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Relative orientation of the carbonyl groups determines the nature of orbital interactions in carbonyl–carbonyl short contacts†

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Carbonyl–carbonyl (CO⋯CO) interactions are emerging noncovalent interactions found in many small molecules, polyesters, peptides and proteins. However, little is known about the effect of the relative orientation of the two carbonyl groups on the nature of these interactions. Herein, we first show that simple homodimers of acetone and formaldehyde can serve as models to understand the effect of relative orientations of the two carbonyl groups on the nature of CO⋯CO interactions. Further, from a comprehensive statistical analysis of molecules having inter- or intramolecular CO⋯CO interactions, we show that the molecules can be broadly categorized into six different structural motifs (I–VI). The analysis of pyramidity of the acceptor carbon atoms in these motifs and natural bond orbital (NBO) analysis suggest that the relative orientation of the two interacting carbonyl groups determines whether the orbital interaction between the two carbonyl groups would be $n \rightarrow \pi^*$ or $\pi \rightarrow \pi^*$ or a combination of both.

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Introduction

Noncovalent interactions mediated by carbonyl groups are important in both chemistry and biology. The carbonyl group is polar and has a strong dipole moment. Therefore, it can participate in electrostatic interactions through its positively charged carbon and negatively charged oxygen centers as well as can involve in dipolar interactions. Further, the lone pairs of oxygen and the antibonding π orbital of the C=O bond of the carbonyl group can be involved in orbital interactions with appropriate acceptor and donor atom orbitals, respectively. These properties make the carbonyl group unique and it can participate in many distinct noncovalent interactions such as hydrogen bonding¹ (C=O⋯H), carbonyl–chalcogen interactions² (C=O⋯X; X = S, Se, and Te) and nucleophile–carbonyl interactions³ (Nu⋯CO) in chemical and biological systems. While hydrogen bonding and carbonyl–chalcogen interactions are well studied, relatively less is known about Nu⋯CO interactions. In their seminal work,⁴ Burgi *et al.* showed that various nucleophilic atoms noncovalently interact with the carbonyl carbon atom in a specific trajectory [\angle Nu–C=O \sim 109°; Burgi–Dunitz (BD) trajectory], which laid the foundation for our current understanding of the mechanisms of nucleophilic addition and substitution reactions at the carbonyl centers. When the nucleophile is another carbonyl group, the

interaction is termed as a carbonyl–carbonyl (CO⋯CO) interaction. This interaction is important as it can control the geometries and functions of molecules, especially when they have multiple carbonyl groups such as polyesters, peptoids, peptides and proteins.

CO⋯CO short contacts were first observed by Bolton in a glucose analogue named alloxan.⁵ He reported intermolecular O⋯C distances much shorter than the sum of van der Waals radii (3.22 Å)⁶ between the oxygen and carbon atoms of the carbonyl groups of two neighboring alloxan molecules in the solid state. Interestingly, in alloxan, the CO⋯CO interaction determined the crystal packing even in the presence of strong hydrogen bond donor (NH) and acceptor (C=O) moieties. Several subsequent structural studies established that two carbonyl groups can interact with each other in three possible ways leading to structural motifs I–III shown in Fig. 1A–C.⁷ Maccallum *et al.* demonstrated the importance of the “*intramolecular*” CO⋯CO interaction between proximal amide groups as an important factor in stabilizing α -helices, β -sheets and the right-hand twist often observed in β -strands of proteins.⁸ Later on, Raines and co-workers showed that perturbations of CO⋯CO interactions could affect the stability of proteins.⁹ Since then, there have been extensive studies on intramolecular CO⋯CO interactions in small molecules, peptides, polyesters and proteins.¹⁰ It is now well established that intramolecular CO⋯CO interactions are predominant in α -helices.^{10j,k} Recently, the “reciprocal” variant of the CO⋯CO interaction has been discovered in small molecules and proteins.¹¹

There are two schools of thought on the nature of CO⋯CO interactions. Bolton, Allen and others considered intermolecular CO⋯CO interactions as dipolar interactions.¹² Maccallum

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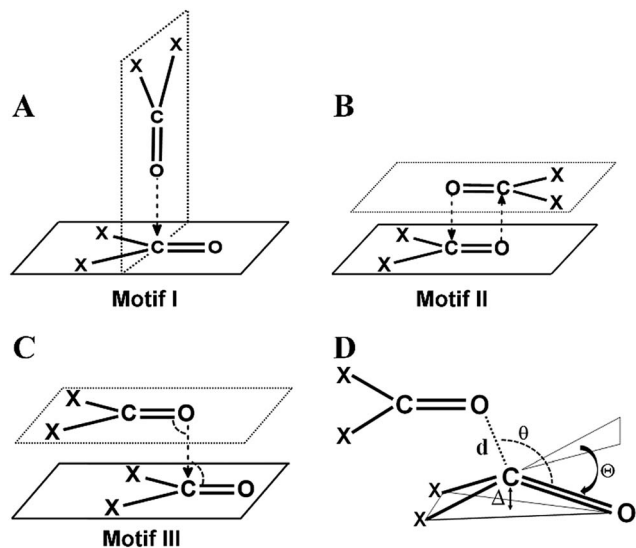


Fig. 1 Different structural motifs previously observed by Allen *et al.* in intermolecular CO \cdots CO interactions in ketone dimers (A) perpendicular motif I, (B) antiparallel motif II, (C) sheared parallel motif III and (D) various structural parameters that are indicative of $n \rightarrow \pi^*$ interactions.

et al. also considered intramolecular CO \cdots CO interactions in proteins and peptides to be dipolar in nature. However, Raines and co-workers showed that intramolecular CO \cdots CO interactions are stabilized by $n \rightarrow \pi^*$ interactions where electron delocalization occurs between the p-character lone pair of a donor carbonyl oxygen atom and the $\pi^*_{C=O}$ orbital of the C=O bond of a nearby acceptor carbonyl group.¹³ The $n \rightarrow \pi^*$ nature of the CO \cdots CO interaction in an intramolecular system was subsequently supported by many other researchers working in this area.¹⁰ The reciprocal variants of intramolecular CO \cdots CO interactions were reported to have contributions from both $n \rightarrow \pi^*$ and $\pi \rightarrow \pi^*$ orbital interactions.¹¹

The questions that then arise are “Are inter- and intramolecular CO \cdots CO interactions different in nature? Do the relative orientations of the two interacting carbonyl groups in inter- and intramolecular systems affect the nature of CO \cdots CO interactions?” To address these questions, it would be necessary to first find out all the possible structural motifs in molecules having CO \cdots CO interactions in both inter- and intramolecular systems. The recent discovery of a reciprocal variant of the CO \cdots CO interaction indicates that we may not yet completely know all the possible structural motifs of CO \cdots CO interactions for inter- and intramolecular systems. However, to the best of our knowledge, there are no systematic studies on CO \cdots CO interactions in both inter- and intramolecular systems. In this paper, we have studied both inter- and intramolecular CO \cdots CO interactions using statistical and theoretical methods. Herein, we have first studied the homodimers of acetone and formaldehyde to understand how different orientations of the two interacting carbonyl groups affect the nature of CO \cdots CO interactions. Further, we have identified the various structural motifs of inter- and intramolecular CO \cdots CO interactions and showed that the nature of orbital interactions for the

stabilization of CO \cdots CO interactions varies with the relative geometrical orientations of the two interacting carbonyl groups in these motifs. Further, from gas-phase computational and solid-state structural studies we show that the nature of orbital interactions in these motifs can be effectively modelled by using appropriate conformations of homodimers of acetone and formaldehyde.

Results and discussion

1 CO \cdots CO interactions in acetone and formaldehyde homodimers

This work is inspired by our observation that the two simple homodimeric carbonyl systems, the acetone dimer and the formaldehyde dimer, have very different arrangements of the carbonyl groups in their most stable conformations in the gas phase. Previous studies have shown that the antiparallel C_{2h} symmetric structure of the acetone dimer that resembles motif II is the global minimum.^{14a} For the formaldehyde dimer, the global minimum is a C_s symmetric structure where the molecular plane of one molecule of formaldehyde is orthogonal to the molecular plane of the second formaldehyde molecule.^{14b-d} In view of their well-established global minimum structures, we avoided carrying out detailed conformational analysis of these dimeric systems and looked only into the C_{2h} symmetric structure of the acetone dimer and the C_s symmetric structure of the formaldehyde dimer for further studies. We optimized these two dimer structures using the MP2/6-311+G(2d,p) level of theory (Fig. 2A–E). Natural bond orbital (NBO)¹⁵ analyses were carried out to investigate the orbital interactions in these molecules. We observed that the major orbital interaction for the stabilization of the acetone dimer comes from the reciprocal $\pi \rightarrow \pi^*$ orbital interactions (2.06 kcal mol $^{-1}$) of the filled π orbital of one acetone molecule with the vacant π^* orbital of the other. There is a relatively small contribution from $n \rightarrow \pi^*$ interactions (0.14 kcal mol $^{-1}$) to the stabilization of the acetone dimer. The BSSE-corrected¹⁶ binding energy of the acetone dimer (5.65 kcal mol $^{-1}$) is much higher than the sum of orbital interaction energies, which suggests that other interactions such as dipole–dipole or electrostatic interactions may also be important for the stabilization of the system. In the formaldehyde dimer on the other hand, the major orbital contribution to the CO \cdots CO interaction comes from $n \rightarrow \pi^*$ electron delocalization of the lone pair of oxygen of one formaldehyde to the π^* orbital of the other formaldehyde C=O bond (3.9 kcal mol $^{-1}$) and a relatively small contribution from $\pi \rightarrow \pi^*$ orbital interactions (0.26 kcal mol $^{-1}$). In addition, there is a C–H \cdots O hydrogen bond that holds the two formaldehyde molecules together (0.27 kcal mol $^{-1}$). The low value of the BSSE-corrected binding energy (3.52 kcal mol $^{-1}$) of the formaldehyde dimer suggests that other interaction energies such as dipole–dipole and electrostatic interactions should contribute little to the stabilization of the system. Atoms in molecules (AIM)¹⁷ analysis of the two dimer complexes clearly shows that the bond path in the acetone dimer is similar to that of a typical $\pi \rightarrow \pi^*$ interaction and that of the formaldehyde dimer is similar to that of the $n \rightarrow \pi^*$ interaction (Fig. 2C and D). In the acetone dimer,





Fig. 2 Optimized structures, NBO overlap diagrams and AIM structures of the homodimers of acetone and formaldehyde. (A) Optimized structure of the acetone dimer; (B) $n \rightarrow \pi^*$ interaction in the acetone dimer; (C) $\pi \rightarrow \pi^*$ interaction in the acetone dimer; (D) AIM structure of the acetone dimer showing the bond paths and various critical points; (E) optimized structure of the formaldehyde dimer; (F) $n \rightarrow \pi^*$ interaction in the formaldehyde dimer; (G) $n \rightarrow \sigma^*$ interaction in the formaldehyde dimer; (H) AIM structure of the formaldehyde dimer showing the bond paths and various critical points.

the $n \rightarrow \pi^*$ interaction involves the sp -character lone pair on the carbonyl oxygen atom. On the other hand, in the formaldehyde dimer, the participating lone pair of the carbonyl oxygen atom is of p -character as seen from the NBO calculations.

These observations suggest that the nature of the orbital interaction in the $\text{CO} \cdots \text{CO}$ interaction is quite different in the homodimers of acetone and formaldehyde and it is clearly dependent on the orientation of the monomers with respect to each other. To understand how the nature of orbital interactions changes with change in relative orientation of the two carbonyl groups, we carried out relaxed potential energy surface (PES) scans on the acetone dimer to assess geometries resembling motifs I, II and III. Further, we looked into the orbital nature of the $\text{CO} \cdots \text{CO}$ interaction in every geometry in the interconversion pathways for motifs I, II and III. The motif II geometry of acetone ($\text{X} = \text{CH}_3$) can be converted to motif III by changing the $\text{C}=\text{O} \cdots \text{C}=\text{O}$ dihedral angle (T) whereas the motif III geometry can be converted to motif I by changing the bond angle θ_3 or θ_4 (Fig. 3; $\text{X} = \text{CH}_3$). First, we scanned the $\text{C}=\text{O} \cdots \text{C}=\text{O}$ dihedral angle (T) starting from the antiparallel geometry [motif II; $T = 0^\circ$] to the parallel geometry [motif III; $T = 180^\circ$] at an increment of 20° at the MP2/6-311+G(2d,p) level of theory. The scans were done for three different intermolecular $\text{O} \cdots \text{C}$ distances (d) of 2.90 Å, 3.00 Å, and 3.09 Å and orbital interactions for each intermediate geometry were analyzed by NBO. NBO analyses indicated that during the interconversion of II to III, $\pi \rightarrow \pi^*$ orbital interactions gradually decreased with a concomitant increase in the $n \rightarrow \pi^*$ orbital interactions as shown in Fig. 3B. This study indicated that the orientation of the two acetone monomers dictates whether the carbonyl groups would be involved in $\pi \rightarrow \pi^*$ or $n \rightarrow \pi^*$ orbital interactions. We then carried out a relaxed potential energy scan by

changing the $\text{C}=\text{O} \cdots \text{C}$ bond angle (θ_3 or θ_4) from 90° (motif I) to 170° (motif III) for three $\text{O} \cdots \text{C}$ distances (d) of 2.90 Å, 3.00 Å, and 3.10 Å and carried out NBO analyses on each geometry along the path. As expected from the geometry, NBO analyses indicated that the $\pi \rightarrow \pi^*$ interaction present in the parallel motif III diminished completely in the T-shaped motif I and the $n \rightarrow \pi^*$ orbital interaction became the sole contributor to the stabilization of the $\text{CO} \cdots \text{CO}$ interaction (Fig. 3C). Taken together, the PES scan and NBO analyses suggest that the $\pi \rightarrow \pi^*$ interaction gradually decreases and the $n \rightarrow \pi^*$ interaction increases from the antiparallel (II) to the parallel (III) motif and it becomes solely the $n \rightarrow \pi^*$ interaction in T-shaped motif I.

During the PES scan of the acetone dimer from the antiparallel (II) to the parallel (III) geometry [$T = 0^\circ$ to $T = 180^\circ$], we observed a unique “L-shaped” structure at an intermediate $\text{C}=\text{O} \cdots \text{C}=\text{O}$ dihedral angle of $T = 90^\circ$ (Fig. 3A, motif IV). To get better insight into this geometry, we carried out geometry optimization of the acetone dimer by freezing the dihedral angle (T) at 90° . The L-shaped local minimum for $T = 90^\circ$ was obtained with one $\text{O} \cdots \text{C}$ (d) contact of 3.109 Å and $\theta = 97.4^\circ$ (Fig. 3D). NBO analysis of the structure showed the presence of both $n \rightarrow \pi^*$ [$0.30 \text{ kcal mol}^{-1}$] and $\pi \rightarrow \pi^*$ [$0.61 \text{ kcal mol}^{-1}$] interactions. This could be a unique dimeric structure for ketones, however, Allen *et al.* in their CSD analysis of ketone dimers did not observe such a motif.

The formaldehyde dimer structure is unique due to the presence of both $n \rightarrow \pi^*$ interactions and $\text{C}-\text{H} \cdots \text{O}$ hydrogen bonding. As their study was restricted to ketone dimers that lacked hydrogen bond donors, Allen *et al.* did not observe such a geometry.^{7d} We envisage that aldehydes, carboxylic acids and amides that have hydrogen bond donors may form structures similar to that of the formaldehyde dimer. The relaxed potential



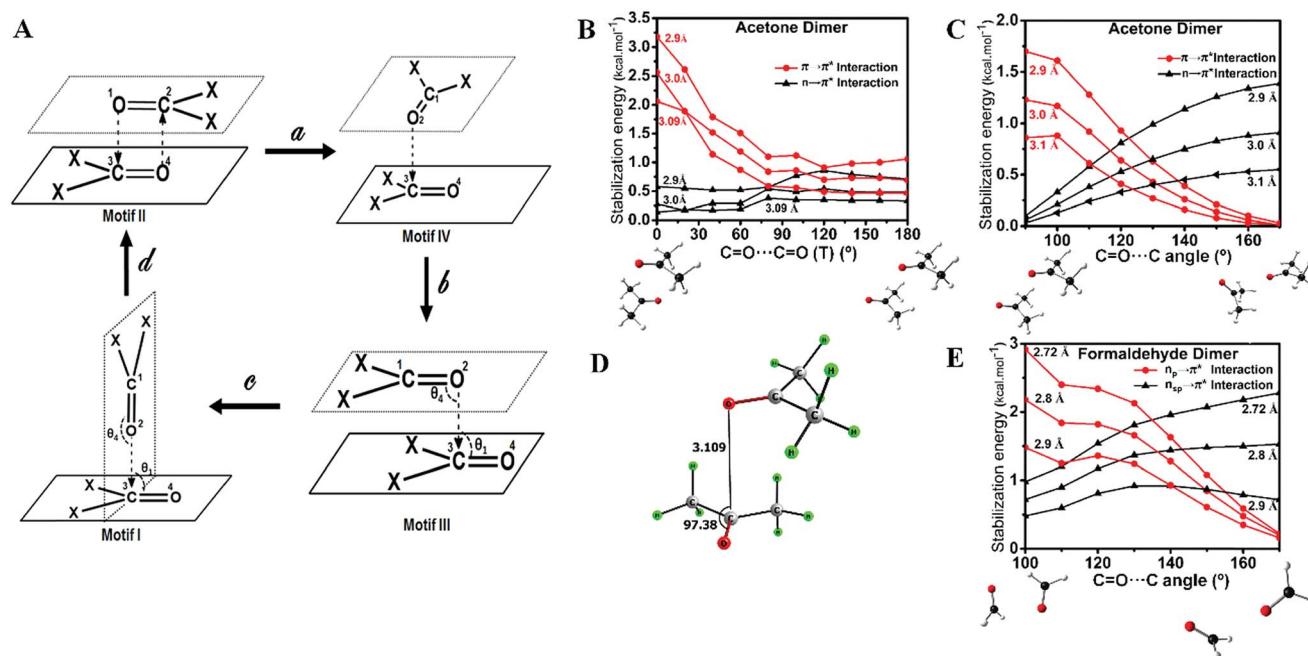


Fig. 3 Relaxed potential energy surface (PES) scans and NBO analyses of the homodimers of acetone and formaldehyde and evaluation of orbital interactions. (A) Mechanism of interconversion between motifs I–III. We observed a unique “L-shaped” structure at an intermediate $C=O\cdots C=O$ dihedral angle of $T = 90^\circ$. (B) $C=O\cdots C=O$ (T) show that the $\pi \rightarrow \pi^*$ interaction gradually decreases and the $n \rightarrow \pi^*$ interaction gradually increases from the antiparallel (II) to the parallel motif (III). (C) NBO analyses of the various geometries of the acetone dimer obtained during relaxed scan of $C=O\cdots C$ angle show that the $\pi \rightarrow \pi^*$ interaction gradually decreases and the $n \rightarrow \pi^*$ interaction gradually increases from the perpendicular T-shaped geometry (I) to the parallel geometry (III). (D) Novel L-shaped geometry of the acetone dimer observed during the relaxed scan of T (motif IV). (E) NBO analyses of various geometries obtained during the relaxed scan of the $C=O\cdots C$ angle of the formaldehyde dimer show that the character of the oxygen lone pair participating in the $n \rightarrow \pi^*$ interaction gradually changes from p to sp-character.

energy surface scan to convert the formaldehyde dimer to a T-shaped motif (II) brought out an interesting observation. NBO analyses indicated that the nature of the oxygen lone pair involved in $n \rightarrow \pi^*$ orbital interactions gradually changed from p- to sp-character (Fig. 3E) suggesting that $n \rightarrow \pi^*$ orbital interactions can involve either the sp- or p-character lone pair of the donor oxygen atom depending on the orientation of the two monomers that participate in interactions.

2 Cambridge structural database analysis

Inspired by the above results, we carried out a Cambridge Structural Database¹⁸ (CSD) search and extracted all the structures having inter- and intramolecular $CO\cdots CO$ interactions for all kinds of carbonyl groups. We searched for organic molecules having inter- and intramolecular $CO\cdots CO$ short contacts by using a restriction of 3.20 Å to the nonbonded $C\cdots O$ atomic distances [sum of the van der Waals radii of C and O is 3.22 Å].⁶ We looked for two different modes of $CO\cdots CO$ short contacts (a) one-sided interaction where one carbonyl donates and the other carbonyl accepts [only one $O\cdots C$ distance (either d_1 or d_2) is ≤ 3.20 Å] and (b) reciprocal $CO\cdots CO$ interaction where both the carbonyl groups work as donors and acceptors (both d_1 and d_2 are ≤ 3.20 Å) (Fig. 4A). For one-sided interaction, the ‘group-group (G–G)’ search facility of the CSD was utilized. Atoms C^1 and O^2 were defined as G1, and C^3 and O^4 were defined as G2 (Fig. 4A). The search for the nonbonded distance between the

centroids of G1 and G2 was repeated three times using $d(G1\cdots G2) \leq 3.20$ Å, 4.00 Å and 5.50 Å. The data were combined and duplicate values were omitted. Only those molecules having the nonbonded distance $O^2\cdots C^3$ (d_1) or $O^4\cdots C^1$ (d_2) ≤ 3.20 Å were considered for this study. For reciprocal interactions, both the nonbonded distance d_1 and d_2 were kept ≤ 3.20 Å and the search was carried out for both intermolecular and intramolecular cases. For the intramolecular system, the software considers the minimum bond path between C and O atoms.

2(a) Classification of structures into different motifs. From the CSD search, we obtained 10 858 molecules out of which 4237 contained intermolecular and 6621 contained intramolecular nonbonded $C\cdots O$ contacts of ≤ 3.20 Å. For the intermolecular case, we observed that out of 4237 molecules, 2897 (68%) contained one-sided interactions where only one of the $C\cdots O$ distances is within 3.20 Å while 1340 (32%) molecules contained reciprocal interactions where both $C\cdots O$ distances were within 3.20 Å (Table 1). In the intramolecular case, 4749 (76%) molecules contained one-sided interactions while 1472 (24%) molecules were found to have reciprocal interactions. The shortest $O\cdots C$ distance in the intermolecular one-sided interactions is ~ 2.72 Å, whereas the shortest $O\cdots C$ distance in the intramolecular one-sided interactions is ~ 2.44 Å. The average $O\cdots C$ distance (2.91 Å) is shorter for intramolecular interactions compared to intermolecular interactions (3.06 Å), which suggests that intramolecular interactions are generally





Fig. 4 Chemical structures of some motifs newly discovered in this study. (A) Fragment having $C=O\cdots C=O$ short contacts used for the CSD search. Torsion angle, T is defined as $C^1=O^2\cdots C^3=O^4$. Chemical structures of newly discovered (B) motif IV (C) motif V and (D) motif VI. (E) Plot showing how the number of molecules decreases for intramolecular one-sided and reciprocal interactions as the bond separation between the two carbonyl groups increases. In $1,x$ -type interaction x represents the size of the ring to be formed due to the interaction. For example, 1,5-type interaction means there are 3 bonds separating the carbonyl groups and a 5-membered ring will be formed due to the noncovalent interaction.

stronger than intermolecular interactions. Also, the average $O\cdots C$ distance in reciprocal interactions is longer than that in one-sided interactions (Table 1). Therefore, the individual interaction of one carbonyl group with the other should be relatively weak for reciprocal interactions compared to one-sided interactions. However, the overall effect of the back and forth interactions between the two carbonyl groups should contribute a significant amount to the stabilization of the molecules. The statistical data are summarized in Fig. S2–S6.† The average value of the $O\cdots C=O$ angle in the intermolecular case is 92.6°

whereas for the intramolecular case it is 94.7° , which are smaller than that in the Burgi–Dunitz trajectory.

The motif structures (motifs I–V) were categorized based on the $C=O\cdots C=O$ dihedral angle T , and the two $C=O\cdots C$ angles θ_3 and θ_4 . For motif VI, the carbonyl planes were also taken into account. The different parameters for an ideal geometry of each motif type are shown in Table S2† and the criteria for their categorization into various motifs are discussed in the ESI.† In the following section, molecules from the motifs I–VI will be divided into two major classes (i) molecules having one-sided $CO\cdots CO$ interactions and molecules with (ii) reciprocal $CO\cdots CO$ interactions.

2(a) (i) *One-sided interaction (motifs I, III and IV)*. Molecules having one-sided interactions could be categorized into three different structural motifs I, III and IV, irrespective of whether the interactions are inter- or intramolecular. In intermolecular cases, most of the 2897 molecules with one-sided $CO\cdots CO$ interactions have geometries represented by either perpendicular motif I [$\angle C=O\cdots C \sim 150\text{--}180^\circ$] or parallel motif III [$C=O\cdots C=O$ (T) = 180° and $\angle C=O\cdots C = 90^\circ$ or 50°] observed previously by Allen and co-workers for ketone dimers. These motifs are often found in the crystal structures where the carbonyl group(s) are attached to bulky groups to avoid steric clashes. We also observed many molecules with a novel L-shaped geometry with $C=O\cdots C=O$ torsion angles (T) $\sim 90^\circ$ as discussed above for the acetone dimer. These molecules cannot be categorized into motif I or III. We identified ~ 400 such molecules, represented by the structural motif IV. In motif IV, one carbonyl group sits over the other carbonyl group in an “L-shaped” manner facilitating the nonbonded $O\cdots C$ short contact. These dimeric structures have angles ($\theta_1\text{--}\theta_4$) similar to that of motif III but unlike motif III where the $C=O\cdots C=O$ torsion angle (T) is $\sim 180^\circ$, they have torsion angles of $\sim 90^\circ$.

For intramolecular cases, 1059 molecules with one-sided interactions were found to have a torsion angle $|T| \sim 180^\circ$, which resembles that of the sheared parallel motif III. In contrast to the intermolecular case, only a few molecules (14) with a T-shaped geometry ($\angle C=O\cdots C = 150\text{--}180^\circ$) were observed having one-sided intramolecular interactions. This is expected given the steric crowding and bond constraints that arise in such geometries. The majority of the molecules (1762) can be categorized into the “L-shaped” motif IV with the $C=O\cdots C=O$ torsion angle $|T|$ close to 90° . Interestingly, most of the intramolecular interactions are 1,5-type in nature where the two carbonyl groups were separated by three covalent bonds. It should be noted that all the backbone carbonyl–carbonyl interactions in proteins are also 1,5-type in nature. As the

Table 1 Summary of various structural parameters for molecules involved in the $CO\cdots CO$ interaction in inter- and intramolecular systems^a

	Total	One-sided	d_{\min} (Å)	d_{avg} (Å)	θ_{avg} (deg)	Reciprocal	d_{\min} (Å)	d_{avg} (Å)	θ_{avg} (deg)
Intermolecular	4237	2897	2.72	3.06	95.5	1340	2.81	3.10	89.8
Intramolecular	6221	4749	2.44	2.91	103.8	1472	2.31	3.00	85.7

^a d_{\min} – minimum $O\cdots C$ distance; d_{avg} – average $O\cdots C$ distance; θ_{avg} – average $O\cdots C=O$ angle.



number of bonds separating the two carbonyl groups increases, the number of molecules steeply decreases (Fig. 4E).

2(a) (ii) Reciprocal interaction (motifs II and V). In the intermolecular system, the majority of the molecules have reciprocal interactions (motif II; $|T| \sim 0^\circ$ and $\angle C=O \cdots C \sim 90^\circ$) with sheared antiparallel orientation, which is similar to the global minimum geometry of the acetone dimer. In intramolecular cases, the reciprocal interactions were observed with a $C=O \cdots C=O$ dihedral angle (T) of $\sim 70^\circ$ and two $\angle C=O \cdots C$ angles $\sim 75^\circ$ (motif V). Rahim *et al.* recently reported that donor oxygen atoms approach the acceptor carbonyl carbon atoms significantly away from the BD trajectory ($\angle O \cdots C=O$ is $\sim 85^\circ$) in these cases.¹¹ Motif V has a close resemblance to motif IV, the major difference being, in motif V, both $O \cdots C$ distances are shorter than 3.20 Å whereas only one of the $O \cdots C$ distances is shorter than 3.20 Å in motif IV. Structurally motif V is more twisted than motif IV due to the restriction on both the $O \cdots C$ distances in motif V (< 3.20 Å).

Note that there is a striking difference in the geometrical arrangements of the two carbonyl groups in intramolecular and intermolecular reciprocal interactions. In the intermolecular case, reciprocal interactions were observed in the antiparallel motif II ($T \sim 0^\circ$) whereas intramolecular reciprocal interactions are observed in “twisted” geometries with $T = \sim 70^\circ$ that we have represented as motif V in this paper. Similar to one-sided interactions, reciprocal interactions can also be categorized as separate 1,x-types in the intramolecular case (Fig. 4E). Interestingly, as the bond separation between the two carbonyl groups increases, the number of molecules decreases and the “twisted” bond type motif V tends to slowly convert to an antiparallel motif II (ESI Fig. S10†).

2(a) (iii) Unprecedented structural motif VI that resembles the formaldehyde dimer. As discussed above, the formaldehyde dimer does not fall in any of the aforementioned categories. We were curious to see if such molecules are also present in the CSD. We could envisage such a motif in aldehydes, carboxylic acids or amides wherein a carbonyl group can engage in one $O \cdots C$ short contact and one $O \cdots H$ short contact as shown in Fig. 4D and ESI Fig. S7.† We have represented this type of structure as motif VI in this paper. Though the carbonyl groups are antiparallel to each other the H-bonding imposes restriction on the O atom to interact with the C attached to the hydrogen producing a separate geometry, where $\angle C=O \cdots C$ is neither 90° nor 180° . For this kind of structure, the carbonyl planes are perpendicular to each other. A total of 17 aldehydes [motif VI(a)], 7 carboxylic acids [motif VI(b)] and 24 amides [motif VI(c)] were found in the CSD search that have a geometry similar to that of motif VI. In the search, the software was allowed to normalize the hydrogens and one of the two carbonyl groups was considered as aldehyde or carboxylic acid or amide in the CSD search fragment. We observed that motif VI is unique to intermolecular systems and we did not find any molecule with the intramolecular $CO \cdots CO$ interaction having a geometry similar to that of motif VI. This could be due to the probable steric restrictions associated with such an intramolecular system.

We observed that motifs III and IV are common to both inter- and intramolecular systems. However, there are very few

molecules having intramolecular $CO \cdots CO$ interactions that can be categorized into motif I or II whereas there is no molecule of motif VI type with intramolecular $CO \cdots CO$ interactions. On the other hand, motif V is only present in the intramolecular system and there are very few molecules with the intermolecular $CO \cdots CO$ interaction that could be categorized into motif V.

2(b) Approach of the donor oxygen to the acceptor carbon atom. The distribution of the $O \cdots C=O$ (θ) angle in inter- and intramolecular cases is shown in Fig. 5A and B. As can be seen from Fig. 5A, θ has a maximum of $\sim 90^\circ$ in the intermolecular cases, while the maximum of θ lies close to $\sim 100^\circ$ for intramolecular interactions (Fig. 5B) suggesting that the approach of donor oxygen to acceptor carbon is closer to the Burgi–Dunitz trajectory for intramolecular cases. A closer inspection of the θ values in individual motifs (I–VI) shows that the majority of the molecules in each motif have a θ value less than 100 except for motif III (Fig. 5E). However, for all types of motifs, as the $O \cdots C$ distance gets shorter (interaction gets stronger), the value of θ tends to approach closer to the BD trajectory ($\sim 109^\circ$) (Fig. 6A–F). This signifies that it is important that the donor atom approaches the acceptor atom in a trajectory closer to the BD trajectory for stronger $CO \cdots CO$ interactions. Interestingly, for the motifs II and V having reciprocal $CO \cdots CO$ interactions, the angle of approach (θ) of the donor oxygen atom to the acceptor carbon atom mostly falls in the 80 – 90° range, which is much smaller than that of the BD trajectory.

2(c) Pyramidalization of the acceptor carbon atom. Previous studies have demonstrated that pyramidalization of the acceptor carbonyl carbon toward a putative electron donor is a crucial piece of experimental evidence supporting a partially covalent $n \rightarrow \pi^*$ interaction rather than a simple favorable dipole–dipole alignment or electrostatic interactions.^{4,10a,j,19} We looked into the pyramidalization (Θ) of the acceptor carbons with respect to the donor oxygen atoms in 50 molecules with $O \cdots C \leq 3.0$ Å for each motif type (I–VI). Of the six motifs (I–VI), only motif III and IV showed positive pyramidalization (Θ) (Fig. 7). In all other cases, the Θ values were spread toward both positive and negative values. The positive values of pyramidalization in motifs III and IV indicate strong $n \rightarrow \pi^*$ interaction components in these motifs. However, some spread in pyramidalization was observed in motifs III and IV at $O \cdots C$ distances greater than 3.0 Å indicating a weaker $n \rightarrow \pi^*$ interaction for $O \cdots C$ distances greater than 3.0 Å (Fig. S9†). In motif I, we expected a strong positive pyramidalization. However, in motif I the acceptor carbonyl carbon interacts with two donor oxygen atoms from opposite directions, and it is likely that the interaction from one side nullifies the effect of the interaction from the other side and hence a positive pyramidalization was not observed.

2(d) Nature of orbital interactions in motifs I–VI. To get insights into the nature of orbital interactions in motifs I–VI, we carried out NBO analyses on ten molecules from each category (five intermolecular and five intramolecular cases) (Table S4†). We observed that for a particular motif, the nature of the orbital interaction is similar irrespective of whether it is an inter- or intramolecular interaction. However, the nature of the orbital interaction strongly depends on the relative orientation of the



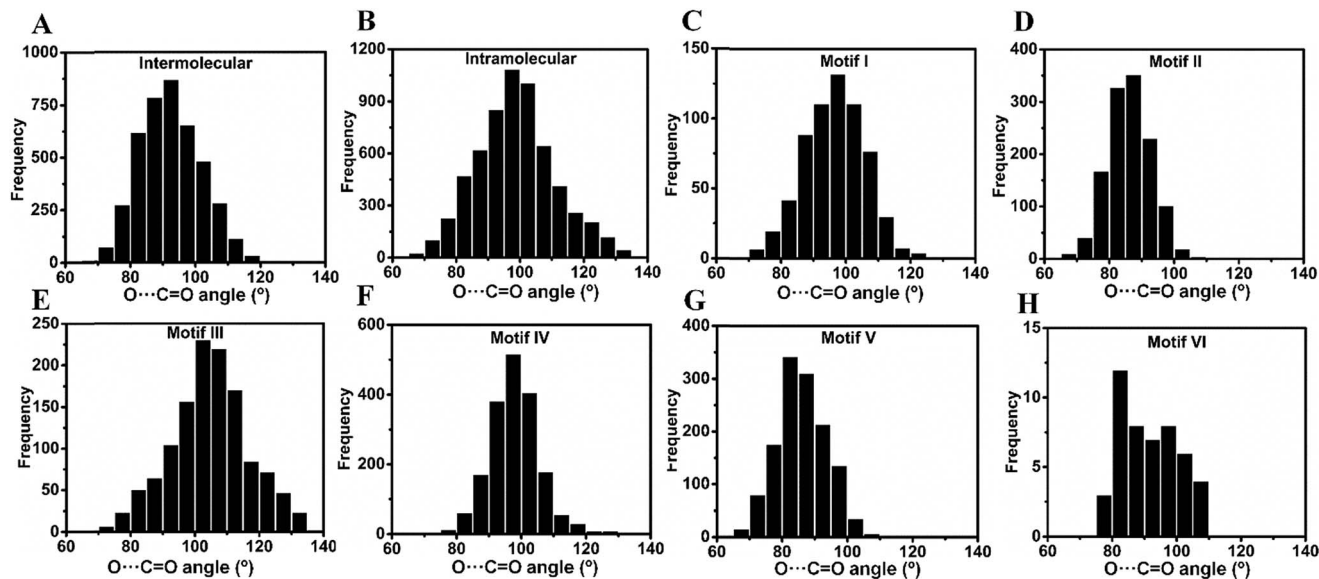


Fig. 5 Distribution of the $O\cdots C=O$ angle (θ) of all the molecules obtained from the CSD search. (A) all molecules having intermolecular $CO\cdots CO$ interactions; (B) for all molecules having intramolecular $CO\cdots CO$ interactions; (C) motif I; (D) motif II; (E) motif III; (F) motif IV; (G) motif V; (H) motif VI.

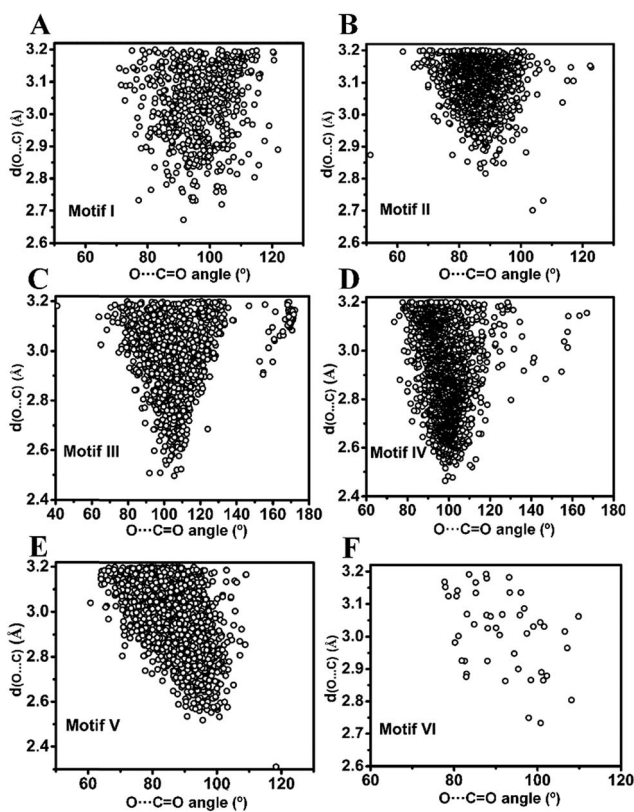


Fig. 6 $O\cdots C=O$ angle (θ) versus $O\cdots C$ distance ($d_{O\cdots C}$) scatter plot of molecules obtained from the CSD search. (A)–(F) motifs I–VI.

carbonyl groups and is different in each motif type (Fig. 8). For the perpendicular motif I, molecules randomly picked from the CSD show that the interaction is stabilized by an $n \rightarrow \pi^*$

interaction involving the sp -character lone pair of the donor carbonyl oxygen and the π^* orbital of the acceptor carbonyl group, irrespective of whether the interaction is inter- or intramolecular (Table S6[†]). The nature of orbital interactions in these molecules resembles that of the perpendicular geometry of the acetone dimer. In molecules having antiparallel geometries (motif II) that resemble the global minimum geometry of the acetone dimer, the $CO\cdots CO$ interactions are stabilized by both $n \rightarrow \pi^*$ and $\pi \rightarrow \pi^*$ interactions, the $\pi \rightarrow \pi^*$ interactions being the major contributor. In addition, as in the acetone dimer, the sp -character lone pair of the donor oxygen atom is involved in the weak $n \rightarrow \pi^*$ interaction. Therefore, it is not surprising that a positive pyramidity was not observed in motif II. In molecules having a motif III geometry, the stabilization of the $CO\cdots CO$ interaction comes from both $\pi \rightarrow \pi^*$ and $n \rightarrow \pi^*$ orbital interactions but the major contribution comes from $n \rightarrow \pi^*$ orbital interactions. In this case, both the p - and sp -character lone pairs participate in $n \rightarrow \pi^*$ orbital interactions and a positive pyramidity was observed in these molecules. In the novel L-shaped motif IV, $CO\cdots CO$ interactions are stabilized by $n \rightarrow \pi^*$ interactions involving the p -character oxygen lone pair (Table S6[†]). Interestingly, the nature of orbital interactions in these motifs (I–IV) can be mimicked by using the corresponding geometries of the acetone dimer as discussed above.

In motif V, significant reciprocal $n \rightarrow \pi^*$ interactions were observed. These interactions can be viewed as donation from the first carbonyl group to the second and back donation from the second carbonyl group to the first. Both p - and sp -character oxygen lone pairs participate in this interaction. We also observed some $\pi \rightarrow \pi^*$ interactions in motif V. This interaction is unique to intramolecular systems and, therefore, cannot be mimicked by using a dimeric model system.



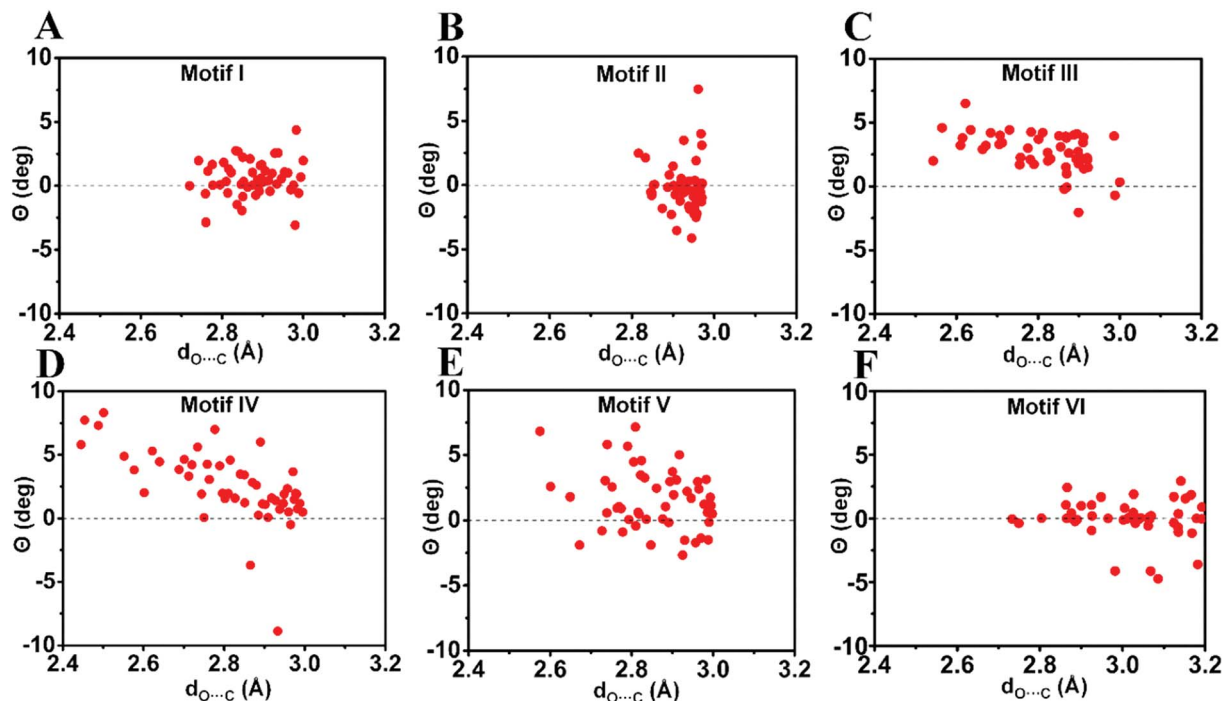


Fig. 7 Pyramidity (θ) of the acceptor carbonyl carbon in motifs I–VI. (A)–(F) Motifs (I)–(VI). Only molecules with $d_{O...C} \leq 3.0$ Å [$d_{O...C} \leq 3.2$ Å for motif VI] were considered. See ESI† for details.

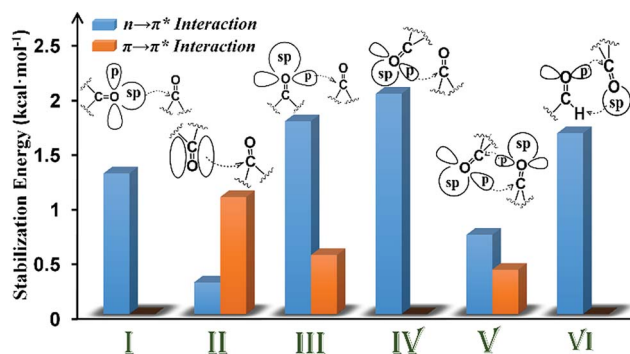


Fig. 8 Plot showing the various orbital interactions in various motifs (I–VI) having CO \cdots CO interactions and the NBO second-order perturbation energy of representative molecules of each motif.

The CO \cdots CO orbital interactions in molecules from motif VI can be explained by using the formaldehyde dimer model wherein the p-character oxygen lone pair donates electrons to the π^* orbital of the acceptor carbonyl to form an $n \rightarrow \pi^*$ interaction. The acceptor carbonyl oxygen is also involved in a hydrogen bonding interaction with the donor XH group (X = C, N, and O).

In conclusion, we have shown that the molecules with inter- and intramolecular CO \cdots CO interactions can be categorized into six different structural motifs (I–VI). Both $n \rightarrow \pi^*$ and $\pi \rightarrow \pi^*$ interactions may contribute to the stabilization of CO \cdots CO interactions. The majority of the intermolecular CO \cdots CO interactions resemble antiparallel motif II that are driven by $\pi \rightarrow \pi^*$ orbital interactions whereas intramolecular CO \cdots CO interactions

are dominated by motifs III and IV with strong $n \rightarrow \pi^*$ interactions and motif V with both $n \rightarrow \pi^*$ and $\pi \rightarrow \pi^*$ interactions. We observed that the relative geometrical arrangement of the two interacting carbonyl groups determines the nature of the orbital interaction for the stabilization of the CO \cdots CO interaction. The nature of the donor oxygen lone pair (sp- or p-character) in the $n \rightarrow \pi^*$ orbital interaction also depends on the relative orientation of the two carbonyl groups. Interestingly, CO \cdots CO interactions in motifs I–VI in both inter- and intramolecular systems can be modelled by using simple homodimeric models of acetone and formaldehyde. As intramolecular CO \cdots CO interactions in large systems are often modelled by using small dimeric systems having intermolecular CO \cdots CO interactions, our study should provide guidance in choosing the correct model system based on the similarity of the nature of CO \cdots CO interactions in the system under investigation and the system used to model it. Although theoretically all six structural motifs I–VI can exist in proteins, the presence of only two of these structural motifs III and V is known to date. As long range CO \cdots CO interactions are probable in proteins, it would be interesting to see if any of the other motifs reported herein is also observed in proteins in the future.

Author contributions

All authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts of interest to declare.



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