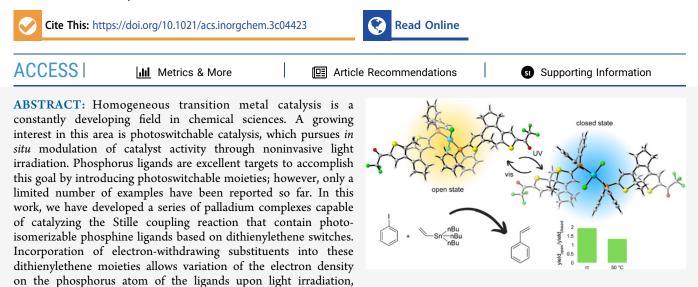
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# Dithienylethene-Based Photoswitchable Phosphines for the Palladium-Catalyzed Stille Coupling Reaction

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which in turn leads to a modulation of the catalytic properties of the formed complexes and their activity in a model Stille coupling reaction. These results are supported by theoretical computations, which show that the energy barriers for the rate-determining steps of the catalytic cycle decrease when the photoswitchable phosphine ligands are converted to their closed state.

# INTRODUCTION

Inspired by the dynamic behavior of enzymes in nature, controlling the operation of artificial molecular catalysts with external stimuli has become an active area of research with potential application in a variety of fields<sup>1–3</sup> (e.g., polymer synthesis,<sup>4,5</sup> bioorthogonal chemistry,<sup>6,7</sup> and additive manufacturing<sup>8</sup>). The use of light to achieve this goal holds great promise, as it opens the way to catalyst regulation on-demand with high selectivity and spatiotemporal precision.<sup>9–11</sup> A major strategy toward this purpose is the design of photoswitchable catalysts, whose activity and selectivity can be reversibly modulated with light by installing photochromic units into their structure, i.e., light-responsive moieties, such as azobenzenes,<sup>12,13</sup> dithienylethenes<sup>14</sup> and stilbenes,<sup>12</sup> which alter the reactivity of nearby catalytic sites by photoisomerizing between two different states.

Because of their fundamental role in modern synthetic chemistry, transition metal complexes are among the principal targets of photoswitchable catalysis.<sup>15–17</sup> Reversible light control of the catalytic activity of these compounds is generally accomplished by introducing photochromic ligands (e.g., photoswitchable phosphines<sup>18</sup>). In most of the cases, the geometrical changes that these ligands undergo upon photo-isomerization cause catalyst reactivity modulation,<sup>15–17</sup> for instance, by distorting the structure around the catalytic site<sup>19–21</sup> or varying the separation distance between two

cooperative active metal centers.<sup>22,23</sup> However, the actual impact of these effects on catalytic activity and selectivity can be detrimentally affected by catalyst conformational flexibility<sup>24</sup> and be dependent on substrate size and geometry. An alternative, less exploited approach to light-control the performance of transition metal catalysts is to capitalize on the electronic changes that occur upon ligand photoisomerization.<sup>15–17</sup> For this strategy, dithienylethenes (DTEs) are the photochromic units of choice<sup>14</sup> because, in contrast to azobenzenes and stilbenes, they undergo a large modification in electronic structure when reversibly toggling between their ring-open (**o**) and ring-closed (**c**) isomers.<sup>25</sup>

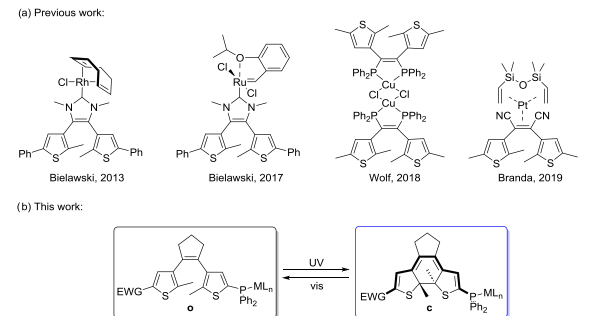
To date, only a small number of DTE-based complexes have been described where photomodulation of catalysis is accomplished via electronic effects.<sup>26–29</sup> All of these systems share a common design principle: they contain DTE ligands that bind to the metal center through the groups installed in their central ethylene bridging moiety (e.g., carbene,<sup>26,27</sup> phosphines<sup>28</sup>), which are either removed<sup>29</sup> or lose electron

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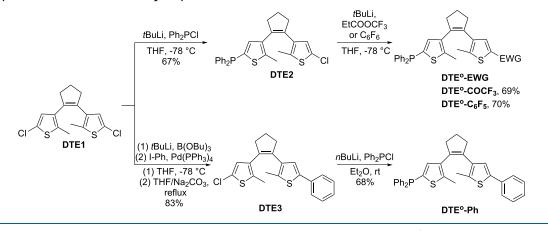


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# Scheme 1. (a) Previously Reported Transition Metal Complexes Incorporating DTE-Based Photoswitchable Units as Ligands and (b) DTE-Based Photoswitchable Complexes Studied in This Work



Scheme 2. Synthetic Route toward Phosphanyl-Substituted DTEs



density $^{26-28}$  upon ring-closing (Scheme 1a). In contrast, other electronic features that accompany DTE photoisomerization have yet to be exploited in metal-based photoswitchable catalysis. In this work, we proposed to explore one of these additional features: the variation in electronic communication between the external thiophene substituents upon photoconversion, which has already been utilized to control chemical reactivity with light-responsive organocatalysts<sup>30,31</sup> and organic substrates.<sup>32,33</sup> In particular, herein, we devised the synthesis of nonsymmetric DTE derivatives as photoswitchable ligands bearing (i) a metal-binding phosphine group at one thiophene ring and (ii) an electron-withdrawing substituent at the other (Scheme 1b). As these two groups must be electronically insulated in the open state of the system and become selectively conjugated upon ring-closing, our molecular design should allow modulating the electron density of the phosphine ligand with light and, eventually, the catalytic activity of metal centers upon coordination. To validate this hypothesis, we prepared palladium(II) complexes of our DTE ligands and tested them as precatalysts in a Stille coupling reaction, a widely employed Pd-catalyzed transformation that is sensitive

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to electronic effects  $^{34-36}$  and has not been tested yet in photoswitchable catalysis.

# RESULTS AND DISCUSSION

Synthesis of Photoswitchable Ligands and Monometallic Complexes. To explore our approach toward photoswitchable catalysts, two different electron-withdrawing groups (EWGs) were introduced in DTE-based phosphine ligands: (i) a trifluoroacetyl group in DTE-COCF<sub>3</sub>, which is a strong EWG according to its Hammett  $\sigma$ -meta and  $\sigma$ -para substituent constants ( $\sigma_m = 0.63$ ,  $\sigma_p = 0.80$ ),<sup>37</sup> and (ii) a pentafluorophenyl group in DTE-C<sub>6</sub>F<sub>5</sub>, which presents a less pronounced electron-withdrawing power ( $\sigma_m = 0.26$ ,  $\sigma_p =$ 0.27).<sup>37</sup> As a reference, we also considered the preparation of a DTE-based phosphine ligand bearing a phenyl substituent (DTE-Ph), which should impart very minor electronic effects ( $\sigma_m = 0.06$ ,  $\sigma_p = -0.01$ ).<sup>37</sup>

The ring-open isomer of the DTE-based ligands was synthesized through asymmetrical stepwise functionalization of 1,2-bis(2-chloro-5-methylthien-4-yl)cyclopentene (DTE1, Scheme 2), which is a common precursor for the preparation

of dithienylethene derivatives via lithiation-mediated processes.<sup>38</sup> For DTE<sup>o</sup>-COCF<sub>3</sub> and DTE<sup>o</sup>-C<sub>6</sub>F<sub>5</sub>, the sequence to introduce the phosphine and EWG groups in DTE1 was governed by their sensitivity toward lithium reagents. For this reason, we first conducted lithiation of DTE1 with tbutyllithium (tBuLi), followed by quenching with chlorodiphenylphosphine to obtain phosphanyl compound DTE2 (67% yield); next, additional lithiation with tBuLi followed by the reaction with ethyl trifluoroacetate (69% yield) or hexafluorobenzene (70% yield) furnished DTE°-COCF3 and DTE°-C<sub>6</sub>F<sub>5</sub>, respectively. As for DTE°-Ph, we inverted the order in which the external thiophene substituents were introduced to the DTE core according to a previously reported procedure.<sup>39</sup> In particular, its phenyl side group was first installed through consecutive lithiation, borylation, and Suzuki coupling steps to produce compound DTE3 (83% yield), which was then subjected to further lithiation with nbutyllithium (nBuLi) and subsequent treatment with chlorodiphenylphosphine (68% yield) to introduce the phosphanyl moiety of DTE°-Ph. All synthesized phosphines presented a singlet in the <sup>31</sup>P{<sup>1</sup>H} NMR spectra at around  $\delta \approx -19.6$ ppm, a chemical shift value that is similar to those reported for (2-methyl-5-thienyl)diphenylphosphine  $(\delta = -21.9 \text{ ppm})^{40}$ and symmetric bis(phosphine) DTE ( $\delta \approx -20.0$  ppm).<sup>41,42</sup> As expected, this result corroborates that the phosphanyl and electron-modulating groups in the nonplanar structure of DTE°-COCF<sub>3</sub>, DTE°-C<sub>6</sub>F<sub>5</sub>, and DTE°-Ph are not conjugated and, therefore, do not significantly affect each other.

When two equivalents of the phosphine ligands were used to react with *trans*-[PdCl<sub>2</sub>(PhCN)<sub>2</sub>], monometallic palladium(II) 1:2 type complexes were formed (Figure 1a). For all of these compounds, an isolated singlet was registered in their  ${}^{31}P{}^{1}H$ NMR spectra that shifted downfield to  $\delta \approx 12$  ppm, thereby corroborating metal complexation and the formation of exclusively one isomer (cis or trans). In addition, negligible differences were found in the chemical shift of this  ${}^{31}P{}^{1}H{}$ NMR signal for the three Pd complexes prepared, which again demonstrates the lack of electronic communication between the external thiophene substituents in the open isomers of the ligands. Furthermore, complexation to the Pd center could also be verified by the downfield shift of the <sup>1</sup>H NMR resonance of the thiophene ring proton next to the phosphanyl group. Single crystals suitable for X-ray structure determination were obtained for the three complexes, which revealed that the Pd complexes have a square-planar geometry with trans orientation of the phosphine ligands (Figure 1b, Figures S1-S3 and Table S1). The main difference observed between the crystal structures of these compounds was the conformation of their DTE ligands. In [PdCl<sub>2</sub>(DTE°-Ph)<sub>2</sub>] and [PdCl<sub>2</sub>(DTE°- $COCF_3)_2$ , the ligands are locked in a distorted parallel openstate conformation with a distance of 4.31 and 5.19 Å, respectively, between the carbon atoms that should react upon ring-closing photoisomerization  $(C_{16}-C_{26})$ . As previously described in the literature,<sup>25</sup> this type of conformation cannot undergo the light-induced conrotatory electrocyclization reaction to produce the corresponding closed isomer and, consequently, no photochromism was observed in the solid state for  $[PdCl_2(DTE^{\circ}-Ph)_2]$  and  $[PdCl_2(DTE^{\circ}-COCF_3)_2]$ . Contrarily, in the case of  $[PdCl_2(DTE^{\circ}-C_6F_5)_2]$ , the DTE ligands are present in an antiparallel open-state conformation with a shorter distance between the reactive carbons  $(C_{16}-C_{26})$ 3.69 Å), two structural features that are compatible with the photoinduced ring-closing reaction.43 Indeed, irradiation at

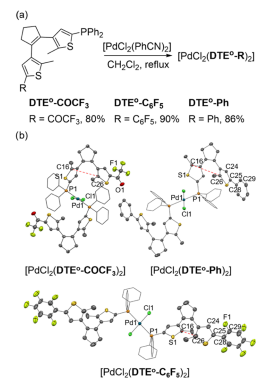


Figure 1. (a) Synthesis of  $[PdCl_2(DTE^\circ-COCF_3)_2]$ ,  $[PdCl_2(DTE^\circ-C_6F_5)_2]$ , and  $[PdCl_2(DTE^\circ-Ph)_2]$ . (b) Molecular structures of  $[PdCl_2(DTE^\circ-Ph)_2]$ ,  $[PdCl_2(DTE^\circ-COCF_3)_2]$ , and  $[PdCl_2(DTE^\circ-C_6F_5)_2]$ . Thermal ellipsoids are set at the 50% probability level. For clarity, P-bound phenyl rings are depicted in wireframe style, and cocrystallized solvent and hydrogen atoms are omitted. Distances between the reactive carbon atoms in DTE photoisomerization are marked with dashed red line.

312 nm yielded a color change from yellow to red, and the process was reversed with irradiation at 520 nm, thus demonstrating solid-state photoswitching for  $[PdCl_2(DTE^\circ-C_6F_5)_2]$  (Figure S4).

Photochemical Behavior of Ligands and Complexes. Irrespective of their behavior in the solid state, all of the DTEbased ligands and complexes prepared should photoisomerize in solution upon irradiation. However, while DTE-COCF<sub>3</sub>, DTE-C<sub>6</sub>F<sub>5</sub>, and DTE-Ph should just photoconvert between their ring-open and ring-closed isomers, a more complex situation is expected for their complexes (Scheme 3). Because of their 2:1 phosphine/metal stoichiometry, these compounds must toggle between three different states, where DTE-based ligands are both ring-open (oo), one of them ring-open and the other ring-closed (oc), or both ring-closed (cc).

To study these photoinduced processes, the UV-vis absorption spectra of the initial ring-open isomer of the synthesized DTE-based ligands and their palladium(II) complexes were first recorded in cyclohexane (Figures 2a,b and S5). In addition, TD-DFT calculations at the CAM-B3LYP-D3/6,31G(d,p) level were performed to further investigate the electronic excitations of these compounds (Tables S2-S5), for which ground state geometries were first computed. For ligands and complexes bearing open-state DTE units, we considered only their photocyclizable antiparallel conformation in our calculations. All open-state phosphine ligands were characterized by a strong absorption in the UV region with maxima around  $\lambda_{abs} \approx 290$  nm, which was reproduced in computations and could be attributed to  $\pi$ - $\pi$ \* Scheme 3. Photoisomerization Processes for DTE-Based Free Ligands and Their Palladium(II) Complexes

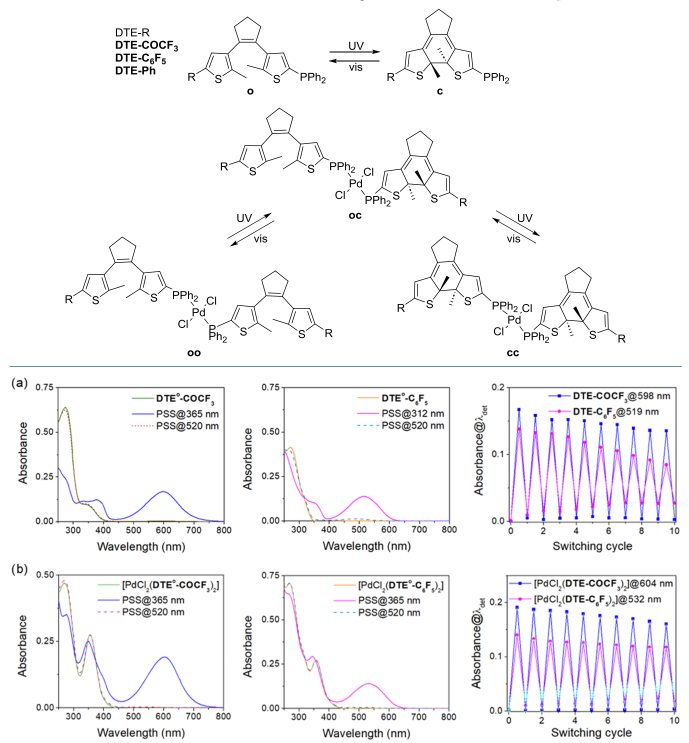


Figure 2. (a) Variation of the absorption spectrum of free ligands  $DTE^{\circ}$ -COCF<sub>3</sub> ( $c = 1.5 \times 10^{-5}$  M) and  $DTE^{\circ}$ -C<sub>6</sub>F<sub>5</sub> ( $c = 1.5 \times 10^{-5}$  M) in cyclohexane upon sequential irradiation with UV ( $\lambda_{exc} = 365$  for 90 s, or  $\lambda_{exc} = 312$  nm for 110 s) and green light ( $\lambda_{exc} = 520$  nm for 120–160 s) until the corresponding ring-closing and ring-opening photostationary states (PSSs) are obtained, respectively. The variation of the absorbance at the spectral maximum of the ring-closed isomer of these compounds ( $\lambda_{det} = 598$  or 519 nm) upon 10 consecutive photoswitching cycles under the same concentration and irradiation conditions is also given. (b) Variation of the absorption spectrum of the open-state complexes [PdCl<sub>2</sub>(DTE°-COCF<sub>3</sub>)<sub>2</sub>] ( $c = 1.5 \times 10^{-5}$  M) and [PdCl<sub>2</sub>(DTE°-C<sub>6</sub>F<sub>5</sub>)<sub>2</sub>] ( $c = 1.5 \times 10^{-5}$  M) in cyclohexane upon sequential irradiation with UV ( $\lambda_{exc} = 365$  nm for 90 and 150 s, respectively) and green light ( $\lambda_{exc} = 520$  nm for 180 s) until the corresponding ring-closing and ring-opening PSSs are obtained, respectively. The variation of the absorbance at the spectral maximum of the ring-closed isomer of these compounds ( $\lambda_{det} = 604$  or 532 nm) upon 10 consecutive photoswitching cycles at the same concentration and irradiation conditions is also given.

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Table 1. Photoc	hemical Properties	s of DTE-Based	Ligands an	d Complexes

	$\lambda_{\mathrm{abs}}^{\mathrm{o}}  [\mathrm{nm}] \; (\varepsilon \; [\mathrm{M}^{-1} \; \mathrm{cm}^{-1}])^a$	$\lambda_{abs}^{c} [nm] (\varepsilon [M^{-1} cm^{-1}])^{b}$	$PSS_{o-c}$ composition [%] <sup>c</sup>	$\Phi_{o-c}{}^d$	$\Phi_{c-o}^{e}$
DTE-COCF <sub>3</sub>	268 (35 673), 339 (6 431)	598 (12 261)	91:9	0.480	0.012
$[PdCl_2(DTE-COCF_3)_2]$	271 (34 149), 355 (19 065)	604 (18 870)	45:45:10	0.148/0.048	0.013/0.014
DTE-C <sub>6</sub> F <sub>5</sub>	272 (28 303)	519 (11 379)	81:19	0.530	0.015
$[PdCl_2(DTE-C_6F_5)_2]$	268 (40 394), 357 (14 367)	532 (19 391)	25:45:30	0.047/0.019	0.015/0.018
DTE-Ph	273 (28 117)	518 (13 673)	39:61	0.485	0.011
$[PdCl_2(\mathbf{DTE-Ph})_2]$	270 (42 700), 359 (15 400)	538 (16 310)	27:44:29	0.060/0.023	0.015/0.014

<sup>*a*</sup>Wavelength and molar absorptivity coefficient of the absorption band maxima of the open isomer (for complexes, **oo** state) in cyclohexane. <sup>*b*</sup>Wavelength and molar absorptivity coefficient of the maximum of the visible absorption band of the closed isomer (for complexes, **cc** state) in cyclohexane. <sup>*b*</sup>Wavelength and molar absorptivity coefficient of the maximum of the visible absorption band of the closed isomer (for complexes, **cc** state) in cyclohexane. <sup>*b*</sup>Composition of the PSS reached for the photocyclization process in toluene-*d*<sub>8</sub> upon irradiation at  $\lambda_{exc} = 365$  nm (**DTE-COCF**<sub>3</sub> and all of the complexes) or 312 nm (**DTE-C**<sub>6</sub>**F**<sub>5</sub> and **DTE-Ph**). DTE<sup>c</sup>/DTE<sup>o</sup> and DTE<sup>cc</sup>/DTE<sup>oc</sup>/DTE<sup>oc</sup> concentration ratios are given for free ligands and complexes, respectively. <sup>*d*</sup>Photocyclization quantum yields measured in cyclohexane at  $\lambda_{exc} = 355$  nm (**DTE-COCF**<sub>3</sub> and all of the complexes) or 312 nm (**DTE-C**<sub>6</sub>**F**<sub>5</sub> and **DTE-Ph**). For the complexes, separate  $\Phi_{o-c}$  values are given for the **oo**  $\rightarrow$  **oc** and **oc**  $\rightarrow$  **cc** ring-closing processes. <sup>*e*</sup>Photocycloreversion quantum yields measured in cyclohexane at  $\lambda_{exc} = 532$  nm. For the complexes, separate  $\Phi_{c-o}$  values are given for the **cc**  $\rightarrow$  **oc** and **oc**  $\rightarrow$  **oo** ring-opening processes.

transitions (HOMO – LUMO+1 or HOMO – LUMO) of their core (Tables S3 and S5, Figures S24–S26). In the case of **DTE-COCF**<sub>3</sub>, a shallower absorption band ranging up to  $\lambda_{abs} \approx 400$  nm was detected due to the lowering of the energy of the LUMO caused by the introduction of a strong EWG (Table S2).<sup>44</sup> As for the UV–vis absorption of the open-state Pd<sup>II</sup> complexes, they not only preserved all of the spectral features of their constituting DTE° ligands but also exhibited a new red-shifted band with a maximum at  $\lambda_{abs} \approx 350$  nm (Table 1). According to our TD-DFT calculations, this additional absorption band can be mainly assigned to a ligand-to-metal charge transfer (LMCT) transition, as the LUMO of all of the open-state complexes is located on the palladium center (Tables S4 and S5 and Figures S27–S29).

Based on the electronic absorption properties of the openstate ligands and complexes, their ring-closing photoisomerization was assayed upon irradiation with UV light. For open-state ligands, a broad, red-shifted absorption band emerged in the visible part of the spectrum upon UV illumination of their cyclohexane solutions, which changed from colorless to deep blue ( $\lambda_{abs}$  = 598 nm for DTE<sup>c</sup>-COCF<sub>3</sub>) or deep pink ( $\lambda_{abs} \approx$ 518 nm for  $DTE^{c}-C_{6}F_{5}$  and  $DTE^{c}-Ph$ ) (Figure 2a, Table 1 and Figure S5). This behavior is characteristic of closed-state DTEs,45 as confirmed by TD-DFT calculations (Table S5, Figures S24-S26) and additional NMR spectroscopic measurements. In particular, we observed the appearance of a new set of signals for the closed isomer in the <sup>1</sup>H, <sup>31</sup>P (proton-coupled and decoupled), and, when applicable, <sup>19</sup>F NMR spectra of the UV-irradiated ligands in toluene- $d_{8}$ , which were upfield- (<sup>1</sup>H NMR) or downfield-shifted (<sup>31</sup>P, <sup>31</sup>P{<sup>1</sup>H}, and <sup>19</sup>F NMR) relative to the NMR resonances of the open state (Figures S7-S14). Analysis of these NMR data also revealed that the photocyclization reaction of DTE-based ligands was not quantitative. Instead, a photostationary state (PSS) composed of an equilibrium mixture of the ligands' o and c states was obtained in all of the cases since both isomers absorb at the UV excitation wavelength used and, therefore, should simultaneously undergo ring-closing and ring-opening reactions. Under our illumination conditions in toluene- $d_{8y}$ better photocyclization conversions were observed with DTE°-**COCF**<sub>3</sub> (91% at  $\lambda_{\text{exc}}$  = 365 nm) and **DTE°-C**<sub>6</sub>**F**<sub>5</sub> (81% at  $\lambda_{\text{exc}}$  = 312 nm) compared to DTE°-Ph (39% at  $\lambda_{exc}$  = 312 nm) (Table 1). This can be ascribed to the lower ring-closing quantum yield and high photodegradation tendency of DTE-Ph, which limited the UV exposure time for photocyclization and led to a lower percentage of the closed isomer.

Similar to free DTE ligands, irradiation of the Pd<sup>II</sup> complexes in cyclohexane with UV light also caused the appearance of an absorption band in the visible part of the spectrum and a concomitant color change (Figure 2b, Table 1 and Figure S5). In combination with NMR spectroscopic measurements in toluene- $d_8$  and TD-DFT calculations, it was demonstrated that DTE photocyclization takes place in the complexes, in contrast to some previously reported DTE-based phosphine-metal compounds.<sup>39,46</sup> However, palladium(II) complexation did have a relevant effect on the UV-induced ring-closing process of the phosphine ligands. First, we could use less energetic UV radiation to promote the photocyclization of  $[PdCl_2(DTE^\circ-C_6F_5)_2]$  and  $[PdCl_2(DTE^\circ-Ph)_2]$  $(\lambda_{exc} = 365 \text{ nm})$  compared to DTE°-C<sub>6</sub>F<sub>5</sub> and DTE°-Ph  $(\lambda_{exc} =$ 312 nm) because of the new red-shifted band measured for the two complexes at  $\lambda_{abs} \sim 350$  nm. Second, a bathochromic shift of the absorption band of the closed isomer in the visible range was registered relative to the free ligands ( $\lambda_{abs} = 604$ , 532, and 538 nm for  $[PdCl_2(DTE^{c}-COCF_3)_2]$ ,  $[PdCl_2(DTE^{c}-C_6F_5)_2]$ , and [PdCl<sub>2</sub>(DTE<sup>c</sup>-Ph)<sub>2</sub>]) (Figure 2a,b, Table 1 and Figures S27-S29). According to TD-DFT calculations, this effect is due to the decrease in the HOMO-LUMO gap of DTE-based phosphines upon Pd<sup>II</sup> complexation, which should be especially important for DTE-C<sub>6</sub>F<sub>5</sub> and DTE-Ph, as experimentally observed (Table S2). On the other hand, the spectral features of the closed DTE absorption band were not influenced by the state of the nearby DTE ligand in the same complex (open or closed), i.e., it did not evolve when photoconverting from the oc species with one open and one closed DTE unit to the fully closed cc complex, which was substantiated by TD-DFT calculations (Figures S27-S29). As a result, these two different photocyclization products could only be discriminated by NMR spectroscopic experiments, which proved that they were sequentially formed upon UV irradiation and allowed us to determine the composition of the equilibrium PSS generated in toluene- $d_8$  (Table 1 and Figures S15-S22). Interestingly, only one set of signals was detected for the cc state of the complexes in the NMR spectra, though it must comprise a mixture of different stereoisomers because of the chiral centers created in their DTE units upon conrotatory photocyclization.<sup>25</sup> Therefore, this suggests that the stereochemistry of these ring-closed units has a minor effect on the chemical environment of the phosphorus atoms of the phosphines, and consequently, it should not affect the catalytic activity of the complexes. Further analysis of the NMR spectra of the complexes allowed us concluding that metal

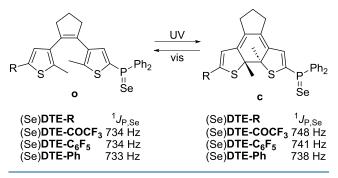
coordination detrimentally affected the photocyclization conversion of DTE°-COCF<sub>3</sub> and DTE°-C<sub>6</sub>F<sub>5</sub>, as only 66 and 48% of the DTE units in  $[PdCl_2(DTE^{\circ}-COCF_3)_2]$  and  $[PdCl_2(DTE^{\circ}-C_6F_5)_2]$  could be ring-cycled. This result is consistent with the lower photocyclization quantum yields  $(\Phi_{o-c})$  measured for both the **oo** and **oc** states of the complexes relative to the free ligands (Table 1), which can be attributed to two main effects: (i) the competition between photoisomerization and ligand-to-metal charge transfer, which, as previously mentioned, gives rise to additional absorption bands in the UV region of the complexes at which they are excited to promote photocyclization, and (ii) the further reduction of  $\Phi_{o-c}$  for the second ring-closing step in the complexes, a behavior already reported for other DTE dimers,<sup>47</sup> which can be ascribed to intramolecular energy transfer between the open and closed DTE units in the oc state upon photoexcitation, i.e., the photocyclization of one of these units severely hinders the ring-closing reaction for the second DTE group.

Once photocyclized with UV light, the closed states of the DTE-based ligands and their palladium(II) complexes were found to be thermally stable in solution, and no spontaneous back-isomerization was detected in the dark at room temperature. Accordingly, visible irradiation ( $\lambda_{exc} = 520$  nm) was applied to promote photoinduced ring-opening of these compounds and to demonstrate the reversibility of their photoswitching behavior in solution. For all of them, complete disappearance of the visible absorption band associated with the closed isomer was registered, which demonstrates quantitative photochemical cycloreversion, regardless of metal complexation (Figure 2a,b and Figures S5, S15, S18, S19, S21, and S22, see SI Materials and Methods for additional details). This is due to the selective photoexcitation of closedstate DTEs with visible light that counterbalances their commonly low ring-opening quantum yields,<sup>25</sup> as we also measured herein for the ligands and the cc and oc states of the complexes ( $\Phi_{c-o}$  < 0.02, Table 1). In spite of this, some residual visible absorption was registered for some of these compounds upon light-induced photocyclization reversion  $(DTE-C_6F_5 \text{ in Figure 2a; } DTE-Ph \text{ and } [PdCl_2(DTE-Ph)_2]$ in Figure S5), which could not be attributed to unreacted closed-state species. Instead, it arose from DTE photodegradation, which is normally associated with the UV irradiation of the closed isomer during photocyclization and leads to a characteristic product that absorbs at  $\lambda_{\rm abs}\approx$  500 nm.<sup>48</sup> This effect was further proven by an examination of the fatigue resistance of ligands and complexes upon repetitive photoinduced ring-closing and ring-opening cycles in cyclohexane (Figure 2a,b and Figures S5 and S6). Although some deterioration of their photoswitching behavior was eventually observed for all of these compounds, the highest photodegradation effects were registered for DTE-Ph, DTE-C<sub>6</sub>F<sub>5</sub>, and  $[PdCl_2(DTE-Ph)_2]$ . We ascribe these results to two main factors that increase the photostability of the remaining ligands and complexes: (i) the presence of the strong trifluoroacetyl EWG at the external position of the thiophene ring, which is known to slow down DTE photodegradation,48 and (ii) the use of less energetic UV light to photoisomerize the palladium(II) complexes relative to the free ligands. Consequently, target compounds  $[PdCl_2(DTE-COCF_3)_2]$  and  $[PdCl_2(DTE-C_6F_5)_2]$  showed the highest resistance to photodegradation.

Photomodulation of the Properties of the Phosphine Ligand. As anticipated by molecular design and demonstrated by <sup>31</sup>P (proton-coupled and -decoupled) NMR spectroscopic measurements discussed above, the phosphanyl and electronmodulating groups of the synthesized DTE-based ligands are electronically decoupled in their open state. As a result, these ligands should present similar binding properties to metals, a behavior that we aimed to modulate upon photoisomerization. In fact, UV-induced ligand ring-closing caused a measurable downfield shift of the <sup>31</sup>P and <sup>31</sup>P{<sup>1</sup>H} NMR signal of these compounds ( $\Delta \delta$  = 9.4, 8.2, and 8.3 ppm for DTE-COCF<sub>3</sub>,  $DTE-C_6F_5$ , and DTE-Ph, respectively), thus suggesting a change in the electronic properties of their phosphine groups that might be dependent on the nature of the external substituent present in the other thiophene of the DTE core. In particular, introduction of electron-withdrawing substituents such as trifluoroacetyl and pentafluorophenyl to phosphines is expected to (i) increase the s character of the lone pair of electrons at the phosphorus atom involved in  $\sigma$  bonding to metals while (ii) stabilizing and enlarging the size of the phosphine's  $\sigma^*$  antibonding orbital participating in metal  $\pi$ backbonding.

A well-established method to assess such an effect for phosphines is to measure the spin–spin coupling constant between <sup>31</sup>P and <sup>77</sup>Se ( ${}^{1}J_{P,Se}$ ) for the corresponding selenide derivatives (Scheme 4),<sup>39,41</sup> which were prepared by heating

Scheme 4. Variation of  ${}^1J_{P,Se}$  in the Open and Closed Isomers of the Selenides of DTE-COCF<sub>3</sub>, DTE-C<sub>6</sub>F<sub>5</sub>, and DTE-Ph



the ligands and gray selenium in CDCl<sub>3</sub>. In these compounds,  $^{1}J_{P,Se}$  values depend on the s character of the P = Se bond, which is related to the electronic character and size of the substituents on the phosphorus atom.<sup>49–51</sup> Consequently, they provide an estimate of the  $\sigma$ -donating ability of phosphines, which is lower for higher values of  ${}^{1}J_{P,Se}$ . For the open-state selenides of DTE-COCF<sub>3</sub>, DTE-C<sub>6</sub>F<sub>5</sub>, and DTE-Ph, measured  ${}^{1}J_{P,Se}$  values are almost identical and match the reported coupling constant for the selenide of (2-methyl-5-thienyl)-diphenylphosphine  $({}^{1}J_{P,Se} = 733 \text{ Hz})^{40}$  (Scheme 4). Again, this result corroborates that the two thiophenes in these DTE structures are electronically isolated, and for that reason, their phosphanyl groups have similar electronic properties. In contrast, as ring-closing extends the conjugation throughout the DTE moiety, the electron density on the phosphorus atom should decrease and can be affected by the external substituent in the other thiophene ring with which it communicates. This behavior was indeed proven by NMR measurements of the closed selenide isomers: higher <sup>1</sup>J<sub>P,Se</sub> values were detected relative to the open isomers with increments  $(\Delta^{c-o}({}^{I}J_{P,Se}))$  that

scaled up with the electron-withdrawing nature of the electronmodulating substituents (Scheme 4, Table 2), i.e., the studied

# Table 2. Experimental and Computed Parameters to Estimate the Photomodulation of the Properties of the Phosphine Ligands DTE-COCF<sub>3</sub>, DTE-C<sub>6</sub>F<sub>5</sub>, and DTE-Ph

	free ligands			Pd <sup>II</sup> complexes <sup>a</sup>	
	$\Delta^{c-o}({}^{1}J_{P_{o}Se})$ [Hz]	$\Delta^{\mathrm{c-o}}(q_{\mathrm{P}}^{\mathrm{Mulliken}})^{c,d}$	$\Delta^{c-o}(\%)$ $s_{p}^{c,e}$	$\Delta^{c-o}(BE_{P-Pd})$ [kcal·mol <sup>-1</sup> ] <sup>c,f</sup>	
DTE-COCF <sub>3</sub>	14	0.014	0.79	-1.98	
DTE-C <sub>6</sub> F <sub>5</sub>	7	0.004	0.42	-0.92	
DTE-Ph	5	< 0.001	0.34	-0.64	

<sup>*a*</sup>trans-[PdCl<sub>2</sub>(**DTE-COCF**<sub>3</sub>)<sub>2</sub>], trans-[PdCl<sub>2</sub>(**DTE-C**<sub>6</sub>F<sub>5</sub>)<sub>2</sub>], and trans-[PdCl<sub>2</sub>(**DTE-Ph**)<sub>2</sub>]. <sup>*b*</sup>Difference in <sup>1</sup>J<sub>P,Se</sub> for the corresponding selenides measured in CDCl<sub>3</sub>. <sup>*c*</sup>Computed at the B3LYP-D3 level in THF (see the Experimental Section for further details). <sup>*d*</sup>Difference in Mulliken charges in electronic units on the phosphorus atom. <sup>*c*</sup>Difference in percentage of *s* character of the phosphorus lone pair of electrons. <sup>*f*</sup>Difference in phosphine-Pd<sup>II</sup> bond energy (per one bond) between the **oo** and **cc** isomers.

phosphines became electron-poorer upon photocyclization and introduction of a stronger EWG in the opposite thiophene ring. Thus, the highest value of  $\Delta^{c-o}({}^1J_{P,Se})$  was registered for the selenide of **DTE-COCF**<sub>3</sub> ( $\Delta^{c-o}({}^1J_{P,Se}) = 14$  Hz), which is the highest reported for DTE-based phosphine ligands<sup>39,41</sup> and mimics the electronic effects caused by substituting one phenyl ring in triphenylphosphine for a *tert*-butyl group.

To further investigate the photomodulation of the electronic features for the prepared DTE-based phosphines, we analyzed a set of properties derived from DFT calculations of their ground state structures. As a first step, we considered the variation of the Mulliken charges on the phosphorus atom  $(\Delta^{c-o}(q_P^{\text{Mulliken}}))$  and the percentage of s character of the lone pair of electrons at phosphorus  $(\Delta^{c-o}(\%s_p))$  for open antiparallel and closed conformations of DTE-COCF<sub>3</sub>, DTE- $C_6F_5$ , and DTE-Ph (Table 2). According to the Mulliken charges, photocyclization decreases electron density on phosphorus, while through NBO analysis,<sup>52</sup> we can anticipate an increase in s orbital participation in the phosphorus lone pair of electrons. More importantly, the variation of these parameters was found to increase with the electron-withdrawing power of the introduced electron-modulating group, thus again validating that DTE-COCF<sub>3</sub> and, to a lesser extent,  $DTE-C_6F_5$  experience the largest change in phosphine's electronic properties upon photoisomerization.

As a second step, the effect of the light-induced modulation of the DTE-based phosphine ligands on the bond energy in their Pd<sup>II</sup> complexes was investigated computationally. For this, we analyzed the difference in phosphine-Pd<sup>II</sup> binding energy ( $\Delta^{c-o}(BE_{P-Pd})$ ) between the **oo** and **cc** isomers of their 2:1 *trans*-phosphine–palladium(II) complexes (Table 2). As expected, due to the loss of the phosphine's  $\sigma$ -donating ability, a varying decrease in BE<sub>P-Pd</sub> for the complexes bearing ringclosed DTE ligands is observed, depending on the nature of the introduced electron-modulating group. Thus, weakening of the phosphine–palladium(II) binding upon photocyclization was observed for the ligands bearing the electron-withdrawing pentafluorophenyl and especially trifluoroacetyl substituents, as we initially devised.

**Catalytic Studies.** Among the vast range of Pd-catalyzed reactions, Stille coupling was chosen to evaluate the activity of

the open and closed states of the prepared metal complexes. For this reaction, previous mechanistic studies have established that bulky phosphines accelerate the rate of oxidative addition while electron-poor phosphines are advantageous for the transmetalation step.<sup>36,53–55</sup> This makes Stille coupling a suitable benchmark process to validate our model toward photoswitchable catalysis, as we have proven above that the electron density on the DTE-based phosphines developed herein can be modulated upon photoisomerization. With this aim, the palladium(II) complexes were tested as precatalysts for the Stille reaction between iodobenzene and tributylvinyltin in THF- $d_8$  at two different temperatures (room temperature and 50 °C) (Table 3). In all cases, catalytic experiments were

 Table 3. Comparison of the Investigated Complexes as

 Precatalysts in the Stille Coupling Reaction

		25 °C		50 °C	
precatalyst [Pd]	state <sup>a</sup>	entry	yield (%) <sup>b</sup>	entry	yield (%) <sup>b</sup>
[PdCl <sub>2</sub> (DTE-COCF <sub>3</sub> ) <sub>2</sub> ]	00	1	4.5	8	36.4
	cc	2	8.8	9	48.9
$[PdCl_2(DTE-C_6F_5)_2]$	00	3	21.2	10	42.5
	cc	4	29.5	11	59.6
$[PdCl_2(DTE-Ph)_2]$	00	5	13.4	12	53.8
	cc	6	14.2	13	43.4
$\left[ PdCl_2(PPh_3)_2 \right]$		7	9.8	14	28.0

<sup>*a*</sup>cc state here is a closed-state enriched precatalyst complex as specified in the text. <sup>*b*</sup>The average yields of two repetitions were determined by <sup>1</sup>H NMR spectroscopy using 1,3,5,-trimethoxybenzene as a standard.

conducted in the dark and separately for the pure oo complexes and for cc-enriched mixtures of isomers, which were observed to be thermally stable even at the highest temperature considered (50 °C; Figures S33-S36). Because of the moderate efficiency of oo-to-cc photocyclization in the complexes (see Table 1), we maximized the relative amount of cc species in such mixtures by first ring-closing the corresponding free ligand and then conducting metal complexation (see the Supporting Information for further details). In this way, more cc-enriched precatalyst mixtures could be prepared, which yet contained a moderate relative concentration of cc isomers: 52, 38, and 49% for [PdCl<sub>2</sub>(DTE- $COCF_3)_2$ ], [PdCl<sub>2</sub>(DTE-C<sub>6</sub>F<sub>5</sub>)<sub>2</sub>], and [PdCl<sub>2</sub>(DTE-Ph)<sub>2</sub>], respectively (Figures S30-S32). To compare the catalytic efficiency of these mixtures with those of their oo samples, we monitored the kinetics of the Stille reaction for 6 h and measured the difference in product formation after this time (entries 1-6 and 8-13 in Table 3; Figure S37). In addition, equivalent measurements were performed using  $[PdCl_2(PPh_3)_2]$  (entries 7 and 14 in Table 3) as a non-lightresponsive reference precatalyst, which, in most of the cases, turned out to be less efficient than our DTE-based complexes.

As shown in Table 3, two different behaviors were observed in the catalytic tests. For complexes bearing DTE units with external electron-withdrawing substituents, a large increase in yields for the Stille coupling at 6 h was registered for the ccenriched mixtures compared to the **oo** complexes: about 2-(from entry 1 to entry 2) and 1.35-fold increase in yield (from entry 8 to entry 9) for  $[PdCl_2(DTE-COCF_3)_2]$  at room temperature and 50 °C, respectively, and about 1.4-fold increase in yield for  $[PdCl_2(DTE-C_6F_5)_2]$  (from entries 3 to 4 and 10 to 11) both at room temperature and 50 °C. By contrast, DTE photocyclization did not enhance the Stille coupling efficiency for  $[PdCl_2(DTE-Ph)_2]$ , which lacks the electron-withdrawing substituent on the DTE core. In this case, similar reaction conversions were measured for the oo complex and cc-enriched mixture at room temperature (entries 5 and 6), while we observed a decrease in reactivity upon ringclosing at 50 °C (entries 12 and 13). Importantly, these results are in agreement with the prediction made to accomplish photoswitchable catalysis by installing external EWGs in DTEbased phosphines. Upon photocyclization, the electron-withdrawing and phosphanyl substituents become selectively conjugated in these compounds, thus varying the electronic properties of the phosphine ligand and affecting the catalytic behavior of its metal complexes. Unfortunately, the catalytic modulation accomplished in this way is limited by the nonquantitative nature of DTE photocyclization, which prevented us from conducting experiments with pure cc precatalysts.

To rationalize the modulation of Stille coupling reactivity determined for the open and closed states of  $[PdCl_2(DTE-COCF_3)_2]$  and  $[PdCl_2(DTE-C_6F_5)_2]$ , we conducted additional DFT calculations. For this, we considered the Stille reaction mechanism,<sup>36,56</sup> which requires previous reduction of the palladium(II) precatalyst used to palladium(0) before the catalytic cycle begins. Similar to other palladium-catalyzed couplings, the catalytic cycle consists of three major steps: oxidative addition, transmetalation, and reductive elimination,<sup>57</sup> among which the first two are typically the rate-determining steps when aryl halides are used as substrates (Figure 3).<sup>35,58</sup> Over the years, various pathways have been

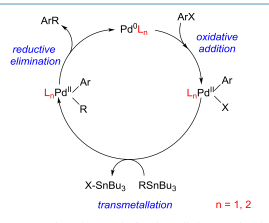


Figure 3. General catalytic cycle for the palladium-catalyzed Stille coupling reaction.

proposed for each of these steps. On the one hand, the oxidative addition of the organic electrophile to Pd<sup>0</sup> can occur through a monophosphine pathway, an associative displacement pathway, and a bisphosphine pathway.<sup>59,60</sup> Principally, the presence of bulky ligands should favor oxidative addition via a monoligated transition state owing to steric repulsion.<sup>61</sup> Nevertheless, it was recently established that intramolecular dispersion forces can serve to stabilize a bisligated transition state for bulky ligands, such as  $PtBu_3$ ,<sup>62</sup> thus demonstrating the importance of incorporating dispersion effects into the analysis. Accordingly, we explored both types of pathways in our computations. As for the transmetalation step, it can also proceed through three different mechanisms: cyclic, open, and ionic.<sup>58</sup> In our calculations, we only considered the first of these cases, as halides are considered to be good bridging

ligands that facilitate the formation of cyclic transition states; in contrast, open and ionic mechanisms are preferred in the case of poorly coordinating anionic ligands and highly polar solvents. Finally, the reductive elimination step can also proceed through bisligated and monoligated transition states.<sup>58</sup> For simplicity, herein, we only explored the second of these options and did not compute the bisligated pathway.

Based on these mechanistic assumptions, we computed the catalytic cycle of the Stille coupling reaction investigated experimentally. Calculations were carried out for complexes formed with ligand DTE-COCF<sub>3</sub>, as it should impart the strongest electronic effects upon photoisomerization. Three possible states of the DTE-COCF<sub>3</sub>-based Pd<sup>0</sup> catalyst were considered in these calculations:  $[Pd(DTE^{\circ}-COCF_3)_2]$ , [Pd- $(DTE^{\circ}-COCF_3)(DTE^{\circ}-COCF_3)]$ , and  $[Pd(DTE^{\circ}-COCF_3)_2]$ (Figure 4, Figures S38-S40 and Table S6). Reaction intermediates and transition states were computed at the B3LYP-D3 level of theory (see the Supporting Information for additional details). The Gibbs energies of the species at 298 K are presented relative to the zero point consisting of the corresponding bisligated palladium(0) complex (0, Figure 4), phenyl iodide, and tributylvinyltin. For the oxidative addition step, the monophosphine pathway was found to be disfavored, as phosphine ligand dissociation was computed to have a high energy requirement (>20 kcal/mol, Figures S39-S40). It is worth noting, however, that the energy barrier for the dissociation of the closed-state ligand from the cc (23.1 kcal/mol) and oc (26.8 kcal/mol) species is lower than that for the dissociation of the open ligand from the **oo** (25.8 kcal/ mol) and oc (27.3 kcal/mol) states of the catalyst, thus corroborating our predictions on the effect of the DTE-COCF<sub>3</sub> isomerization state on the stability of the phosphinepalladium bond. The bisphosphine oxidative addition pathway was found to be preferred as it proceeds through the less energetic three-center transition state TS1, which is accessed through the previous intermediate 1 (Figure 4). Among the transition states TS1 computed for the three different catalytic systems, the lowest energy values are associated with the cc and oc states, at 7.4 and 7.8 kcal/mol, respectively, while the oo isomer requires 10.6 kcal/mol. As the transition state for the oxidative addition of the bisligated complex leads to the formation of the fully *cis*-coordinated Pd<sup>II</sup> species 2, ligand dissociation is a prerequisite for subsequent steps, which was found to be slightly less energy demanding for the cc system (13.2 kcal/mol for the cc species vs 14.4 kcal/mol for the oo species). Then, the formed intermediate 3 isomerizes through the transition state TS3 (energy barrier  $\approx 2.8-4.7$  kcal/mol), and upon tin coordination, transmetalation takes place with the highest energy barrier among all of the steps. In particular, the energy barrier for the cyclic transition state TS5 involving the closed-state Pd<sup>0</sup> species is 13.5 kcal/mol, which is lower than that for the open-state catalytic system (15.4 kcal/mol). Lastly, reductive elimination of intermediate 6 was observed to take place through transition state TS7, with an energy barrier difference of only 0.3 kcal/mol between the closed- and openstate isomers. In conclusion, our computational analysis revealed that the barriers for the most energy-demanding steps of the investigated Stille coupling reaction, i.e., oxidative addition, ligand dissociation, and transmetalation, are lower for the catalytic palladium species bearing closed-state DTE-COCF<sub>3</sub> ligands. This is in agreement with our experimental results, which showed higher catalytic activity for the ccenriched state of the  $[PdCl_2(DTE-COCF_3)_2]$  (and

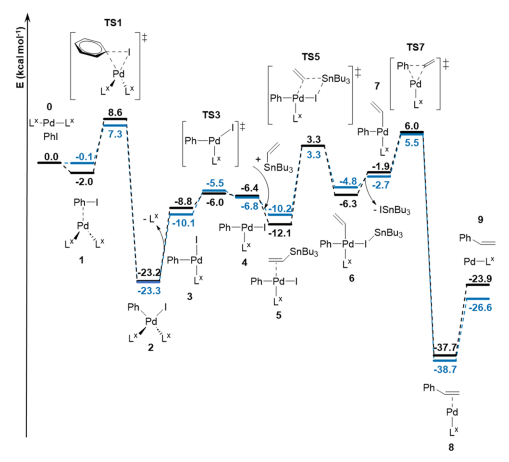


Figure 4. Calculated Gibbs energy profile for the Stille coupling reaction in solution (THF) using  $[PdCl_2(DTE^\circ-COCF_3)_2]$  or  $[PdCl_2(DTE^\circ-COCF_3)_2]$  as a precatalyst, where L<sup>x</sup> is DTE<sup>o</sup>-COCF<sub>3</sub> (black line) or DTE<sup>c</sup>-COCF<sub>3</sub> (blue line).

 $[PdCl_2(DTE-C_6F_5)_2]$ ) precatalyst. Therefore, these results prove that reducing the σ-donation ability of the phosphines of these complexes by making them conjugated with electronwithdrawing groups upon DTE photocyclization favors their catalytic activity in the Stille coupling reaction. Nevertheless, the differences in energy barriers computed for the open- and closed-state isomers of the system along the catalytic cycle are moderate (≈1−3 kcal/mol), which in combination with incomplete DTE photocyclization should account for the limited modulation of reactivity accomplished in our experiments.

## CONCLUSIONS

Dithienylethenes were used to prepare the photoisomerizable phosphine ligands DTE-COCF<sub>3</sub> and DTE-C<sub>6</sub>F<sub>5</sub>, where the electronic communication between the phosphorus atom and electron-withdrawing groups is switched on and off by reversible open- to closed-state conversion. As a result, the  $\sigma$ -donating ability of these phosphines can be efficiently modulated upon light irradiation, in agreement with DFT calculations, improving the behavior observed for the analogous DTE-Ph ligand bearing an EWG-free dithienylethene moiety. Interestingly, when coordinated to palladium-(II), the resulting bisphosphine complexes preserve the ligand's capacity to undergo ring cyclization under illumination, albeit resulting in incomplete phototransformation into their dual closed-state isomer. Finally, when testing the synthesized complexes as precatalysts in the Stille coupling reaction between phenyl iodide and tributylvinyltin, a clear modulation in reaction rate was observed upon photoisomerization of the compounds bearing DTE-COCF<sub>3</sub> and DTE-C<sub>6</sub>F<sub>5</sub> ligands. In particular, the catalytic activity of the complexes increased in the closed state of these ligands, which could be rationalized by DFT calculations. Therefore, these results validate our molecular design toward photoswitchable catalysis, which can be extended to other metals and reactions in the future.

#### EXPERIMENTAL SECTION

**Synthesis.** A detailed description of the synthesis of ligands and complexes is given in the Supporting Information. No uncommon hazards are noted.

**Single-Crystal X-ray Diffraction Analysis.** The data were collected on a Gemini diffractometer (Rigaku Oxford Diffraction) using Mo K $\alpha$  radiation and  $\omega$ -scan rotation. Data reduction was performed with CrysAlisPro,<sup>63</sup> including the program SCALE3 ABSPACK for an empirical absorption correction. As a result of the extremely thin crystal, for  $[PdCl_2(DTE^\circ-C_6F_5)_2]$ , a numerical absorption correction was applied as well using a multifaceted crystal model based on expressions derived by Clark and Reid.<sup>64</sup> All structures were solved by dual space methods with SHELXT<sup>65</sup> and the refinement was performed with SHELXL.<sup>65</sup> Hydrogen atoms were calculated on idealized positions by using the riding model. Structure figures were generated with DIAMOND-4<sup>66</sup> and Mercury (version 2022.2.0).<sup>67</sup>

**Photochemical Characterization.** Photoswitching was monitored by UV–vis and NMR spectroscopy. The photostationary state PSS<sup>oc</sup> composition was determined through <sup>31</sup>P or <sup>19</sup>F NMR spectroscopy from a PSS<sup>oc</sup> state produced by irradiating a toluene $d_8$  solution in an NMR tube with the appropriate wavelength. Spectra of the closed-state isomers shown in Figures S24-S29 were estimated from the PSS<sup>oc</sup> and open-state UV-vis spectra. Photoisomerization quantum yields were determined by monitoring the variation of the UV-vis absorption spectra of ligands and complexes in cyclohexane upon irradiation with UV (for photocyclization,  $\lambda_{exc} = 312$  or 355 nm) or visible light (for photocycloreversion,  $\lambda_{\text{exc}} = 532$  nm). In the case of the free ligands bearing one DTE unit, spectral data were fitted to a simple kinetic model previously reported.<sup>68</sup> For the complexes containing two DTE groups, a more complex kinetic model had to be used to separately determine the photoisomerization quantum yields of their oo, oc, and cc isomers.<sup>69</sup> To apply this model, we assumed the UV-vis absorption spectrum of each DTE unit in the complexes to be independent of the isomerization state of the other, i.e., the extinction coefficients of open DTE units are the same in the oo and oc states, while those of closed DTE groups are equal in the oc and cc states, as suggested by our  $\ensuremath{\text{TD-DFT}}$  calculations and observed in previous works on DTE dimers.<sup>70</sup> In all of the cases, the irradiation intensities used in our photoisomerization quantum yield experiments were determined by monitoring the photocyclization and photocycloreversion processes of 1,2-bis(2-methyl-5-trifluoroacetylthien-3-yl)cyclopentene in toluene as a reference ( $\Phi_{oc}$  = 0.37 and  $\Phi_{co}$  = 0.031).33

Computational Details. DFT calculations were carried out using the Gaussian16 program package.<sup>71</sup> Geometry optimizations were conducted without any constraints using the B3LYP functional<sup>72-</sup> with Grimme's D3 correction to account for dispersion effects.75 Optimizations were performed in THF using the solvation model density (SMD) continuum model<sup>76</sup> with basis set 1 (BS1). BS1 <sup>(,78</sup> (H, included the 6-31G(d,p) basis set for the main group atoms<sup>77</sup> C, O, F, P, S) and the Stuttgart-Dresden SDD effective core potential (ECP) and its corresponding double- $\zeta$  basis set,<sup>79</sup> with a set of d polarization functions<sup>80</sup> for I and Sn and f polarization functions<sup>81</sup> for Pd. Frequency calculations were performed for all of the optimized geometries to determine the stationary points as either minima or transition states. Energies in THF were refined through single-point calculations of the optimized BS1 geometries with an extended basis set (BS2). BS2 consisted of def2-TZVP for main group atoms and the quadruple- $\zeta$  def2-QZVP basis set for Pd, together with the def2 ECP.<sup>82</sup> Gibbs energies in THF were calculated by adding thermal and entropic correction from BS1 to BS2 energies in THF. An additional correction of 1.9 kcal/mol was applied to all of the Gibbs energies to change the standard state from the gas phase (1 atm) to the condensed phase (1 M) at 298.15 K.83 Frontier molecular orbitals and natural bond orbital (NBO)<sup>84</sup> analysis were calculated at the B3LYP-D3/BS1 level in THF using the SMD continuum model. TD-DFT calculations were carried out using the CAM-B3LYP functional<sup>85</sup> with Grimme's D3 correction to account for dispersion effects.<sup>75</sup> The first 15 electronic transitions were calculated in cyclohexane using the SMD continuum model with BS1 described above.

**General Procedure for Catalytic Studies.** In an NMR tube, 0.033 mL of iodobenzene (0.30 mmol, 1.0 equiv), 0.097 mL of tributylvinyltin (0.33 mmol, 1.1 equiv), 5 mol % [Pd] catalyst, and 0.05 g of (0.03 mmol, 0.1 equiv) 1,3,5-trimethoxybenzene were dissolved in 1 mL dry, degassed THF- $d_8$ . The reactions were carried out in the dark and monitored every 2 h by <sup>1</sup>H, <sup>31</sup>P{<sup>1</sup>H}, and, when applicable, <sup>19</sup>F NMR spectroscopy. The reaction yields are an average of two replicates.

#### ASSOCIATED CONTENT

#### **Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.inorgchem.3c04423.

Synthetic procedures, additional experimental details, photochemical and catalytic studies, additional theoretical data, and NMR spectra (PDF)

# **Accession Codes**

CCDC 2310549-2310551 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data\_request/cif, or by emailing data\_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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#### **Author Contributions**

A.S. performed the experimental work, DFT calculations, and data analysis and prepared the original draft; J.H. performed data analysis, supervised the project, and prepared the original draft; P.L. performed single-crystal X-ray analysis and interpreted the structural data; A.L. supervised computational studies and revised the draft; R.M.S. and E.H.-H. supervised the project, revised the draft, and acquired funding. All authors have read and approved the final version.

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# Notes

The authors declare no competing financial interest.

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#### REFERENCES

(1) Blanco, V.; Leigh, D. A.; Marcos, V. Artificial switchable catalysts. *Chem. Soc. Rev.* 2015, 44 (15), 5341–5370.

(2) Aubert, S.; Bezagu, M.; Spivey, A. C.; Arseniyadis, S. Spatial and temporal control of chemical processes. *Nat. Rev. Chem.* **2019**, *3* (12), 706–722.

(3) Thaggard, G. C.; Haimerl, J.; Fischer, R. A.; Park, K. C.; Shustova, N. B. Traffic Lights for Catalysis: Stimuli-Responsive Molecular and Extended Catalytic Systems. *Angew. Chem., Int. Ed.* **2023**, *62* (29), No. e2023028.

(4) Teator, A. J.; Lastovickova, D. N.; Bielawski, C. W. Switchable Polymerization Catalysts. *Chem. Rev.* **2016**, *116* (4), 1969–1992.

(5) Ihrig, S. P.; Eisenreich, F.; Hecht, S. Photoswitchable polymerization catalysis: state of the art, challenges, and perspectives. *Chem. Commun.* **2019**, *55* (30), 4290–4298.

(6) Wang, F.; Zhang, Y.; Du, Z.; Ren, J.; Qu, X. Designed heterogeneous palladium catalysts for reversible light-controlled bioorthogonal catalysis in living cells. *Nat. Commun.* **2018**, *9* (1), No. 1209.

(7) Jemas, A.; Xie, Y.; Pigga, J. E.; Caplan, J. L.; am Ende, C. W.; Fox, J. M. Catalytic Activation of Bioorthogonal Chemistry with Light (CABL) Enables Rapid, Spatiotemporally Controlled Labeling and No-Wash, Subcellular 3D-Patterning in Live Cells Using Long Wavelength Light. J. Am. Chem. Soc. **2022**, 144 (4), 1647–1662.

(8) Ahn, D.; Stevens, L. M.; Zhou, K.; Page, Z. A. Rapid High-Resolution Visible Light 3D Printing. ACS Cent. Sci. 2020, 6 (9), 1555–1563.

(9) Neilson, B. M.; Bielawski, C. W. Illuminating Photoswitchable Catalysis. ACS Catal. 2013, 3 (8), 1874–1885.

(10) Göstl, R.; Senf, A.; Hecht, S. Remote-controlling chemical reactions by light: towards chemistry with high spatio-temporal resolution. *Chem. Soc. Rev.* **2014**, 43 (6), 1982–1996.

(11) Kondo, M.; Nakamura, K.; Krishnan, C. G.; Sasai, H.; Takizawa, S. Photoswitchable Chiral Organocatalysts: Photocontrol of Enantioselective Reactions. *Chem. Rec.* **2023**, *23* (7), No. e202300040.

(12) Dorel, R.; Feringa, B. L. Photoswitchable catalysis based on the isomerisation of double bonds. *Chem. Commun.* **2019**, 55 (46), 6477–6486.

(13) Liu, R.; Zhang, X.; Xia, F.; Dai, Y. Azobenzene-based photoswitchable catalysts: State of the art and perspectives. *J. Catal.* **2022**, *409*, 33–40.

(14) Majee, D.; Presolski, S. Dithienylethene-Based Photoswitchable Catalysts: State of the Art and Future Perspectives. *ACS Catal.* **2021**, *11* (4), 2244–2252.

(15) Freixa, Z. Photoswitchable catalysis using organometallic complexes. *Catal. Sci. Technol.* **2020**, *10* (10), 3122–3139.

(16) Lunic, D.; Bergamaschi, E.; Teskey, C. J. Using Light to Modify the Selectivity of Transition Metal Catalysed Transformations. *Angew. Chem., Int. Ed.* **2021**, *60* (38), 20594–20605.

(17) Galangau, O.; Norel, L.; Rigaut, S. Metal complexes bearing photochromic ligands: photocontrol of functions and processes. *Dalton Trans.* **2021**, *50* (48), 17879–17891.

(18) Medici, F.; Goual, N.; Delattre, V.; Voituriez, A.; Marinetti, A. Photoswitchable phosphines in catalysis. *ChemCatChem* **2020**, *12* (22), 5573–5589.

(19) Sud, D.; Norsten, T. B.; Branda, N. R. Photoswitching of stereoselectivity in catalysis using a copper dithienylethene complex. *Angew. Chem., Int. Ed.* **2005**, 44 (13), 2019–2021.

(20) Kean, Z. S.; Akbulatov, S.; Tian, Y.; Widenhoefer, R. A.; Boulatov, R.; Craig, S. L. Photomechanical Actuation of Ligand Geometry in Enantioselective Catalysis. *Angew. Chem., Int. Ed.* **2014**, 53 (52), 14508–14511.

(21) Zhao, D.; Neubauer, T. M.; Feringa, B. L. Dynamic control of chirality in phosphine ligands for enantioselective catalysis. *Nat. Commun.* **2015**, *6* (1), No. 6652.

(22) Cacciapaglia, R.; Di Stefano, S.; Mandolini, L. The Bis-Barium Complex of a Butterfly Crown Ether as a Phototunable Supramolecular Catalyst. J. Am. Chem. Soc. 2003, 125 (8), 2224–2227. (23) Arif, T.; Cazorla, C.; Bogliotti, N.; Saleh, N.; Blanchard, F.; Gandon, V.; Métivier, R.; Xie, J.; Voituriez, A.; Marinetti, A. Bimetallic gold(i) complexes of photoswitchable phosphines: synthesis and uses in cooperative catalysis. *Catal. Sci. Technol.* **2018**, *8* (3), 710–715.

(24) Gallarati, S.; Fabregat, R.; Juraskova, V.; Inizan, T. J.; Corminboeuf, C. How Robust Is the Reversible Steric Shielding Strategy for Photoswitchable Organocatalysts? *J. Org. Chem.* **2022**, *87* (14), 8849–8857.

(25) Irie, M.; Fukaminato, T.; Matsuda, K.; Kobatake, S. Photochromism of diarylethene molecules and crystals: memories, switches, and actuators. *Chem. Rev.* **2014**, *114* (24), 12174–12277.

(26) Neilson, B. M.; Bielawski, C. W. Photoswitchable Metal-Mediated Catalysis: Remotely Tuned Alkene and Alkyne Hydroborations. *Organometallics* **2013**, *32* (10), 3121–3128.

(27) Teator, A. J.; Shao, H.; Lu, G.; Liu, P.; Bielawski, C. W. A Photoswitchable Olefin Metathesis Catalyst. *Organometallics* **2017**, *36* (2), 490–497.

(28) Xu, Z.; Cao, Y.; Patrick, B. O.; Wolf, M. O. Photoswitching of Copper(I) Chromophores with Dithienylethene-Based Ligands. *Chem. - Eur. J.* **2018**, *24* (41), 10315–10319.

(29) Kaur, B.; Raza, R.; Stashick, M. J.; Branda, N. R. Using light to control the inhibition of Karstedt's catalyst. *Org. Chem. Front.* **2019**, *6* (8), 1253–1256.

(30) Wilson, D.; Branda, N. R. Turning "on" and "off" a pyridoxal 5'-phosphate mimic using light. *Angew. Chem., Int. Ed.* **2012**, *51* (22), 5431–5434.

(31) Majee, D.; Ramanauskaite, G.; Presolski, S. Electronic Influences on the Dynamic Range of Photoswitchable Dithienylethene–Thiourea Organocatalysts. *J. Org. Chem.* **2023**, *88* (7), 4372– 4378.

(32) Kathan, M.; Kovaříček, P.; Jurissek, C.; Senf, A.; Dallmann, A.; Thünemann, A. F.; Hecht, S. Control of Imine Exchange Kinetics with Photoswitches to Modulate Self-Healing in Polysiloxane Networks by Light Illumination. *Angew. Chem., Int. Ed.* **2016**, 55 (44), 13882–13886.

(33) Villabona, M.; Wiedbrauk, S.; Feist, F.; Guirado, G.; Hernando, J.; Barner-Kowollik, C. Dual-Wavelength Gated oxo-Diels-Alder Photoligation. *Org. Lett.* **2021**, *23* (7), 2405–2410.

(34) Stille, J. K. The Palladium-Catalyzed Cross-Coupling Reactions of Organotin Reagents with Organic Electrophiles [New Synthetic Methods (58)]. *Angew. Chem., Int. Ed.* **1986**, *25* (6), 508–524.

(35) Espinet, P.; Echavarren, A. M. The mechanisms of the Stille reaction. Angew. Chem., Int. Ed. 2004, 43 (36), 4704-4734.

(36) Cordovilla, C.; Bartolomé, C.; Martínez-Ilarduya, J. M.; Espinet, P. The Stille Reaction, 38 Years Later. *ACS Catal.* 2015, 5 (5), 3040–3053.

(37) Hansch, C.; Leo, A.; Taft, R. W. A survey of Hammett substituent constants and resonance and field parameters. *Chem. Rev.* **1991**, *91* (2), 165–195.

(38) Sponza, A. D.; Liu, D.; Chen, E. P.; Shaw, A.; Diawara, L.; Chiu, M. Synthesis strategies for non-symmetric, photochromic diarylethenes. *Org. Biomol. Chem.* **2020**, *18* (37), 7238–7252.

(39) Bianchini, G.; Strukul, G.; Wass, D. F.; Scarso, A. Photomodulable phosphines incorporating diarylethene moieties. *RSC Adv.* **2015**, 5 (14), 10795–10798.

(40) Chevykalova, M. N.; Manzhukova, L. F.; Artemova, N. V.; Luzikov, Yu. N.; Nifant'ev, I. E.; Nifant'ev, E. E. Electron-donating ability of triarylphosphines and related compounds studied by <sup>31</sup>P NMR spectroscopy. *Russ. Chem. Bull.* **2003**, *52* (1), 78–84.

(41) Sud, D.; McDonald, R.; Branda, N. R. Synthesis and coordination chemistry of a photoswitchable bis(phosphine) ligand. *Inorg. Chem.* **2005**, *44* (17), 5960–5962.

(42) Yin, J.; Lin, Y.; Cao, X.; Yu, G.-A.; Tu, H.; Liu, S. H. The synthesis and photochromic properties of two bis(phosphine) ligands based on dithienylethene backbone and their oxides, sulfurets and selenides. *Dyes Pigm.* **2009**, *81* (2), 152–155.

(43) Kobatake, S.; Uchida, K.; Tsuchida, E.; Irie, M. Singlecrystalline photochromism of diarylethenes: reactivity-structure relationship. *Chem. Commun.* **2002**, No. 23, 2804–2805.

(44) Herder, M.; Eisenreich, F.; Bonasera, A.; Grafl, A.; Grubert, L.; Pätzel, M.; Schwarz, J.; Hecht, S. Light-Controlled Reversible Modulation of Frontier Molecular Orbital Energy Levels in Trifluoromethylated Diarylethenes. *Chem. - Eur. J.* **2017**, 23 (15), 3743–3754.

(45) Gilat, S. L.; Kawai, S. H.; Lehn, J.-M. Light-triggered electrical and optical switching devices. J. Chem. Soc., Chem. Commun. 1993, No. 18, 1439–1442.

(46) Liang, J.; Yin, J.; Li, Z.; Zhang, C.; Wu, Di.; Liu, S. H. Synthesis and properties of dithienylethene-based binuclear gold complexes and a palladium chlorine-bridged macrocycle. *Dyes Pigm.* **2011**, *91* (3), 364–369.

(47) Lasorne, B.; Fihey, A.; Mendive-Tapia, D.; Jacquemin, D. A curve-crossing model to rationalize and optimize diarylethene dyads. *Chem. Sci.* **2015**, *6* (10), 5695–5702.

(48) Herder, M.; Schmidt, B. M.; Grubert, L.; Pätzel, M.; Schwarz, J.; Hecht, S. Improving the fatigue resistance of diarylethene switches. *J. Am. Chem. Soc.* **2015**, *137* (7), 2738–2747.

(49) Allen, D. W.; Taylor, B. F. The Chemistry of Heteroarylphosphorus Compounds. Part 15, Phosphorus-31 Nuclear Magnetic Resonance Studies of the Donor Properties of Heteroarylphosphines towards Selenium and Platinum(II). J. Chem. Soc., Dalton Trans. 1982, No. 1, 51–54.

(50) Bent, H. A. An Appraisal of Valence-bond Structures and Hybridization in Compounds of the First-row elements. *Chem. Rev.* **1961**, *61* (3), 275–311.

(51) Levin, C. C. Qualitative molecular orbital picture of electronegativity effects on  $XH_3$  inversion barriers. *J. Am. Chem. Soc.* **1975**, 97 (20), 5649–5655.

(52) Glendening, E. D.; Landis, C. R.; Weinhold, F. Natural bond orbital methods. *Wiley Interdiscip. Rev. Comput. Mol. Sci.* 2012, 2 (1), 1–42.

(53) Casado, A. L.; Espinet, P.; Gallego, A. M. Mechanism of the Stille Reaction. 2. Couplings of Aryl Triflates with Vinyltributyltin. Observation of Intermediates. A More Comprehensive Scheme. J. Am. Chem. Soc. 2000, 122 (48), 11771–11782.

(54) Casado, A. L.; Espinet, P. Mechanism of the Stille Reaction. 1. The Transmetalation Step. Coupling of  $R_1I$  and  $R_2SnBu_3$  Catalyzed by trans-[PdR<sub>1</sub>IL<sub>2</sub>] ( $R_1 = C_6Cl_2F_3$ ;  $R_2 =$  Vinyl, 4-Methoxyphenyl; L = AsPh<sub>3</sub>). J. Am. Chem. Soc. **1998**, 120, 8978–8985.

(55) Meijere, A. de.; Diederich, F. Metal-catalyzed cross-coupling Reactions, 2nd, completely rev. and enl. ed.; Wiley-VCH, 2004.

(56) Nova, A.; Ujaque, G.; Maseras, F.; Lledós, A.; Espinet, P. A critical analysis of the cyclic and open alternatives of the transmetalation step in the stille cross-coupling reaction. *J. Am. Chem. Soc.* **2006**, *128* (45), 14571–14578.

(57) García-Melchor, M.; Braga, A. A. C.; Lledós, A.; Ujaque, G.; Maseras, F. Computational perspective on Pd-catalyzed C-C crosscoupling reaction mechanisms. *Acc. Chem. Res.* **2013**, *46* (11), 2626– 2634.

(58) Echavarren, A. M.; Cárdenas, D. J. Mechanistic Aspects of Metal-Catalyzed C,C- and C,X-Bond-Forming Reactions: 1. In *Metal-Catalyzed Cross-Coupling Reactions*; John Wiley & Sons, Ltd, 2004; pp 1–40 DOI: 10.1002/9783527619535.ch1.

(59) McMullin, C. L.; Fey, N.; Harvey, J. N. Computed ligand effects on the oxidative addition of phenyl halides to phosphine supported palladium(0) catalysts. *Dalton Trans.* **2014**, *43* (36), 13545–13556.

(60) McMullin, C. L.; Jover, J.; Harvey, J. N.; Fey, N. Accurate modelling of Pd(0) + PhX oxidative addition kinetics. *Dalton Trans.* **2010**, 39 (45), 10833–10836.

(61) Lam, K. C.; Marder, T. B.; Lin, Z. DFT Studies on the Effect of the Nature of the Aryl Halide  $Y-C_6H_4-X$  on the Mechanism of Its Oxidative Addition to  $Pd^0L$  versus  $Pd^0L_2$ . Organometallics **2007**, 26 (3), 758–760.

(62) Lyngvi, E.; Sanhueza, I. A.; Schoenebeck, F. Dispersion Makes the Difference: Bisligated Transition States Found for the Oxidative Addition of  $Pd(PtBu_3)_2$  to Ar-OSO<sub>2</sub>R and Dispersion-Controlled Chemoselectivity in Reactions with  $Pd[P(iPr)(tBu_2)]_2$ . Organometallics **2015**, 34 (5), 805–812.

(63) Rigaku Oxford Diffraction. CrysAlisPro Software system; Rigaku Corporation: Wroclaw, Poland, 1995–2023.

(64) Clark, R. C.; Reid, J. S. The analytical calculation of absorption in multifaceted crystals. *Acta Crystallogr. A* **1995**, *51* (6), 887–897.

(65) Sheldrick, G. M. SHELXT - Integrated space-group and crystalstructure determination. *Acta Crystallogr. A* **2015**, *71* (1), 3–8.

(66) DIAMOND 4; K. Brandenburg, Crystal Impact GbR: Bonn, Germany.

(67) Macrae, C. F.; Bruno, I. J.; Chisholm, J. A.; Edgington, P. R.; McCabe, P.; Pidcock, E.; Rodriguez-Monge, L.; Taylor, R.; van de Streek, J.; Wood, P. A. Mercury CSD 2.0 - new features for the visualization and investigation of crystal structures. *J. Appl. Crystallogr.* **2008**, *41* (2), 466–470.

(68) Lees, A. J. A Photochemical Procedure for Determining Reaction Quantum Efficiencies in Systems with Multicomponent Inner Filter Absorbances. *Anal. Chem.* **1996**, *68* (1), 226–229.

(69) Ordronneau, L.; Aubert, V.; Métivier, R.; Ishow, E.; Boixel, J.; Nakatani, K.; Ibersiene, F.; Hammoutène, D.; Boucekkine, A.; Le Bozec, H.; Guerchais, V. Tunable double photochromism of a family of bis-DTE bipyridine ligands and their dipolar Zn complexes. *Phys. Chem. Chem. Phys.* **2012**, *14* (8), 2599–2605.

(70) Fihey, A.; Perrier, A.; Browne, W. R.; Jacquemin, D. Multiphotochromic molecular systems. *Chem. Soc. Rev.* 2015, 44 (11), 3719–3759.

(71) Frisch, M. J.; Trucks, G. W.; Schlegel, H. B.; Scuseria, G. E.; Robb, M. A.; Cheeseman, J. R.; Scalmani, G.; Barone, V.; Petersson, G. A.; Nakatsuji, H.; Li, X.; Caricato, M.; Marenich, A. V.; Bloino, J.; Janesko, B. G.; Gomperts, R.; Mennucci, B.; Hratchian, H. P.; Ortiz, J. V.; Izmaylov, A. F.; Sonnenberg, J. L.; Williams-Young, D.; Ding, F.; Lipparini, F.; Egidi, F.; Goings, J.; Peng, B.; Petrone, A.; Henderson, T.; Ranasinghe, D.; Zakrzewski, V. G.; Gao, J.; Rega, N.; Zheng, G.; Liang, W.; Hada, M.; Ehara, M.; Toyota, K.; Fukuda, R.; Hasegawa, J.; Ishida, M.; Nakajima, T.; Honda, Y.; Kitao, O.; Nakai, H.; Vreven, T.; Throssell, K.; Montgomery, J. A., Jr.; Peralta, J. E.; Ogliaro, F.; Bearpark, M. J.; Heyd, J. J.; Brothers, E. N.; Kudin, K. N.; Staroverov, V. N.; Keith, T. A.; Kobayashi, R.; Normand, J.; Raghavachari, K.; Rendell, A. P.; Burant, J. C.; Iyengar, S. S.; Tomasi, J.; Cossi, M.; Millam, J. M.; Klene, M.; Adamo, C.; Cammi, R.; Ochterski, J. W.; Martin, R. L.; Morokuma, K.; Farkas, O.; Foresman, J. B.; Fox, D. J. Gaussian 16, Revision B.01 2016.

(72) Lee, C.; Yang, W.; Parr, R. G. Development of the Colle-Salvetti correlation-energy formula into a functional of the electron density. *Phys. Rev. B* **1988**, 37 (2), 785–789.

(73) Miehlich, B.; Savin, A.; Stoll, H.; Preuss, H. Results obtained with the correlation energy density functionals of Becke and Lee, Yang and Parr. *Chem. Phys. Lett.* **1989**, *157* (3), 200–206.

(74) Becke, A. D. Density-functional thermochemistry. III. The role of exact exchange. J. Chem. Phys. **1993**, 98 (7), 5648–5652.

(75) Grimme, S.; Antony, J.; Ehrlich, S.; Krieg, H. A consistent and accurate ab initio parametrization of density functional dispersion correction (DFT-D) for the 94 elements H-Pu. *J. Chem. Phys.* 2010, 132 (15), No. 154104.

(76) Marenich, A. V.; Cramer, C. J.; Truhlar, D. G. Universal Solvation Model Based on Solute Electron Density and on a Continuum Model of the Solvent Defined by the Bulk Dielectric Constant and Atomic Surface Tensions. *J. Phys. Chem. B* 2009, *113* (18), 6378–6396.

(77) Hehre, W. J.; Ditchfield, R.; Pople, J. A. Self—Consistent Molecular Orbital Methods. XII. Further Extensions of Gaussian— Type Basis Sets for Use in Molecular Orbital Studies of Organic Molecules. J. Chem. Phys. **1972**, 56 (5), 2257–2261.

(78) Francl, M. M.; Pietro, W. J.; Hehre, W. J.; Binkley, J. S.; Gordon, M. S.; DeFrees, D. J.; Pople, J. A. Self-consistent molecular orbital methods. XXIII. A polarization-type basis set for second-row elements. J. Chem. Phys. **1982**, 77 (7), 3654–3665.

(79) Andrae, D.; Häußermann, U.; Dolg, M.; Stoll, H.; Preuß, H. Energy-adjusted *ab initio* pseudopotentials for the second and third row transition elements. *Theor. Chem. Acc.* **1990**, 77 (2), 123–141.

(80) Höllwarth, A.; Böhme, M.; Dapprich, S.; Ehlers, A. W.; Gobbi, A.; Jonas, V.; Köhler, K. F.; Stegmann, R.; Veldkamp, A.; Frenking, G. A set of d-polarization functions for pseudo-potential basis sets of the main group elements Al-Bi and f-type polarization functions for Zn, Cd, Hg. Chem. Phys. Lett. **1993**, 208 (3), 237–240.

(81) Ehlers, A. W.; Böhme, M.; Dapprich, S.; Gobbi, A.; Höllwarth, A.; Jonas, V.; Köhler, K. F.; Stegmann, R.; Veldkamp, A.; Frenking, G. A set of f-polarization functions for pseudo-potential basis sets of the transition metals Sc-Cu, Y-Ag and La-Au. *Chem. Phys. Lett.* **1993**, 208 (1), 111–114.

(82) Weigend, F.; Ahlrichs, R. Balanced basis sets of split valence, triple zeta valence and quadruple zeta valence quality for H to Rn: Design and assessment of accuracy. *Phys. Chem. Chem. Phys.* **2005**, 7 (18), 3297–3305.

(83) Bryantsev, V. S.; Diallo, M. S.; Goddard, W. A., III Calculation of Solvation Free Energies of Charged Solutes Using Mixed Cluster/ Continuum Models. J. Phys. Chem. B 2008, 112 (32), 9709–9719.

(84) Glendening, E. D.; Reed, A. E.; Carpenter, J. E.; Weinhold, F. NBO Version 3.1.

(85) Yanai, T.; Tew, D. P.; Handy, N. C. A new hybrid exchange– correlation functional using the Coulomb-attenuating method (CAM-B3LYP). *Chem. Phys. Lett.* **2004**, 393 (1), 51–57.