Dual optical cycling centers mounted on an organic scaffold: New insights from quantum chemistry calculations and symmetry analysis

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Molecules cooled to the ultracold temperatures are desirable for applications in fundamental physics and quantum information science. However, cooling polyatomic molecules with more than six atoms has not yet been achieved. Building on the idea of an optical cycling center (OCC), a moiety supporting a set of localized and isolated electronic states within a polyatomic molecule, molecules with two OCCs (bi-OCCs) may afford a better cooling efficiency by doubling the photon scattering rate. By using quantum chemistry calculations, we assess the extent of the coupling of the two OCCs with each other and with the molecular scaffold. We show that promising coolable bi-OCC molecules can be proposed following chemical design principles.

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Ultracold molecules are attractive for novel applications^{[1–](#page-14-0)[3](#page-14-1)}. Molecules can be cooled to the ultracold temperatures by laser cooling, a repeated absorption and emission of photons exploiting the Doppler effect^{[4](#page-14-2)}. When excitation laser is tuned to energy slightly below the electronic transition, only molecules moving towards the photon source can be excited. The excited molecule then undergoes spontaneous decay by which it emits recoil momentum in all directions, slowing the average momentum over time. The key to laser-cooling molecules is the ability to prevent decay to unwanted decay channels, such as excited vibrational states. The process of scattering many photons using the same closed loop is called optical cycling. The undesirable population leakage into vibrationally excited states is minimized when ground and electronically excited states have the same structures, thereby resulting in the diagonal Franck-Condon factors (FCFs; vibrational overlaps). This goal can be achieved by designing molecules with localized atomic-like transitions, as was first described by Isaev and Berger^{[5](#page-14-3)}, who also proposed several promising candidates—all based on a metal-oxygenligand (M-O-L) framework with a second-group metals. The essential features of these systems are a strongly ionic M-O bond and a localized unpaired electron on the metal, giving rise to localized atomic-like transitions. These transitions can be used as the main photon cycling loop; hence, the name optical cycling center (OCC).

Several molecules featuring such electronic structure have been successfully laser-cooled^{6-[8](#page-14-5)}. Cooling larger molecules could be advantageous for quantum information and sensing ap-plications^{[9](#page-14-6)[–11](#page-14-7)}. Although initially the enthusiasm was dampened by the concerns that the increased complexity is likely to result in more vibrational branching channels, several theoretical studies have shown that such metal-based OCCs can in fact operate in larger molecules. Following this strategy, more polyatomic molecules were proposed computation-ally as prospective candidates for laser cooling^{[12–](#page-14-8)[21](#page-15-0)}. The screening of the candidate molecules entails finding a low-lying bright electronic transition, assessing that it is electronically uncoupled from the rest of the molecule, and computing FCFs (vibrational overlaps). The electronic transitions and FCFs can be tuned using chemical intuition. As recent studies show, strongly diagonal FCFs are possible in large molecules, leading to vibrational branching ratios rivaled that of CaOH^{17-[19](#page-15-2)}.

It is desirable to further optimize photon cycling in the candidate molecules, such that one can use fewer repump lasers, introduce additional optical controls, and explore more varied types of molecules. One way to increase cycling efficiency is to install multiple OCCs

on one scaffold, which can increase the oscillator strength, potentially doubling the photon scatters^{[13,](#page-14-9)[15](#page-15-3)[,22,](#page-15-4)[23](#page-15-5)}. Here, we explore this idea further, by considering a larger set of molecular scaffolds augmented by two OCCs (we refer to them as bi-OCCs). We analyze the effect of symmetry and use chemical design to control the optical properties of the prospective bi-OCCs and the respective FCFs (FCFs control vibrational branching ratios in electronic transitions). α ansitions).

in bi-OCCs. Configuration (i) is the high-spin $(M_s=1)$ triplet state and configurations (ii)-(v) are low-spin $(M_s=1)$ states. Case 1 corresponds to perfectly delocalized frontier orbitals (such as bonding and antibonding combinations of atomic orbitals) and Case 2 corresponds to the molecular orbitals perfectly localized on the radical centers. FIG. 1. Diradicals: Different electronic configurations derived from distributing two electrons in two orbitals and their characters. M_1 and M_2 denote two radical centers, e.g., the two metal atoms external magnetic field, and thus, where α FIG. 1. Diradicals: Different electronic configurations derived from distributing two electrons in

Although bi- \overline{OCC} molecules may seem exotic, many features of their electronic structure can be understood by revisiting diradicals $\frac{1}{\sqrt{2}}$ and fCu2 denote orbitals localized or by distributing two electrons in two orbitals. Fig. [1](#page-2-0) shows resulting electronic configuraincluding small contributions from the nearest ligand atoms). tions. Ine character of the wave-functions lized and can be described as (nearly degenerate) bonding and and to complete. The displacement of case α is α trons localized on the two metal centers whereas the singlets can be either covalent, ionic $(M_1^+M_2^- \pm M_2^-M_1^+)$, or a mixture of covalent and ionic (charge-resonance) contributions. state can vary, depending on the exact weights of configuration $\mathcal{L}_\mathbf{z}$ tions (case 1 in F ϵ or equal orthonormal configuration with two units covalent an adical comiguration with two and ture of ionic and covalent configurations when $\lambda=0$. The extent of diradical character of the between the two centers stabilizes the bonding MO relative to wave-function can be quantified by computing the number of effectively unpaired electrons, Although bi-OCC molecules may seem exotic, many features of their electronic structure ϵ Directicals are species in which the two from Diradicals are species in which the two holitier molecular orbitals are (nearly)-degenerate and the low-lying electronic states derived methods as singlet–triplet energy gaps. This approach can be multiple polyradical centers.18,19 ends whether the orbitals are delocalized m_{max} they come and to the unperiod close when $\sum_{i=1}^{\infty}$ increases the experience of $\sum_{i=1}^{\infty}$ Slater determinant and, therefore, their energies can be reliably cluster theory or DFT. In contrast, wave functions of low-spin Considering delocalized orbitals (case 1 in Fig. [1\)](#page-2-0), configuration (iv) corresponds to purely $\frac{1}{2}$ are positive are poorly described by single-reference methods. covalent diradical configuration with two unpaired electrons when $\lambda=1$ and to an equal mix- \mathcal{L} and \mathcal{L} is singlet and triplet states. In the state states states and \mathcal{L} In these expressions, fCu2 denotes $\mathcal{L}_{\mathcal{A}}$ and fCu2 denotes orbitals localized or lized and can be described as (nearly degenerate) bonding and and the combination of the wave-functions of α is the MMs studied function of \mathbb{R}^n in the MMs studied here, the triplet states are \mathbb{R}^n of focalized. The diplor states are always state can variety on the exact we increase with α $\binom{1}{1}$ $\binom{1}{2}$ \cong $\binom{1}{2}$ $\binom{1}{1}$, or we have to covar into the wave function. This happens when the interaction. This happens when the interaction \mathcal{L}_1 the singlet wave functions due to mixing with configuration wave-function can be quantified by computi can be understood by revisiting diradicals.^{[24–](#page-15-6)[28](#page-16-0)} Diradicals are species in which the two fronmultiple polyradical centers.18,19 multiconfigurational character of the low-spin wave functions, tions. The character of the wave-functions depends whether the orbitals are delocalized s_{short} they correspond to the unperiod close or localized. The triplet states are always covalent—they correspond to the unpaired elecrose the singlets can be either covelent ienig trons localized on the two metal centers whereas the singlets can be either covalent, ionic t and jonic (charge-resonance) contributions. $(M_1^+M_2^- \pm M_2^-M_1^+)$, or a mixture of covalent and ionic (charge-resonance) contributions. states results in the computed singlet \mathcal{L} the most popular are the broken symmetry (BS) methods20–22 wave-function can be quantified by computing the number of effectively unpaired electrons,

 $n_{u,nl}$, using Head-Gordon's formula^{[29](#page-16-1)}. In the triplet state (configurations (i) and (ii)), the number of unpaired electrons is 2. For configuration (iv) and delocalized orbitals (case 1 in Fig. [1\)](#page-2-0), the number of unpaired electrons can be computed from the parameter λ as follows^{[28](#page-16-0)}:

$$
n_{u,nl} = \frac{32\lambda^4}{(1+\lambda^2)^4}.\tag{1}
$$

Energy gap between the frontier orbitals controls parameter λ and the relative energies of the lowest singlet and triplet states—lager gaps favor ground singlet states. If the energy separation between ϕ_1 and ϕ_2 states is very small, the triplet can become the ground state. The gap depends on the through-space and through bond interactions. Through-space interaction depends on the overlap of the contributing atomic orbitals whereas throughbong interaction depends on the molecular scaffold and the relative placements of the radical centers. This may lead to flipping the ground-state multiplicity in structural isomers—for example, meta-xylylenes have triplet ground state whereas para-xylylene have singlet ground state.

Previous studies of bi-OCCs considered the following motifs—M1-CC-M2, with M1/M2=Mg,Ca,Sr (Ref. [13\)](#page-14-9) and $M1=Yb$ and $M2-Ca/Al$ (Ref. [22\)](#page-15-4), benzenes functionalized by Ca in ortho-, meta-, and para- positions^{[15](#page-15-3)}, as well as Sr and OSr installed on $(CH_2)_n$ chains and fullerenes (Ref. [23\)](#page-15-5). These studies revealed the following essential features of bi-OCCs:

- Even in the systems with a short acetylene linker, the frontier orbitals are localized on metal centers and the low-lying electronic states are largerly uncoupled from the scaffold; this can be explained by the large energy gap between the metal and linker orbitals and the diffuse nature of the former.
- Both through-space and through-bond interactions between the two centers are weak, leading to nearly or almost degenerate singlet and triplet states.
- Because of the weak interaction between the OCCs, the characters of the singlet and triplet states are similar—both are covalent (i.e., $n_{u,nl} \geq 1.8$ for the lowest singlet states). Furthermore, the low-lying excited states in each multiplicity follow nearly identical patterns. The energy spacing follows that of a single OCC very closely, with very small excitonic splitting.
- In bi-OCCs with the short acetylenic linker, the coupling between the centers and the

linker and between the centers themselves is smaller for heavier metals for which the two manifolds become even more similar. However, in phenoxide with a single OCC, Sr resulted in worse FCFs than Ca.^{[30](#page-16-2)}

In high-symmetry cases, excited states show excitonic patterns, with one state dark and one carrying double oscillator strength of a single OCC. Depending on the structure, the lowest excited state can be either bright (desirable for cycling) or dark (undesirable). This can be understood in the framework of the exciton model^{[31,](#page-16-3)[32](#page-16-4)}—if the transition moments of the two chromophores are parallel but not aligned, one obtains an H-dimer in which the lowest state is dark. If the two moments are aligned along the same axis, one obtains a J-dimer in which the lowest state is bright (super-radiant).

The lowest transitions in the metal-based OCCs are of $s \to p$ types, with the *p*-orbital aligned along the MO bond being hybridized with the respective d-orbital. The transition moments are aligned along the direction of the corresponding p-orbitals. Hence, the excitonic pattern can be controlled by splitting the degeneracy of the p-orbitals and their relative orientation in a molecule. Specifically, in p-benzyne bi-OCC, the lowest transition is dark and in m- and o-benzynes it is bright (but not super-radiant).

Here we use M-O-X framework, in which the metal centers are even further apart, thus improving the decoupling relative to the M-X motifs. Fig. [2](#page-4-0) shows the bi-OCC molecules studied in this work.

FIG. 2. Structures of bi-OCC molecules considered in this work.

The type of electronic structure featured by OCCs can be accurately described by the double electron attachment variant^{[33](#page-16-5)} of equation-of-motion coupled-cluster methods^{[34](#page-16-6)} in which the ground and excited target states are described by attaching two electrons to a dicationic reference, as was done in Refs. [12](#page-14-8)[,13.](#page-14-9) Here we use EOM-DEA-CCSD with the cc-pVDZ basis on H,C,O and aug-cc-pwCVDZ-PP on Ca and Sr with the ECP10MDF and ECP28MDF pseudopotentials used for the respective metal atoms. EOM-DEA-CCSD calculations were carried out using Q-Chem^{[35](#page-16-7)}.

For extensive screening involving calculations of FCFs for relatively large molecules it is desirable to develop a less expensive protocol. Here we take advantage of the fact that lowest singlet and triplet manifolds of bi-OCCs have very similar character. Hence, one can use triplet manifold as a proxy for singlet states. Whereas low-spin singlet and triplet states are multi-configurational, the high-spin triplet states can be well described by a single-reference method (HF, CCSD, DFT), which is exploited, for example, in spin-flip approach $36,37$ $36,37$. Following the same logic, here we compute the triplet-state manifold from the lowest high-spin triplet state using density functional theory (DFT) and its extension to the excited states, TD-DFT. We use the PBE0-D3/def2-TZVPPD level of theory for geometry optimizations, excitations, frequencies, and FCFs, as implemented in Gaussian 16^{38-41} 16^{38-41} 16^{38-41} . The validate this approach, we benchmarked it against EOM-DEA-CCSD; the results are given in the SI.

We begin by discussing benzenes functionalized by two Ca-O OCCs installed in meta- $((1))$ and para- $((2))$ positions (see Fig. [2\)](#page-4-0). This builds upon previous work where we experimentally realized the single OCC case, $CaOP¹⁷$ $CaOP¹⁷$ $CaOP¹⁷$.

The natural orbitals hosting the unpaired electrons are in-phase and out-of-phase combinations of the s- and p-orbitals of the to Ca atoms (Fig. [3\)](#page-6-0). The ground state is derived by distributing two electrons in the molecular orbitals derived by linear combinations of the two s-orbitals. The lowest triplet states are ${}^{3}B_{1}$ in (1) and ${}^{3}B_{3u}$ in (2); they appear to be degenerate with the lowest singlet states $({}^{1}A_{1}$ and ${}^{1}A_{g})$ at the EOM-DEA-CCSD level of theory. The wave-function analysis yields $n_{u,nl}=2$ for the singlet states, confirming that the singlet state has pure diradical character. We note that for the analogous Ca-substituted benzenes^{[15](#page-15-3)}, $n_{u,nl}$ values were smaller—1.83, 1.92, and 2.00 for ortho-, meta-, and paraisomers, respectively. Hence, linking metals through the oxygen improves the decoupling of the unpaired electrons and reduces ionic contributions, which should improve the FCFs. This effective decoupling also makes singlets electronically very similar to the triplets. Hence, we expect that the cycling efficiency should not be affected by the transitions between the two spin manifolds, which can be induced by spin–orbit coupling.

Excited-state properties of (1) and (2) can be rationalized using symmetry analysis.

FIG. 3. Frontier orbitals and their symmetries for the two simplest bi-OCC molecules — Ca-O-Ph-3-O-Ca with C_{2v} point group (1) and Ca-O-Ph-4-O-Ca with D_{2h} point group (2). Note: Symmetry labels correspond to Q-Chem's standard molecular orientation^{[42](#page-18-1)}. EOM-DEA-CCSD/ccpVDZ[H,C,O]/aug-cc-pwCVDZ-PP[Ca]/ECP10MDF[Ca]; isovalue 0.03.

The OCC transitions are derived by promoting electrons to the molecular orbitals derived from a pair of p-orbitals. The degeneracy between the atomic p-orbitals is lifted such that the lowest p-orbitals are in-plane, perpendicular to the Ca-O bond, followed by the out-ofplane orbitals, and p-orbitals parallel to the Ca-O bond (these highest pair of orbitals is not shown in Fig. [3\)](#page-6-0). Using point-group symmetry, we can then determine the number of bright transitions for a given OCC molecule. For example, in (1) (C_{2v}) combinations of s-orbitals have a_1 and b_1 symmetries, and combinations of p-orbitals have a_1 , b_1 , b_2 , and a_2 symmetries. One of the in-plane excited states is derived by $a_1 \rightarrow a_1$ and $b_1 \rightarrow b_1$ transitions, yielding bright transition of A₁ symmetry. Transition $a_1 \rightarrow b_1$ (of B₁ symmetry) is also optically allowed. Similarly, two out-of-plane transitions have B_1 and A_2 symmetries, rendering the latter one dark. We remind that for the transitions from ${}^{1}A_{1}$, the symmetry of the target state is the same as the symmetry of the transition whereas for the transitions from the open-shell triplet state, the symmetry of the target state is the product of the symmetry of the transition and the initial triplet state $({}^{3}B_{1}$ in (2)). Following the same analysis, in (2), (D_{2h}) , we anticipate two bright transitions—B_{1u} and B_{2u}. Unfortunately, as anticipated from the orbital pictures, the lowest states correspond to H-aggregate type, rendering the lowest excited states dim in (1) and optically forbidden in (2).

Table [I](#page-7-0) shows the results for (1) and (2) for the lowest transitions in the triplet manifold and the respective oscillator strength. As anticipated from our analysis, (1) has three bright transitions with symmetries B_1 , A_1 , and B_2 (corresponding to 3A_1 , 3B_1 , and 3A_2 states). The lowest state is optically allowed, but not very bright $(f_l=0.126)$. The respective FCF, ${}^{3}A_{1}(\nu = 0) \rightarrow {}^{3}B_{1}(\nu = 0)$, is almost twice smaller than that of the fundamental transitions from the ${}^{3}B_1$ and ${}^{3}A_2$ states (0.5051 versus 0.9176 and 0.9604). Previously, some of us investigated this effect and found that the diagonality of FCFs is improving proportionally to the distance between the optical centers.^{[43](#page-18-2)}

TABLE I. Properties of four lowest excited states^{*a*} of (1) and (2) in the triplet manifold; the lowest triplet state is ${}^{3}B_{1}$ and ${}^{3}B_{3u}$, respectively.

(1)	(2)			
$Ca-O-Ph-3-O-Ca$	$Ca-O-Ph-4-O-Ca$			
State E_{ex} , eV (f_l) FCF	State E_{ex} , eV (f_l) FCF			
$13A1$ 2.036 (0.126) 0.5051	1^3B_{2u} 2.044 (-)			
1^3B_2 2.058 (-)	1^3B_{1u} 2.060 (-)			
2^3B_1 2.085 (0.401) 0.9176	$13B1q$ 2.066 (0.518) 0.9663			
	1^3A_2 2.097 (0.543) 0.9604 1^3B_{2g} 2.085 (0.544) 0.9792			
^{<i>a</i>} Excitation energies and oscillator strengths:				

EOM-DEA-CCSD/cc-pVDZ[H,C,O]/aug-cc-pwCVDZ-PP[Ca]/ECP10MDF[Ca]; FCFs: PBE0-D3/def2-TZVPPD.

(2) has two highly diagonal bright states of symmetries B_{1g} and B_{2g} with FCFs 0.9663 and 0.9792, respectively. The oscillator strengths of (2) 's bright transitions are roughly twice as large as the oscillator strength of the single-OCC CaOPh molecule (0.518 and 0.544 in bi-OCC versus 0.2075 and 0.2155 in single-OCC) and two subradiant states^{[44](#page-18-3)}, following the same pattern as in $Ca-X$ species^{[13](#page-14-9)}. Unfortunately, the lowest transition is dark because of the H-dimer arrangement. In order to obtain superradiant lowest states in these two model systems, the energies of the two highest molecular orbitals, which are derived from the hybridized p and d orbitals parallel to the Ca-O bond, have to be brought below the other orbitals in the p-manifold, which might be possible to achieve by using bulky substituents that destabilize the other four orbitals. However, given the small size of the benzene scaffold, this route is impractical as such substituents would deteriorate FCFs.

Overall, bi-OCC molecules are good candidates for laser cooling purposes because of good FCFs and the flexibility they afford to tune excited states. Because the phenyl ring does not allow for complex functionalization—even a small substituents (such as a CH_3 group) interferes with the Ca-centered orbitals, reducing FCFs and increasing orbital mixing (see Supplementary Information)—we turn to the larger scaffolds, such as naphthalene (systems (3)-(11) in Fig. [2\)](#page-4-0). Naphthalene and its structural isomer azulene, were previously considered in its unfunctionalized version for buffer-gas cooling.[45](#page-18-4) Below we demonstrate how various structural modifications affect optical properties of bi-OCCs, allowing one to enhance or disentangle relevant electronic transitions.

$(3) 1^3B_u$			(8) 1 ³ A'			
2^3B_u	1.991 (-)		$2^3\mathrm{A}^{\prime}$		2.092(0.077)	0.9145
1^3A_g	2.005(0.518)	0.9668	$1^3A''$		2.099(0.013)	0.9603
1^3A_u	2.015 (-)		$3^3A'$		2.120(0.417)	0.9657
1^3B_g	2.030(0.550)	0.9815	$2^3A''$		2.123(0.513)	0.9831
(4) 1 ³ B ₁			(9) 1 ³ B _u			
1^3A_1	1.987(0.123)	${0.9264}$	1^3A_u	$2.123 (-)$		
1^3B_2	2.013 (-)		2^3B_u	$2.130 (-)$		
$2^{3}B_{1}$	2.013(0.400)	0.9646	1^3B_q		2.145(0.523)	0.9855
1^3A_2	2.035(0.550)	0.9796	1^3A_q		2.149(0.493)	0.8048
$(5) 1^{3}B_{u}$			(10) 1^3 A'			
2^3B_u	1.994 (-)		$2^3A'$		2.074 (0.0002) 0.9619	
1^3A_g	2.014(0.504)	0.9631	$3^3A'$		2.087 (0.507) 0.9721	
1^3A_u	2.020 (-)		1 ³ A''		2.089(0.015)	0.9793
1^3B_q	2.044 (0.540) 0.9817		$2^3A''$		2.105 (0.525) 0.9846	
(6) 1 ³ B ₁			(11) 1 ³ B _u			
1^3A_1	1.979 (0.0001) 0.9409		2^3B_u	2.090 (-)		
$2^{3}B_{1}$	2.006 (0.503) 0.9649		1^3A_u	$2.100 (-)$		
1^3B_2	$2.014 (-)$		1^3A_q		2.103(0.505)	0.951
1^3A_2	2.039 (0.540) 0.9354		1^3B_g		2.115(0.536)	0.970
(7) $1^3A'$						
$2^3A'$	1.984(0.143)	0.8583				
$1^3A''$	2.013(0.007)	0.9631				
$3^3A'$	2.015(0.381)	0.9547				
$2^3A''$	2.042(0.538) ^a Excitation energies and oscillator strengths:	0.9845				

TABLE II. Excitation energies (eV), oscillator strengths (in parentheses), and FCFs for the $0\rightarrow 0$ transition for different excited states in the triplet manifold for (3) - $(11)^{a}$. The symmetry of the lowest triplet state for each system is given next to the system number.

EOM-DEA-CCSD/cc-pVDZ[H,C,O,F]/aug-cc-pwCVDZ-PP[Ca]/ECP10MDF[Ca]; FCFs:

PBE0-D3/def2-TZVPPD.

Table [II](#page-9-0) presents the results for naphthalene-based bi-OCCs (see Fig. [2\)](#page-4-0). The naphthalene molecule has three types of symmetry non-equivalent carbon atoms where we can install an OCC, giving rise to the following five bi-OCC structures: four molecules with OCCs on equivalent carbon sites (systems (3) , (4) , (5) , and (6)) and one molecule with OCCs on non-equivalent sites (system (7)). In these systems, the lowest singlet and triplet are also degenerate (at the EOM-DEA-CCSD level), and the number of effectively unpaired electrons in the lowest singlet state is two.

With these arrangements, we observe different patterns of how the combined oscillator strength derived from the $s \to p$ transitions is distributed among low-lying manifold of excited states. In systems with high symmetry, the oscillator strength is concentrated in a smaller number of states whereas in low-symmetry cases it is distributed among more states so that in non-symmetric cases the number of optically allowed states is the largest. As for (1) and (2), symmetry considerations explain the results for these larger systems. Systems (3) and (5) have the C_{2h} symmetry and two bright states. (4) and (6) have the C_{2v} symmetry and three bright states whereas (7) has the C_s symmetry and four bright states.

For C_{2h} systems, both bright states are superradiant (the oscillator strength is on the order of 0.5, which is twice larger than that in single-OCC molecules), whereas for C_{2v} systems there are two super- and one sub-radiant states, with the latter having low oscillator strength $(0.123$ for (4) , and $(0.0001$ for (6)). For system (7) , both super- and sub- radiant states are bright. Here again the brighter states appear higher in energy, which is not suitable for optical cycling.

As was previously shown, electron-withdrawing groups with high Hammett parameters improve FCFs for single-OCC transitions for benzenes and naphthalenes^{[17,](#page-15-1)[46](#page-19-0)}. We extend this idea to their bi-OCC counterparts by functionalizing two of the five systems with electronwithdrawing CF_3 groups (systems (8)-(11)). We functionalized systems (3) and (5) because the substituents are positioned at a distance away from the OCCs such that they do not disrupt the OCC transitions. We then functionalized (3) and (5) by one or two substituents to produce systems (10) , (11) and (8) , (9) , respectively, to investigate stronger electronwithdrawing effects.

For singly CF_3 functionalized systems, (8) and (10), all four excited states are bright due to C_s symmetry point group, similar to system (7). The first excited A' and A'' states are subradiant (with oscillator strengths less than (0.1)). For (8) , the superradiant transitions (to the third and fourth excited states) have FCFs of 0.9657 and 0.9831, an improvement over the corresponding ground to excited state transitions of their unfunctionalized counterpart, (5) (0.9631 and 0.9817). Similarly, (10)'s superradiant transitions have FCFs of 0.9721 and 0.9846, an improvement over (3)'s (FCFs of 0.9668 and 0.9815).

Double functionalization with CF_3 (systems (9) and (11)) recovers the symmetry of the parent species and, consequently, their numbers of bright excited states. However, the FCFs for doubly functionalized species deteriorate slightly. This is because functionalization often introduces more vibrational degrees of freedom that electronic states can couple to, even while improving on the order of this coupling by utilizing electronic effects. For example, the first two decay channels for the B_u transition of (5) are decays to the stretching and bending modes with FCFs of 0.0132 and 0.0124, respectively. For the B_u transition of (9) the two main channels are also stretching and bending, but with FCFs of 0.0041 and 0.0035, respectively. The same characteristic applies to (11) and (3) . This means that we succeed in improving the diagonality of the transition electronically, however, in this particular case the introduced vibrational degrees of freedom quench that effect and ultimately result in worsening of FCF. This suggests to consider exploring heteroatomic motifs in future work e.g., introducing nitrogen or boron in the scaffold can significantly affect electronic states (as was observed in organic di- and tri-radicals^{$47-49$ $47-49$}), but without introducing more degrees of freedom.

state is $1^{\circ}A^{\circ}$ and $1^{\circ}A_1$, respectively.				
(12)	(13)			
$Ca-O-Ph-3-O-Sr$	$Ca-O-Ph-4-O-Sr$			
State E_{ex} , eV (f_l) FCF	State E_{ex} , eV (f_l) FCF			
2^3 A' 1.762 (0.261) 0.8122	$13B1$ 1.627 (0.274) 0.9057			
$13A''$ 1.784 (0.251) 0.9293	$13B2$ 1.656 (0.283) 0.9279			
3^3 A' 2.045 (0.378) 0.9256	2^3A_1 1.918 (0.278) 0.9325			
2^3 A" 2.066 (0.165) 0.9594 2^3B_1 2.071 (0.275) 0.9538				

TABLE III. Properties of four lowest excited states of (12) and (13) in the triplet manifold^a; the lowest triplet state is 1^3 A' and 1^3 A

^a Excitation energies and oscillator strengths: $\overline{EOM-DEA-CCSD/cc-pVD}Z[H,C,O]/aug-cc$ pwCVDZ-PP[Ca,Sr]/ECP10MDF[Ca]/ECP28MDF[Sr]; FCFs: PBE0-D3/def2-TZVPPD.

Additional flexibility in laser-coolable bi-OCCs is afforded by using different metals, 13,22 13,22 13,22 13,22 which makes the two centers distinguishable and independently addressable. Table [III](#page-11-0) and Figs. [4](#page-12-0) and [5](#page-13-0) show the results for the systems with one Ca and one Sr OCCs (systems (12) and (13) in Fig. [2\)](#page-4-0). As we can see from the natural orbitals, electronic transitions are localized, so that they can be addressed individually. This is different from the bimetallic bi-OCCs linked through the short acetylene bridge^{[13](#page-14-9)} where the transitions showed some degree of delocalization, consistent with stronger interactions between the two centers.

FIG. 4. Natural orbitals and their occupations for the lowest triplet states of a mixed bi-OCC CaO-Ph-3-OSr (12). EOM-DEA-CCSD/cc-pVDZ[H,C,O]/aug-cc-pwCVDZ-PP[Ca,Sr]/ECP10MDF[Ca]/ECP28MDF[Sr]; isovalue 0.03.

In this contribution, we investigated details of the electronic structure of bi-OCCs. We have shown that in bi-OCCs built upon benzene and naphthalene scaffolds by augmenting them with two OM groups $(M=Ca, Sr)$, the two optical centers are decoupled from each other and from the organic scaffold, giving rise to degenerate perfect diradical singlet and triplet ground states. In the future work, we will investigate whether this degeneracy is lifted when higher correlation treatment is employed and spin–orbit interaction is accounted for.

FIG. 5. Natural orbitals and their occupations for the lowest triplet states of a mixed bi-OCC CaO-Ph-3-OSr (13). EOM-DEA-CCSD/cc-pVDZ[H,C,O]/aug-cc-pwCVDZ-PP[Ca,Sr]/ECP10MDF[Ca]/ECP28MDF[Sr]; isovalue 0.03.

We expect that the multiplicity of the ground states in these systems is not important, as the lowest singlet and triplet states are very similar electronically. Moreover, the singlet and triplet manifolds of excited states are also very similar. Thus, we expect that the transitions between the two manifolds should not pose problems for cycling.

This study also shows that increasing complexity of the system provides several avenues for a chemical control over the electronic properties of OCC. Using chemical design principles, one can control the amount of the excited states in bi-OCC molecules and through symmetry and the degree of coupling between two metal centers. Since chemist's toolbox is essentially endless, many more handles and switches can be introduced into the system to control its electronic properties.

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