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Structural and FTIR spectroscopic studies of matrix-isolated 3-thio-1,2,4-triazole complexes with carbon dioxide. The UV-induced formation of thiol···CO₂ complexes†

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Matrix isolation FTIR spectroscopy was combined with quantum chemical calculations to characterize complexes of 3-thio-1,2,4-triazole (ST) with carbon dioxide. Geometries of the possible 1:1 and 1:2 complexes were optimized at the DFT (B3LYPD3) level of theory with the 6-311++G(3df,3pd) basis set. The computational results show that ST interacts specifically with carbon dioxide through different hydrogen bond and van der Waals interactions. For the 1:1 complexes of the most abundant ST thione tautomer, four stable minima, $STn \cdots CO_2$, have been located on the potential energy surface. In contrast, for the ST thiol tautomer, three STI····CO₂ structures were optimized. Experimentally, the two most stable 1:1 complexes of STn with CO2, characterized by the presence of the N-H···O hydrogen bridge and an additional S6...C10 interaction, were identified in solid argon upon deposition. Annealing of the matrix at 32 K proved that one 1:2 structure is also present, resulting from the addition of a second CO2 molecule to the 1:1 complexes. The laser irradiation at $\lambda = 270$ nm, apart from generating the thiol tautomer of ST, also leads to the formation of three thiol···CO2 complexes. Furthermore, the presence of CO2 in the argon matrix was found to influence the efficiency of the UV-induced thione-thiol tautomerization, though to a lesser extent than nitrogen. This suggests that while CO2 forms stronger intermolecular interactions with ST, its impact on tautomerization kinetics is less pronounced, highlighting the nuanced role of specific gas-phase interactions in modulating photochemical transformations in low-temperature matrices. The findings presented in this work not only enhance the fundamental understanding of weak intermolecular interactions but also provide new insights into the role of CO₂-specific effects in photochemical and structural transformations of heterocycles.

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Introduction

Intermolecular interactions have been the subject of study for many years and have been examined by a wide variety of techniques. The interest in this area is related to the prevalence of this behavior in nature (e.g. biochemical processes, 1,2 the atmosphere, 3,4 surface science, 5 etc.). Moreover, in recent years, there has been a great deal of interest in the interactions of various heterocyclic molecules with carbon dioxide. This is related to the role that CO₂ plays in the greenhouse effect and the efforts to establish efficient methods for capturing various atmospheric pollutants. Heterocycles, with their diverse electronic and structural properties, offer a promising foundation for developing functional frameworks that selectively bind and

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 \dagger Electronic supplementary information (ESI) available: Selected AIM parameters, selected wavenumber shifts, and band intensities of ST···CO $_2$ complexes calculated at B3LYP-D3/6-311++G(3df,3pd) level. See DOI: https://doi.org/10.1039/d5ra02230d

sequester CO₂. A group of compounds that are particularly suited for this purpose are azoles, whose applications have been proven by numerous scientific articles regarding the use of those molecules as MOF, MOP, ZIF, or other frameworks' components. 6-10 The subject of this work, 3-thio-1,2,4-triazole (ST), makes a relevant candidate for capturing CO₂ and other pollutants due to the presence of electronegative nitrogen atoms in the ring as well as a thiol group, which creates several possibilities for the acceptor molecule to attach. Another interesting feature of this molecule is its ability to undergo thione-thiol tautomerization, which has been the subject of research due to the importance of heterocyclic tautomerism in various processes related to bioactivity. 11-14 Recent studies have demonstrated that photo-induced tautomerization, such as keto-enol transitions, can alter the electron density distribution of a molecule and subsequently modify the binding affinity and geometry of CO2 complexes, as shown for carbonyl systems under UV exposure.15

Based on our previous work on 3-thio-1,2,4-triazole (ST) complexes with dinitrogen, ¹⁶ where we demonstrated different

ways of binding N2 molecules by the azole and identified particular geometries spectroscopically, we decided to study ST-CO₂ complexes in a similar way and compare their interaction energies, geometries, and vibrational spectra. Unlike N2, which is chemically inert and interacts primarily through dispersion forces and weak hydrogen bonding, CO2 exhibits a higher degree of electronic polarizability, leading to stronger and more directional interactions with ST. Therefore, CO2 may serve as a modulator of reaction pathways by stabilizing particular conformations or influencing the energy barriers for tautomerization and photolysis. Changes in the matrix environment may influence the tautomeric equilibrium of heterocyclic compounds as well as the efficiency of UV-induced transformations, which has been documented in the literature in previous years. 17-19 The investigation of ST-CO2 complexes introduces new aspects that significantly extend our understanding of intermolecular interactions in matrix isolation.

2. Experimental

2.1. Theoretical studies

All computations in this work were performed using the Gaussian 16 suite of quantum chemical programs.20 The structures of the monomers and the 3-thio-1,2,4-triazole complexes with CO2 were created in the GaussView 6.0 program²¹ and then optimized at the B3LYP-D3/6-311++G(3df,3pd) level of theory²²⁻²⁶ with the Boys-Bernardi full counterpoise method by Dannenberg et al.27,28 Interaction energies (E_{int}) were calculated by subtracting the total electronic energies of the monomers from the energy of the complex while preserving the geometry of the complex. Vibrational wavenumbers were determined using the harmonic approximation to verify that the identified structures correspond to minima on the potential energy surfaces and to monitor spectral changes resulting from the complex formation. To account for anharmonicity, the calculated wavenumbers were scaled by factors of 0.951 for the 4000-2800 cm⁻¹ range and 0.982 for the 2800-400 cm⁻¹ range. Interactions between ST and CO₂ in complexes were characterized using topological analysis of electron density (AIM), 29,30 which was conducted at the B3LYP-D3/6-311++G(3df,3pd) level with the AIM Studio program (Version 19.10.12, Professional).31

2.2. Matrix isolation experiments

The preparation of cryogenic matrices was carried out using methods and equipment as described in our previous paper on ST complexes. ¹⁶ Carbon dioxide used to prepare CO_2/Ar matrix gas mixtures was synthesized in the laboratory, and the procedure was described in ref. 32 in section 2.2. The CO_2/Ar concentrations obtained in the matrix were 1/500 and 1/1000. Photochemical processes were induced in ST/ CO_2/Ar matrices by UV radiation ($\lambda = 270$ nm) of a pulsed optical parametric oscillator Vibrant (Opotek, Inc.) pumped with a pulsed Nd:YAG laser (Quantel). The progress of the deposition and UV-induced reactions were monitored by recording successive IR spectra.

3. Results and discussion

The monomer of 3-thio-1,2,4-triazole (ST) in the matrix isolation conditions appears in the form of thione tautomer (ST*n*) with trace amounts of the thiol tautomer (STl), as described by Rostkowska *et al.*³³ Therefore, considering the complexation of ST with carbon dioxide, only those two structures were taken into account. They are presented in Fig. 1 below, together with the adopted numbering of atoms.

3.1. Quantum chemical calculations

3.1.1. Complexes of the 1:1 stoichiometry. Using the B3LYP-D3/6-311++G(3df,3pd) method, four $STn \cdots CO_2$ structures and three STI···CO2 structures were optimized. They are presented in Fig. 2 together with the atom numbering as well as calculated values of the interaction energy E_{int} and relative electronic energy ΔE . Selected geometric parameters: bond lengths and angle values and key topological data generated by AIM calculations are given in Tables S1 and S2 in the ESI.† Since the thione and thiol forms differ only in the position of one of the hydrogen atoms, the optimized geometries of the complexes of both tautomers are similar. Therefore, it is possible to distinguish analogous structures in both groups. They are marked with the same number (STn-C1 and STl-C1, STn-C2 and STl-C2, STn-C4 and STl-C4). The thiol equivalent of the STn-C3 complex could not be optimized, as it appeared as a very shallow minimum that undergoes a barrierless transformation into the more stable STl-C4 structure. In the first pair of complexes, STn-C1 and STl-C1, the subunits are connected by a hydrogen bond bridge N1-H9···O11, the angle of which equals 148.6 and 139.0°, respectively. The difference in the values of this angle results from the different position of the CO₂ molecule relative to the triazole ring. In the structure of ST*n*-C1, it lies in the plane, while in the case of STl-C1 it is tilted out of the plane.

In both complexes, there are also additional van der Waals interactions formed between the sulfur atom of the substituent and one of the atoms of the CO_2 molecule. In the case of STn-C1, it is a carbon atom, and in STl-C1, it is an oxygen atom. Interestingly, the interaction energy value calculated for the thione form complex is significantly higher than for the second complex $(-22.1 \text{ kJ mol}^{-1} \text{ compared to } -14.8 \text{ kJ mol}^{-1})$. This is most likely because in STn-C1, the geometry is stabilized by

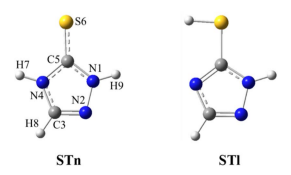


Fig. 1 Structures of thione and thiol tautomers of ST.

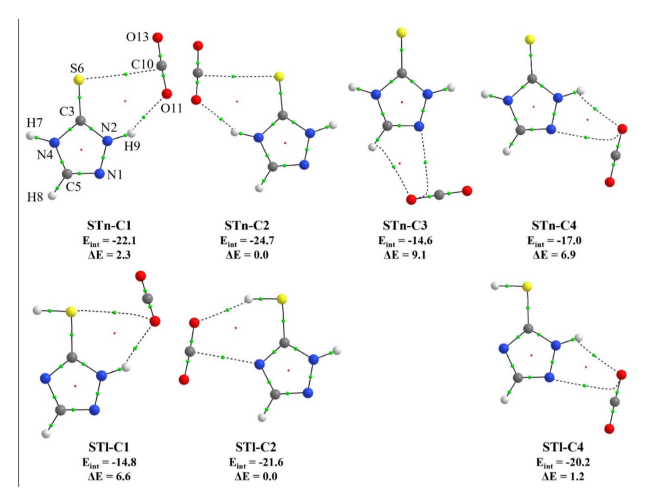


Fig. 2 Structures of 1:1 complexes of both ST tautomers with CO_2 molecule. Bond and ring critical points are marked with green and red dots, respectively. The values of interaction energy (E_{int}) and relative electronic energy (ΔE) are given below the structures, both in kJ mol⁻¹.

interactions involving two atoms of the CO2 molecule (C10 and O11), while in the case of STl-C1, it is only an oxygen atom, whose electron density is distributed over two interactions. The next pair of structures are STn-C2 and STl-C2, which are also stabilized by two connections - a hydrogen bond and a van der Waals contact. Due to the presence of two NH groups in STn-C2, the subunits are linked by an N4-H7···O11 hydrogen bond and an additional S6···C10 interaction. In the case of STl-C2, it is an S6-H7···O11 hydrogen bond and an N4···C10 interaction. It is worth mentioning that these complexes turned out to be the most stable in their groups (thione and thiol tautomers). The structures STn-C4 and STl-C4 are characterized by the presence of a hydrogen bridge N1-H9···O11 and an additional N2···O11 connection. Both geometries are planar, and the NHO angle is 120.0 and 120.9⁰, respectively. Despite their very similar structure, the interaction energy values of these complexes differ by 4.2 kJ mol⁻¹ in favor of the STl-C4 form. The last of the optimized structures, STn-C3, is the least stable, with the interaction energy equal to $-14.6 \text{ kJ} \text{ mol}^{-1}$. It is caused by the fact that, instead of the NH group, the CH group of the ring is involved in the interaction, forming a very weak hydrogen bond.

Worth mentioning is a certain relationship between the length of hydrogen bridges of individual structures and their energetic stability. In the thione tautomer group, the most stable ST*n*–C2 complex has a hydrogen bridge involving the N4–H group. The H7···O11 distance in this bridge is smaller than the analogous value of H9···O11 in the ST*n*-C1 structure where the N2–H group is involved (2.093 Å compared to 2.149 Å, respectively). This suggests that the hydrogen bonds formed by the N4–H group are stronger than those formed by the N2–H group. This observation is consistent with the results of the structural analysis of ST complexes with N₂.¹⁶

To better understand the origin of complex stability, we analyzed the hydrogen bond lengths and AIM topological parameters across all structures. The most stable complexes, STn-C2 and STl-C2, exhibit notably shorter N-H···O hydrogen bond distances and higher electron densities at the corresponding bond critical points, indicating stronger and more directional interactions. Furthermore, cooperative van der Waals contacts involving the sulfur atom (e.g., S···C interactions) enhance stabilization by supporting the binding geometry. These features correlate well with the calculated interaction energies, highlighting the synergistic role of hydrogen bonding and dispersion forces in determining the relative stabilities of the complexes.

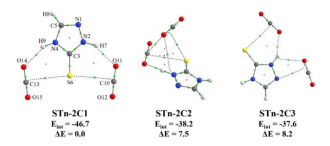


Fig. 3 Structures of 1:2 complexes of the STn tautomer with CO_2 . Bond, ring, and cage critical points are marked with green, red, and blue dots, respectively. The values of interaction energy (E_{int}) and relative electronic energy (ΔE) are given below the structures, both in kJ mol⁻¹.

3.1.2. Complexes of the 1:2 stoichiometry. Although the main goal of the study was to detect and analyze ST–CO $_2$ complexes of 1:1 composition, the formation of 1:2 aggregates could not be excluded under the experimental conditions used. The performed B3LYP-D3/6-311++G(3df,3pd) calculations revealed three stable structures formed between the ST thione tautomer and two CO $_2$ molecules. They are presented in Fig. 3, together with the values of interaction energy and relative energy.

Out of three 1:2 geometries, the most stable STn-2C1 is characterized by two hydrogen bonds: N2-H9···O11 and N4-H7···O14, as well as two additional stabilizing van der Waals interactions connecting the S6 atom with the C10 and C13 atoms of two CO₂ molecules. The H7···O14 distance in the hydrogen bridge formed by the N4-H group is smaller than the H9···O11 distance of the N2-H9···O11 bond (2.155 Å compared to 2.096 Å). This is consistent with the data obtained for ST

complexes with CO_2 of the 1:1 stoichiometry. In turn, the STn-2C2 and STn-2C3 complexes are characterized by higher energy values (by 7.5 and 8.2 kJ mol $^{-1}$, respectively), as well as significantly lower interaction energies. The first one has an interesting geometry, in which, according to the AIM results, the arrangement of bond paths allowed for the formation of as many as four rings, which together form a cage. The AIM topological parameters of individual BCPs of the 1:2 structures, together with their selected geometric parameters, are available in Table S3 of the ESI.†

3.2. Matrix isolation infrared spectra of ST/CO₂/Ar samples

In agreement with the previously published results¹6 the ST monomers isolated in an argon matrix are present in the STn thione form (see Fig. 1). The introduction of the CO₂ dopant into the matrix gas resulted in the appearance of a series of bands that had not been present in the spectrum of ST in pure argon. To identify the particular geometries of the 1:1 complexes formed between ST and CO₂ in the matrix, positions of the new bands were compared with the theoretical wavenumbers calculated at the B3LYP-D3/6-311++G(3df,3pd) level. The values of the predicted band shifts of the four optimized 1:1 geometries (relative to the analogous monomer absorptions), together with their intensities and the corresponding experimental shifts, are summarized in Table 1.

Six selected fragments of the spectra of ST/CO₂/Ar matrices are presented in Fig. 4 compared with the ST spectrum in pure argon. In the NH stretching mode range (3510–3430 cm⁻¹), in addition to two intense absorptions caused by the $\nu(N_2H)$ and $\nu(N_4H)$ vibrations of the ST thione located at 3498.0 and 3491.0 cm⁻¹, four new bands at lower wavenumbers appeared as a result of the complexation of the ST monomer with the CO₂

Table 1 Selected shifts of wavenumbers $\Delta \nu$ (cm⁻¹) and band intensities (km mol⁻¹, in brackets) calculated at the B3LYP-D3/6-311++G(3df,3pd) level for the 1:1 complexes of the thione tautomer (STn) with a CO₂ molecule, compared with the experimental shifts

Theoretical shifts $(\Delta \nu)$						
STn-C1	STn-C2	STn-C3	STn-C4	$Mode^{16,a}$	Experimental shifts ^b	Assignment
-10 (84)	-1 (104)	-1 (89)	-10 (67)	vN_2H	n.o.	
-29 (228)	-53(249)	0 (103)	-7 (1 77)	vN_4H	-26.5	STn-C1
					-52.0	STn-C2
+1 (81)	+2 (61)	0 (50)	+2 (58)	ν CN + δ NH	+1.0	STn-C1
					+3.0	STn-C2
+4 (402)	+6 (430)	+1 (457)	+1 (507)	ν CN + δ NH + δ _{ring}	+4.5, +6.0	STn-C1
				_	+7.0	STn-C2
+3 (15)	+5 (16)	-1 (21)	+1 (16)	ν CN + δ_{ring}	+7.0	STn-C2
				_	+9.0	STn-C1
-1 (77)	+3 (84)	+1 (100)	+1 (96)	$\nu C = S + \nu CN + \delta_{ring}$	-1.5	STn-C1
					+2.0	STn-C2
+4 (26)	+5 (34)	+4 (37)	+3 (42)	$\delta_{ m ring}$	+2.5	STn-C1
					+4.5	STn-C2
-33 (89)	-36 (64)	-15 (80)	-13 (71)	$\delta \mathrm{CO}_2$	-17.0	STn-C1
					-19.0	STn-C2
+40 (81)	+65 (101)	+3 (138)	+16 (120)	$\gamma NH + \gamma_{ring}$	+28.5	STn-C1
					+30.5	STn-C2

^a Abbreviations: ν – bond stretching, δ – in-plane bending, γ – out-of-plane bending. ^b Positions of the ST monomer bands: 3498.0, 3491.0, 1558.0, 1474.5, 1309.0, 1192.5, 932.5, 662.0, 566.0 cm⁻¹.

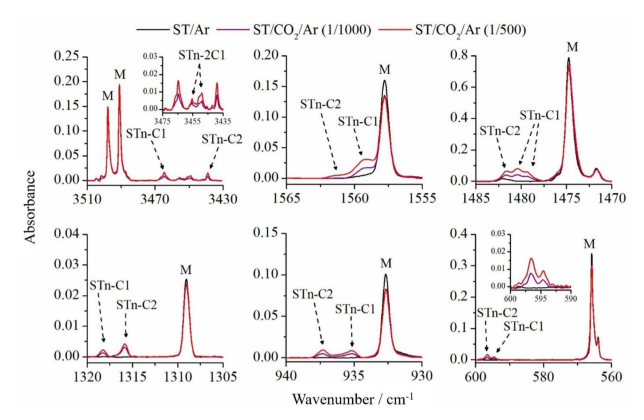


Fig. 4 Selected spectral ranges of $ST/CO_2/Ar$ matrices, where the CO_2/Ar ratio was 1/500 (red line) and 1/1000 (purple line) and, for comparison, the spectrum of the ST/Ar matrix (black line). The letter M shows the ST monomer bands

molecule. Two of them, at 3464.5 and 3439.0 cm⁻¹, are characterized by a constant intensity ratio regardless of the CO2 concentration in the system, which suggests that they can be assigned to 1:1 complexes. The observed wavenumber shifts relative to the STn monomer are equal to -26.5 and -52.0 cm^{-1} . On the other hand, the doublet located at 3455.5/ 3449.0 cm⁻¹ is more intense at the CO₂/Ar mixture concentration of 1/500. This most likely indicates the presence of aggregates of 1:2 stoichiometry.

Based on the results of DFT calculations, the two most stable 1:1 complexes of STn with CO₂, namely STn-C1 and STn-C2, are characterized by −29 and −53 cm⁻¹ shifts in the v(N₄H) stretching region, respectively. These values are in very good agreement with the experimental observation. Both geometries are characterized by the presence of the N-H···O hydrogen bridge and an additional S6···C10 interaction (see Fig. 2). In the remaining two STn···CO₂ complexes (STn-C3 and STn-C4), their NH groups do not participate in the formation of interactions with carbon dioxide, therefore their theoretical shifts of the vN_4H mode are much smaller (0 and -7 cm⁻¹, respectively).

Other vibrations of the ST molecule are also sensitive to complexation, as shown in the remaining spectral regions of Fig. 4 and Table 1. In each region, new bands are present, shifted by a few cm⁻¹ towards higher wavenumbers, and their positions correspond to the predicted shifts for STn-C1 and STn-C2 species. In the range of $1485-1470 \text{ cm}^{-1}$, containing the most intense band of the ST monomer originating from complex stretching and deformation vibrations of the triazole ring, the band of the STn-C1 complex appears as a doublet separated by +4.5/+6.0 cm⁻¹, and the third band (+7.0) can be assigned to the STn-C2 structure. In the remaining parts of the spectrum, the bands of both complexes appear as singlets. The observed shifts of the modes presented in Table 1 fit very well to the corresponding theoretical values obtained for ST-C1 and ST-C2 structures, indicating that both complexes are present in solid argon. It should be noted that the presence of the STn-C3 complex in the matrix cannot be completely ruled out because the DFT-predicted wavenumber shifts are, in many cases, very small or zero, and the bands originating from this structure can be obscured by the ST monomer peaks. On the other hand, much lower energetic stability of this form (9.2 kJ mol⁻¹ difference relative to the most stable complex) indicates that the formation of this structure is unlikely.

As mentioned earlier, two bands were also formed in the vNH range of the spectrum after deposition, which could be assigned to one of the complexes with stoichiometry higher than 1:1. To confirm this interpretation and to reveal more bands originating from these aggregates, the matrix temperature was increased to 32 K and cooled back to 10 K. A change in the intensity of the bands present in the spectrum and the appearance of new ones were observed. As shown in paragraph 3.1.2, the performed calculations reveal three $STn \cdots (CO_2)_2$ structures resulting from the addition of a second CO2 molecule to the 1:1 complexes (see Fig. 2). Fig. 5 shows four spectrum

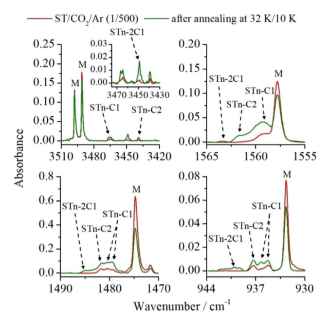


Fig. 5 Selected spectral ranges of ST/CO $_2$ /Ar matrices (CO $_2$ /Ar = 1/500) after deposition (red line) and after annealing the matrix to 32 K/10 K (green line). The letter M shows the ST monomer bands.

fragments, based on which the analysis and discussion on the structure of 1:2 complexes was carried out. The previously unassigned bands in the νNH range are located at 3445.5 and

3449.0 cm $^{-1}$. Upon annealing, the latter was observed to increase its intensity more significantly, which allowed us to assume that both bands originate from the same structure isolated in two different matrix cavities. The second possibility is the presence of overlapping $vN_{(2)}H$ and $vN_{(4)}H$ bands of one of the 1:2 complexes. Comparison of the experimental and theoretical frequencies and intensities allowed us to conclude that the band located at 3449.0 cm $^{-1}$ comes from the $vN_{(4)}H$ vibration, and the other from the $vN_{(2)}H$ vibration of the ST*n*-2C1 form. This is also confirmed by the fact that in other spectral ranges, no new absorptions were observed that could be assigned to another structure with the 1:2 composition. The data is available in Table S4 in the ESI.†

The described analysis at this point does not include complexes of the ST thiol tautomer. This is because this form is present in the matrix in a very small amount after deposition, and the observation of its complexes with the CO_2 molecule is impossible. However, information on the optimized $STI\cdots CO_2$ structures, their geometry, energetics, and spectroscopic parameters (see paragraph 3.1.2.) are important in the context of the next paragraph, where the photoinduced tautomerization of STn and its complexes are discussed.

3.3. Photochemical generation of the thiol form STI complexes with CO₂

Laser irradiation of matrices with UV light induced interesting changes in the ST/CO₂/Ar spectra. Fig. 6 presents several regions

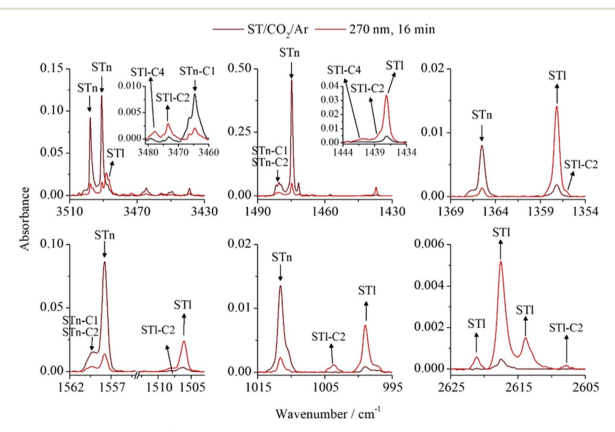


Fig. 6 Selected spectral ranges of the ST/CO₂/Ar matrix (CO₂/Ar = 1/500) after deposition (burgundy line) and after 16 min of irradiation with radiation of wavelength λ = 270 nm (red line).

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of the spectra obtained after 16 min irradiation at 270 nm compared to the spectrum of the matrix after deposition. The formation of the monomer thiol form STl as a result of tautomerization was accompanied by the appearance of additional bands in several spectral ranges. These bands were assigned to the complexes of the thiol form (STI···CO₂), whose structures are presented in Fig. 2. Analyzing individual geometries, it could be predicted that the STn-C1 complex present in the matrix would transform into the STI-C1 form upon irradiation, as this process involves a change in the position of the hydrogen atom H7. In turn, irradiation of the STn-C2 complex will probably lead to the formation of the STl-C2, which is characterized by the lowest energy among all thiol STI···CO₂ complexes. However, despite the lack of the STn-C4 structure after deposition, the appearance of the STl-C4 structure under the influence of irradiation was also considered as a result of attaching a CO2 molecule to the STI monomer formed in the matrix cavity.

The above considerations were assessed by comparing theoretical and experimental wavenumber shifts of the new bands (see Table 2), which are located near the analogous absorptions of the STl monomer. In the range of 3510-3430 cm⁻¹, bands originating from the following structures are present before irradiation: STn-C1 (3464.5 cm⁻¹), STn-C2 (3439.0 cm^{-1}) , and STn-2C1 (doublet at 3455.5/3449.0 cm⁻¹). All of them decreased in intensity upon irradiation and, in addition to a band located at 3488.0 cm⁻¹ assigned to the ST1 monomer, a doublet appeared, which is shifted relative to the monomer by -10.0/-15.0 cm⁻¹. These values correspond to the theoretical predictions for the vNH vibrations of the STl-C1 and STI-C4 complexes. The NH group of the third structure (STI-C2) is not involved in the interaction, therefore, the calculated vNH shift is equal to zero, and this band is not observed. In the range of 1490-1430 cm⁻¹, where there are bands from complex stretching and deformation vibrations of the triazole ring, two new bands of low intensity appear. One of them occurs in the form of a shoulder on the STI monomer band ($\Delta \nu = +1.0 \text{ cm}^{-1}$), while the other is located away from the monomer absorption by +4.0 cm⁻¹. These bands can be assigned to the STl-C2 and STI-C4 structures, respectively, based on the calculated shift values. The remaining spectral regions contain one new band

each, whose shifts best correspond to the values predicted for the STI-C2 structure. This suggests that this complex is predominant in the matrix after irradiation. It should also be mentioned that despite the presence of one STn geometry with a stoichiometry of 1:2 in the matrix after deposition, no bands were identified in the spectrum after irradiation that could originate from its thiol counterpart, most likely due to the low efficiency of photochemical conversion or a small amount of these species in the matrices.

3.4. Effect of ST complexation with CO₂ on the efficiency of thione-thiol tautomerization

Hydrogen atom transfer upon UV radiation is quite often observed in low-temperature matrices. In their latest review on this topic, Lapinski et al. 19 drew attention to the influence of matrix material on the H-atom transfer phenomenon. What is more, results of our recent paper on ST···N2 complexes16 showed that the efficiency of hydrogen atom transfer in ST complexed with N2 is lower than in the monomeric form. Moving forward, it seemed interesting to investigate how the specific 1:1 interaction between STn and CO_2 influences the described thione \rightarrow thiol transformation. For this purpose, the experimental integrated intensity ratios of the vN2H bands were determined for STn and STl monomers as well as for the STn-C2 and STl-C2 complexes. The ST/Ar and ST/CO2/Ar spectra recorded before and after 16 min of irradiation with radiation of $\lambda = 270$ nm were used. As can be seen in Table S5 (ESI),† the vN₂H band intensity ratio equals 0.400 for monomers and 0.318 for complexes. Thus, the specific interaction of ST with the CO₂ molecule causes the transfer of the hydrogen atom in the complex to be about 20% less efficient than in the case of the monomer. This difference is smaller than in the case of the interaction with the nitrogen molecule, which indicates that despite the formation of stronger interactions between the ST and CO₂ subunits, they disturb the kinetics of the photochemical transformation to a lower extent. Changes in the concentration of individual monomers and complexes are presented in Fig. 7.

Analysis of the ST/Ar and ST/CO2/Ar spectra before and after irradiation revealed additional insights into how changes in bulk interactions, caused by the admixture of CO2 in the argon

Table 2 Selected shifts of wavenumbers $\Delta \nu$ (cm⁻¹) and band intensities (km mol⁻¹, in brackets) calculated by the B3LYP-D3/6-311++G(3df,3pd) method for the 1:1 complexes of the thiol tautomer ST (STI) with a CO₂ molecule, compared with the experimental shifts. The values of the wavenumbers of the monomer bands, with respect to which the shifts were calculated, are given in brackets in the rightmost column

Theoretical shifts $(\Delta \bar{ u})$					
STl-C2	STl-C4	STl-C1	$Mode^{16,a}$	Experimental shifts ^b	Assignment
0 (87)	-16 (130)	-7 (147)	vN_2H	-10.0	STl-C2
,	, ,	,	-	-15.0	STl-C4
-7(48)	0 (8)	+1 (7)	ν SH	-9.5	STl-C2
+2 (43)	-1(52)	+1 (61)	$vCN + \delta NH + \delta CN$	+2.0	STl-C2
+2 (87)	+4 (131)	+2 (77)	$vCN + \delta NH + \delta CN + \delta_{ring}$	+4.0	STl-C2
+1 (36)	-3(39)	-1(37)	v CN + δ_{ring}	-1.0	STl-C2
+4 (40)	+1 (59)	+2 (35)	δ CN + δ _{ring}	+4.5	STl-C2

^a Abbreviations: ν – bond stretching, δ – in-plane bending. ^b Positions of the ST monomer bands: 3488.0, 2617.5, 1506.0, 1437.0, 1357.0, 999.0 cm⁻¹.

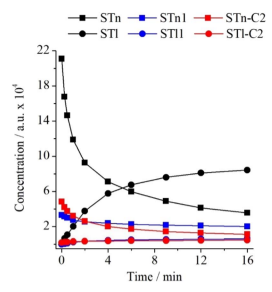


Fig. 7 Changes in the intensity of the νN_2H bands of ST monomers in the argon matrix and their complexes in the matrix doped with N_2 or CO_2 as a function of irradiation time at $\lambda=270$ nm.

matrix, affect the thione \rightarrow thiol transformation of the ST monomer. The intensity ratios calculated based on the spectra of the ST/Ar and ST/CO₂/Ar matrices (CO₂/Ar = 1/500) are presented in Table S6 (ESI).† It presents that, in comparison to the nitrogen admixture, the presence of CO₂ molecules in the STn environment has a less significant effect on the discussed photochemical reaction in the unbound ST. As shown in Fig. 8 below, one can notice that the decrease of STn and the simultaneous increase of STl are less substantial than in the case of the N₂-containing matrices.

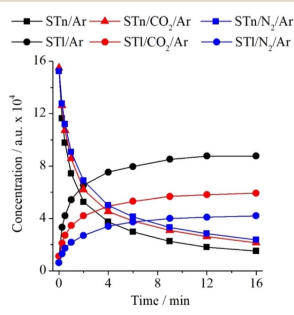


Fig. 8 Changes in the intensity of the most intense bands of STn and STl monomers (1474.5 cm $^{-1}$ and 1437.0 cm $^{-1}$, respectively) in ST/Ar, ST/N₂/Ar, and ST/CO₂/Ar matrices as a function of irradiation time at λ = 270 nm.

4. Conclusions

Building on our previous work on 3-thio-1,2,4-triazole (ST) complexes with dinitrogen,16 the study of ST complexes with carbon dioxide offers a compelling extension to explore the role of non-covalent interactions in atmospheric and environmental chemistry. Unlike N2, which primarily interacts via weak van der Waals forces and specific hydrogen bonds, CO₂'s linear geometry and quadrupolar nature allow for diverse interaction modes, including C=O···H hydrogen bonding and Lewis acidbase interactions with electron-rich sites on ST. Our findings reveal that the presence of CO2 in the matrix influences the photochemical behavior of ST, particularly its thione-thiol tautomerization. While both N2 and CO2 reduce the efficiency of this transformation compared to the isolated monomer, the effect of CO2 is notably less pronounced despite its stronger binding interactions. This suggests that the extent to which a gas-phase species modulates photochemical processes in lowtemperature matrices depends not only on interaction energy but also on the geometry and specificity of intermolecular contacts. In this context, the current findings provide important insight into how CO2 modulates tautomerization via directional, non-covalent interactions. Although ST···CO2 complexes exhibit stronger binding than their N2 counterparts, the reduction in tautomerization efficiency remains modest, underscoring that interaction geometry plays a more decisive role than energetic strength alone. Complexes such as STn-C2, stabilized by hydrogen bonds and S...C van der Waals contacts, subtly hinder hydrogen transfer without entirely suppressing reactivity. These results highlight that photochemical reactivity in matrix-isolated systems is governed by the orientation and distribution of interactions, not solely their binding strength. Given the importance of CO₂ in environmental chemistry and the growing interest in heterocyclic frameworks for CO2 capture, the findings presented here not only deepen our understanding of weak intermolecular forces but also shed light on CO2-specific effects in photochemical and structural transformations of heterocycles. This study, therefore, expands the scope of previous investigations by demonstrating how differences in gas-matrix interactions can reshape molecular reactivity and photochemistry.

Data availability

Data for this article, including spectra (.dat) and optimized structures (.log), are available at the Zenodo repository at https://doi.org/10.5281/zenodo.15101158.

Author contributions

K. Mucha: conceptualization, methodology, data curation, investigation, visualization, writing – original draft, writing – eeview & editing. M. Pagacz-Kostrzewa: investigation, formal analysis, visualization; M. Wierzejewska: writing – original draft, supervision.

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Conflicts of interest

There are no conflicts to declare.

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