Observation of alkaline earth complexes M(CO)\(_8\) (M = Ca, Sr, or Ba) that mimic transition metals

Xuan Wu\(^1\), Lili Zhao\(^2\), Jiaye Jin\(^1\), Sudip Pan\(^2\), Wei Li\(^1\), Xiaoyang Jin\(^1\), Guanjun Wang\(^1\), Mingfei Zhou\(^1\), Gernot Frenking\(^2,3\)

The alkaline earth metals calcium (Ca), strontium (Sr), and barium (Ba) typically engage in chemical bonding as classical main-group elements through their ns and np valence orbitals, where n is the principal quantum number. Here we report the isolation and spectroscopic characterization of eight-coordinate carbonyl complexes M(CO)\(_8\) (where M = Ca, Sr, or Ba) in a low-temperature neon matrix. Analysis of the electronic structure of these cubic \(O_\text{h}\)-symmetric complexes reveals that the metal–carbon monoxide (CO) bonds arise mainly from \([M(d_{\text{A}})] \rightarrow \text{CO} \equiv \pi\) backdonation, which explains the strong observed red shift of the C–O stretching frequencies. The corresponding radial cation complexes were also prepared in gas phase and characterized by mass-selected infrared photodissociation spectroscopy, confirming adherence to the 18-electron rule more conventionally associated with transition metal chemistry.

The periodic table of the elements is conventionally divided according to the valence atomic orbitals (AOs) into main-group s and p blocks, a transition metal d block, and a lanthanide and actinide f block. A useful set of guidelines for understanding the structures and stabilities of molecules encompasses the associated 8-, 18-, and 32-electron rules introduced by Langmuir (1, 2) before the advent of quantum theory. These rules were later explained by attributing particular stability to filled sp, spd, or spdf valence shells, respectively (3).

The alkaline earth elements beryllium, magnesium, calcium, strontium, and barium have a ns\(^2\) valence-shell configuration, where n is the principal quantum number, and, as such, typically engage in chemical bonding as ionic salt compounds or in polar bonds via their two ns valence electrons in divalent M(II) species (4), where M is an alkaline earth metal. Earlier studies suggested that the heaviest atom barium may use its 5d AOs to some extent in chemical bonds (5), which led to the suggestion that barium be designated an “honorary transition metal” (6). Previously, we reported the experimental observation of barium carbonyl ions Ba(CO)\(^9\) (where charge \(q = +1\) and \(-1\)) (7). The analysis of the electronic structure showed that the cation binds the ligand mainly through Ba\(^{5d_{5/2}}\) \(\rightarrow\) CO(\(\pi\) LUMO) backdonation (LUMO, lowest unoccupied molecular orbital), with Ba\(^+\) in the excited \(^3\text{D}(5d^1)\) electronic reference state. In that respect, the Ba(CO)\(^+\) complex behaves similarly to a transition metal carbonyl. The Ba-CO interactions in the radical anion Ba(CO)\(^-\) were consistent with dominant contributions of Ba\(^{5d}\) \(\rightarrow\) CO(\(\pi^*\) SOMO) donation (SOMO, singly occupied molecular orbital) and Ba(5d\(_{5/2}\)/6s) \(\rightarrow\) CO(\(\sigma\) HOMO) \(\sigma\) donation (HOMO, highest occupied molecular orbital). The most important valence functions of barium in Ba(CO)\(^+\) cation and Ba(CO)\(^-\) anion thus appeared to be the 5d orbitals (7).

These findings inspired us to search for the 18-electron octacarbonyl complex Ba(CO)\(_8\). Surprisingly, we found that not only barium but also the lighter homologs strontium and calcium formed octacarbonyl complexes M(CO)\(_8\) (M = Ca, Sr, or Ba) that can be stabilized in a low-temperature neon matrix. The neutral alkaline earth–carbonyl complexes were prepared by the reactions of pulsed laser-evaporated metal atoms and carbon monoxide (CO) in solid neon and were investigated using Fourier transform infrared absorption spectroscopy. The experiments were carried out with a wide range of CO concentrations (from 0.02 to 2% relative to Ne on the basis of volume). In the experiments with relatively low CO concentrations, terminally bonded mononuclear low-coordinate carbonyl complexes with C=O stretching frequencies in the 2050 to 1800 cm\(^{-1}\) region were observed. Experiments with isotopically substituted CO samples allowed the unambiguous identification of some low-coordinate complexes through isotope shifts and splittings. The barium di-, tri-, and tetracarbonyls can clearly be identified on the basis of spectra in the experiments with 0.03% \(^{12}\text{C}^{16}\text{O}\) (fig. S1); 0.05% \(^{13}\text{CO}\) and 0.05% \(^{16}\text{CO}\) (fig. S2); and 0.05% \(^{16}\text{O}\) and 0.05% \(^{18}\text{O}\) (fig. S3). The monocarbonyls were theoretically predicted to be unstable (8) and were not observed in the experimental vibrational spectra. Intense absorption bands centered at 1987 cm\(^{-1}\) for Ca, 1995 cm\(^{-1}\) for Sr, and 2014 cm\(^{-1}\) for Ba were observed upon progressive annealing of the samples to temperatures of 10 to 13 K under relatively high CO concentrations (Table 1). These absorptions become the dominant features in the spectra with high CO concentrations (see Fig. 1A for Ca and figs. S4 and S5 for Sr and Ba), suggesting that the absorber is the coordinatively saturated 18-electron octacarbonyl complex. The observation of only one carbonyl stretching band suggests that these neutral octacarbonyls have the highest cubic \(O_\text{h}\) symmetry. Experiments with mixtures of \(^{12}\text{C}^{16}\text{O}\) and \(^{13}\text{C}^{16}\text{O}\) and also \(^{12}\text{C}^{18}\text{O}\) and \(^{13}\text{C}^{18}\text{O}\) provided conclusive identification of these cubic octacarbonyl complexes. Although the bands of the Sr and Ba complexes are too broad to resolve isotopic splittings, the band of the Ca complex is sharp and intense in the spectra with relatively low CO concentrations (Fig. 1A); well-resolved mixed isotopic spectra could therefore be compared with calculations. The experimentally observed spectra are in good agreement with the simulated isotopic spectral features shown in figs. S6 and S7.

The radical cations of the alkaline earth–carbonyl complexes were prepared in the gas phase by using a pulsed laser vaporization–supersonic-expansion ion source and studied by mass-selected infrared photodissociation spectroscopy in the carbonyl stretching-frequency region. Typical mass spectra are shown in figs. S8 to S10. Mononuclear metal–carbonyl cation complexes [M(CO)\(_n\)]\(^+\) (M = Ca, Sr, or Ba) with n as high as 10 to 15 were observed. These larger complexes contain both strongly bound CO ligands directly coordinated to the central metal ion and weakly bound, or tagged, CO ligands (9, 10). All of these complexes dissociated by elimination of a CO ligand after photodissociation at the \(\text{C}=\text{O}\) stretching vibrational frequencies. The infrared spectra of [Sr(CO)\(_n\)]\(^+\) (n = 6 to 9) are shown in Fig. 1B and the corresponding spectra of the Ca and Ba complexes in figs. S11 and S12. All the spectra for the [M(CO)\(_n\)]\(^+\) complexes with n = 6 to 8 feature a broad single band (full width at half maximum (FWHM) of 49 cm\(^{-1}\) for Ca, 52 cm\(^{-1}\) for Sr, and 29 cm\(^{-1}\) for Ba for the n = 8 complexes) that is slightly red shifted relative to the free-CO absorption at 2143 cm\(^{-1}\). The dissociation efficiency increases substantially (about 70% for Ca, 220% for Sr, and 107% for Ba) from n = 8 to 9, and the n = 9 complexes have a much narrower bandwidth (FWHM of 36 cm\(^{-1}\) for Ca, 10 cm\(^{-1}\) for Sr, and 16 cm\(^{-1}\) for Ba). Along with the intense band around 2100 cm\(^{-1}\), an additional weak band in the 2160 to 2180 cm\(^{-1}\) region is also observed for the n = 9 complexes. The bands in this latter frequency region are assigned to the vibrations of weakly tagged external CO ligands (9, 10). The appearance of the tagged CO band at n = 9 indicates that the n = 8 complexes are coordinatively saturated.

We carried out quantum chemical calculations using density functional theory and ab initio
methods to support the assignments of the vibrational spectra to the observed species and to examine the electronic structure of the carbonyl complexes. Figure 2A shows the optimized geometries of the neutral octacarbonyls calculated at the M06-2X-D3/def2-TZVPP level. The molecules have cubic (O₈h) symmetry and a triplet (3A₁g) electronic ground state with the valence electron configuration α⁸_s⁵_2t_12g_8_3g. Calculations of the singlet state (fig. S13) gave structures with D₆h (Ca and Sr) or D₄h symmetry (Ba), which were between 6.5 and 7.5 kcal/mol higher in energy than the corresponding triplet state species. Figure 2, B and C, shows the equilibrium geometries of the cations [M(CO)₈]+, which resemble the neutral species in the singlet state. Thus, [Ca(CO)₈]+ and [Sr(CO)₈]+ possess a D₆h structure and a 2A₁ electronic ground state, whereas [Ba(CO)₈]+ has D₄h symmetry and a 2B₂g electronic state. Figure 2 also shows the calculated zero-point energy (ZPE) corrected bond dissociation energies (D₀) of the octacarbonyls for loss of one and eight CO ligands. At the M06-2X-D3/def2-TZVPP level, the D₀ values for the dissociation of one CO lie between 9.1 kcal/mol for Ca and 11.5 kcal/mol for Sr for the M(CO)₈ neutral complexes and between 8.4 kcal/mol for Ba and 9.6 kcal/mol for Sr for the [M(CO)₈]+ cation complexes. The theoretical D₀ values for loss of eight CO ligands yielding M/M+ in the electronic ground state were between 58.8 kcal/mol for Sr and 63.3 kcal/mol for Ca for the neutrals and between 73.5 kcal/mol for Ba and 87.6 kcal/mol for Ca for the cations. The calculated values at the CCSD(T)/def2-TZVPP level, the values in parentheses, are the corresponding energies for the loss of eight COs.

The theoretical wave numbers for the C=O stretching modes of M(CO)₈ and [M(CO)₈]+ are shown in Table 1 along with the experimental values. The calculated values refer to the harmonic antisymmetric stretching frequencies scaled by a factor of 0.941, which comes from the ratio of the calculated stretching mode of free CO (2278 cm⁻¹) to the experimental value (2143 cm⁻¹) (JH). The calculated harmonic wave numbers are slightly higher than the experimental anharmonic values, but the trends for the different metals and for the isotopes of the neutral systems are in excellent agreement with the recorded values. The calculations suggest only one infrared (IR) active mode for the neutral complexes M(CO)₈ and two closely lying IR active modes for the cations [M(CO)₈]+. The latter splitting is too small to be experimentally observed. Theory and experiment indicate a considerable red shift of the C=O frequency for the neutral systems and a much smaller red shift for the cations. The calculated red shifts and the isotopic variations of the calculated and experimental wave numbers match each other.

Figure 3 shows the splitting of the valence orbitals of atoms with a spd valence shell in an octacoordinate cubic field with O₈ symmetry (I2). The breakdown of the AO into irreducible representations of the O₈ point group in an eight-coordinate ligand field is similar to the splitting in a six-coordinate octahedral field (I₃, I₄), but there are two important differences. One concerns the splitting of the (n − 1)d₄ AOs of the metal. The degenerate eₗ AOs in the octacoordinate cubic field are the (n − 1)d₄ AOs, and the triply degenerate t₂g AOs are the (n − 1)d₅ AOs. By contrast, in the six-coordinate octahedral field, the degenerate eₗ AOs are the (n − 1)d₅ AOs, and the triply degenerate t₂g AOs have (n − 1)d₄ character. The second difference concerns the appearance of the a₂u molecular orbital (MO) in the cubic field (fig. 3), which is absent in the octahedral field. The a₂u MO is a ligand-only orbital, because there is no valence AO of the spd shell that possesses this symmetry. This has consequences for the number of electrons that are required to fulfill the 18-electron rule. Because two valence electrons of the ligands in a cubic field are not available for donation to the central metal atom M in ML₈, where L is a ligand, the complex must provide a total of 20 electrons to fully occupy the metal’s valence shell. This scenario has recently been explored in the transition series.

Fig. 1. IR spectra of alkaline earth carbonyl complexes. (A) IR absorption spectra of calcium-carbonyl complexes in the 2150 to 1850 cm⁻¹ region from co-deposition of laser-evaporated calcium atoms with 0.1% CO in neon. Spectral lines: (i) after 30 min of sample deposition at 4 K, (ii) after a 12-K annealing period, (iii) after a 13-K annealing period, (iv) after 15 min of visible light irradiation, and (v) after another 12-K annealing period. (B) IR photodissociation spectra of the Sr(CO)₈⁺ (n = 6 to 9) complexes in the 2300 to 1800 cm⁻¹ region.

Fig. 2. Calculated equilibrium geometries of alkaline earth octacarbonyls. (A) M(CO)₈ (M = Ca, Sr, or Ba), (B) [M(CO)₈]⁺ (M = Ca or Sr), and (C) [Ba(CO)₈]⁺. Bond lengths are in angstroms. The D₀ values in roman type are the ZPE-corrected bond dissociation energies for loss of one CO ligand; the italicized values are the corresponding energies for the loss of eight CO ligands and M/M⁺ in the ground state. The values without parentheses are from M06-2X-D3/def2-TZVPP calculations; the values in parentheses are from CCSD(T)/def2-TZVPP using M06-2X-D3/def2-TZVPP optimized geometries.
metal octacarbonyl anions [TM(CO)₆]⁺ (TM = Sc, Y, or La) (15). Only 18 electrons are available in M(CO)₆ (M = Ca, Sr, or Ba), so the degenerate e₂ MO is occupied by two electrons with the same spin, giving a triplet (3A₂g) electronic ground state. Because the e₂ correlates with the (n – 1)dⁿ electron configuration of Ca, Sr, or La (19), the degenerate MO is occupied by two electrons with the same spin, giving a triplet electronic configuration. The metal center has a zero formal oxidation state.

We analyzed the nature of the metal-CO bonds with the energy decomposition analysis—natural orbitals for chemical valence (EDA-NOCV) (16) method, a powerful tool that provides detailed insight into chemical bonding (17). A description is given in the methods section. Table 2 shows the calculated results for the interactions between the metal atom M in the triplet electronic reference state with a ns⁶(n – 1)d² electron configuration and the (CO)₆ cage at the frozen geometry of M(CO)₆. The interaction energies ΔEₑₓₐₜ suggest that the intrinsic attraction M-(CO)₆ is strong and varies in the order Ca > Sr >> Ba. The dominant contribution to the total interaction ΔEₑₓₐₜ comes from the orbital term ΔEₒₐₜ. The preparation energies (ΔEₑₓᵣₑₓ) involved in the formation of the (CO)₆ cage from free CO molecules are low, ranging from 3.3 kcal/mol for Ba to 13.9 kcal/mol for Ca, whereas the electronic excitation energies for the atoms into the spherical symmetric ns² → ns²(n – 1)d² triplet state of M are quite high, lying between 68.2 kcal/mol for Ba and 159.5 kcal/mol for Ca. The strongly stabilizing interaction energies ΔEₑₓₐₜ caused by the attraction between the CO ligands and electronically excited M overcompensate the ΔEₑₓᵣₑₓ values. The adiabatic interaction energy ΔE (≡Dₑₓₐₜ) where Dₑₓ is the dissociation energy without zero-point vibrational energy correction) with respect to the electronic ground state of M and eight CO is between −65.5 kcal/mol for Sr and −73.7 kcal/mol for Ba. The Dₑₓ values in Table 2 exhibit a similar trend as the Dₑₓ data in Fig. 2.

The most important insight from the EDA-NOCV calculations is the breakdown of ΔEₑₓₐₜ into pairwise orbital interactions. Table 2 shows that the metal-CO bonding comes mainly from the (M(dₓ²) → (CO)₆ π backdonation of the degenerate (e₂) set of singly occupied (n – 1)d AO of the metal into the antibonding π* MOs of CO. This explains the large red shift of the C=O stretching mode of the octacarbonyls. The contribution of the [M(n)] ← (CO)₆ σ donation into x, where x denotes the valence acceptor AO of M, is much smaller than the [M(dₓ²) → (CO)₆ π backdonation. The order of the acceptor AOs of the metal atoms for the interaction energy is (n – 1)dₓ² > ns > np. There is also a small stabilizing contribution of the aₙ MO, which is due to the polarization of the CO ligand orbitals.

The dominating orbital interactions by the valence d electrons of M can be explained with the energetically very-high-lying occupied orbitals, which make M atoms excellent donor species. In our previous study of Ba(CO)₆ (19), we found that the cation Ba⁺(5d¹) is a donor to neutral CO; the atomic partial charge of Ba in Ba(CO)₆ was calculated as +1.39 e. The experimental values for the energetically lowest-lying σ* orbital in the electron configuration ns²(n – 1)d² are not far from the ionization limit (18). The valence d electrons of the M atoms are only weakly bonded to the atoms.

Figure 3 also shows the occupied valence MOs of Ca(CO)₆ that are relevant for the metal-CO bonding. The shape of the five (19) MOs reveals that the contributions of the metal AOs are very small, except in the SOMO, which clearly exhibits the shape of the (n – 1)dₓ² AOs. The valence MOs of Ca(CO)₆ are shown in Fig. 3.

### Table 1. Calculated and experimental wave numbers

<table>
<thead>
<tr>
<th>Complex</th>
<th>CO</th>
<th>Δ*</th>
<th>¹³CO</th>
<th>Δ†</th>
<th>C¹⁸O</th>
<th>Δ†</th>
<th>CO</th>
<th>Δ*</th>
<th>¹³CO</th>
<th>Δ†</th>
<th>C¹⁸O</th>
<th>Δ†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ca(CO)₆</td>
<td>2018</td>
<td>−125</td>
<td>1974</td>
<td>−44</td>
<td>1969</td>
<td>−49</td>
<td>1987</td>
<td>−156</td>
<td>1944</td>
<td>−43</td>
<td>1939</td>
<td>−48</td>
</tr>
<tr>
<td>[Ca(CO)₆]⁺</td>
<td>2116</td>
<td>−27</td>
<td>2094</td>
<td>−49</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>[Sr(CO)₆]⁺</td>
<td>2113</td>
<td>−30</td>
<td>2096</td>
<td>−47</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>[Ba(CO)₆]⁺</td>
<td>2127</td>
<td>−16</td>
<td>2113</td>
<td>−30</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Frequency shift relative to free CO. †Isotopic frequency shift.

Wu et al., Science 361, 912–916 (2018) 31 August 2018
Table 2. EDA-NOCV results. EDA-NOCV results for triplet state M(CO)₈ (M = Ca, Sr, or Ba) complexes at the M06-2X/TZ2P-ZORA level using M06-2X-D3/def2-TZVP optimized geometries, taking (CO)₈ in singlet ground state and M in triplet excited state with a nsⁿ(n – 1)δ² valence electronic configuration as interacting fragments. Energy values are given in kcal/mol.

<table>
<thead>
<tr>
<th>Energy term</th>
<th>Assignment</th>
<th>Ca + (CO)₈</th>
<th>Sr + (CO)₈</th>
<th>Ba + (CO)₈</th>
</tr>
</thead>
<tbody>
<tr>
<td>ΔE_total</td>
<td>(M(d)) → (CO)₈ π backdonation</td>
<td>-206.2 (86.2%)</td>
<td>-206.4 (86.2%)</td>
<td>-95.0 (69.8%)</td>
</tr>
<tr>
<td>ΔE_total‡</td>
<td>(M(d)) → (CO)₈ σ donation</td>
<td>-21.3 (9.0%)</td>
<td>-20.7 (8.7%)</td>
<td>-22.8 (16.8%)</td>
</tr>
<tr>
<td>ΔE_total†</td>
<td>(M(s)) → (CO)₈ σ donation</td>
<td>-2.4 (1.0%)</td>
<td>-2.9 (1.2%)</td>
<td>-3.2 (2.4%)</td>
</tr>
<tr>
<td>ΔE_total‡</td>
<td>(M(p)) → (CO)₈ σ donation</td>
<td>-0.9 (0.3%)</td>
<td>-0.9 (0.3%)</td>
<td>-2.7 (2.1%)</td>
</tr>
<tr>
<td>ΔE_total†</td>
<td>(CO)₈ polarization</td>
<td>-0.6 (0.3%)</td>
<td>-1.0 (0.4%)</td>
<td>-2.3 (1.7%)</td>
</tr>
<tr>
<td>ΔE_total‖</td>
<td>8 CO → (CO)₈</td>
<td>13.9</td>
<td>7.7</td>
<td>3.3</td>
</tr>
<tr>
<td>ΔE_total‖</td>
<td>M, nsⁿ → nsⁿ(n – 1)δ² (T)</td>
<td>159.5</td>
<td>150.9</td>
<td>68.2</td>
</tr>
<tr>
<td>ΔE (ΔE_total + ΔE_elstat)</td>
<td>= –D₀</td>
<td>M(CO)₈ → M(S) + 8 CO</td>
<td>-65.5</td>
<td>-73.7</td>
</tr>
</tbody>
</table>

*Contribution of the metahybrid term in M06-2X. ΔE_total is the electrostatic interaction energy. †The values in parentheses show the contribution to the total attractive interactions ΔE_total plus ΔE_elstat, where ‡The values in parentheses show the contribution to the total orbital interaction, ΔE_orb. §The sum of the two (e₈) or three (t₈ or t₆) components is given. ‖The EDA calculations give a triplet state with spherically symmetrical distribution of the d electrons. The experimental values for excitation into the energetically lowest-lying 2F state with nsⁿ(n – 1)δ² configuration are 124.2 and 59.8 kcal/mol for Ca and Ba, respectively. There is no experimental value for the relevant 2F state of Sr. The data are taken from (18).

REFERENCES AND NOTES
8. Geometry optimizations of M(CO)₈ using density functional theory and ab initio methods gave only weakly bonded van der Waals complexes with long M-CO distances.
19. The doubly or triply degenerate orbitals SOMO (e₈), HOMO-1 (t₈), and HOMO-2 (t₆) are single MOs, which have two and three components, respectively.

ACKNOWLEDGMENTS
Funding: The authors gratefully acknowledge financial support from the National Natural Science Foundation of China (grant nos. 21888102, 21433005, and 21703059), Nanjing Tech University and SICAM Fellowship from Jiangsu National Synergetic Innovation Center for Advanced Materials (L.Z. and G.F., grant nos. 39837132 and 39837133), and the National Science Foundation of Jiangsu Province for Youth (grant no. BK20170794). S.P. thanks Nanjing Tech University for a postdoctoral fellowship and the High Performance Computing Center of Nanjing Tech University for supporting the computational resources. Author contributions: X.W. did the matrix isolation experiments; J.J., W.L., X.J., and G.W. did the gas-phase infrared photodissociation experiments. M.Z. contributed to conception and design of experiments.
data analysis and interpretation; and drafting and critical revision of the manuscript. S.P. carried out the calculations and contributed to data collection and analysis of the results. L.Z. and G.F. contributed to conception and analysis of the theoretical results and drafting and critical revision of the manuscript. Competing interests: The authors have no competing interests. Data and materials availability: All data for this work are provided in the manuscript and in the supplementary materials.

SUPPLEMENTARY MATERIALS
www.sciencemag.org/content/361/6405/912/suppl/DC1
Materials and Methods

Figs. S1 to S19
Tables S1 to S3
References (22–40)
Data S1
6 May 2018; accepted 13 July 2018
10.1126/science.aau0839
Observation of alkaline earth complexes M(CO)$_8$ (M = Ca, Sr, or Ba) that mimic transition metals

Xuan Wu, Lili Zhao, Jiaye Jin, Sudip Pan, Wei Li, Xiaoyang Jin, Guanjun Wang, Mingfei Zhou and Gernot Frenking

*Science* **361** (6405), 912-916.
DOI: 10.1126/science.aau0839

**Carbonyls in the s block**

Conventional wisdom in chemistry distinguishes transition metals from other elements by their use of d orbitals in bonding. Wu *et al.* now report that alkaline earth metals can slide their electrons from s- to d-orbital bonding motifs as well (see the Perspective by Armentrout). Calcium, strontium, and barium all form coordination complexes with a cubic arrangement of eight carbonyl ligands and an 18-electron valence shell. The compounds were characterized in frozen neon matrices by vibrational spectroscopy and in gas phase by mass spectrometry. *Science*, this issue p. 912; see also p. 849.