) antibonding orbitals. Effects of different substituents X on the XS···N interaction \(^{40}\) have been recently studied. It was learned that the most electronegative substituents like F and Cl enhance the interaction by the largest amount, just as electron-releasing groups such as CH\(_3\) and NH\(_2\) have the opposite effect. Regardless of the substituent, the components of binding remain the same although induction plays a more dominant role as the complexes are strengthened.

The presence of charge on either of the interacting units has been shown to have profound effect on HBs. For instance, the water dimer is bound by a OH···O HB with binding energy of some 5 kcal/mol. But the presence of either positive or negative charge on one of the H\(_2\)O molecules enhances the binding energy \(^{41}\text{-}^{43}\) by a factor of 5-8. Even much weaker HBs are strengthened by a charged subunit. One striking example arises in lysine methyl transferase enzymes where S-adenosylmethionine, containing a sulfonium cation, magnifies the usually weak CH···O HB to the point where it becomes an important part of the enzymatic mechanism \(^{44}\). Another recent study shows that the positive charge in trialkyl sulfonium and tetraalkyl ammonium enhances the \((S^+/(\text{N}^+))\)-CH···O HB energy \(^{45}\) by a factor of 4-9. The magnifying effect
of this charge remains in effect even when the \((S^+/N^+\) center and the CH groups are separated by a long hydrocarbon chain.

Given the strong effects of charge upon HBs, the obvious question arises as to whether chalcogen bonds are subject to similar changes. There are some indications in the literature that this might be the case. There are a number of indications that halogen bonds which involve an anionic electron donor can be rather strong \(46-54\), and similarly for tetrel bonds \(55\). Cationic systems are less extensively studied. In one related work, a direct \(Z\cdots\cdot N\) interaction was preferred over \(Z-H\cdots\cdot N\) H-bonds in certain cases \(56\) where the subunit containing the pnictogen atom \(Z\) was positively charged. An ion which does not itself participate in a halogen bond, but instead is positioned nearby, can nonetheless exert a strengthening influence as well \(57,58\). Data for the chalcogen bonds are rare, although a charge-assisted chalcogen bond was identified between chalcogen atoms S and Se and halide anions \(59-61\) so there is reason to believe that charge may very well strengthen these bonds. There does not appear to be anything in the literature dealing with systems in which it is the chalcogen-containing subunit that bears the charge.

The present work is aimed at filling this important gap in our understanding of the chalcogen bond. The \(S\cdots N\) interaction is the focus of our attention, particularly the effect of adding a positive charge to the electron-accepting S subunit. After first comparing \(H_3S^+\) with \(H_2S\), the H atoms are replaced by methyl groups which adds the possibility of a \(^+\)CH···N HB which could either supplement or compete with the \(S\cdots N\) interaction. These methyl groups also serve as a model of the alkyl chains that might be found in the sulfones in certain biological situations. Various substituents \(X\) (CH\(_3\), H, NH\(_2\), CF\(_3\), NO\(_2\), OH, Cl and F) are added to the electron acceptors to examine the combination of substituent effects with charge enhancement.

### COMPUTATIONAL METHODS

All calculations were performed using the Gaussian-09 package \(62\) at the MP2/aug-cc-pVDZ level of theory. This level of theory has been found to be consistent with the so-called gold standard CCSD(T) with larger basis sets, and with available experimental quantities, especially for the sorts of systems considered here \(17,28,34,63-70\). All minima were verified as having no imaginary frequencies. Binding energies were calculated as the difference in energy between the optimized complex and the sum of energies of separately optimized monomers. Basis set
superposition error (BSSE) was removed by the counterpoise correction.\textsuperscript{71} Natural Bond Orbital (NBO) analyses\textsuperscript{72,73} were performed to evaluate second-order perturbation energies E(2) using procedures contained within Gaussian within the context of the aug-cc-pVDZ basis set. Symmetry Adapted Perturbation Theory (SAPT)\textsuperscript{74,75} of the Hartree-Fock variety was used to decompose the total binding energy into several components via the Molpro\textsuperscript{76} set of codes, also using aug-cc-pVDZ.

\textbf{RESULTS}

\textbf{Neutral X(Me)S···NH\textsubscript{3} Complexes}

Neutral complexes were constructed by permitting X(Me)S to interact with universal electron donor NH\textsubscript{3}, with X pointing away from the S···N axis. NH\textsubscript{3} was chosen for this purpose for a number of reasons. In the first place, it has been used extensively for this purpose by our group as well as others, which facilitates comparison between different systems and published reports. Secondly, its lone pair provides a ready source of electrons. The presence of only a single lone pair avoids geometrical complications that would arise if there were multiple lone pairs. The absence of substituents also avoids secondary interactions that might complicate the analysis of the chalcogen bond. Finally, the small size of this molecule permits higher level and more accurate calculations to be applied. In addition to the S···N noncovalent bond, the methyl group in the electron acceptor offers a second alternative interaction; viz. a CH···NH\textsubscript{3} H-bond. The fully optimized X(Me)S···NH\textsubscript{3} geometries with various substituents X are presented in Fig 1, along with some of the most important geometrical parameters. The first two rows of Table 1 report the binding energies both without and with counterpoise correction. This quantity increases in the order of X=NH\textsubscript{2}<CH\textsubscript{3}<H<OH<CF\textsubscript{3}<Cl<NO\textsubscript{2}< F and ranges from 2.32 kcal/mol for X=NH\textsubscript{2} to 4.42 kcal/mol for X=F. The S···N distance varies between 2.792 and 3.308 Å. The XS···N arrangement is within 4° of linearity in all cases.

As indicated above, these structures are stabilized primarily by two possible attractive forces. Charge transfer from the N lone pair can flow to either the S-X σ* antibonding orbital or to a methyl C-H σ*. In either case, this transfer will be accompanied and revealed by a number of factors. Transfer into an antibonding orbital should lengthen the relevant bond, and NBO analysis would quantify this stabilizing force via second-order perturbation energy E(2), as well as the amount of charge transferred Δq, and the change in occupancy of the pertinent antibonding orbital, Δocc.
These quantities are all presented in Table 1 and exhibit some interesting trends. The binding energies are depicted by the black line in Fig 2. The broken blue line represents E(2) for the S···N interaction which is very small on the left, but grows very rapidly from left to right. The CH···S HB obeys a nearly opposite pattern, as E(2) is fairly large on the left side of Fig 2, and diminishes by only a small extent as the total bonding is enhanced (see red broken curve). These trends in E(2) are echoed by Δq and Δocc (see Table 1). One may conclude that there is a CH···N HB present in all of these complexes that is relatively immune to the nature of X, but that the growing strength of the binding in the complex can be traced primarily to the S···N bond.

Comparison with Neutral XHS···NH₃ complexes

Previous calculations ⁴⁰ have extracted comparable information for complexes wherein the methyl group is replaced by the simpler H atom. The optimized geometries are similar to those in Fig 1 with some small differences. For one thing, the R(S···N) distances are a bit shorter for XHS···NH₃, and the binding energies somewhat larger, particularly for the more strongly bound complexes on the right side of Table 1. Given the fact that the methyl groups provide a second binding option, of a CH···N as well as a S···N bond, this distinction might seem contradictory. The underlying source of this difference may be traced to the electrostatic potentials of the XHS vs XMeS subunits. Whereas the major lobe of the positive region around XHS lies directly along the X-S axis, this lobe is redirected more toward the methyl group in XMeS, as illustrated in Figs 3a and 3b for X=H and X=F, respectively. It is for this reason that the C₃ axis of the NH₃ molecule, corresponding to the N lone pair, is turned up toward the methyl group in most of the structures of Fig 1, whereas this lone pair is more closely aligned with the S···N axis ⁴⁰ for XHS complexes. This misalignment in the XMeS geometries results in a reduced charge transfer energy from the N lone pair to the SX σ* orbital. For example, while E(2) for Nlp→σ*(S-X) ranges to as high as 28 kcal/mol for FHS···NH₃, the corresponding quantity in FMeS···NH₃ is only 9 kcal/mol. The additional E(2) for the CH···N HB in FMeS···NH₃ cannot compensate for this difference, as it only amounts to about 1 kcal/mol. As a consequence the interaction energy in FMeS···NH₃ is only about half that in FHS···NH₃. The importance of the different electrostatic potentials of FHS and FMeS is verified by a decomposition of the interaction energies in the two corresponding complexes, in that the electrostatic component of FMeS···NH₃ is only about half that of FHS···NH₃.
XH$_2$S$^+$···NH$_3$ Complexes

Adding a proton to each of the XHS subunits leads to a positively charged XH$_2$S$^+$ and facilitates an examination of the effect of such a charge upon the properties of the various complexes. The optimized geometries in Fig 4 show that the X substituent lies some 162°-171° from the N atom, very similar to their neutral XHS···NH$_3$ analogues. On the other hand, the charged complexes are much more strongly bound. Whereas the binding energies of the neutrals vary from 1.5 to 7.9 kcal/mol, the charged complexes range between a minimum of 12.9 kcal/mol up to 36.7 kcal/mol. On the weaker side of the spectrum, neutral H$_2$S does not form a true minimum to NH$_3$ bound by a S···N bond. In contrast, H$_3$S$^+$, HMeS$^+$, and HMe$_2$S$^+$ all engage in a strong bond of this type, bound by 17.6, 15.0, and 12.7 kcal/mol, respectively. The latter charged S··N bonds are stronger than any of the neutral analogues. On the strong end, the neutral FS··N bond of 7.9 kcal/mol is magnified by a factor of 4.6 when charged. This magnification is even more pronounced for X=NH$_2$ and CH$_3$ where the charged complex is bound more strongly than the neutral system by a factor of 8 and 10, respectively.

Table 2 reports the various energetic and geometric properties of the charged complexes, where it may be observed that the R(S··N) distance varies from 2.19 to 2.87 Å, as compared to the 2.47 - 3.25 Å range for the neutrals, a charge-induced contraction of roughly 0.3 Å. The N$_{lp}$$rightarrow$$\sigma^*(SX)$ charge transfer is also enhanced, with E(2) varying from a minimum of 6.8 kcal/mol up to more than 66 kcal/mol. One notes also large charge transfers as measured both by $\Delta q$ and $\Delta occ$, both up around 150-170 me for the stronger bonds. The charge transfer into the antibonding S-X orbital lengthens this bond in most cases, by as much as 66 mÅ. (Exceptions are discussed below.)

Given the fact that a positive charge would enhance the acidity of the XSH$_2^+$ species, it might be tempting to suggest that there are SH$^+$···N HBs present in these ionic complexes that add to the binding energy. However, the R(H··N) distances are rather long, all exceeding 2.4 Å, and the $\theta$(SH··N) angles very far from linearity. Nor does the NH$_3$ molecule rotate its lone pair up toward the SH protons to any significant degree. The absence of significant NBO E(2) values for N$_{lp}$$rightarrow$$\sigma^*(SH)$ further confirms the absence of any such HBs.

SAPT decomposition of the ionic XH$_2$S$^+$···NH$_3$ complexes reported in Table 3 shows a strong enhancement in each of the components when compared to the neutrals. For example, ES for the neutrals ranged up to a maximum of -24 kcal/mol in FHS··NH$_3$, and this same quantity
is equal to -64 kcal/mol for FH$_2$S$^+$$\cdots$NH$_3$. The induction energy is even more enlarged, from -34 to -109 kcal/mol. Indeed, whereas ES and IND are roughly equal for the neutral complexes, the latter quantity exceeds ES for the more strongly bound cationic systems by a sizable margin. Even though DISP is enhanced by the charge, it remains considerably smaller than the two other components mentioned above, roughly 15-30% of their magnitudes. SAPT decompositions for all complexes are displayed in Tables S1-S3 of the Supplementary Information. It may be seen that the same enhancement effects are observed as well for the methyl-substituted complexes.

**X(Me)$_2$S$^+$$\cdots$NH$_3$ Complexes**

Evaluation of X(Me)$_2$S$^+$ as electron acceptor may be considered in one of two ways. It may be thought of as either the addition of a positive charge to XMeS, or as the replacement of the two H atoms of XH$_2$S$^+$ by methyls, with no change in charge state. The optimized X(Me)$_2$S$^+$$\cdots$NH$_3$ complexes are illustrated in Fig 5, with other details provided in Table 4. Comparison with Fig 4 and Table 2 reveals first that the replacement of the H atoms of XH$_2$S$^+$ by methyls weakens the S$\cdots$N bond by a surprising amount. For example, the binding energy of FH$_2$S$^+$$\cdots$NH$_3$ is cut in half by this disubstitution, although the diminutions in the other systems are less dramatic. Along with this weakening, there is a concomitant stretch of the R(S$\cdots$N) distance by some 0.2-0.3 Å and the $\theta$(XS$\cdots$N) angle becomes a bit more linear. There is a sizable drop in the N$_{lp}$$\rightarrow$$\sigma^*$(SX) charge transfer, as measured by E(2), $\Delta q$, and $\Delta$occ, as well as a diminished S-X bond stretch. (A primary reason for the reduction in molecular interaction strength arising from methyl substitution is elaborated below.)

Unlike XH$_2$S$^+$, the methyl groups of X(Me)$_2$S$^+$ extend toward the N atom, to the point where they seem capable of forming SCH$^+$$\cdots$N HBs, albeit weak ones. The characteristics of these HBs are listed in the lower portions of Table 4, where bond lengths are in the range of 2.6 - 2.7 Å. The $\theta$(CH$\cdots$N) angles are substantially distorted from linearity, but there is nonetheless significant N$_{lp}$$\rightarrow$$\sigma^*$(CH) charge transfer, as evident by E(2), $\Delta q$, and a contraction of r(CH) which is characteristic of many CH$\cdots$N H-bonds. Although generally weaker than the complexes with XH$_2$S$^+$, these X(Me)$_2$S$^+$ ions form much stronger interactions with NH$_3$ than do their neutral XMeS counterparts, by a factor of 3-5, with corresponding increases in the various indicators of charge transfer from NH$_3$ to $\sigma^*$(S-X). It is intriguing to note that despite the positive charge on the proton donor X(Me)$_2$S$^+$, there is what would appear to be a weakening of the CH$\cdots$N HBs relative to neutral XMeS, in terms of E(2) and $\Delta q$, as well as geometrical indicators R(H$\cdots$N) and
\( \theta(\text{CH} \cdot \cdot \text{N}) \). In summary, the addition of positive charge on the S-electron acceptor very substantially strengthens the S \cdot \cdot N bond, at the expense of a small reduction in CH \cdot \cdot N HB strength.

In terms of accuracy, one may always question the validity of a given level of theory. As mentioned earlier, the MP2/aug-cc-pVDZ level has proven its fidelity on a number of occasions in the past. In order to check its accuracy on the particular systems examined here, correlation was included via the CCSD(T) approach, using the larger aug-cc-pVTZ basis set for a representative sampling of systems. As may be seen in Table 5, complexes were chosen that were both neutral and cationic, methylated and unmethylated, using both H and F as the two substituents which represent opposite extremes of binding strength. The comparisons in Table 5 make it clear that there is little difference between the MP2/aug-cc-pVDZ and CCSD(T)/aug-cc-pVTZ binding energies. Values are all within 7\% of one another, some differences less than 1\%.

**Competition between CH\cdot\cdotN and S\cdot\cdotN interactions**

As described above, the CH\(_3\) groups in X(Me)S and X(Me)\(_2\)S\(^+\) units offer the N lone pair an alternative with which to interact. That is, this lone pair may donate charge to the S-X \( \sigma^* \) orbital in a S \cdot \cdot N bond, or may interact instead with a CH bond of a methyl group, donating charge into its \( \sigma^*(\text{CH}) \) antibonding orbital. The orientation of the NH\(_3\) molecule is heavily influenced by this competition, as the former is favored when the lone pair of NH\(_3\) lies along the S \cdot \cdot N axis, while a displacement upward toward a methyl group would facilitate the CH \cdot \cdot N HB. One can measure this displacement with the simple assumption that the lone pair is collinear with C\(_3\) symmetry axis of NH\(_3\).

For complexes without methyl groups, XH\(_2\)S\(^+\) and neutral XHS in which there is no CH\cdot\cdotN bond, the lone pair lies within 6-9° of the S \cdot \cdot N axis. In contrast, the methylated systems vividly illustrate the competition between the S \cdot \cdot N and CH \cdot \cdot N interactions. For those complexes containing the neutral X(Me)S, the N lone pair is displaced from S \cdot \cdot N by a large amount, 66° for HMeS, facilitating the CH \cdot \cdot N bond. But as the substituent becomes progressively more electron withdrawing, and as the S \cdot \cdot N interaction strengthens, this deviation diminishes, dropping down below 30° for FHS \cdot \cdot NH\(_3\) and 18° for FMeS \cdot \cdot NH\(_3\). A similar pattern is observed in the dimethyl FMe\(_2\)S\(^-\) cases, although these deviations are systematically smaller here. These angular characteristics affirm the idea that CH \cdot \cdot N HBs outweigh S \cdot \cdot N for the more weakly bound complexes, but it is S \cdot \cdot N which dominates as the interaction grows in strength.
Another important factor in controlling the NH$_3$ orientation is the electrostatic potential around the electron acceptor molecule. Besides acting as the principal source of electron density for transfer, the NH$_3$ lone pair is also coincident with the most negative potential around this molecule. As such, this lone pair ought to be drawn toward the most positive region of the potential around the partner molecule. As was mentioned earlier, the replacement of a H atom in XHS by a methyl group brings the most positive region of the electrostatic potential up toward the methyl group, and with it the N lone pair. This upward displacement of the most positive portion of the potential for both HMeS and FMeS are evident by the blue regions in Fig 3a and 3b. When a second methyl group is added, leading to the XMe$_2$S$^+$ cations, however, the displacement is much smaller in that the blue regions in Fig 3c and 3d lie nearer to the S···N axis, resulting in the smaller upward rotations of the NH$_3$ molecule in these dimethylated systems, none of which are more than 20°.

The electrostatic potentials also offer a ready explanation for the drop in binding energy when XH$_2$S$^+$ is replaced by its dimethylated XMe$_2$S$^+$ analogue. Comparison of Fig 3c with 3e shows a strong increase in the positive electrostatic potential to the right of the S atom when the methyl groups of HMe$_2$S$^+$ are both replaced by H. This same effect arises when the F atom is the substituent, as in Figs 3d and 3f.

The shifts of electron density that accompany the formation of a complex can frequently provide insights about the nature of the interaction. These shifts are displayed in Fig 6 as the difference in density between the complex on one hand, and the sum of isolated monomers on the other hand, in the same positions as in the complex. Purple areas indicate an increased density in the complex, and losses are represented by the green regions. Focusing first on the neutral complexes of Figs 6a and 6b, both complexes show a purple gain of density in the region of the N lone pair. The green loss around the methyl proton in Fig 6a is characteristic of H-bonds. (There is no such density loss around the SH proton of the unmethylated counterparts, verifying the absence of a SH···N HB.). One may note the absence of a green region to the right of S in Fig 6a, consistent with the idea that MeHS is bound to NH$_3$ primarily by a CH···N HB. The green density-loss region encompasses both the S and CH proton in FHS···NH$_3$ in Fig 6b, verifying the growing strength of the S···N bond. A gain of density appears also around the F substituent, consonant with the idea of charge transfer into the S-F σ* orbital.
It is immediately clear that there is much greater density rearrangement in the charged complexes in Figs 6c and 6d than in their neutral counterparts. (Note also the larger value of the isocontour used for the ionic complexes.) This enhancement is consistent with the stronger binding in the ionic complexes, and the idea that a charged species can more powerfully affect the charge distribution within its partner. In these ionic complexes, the green charge-loss region to the right of S dwarfs any density changes around the CH protons, further attesting to the dominance of S···N over CH···N interactions in charged dimers.

**Anomalous Behavior of NO$_2$ and CF$_3$ Substituents**

The reader has probably noticed that complexes containing the NO$_2$ group have behaved differently than most of the others. For example, the last column of Table 1 shows that even though there is a sizable charge transfer into the S-N σ* orbital of (NO$_2$)MeS··NH$_3$, the net occupancy of this orbital drops by 2 me, and the S-N bond shortens by 5 mÅ; these anomalous effects are magnified in the ionic (NO$_2$)H$_2$S$^+$··NH$_3$ and (NO$_2$)Me$_2$S$^+$··NH$_3$ complexes in Tables 2 and 4, respectively. Indeed, similar contrary behavior, albeit not as large in magnitude, is observed for the -CF$_3$ substituent. The anomalous behavior cannot be attributed to any conformational change within the monomer in question, as the overall geometry remains unchanged upon complexation.

What these two substituents have in common is the presence of more than one non-hydrogen atom. And it would appear it is this feature which permits this counterintuitive behavior. In particular, charge is indeed transferred from the N lone pair to the S-N σ* antibonding orbital of (NO$_2$)H$_2$S$^+$, as evident from the NBO analysis. However, this charge does not remain there, continuing on to the O lone pairs in the case of NO$_2$. Not only this charge, but also a certain amount of density that originally resided in the N-S σ* orbital in the monomer moves over to the O lone pairs as well, making the O atoms more negatively charged. The net decrease in σ* population is reflected in the values of Δocc, as well as the shortening of the S-N bond. These same principles are in operation within the -CF$_3$ substituent wherein charge is transferred from the S-C σ* antibonding orbital to the F lone pairs. And in both the NO$_2$ and CF$_3$ substituent cases, the increased density in the peripheral atom lone pairs is visible when viewing the density difference maps.
SUMMARY

The effects of methylation and of addition of a charge are summarized in Fig 7 which displays the total binding energy for the four sorts of systems, and how this quantity varies as the substituent is changed. For each type of interaction, whether neutral or charged, the interaction grows stronger as the substituent becomes more electron-withdrawing (with the exception of X=NO$_2$). The two neutral types containing either XHS or XMeS are generally similar to one another, although the former is a bit stronger for the more electronegative groups on the right side. The ionic complexes are much more strongly bound, in particular XH$_2$S$^+$. For example, FH$_2$S$^+\cdots$NH$_3$ is bound by some 37 kcal/mol, which represents a nearly five-fold magnification compared to its neutral FHS$\cdots$NH$_3$ analogue. Accompanying this charge-induced strengthening is a reduction in the intermolecular separation by some 0.5 Å on average.

The total binding energy is strongly correlated with the charge transfer from the N lone pair into the S-X σ* antibonding orbital, as verified by a number of markers, including E(2), the change in occupation of the σ* orbital, and the stretch of the S-X bond length. For example, the correlation coefficient relating E(2) to the binding energy is equal to 0.91 for complexes involving XMeS, 0.94 for XMe$_2$S$^+$, and 0.80 for XH$_2$S$^+$. The induction energy is comparable to, and in some cases larger than, the electrostatic interaction energy. The dispersion energy is considerably smaller, but cannot be ignored, amounting to as much as 11 kcal/mol in the most strongly bound system.

In addition to the S$\cdots$N bond, there is a certain amount of CH$\cdots$N H-bonding involving a methyl group on the S atom. This bond makes a fixed, but small, contribution to the total interaction energy, fairly insensitive to the nature of other substituents on the S. Consequently, the weakest complexes rely to a large degree on the CH$\cdots$N HB for their stability, whereas it is S$\cdots$N which predominates for the stronger complexes. On the other hand, the replacement of a H atom on S by a methyl group displaces the positive portion of the electrostatic potential of the monomer away from the S atom. This displacement tends to draw the NH$_3$ lone pair away from the S-X σ* orbital, thereby weakening the interaction.

The NO$_2$ substituent behaves somewhat differently than the other substituents in that the charge the monomer acquires from the NH$_3$ lone pair does not remain in the σ* S-N antibonding orbital, but continues on into the O lone pairs. It is for this reason that the S-N bond contracts, and the O atoms become more negatively charged. This anomalous behavior also accounts for
the drop in binding energy for the NO₂ substituent in comparison to those of comparable
electron-withdrawing potency (see Fig 7).

It might be noted that not all of the minima studied here represent the global minimum on the
_corresponding potential energy surface. The OH substituent for example, is such a strong proton
donor that a OH··N H-bond will typically be energetically preferable to a S··N interaction. And
since NH₃ is a more powerful proton acceptor than is XHS, an additional proton would prefer the
former site, resulting in a XHS···NH₄ complex, rather than XH₂S···NH₃. But all of the
structures considered here represent true minima nonetheless. The need to study complexes of
this type is underscored by examples of biological systems where a sulfonium cation is an
essential component in the mechanism of certain enzymes. Systems such as these also illustrate
the importance of replacing SH protons by a methyl group in the model systems, to better
represent these biological situations.

In summary, the cationic S⁺···N interaction represents a very strong force, several-fold
greater than its neutral counterparts or uncharged hydrogen and halogen bonds. The addition of a
charge does not fundamentally alter the nature of bonding, affecting primarily the magnitudes of
the components. Nor does the charge alter the ratio of electrostatic and induction components of
the total interaction energies.

SUPPORTING INFORMATION
SAPT dissections of complexes and optimized coordinates of complexes. This material is
available free of charge via the Internet at http://pubs.acs.org

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(75) Moszynski, R.; Wormer, P. E. S.; Jeziorski, B.; van der Avoird, A. Symmetry-Adapted Perturbation

(76) Werner, H.-J.; Knowles, P. J.; Knizia, G.; Manby, F. R.; Schütz, M.; Celani, P.; Korona, T.; Lindh,
R.; Mitrushenko, A.; Rauhut, G.; Shamansundar, K. R.; Adler, T. B.; Amos, R. D.; Bernhardsson,
May, A. J.; McNicholas, S. J.; Meyer, W.; Mura, M. E.; Nicklass, A.; O'Neill, D. P.; Palmieri, P.;
Wang, M.; Wolf, A. Molpro, Version 2010.1, a Package of Ab Initio Programs.
Table 1. Energetic (kcal/mol) and geometrical properties of X(Me)S···NH₃ complexes

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<tr>
<th>X</th>
<th>H</th>
<th>CH₃</th>
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<th>CF₃</th>
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<td>1.87</td>
<td>1.77</td>
<td>1.23</td>
<td>2.08</td>
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a) counterpoise correction
b) change in X-S bond that accepts charge into σ* antibond
c) computed as 2*(F/Δε)²
d) Change of occupation of S-X σ* orbital

Table 2. Energetic (kcal/mol) and geometrical properties of XH₂S⁺···NH₃ complexes

<table>
<thead>
<tr>
<th>X</th>
<th>H</th>
<th>CH₃</th>
<th>NH₂</th>
<th>CF₃</th>
<th>OH</th>
<th>Cl</th>
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<td>66.1</td>
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<td>88.34</td>
<td>110.44</td>
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Table 3. SAPT energy decomposition (kcal/mol) for XH$_2$S$^+$···N interaction

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<th>EX</th>
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Table 4. Energetic (kcal/mol) and geometrical properties of X(Me)$_2$S$^+$···NH$_3$ complexes

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<th>X</th>
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<th>NH$_2$</th>
<th>CF$_3$</th>
<th>OH</th>
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<td>13.75</td>
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<td>37.3</td>
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<td>4.66</td>
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<td>14.55</td>
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<td>Δq$^\text{d}$, me</td>
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<td>Δr(C-H), mÅ</td>
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<td>E(2) N$^\text{lp}$→σ*(CH)</td>
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$^1,^2$ for two different CH$_3$ groups
Table 5: Counterpoise-corrected binding energies (kcal/mol) of some representative complexes with NH$_3$ computed at the CCSD(T)/aug-cc-pVTZ level of theory. MP2/aug-cc-pVDZ values are shown for comparison.

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<th>XMe$_2$S$^+$</th>
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<td>F</td>
<td>7.52</td>
<td>7.92</td>
<td>36.77</td>
<td>36.71</td>
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</table>

$\Delta E + CC$, kcal/mol
Figure Captions

Fig 1. Optimized geometries of X(Me)S···NH₃ complexes for X= a) H, b) CH₃, c) NH₂, d) CF₃, e) OH, f) Cl, g) F, and h) NO₂. Distances are in Å and angles are in degrees. Counterpoise-corrected binding energies in kcal/mol are represented by blue numbers.

Fig 2. Total binding energies -ΔE and NBO values of E(2) for X(Me)S···NH₃ complexes. E(2) S-X refers to Nlp→σ*(SX) and E(2) C-H to Nlp→σ*(CH).

Fig 3. Electrostatic potential maps of monomers a) HMeS, b) FMeS, c) HMe₂S⁺, d) FMe₂S⁺, e) H₃S⁺ and f) FH₂S⁺, on a surface lying twice the vdW radius of each atom. Red and blue regions of a and b indicate -0.015 and +0.015 au, respectively, while they refer to +0.015 and +0.018 au in the cations in c-f.

Fig 4. Optimized geometries of XH₂S⁺···NH₃ complexes for X= a) H, b) CH₃, c) NH₂, d) CF₃, e) OH, f) Cl, g) F, and h) NO₂. Distances are in Å and angles in degrees. Counterpoise-corrected binding energies in kcal/mol are represented by blue numbers.

Fig 5. Optimized geometries of X(Me)₂S⁺···NH₃ complexes for X= a) H, b) CH₃, c) NH₂, d) CF₃, e) OH, f) Cl, g) F, and h) NO₂.

Fig 6. Electron density shifts occurring upon complexation of NH₃ with a) HMeS, b) FMeS, c) HMe₂S⁺, and d) FMe₂S⁺. Purple and green regions indicate gain and loss of density, respectively, in dimer relative to monomers. The 0.001 au contour is depicted in a and b, 0.002 au in c and d.

Fig 7. Variation of binding energy (kcal/mol) of neutral and cationic S···N complexes with various substituents.