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Analysis of bonding motifs in unusual molecules I: planar hexacoordinated carbon atoms†

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The bonding structures of CO_3Li_3^+ and CS_3Li_3^+ are studied by means of oriented quasi-atomic orbitals (QUAOs) to assess the possibility of these molecules being planar hexacoordinated carbon (phC) systems. CH_3Li and CO_3^{2-} are employed as reference molecules. It is found that the introduction of Li^+ ions into the molecular environment of carbonate has a greater effect on the orbital structure of the O atoms than it does on the C atom. Partial charges computed from QUAO populations imply repulsion between the positively charged C and Li atoms in CO_3Li_3^+ . Upon the transition from CO_3Li_3^+ to CS_3Li_3^+ , the analysis reveals that the substitution of O atoms by S atoms inverts the polarity of the carbon–chalcogen σ bond. This is linked to the difference in s- and p-fractions of the QUAOs of C and S, as element electronegativities do not explain the observed polarity of the $\text{CS}\sigma$ bond. Partial charges indicate that the larger electron population on the C atom in CS_3Li_3^+ makes C–Li attraction possible. Upon comparison with the C–Li bond in methyllithium, it is found that the C–Li covalent interactions in CO_3Li_3^+ and CS_3Li_3^+ have about 14% and 6% of the strength of the C–Li covalent interaction in CH_3Li , respectively. Consequently, it is concluded that only CS_3Li_3^+ may be considered to be a phC system.

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1. Introduction

Exceptions to the conventional bonding structures of carbon that are commonly taught in introductory chemistry courses, traditionally characterized by so-called hybridization states, have been of great interest to chemists. The study of such exceptions can provide insight into the nature of the interactions that carbon may establish outside of the well-known bonding patterns observed in most organic molecules. An example of such an exception was proposed in 1970 by Hoffmann and collaborators,¹ when they studied a variety of possible planar tetracoordinated carbon (ptC) structures by means of Extended Hückel^{2–5} (EH), Complete Neglect of Differential Overlap^{6–10} (CNDO), and *ab initio* calculations. Their work elucidated a set of strategies to stabilize such ptC geometries. Later, in 1976, Schleyer, Pople and collaborators conducted an extensive analysis of possible ptC systems.¹¹ In their work, they reported 1,1-dilithiocyclopropane and similar three-membered ring systems that contained a C atom that displayed a preference for a ptC structure over a tetrahedral one. Multiple

computational studies on ptC systems succeeded the aforementioned works.^{12,13}

The interest in unusual carbon bonding motifs has extended beyond tetracoordinate species to planar penta¹⁴ and hexacoordinated^{15–25} carbon (ppC and phC, respectively) systems. In 2012, Wu and collaborators proposed D_{3h} CO_3Li_3^+ as a viable phC system.²³ The D_{3h} geometry was identified as the global minimum on the potential energy surface. Nonetheless, the main argument to consider this molecule as a phC system, and not just a planar carbonate dianion stabilized by three Li^+ ions, was the relatively short C–Li distance (2.212 Å at the coupled cluster CCSD(T)/aug-cc-pVTZ level of theory), despite computing significantly low Wiberg Bond Indices²⁶ (WBI) between these atoms. More recently, in 2021, Leyva-Parra and collaborators indicated that the criterion employed by Wu and collaborators to determine hexacoordination was purely geometrical and reported strongly repulsive interactions between the carbon and lithium atoms due to their partial positive charges in the molecule.¹⁵ In their work, Leyva-Parra and coauthors suggested a series of alternative molecules in which the oxygen atoms were replaced by heavier, less electronegative chalcogens. It was observed that the central C atom in such systems was covalently bonded to the chalcogens and electrostatically attracted to the alkali metal atoms.

The aim of the present study is to analyze the unusual bonding structures of unique chemical bonds in potentially hexacoordinated carbon species. In order to analyze the

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bonding structures of these molecules, the quasi-atomic orbital (QUAO) analysis, developed by Ruedenberg and collaborators,^{27–38} is employed. The bonding interactions in CH₃Li are analyzed as well and used as reference for C–Li interactions.

An important aspect of the QUAO analysis is that it is based on the actual molecular wave function, and thus avoids the introduction of inherently biased information based on pre-conceived notions. Moreover, given the nature of the analysis, QUAOs can be conceptualized as the *ab initio* counterparts of the early concept of hybrid atomic bond orbitals.²⁹ QUAOs provide a wave function-inherent perspective on the bonding that is consistent with the chemical intuition of the reader.

The details of the QUAO analysis are extensively discussed in ref. 28–31 and thus only the most relevant aspects of the theory are outlined in Section 2 of this work. In Section 3, the computational details of the reported calculations are given. In Section 4, the computed structures and the results of the QUAO analysis are reported and discussed. Finally, the main conclusions of this work are summarized in Section 5.

2. Quasi atomic orbital analysis

In this section an overview of the most relevant aspects and basic concepts of the QUAO analysis is provided. As mentioned in the Introduction, the quasi-atomic orbital analysis, available in the GAMESS electronic structure software,^{39–42} has been extensively discussed in ref. 28–31. The QUAO analysis reported in this work was done employing RHF wave functions. In general, QUAOs can be conceptualized as minimal basis set orbitals that have been deformed relative to the isolated atom to exhibit the bonding interactions in a molecule. QUAOs can be conceived as the *ab initio* analogs of the early qualitative concept of “hybrid atomic bond orbitals”.²⁹

2.1 Conceptual framework

The QUAO analysis is based on the conceptual partitioning of the full molecular orbital (MO) space (for HF wave functions, the HF orbital space) into three distinct subspaces: the chemical

core, the valence, and the external space, as shown in Fig. 1. The chemical core and valence spaces are the union of the chemical core and valence spaces of the individual atoms in the molecule, respectively (e.g., in minimal-basis C₂, the chemical core space contains the 1s orbitals on each C atom, while the valence space contains the 2s and 2p orbitals on each C atom). The external space is the orthogonal complement to the core and valence spaces. For the study of bonding in molecules, the valence space is the primary interest.²⁹

Typically, the Hartree–Fock (HF) self-consistent field (SCF) procedure only yields the bonding and non-bonding combinations of the valence atomic orbitals (AOs). The antibonding combinations are often missing since the virtual (*i.e.*, unoccupied) orbitals are generated as just an orthogonal complement to the occupied set. Thus, in general, the occupied HF space does not span the full valence space. Nonetheless, Ruedenberg and collaborators have shown that the antibonding combinations of the AOs may be extracted from the HF virtual space *via* a projection onto a set of orthogonal minimal basis set atomic orbitals.^{43–45} This extraction employs a set of accurate atomic minimal basis set (AAMBS) orbitals, discussed in Section 2.2, and the resulting (antibonding) molecular orbitals are called the valence virtual orbitals (VVOs).⁴⁶ The VVOs constitute an *ab initio* analog of the lowest unoccupied molecular orbitals (LUMOs)^{47–49} and complement the HF filled orbitals to fully specify the valence space in HF SCF calculations, as shown in Fig. 1.

2.2 Accurate atomic minimal basis set

Minimal basis set (MBS) orbitals, *i.e.*, the orbitals obtained from atomic closed- or open-shell HF SCF or multi-configuration self-consistent-field (MCSCF) calculations, are the fundamental basis for the description of the physical nature of the electronic structure of atoms. Concomitantly, MBS orbitals for atoms are the core of a qualitative understanding of chemistry, essentially accounting for the structure of the periodic table.⁵⁰ Ruedenberg, Gordon and collaborators have shown in previous works^{27–38,43–46,50–54} the utility of MBS orbitals for the extraction of qualitative and quantitative understanding of the results of molecular electronic

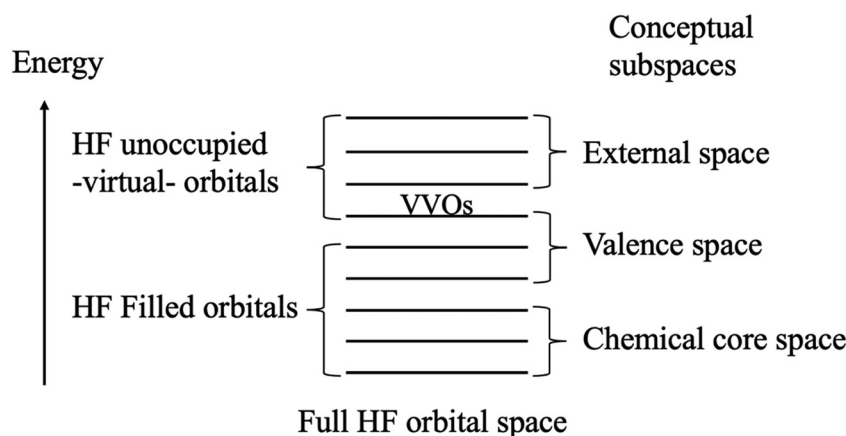


Fig. 1 Conceptual partitioning of the full Hartree–Fock (HF) orbital space. The lowest virtual orbitals are replaced by the valence virtual orbitals (VVOs) to fully specify the valence space.



structure calculations. This is rooted in the notion that atom-like entities are preserved in molecular electronic wave functions, albeit distorted relative to the free atom wave functions. This notion, although intuitive to chemists, is not usually fundamental to physical theory, in which the conception of molecules is primarily based on the many-electron many-nuclei model.^{50,55}

To obtain accurate results from the extractions mentioned above, an accurate representation of the atomic MBS orbitals is important. To this end, a set of AAMBS orbitals has been developed for most atoms in the periodic table.^{46,56,57} These AAMBS orbitals are embedded in the GAMESS electronic structure program.^{39–42} The AAMBS orbitals have been chosen to be the HF SCF ground state neutral atom orbitals for the s- and p-blocks. For transition metals, for which multiple low-lying configurations may play a role in molecular interactions, the MBS orbitals of the neutral atoms are obtained from state-averaged (SA) MCSCF calculations involving the two lowest energy atomic configurations.⁴⁶ In all cases, the core orbitals are included. The AAMBS orbitals are expanded using a large number of primitive Gaussian functions, effectively at the atomic basis set limit, thus providing an effectively exact representation of the (HF or SA-MCSCF) atomic MBS orbitals. For the s- and p-block elements up to Ne, even-tempered Gaussians are used.⁵⁸ After Ne and up to Xe, well-tempered Gaussians^{59,60} are employed. A detailed discussion of the AAMBS orbitals can be found in ref. 46 and 57.

2.3 QUAO extraction and orientation

The projection of the AAMBS orbitals onto the valence space is obtained *via* a singular value decomposition (SVD),^{29,61,62} of the corresponding orbital overlap matrix. This projection produces the molecular orbitals that are most optimally aligned with the free atom AAMBS orbitals, thus capturing the physical nature of each atom within the molecule. For each atom, the number of projected orbitals is equal to the number of minimal basis valence orbitals of that atom. Once the projection of each atom has been computed independently, the complete set of projected orbitals is orthogonalized. The resulting orthogonal QUAOs represent the set of molecular orbitals that are closest to the free atom AOs.

To clearly elucidate the bonding patterns in molecules, the orthogonal QUAOs are adapted to the molecular environment by allowing the QUAOs on each atom to mix with each other, thereby resulting in hybridized orbitals.^{50,52} In the corresponding one-electron density matrix, the diagonal elements represent orbital populations, while the off-diagonal elements represent bond orders, which are indicative of the covalent bonding interactions in the molecule. To orient the orbitals, the linear combinations of the QUAOs on each atom are taken to be those for which the off-diagonal blocks of the one-electron density matrix have as few elements with large magnitudes as possible.⁵² The resulting orbitals are called oriented QUAOs, as they capture the covalent interactions in the molecule. The s- and p-characters of the oriented QUAOs are obtained by directly computing their s- and p-fractions. The terms oriented QUAO and QUAO are henceforth used interchangeably.

It is worth noting that, while chemical intuition may allow one to anticipate the bonding patterns in a molecule, the orientation procedure (discussed in detail in ref. 50) makes no prior assumptions about the bonding in any given system, thus providing an unbiased, molecule-inherent picture of bonding. Such an unbiased approach may be of particular relevance in molecules in instances for which intuition/prior assumptions are not good guides.

2.4 Bond orders and kinetic bond orders

For diatomic molecules positive bond orders (BOs) can be associated with bonding interactions, and negative bond orders with antibonding interactions. However, in polyatomic molecules, the signs of the bond orders depend on the phases of the QUAOs. In addition, BOs are not energy quantities. Therefore, Ruedenberg and collaborators introduced kinetic bond orders (KBOs),^{28,30} which include the kinetic contribution to the interference energy as an approximate energetic measure of the strength of a covalent bonding interaction. This is rooted in the (now widely accepted)^{63–67} identification of the interference kinetic energy as the fundamental origin of the covalent bond.^{29–32,68–71} KBOs are then defined by:

$$k_{Aa,Bb} = 0.1 \times p_{Aa,Bb} \left\langle Aa \left| -\frac{1}{2} \nabla^2 \right| Bb \right\rangle \quad (1)$$

In eqn (1) $|Aa\rangle$ represents QUAO a on atom A, and $p_{Aa,Bb}$ represents the bond order between QUAO b on atom B and QUAO a on atom A. The scale factor of 0.1 was introduced to compensate for the omission of a potential energy contribution, which is typically antibonding.^{55,71} Moreover, KBOs have been found to be negative in every situation in which covalent bonding occurs, and have been previously employed to elucidate and compare bonding interactions in various molecules and molecular clusters.^{27,28,33–36}

2.5 Basis set independence

Previous studies by Ruedenberg and collaborators have shown that MOs extracted from electronic wave functions employing atomic minimal basis set (AAMBS) orbitals are consistently basis set independent; *i.e.*, the resulting projections are independent of the working basis used to expand the MOs.^{43,44} Moreover, it has been observed that, even when large AO bases are employed, the one-electron density matrix is dominated by the terms that involve the minimal basis set orbitals on all atoms.⁴³ Thus, despite the well-known need for extended AO basis sets for the computation of accurate wave functions (and the associated energetics), the AAMBS basis representation has been observed to contain the essential information about bonding in molecules.

Given the nature of the AAMBS employed for the extraction of QUAOs, (*i.e.*, the AAMBS AOs represent the exact, basis set-independent, optimal atomic minimal basis set orbitals),⁴⁶ the overlaps with the working basis remain consistent across different types of extended basis sets, as long as the working basis provides an appropriate description of the electronic



structure of the molecule at hand. Such basis set independence has also been observed in the extraction of the valence virtual orbitals^{29,43} (VVOs) implemented by Schmidt, Hull and Windus,⁴⁶ which employs the AAMBS orbitals discussed in Section 2.2 as well. An in-depth analysis of the basis set independence of the VVOs obtained employing the AAMBS can be found in ref. 46.

3. Computational details

Previous works have shown that the 2p orbitals in Li may play an important role in the bonding interactions of Li,^{64,72–74} and that they can be regarded as valence orbitals. Therefore, the version of the AAMBS that includes the 2p orbitals on Li was employed for the extraction of the VVOs and QUAOs in this work.⁵⁷

The geometries of all molecules discussed in this work were optimized at the RHF/6-31G(d), MP2/aug-cc-pVDZ and CCSD(T)/aug-cc-pVTZ levels of theory. For conciseness, the aug-cc-pVDZ and aug-cc-pVTZ basis sets will be referred to as ACCD and ACCT, respectively, throughout this work. The geometry of CH₃Li was optimized enforcing *C*_{3v} symmetry, while the geometries of CO₃Li₃⁺ and CS₃Li₃⁺ were optimized enforcing *D*_{3h} symmetry. The corresponding RHF and MP2 Hessians were computed to establish the stationary point nature of every geometry. Localized quasi-atomic orbitals were obtained using RHF/6-31G(d) wave functions at the RHF/6-31G(d) and CCSD(T) optimized geometries. The QUAO results are essentially the same at both the RHF and CCSD(T) geometries, thus only the RHF/6-31G(d) results are reported. To compare the results obtained with RHF to those obtained with a higher level of theory, the QUAOs of CH₃Li were computed using a full-valence (8,11)-CASSCF/6-311++G(d,p)//RHF/6-311++G(d,p) wave function. It was found that the results did not change significantly when the CASSCF wave function was employed. The comparison of the results obtained at both the RHF and CASSCF levels of theory can be found in Section S4 of the ESI.[†]

All calculations were done using the GAMESS software^{39–42} and QUAOs were plotted using the MacMolPlt visualization software⁷⁵ with contours of 0.1 bohr^{−3/2} for all QUAOs, except

for those centered on Li. For the QUAOs of Li, contours of 0.07 bohr^{−3/2} were used given their large spatial extent.

4. Hexacoordinated carbon species

4.1 Geometries

The geometries of CH₃Li (*C*_{3v}), CO₃Li₃⁺ (*D*_{3h}) and CS₃Li₃⁺ (*D*_{3h}) optimized at the RHF/6-31G(d) level of theory are shown in Fig. 2. The geometric parameters of all three molecules at the MP2/ACCD and CCSD(T)/ACCT levels of theory can be found Section S1 of the ESI.[†] A comparison between the RHF, MP2 and CCSD(T) geometries as well as with previously reported results can be found in Section S1 of the ESI.[†] as well.

4.2 Quasi-atomic bonding analysis

The QUAOs presented in this section are labeled according to their role in their respective molecules. The labeling works as follows: the atomic symbol of the atom on which the orbital is centered is listed first, with the first letter capitalized. If the orbital participates in a (covalent) bonding interaction, the symbol of the complementary atom is listed second in lower case. The third component of the label indicates the kind of bonding in which the orbital participates; e.g., σ or π. Orbitals with occupations close to 2, have the labels sl and pl to denote whether they are primarily an s-type or a p-type lone pair, respectively. For those orbitals with occupations below 0.2, that are not involved in bonding interactions, the label nv is employed to denote that the orbital is nearly vacant. For example, an orbital centered on carbon that participates in a C–O π bonding interaction would be labeled Cοπ; an orbital centered on O with a population close to 2 and predominantly p-character would be labeled Opl.

The QUAOs obtained for CH₃Li are displayed in Fig. 3, those obtained for CO₃Li₃⁺ are shown in Fig. 4, and those corresponding to CS₃Li₃⁺ are depicted in Fig. 5. For CH₃Li, the most relevant bonding interactions are summarized in Table 1; the orbital occupations, as well as s- and p-characters, are presented in Table 2. Partial charges on each atom in every molecule in the present study, computed from QUAO populations, are listed in

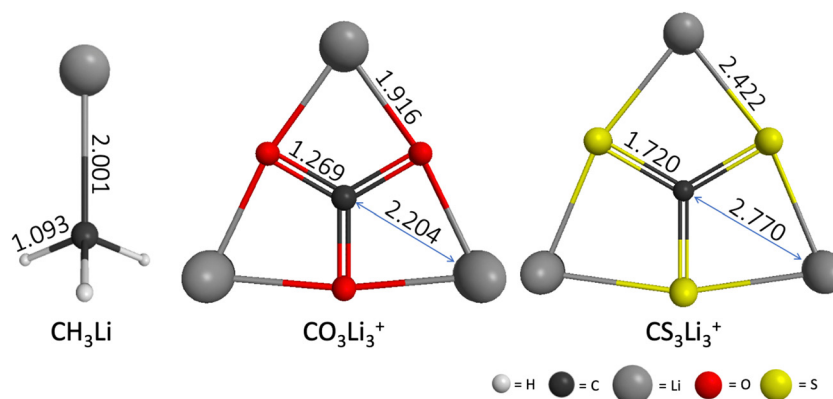


Fig. 2 CH₃Li (*C*_{3v}), CO₃Li₃⁺ (*D*_{3h}) and CS₃Li₃⁺ (*D*_{3h}) equilibrium structures computed at the RHF/6-31G(d) level of theory. Interatomic distances shown in Angstroms.

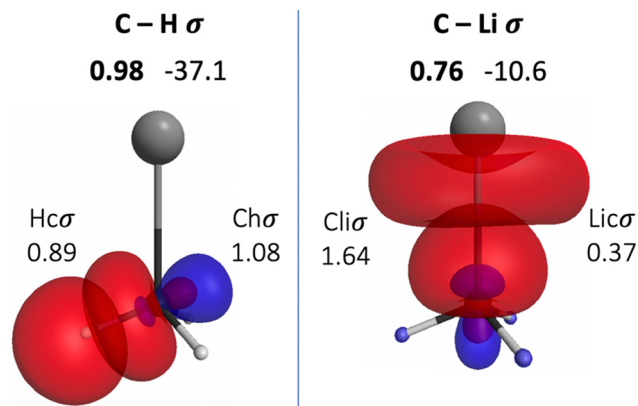


Fig. 3 Interactions between the QUAOs in CH_3Li . Bond orders are shown in bold above the displayed orbitals. Kinetic bond orders (in kcal mol^{-1}) are shown to the right of the corresponding bond order. The labels of the orbitals involved in the interactions are shown to the sides of the orbitals, and orbital populations are shown below the orbital label.

Table 3. For CO_3Li_3^+ , the most relevant bonding interactions are summarized in Table 4; the orbital occupations, as well as their s- and p-characters, are given in Table 5. The results for CS_3Li_3^+ are summarized in Tables 6 and 7, which are analogous to Tables 4 and 5. In general, only interactions with bond orders larger than 0.15 and kinetic bond orders lower than $-1.0 \text{ kcal mol}^{-1}$ are reported, with the exception of very weak interactions

that may aid in the determination of the coordination number of a C atom.

4.2.1 CH_3Li . In CH_3Li , the central carbon atom is bonded to the three H atoms *via* three symmetrically equivalent C σ orbitals, and to the Li atom *via* a Cl σ orbital. Each H atom is bonded to the C atom *via* a Hc σ orbital, while the Li atom is bonded to the C atom *via* a Li σ orbital. The orbital populations show a small charge transfer from H to C; the transfer of charge is expected given the difference in electronegativity between H and C. The s- and p-characters of the C σ orbital are 0.19 and 0.81, respectively, roughly sp^3 hybridization. The Hc σ orbital has an entirely s-type character.

The BO of the Cl σ interaction is 0.76 and the KBO is $-10.6 \text{ kcal mol}^{-1}$. The occupation numbers of the Cl σ (1.64) and Li σ (0.37) orbitals imply that a transfer of charge, much larger than the one occurring from H to C, occurs from Li to C. This transfer is also consistent with the difference in the electronegativities of Li and C. Moreover, the transfer of charge from H and Li to C lead to the appearance of relatively large opposite partial charges on the Li (+0.56) and C (-0.89) atoms. The presence of these charges is consistent with previous assertions regarding the C–Li interaction in CH_3Li as “80–90% ionic”.^{76,77} Nonetheless, previous studies have also shown that covalent character plays a non-negligible role in the C–Li bond.^{72,76,78} The covalent nature of the Cl σ interaction is reflected in the corresponding KBO, which suggests a

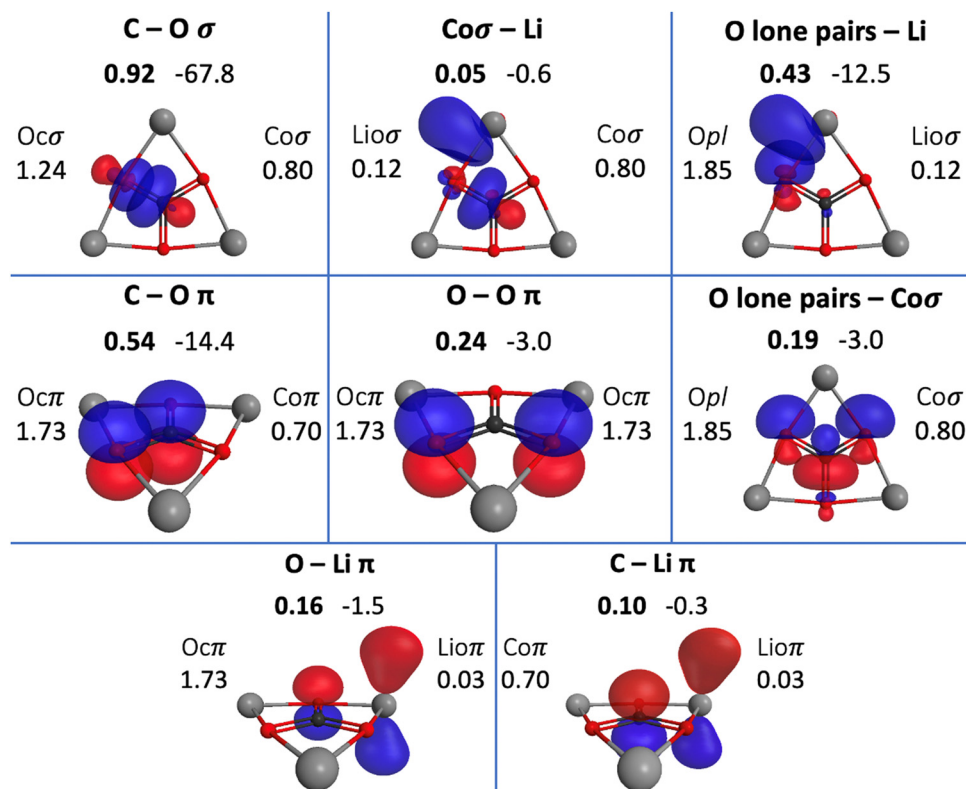
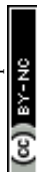


Fig. 4 Interactions between the QUAOs in CO_3Li_3^+ . Bond orders are shown in bold above the displayed orbitals. Kinetic bond orders (in kcal mol^{-1}) are shown to the right of the corresponding bond order. The labels of the orbitals involved in the interactions are shown to the sides of the displayed orbitals, and orbital populations are shown below the corresponding orbital label.



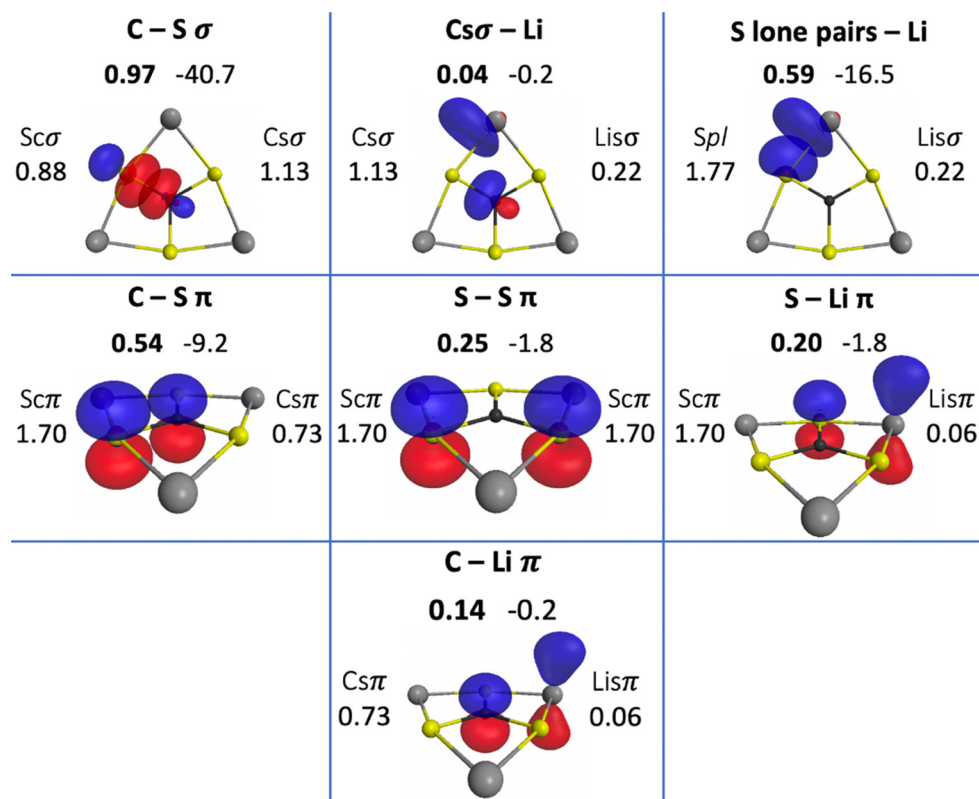


Fig. 5 Interactions between the QUAOs in CO_3Li_3^+ . Bond orders are shown in bold above the displayed orbitals. Kinetic bond orders (in kcal mol^{-1}) are shown to the right of the corresponding bond order. The labels of the orbitals involved in the interactions are shown to the sides of the orbitals, and orbital populations are shown below the corresponding orbital label.

Table 1 Bonding interactions and characteristics in CH_3Li . Only interactions with $\text{BO} > 0.15$ and $\text{KBO} < -1 \text{ kcal mol}^{-1}$ are listed

Bond	Orbital I	Orbital J	BO	KBO (kcal mol^{-1})
$\text{CH}\sigma$	$\text{Ch}\sigma$	$\text{Hc}\sigma$	0.98	−37.1
$\text{CLi}\sigma$	$\text{Cli}\sigma$	$\text{Lic}\sigma$	0.76	−10.6

Table 2 Orbital occupations and s- and p-character fractions in CH_3Li

Orbital	Occupation	Fraction s	Fraction p
$\text{Ch}\sigma$	1.08	0.19	0.81
$\text{Hc}\sigma$	0.89	1	0
$\text{Cli}\sigma$	1.64	0.26	0.74
$\text{Lic}\sigma$	0.37	0.81	0.19
Linv	0.02	0.09	0.91

non-trivial degree of covalent character in the interaction. The s- and p-characters of the $\text{Cli}\sigma$ orbital are 0.26 and 0.74, respectively. The $\text{Lic}\sigma$ orbital has s- and p-characters of 0.81 and 0.19, respectively, reflecting a small but significant degree of s and p hybridization in Li, consistent with previous observations.^{64,72–74} The inclusion of the 2p orbitals in the Li AAMBS leads to three Linv orbitals, which are not involved in any bonding interactions in the molecule, and are essentially vacant (Fig. S1, ESI[†]). Thus, the bonding of Li in CH_3Li is

Table 3 Atomic partial charges computed from QUAO populations in CH_3Li , CO_3^{2-} (obtained from ref. 28), CO_3Li_3^+ and CS_3Li_3^+

Molecule	CH_3Li			CO_3^{2-}		
Atom	C	H	Li	C	O	
Charge	−0.89	+0.11	+0.56	+0.84	−0.95	

Molecule	CO_3Li_3^+			CS_3Li_3^+		
Atom	C	O	Li	C	S	Li
Charge	+0.90	−0.68	+0.72	−0.12	−0.12	+0.49

dominated by the 2s orbital; the mixing with the 2p functions allows the polarization of the resulting $\text{Lic}\sigma$ orbital.

4.2.2 CO_3Li_3^+ . There are three symmetrically equivalent $\text{CO}\sigma$ bonds in CO_3Li_3^+ (Fig. 4). Each such interaction has a KBO of $-67.8 \text{ kcal mol}^{-1}$ and a BO of 0.92. Each O atom is involved in one $\text{CO}\sigma$ bond, via one $\text{Oc}\sigma$ orbital. The occupation numbers for the $\text{Co}\sigma$ and $\text{Oc}\sigma$ orbitals are 0.80 and 1.24, respectively, indicating that the $\text{CO}\sigma$ bond is polarized toward the O atom, as expected given the relative electronegativities of C and O. The $\text{Co}\sigma$ orbital has an s-character of 0.29 and a p-character of 0.71, roughly sp^2 hybridization. The $\text{Oc}\sigma$ orbital has an s-character of 0.28 and a p-character of 0.72.

There are three symmetrically equivalent $\text{CO}\pi$ bonds, each with a KBO of $-14.4 \text{ kcal mol}^{-1}$ and a BO of 0.54, in which the



Table 4 Bonding structures and characteristics in CO_3^{2-} obtained from ref. 28 and CO_3Li_3^+ . Only interactions with $\text{BO} > 0.15$ and $\text{KBO} < -1 \text{ kcal mol}^{-1}$ are listed. The Li–C interactions are listed despite being below the set threshold

Bond	Orbital I	Orbital J	CO_3^{2-28}		CO_3Li_3^+	
			BO	KBO	BO	KBO
CO σ	Co σ	Oc σ	0.93	−66.0	0.92	−67.8
CO π	Co π	Oc π	0.56	−16.1	0.54	−14.4
OO π	Oc π	Oc π	0.24	−3.4	0.24	−3.0
Opl–Co σ	Opl	Co σ	0.21	−5.4	0.19	−3.0
Opl–Lio σ	Opl	Lio σ			0.43	−12.5
Oc π –Lio π	Oc π	Lio π			0.16	−1.5
Lio σ –Co σ	Lio σ	Co σ			0.05	−0.6
Lio π –Co π	Lio σ	Co π			0.10	−0.3

C atom employs a single Co π orbital to interact with the three symmetrically equivalent Oc π orbitals in the molecule. The occupation numbers of the Co π and Oc π orbitals are 0.70 and 1.73, respectively, reflecting the polarization of the π -system toward the O atoms. The C–O interactions in the π -system are complemented by bonding interactions between the symmetrically equivalent Oc π orbitals. This interaction has a BO of 0.24 and a KBO of $-3.0 \text{ kcal mol}^{-1}$.

There are several weak interactions in CO_3Li_3^+ that are worth mentioning. There are, for example, two symmetrically equivalent Opl orbitals on each O atom, each with a population of 1.85. The Opl orbitals have s- and p-characters of 0.33 and 0.67, respectively. There are hyperconjugative interactions between the Opl and Co σ orbitals; as shown in Fig. 4, a single Co σ orbital is hyperconjugated with two different Opl orbitals, each located on the two O atoms that the Co σ orbital is not oriented toward. For each Opl–Co σ orbital pair, the hyperconjugative interaction has a BO of 0.19 and KBO of $-3.0 \text{ kcal mol}^{-1}$.

Each lithium atom has two symmetrically equivalent Lio σ orbitals that interact with the lone pairs of the two closest O atoms. Each Opl–Lio σ interaction has a BO of 0.43 and a KBO of $-12.5 \text{ kcal mol}^{-1}$. The Lio σ orbitals have s and p characters of 0.28 and 0.72, indicating that the mixing of the Li 2s and 2p functions is larger in CO_3Li_3^+ than in CH_3Li . Interactions among ligands have been previously suggested to contribute to the stability of planar hypercoordinated species.^{15,23} Covalent bonding between ligands contributes to the kinetic energy lowering that drives molecule formation; the KBOs indicate that the Opl–Lio σ interactions make non-negligible contributions to that

Table 6 Bonding structures and characteristics in CS_3Li_3^+ . Only interactions with $\text{BO} > 0.15$ and $\text{KBO} < -1 \text{ kcal mol}^{-1}$ are listed. The Li–C interactions are listed despite being below the set threshold

Bond	Orbital I	Orbital J	BO	KBO (kcal mol^{-1})
CS σ	Cs σ	Sc σ	0.97	−40.7
CS π	Cs π	Sc π	0.54	−9.2
SS π	Sc π	Sc π	0.25	−1.8
Spl–Lis σ	Spl	Lis σ	0.59	−16.5
Sc π –Lis π	Sc π	Lis π	0.20	−1.8
Lis σ –Cs σ	Lis σ	Cs σ	0.04	−0.2
Lis π –Cs π	Lis π	Cs π	0.14	−0.2

Table 7 Orbital occupations and s- and p-character fractions in CS_3Li_3^+

Orbital	Occupation	Fraction s	Fraction p
Cs σ	1.13	0.30	0.70
Cs π	0.73	0.00	1.00
Sc σ	0.88	0.11	0.89
Sc π	1.70	0.00	1.00
Spl	1.77	0.41	0.59
Lis σ	0.22	0.31	0.69
Lis π	0.06	0.00	1.00
Linv	0.02	0.35	0.65

energy lowering.^{71,79} The Opl–Lio σ interactions are complemented by weak π interactions between O and Li, with a BO and KBO of 0.16 and $-1.5 \text{ kcal mol}^{-1}$, respectively. Thus, the six Opl–Lio σ and Oc π –Lio π interactions have a total KBO of $-84 \text{ kcal mol}^{-1}$, indicating that they make a net contribution to the stabilization of the molecule even larger than that of a single CO σ bond.

Wu and collaborators have suggested that CO_3Li_3^+ may be conceptualized as simply a CO_3^{2-} moiety stabilized by three Li^+ ions.²³ As shown in Table 4, the BOs and KBOs for the C–O and O–O interactions are similar to the values reported previously by West and collaborators for the carbonate ion.²⁸ The similarity of the KBOs of the CO σ and CO π interactions suggests that, as proposed by Wu *et al.*, the bonds between C and O are not significantly affected by the presence of the three lithium ions in CO_3Li_3^+ with respect to CO_3^{2-} . As shown in Table 5, the main differences between CO_3^{2-} and CO_3Li_3^+ are observed in the lone pairs of the O atoms, where the introduction of the Li^+ ions is observed to increase the mixing of s and p functions.

The C–Li interaction in CO_3Li_3^+ occurs between each Co σ orbital and the two closest Lio σ orbitals (each Lio σ on a

Table 5 Orbital occupations and s- and p-character fractions in CO_3^{2-} obtained from ref. 28 and CO_3Li_3^+

Orbital	CO_3^{2-28}			CO_3Li_3^+		
	Occupation	Fraction s	Fraction p	Occupation	Fraction s	Fraction p
Co σ	0.81	0.32	0.68	0.80	0.29	0.71
Co π	0.73	0.00	1.00	0.70	0.00	1.00
Oc σ	1.27	0.38	0.62	1.24	0.28	0.72
Oc π	1.76	0.00	1.00	1.73	0.00	1.00
Opl	1.92	0.00	1.00	1.85	0.33	0.67
Osl	2.00	0.60	0.40			
Lio σ				0.12	0.28	0.72
Lio π				0.03	0.00	1.00
Linv				0.01	0.36	0.64



different atom), and between the $\text{Li}\sigma$ and $\text{Co}\pi$ orbitals. The BO and KBO for the $\text{Li}\sigma$ – $\text{Co}\sigma$ interaction are 0.05 and $-0.6 \text{ kcal mol}^{-1}$, respectively, and the BO and KBO for the $\text{Li}\sigma$ – $\text{Co}\pi$ interaction are 0.10 and $-0.3 \text{ kcal mol}^{-1}$. The sum of these interactions gives a total C–Li KBO of $-1.5 \text{ kcal mol}^{-1}$. This KBO reflects a very weak C–Li covalent interaction. This total KBO of $-1.5 \text{ kcal mol}^{-1}$ suggests that the C–Li covalent interaction in CO_3Li_3^+ has about 14% of the strength of the same interaction in CH_3Li . So, no significant C–Li covalent bonds exist in CO_3Li_3^+ .

Table 3 shows partial positive charges on the C and Li atoms, and partial negative charges on the O atoms. These repelling positive charges on C and Li imply that C–Li ionic attraction is not likely, in agreement with what has been previously observed by Leyva-Parra and collaborators.¹⁵ The most significant interactions of each of the three Li atoms are established with the two closest O atoms, *via* a combination of covalent interactions and ionic attraction.

4.2.3 CS_3Li_3^+ . There are three $\text{CS}\sigma$ bonds in CS_3Li_3^+ ; each of these bonds has a KBO of $-40.7 \text{ kcal mol}^{-1}$ and a BO of 0.97. The central C atom uses three symmetrically equivalent $\text{C}\sigma$ orbitals for such interactions, while each sulfur atom is involved in only one $\text{CS}\sigma$ bond and uses a single $\text{S}\sigma$ orbital for this interaction. The populations of the $\text{C}\sigma$ and $\text{S}\sigma$ orbitals are 1.13 and 0.88, respectively, showing that the $\text{CS}\sigma$ bond is slightly polarized toward the C atom. Depending on the choice of electronegativity values⁸⁰ this bond is not expected to be polarized, or to be only slightly polarized toward the S atom.

A possible justification for the slight polarization toward C is based on the orbital s- and p-fractions in C vs. S: the $\text{C}\sigma$ orbital has an s-character of 0.30 and a p-character of 0.70, while the partner $\text{S}\sigma$ orbital has an s-character of 0.11 and a p-character of 0.89. The larger s-character of the $\text{C}\sigma$ orbital with respect to the partner $\text{S}\sigma$ orbital is not surprising as the 2s and 2p orbitals have a well-known tendency to mix, especially in C. The evident lack of s- and p-orbital mixing in the $\text{S}\sigma$ orbital can be attributed to the different spatial extents of the 3s and 3p atomic orbitals, as was noted by Kutzelnigg.^{81,82} Based on previous QUAO analyses of $\text{CH}\sigma$, $\text{CO}\sigma$ and $\text{CP}\sigma$ bonds,^{28,38} higher s-orbital characters are accompanied by higher orbital populations. For $\text{CS}\sigma$ bonds, given the similarity in the electronegativities of C and S, the difference in the s- and p-characters of the $\text{C}\sigma$ and $\text{S}\sigma$ orbitals might determine the polarity of the $\text{CS}\sigma$ bond. This can also be understood in terms of the early concept of orbital electronegativity,^{83,84} which has been observed to increase as the s-character of an orbital increases.^{85–87} In the present and previous QUAO analyses,^{28,38} this effect has been observed to be reflected in orbital populations.

The three $\text{CS}\pi$ bonds in CS_3Li_3^+ each have a BO of 0.54 and a KBO of $-9.2 \text{ kcal mol}^{-1}$. The occupation numbers of the $\text{C}\pi$ and $\text{S}\pi$ orbitals are 0.73 and 1.70, respectively. All of the QUAOs involved in π bonding have p-characters of 1.00. So, the orbital structure of the C atom resembles sp^2 hybridization. The $\text{S}\pi$ orbitals are also observed to interact with each other, with a BO of 0.25 and a KBO of $-1.8 \text{ kcal mol}^{-1}$.

There are two Spl orbitals on each S atom. Each of these orbitals has an occupation of 1.77 and s- and p-characters of

0.41 and 0.59, respectively. Thus, they represent lone pairs on the S atoms with predominantly p-character and have been labeled as such in the figures and tables.

Each lithium atom employs two symmetrically equivalent $\text{Lis}\sigma$ orbitals to interact with the lone pairs of the two closest S atoms. Each Spl – $\text{Lis}\sigma$ interaction has a BO of 0.59 and a KBO of $-16.5 \text{ kcal mol}^{-1}$. The $\text{Lis}\sigma$ orbitals have s- and p-characters of 0.31 and 0.69. A complementary interaction occurs between the $\text{S}\pi$ and $\text{Lis}\pi$ orbitals, with a BO and KBO of 0.20 and $-1.8 \text{ kcal mol}^{-1}$, respectively. The six Spl – $\text{Lis}\sigma$ and $\text{S}\pi$ – $\text{Lis}\pi$ interactions have a total KBO of $-109.8 \text{ kcal mol}^{-1}$, making the stabilization effect of the covalent interactions between the S and Li atoms comparable to that of the three $\text{CS}\sigma$ bonds ($-122.1 \text{ kcal mol}^{-1}$).

The C–Li interaction in CS_3Li_3^+ occurs between each $\text{C}\sigma$ orbital and the two closest $\text{Lis}\sigma$ orbitals (on two different Li atoms), and between the $\text{C}\pi$ and $\text{Lis}\pi$ orbitals. The $\text{C}\sigma$ – $\text{Lis}\sigma$ interaction has a BO and KBO of 0.04 and $-0.2 \text{ kcal mol}^{-1}$, and the $\text{C}\pi$ – $\text{Lis}\pi$ interaction has a BO and KBO of 0.14 and $-0.2 \text{ kcal mol}^{-1}$. Since there are two $\text{C}\sigma$ – $\text{Lis}\sigma$ and one $\text{C}\pi$ – $\text{Lis}\pi$ interactions between the C atom and each Li atom, there is a total KBO of $-0.6 \text{ kcal mol}^{-1}$ for each C–Li interaction. This total KBO suggests that the covalent C–Li interaction in CS_3Li_3^+ has about 6% of the strength of the same interaction in CH_3Li . Consequently, it is concluded that no significant C–Li covalent bond exists in CS_3Li_3^+ .

The computed charges, displayed in Table 3, show that there are small negative charges on the S and C atoms, and a positive charge on the Li atoms. The opposite charges on the C and Li atoms suggest the possibility of ionic attraction between them, consistent with what has been suggested by Leyva-Parra and collaborators.¹⁵ The relative charges also imply an ionic attraction between the S and Li atoms.

The occupation numbers of the orbitals involved in the $\text{CS}\sigma$ and $\text{CO}\sigma$ interactions show that the substitution of O by S has the effect of inverting the polarity of the carbon–chalcogen sigma bond. The effect of this polarity inversion of the carbon–chalcogen sigma bond is an increase in the negative charge on the C atom. Each $\text{CS}\sigma$ bond increases the electron population on the C atom by about 0.33, relative to CO_3Li_3^+ . Consequently, the three $\text{CS}\sigma$ bonds lead to the presence of 0.99 more electrons on the C atom in CS_3Li_3^+ than in CO_3Li_3^+ . This extra electron on C is consistent with the ionic attraction between the C and Li atoms in CS_3Li_3^+ that was previously noted by Leyva-Parra and coauthors.¹⁵ There is little change in the orbitals involved in π bonding upon transition from CO_3Li_3^+ to CS_3Li_3^+ . The C atom has a similar orbital structure in both CO_3Li_3^+ and CS_3Li_3^+ , closely resembling sp^2 hybridization.

Upon substitution of O by S, the partial charges on the chalcogen atoms become less negative, while the partial charge on the Li atom becomes only slightly less positive. Thus, the ionic attraction between the chalcogens and the Li atoms is expected to be weaker in CS_3Li_3^+ than in CO_3Li_3^+ . Moreover, the substitution of O by S decreases the strength of the covalent C–Li interaction from 14% to 6% relative to the C–Li interaction in CH_3Li . Therefore, no chemically meaningful covalent C–Li bonds exist in either CO_3Li_3^+ or CS_3Li_3^+ . Nonetheless, covalent



bonds alone do not determine the coordination number of a given atom. The IUPAC definition for coordination number reads: “the coordination number of a specified atom in a chemical species is the number of other atoms directly linked to that specified atom”.⁸⁸ While the definition of “linked” can be subject to interpretation, it is reasonable to conclude that it refers to an interaction capable of holding two atoms together. Consequently, the C atom in a given molecule should establish non-negligible interactions with six other atoms to be considered to be hexacoordinated. The present study shows that in the molecules discussed here, this may only occur (marginally) in CS_3Li_3^+ , where C–Li ionic attraction is plausible. On the other hand, the lack of C–Li covalent or ionic attractive interactions indicates that CO_3Li_3^+ cannot be considered to be a phC species.

Note that geometry alone is not always a good indicator of the bonding in a molecule. There is a variety of instances in which short interatomic distances do not correlate with favored bonding interactions. One example is the weakened C–C bonds in strained molecules, despite their frequently shorter C–C interatomic distances, relative to their unstrained counterparts.^{89–91} Despite the useful notions that geometries provide about the interactions that may occur in a molecule, it is evident that electronic structure theory tools, such as the QUAO analysis employed in the present work, enable a more rigorous study of the actual bonding structures in molecules. The importance of such tools is particularly highlighted for unusual molecules.

5. Conclusions

The bonding structures of CO_3Li_3^+ and CS_3Li_3^+ have been analyzed in terms of Quasi Atomic Orbitals. The bonding in CH_3Li has been analyzed and used as a reference along with previously reported QUAO results for the carbonate ion. The analysis shows that the bonding interactions between C and O in CO_3Li_3^+ closely resemble those in carbonate. The most notable difference between carbonate and the CO_3^{2-} moiety in CO_3Li_3^+ is observed to occur in the orbital structure of the lone pairs of the O atoms. These lone pair orbitals have increased s- and p-mixing in CO_3Li_3^+ , due to the inclusion of Li^+ ions into the molecular environment of CO_3^{2-} . The main bonding interactions of the Li atoms in CO_3Li_3^+ are with the O atoms, indicating covalent bonding between the “ligands” of the central carbon atom that contributes to the stabilization of the molecule. It is concluded that the introduction of Li^+ ions into the molecular environment of CO_3^{2-} has a greater effect on the O atoms than on the C atom, and that this introduction has little effect on the $\text{CO}\sigma$ and $\text{CO}\pi$ bonds.

Partial charges computed from QUAO populations indicate that ionic attraction between the C and Li atoms in CO_3Li_3^+ is not plausible, as both the C and Li atoms have positive partial charges.

The QUAO analysis shows that the substitution of the O atoms by S atoms slightly inverts the polarity of the carbon–chalcogen σ bond in the molecule, with respect to CO_3Li_3^+ . Similar to CO_3Li_3^+ , the main bonding interaction of the Li atoms in CS_3Li_3^+ are established with the S atoms.

The computed partial charges suggest that the ionic attraction between the Li and S atoms in CS_3Li_3^+ is weaker than that between the Li and O atoms in CO_3Li_3^+ . The partial charges also indicate the possibility of ionic attraction between the C and Li atoms in CS_3Li_3^+ , unlike in CO_3Li_3^+ .

There are extremely weak C–Li covalent interactions in both CO_3Li_3^+ and CS_3Li_3^+ that are estimated to have about 14% and 6% of the strength of the C–Li covalent interaction in CH_3Li , respectively. It is thus concluded that no chemically meaningful C–Li covalent bonds exist in either CO_3Li_3^+ or CS_3Li_3^+ .

It is concluded that because of the possibility of C–Li ionic attraction in CS_3Li_3^+ , it can indeed be considered as a planar hexacoordinated carbon species, while CO_3Li_3^+ cannot. This leads to the further conclusion that despite the useful notions provided by molecular geometries, electronic structure theory tools, such as the QUAO analysis employed in this work, are better suited for the study of the bonding in unusual molecules.

The insights presented in this work serve to emphasize the power of the quasi-atomic orbital analysis for providing understanding of the widely varying nature of the chemical bond. This is especially satisfying since these insights are drawn entirely from the wave function, with no implicit or explicit bias.

Data availability

All relevant data is available in the ESI.†

Conflicts of interest

There are no conflicts to declare.

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