How $\pi$ back-donation quantitatively controls the CO stretching response in classical and non-classical metal carbonyl complexes†

Giovanni Bistoni,*ab Sergio Rampino,*b Nicola Scafuri,c Gianluca Ciancaleoni,b Daniele Zuccaccia,d Leonardo Belpassi*b and Francesco Tarantelli*ab

The CO stretching response upon coordination to a metal M to form [(L)nM(CO)]n complexes (L is an auxiliary ligand) is investigated in relation to the $\sigma$ donation and $\pi$ back-donation components of the M−CO bond and to the electrostatic effect exerted by the ligand−metal fragment. Our analysis encompasses over 30 carbonyls, in which the relative importance of donation, back-donation and electrostatics are varied either through the ligand in a series of [(L)Au(CO)]0/+ gold(I) complexes, or through the metal in a series of anionic, neutral and cationic homoleptic carbonyls. Charge-displacement analysis is used to obtain well-defined, consistent measures of $\sigma$ donation and $\pi$ back-donation charges, as well as to quantify the $\sigma$ and $\pi$ components of CO polarization. It is found that all complexes feature a comparable charge flow of $\sigma$ symmetry (both in the M−CO bonding region and in the CO fragment itself), which is therefore largely uncorrelated to CO response. By contrast, $\pi$ back-donation is exceptionally variable and is found to correlate tightly with the change in CO bond distance, with the shift in CO stretching frequency, and with the extent and direction (C → O or C ← O) of the CO π polarization. As a result, we conclusively show that $\pi$ back-donation can be an important bond component also in non-classical carbonyls and we provide the framework in which the spectroscopic data on coordinated CO can be used to extract quantitative information on the $\pi$ donor properties of metal−ligand moieties.

1 Introduction

The high affinity of carbon monoxide (CO) towards metals (M) has been known since the end of the nineteenth century† and its relevance has kept growing thereafter, both in pure and applied chemistry.‡ This has led many chemists to study in detail the coordination bond between M and CO in metal−carbonyl complexes, which is commonly described in terms of the Dewar−Chatt−Duncanson (DCD) model.§ According to this scheme, the interaction between M and CO involves the donation of electron charge from the carbon’s lone pair to the empty M orbitals of $\sigma$ symmetry ($M \rightarrow CO \sigma$ donation), and a back-donation from filled M to empty CO orbitals of $\pi$ symmetry ($M \rightarrow CO \pi$ back-donation). The effectiveness of this model for the description of the M−CO bond has been consolidated over the years by a large number of theoretical studies based on a variety of techniques, including energy decomposition schemes, Natural Bond Orbitals (NBO) analysis,$\dagger$ Electron Localization Function (ELF) approaches$\ddagger$ and the Quantum Theory of Atoms In Molecules (QTAIM).$\S\H$

On the experimental side, discussions on the nature of the M−CO bond are mostly based on the analysis of the variation in the CO stretching frequency $\nu_{CO}$ (via IR spectroscopy) and bond distance $r_{CO}$ (via X-ray crystallography) with respect to free CO ($r_{\text{free-CO}} = 1.12822 \, \text{Å}$). In most metal−carbonyl complexes the CO bond appears weakened, i.e., the stretching frequency decreases ($\Delta \nu_{CO} = \nu_{CO} - \nu_{\text{free-CO}} < 0$) and the bond distance increases ($\Delta r_{CO} = r_{CO} - r_{\text{free-CO}} > 0$), but in a minority of (mainly late-metal cationic) complexes, which are sometimes termed “non-classical”, the CO bond appears strengthened ($\Delta \nu_{CO} > 0$ and $\Delta r_{CO} < 0$). These differences in the CO stretching response to the M−CO bond formation in metal carbonyl complexes are commonly explained in terms of the relative importance of the DCD constituents of the M−CO bond. In particular, M → CO $\pi$ back-donation is represented as exerting a bond-weakening effect on CO, while M → CO $\sigma$...
donation is thought to act in the opposite way.\textsuperscript{19-20} This view relies on a molecular-orbital picture in which both the π acceptor and σ donor CO orbitals have a C–O anti-bonding character. However, while there is general agreement on the effect thus played by π back-donation, the role of σ donation has been brought into question in the last fifteen years.\textsuperscript{21-23} In particular, these studies suggest that the σ CO donor orbital has, rather, a weak bonding character and that the CO bond strengthens in non-classical complexes is an electrostatic effect due to the (positively charged) ligand–metal moiety, whereby the CO bonding orbitals of both σ and π symmetry are polarized in the C ← O direction, thus enhancing the covalency of the CO bond.

One way to schematically depict M(CO) bonding resorts to a simple Valence Bond (VB) picture. Focusing on the M(CO) moiety of a generic [(L)mM(CO)]\textsuperscript{m} complex, three VB structures differing for the extent of π back-donation can be written:

(a) \( -\text{M} - \text{C} \equiv \text{O}^+ \leftrightarrow \text{b} \) \( \text{M} = \text{C} \equiv \text{O} \leftrightarrow \text{c} \) \( +\text{M} \equiv \text{C} - \text{O}^- \)

In going from structure (a) to structure (b) and (c), where one has zero, one and two π* orbitals of CO engaged in back-bonding, the CO bond multiplicity goes from three to two to one. The relative weight of each structure will depend on the π donor properties of the specific [(L)mM(CO)]\textsuperscript{m} fragment. At the same time, the electronic structure of CO is also affected by the electric field generated by this fragment, especially in those cases when \( m \neq 0 \). For CO in the presence of an electric field generated, for instance, by a positively charged metal fragment (exemplified here with the symbol \( \oplus \)), three analogue VB structures can be written:

(d) \( -\oplus \) \( \text{C} \equiv \text{O}^+ \leftrightarrow \text{e} \) \( \oplus \text{C} \equiv \text{O} \leftrightarrow \text{f} \) \( \oplus +\text{C} - \text{O}^- \)

The presence of such electric field would in this case favour the triple bonded structure (d) over structures (e) and (f), featuring a double and single bond, respectively (an opposite weight, so that their interplay in driving CO stretching response may be difficult to disentangle.

Still, however, carbonyl complexes showing blue shifted (\( \Delta_{\text{CO}} > 0 \)) CO stretching frequencies are most often assumed to lack back-donation from the metallic fragment.\textsuperscript{19,24,25} Exemplary in this respect is the set of complexes [(L)\textsubscript{m}Au(CO)]\textsuperscript{m+} of gold(i) that have been experimentally characterized.\textsuperscript{24,26-31} Until last year, to our knowledge, nine gold(i) carbonyl complexes had been spectroscopically characterized: the ligand free [Au(CO)]\textsuperscript{+} (observed in neon matrix\textsuperscript{18}) and its derivatives with ligands Cl\textsuperscript{−}, Br\textsuperscript{−}, CF\textsubscript{3}\textsuperscript{−}, CO\textsuperscript{2−}, Mes\textsubscript{3}P,\textsuperscript{29} Stidipp,\textsuperscript{30} Idipp\textsuperscript{30} and [HB(3,5-(CF\textsubscript{3})\textsubscript{2}Pz)]\textsuperscript{4−},\textsuperscript{34} where Mes stands for 2,4,6-Me\textsubscript{3}C\textsubscript{6}H\textsubscript{2}, Stidipp for 1,3-bis(2,6-diisopropylphenyl)imidazolin-2-ylidene, Idipp\textsuperscript{30} for 1,3-bis(2,6-diisopropylphenyl)imidazol-2-ylidene and [HB(3,5-(CF\textsubscript{3})\textsubscript{2}Pz)]\textsuperscript{4−}\textsuperscript{31} is a fluorinated tris(pyrazol)borate ligand. They all exhibit blue shift of the CO frequency and therefore are classified as non classical. This has been taken by some authors as proof that the gold fragment gives poor or no back-donation.\textsuperscript{32,33} However, in apparent contradiction, both theoretical and experimental studies have shown that the π donor character of gold is usually far from negligible\textsuperscript{34-36} (especially toward carbon monoxide\textsuperscript{37}) with important effects in catalysis.\textsuperscript{38,39} Recently, furthermore, a gold(i) complex showing \( \Delta_{\text{CO}} < 0 \) has been fully characterized,\textsuperscript{40} bearing a neutral o-carbaborane diphosphine (DCPb) as an ancillary ligand. Such an “exception”, which is even more singular when considering that the formal positive charge should strengthen the CO bond, made the authors speak of “enhanced π back-donation” from the [(DCPb)Au\textsuperscript{+}] fragment.

For the reader’s convenience, an overview of the experimentally characterized systems, with the reported \( \Delta_{\text{CO}} \) values and reference to the original papers, is displayed in Fig. 1. An additional gold(i) system, [[MeB[3-(Mes)Pz]\textsubscript{2}]Au(CO)], has been preliminarily reported as red-shifted in ref. 41.

The relationship of the DCD constituents of coordination bonds, determined unambiguously via charge-displacement (CD) analysis,\textsuperscript{35,42} with spectroscopic observables has been the subject of some of our recent work,\textsuperscript{36,37,43} and in the present work we have used this analysis to systematically study an extensive series of carbonyl compounds. The unique power of CD analysis lies in the fact that it provides a complete picture, across the entire molecular space, of the charge flow of σ and π.

\[ \text{Fig. 1. The experimentally characterized gold(i) carbonyl complexes discussed in this work, with the observed carbonyl stretching frequency shifts, } \Delta_{\text{CO}} \text{ (in cm}^{-1}) \text{, and literature references.} \]
character accompanying the formation of a coordination bond, and it permits a well-defined, consistent measure of the charge transfer (CT) associated both with the DCD components of the M–CO bond and with the σ and π components of the polarization taking place at the CO ligand itself. As a result, as we hope will we have convinced the reader by the end of the paper, this work provides a definitive and quantitative account of the role and interplay of the DCD components of the M–CO bond and of CO polarization in driving CO stretching response to coordination.

We thus investigate the relation between Δρ_{CO} and Δρ_{M:C} and the charge displacements of σ and π symmetry along the M–C–O axis in response to the M–CO bond formation in metal carbonyl complexes. We carry out our analysis first on an exhaustive series of 23 gold(I) carbonyls of formula [(L)Au(CO)]^{0/+}, where L is a varying auxiliary ligand (including none), which includes 8 of the experimentally characterized complexes and which is evenly partitioned between charged and neutral complexes, as well as between classical (CO bond elongated and frequency red-shifted) and non-classical (CO bond shortened and frequency blue-shifted). The choice of binary gold complexes seems to be particularly simple and useful, as it permits to isolate and study systematically the effect of the trans ligand across a wide variety of metal binding properties and electronic effects. We begin our analysis (Section 3.1) by studying in greater detail the two extreme cases of “naked” Au^+, [Au(CO)]^+, which displays the experimentally largest blue-shift, and of [(DPCb)Au(CO)]^+, which is the only known case of a positively charged but significantly red-shifted gold(I) complex. Having thus highlighted the main findings, we then thoroughly confirm them by extending the study to the whole series of complexes (Section 3.2). To complete the work we then also investigate the role of the metal itself in driving CO response to coordination, by studying a series of homoleptic [(CO)_nM(CO)]^{m/+} complexes, with M including Hg, Ir, Ni, Fe, Cr, Mo, Co, Ru (Section 3.3). Finally, an ad hoc study of CO in a uniform axial electric field (Section 3.4) concludes the work, in order to isolate the impact of CO polarization and of its σ and π components on CO stretching response.

2 Methodology and computational details

In the charge-displacement (CD) analysis framework, a chemical bond A–B is analyzed in terms of the difference Δρ(x,y,z) between the electron density of the adduct AB and that of the two non-interacting fragments A and B frozen at their in-adduct geometries. A partial progressive integration of Δρ(x,y,z) along a suitably chosen bond axis z yields the so called charge-displacement function (CDF)^42

$$\Delta \rho (z) = \int_{-\infty}^{x} dx' \int_{-\infty}^{y} dy' \int_{-\infty}^{z} \Delta \rho (x, y, z') dx dy dz . \tag{1}$$

The CDF defines, at each point z, the exact amount of electron charge displaced from right to left (the direction of decreasing z) upon bond formation through a plane perpendicular to the z axis through the point z (negative CDF values indicate a charge flow in the opposite direction). If both the adduct and its constituting fragments have proper symmetry, Δρ(x,y,z) can be decomposed into additive components of σ and π symmetry with respect to the bond axis z (see ref. 35 for further details).

All of the complexes studied in this work have general formula [(L)M(CO)]^{m/+}. Since the M–CO bond is under investigation, the appropriate fragments are the ligand–metal moiety [(L)M]^{m/+} and carbon monoxide CO, and the z reference axis joins the M and C centres. For the purpose of separating the σ and π components of Δρ(x,y,z), we group the orbitals of adduct and fragments according to the irreducible representations of the complex (and fragments) symmetry groups, which in the present cases are either the C_{2v} group (where the A1 representation corresponds to σ, while B1 and B2 correspond to π) or the C_{3v} group (A1 corresponding to σ donation, and E1 to π back-donation). No CO orbital is of A2 symmetry, therefore this representation is not relevant for the DCD analysis of the M–CO bond and is found to represent only a (minor) rearrangement internal to the ligand–metal fragment. Among the gold(I) complexes considered, [(PF_6)Au(CO)]^+, [(PPh_3)Au(CO)]^+, [(P(CH_3)_3)Au(CO)]^+, [(CF_3)Au(CO)] and [(CH_3)Au(CO)] belong to the C_{3v} point group. All others have C_{2v} symmetry. The symmetry point groups of the homoleptic complexes considered in Section 3.3 are listed in Table 2. The reduced symmetries C_{2v} and C_{3v} have been used to separate the σ and π components of the electron density difference also for these complexes.

The CDFs of the σ and π components of Δρ(x,y,z) provide a thorough, spatially detailed picture of the DCD donation and back-donation charge flows.^{38} Well-defined measures of the net charge transfer and of its donation and back-donation contributions (hereafter CT_{nets}, CT_{don} and CT_{back}, respectively) can be obtained by taking the CDFs values at a plausible inter-fragment boundary, which we take to be the z point where equal-valued isodensity surfaces of the fragments become tangent.^{35,36}

As mentioned in the Introduction, in the present context the CDF also provides precious additional information concerning CO polarization. Since the C–O bond is collinear with the M–C z axis of integration, the CDF in the C–O bond region represents the electron displacement within CO with respect to free CO in response to the M–CO bond formation. The amount of charge flowing across a plane normal to the CO bond through its midpoint (i.e., the CDF value at z = r_{CT}/2) can be usefully taken as a quantitative estimate of such polarization, and the total value can again itself be decomposed in σ and π components. We shall refer to these values as to CT_{f_{σ}/2}, CT_{f_{π}/2} and CT_{r_{CT}/2} respectively.

Geometry optimizations and the calculation of harmonic frequencies and electron densities were carried out by means of Density Functional Theory (DFT) with the ADF package.^{45–47} Becke’s exchange functional^{48} in combination with the Lee–Yang–Parr correlation functional^{49} (BLYP) was adopted. We used an all electron triple-zeta basis set with two polarization functions (TZ2P) and a small frozen core for all atoms. Relativistic effects were included via the zeroth-order regular approximation (ZORA) Hamiltonian.^{58–52} An assessment of the
3 Results and discussion

As mentioned in the Introduction, we first describe here a detailed investigation of the M–CO bond in [(L)Au(CO)] and [(DPCb)Au(CO)] (Section 3.1). We then extend the analysis to a series of 21 [(L)Au(CO)] complexes (Section 3.2) and, finally, to a series of nine homoleptic complexes of general formula [(CO)ₙM(CO)] (Section 3.3). The full list of complexes considered is in Tables 1 and 2. The purely electrostatic effect is investigated in the last Section (3.4) where an analysis of CO in a uniform axial electric field is carried out.

Table 1: Computed Δr̃_{CO} (Å) and Δr̃_{CO} (Å) and charge-transfer results (e) obtained from the CD analysis for the considered series of [(L)Au(CO)] complexes. Boldface, data for the experimentally observed complexes. Reference values are r_{free-CO} = 1137 Å. Δr̃_{CO} is the vibrational coupling between the CO and the ligand has been eliminated through isotopic substitution [(CF₃)Au(CO)] and [(Cl)Au(CO)] and [(Br)Au(CO)]. The other two have been excluded from our analysis because they are much more asymmetric and to constrain them to C₃v symmetry would probably alter their properties significantly.

Classical behavior

| Complex | Δr̃_{CO} (Å) | Δr̃_{CO} (Å) | CT_{net} | CT_{don} | CT_{back} | CT_{iso/2} | CT_{iso/2} |
|---------|-------------|-------------|---------|---------|----------|-----------|-----------|
| [(CF₃)Au(CO)] | 2 (51 ref. 24) | 0.002 | -0.02 | 0.22 | -0.24 | 0.05 | 0.05 |
| [(CN)Au(CO)] | 6² | 0.003 | -0.06 | 0.20 | -0.26 | 0.05 | 0.05 |
| [(H)Au(CO)] | 23 | 0.004 | -0.07 | 0.19 | -0.26 | 0.05 | 0.05 |
| [(CH₃)Au(CO)] | 42 | 0.007 | -0.07 | 0.21 | -0.28 | 0.01 | 0.04 |
| [(C₂H₅)Au(CO)] | 45 | 0.007 | -0.06 | 0.22 | -0.28 | 0.02 | 0.04 |
| [(I)Au(CO)] | 45 | 0.008 | -0.08 | 0.24 | -0.32 | 0.02 | 0.05 |
| [(Cl)Au(CO)] | 29 (13 ref. 27) | 0.008 | -0.11 | 0.23 | -0.33 | 0.02 | 0.05 |
| [(Br)Au(CO)] | 40 (10 ref. 27) | 0.008 | -0.09 | 0.24 | -0.33 | 0.02 | 0.05 |
| [(F)Au(CO)] | 23 | 0.009 | -0.13 | 0.22 | -0.35 | 0.02 | 0.05 |
| [(DPCb)Au(CO)] | -76 (30 ref. 40) | 0.010 | -0.06 | 0.26 | -0.32 | 0.03 | 0.05 |

Non classical behavior

| Complex | Δr̃_{CO} (Å) | Δr̃_{CO} (Å) | CT_{net} | CT_{don} | CT_{back} | CT_{iso/2} | CT_{iso/2} |
|---------|-------------|-------------|---------|---------|----------|-----------|-----------|
| [(CO)Au(CO)] | 72 (74 ref. 28) | -0.010 | 0.08 | 0.21 | 0.15 | 0.15 | 0.06 |
| [(PF₆)Au(CO)] | 80 | -0.009 | 0.09 | 0.22 | -0.13 | 1.15 | 0.06 |
| [(Au(CO)] | 75 (94 ref. 26) | -0.008 | 0.15 | 0.34 | -0.18 | 0.16 | 0.07 |
| [(Ne)Au(CO)] | 76 | -0.008 | 0.15 | 0.33 | -0.18 | 0.16 | 0.07 |
| [(Cl)Au(CO)] | 63 | -0.007 | 0.08 | 0.24 | -0.16 | 0.13 | 0.06 |
| [(Br)Au(CO)] | 60 | -0.007 | 0.06 | 0.22 | -0.16 | 0.12 | 0.05 |
| [(C₂H₅)Au(CO)] | 63 | -0.007 | 0.06 | 0.23 | -0.17 | 0.13 | 0.06 |
| [(Xe)Au(CO)] | 62 | -0.006 | 0.08 | 0.27 | -0.19 | 0.13 | 0.06 |
| [(P(CH₃)₃)Au(CO)] | 38 | -0.004 | 0.05 | 0.22 | -0.17 | 0.10 | 0.05 |
| [(NH₃)Au(CO)] | 39 | -0.004 | 0.00 | 0.20 | -0.20 | 0.10 | 0.05 |
| [(C₂H₅)₃N]Au(CO)] | 46 | -0.003 | 0.01 | 0.23 | -0.22 | 0.10 | 0.05 |
| [(Snipp)Au(CO)] | 23 (54 ref. 30) | -0.002 | -0.01 | 0.20 | -0.21 | 0.08 | 0.05 |
| [(Iddipp)Au(CO)] | 17 (49 ref. 30) | -0.001 | -0.02 | 0.20 | -0.22 | 0.08 | 0.05 |

As Table 1 shows, the experimental CO stretching frequency for the three complexes [(CF₃)Au(CO)], [(Cl)Au(CO)] and [(Br)Au(CO)] (the first of which is measured in the solid state and the others in solution) is actually blue-shifted rather than red-shifted as the calculations consistently suggest for all the neutral systems (the computed v_{free-CO} is 2143 cm⁻¹). Regarding this apparent inconsistency, Frenking et al. recently found that the experimental blue shift is actually due to the presence of intermolecular interactions and not to the properties of the single molecule. They proved this by computing the CO frequency of small aggregates of [(CF₃)Au(CO)] and of [(Cl)Au(CO)] and finding that the frequency increases from smaller to larger values than that of free CO. Indeed, Au–Au interactions have been experimentally observed for these two complexes in the solid state and are likely to occur also in solution, especially for ligands with little steric hindrance. For this reason, and since experimental data are available only for a small subset of the complexes considered here, we shall base our discussion on the DFT values of Δr̃_{CO} and Δr̃_{CO} (computed r_{free-CO} = 1137 Å). In fact, we shall most often refer
to the latter parameter only, because the non-uniform influence of vibrational mode coupling, and the more complicated CO vibration modes in the homoleptic carbonyls, make $\Delta r_{\text{CO}}$ a less reliable parameter than $\Delta r_{\text{CO}}$ for a quantitative analysis of its relation with the M–CO bond characteristics.

### 3.1 $[\text{Au(CO)}]^+$ and $[(\text{DPCb})\text{Au(CO)}]^+$

We start our analysis with an in-depth investigation of the gold carbon coordination bond in $[\text{Au(CO)}]^+$ and $[(\text{DPCb})\text{Au(CO)}]^+$. As mentioned in the Introduction, among the experimentally characterized gold carbonyl complexes, these two systems display the most different spectroscopic properties. $[\text{Au(CO)}]^+$ (observed in neon matrix$^{29}$) shows a CO stretching frequency much higher than that of free CO (experimental $\Delta r_{\text{CO}} = 94$ cm$^{-1}$) while $[(\text{DPCb})\text{Au(CO)}]^+$ (ref. 40) represents a unique case of cationic complex with red-shifted CO stretching frequency (experimental $\Delta r_{\text{CO}} = -30$ cm$^{-1}$). The computed values ($\Delta r_{\text{CO}} = 75$ cm$^{-1}$ and $-76$ cm$^{-1}$, respectively, see Table 1) reflect this opposite behavior.

We focus first on $[\text{Au(CO)}]^+$, showing in Fig. 2 the CDFs for the overall density difference and its symmetry-separated components. We recall here that, at a given point $z$, a positive CDF value corresponds to a charge flow from right to left (i.e., in the Au$^+$ $\rightarrow$ CO direction) while a negative value corresponds to a charge flow in the opposite (Au$^+$ $\rightarrow$ CO) direction. The total CDF is positive over both the Au–C and C–O bond regions and also at the oxygen far side of CO, indicating a continuous flow of electrons in the direction from CO towards gold. The negative values of the curve on the left side of Au$^+$ indicate a rearrangement in the opposite direction, which was shown in ref. 42 to be due to gold sd hybridization. The total CDF results from an $A_1$ component which is large and positive in the Au–carbon region (identifying $\sigma$ donation) and a $B_1 + B_2$ component which is negative in the same zone (identifying $\pi$ back-donation) plus a negligible $A_2$ component. These components are easily recognized in the isodensity plots of the respective density difference shown at the top of the figure.

The net charge transfer $\Delta r_{\text{net}}$ from CO to Au$^+$ (the CDF value at the boundary solid vertical line) amounts to 0.16e resulting from a donation component $\Delta r_{\text{don}}$ of 0.34e and a back-donation component $\Delta r_{\text{back}}$ of 0.18e. The first important comment here is that, in a system like this showing a large blue-shift of the CO stretching frequency, back-donation is actually a significant component of the interaction, estimated to be more than half as large as the donation.

An analogous significant contribution from the electron charge rearrangement of $\pi$ symmetry was also recently highlighted in ref. 29 through a Natural Orbitals for Chemical Valence-Extended Transition State (NOCV-ETS)$^{34}$ energy decomposition analysis. In particular, the $\pi$ contribution to the overall orbital interaction energy $\Delta E_{\text{orb}}$ was found to be surprisingly large (32.5% of the overall $\Delta E_{\text{orb}}$). The authors were cautious, however, in attributing such contribution exclusively to $\pi$ back-donation, as $\Delta E_{\text{orb}}$ not only accounts for genuine inter-fragment orbital interactions but also for the polarization of the orbitals within each fragment.

This uncertainty may be dissolved here, because, as discussed in Section 2, the interfragment charge transfer and its components are automatically separated from the corresponding components of CO polarization in the CDF picture. Inspection of Fig. 2 is in fact particularly revealing in this respect. Focusing on the CDFs in the carbonyl region, we notice immediately that the positive value of the total function indicates that the CO bond is on the whole polarized in the C $\rightarrow$ O direction. Remarkably, this polarization results from the concordant positive contributions of both the $\sigma$ and $\pi$ components. We indeed see that, while the $\sigma$ CDF keeps its (positive) sign on the right hand side of C and even beyond the oxygen site, an inversion (from negative to positive) is seen to occur for the $\pi$ component precisely at the carbon site, leading to a maximum located at about the mid-point of the C–O bond. In both cases, therefore, there is a displacement of electrons from oxygen towards carbon, which is due to the presence of the positively charged metal fragment. As discussed in Section 2, we can quantify the extent of CO polarization by taking the CDFs values at the mid-point of the CO bond (dashed vertical line in Fig. 2). For the case under examination, the $C \rightarrow O$ polarization amounts to $\Delta r_{\text{C/O}} = 0.16e$, resulting from a $\sigma$ contribution $\Delta r_{\text{C/O}}$ of 0.07e and a $\pi$ contribution $\Delta r_{\text{C/O}}$ of 0.09e.

We now turn to $[(\text{DPCb})\text{Au(CO)}]^+$, with its CDFs reported in Fig. 3. This is analogous to Fig. 2 except that here the $B_1$ (dashed blue line) and $B_2$ (dotted-dashed line) components are not identical and are shown separately in the plot. We notice an immediate striking contrast with the previous $[\text{Au(CO)}]^+$ case, in that the back-donation components globally dominate over $\sigma$

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**Fig. 2** Total CDF and its symmetry ($C_{\text{av}}$) components for the Au–CO bond in the complex $[\text{Au(CO)}]^+$. Black dots indicate the $z$ position of the atomic nuclei. A solid vertical line marks the boundary between the Au$^+$ and the CO fragments (see Section 2 for its definition). A dashed vertical line indicates the midpoint of the C–O bond ($z = r_{\text{CO}}/2$). At the top: isodensity surfaces ($\pm$0.0025 e a.u.$^{-1}$ (ref. 3)) for the $A_1$, $B_1$ and $B_2$ components of $\Delta \rho(x,y,z)$. Red surfaces identify charge depletion areas, blue surfaces identify charge accumulation areas.
donation in the coordination bond region, so that the total CDF is negative everywhere, indicating a continuous, though modest, flow of electrons from [[DPCb]Au]+ to CO. This confirms the already cited findings of ref. 40. We note that π back-donation is in turn largely dominated by the B2 component. The net charge transfer at the inter-fragment boundary is −0.06e, resulting from a σ donation component of 0.26e (A1) and a π back-donation component of −0.32e (−0.07 due to the B1 component and −0.25 due to the B2 component).

The polarization of the electron cloud in the carbonyl region also differs remarkably from that in [Au(CO)]2−. In analogy with [Au(CO)]2−, the σ CDF remains positive in the CO region and the B1 component turns positive at the C site, reflecting the polarization of the CO bonding orbitals due to the electrostatic effect of the metal fragment. However, by contrast, the B2 component maintains its negative sign also in the CO region, i.e. the back-donation it represents is so pronounced that it penetrates the CO region and extends even beyond the oxygen. As a consequence, the CO bond is on the whole slightly polarized in the C ← O direction (CTσ_COO/2 = 0.03e), resulting from a σ polarization in the same direction (CTσ_COO/2 = 0.05e) and a π polarization in the opposite C → O direction (CTπ_COO/2 = −0.02e).

It is worth deepening the comparison between the two complexes examined so far. In both, the metallic fragment bears a formal positive charge. However, [Au(CO)]2− behaves non-classically [blue-shifted ΔrCO], while [[DPCb]Au(CO)]2− behaves classically [red-shifted ΔpCO]. The CD analysis reveals that the σ donation component of the metal-CO bond is roughly comparable in the two cases (CTσ_don 0.34 vs. 0.26e), while π back-donation is almost twice as large in [[DPCb]Au(CO)]2− (CTπ_back 0.32 vs. 0.18e) and its extent substantially reduces the C ← O polarization of the CO bond. The polarization of the CO σ bonding orbitals is comparable in the two complexes (CTσ_COO/2 0.07 vs. 0.05e), but that of the π bonding orbitals is not (CTπ_COO/2 0.09 vs. −0.02e). These findings suggest that π electron displacement upon coordination is the main factor driving CO bond response. In particular, if the presence of the metal fragment, especially if positively charged, is capable of polarizing the π CO bonding orbitals, even in the presence of a significant back-donation, the CO bond is strengthened; if, on the other hand, π back-donation is strong and extended enough to contrast CO polarization, even in the presence of an equally cationic metal fragment, the CO bond is weakened.

### 3.2 The complete [[L]Au(CO)]0/+ series

We now need to verify if the above preliminary surmise stands the test of a wider series of carboxyl compounds. To this end, we have extended the analysis to all 23 [[L]Au(CO)]0/+ complexes listed in Table 1, which collects the spectroscopic data for ΔrCO and ΔpCO as well as the various computed CT figures. The complexes are listed in order of increasing ΔrCO and the experimentally characterized compounds are those shown in boldface. As briefly discussed at the beginning of Section 3, it is seen that, according to our computed shifts, the neutral complexes plus [[DPCb]Au(CO)]2− behave classically, while the remaining cationic complexes behave non-classically. The σ donation and π back-donation CDFs for these complexes are collected, respectively, in the top and bottom panel of Fig. 4. Red lines are for the complexes showing red shift of rCO, blue lines are for those showing blue shift.

Two eye-catching features emerge upon inspection of Fig. 4. The first is that all systems exhibit a surprisingly similar σ charge rearrangement (top panel) in the CO fragment region, in contrast with a much wider variability on the metal fragment side and despite the fact that some of the complexes are neutral and some cationic. In fact, as Table 1 shows, if one excludes the special cases of the naked Au+, of the inert ligands Ne and Xe, and of the anomalous [[DPCb]Au(CO)]2−, even the net ligand-to-metal σ donation, CTσ_don, varies by only 0.05e across the whole series of ligands. On the contrary, the π CDF (bottom panel of Fig. 4) appears to be strongly influenced by the nature of the ligand over the whole molecular region, and CTπ_back varies by 0.22e over the ligand series. The second important observation is that, in the CO region, the complexes showing a blue-shifted rCO (blue lines) all invariably exhibit a flow of π electrons in the C ← O direction (CTπ_COO/2 > 0), due to the positively charged metallic fragment, while the complexes showing red-shifted rCO (red lines) exhibit a negative CTπ_COO/2, i.e., charge flows in the opposite C → O direction (with the exception of two complexes for which CTπ_COO/2 is essentially vanishing and the red-shift is also negligibly small).

It thus appears quite clearly that in the series of gold(l) carboxyls: (i) σ donation is much less tunable than π back-
donation, being very little dependent on the nature and the charge of the ligand; (ii) whereas the net CO bond polarization turns out to be invariably oriented in the C–O direction (CT_{\text{net}} > 0), the direction of its \( \pi \) density component can vary and appears to be tightly connected with the direction of the CO stretching shift and bond-length change. These findings are given a definitive illustration in Fig. 5 and 6 where the correlation of \( \Delta r_{\text{CO}} \) with CT_{\text{net}}, CT_{\text{don}} and CT_{\text{back}} is reported. In both figures, black triangles are used for the overall CT, red squares for its \( \sigma \) component and blue circles for its \( \pi \) component. Empty symbols are for the neutral species, filled ones for the cationic species.

Focusing first on Fig. 5, no correlation is found, as expected, between \( \Delta r_{\text{CO}} \) and CT_{\text{don}}, while a good inverse correlation \( (R^2 = 0.945) \) can be seen between \( \Delta r_{\text{CO}} \) and CT_{\text{back}}, a trace of which remains in the plot of \( \Delta r_{\text{CO}} \) vs. CT_{\text{net}}. The same bond weakening effect of \( \pi \) back-donation is also evident in the plot of \( \Delta r_{\text{CO}} \) vs. CT_{\text{net}}.

3.3 Homoleptic complexes: the \([\text{CO}]_n\text{M(CO)}_m\) series

In the previous sections we considered a series of gold(I) complexes where the donor/acceptor properties of the M–CO bond were varied through the ligand L. We now extend the analysis to a series of homoleptic carbonyls of formula \([\text{CO}]_n\text{M(CO)}_m\), where the relative extent of the DCD constituents of CT_{\text{back}} (see ESI†), though correlation, as mentioned above, is made worse by mode coupling \( (R^2 = 0.849) \). Fig. 6 shows the correlation of \( \Delta r_{\text{CO}} \) with CT_{\text{net}} and its components CT_{\text{\sigma}} and CT_{\text{\pi}}. Not surprisingly, as these quantities are all directly related to the charge rearrangement of the CO bond itself, correlations are here quantitatively better \( (R^2 = 0.970 \text{ for that with CT}_{\text{\pi}}) \). Obviously, as \( \Delta r_{\text{CO}} \) correlates well with both \( \pi \) back-donation and CO \( \pi \) electron polarization, the latter two quantities are also in mutual correlation.
the M–CO bond and CO polarization are varied essentially by changing the metal. The full list of the considered homoleptic complexes is in Table 2, reporting their spectroscopic shifts and CD decomposition results. Complexes are listed in order of increasing value of \( \Delta \text{MCO} \). We omit for brevity a presentation of the complete CDFs. The computed structures for these systems are in agreement with experimental X-ray data where available.\(^{62-66}\) Hg(CO)\(_2\)\(^{2+}\) and Ir(CO)\(_6\)\(^{3+}\), both cationic, behave non-classically, with experimental blue-shifted \( r\text{MCO} \) at 2279.5 cm\(^{-1}\) for the former and at 2254, 2276 and 2298 cm\(^{-1}\) for the latter.\(^{65,66}\) On the opposite side, the anionic complexes show exceptionally low CO stretching frequency, the most red-shifted being that of Fe(CO)\(_4\)\(^{2-}\) at 1730 cm\(^{-1}\) (this is the first anionic carbonyl complex spectroscopically observed\(^{67,68}\)). In between are Mo(CO)\(_6\), Fe(CO)\(_5\) (for which both the axial and equatorial M–CO bonds have been investigated),\(^{69}\) Ni(CO)\(_4\) and Cr(CO)\(_6\). The complexes present therefore a wide range of \( r\text{MCO} \) variation but \( r\text{MCO} \) turns out not to be a good parameter for analyzing correlations with the CD data because normal-mode coupling varies significantly with the different structure of the complexes. We therefore base our discussion, as already done for the gold(I) complexes, on the computed \( \Delta \text{MCO} \). This varies in a range of 0.087 \( \text{Å} \) over the series, from \(-0.018\) to \(0.069\) \( \text{Å} \) (Table 2).

The tables shows that also in this series of compounds the range of variation in \( \pi \) back-donation (0.69e) is much larger than that of \( \sigma \) donation (0.15e). In particular, almost no back-donation is found for Hg(CO)\(_2\)\(^{2+}\) while \( CT_{\text{back}}^{\text{Fe(CO)\(_4\)}} \) is as high as 0.71e. This picture is consistent with the simple VB view discussed in the Introduction, in that we go from a purely \( \pi \) M–CO bond (structure a) for Hg(CO)\(_2\)\(^{2+}\) to a situation in which all \( \pi^* \) CO orbitals are engaged in back-bonding (structure c) for Fe(CO)\(_4\)^{2–}. Also the charge rearrangement (polarization) in the carbonyl region is seen to follow a similar trend, with a much narrower range of \( CT_{\text{polar}}^{\text{Fe(CO)\(_4\)}} \) values (between 0.02 and 0.10e) than that of \( CT_{\text{polar}}^{\text{Ir(CO)\(_6\)}} \) (from 0.21 to \(-0.30e\)). As before, no clear correlation can be discerned between \( \Delta \text{MCO} \) and \( \sigma \) \( \text{CT} \), while \( CT_{\text{back}}^{\text{Fe(CO)\(_4\)}} \) and \( CT_{\text{polar}}^{\text{Ir(CO)\(_6\)}} \) values are seen to decrease almost monotonically as \( \Delta \text{MCO} \) increases.

A plot of \( \Delta \text{MCO} \) vs. either \( CT_{\text{back}}^{\text{Fe(CO)\(_4\)}} \) or \( CT_{\text{polar}}^{\text{Ir(CO)\(_6\)}} \) for the whole set of complexes studied, including the present homoleptic carbonyls in addition to the gold(I) series, appears in fact to suggest, because the range of variation is now significantly enlarged, that a quadratic, rather than a linear one, may better represent the actual correlation (an evident non-linear relationship has already been found between the electric field strength and \( \Delta \text{MCO} \)). Fig. 7 very clearly shows this to be the case, with the accuracy of all fits improved with respect to the sole subset of gold complexes.

Once again, in the homoleptic series, the carbonyl complexes featuring CO bond strengthening (blue-shifted \( \Delta \text{MCO} \) and negative \( \Delta \text{MCO} \)), \( i.e. \) the cationic Hg(CO)\(_2\)\(^{2+}\) and Ir(CO)\(_6\)\(^{3+}\), show a flow of \( \pi \) electrons in the \( C \leftrightarrow O \) direction. All other complexes, where the CO bond weakens (red-shifted \( \Delta \text{MCO} \) and positive \( \Delta \text{MCO} \)) show opposite-direction flows.

### 3.4 CO in a uniform axial electric field

The observation that the CO bond is lengthened or shortened upon formation of the M–CO bond according to whether the CO bonding orbitals of \( \pi \) symmetry are polarized in the \( C \rightarrow O \) or \( C \leftarrow O \) direction, respectively, is certainly remarkable. To verify that this is a general fact, actually independent of CO coordination, we discuss in this last section an \textit{ad hoc} study of the

| Sym. | \( \text{Exp.} \Delta \text{MCO} \) | \( \Delta \text{MCO} \) | \( CT_{\text{net}} \) | \( CT_{\text{don}} \) | \( CT_{\text{back}}^{\text{Fe(CO)\(_4\)}} \) | \( CT_{\text{polar}}^{\text{Ir(CO)\(_6\)}} \) | \( CT_{\text{polar}}^{\text{Ir(CO)\(_6\)}} \) |
|------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Hg(CO)\(_2\)\(^{2+}\) | \( C_{\text{v}} \) | 136 (ref. 66) \( \{A_1\} \) | \(-0.018\) | 0.29 | 0.31 | \(-0.02\) | 0.13 | 0.08 | 0.21 |
| Ir(CO)\(_6\)\(^{3+}\) | \( O_h \) | 155 (ref. 65) \( \{A_2\} \) | \(-0.015\) | 0.17 | 0.31 | \(-0.13\) | 0.08 | 0.10 | 0.18 |

**Classical behavior**

| Sym. | \( \text{Exp.} \Delta \text{MCO} \) | \( \Delta \text{MCO} \) | \( CT_{\text{net}} \) | \( CT_{\text{don}} \) | \( CT_{\text{back}}^{\text{Fe(CO)\(_4\)}} \) | \( CT_{\text{polar}}^{\text{Ir(CO)\(_6\)}} \) | \( CT_{\text{polar}}^{\text{Ir(CO)\(_6\)}} \) |
|------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Ni(CO)\(_4\) | \( T_d \) | \(-23\) (ref. 70) \( \{A_1\} \) | 0.012 | \(-0.16\) | 0.16 | \(-0.32\) | 0.01 | 0.04 | \(-0.05\) |
| Fe(CO)\(_5\) (ax.) | \( D_{3h} \) | \(-22\) (ref. 69) \( \{A_1\} \) | 0.014 | \(-0.18\) | 0.23 | \(-0.41\) | 0.02 | 0.06 | \(-0.08\) |
| Cr(CO)\(_6\) | \( O_h \) | \(-24\) (ref. 71) \( \{A_2\} \) | 0.016 | \(-0.21\) | 0.17 | \(-0.37\) | 0.02 | 0.05 | \(-0.07\) |
| Mo(CO)\(_6\) | \( O_h \) | \(-22\) (ref. 71) \( \{A_2\} \) | 0.016 | \(-0.23\) | 0.14 | \(-0.37\) | 0.02 | 0.04 | \(-0.06\) |
| Fe(CO)\(_5\) (eq.) | \( D_{3h} \) | \(-22\) (ref. 69) \( \{A_1\} \) | 0.018 | \(-0.20\) | 0.20 | \(-0.40\) | 0.03 | 0.05 | \(-0.08\) |
| Co(CO)\(_5\) | \( T_d \) | \(-255\) (ref. 72) \( \{A_1\} \) | 0.038 | \(-0.36\) | 0.16 | \(-0.52\) | 0.15 | 0.03 | \(-0.18\) |
| Ir(CO)\(_4\) | \( T_d \) | \(-248\) (ref. 73) \( \{A_1\} \) | 0.039 | \(-0.28\) | 0.30 | \(-0.58\) | 0.14 | 0.04 | \(-0.18\) |
| Ru(CO)\(_4\)\(^{2+}\) | \( T_d \) | \(-407\) (ref. 74) \( \{A_1\} \) | 0.066 | \(-0.42\) | 0.25 | \(-0.67\) | 0.24 | 0.03 | \(-0.27\) |
| Fe(CO)\(_4\)\(^{2+}\) | \( T_d \) | \(-413\) (ref. 75) \( \{A_1\} \) | 0.069 | \(-0.55\) | 0.16 | \(-0.71\) | 0.28 | 0.02 | \(-0.30\) |
electron cloud rearrangement and stretching response of CO in an external uniform axial electric field oriented along the C-O bond axis.

In Fig. 8, we show the computed CO stretching $\Delta r_{CO}$ reported versus the $\pi$ and $\sigma$ components of $CT_{\pi\sigma/2}$. The latter vary as a result of the applied field in the same figure. The points representing the computed $\Delta r_{CO}$ and $\pi$ and $\sigma$ components of $CT_{\pi\sigma/2}$ are reported for the whole series of carbonyl complexes studied in this work.

Let us focus first on the stretching response to the electric field. When the field is absent, the system corresponds to free CO and $\Delta r_{CO}$, $CT_{\pi\sigma/2}$ and $CT_{\sigma\pi/2}$ are all zero. As the field increases on the left, in the direction that induces (linearly) $C \rightarrow O$ (negative) polarization, C-O bond length increases quadratically and $\pi$ polarization is seen to increase much more rapidly than $\sigma$ polarization. Conversely, as the field increases on the right, inducing $C \leftarrow O$ polarization, the C-O bond shortens (much less rapidly).

![Fig. 7](image_url) Correlation between CO bond-length change upon coordination, $\Delta r_{CO}$, and (a) $M \rightarrow CO$ $\pi$ back-donation $CT_{\pi\pi/2}$ (upper panel), (b) $CO$ $\pi$ electron polarization $CT_{\pi\pi/2}$ (lower panel), for the whole set of complexes studied. Black-colored points refer to the gold(I) complexes, empty circles for neutral or negatively charged ones. Both the linear fit ($R^2 = 0.959$ for the upper panel, $R^2 = 0.945$ for the lower panel) and the quadratic fit ($R^2 = 0.984$ for the upper panel and $R^2 = 0.996$ for the lower panel) curves are shown.

![Fig. 8](image_url) The dashed lines show $\Delta r_{CO}$ versus the $\sigma$ (red color) and $\pi$ (blue) CO polarization $CT_{\pi\pi/2}$ for a CO molecule placed in a uniform axial electric field of magnitude ranging from $-0.1$ to $0.07$ a.u. in steps of $0.02$ a.u. (colored square points). For comparison, also shown is the correlation between $\Delta r_{CO}$ and $\sigma$ and $\pi$ CO polarization (red and blue circles, respectively) for the whole series of Au and homoleptic complexes studied. The empty circles are for neutral or negatively charged complexes, the filled circles for cationic ones.

When we now compare these curves with the relation observed between $\Delta r_{CO}$ and the $\sigma$ and $\pi$ components of CO polarization induced by metal coordination, rather than by an applied field (disconnected circles in the figure), we notice immediately that the $\pi$ circles follow quite closely the correlation between field-induced polarization and stretching, while, in striking contrast, the $\sigma$ circles deviate from the field-induced line (a clear indication of a much more pronounced “chemical” signature) and, moreover, span a very narrow range of (positive) polarization, essentially without any correlation with the widely varying $\Delta r_{CO}$. This is indeed a very strong confirmation that the CO stretching response to any solicitation causing electron charge rearrangement, be it the formation of a M-CO coordination bond or the effect of an external electric field, is driven essentially exclusively by the charge rearrangement of $\pi$ symmetry: whether induced by an external electric field or by metal coordination, $C \rightarrow O$ [$C \leftarrow O$] polarization of the $\pi$ bond orbitals invariably and tightly correlates with bond shortening (shortening).

4 Conclusions

In this work we have carried out an in-depth analysis of the M-CO bond in $[(L)_{n}(M(CO))_{m}]^{n}$ metal carbonyl complexes, with the aim of elucidating on quantitative grounds the $\sigma$ donation and $\pi$ back-donation effects on the CO stretching response, in particular the change in bond length $\Delta r_{CO}$, to the M-CO bond formation. The analysis was carried out for a large variety of carbonyls, in which the relative extent of the DCD constituents were varied both through $L$ in a series of $[(L)_{n}(Au(CO))_{m}]^{n+}$ gold(i) carbonyl complexes and through $M$ in a series of anionic, neutral and cationic $[(CO)_{n}(M(CO))_{m}]^{n}$ homoleptic carbonyls. Crucially, for the purpose of this investigation, reliable and consistent measures, not only of $\sigma$ donation and $\pi$ back-
donation charges but also of the \(\sigma\) and \(\pi\) components of CO polarization were obtained by the well-established charge-displacement analysis of electron-density differences, as resulting from accurate DFT calculations. The nature of the M-CO bond in the considered complexes was found to range smoothly between the two extreme cases of an almost purely \(\sigma\) bonded complex (\(\text{Hg(CO)}_2^+\), \(\Delta r_{\text{CO}} = 0.02\)) and of a strongly back-bonded complex (\(\text{[Fe(CO)}_3]^{-}\), \(\Delta r_{\text{CO}} = 0.71\)). Conversely, all complexes were found to feature a narrowly comparable \(\sigma\) donation component, with \(\Delta r_{\text{don}}\) values ranging from 0.14 to 0.34e. The same picture holds accurately for the electron cloud rearrangement over the carbonyl region: all considered complexes feature a comparable \(\sigma\) polarization of CO and a much more variable \(\pi\) polarization. Quite remarkably, no correlation is found between \(\Delta r_{\text{CO}}\) and the \(\sigma\) displacements, while \(\Delta r_{\text{CO}}, \pi\) back-donation and CO \(\pi\) polarization all correlate tightly with one another.

These results show that the driving force of the CO stretching response to the M-CO bond formation is provided exclusively by the changes taking place in the \(\pi\) electron density. In the complexes studied, such \(\pi\) charge rearrangement is found to result from the interplay between \(\pi\) back-donation (structures \(a-c\) of the Introduction) and the electrostatic effect (structures \(d-f\)) exerted by the metal-ligand fragment. In particular, cationic metal–ligand fragments polarize the \(\pi\) CO bonding orbitals in the \(C \rightarrow O\) direction, thus shortening the bond and enhancing the covalency, as highlighted in ref. 23. This effect, on the other hand, is contrasted by \(\pi\) back-donation shifting charge in the opposite direction. The net direction \(C \leftarrow O\) or \(C \rightarrow O\) of the polarization of \(\pi\) CO bonding orbitals is found to invariably determine whether the CO bond is strengthened or weakened, respectively. This is most evident in the \(\text{[DPCb}}\) \(\text{Au(CO)}_2^+\) complex, where \(\pi\) back-donation is so strong as to invert the polarization of the \(\pi\) CO bonding orbitals in the \(C \rightarrow O\) direction despite the formal positive charge on the ligand–metal fragment, making it the only example of a cationic gold(i) carbonyl complex with classical behavior (\(\Delta r_{\text{CO}} > 0\)). An \textit{ad hoc} study of CO in a uniform axial electric field demonstrates that it is indeed the polarization of the \(\pi\) CO bonding orbitals, no matter how induced (whether by the coordination bond to M or by an electric field), that drives direction and magnitude of the CO stretching response to the M-CO bond formation.

Regarding the fundamental question of what can be inferred on the nature of the M-CO bond from the analysis of \(\Delta r_{\text{CO}}\) (and less reliably, due to mode coupling, \(\Delta r_{\text{CO}}\)) in metal carbonyl complexes, we conclude that the value of \(\Delta r_{\text{CO}}\) quantifies to an excellent extent the \(\pi\) back-donation component of the M-CO bond, since such component directly correlates with the \(\pi\) polarization. In particular, where \(CT_{\text{CO}}\) changes its sign (i.e. the polarization of \(\pi\) CO bonding orbitals changes direction determining whether the CO bond is weakened or strengthened), \(CT_{\text{back}}\) is approximately as high as the average extent of \(\sigma\) donation among the complexes herein considered. This indicates that \(\pi\) back-donation is an important component also in the class of non-classical complexes, as those of gold(i) considered in this work.

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References

1 A. Roberts, \textit{Chem. News}, 1898, 78, 260.
2 F. Calderazzo, \textit{Anorg. Chem., Int. Ed.}, 1977, 16, 299–311.
3 J.-K. Shen, Y.-C. Gao, Q.-Z. Shi and F. Basolo, \textit{Coord. Chem. Rev.}, 1993, 128, 69–88.
4 F. Fischer and H. Tropsch, \textit{Brennst.–Chem.}, 1923, 4, 276–285.
5 J. L. G. Fierro, \textit{Catal. Lett.}, 1993, 22, 67–91.
6 M. J. S. Dewar, \textit{Bull. Soc. Chim. Fr.}, 1951, 18, C71–C79.
7 J. Chatt and L. A. Duncanson, \textit{J. Chem. Soc.}, 1953, 2939–2947.
8 \textit{Modern Coordination Chemistry: The Legacy of Joseph Chatt}, ed. N. Winterton and J. Leigh, The Royal Society of Chemistry, 2002, pp. X001–X004.
9 T. Ziegler and A. Rauk, \textit{Inorg. Chem.}, 1979, 18, 1755–1759.
10 K. Nakashima, X. Zhang, M. Xiang, Y. Lin, M. Lin and Y. Mo, \textit{J. Theor. Comput. Chem.}, 2008, 7, 639–654.
11 S. Dapprich and G. Frenking, \textit{J. Phys. Chem.}, 1995, 99, 9352–9362.
12 F. Weinhold and C. R. Landis, \textit{Valency and bonding: a natural bond orbital donor–acceptor perspective}, Cambridge University Press, 2005.
13 E. Matito and M. Solà, \textit{Coord. Chem. Rev.}, 2009, 253, 647–665.
14 P. Macchi and A. Sironi, \textit{Coord. Chem. Rev.}, 2003, 238, 383–412.
15 J. Pilme, B. Silvi and M. E. Alikhani, \textit{J. Phys. Chem. A}, 2003, 107, 4506–4514.
16 F. Cortés-Guzmán and R. F. Bader, \textit{Coord. Chem. Rev.}, 2005, 249, 633–662.
17 D. Tiana, E. Francisco, M. Blanco, P. Macchi, A. Sironi and A. Martín Pendás, \textit{J. Chem. Theory Comput.}, 2010, 6, 1064–1074.
18 A. J. Lupinetti, S. H. Strauss and G. Frenking, \textit{Prog. Inorg. Chem.}, 2001, 49, 1–112.
19 P. K. Hurlburt, J. J. Rack, J. S. Luck, S. F. Dec, J. D. Webb, O. P. Anderson and S. H. Strauss, \textit{J. Am. Chem. Soc.}, 1994, 116, 10003–10014.
20 H. Willner and F. Aubke, \textit{Anorg. Chem., Int. Ed.}, 1997, 36, 2402–2425.
21 A. S. Goldman and K. Krogh-Jespersen, \textit{J. Am. Chem. Soc.}, 1996, 118, 12159–12166.
22 A. J. Lupinetti, S. Fau, G. Frenking and S. H. Strauss, \textit{J. Phys. Chem. A}, 1997, 101, 9551–9559.
23 G. Frenking, C. Loschen, A. Krapp, S. Fau and S. H. Strauss, \textit{J. Comput. Chem.}, 2007, 28, 117–126.
24 S. Martinez-Salvador, J. Forniés, A. Martín and B. Menjón, \textit{Angew. Chem., Int. Ed.}, 2011, 50, 6571–6574.
25 R. H. Crabtree, \textit{The organometallic chemistry of the transition metals}, Wiley, 2005.
26 B. Liang and L. Andrews, \textit{J. Phys. Chem. A}, 2000, 104, 9156–9164.
