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Non-bonding pairs in cyclic thioethers: Electrostatic modelling and *ab initio* calculations for complexes of 2,5-dihydrothiophene, thietane and thiirane with hydrogen fluoride

J. Grant Hill¹ and Anthony C. Legon²

¹Department of Chemistry, University of Sheffield, Sheffield S3 7HF, U.K.,
grant.hill@sheffield.ac.uk

²School of Chemistry, University of Bristol, Cantock's Close, Bristol BS8 1TS, U.K.,
a.c.legon@bristol.ac.uk

Abstract.

Locations of nucleophilic regions of Lewis bases play important roles in the geometry of intermolecular complexes and can be probed by small molecules with an electrophilic region (e.g. HF). The electrostatic potential energy $V(\phi)$ of a non-perturbing, protonic charge at a fixed distance r from the S atom in three cyclic thioethers was examined as a function of the angle ϕ made by the r -vector with the C_2 axis (in thiirane and 2,5-dihydrothiophene) or the local C_2 axis associated with the C–S–C plane of the thietane ring. The electrostatic PE $V_{\text{HF}}(\phi)$ of HF, when HF is modelled as an extended electric dipole, was also calculated as a function of ϕ and the results compared with the geometries of the complexes thiirane \cdots HF, thietane \cdots HF and 2,5-dihydrothiophene \cdots HF calculated at the CCSD(T)-F12c/cc-pVTZ-F12 level. The calculations reveal angular deviations $\theta \sim 10\text{-}20^\circ$ of the S \cdots H–F nuclei from collinearity in a direction suggesting secondary interactions of the nucleophilic region of F with electrophilic H atom(s) of the rings. They also reveal the angles ϕ made by the S \cdots H hydrogen bond with the C_2 (or local C_2) axes in the thioether \cdots HF complexes are systematically larger ($\sim 4\text{-}9^\circ$) than those at the minima in the $V_{\text{HF}}(\phi)$ functions, but the latter do agree with the directions of maximum nucleophilicity shown by MESP diagrams of the thioethers. The deviation between these geometries can be explained with a simple model for complex formation. In the simple electrostatic model, the minima $|\phi_{\text{min}}|$ in the $V(\phi)$ versus ϕ functions occur at values smaller ($\sim 5\text{-}10^\circ$) than those in the $V_{\text{HF}}(\phi)$ curves.

1. Introduction.

Non-covalent interactions perform a vital role in numerous chemical and biological processes, from small gas-phase complexes through to the structure, dynamics and activity of macromolecules, including DNA. Several of these non-covalent interactions, including the hydrogen bond and the halogen bond, have been

formally defined by IUPAC working parties,^{1,2} and simple models can be applied to account for the bond. The basis of the simple model is that an electrophilic region of the molecule or functional group acting as a Lewis acid (the hydrogen atom in a hydrogen bond, or the halogen atom in a halogen bond) interacts with the nucleophilic region of a Lewis base B (typically non-bonding or π -bonding electron pairs). The terminology σ -hole or π -hole is often attached to the electrophilic region,^{3,4} and the simple models provide an almost intuitive methodology for predictions of the angular geometries of supramolecular complexes. It is important to note that simple models typically sacrifice some of the subtleties of the underlying nature of the interactions, which has led to great and occasionally heated debate surrounding, for example, the role of charge transfer / degree of covalency within these bonds.⁵⁻¹¹

The relative orientations of hydrogen-bonded complexes in the gas-phase has been the subject of a number of systematic studies, both experimentally (by microwave and infrared spectroscopy) and theoretically, see refs. 9,12-14, for examples and further references. In particular, the use of hydrogen fluoride as a probe molecule has allowed for explorations of the nucleophilic regions of Lewis bases that act as hydrogen bond acceptors. Extending the choice of probe molecules to encompass HX, where X is a halogen atom, CN group or $C\equiv CH$, led to the formulation of simple rules for gas-phase geometries of complexes.^{12,13} A key component of these geometries is the directionality of the interaction, which can be expressed in terms of the angle ϕ (see Figure 1). These angular geometry prediction rules have been demonstrated to be electrostatic in origin and thus can be tested by careful examination of the electrostatic potential, and its angular dependence, in the region surrounding the atom or centre of the Lewis base directly involved in the hydrogen or halogen bond.¹³ Similar arguments apply to the Lewis acid partner in the complex, and logically the sensitivity of the strength of the interaction to the directionality has been linked to the location (or absence of) non-bonding electron pairs in the valence shell of the hydrogen or halogen in the Lewis acid.¹⁵

One of the most suitable Lewis bases to consider in terms of the angular dependence of the electrostatic potential is the water molecule. Water and its congener hydrogen sulphide form hydrogen-bonded complexes $H_2O\cdots HX$ ¹⁶⁻²² and $H_2S\cdots HX$ ²³⁻²⁷ with the hydrogen halides HX (X = F, Cl, Br, I) in the gas phase. They have all been investigated by means of their rotational spectra. Of these, only $H_2O\cdots HF$ could be detected in equilibrium gas mixtures of water and the hydrogen halide at moderately low temperatures (~ 200 K), under which conditions vibrational satellites associated with low-frequency intermolecular modes were observed. The satellites correspond to the same rotational transition but in vibrationally excited states and their analysis allowed, *inter alia*, the potential energy function associated with the low-frequency intermolecular bending mode to be determined.¹⁸ This function, which is shown in Figure 1(a) (blue curve and labels) together with energy levels of the low frequency, intermolecular bending mode deduced from it, is of the double-minimum type having a low barrier at the planar configuration and two minima

corresponding to equivalent pyramidal arrangements at O. The barrier to inversion lies below the zero-point energy level of the molecule. Thus, although the configuration at O is pyramidal at equilibrium, the molecule is effectively planar in the zero-point state (i.e. the vibrational wavefunctions have C_{2v} symmetry). The separation between the $\nu = 0$ and 1 states associated with the inversion mode is large relative to rotational energy level spacings. Other molecules $\text{H}_2\text{O}\cdots\text{HX}$ ^{19–22} were studied in the vibrational ground state by pulsed-jet, Fourier-transform microwave spectroscopy, but the equilibrium geometry was deduced by various observations to be again pyramidal at O, with a low barrier to inversion, as confirmed by *ab initio* calculations. A similar conclusion obtains for several halogen-bonded complexes $\text{H}_2\text{O}\cdots\text{XY}$ ^{28–34} (XY is a homo- or hetero-nuclear di-halogen molecule). Equilibrium values ϕ_e were estimated to lie in the range 35–55° from the rotational constants of each $\text{H}_2\text{O}\cdots\text{HX}$ and $\text{H}_2\text{O}\cdots\text{XY}$ investigated

The situation for $\text{H}_2\text{S}\cdots\text{HX}$ complexes is quite different.^{23–27} Although studied only in the zero-point vibrational state in a supersonic expansion at ~ 2 K, it was possible to deduce that the configuration at S is rigidly pyramidal on the microwave timescale in all but one case, with undetectable inversion splitting between the zero-point state and the first vibrationally excited state and therefore a high barrier to the inversion motion. This conclusion was again confirmed by *ab initio* calculations. The exception was $\text{H}_2\text{S}\cdots\text{HI}$,²⁷ which also exhibited a rotational spectrum in a low-lying vibrationally excited state even at ~ 2 K and therefore showed evidence of conformational non-rigidity. Angles ϕ_e were found to be close to 90° in all other cases investigated. Similar results, with $\phi_e \sim 90^\circ$ and conformational rigidity, were established for several halogen-bonded complexes $\text{H}_2\text{S}\cdots\text{XY}$.^{35–40}

The different experimental conclusions for $\text{H}_2\text{O}\cdots\text{HX}$ and $\text{H}_2\text{S}\cdots\text{HX}$ complexes suggest that the equilibrium conformations can be predicted by assuming that in the equilibrium geometry the HX molecule lies along the axis of one of the non-bonding (n) electron pairs carried by O or S.^{12,13} The model commonly used by chemists invokes sp^3 hybridisation at O in H_2O , with the half-angle of $\sim 50^\circ$ between two equivalent n-pairs, as shown in Figure 2 in the schematic, exaggerated form commonly employed in chemistry. For H_2S , the S–H bonds can be understood if each H 1s orbital overlaps with one of the two orthogonal p orbital on S (hence an angle HSH $\sim 90^\circ$) and if two sp hybrid orbitals at 180° contain one n-pair each (see Figure 2). This model can be validated by calculating the electrostatic potential energy of a non-perturbing, positive charge maintained at a fixed distance r from S. Figures 1 and 2 show the results for $V(\phi)$ as a function of ϕ , the angle made by a protonic charge with the plane of the three H_2Z nuclei ($Z = \text{O}$ and S , respectively), when $r = 1.74 \text{ \AA}$ (the $\text{O}\cdots\text{H}$ distance in $\text{H}_2\text{O}\cdots\text{HF}$)¹³ and 2.31 \AA (the $\text{S}\cdots\text{H}$ distance in $\text{H}_2\text{S}\cdots\text{HF}$)¹³, respectively, and when r is confined to the plane perpendicular to the nuclear plane.¹³ The electric charge distribution employed for H_2O was determined by using the CCSD/cc-pV5Z wavefunction. Figure 2 displays two such $V(\phi)$ versus ϕ curves for H_2S . The blue curve results from a distributed multipole analysis (DMA) carried out by Buckingham and Fowler⁴¹ using a wavefunction calculated at the SCF level with

large basis sets. Also shown in Figure 2 is the potential energy curve (red) recalculated by using the DMA of H₂S obtained from a CCSD/cc-pV(5+d)Z wavefunction. The difference between the two curves in Figure 2 is very small. The two minima at $\phi \approx \pm 80^\circ$ seen in Figure 2 clearly correspond with the most nucleophilic regions of H₂S and the two n-pair directions. The corresponding diagram for H₂O exhibits two minima at $\phi \approx \pm 30^\circ$. This methodology of probing the electrostatic potential of molecules with a view to predicting and understanding how intermolecular interactions may form is thus complementary to other techniques such as visualising the non-covalent interactions index (NCI).^{42,43}

A more sophisticated model of H₂O...HX, for example, involves a non-perturbing, extended electric dipole moment μ to represent HX, rather than a unit positive charge. HF can be represented as two point charges $\delta+$ and $\delta-$ separated by the HF bond length, with δ chosen from the DMA in ref. 41. The resulting potential energy $V(\phi)$ for H₂O...HF calculated at the appropriate distance $r(\text{O}\cdots\text{H})$ is shown in blue as curve (c) in Figure 1 and is close to that [curve (a)] determined from experiment, with minima at $\phi \approx \pm 55^\circ$. This value of ϕ_{min} is encompassed by the experimental value¹⁸ [$\pm 46(8)^\circ$] and is consistent with tetrahedrally disposed n-pairs on O.¹³ The barrier height (2.9 kJ mol⁻¹) for curve (c) in Figure 1 is also similar to that (1.5 \pm 0.8 kJ mol⁻¹)¹⁸ determined spectroscopically. The same model has been applied recently¹⁴ to the series of cyclic ethers 2,5-dihydrofuran, oxetane and oxirane and their complexes with HF. Along this series, the internal ring angle COC decreases from 108 $^\circ$ through $\sim 90^\circ$ to $\sim 60^\circ$. The extended electric dipole moment model of HF suggested¹⁴ that the angle 2ϕ between the n-pairs on O increases as the ring angle COC decreases.

We present here an investigation of the analogous series of thio-ethers, namely 2,5-dihydrothiophene, thietane and thiirane, in which the CSC ring angle also decreases from $\sim 95^\circ$ through $\sim 77^\circ$ to $\sim 48^\circ$, respectively. Molecular diagrams of the three cyclic thioethers are set out in Figure 3. These were calculated at the CCSD(T)-F12c/cc-pVTZ-F12 level of theory, as described in Section 2. Given that the result described above for H₂S suggests an inter-n-pair angle of $\sim 180^\circ$, the aim of the work described here is to determine how the inter-n-pair angle varies in the thioether series. The tools for the investigation are: (1) the electrostatic potential energy of a non-perturbing, unit positive around the S atom in each molecule, (2) the potential energy of the extended electric dipole moment model of HF around S, and (3) *ab initio* calculations of the geometries of the thioether...HF complexes.

2. Computational details

The geometries of the interacting complexes and isolated molecules were optimised using the MOLPRO system^{44,45} of *ab initio* programs with the explicitly correlated CCSD(T)-F12c method⁴⁶ [this is also sometimes referred to as CCSD(T)(F12*)]. The triple-zeta correlation consistent basis sets for explicitly correlated methods, cc-pVTZ-F12, were used,⁴⁷ along with the cc-pVTZ-F12/MP2Fit,⁴⁸ cc-pVTZ/JKFit⁴⁹

and cc-pVTZ-F12/OptRI⁵⁰ auxiliary basis sets. The geminal Slater exponent was $1.0 a_0^{-1}$, and the geometries are provided as Supplementary Material.

The optimised geometries of the isolated molecules were then used in distributed multipole analyses (DMAs),⁵¹ with detailed results of the latter available in the Supplementary Material. The first-order density matrix required for a DMA was calculated at the CCSD/cc-pV(5+d)Z level of theory,^{52,53} where the +d denotes that the basis set includes additional “tight” d functions for sulfur.⁵⁴ Symmetry-adapted perturbation theory (SAPT)⁵⁵ calculations were carried out on some complexes in order to decompose the interaction energy into electrostatic, exchange, induction and dispersion components. The so-called “chemist’s grouping” was used to assemble the individual SAPT2+(3) δ MP2 components into the four above physical components.⁵⁶ All SAPT calculations were carried out with the Psi4 V1.2 program⁵⁷ using the aug-cc-pV(T+d)Z orbital basis set,^{54,58,59} with density fitting enabled by the corresponding auxiliary basis sets.⁶⁰ Molecular electrostatic potential maps (MESPs) were generated at the MP2/6-311++G** level⁶¹ by the SPARTAN package⁶² with an iso-density surface of 0.001 e/bohr³.

3. Results.

3.1 Electrostatic potential energy curves for 2,5-dihydrothiophene, thietane and thiirane

Figure 3 shows diagrams (drawn to scale) of the geometries of the three cyclic thioethers 2,5-dihydrothiophene, thietane and thiirane, as optimised at the CCSD(T)-F12c/cc-pVTZ-F12 level of theory. Details are available in the Supplementary Material. The ring atoms are co-planar in the equilibrium conformations of 2,5-dihydrothiophene and thiirane, but the four membered ring is puckered in thietane, both conclusions in accord with those from rotational spectroscopy.⁶³⁻⁶⁵ The angle of puckering of the ring in thietane (36°) is in agreement with that determined from the rotation-inversion spectrum.⁶⁵

The electrostatic potential energy of a non-perturbing protonic charge $V(\phi)$ at a fixed distances $r = 2.169$ Å, 2.155 Å and 2.162 Å from the S atom as a function of the angle ϕ is shown in Figure 4 for thiirane, 2,5-dihydrothiophene and thietane, respectively. These are the equilibrium values from the CCSD(T)-F12c/cc-pVTZ-F12 calculations. The angle ϕ is that made by the vector r with the heavy-atom plane in 2,5-dihydrothiophene and thiirane and corresponds to the angle defined in Figures 1 and 2. For thietane, the plane in question is that made by S and its two contiguous C atoms.

The two equivalent minima in Figure 4 for thiirane and 2,5-dihydrothiophene occur at $\pm 80^\circ$ and $\pm 75^\circ$, respectively. The hydrogen-bonded complex thiirane \cdots HF has been investigated in detail through its rotational spectrum,⁶⁶ the interpretation of which led to the angle $\phi = 86.5(11)^\circ$, in reasonable agreement with the notion that HF lies along a n-pair direction, if the minima in Figure 4 correspond to n-pair

directions. According to Figure 4, the complex 2,5-dihydrothiophene \cdots HF should have HF forming a hydrogen bond to S with an angle ϕ close to 80° , but this complex has yet to be investigated experimentally. Thietane has the four-membered ring puckered by 36° in the equilibrium geometry and therefore has C_s rather than C_{2v} symmetry. This means that the two n-pairs on S are no longer equivalent, a conclusion reflected in Figure 4, in which the $V(\phi)$ versus ϕ curve for thietane shows two inequivalent minima at approximately -70° and $+65^\circ$ but different in energy by only 9 kJ mol^{-1} . If these minima can be identified with the directions of n-pairs, we might refer to them as axial and equatorial n-pairs, respectively. In fact, Alonso and co-workers⁶⁷ have characterised two conformers of the complex thietane \cdots HF in a supersonic expansion of the two component in helium, with angles $\phi_{\text{axial}} = -88.8(2)^\circ$ and $\phi_{\text{equat.}} = 89.8(3)^\circ$ and with the axial form having the greater population and therefore the low energy at the end of the expansion. The energy order of the minima in the $V(\phi)$ versus ϕ plot for thietane has the equatorial minimum lower in energy than that of the axial minimum and the ϕ_{min} values are somewhat smaller in magnitude than those from experiment. This is confirmed by a high-resolution, infrared spectroscopy investigation of thietane \cdots HF.⁶⁸

The conclusions from Figure 4 are broadly consistent with those that can be drawn from the molecular electrostatic surface potentials of the three cyclic thioethers, which are set out in Figure 5. The MESP in each case corresponds to an iso-density surface of 0.001 e/bohr^3 and was obtained by a SPARTAN calculation at the MP2/6-311++G** level of theory. Half the MESP has been cut away to reveal a tubular model of the molecule in each case. The MESPs of thiirane and 2,5-dihydrothiophene show two equivalent red spots above and below the S atom which make angles of $\phi \sim \pm 70\text{-}80^\circ$ at S with the C_2 axis and correspond to the most nucleophilic regions of these molecules, presumably those along the n-pair axes. This reflects the two equivalent potential energy minima shown in Figure 4 for each of these molecules. On the other hand, the MESP for thietane shows clearly that the equatorial red spot (n-pair) is more nucleophilic than the axial n-pair for which, at the colour resolution employed, no red spot is evident. This observation is consistent with the result for the thietane $V(\phi)$ versus ϕ potential curve (Figure 4) which reveals that the equatorial position has the lowest electrostatic potential energy. The angle ϕ made by the equatorial red spot appears to be smaller than the corresponding angles in thiirane and 2,5-dihydrothiophene, in agreement with the conclusion from Figure 4.

3.2 Potential energy of an extended electric dipole moment model for HF interacting with 2,5-dihydrothiophene, thietane and thiirane

It was suggested some years ago that the geometry of a hydrogen-bonded complex $B\cdots\text{HX}$, where B is a Lewis base and X is a halogen atom or a pseudo-halogen group, can be predicted by assuming that in the

equilibrium geometry the HX molecule lies along the axis of a n-pair carried by the acceptor atom of B.^{12,13} As a corollary to this rule, it was suggested that, e.g. HF might act as a probe of n-pair directions. Persevering with electrostatics, a slightly more sophisticated approach to probing n-pair directions via electrostatic potential energy consists of using a non-perturbing extended electric dipole moment model for hydrogen fluoride in place of merely a non-perturbing, single protonic charge. Guided by the DMA for HF given in ref. 41, we choose a model of HF that consists of two charges $+0.540|e|$ and $-0.540|e|$ on H and F, respectively, separated by 0.9256 Å (the r_0 bond length of HF calculated from the rotational constant B_0).⁶⁹ This model ignores small electric dipoles and quadrupoles centred on H and F in ref. 41 and therefore corresponds to the zeroth-order approximation to the charge distribution of hydrogen fluoride as represented by its DMA. HF is assumed to lie on the line that subtends an angle ϕ at the S atom of the cyclic thioether, as defined earlier when evaluating electrostatic potential energy. Then, with the H atom of HF at an appropriate distance from S, the electrostatic potential energy of both charges is evaluated with the aid of the DMA of the cyclic ether and summed. The results of the energy $V_{\text{HF}}(\phi)$ as a function of ϕ are given in Figure 6. The S \cdots H distances were 2.169 Å, 2.155 Å and 2.162 Å as previously used for the calculation of the electrostatic potential.

The angles ϕ at the minima in Figure 6 are $\pm 85.4(10)^\circ$ for thiirane and $\pm 80.4(10)^\circ$ for 2,5-dihydrothiophene. The two inequivalent minima for thietane are now at $\phi = 68.1(10)^\circ$ and $79.5(10)^\circ$ for the equatorial and axial positions, respectively, with the equatorial minimum again the lower in energy, but by only about 1.7 kJ mol⁻¹. It is worth noting that the angles ϕ when the extended electric dipole model for HF is used are larger and in somewhat better agreement with those of the MESP's in Figure 5 and with the experimental geometries of thiirane \cdots HF⁶⁶ and thietane \cdots HF⁶⁷ than when the simple electrostatic potential energy is used in the comparison. The angle ϕ at the equatorial position is again smaller than that at the axial position. A feature of Figure 6 is the rapid drop of the potential energy of HF around thietane at the angle $\phi \simeq +100^\circ$ and greater. This arises because, when HF goes beyond the axial minimum, the nucleophilic F atom gets close to an electrophilic H atom [blue region in Figure 5(c) on C(2)]. A secondary attractive interaction of F and H-C(2) then becomes significant and causes the noted drop in energy. A secondary interaction is clearly present in the results from *ab initio* calculations to be discussed in Section 3.3 and shown in Figure 7 (c), but it is then moderated by the presence of exchange repulsion, which is absent from the extended electric dipole model of HF. Indeed, SAPT calculations on the two thietane \cdots HF conformers show that a full *ab initio* treatment of the interaction energy places the axial conformer slightly lower in energy, with an electrostatic contribution of -48.57 kJ mol⁻¹, compared to -44.51 kJ mol⁻¹ for the equatorial conformer.

3.3 Geometries of cyclic thioether \cdots HF complexes from *ab initio* calculations

The equilibrium geometries of thiirane···HF, thietane···HF and 2,5-dihydrothiophene···HF calculated at the CCSD(T)-F12c/cc-pVTZ-F12 level of theory are represented as the scale models of the complexes shown in Figure 7. The values of the angle ϕ made by the red-dotted line from H to S (the hydrogen bond) with the C_2 axis in each of thiirane and 2,5-dihydrothiophene or with the local C_2 axis of the CSC fragment of thietane are indicated and given in the caption, together with the angle θ , which measures the angular deviation of the S···H–F nuclei from collinearity. Two low-energy conformers of thietane···HF were detected, namely the axial conformer having $\phi = 88.5^\circ$ in Figure 7(c) and the equatorial conformer having $\phi = 76.6^\circ$, with the former the lower in energy by 1.41 kJ mol⁻¹. This is the energy order deduced by Alonso and co-workers⁶⁷ from their analysis of the rotational spectra of the two forms. These authors report r_0 values of 88.8(2)° and 89.8(3)° for the axial and equatorial conformers, respectively. The equilibrium values of ϕ for the two conformers (see caption to Figure 7) are both larger by about 8-9° than those [79.5(1)° and 68.1(10)°] obtained by using the extended electric dipole model of HF in Section 3.2. The ϕ_e values for thiirane and 2,5-dihydrothiophene (93.1° and $\phi = 84.5^\circ$, respectively) are likewise larger by about 7° and 4° than those [85.4(10)° and 80.4(10)°, respectively] from the extended electric dipole model of HF. A possible reason for this under-estimation of the angles ϕ by the model can be identified by considering the angular non-linearities θ .

All geometries shown in Figure 7 exhibit non-linear hydrogen bonds formed by HF to a n-pair on S. The angular non-linearities θ clearly arise from secondary interactions of F (the nucleophilic end of HF) with the regions near H atoms of the cyclic thioether. In a thought experiment, we might envisage that the HF molecule is initially constrained to approach along an axis of a n-pair on S and then the secondary interaction is switched on. The F atom moves towards an electrophilic H atom, the movement largely pivoted at H of HF but presumably there is some movement of the H (of HF) in the same direction. The larger the secondary interaction, the greater the movement of H and the greater the non-linearity. Thus, as well as causing the hydrogen bond to be non-linear, this model leads ϕ to increase from that defined by the n-pair direction. In this context, we note that the angles ϕ and the non-linearities θ are $\phi = 93.1^\circ$ and $\theta = 21.1^\circ$ for thiirane···HF, $\phi = 84.5^\circ$ and $\theta = 10.3^\circ$ for 2,5-dihydrothiophene···HF, (c) $\phi = 88.5^\circ$ and $\theta = 14.1^\circ$ for *axial*-thietane···HF. Larger deviations of θ from zero tend to be associated with larger positive deviations $\Delta\phi$ of ϕ from the n-pair angle, which (according to the extended electric dipole model of HF as a probe) is in the range 80-85° for these three cases. The values of the ($\Delta\phi$, θ) pairs are (7.6°, 21.1°), (9.0°, 14.1°) and (4.1°, 10.3°) for thiirane···HF, *axial*-thietane···HF and 2,5-dihydrothiophene···HF, respectively. For *equatorial*-thietane···HF, the same model gives the angle $\phi = 68.1^\circ$ for the other n-pair of S, which, because the *ab initio* value is $\phi = 76.6^\circ$, implies that a positive deviation $\Delta\phi = 8.5^\circ$ accompanies the non-linearity $\theta = 17.5^\circ$ for this conformer. The values of θ have been measured experimentally to be 16.8(13)°, 8(4)°, and 12(6)° for the first three members of the series.^{66,67}

4. Conclusions.

To probe the nucleophilic regions of cyclic thioethers in connection with predicting geometries of intermolecular complexes, the electrostatic potential energy $V(\phi)$ of a non-perturbing, protonic charge at a fixed distance r from the S atom in three cyclic thioethers has been examined as a function of the angle ϕ made by the r -vector with the C_2 axis (in thiirane and 2,5-dihydrothiophene) or the local C_2 axis associated with the C–S–C moiety of the thietane puckered ring. The results have been compared with molecular electrostatic surface potentials of the three cyclic thioethers. The electrostatic potential energy $V_{\text{HF}}(\phi)$ of HF modelled as an extended electric dipole has also been calculated as a function of the angle ϕ . Finally, the geometries of the complexes thiirane \cdots HF, thietane \cdots HF and 2,5-dihydrothiophene \cdots HF were optimised at the CCSD(T)-F12c/cc-pVTZ-F12 level of theory. The positions of the minima in the $V_{\text{HF}}(\phi)$ versus ϕ functions are in good agreement with the directions of maximum nucleophilicity exhibited in the MESP diagrams of the cyclic thioethers. In the simplest electrostatic model, the minima $|\phi_{\text{min}}|$ in the $V(\phi)$ versus ϕ functions occur at smaller values, the reason for which has been discussed elsewhere.¹⁴ In the thioether \cdots HF complexes, the *ab initio* calculations reveal that atoms S \cdots H–F deviate from collinearity by an angle θ in a direction that suggests a secondary interaction of the nucleophilic region of F with electrophilic H atoms attached to ring carbon atoms. The *ab initio* calculations also find that the angles ϕ made by the S \cdots H hydrogen bond with the C_2 (or local C_2) axes are systematically larger than those at the minima in the $V_{\text{HF}}(\phi)$ versus ϕ functions. This systematic difference can be explained by using a simple model of complex formation that accounts for secondary interactions between the nucleophilic F and regions near hydrogen atoms of the cyclic thioether.

If it can be assumed that the minima of the $V_{\text{HF}}(\phi)$ versus ϕ potential energy function (resulting from the model in which HF is treated as an extended electric dipole) provide a method of locating non-bonding electron pairs, there is no evidence that the half-angle between the n-pairs on S increases along the series 2,5-dihydrothiophene, *axial*-thietane, thiirane as the ring angle CSC decreases from 94.9° through 76.7° to 47.9°. Indeed, the values of ϕ_{min} are 80.4(10)°, 79.5(10)° and 85.4(10)°, respectively, so that 2,5-dihydrothiophene has the largest ring angle but an inter-n-pair angle that is insignificantly different from that of *axial*-thietane. There is evidence of a small increase when the ring angle is very small in thiirane. (The *equatorial* n-pair of thietane has been excluded from this discussion because of the distorting effect that the secondary interaction mentioned in Section 3.2 has on the potential curve when unmoderated by exchange repulsion). In the corresponding series in which the ring heteroatom is oxygen, namely 2,5-dihydrofuran, oxetane and oxirane, and in which a similar decrease of ring angle obtains, there is evidence of a significant increase in the inter-n-pair angle as the ring angle decreases.¹⁴ Presumably, if the n-pairs on S are in two sp hybrids and the CS bonds involve two orthogonal p orbitals on S, there is only small scope for changing the hybridisation at S as the ring angle decreases. Accordingly, the inter-n-pair angle changes

only slightly in the series of cyclic thioethers. This explanation implies increasingly bent S–C bonds as the ring angle at S decreases.

5 Acknowledgements

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6 References

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Figures

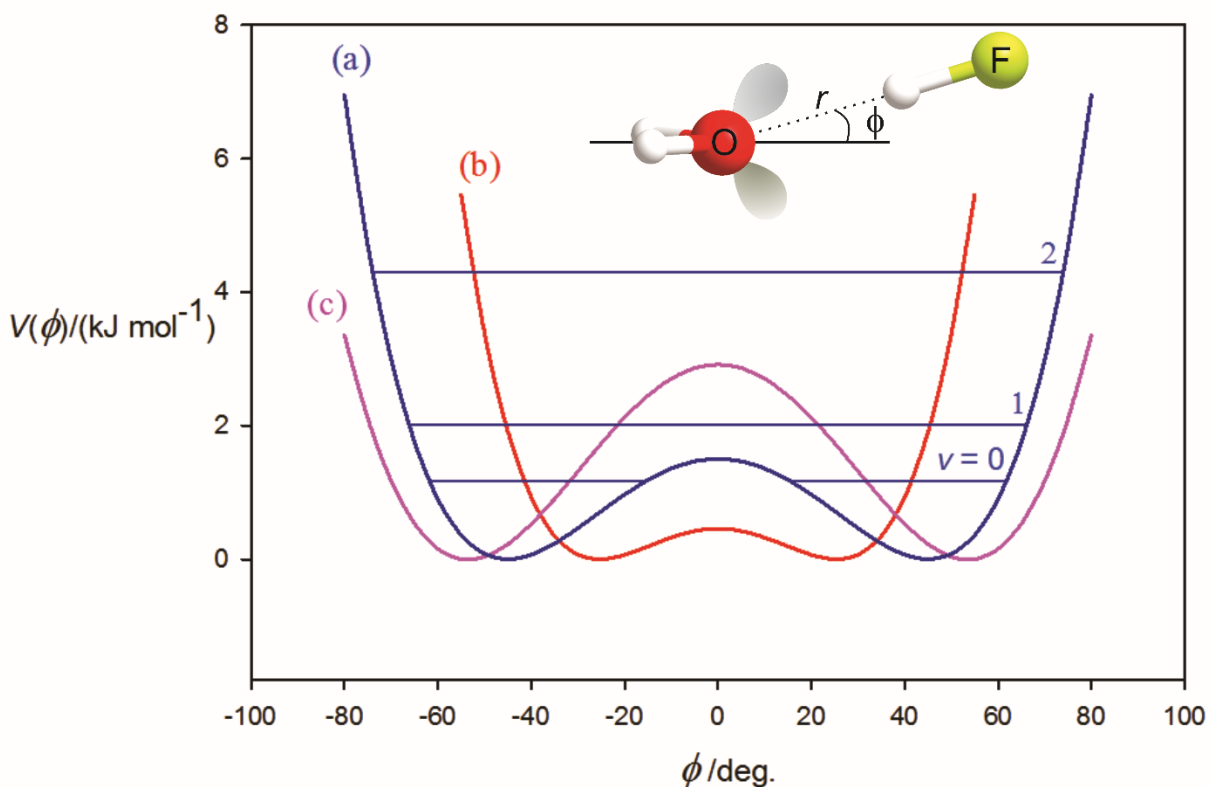


Figure 1. Curve (a) is the experimentally determined potential energy function associated with the low-frequency bending mode of $\text{H}_2\text{O}\cdots\text{HF}$ (blue) and associated energy levels (see ref. 18) The configuration at O is rapidly inverting so that in the zero-point ($v=0$) state the vibrational wavefunction has C_{2v} symmetry and the molecule is effectively planar. At equilibrium the configuration is pyramidal at O. The red curve (b) is the electrostatic potential energy of a non-perturbing, protonic charge placed at a distance of $r=1.74$ Å from O (the experimental $r(\text{O}\cdots\text{H})$ distance in $\text{H}_2\text{O}\cdots\text{HF}$) calculated by using the DMA obtained from the CCSD/cc-pV5Z wavefunction to represent the H_2O charge distribution. The pink curve (c) is the potential energy obtained by using the same charge distribution for H_2O , but with the extended electric dipole model for hydrogen fluoride (see text for discussion) at the experimental distance.

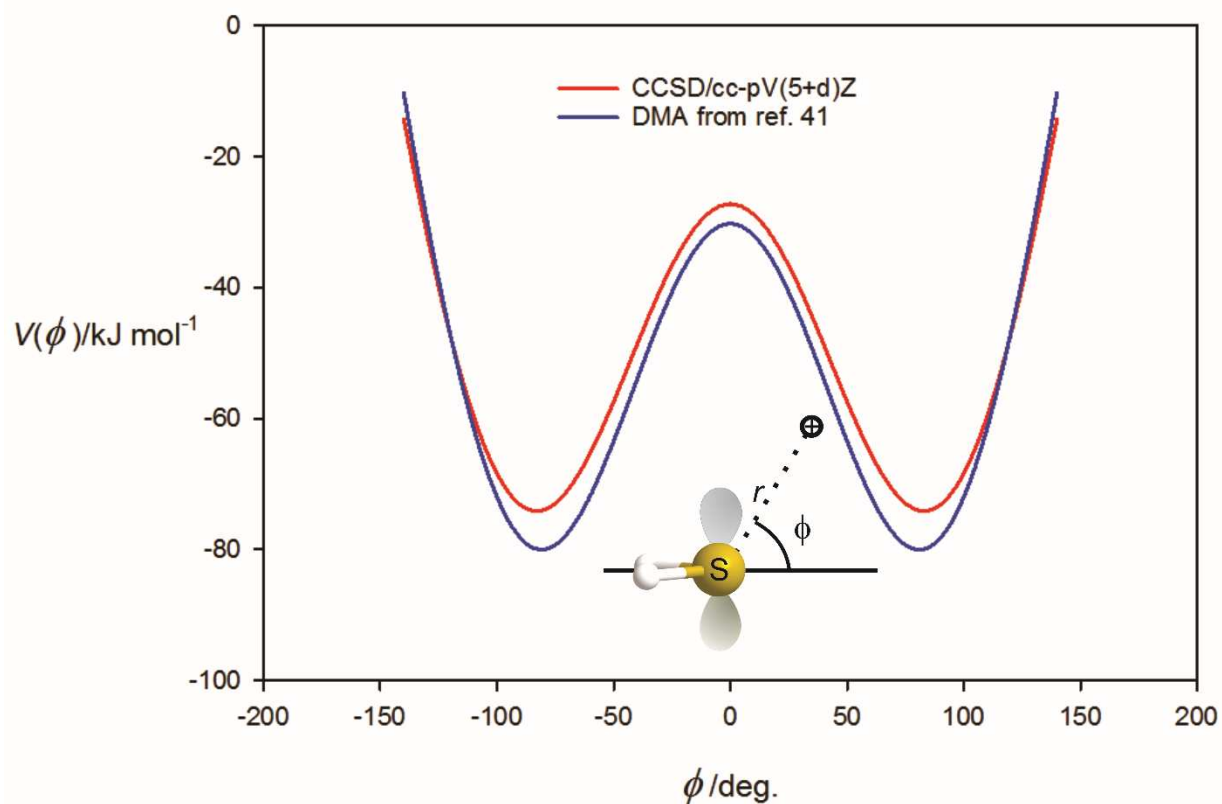


Figure 2. Electrostatic potential energy $V(\phi)$ of a non-perturbing, protonic charge in the plane perpendicular to plane of the H_2S nuclei as a function of the angle ϕ at a fixed distance $r = 2.31 \text{ \AA}$ from S [the experimental distance $r(\text{S}\cdots\text{H})$ in $\text{H}_2\text{S}\cdots\text{HF}$]. The blue (lower) curve results from use of the DMA given in ref. 41 to calculate $V(\phi)$. The red curve was generated by using the DMA of H_2S obtained from the CCSD/cc-pV(5+d)Z wavefunction.

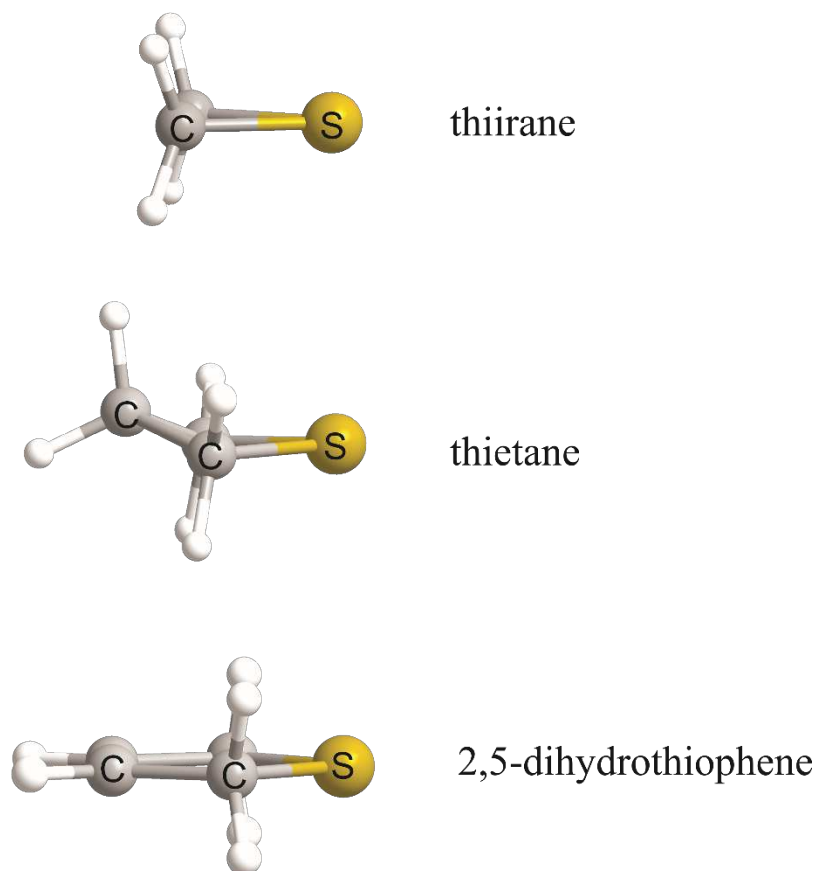


Figure 3. Models of the geometries of the three cyclic thioethers thiirane, thietane and 2,5-dihydrothiophene (to scale). The geometries are those that result from calculations at the CCSD(T)-F12c/cc-pVTZ-F12 level of theory.

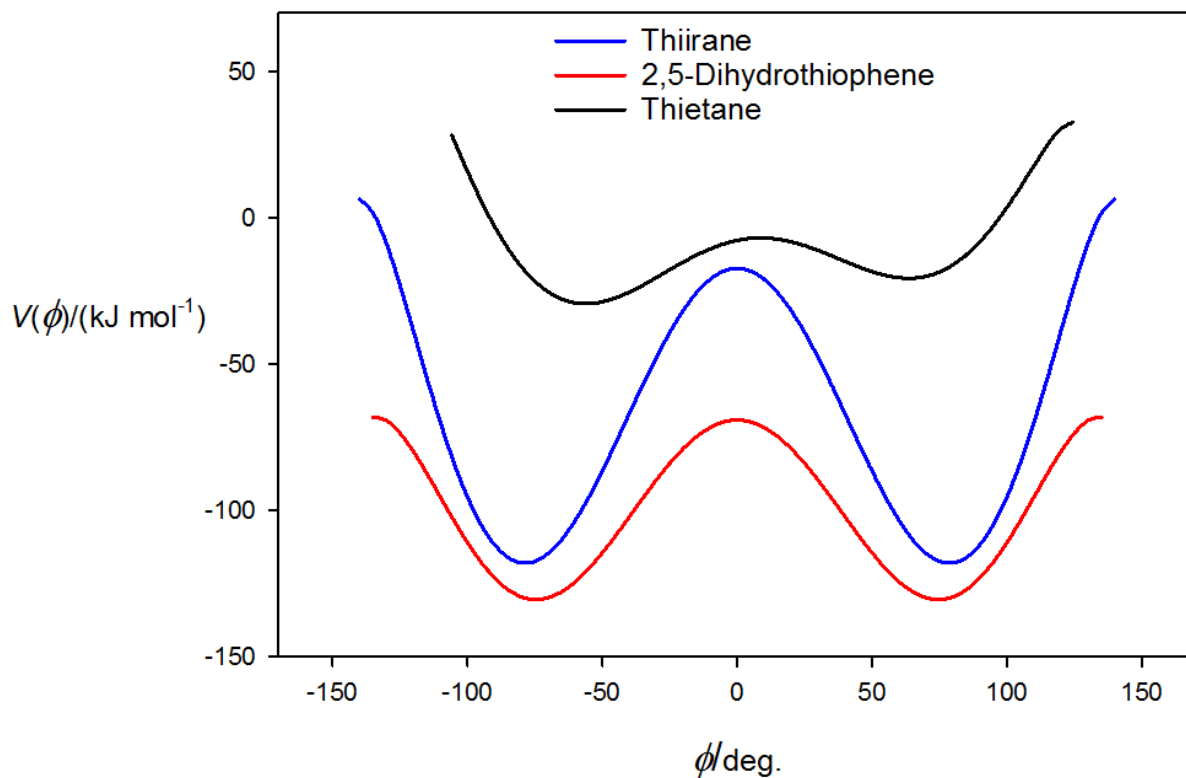


Figure 4. Electrostatic potential energy of a non-perturbing protonic charge at constant distances $r = 2.169 \text{ \AA}$, 2.162 \AA and 2.155 \AA from the S atom as a function of the angle ϕ in thiirane, thietane and 2,5-dihydrothiophene, respectively. In thiirane and 2,5-dihydrothiophene the angle ϕ is the angle made by the vector r with the plane of the ring atoms, as defined in Figure 1. In thietane, ϕ is the angle made by r with the plane defined by the S atom and its two contiguous C atoms. For clarity, 50 kJ mol^{-1} has been added to each point on the thietane curve.

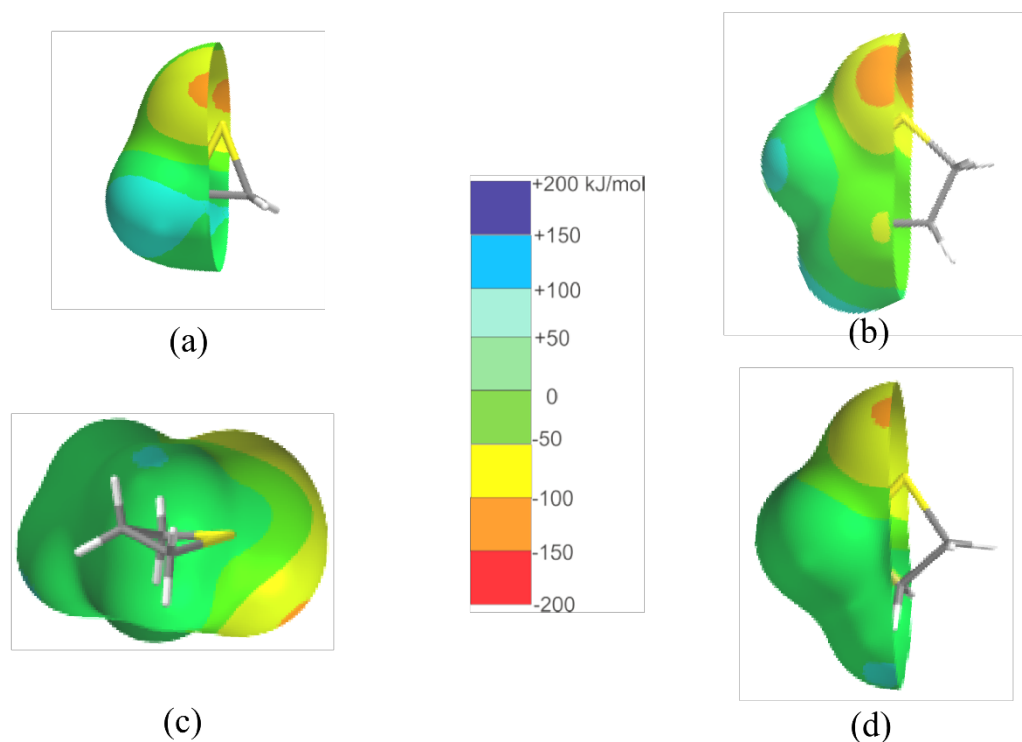


Figure 5. Molecular electrostatic surface potentials (MESPs) of (a) thiirane, (b) 2,5-dihydrothiophene, (c) and (d) thietane, calculated at iso-surfaces having a density of 0.001 e/bohr^3 . In each case, half of the MESP has been cut away to reveal a tubular model of the molecule within. The thiirane and 2,5-dihydrothiophene molecules have been rotated slightly about their C_2 axes anti-clockwise to show some of the inside of the MESP. The red spots on the upper and lower surfaces apparent in both thiirane and 2,5-dihydrothiophene represent the most nucleophilic regions of the molecules and may be identified with directions of the n-pairs on S. In (c), the thietane molecule is viewed from the side, while in (d) the view is from underneath. The red spot in (d) corresponds to the equatorial n-pair; note that, for the resolution of colours employed, there is no corresponding red spot on the inside, under-surface for the axial n-pair. The equatorial red spot is also visible in (c). This indicates that the equatorial n-pair is more nucleophilic than the axial n-pair.

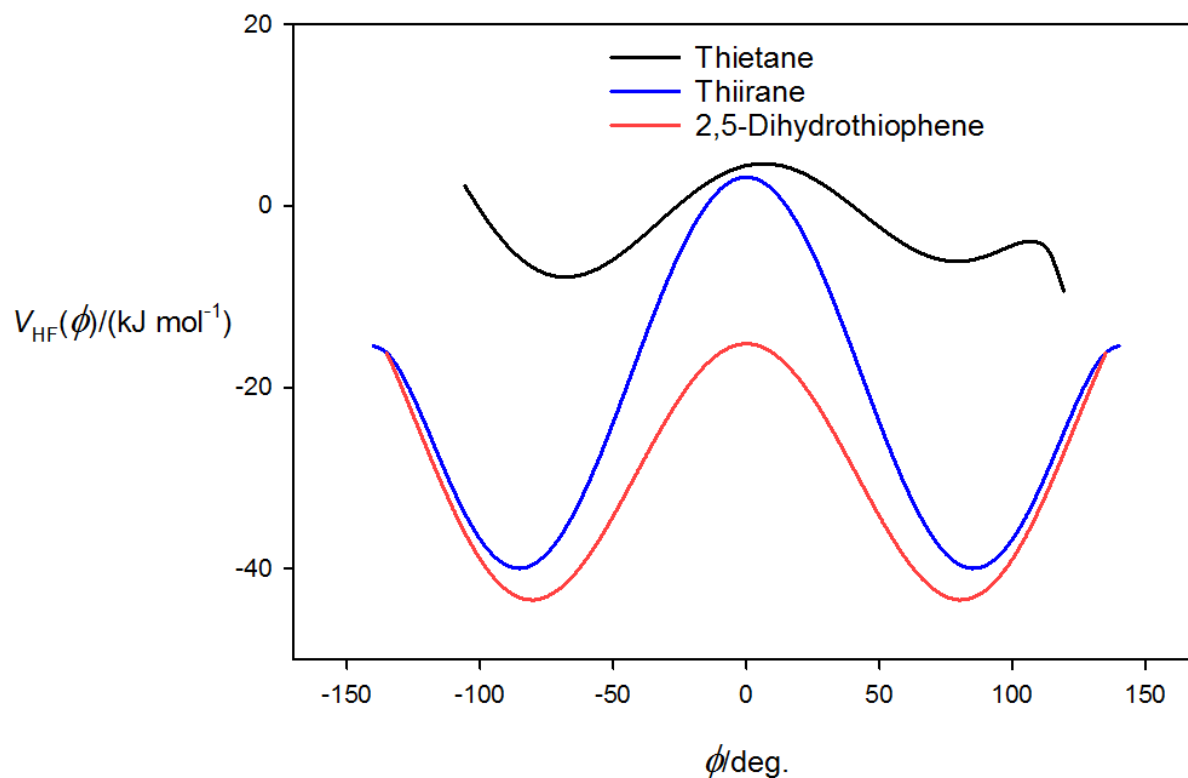


Figure 6. Plot of the potential energy $V(\phi)$ of the extended electric dipole model of HF (see text for model details) as a function of the angle ϕ made by the HF molecule at S with the C_2 axis (for thiirane and 2,5-dihydrothiophene) or the plane of S and its contiguous C atoms (thietane). The distances r of H from S are those given in the caption to Figure 4. For clarity, 15 kJ mol^{-1} has been added to each point of the thietane curve.

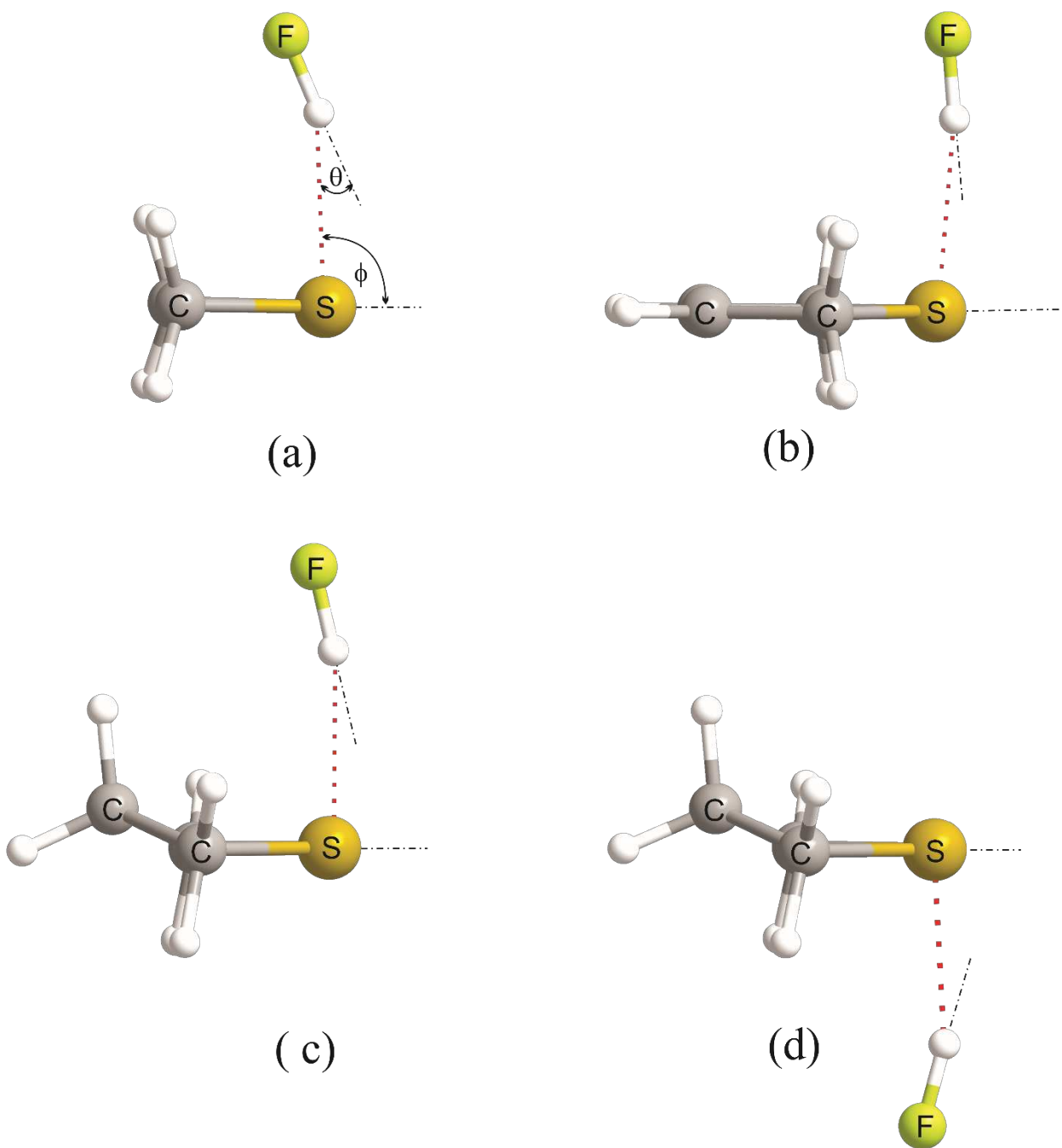


Figure 7. *Ab initio* geometries of (a) thiirane...HF, (b) 2,5-dihydrothiophene...HF, (c) the axial conformer of thietane...HF, and (d) the equatorial conformer of thietane...HF determined in optimisations carried out at the CCSD(T)-F12c/cc-pVTZ-F12 level of theory. Diagrams are to scale. The angles ϕ and θ [defined in (a)] are as follows: (a) $\phi = 93.1^\circ$ and $\theta = 21.1^\circ$, (b) $\phi = 84.5^\circ$ and $\theta = 10.3^\circ$, (c) $\phi = 88.5^\circ$ and $\theta = 14.1^\circ$, and (d) $\phi = 76.6^\circ$ and $\theta = 17.5^\circ$. Further geometrical details are available in the Supplementary Material. The energy difference of the axial (c) and equatorial (d) conformers of thietane...HF is 1.41 kJ mol^{-1} when calculated at the CCSD(T)-F12c/cc-pVTZ-F12 level.