RSC Advances



PAPER

Check for updates

Cite this: RSC Adv., 2023, 13, 29689

Received 4th July 2023 Accepted 19th September 2023

DOI: 10.1039/d3ra04460b

rsc.li/rsc-advances

Introduction

Among multi-palladium composed complexes, aromatic triangular tri-palladium complexes $[LPd]_3^+$ (L = ligand)¹⁻⁵ and their heteroaromatics^{6,7} are excellent species as the metallic analogs of cyclopropene positive ion $[C_3H_3]^+$. The unique allmetal aromaticity and excellent stability endow such three-membered palladium cores with excellent molecular stability and Lewis alkalinity, which has brought them much attention in theoretical chemistry,⁸⁻¹¹ coordination chemistry,¹² catalytic reactions¹³⁻²² and nanomaterials.²³

Compared with the usual negative ions like SbF₆⁻, PF₆⁻, BF₄⁻ and CF₃COO⁻, from the perspective of electronic factors of ion pairs for organometallic complexes, aryl sulfonate anions (ArSO₃⁻) were recognized as much weaker σ -donors and usually present milder coordination abilities with transition metals,²⁴⁻²⁶ which could potentially bring distinctive structural and characteristic properties for aromatic tripalladium frameworks. As the foundation of this work, the Zwitterionic analogue reflected high stability, catalytic activity and stereoselectivity in the selective reduction of alkynes.²⁷ While, the full crystal structures and properties were still undiscovered perhaps owing to the reduced symmetry. In this work, we designed and synthesized a series of *C*₃-symmetry aromatic triangular cations 3–7 (Scheme 1)

Aryl sulfonate anion stabilized aromatic triangular cation [Pd₃]⁺: syntheses, structures and properties †

Miaomiao Wang, Zhixin He, Meng Chen and Yanlan Wang D*

A series of sulfonate anions paired aromatic triangular palladium clusters **3**–**7**, abbreviated as $[Pd_3]^+[ArSO_3]^-$, were synthesized using a simple "one pot" method, and gave excellent isolated yields (90-95%). Their structures and properties have been fully characterized and further investigated by fluorescence, single crystal X-ray Diffraction (XRD) and X-ray Photoelectron Spectroscopy (XPS). In varying organic solvents, they presented apparently stronger absorption and emission in MeOH, driven by the combined interactions of hydrogen bonds and polarity. The crystallographic data demonstrated that the methyl orange ion stabilized complex **7** possessed a D_{3h} symmetric metallic core which was still coplanar and almost equilateral, jointly influenced by the giant hindrance and milder donating effect from the sulfonate. The binding energies for $Pd^{n+} 3d_{5/2}$ and $Pd^{n+} 3d_{3/2}$ measured by XPS presented at 336.55 and 342.00 eV, respectively. These data were much lower than that of a usual $Pd^{2+} 3d$ and significantly higher than that of a Pd^0 species, further proving the unified palladium valence state (+4/3) in the tri-palladium core and its aromaticity featured by the cyclic electron delocalization.

stabilized by substituted aryl sulfonate anions and revealed their extraordinary structures and properties.

Results and discussion

Aryl silver sulfonates 2

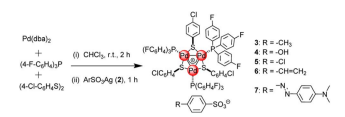
Even the aromatic sulfonates sodium salts are more commercially available, their corresponding silver sulfonates analogues are experimentally proved to be much more effective under identical conditions. Besides the commercial silver 4-methylbenzenesulfonate (**2e**), we synthesized the other employed substituted silver benzenesulfonates (**2a–d**) through two pathways (see Scheme S1†) according to literature,^{28–31} The employed silver salts were prepared either by mixing corresponding sulfonic acids with Ag₂CO₃ under dark (for **2a**, **2b**) or reacting their sodium sulfonates with AgNO₃ (for **2c**, **2d**), gave the isolated yields in the range of 74–99%.

Tri-palladium complexes 3-7

The aryl sulfonate anion stabilized new aromatic triangular tripalladium complexes 3–7 were synthesized using a facile "one pot" method (Scheme 1). Simply mixed $[Pd(dba)_2]$ (1.0 equiv.), tri-4-fluorophenylphosphate (1.0 equiv.), and 4-chlorophenyldisulfide (0.5 equiv.) in chloroform and kept stirring for two hours under N₂. Hitherto, the original purple solution became deep red and the trimetallic cationic core was proposed to be formed by then, while, adding the prepared silver benzenesulfonates (2a–e) (0.033 equiv.) was critical for the ions exchanges which were crucial for the subsequent facile purification and high isolated yields (90–95%). For comparisons, we prepared five

Department of Chemistry and Chemical Engineering, Liaocheng University, 252059 Liaocheng, China. E-mail: wangyanlan@lcu.edu.cn

[†] Electronic supplementary information (ESI) available. CCDC 2270588. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d3ra04460b



Scheme 1 Syntheses of substituted aryl sulfonates stabilized tripalladium complexes 3–7.

analogues tri-palladium complexes, including 3 stabilized by 4methylbenzenesulfonate, 4 stabilized by 4-hydroxybenzenesulfonate, 5 stabilized by 4-chlorobenzenesulfonate, 6 stabilized by 4vinylbenzenesulfonate and 7 stabilized by (*E*)-(((4-((dimethylamino)phenyl)diazenyl)phenyl)sulfonyl)oxy), abbreviated as methyl orange anion. These air-stable deep red solids were obtained by filtration and repeated precipitations using CHCl₃/hexane. The purified products 3–7 were fully characterized by ¹H, ¹³C, ³¹P, ¹⁹F NMR, ESI-MS, UV-visible, X-ray Photoelectron Spectroscopy (XPS) and single crystal X-ray Diffractions (XRD).

High resolution mass spectroscopies (HRMS)

The calculated high resolution mass for the tri-palladium cation in complexes 3–7 was 1698.7048, for example, the founded mass for 3 was 1698.7062 by ESI-MS, which was perfectly consistent with the calculated value (Fig. 1a). As the cationic fragment was identical and the found mass for this part were extremely close for all these complexes. In addition, the counter anions 4-

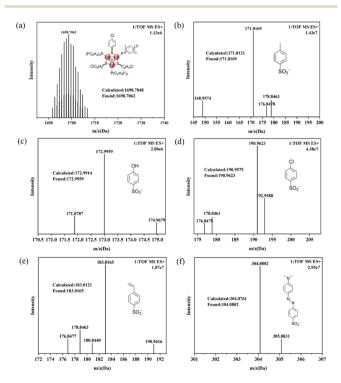


Fig. 1 (a) The ESI-MS for cations of complexes 3; (b) anion for complex 3; (c) anion for complex 4; (d) anion for complex 5; (e) anion for complex 5; (f) anion for complex 7.

methylbenzenesulfonate in 3 (Fig. 1b), 4-hydroxybenzenesulfonate in 4 (Fig. 1c), 4-chlorobenzenesulfonate in 5 (Fig. 1d), 4-vinylbenzenesulfonate in 6 (Fig. 1e) and methyl orange anion in 7 (Fig. 1f) were found at 171.0169, 172.9959, 190.9623, 183.0165 and 304.0802, respectively, which were also fully consistent with calculated values. The above perfect cationic and anionic modes for complexes 3–7 by ESI-MS definitely proved the correct combination and structural integrity.

Single crystal X-ray diffraction

The deep-red single crystal suitable for X-ray diffraction for the most representative complex 7 was obtained by vapor diffusion method. From the plane defined by the three Pd atoms and the plane defined by the aromatic ring in the counter ion (Fig. 2), the structure and orientation for the triangular tri-palladium cation and the methyl orange anion could be undoubtedly confirmed. Noteworthy, the Pd-Pd-Pd bond angles for 7 are 61.11(2)°, 59.24(2)° and 59.65(2)° which were slightly deviated from the examples in the literature (59.5–60.5°).^{5,6} This deviation might be caused by the first introduced anionic benzenesulfonate which probably contributed larger spatial site resistance and milder coordination modes. Even though, the palladium triangles were almost equilateral. The three Pd-Pd bond lengths are 2.8458(10) Å. 2.8577(9) Å and 2.8996(10) Å, respectively, which were much below the sum of van der Waals radii (3.26 Å) for metal-metal bonding and confirmed the Pd-Pd bonding modes.³² Compared to the tri-palladium analogue stabilized by SbF_6^{-6} , the cationic part for complex 7 did not exactly coincide with C_3 symmetry and possessed a D_{3h} symmetric core. Some of the benzene rings on the coordinated S/P ligands in complex 7 tilted to the opposite direction to the negative methyl orange ion mainly due to low energy interactions are preferred when the cluster contains bulky aromatic moieties (Fig. S6[†]). Combined with the NMR and ESI-MS values, these crystal data further proved that our synthetic strategy allowed for the successful construction and isolation for these aryl sulfonate anion paired aromatic triangular tri-palladium cations.

Absorption and emission

In order to understand the luminescent properties for the aryl sulfonate anion stabilized new aromatic triangular tripalladium complexes 3–7, we investigated their absorptions and fluorescence emissions. Under UV-visible light

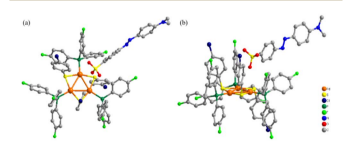


Fig. 2 X-ray diffraction for complex 7 with 30% probability ellipsoids. Hydrogen atoms have been omitted for the sake of clarity. (a) Plane defined by the three Pd atoms, (b) plane defined by the aromatic ring in the counterion.

irradiation, all these complexes showed two distinct absorption peaks at around 245 nm and 400 nm in CH_2Cl_2 solutions, which could be ascribed to the occurrence of π - π^* and n- π^* transitions of phosphine and sulfur ligands (Fig. 3a).^{33,34} For example, 4-methylbenzenesulfonate paired complex 3 presented the maximum absorption at the wavelength of 399 nm. While, the presence of N=N group and the enlarged conjugation area in the methyl orange anion stabilized complex 7 caused red shift (to 404 nm), and the absorption intensity became much stronger than complex 3 (increased about 30%). So, the substituted aryl sulfonate anions possessing different types of chromophores and auxochromes indeed brought variations in the maximum absorption wavelength and absorption intensities under identical conditions.³⁵ For comparison, we further measured the absorption for

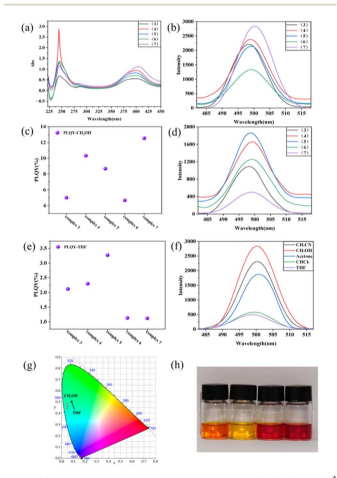


Fig. 3 (a) UV-visible spectra for complexes 3–7 in CH₂Cl₂ (1×10^{-4} M); (b) fluorescence emissions for complexes 3–7 in CH₃OH (1×10^{-4} M); (c) PLQY for complexes 3–7 (5.00%, 10.33%, 8.68%, 4.66%, 12.55%, respectively) in CH₃OH at room temperature (1×10^{-4} M); (d) fluorescence emissions for complexes 3–7 in THF (1×10^{-4} M); (e) PLQY for complexes 3–7 (2.12%, 2.30%, 3.27%, 1.13%, 1.12%, respectively) in THF at room temperature (1×10^{-4} M); (f) fluorescence emissions for complexe 3–7 (2.12%, 2.30%, 3.27%, 1.13%, 1.12%, respectively) in THF at room temperature (1×10^{-4} M); (f) fluorescence emissions for complex 7 (1×10^{-4} M) in different organic solvents (CH₃OH, CH₃-COCH₃, CHCl₃, CH₃CN, THF); (g) the CIE coordinates for complex 7 (1×10^{-4} M) from THF to CH₃OH; (h) from left to right: color of methyl orange-sodium (1d), methyl orange-silver (2d), complex 3, complex 7 in DMF.

complexes (3–7) in MeOH (Fig. S8†), their absorption intensity presented obvious solvent dependence. Interestingly, complex 7 also presented red shift compared with complex 3 when MeOH was used as the solvent.

For better understanding their luminescence properties, we further examined the fluorescence emission for all complexes 3-7 in polar/protic CH₃OH (Fig. 3b), or less polar/ nonprotic THF (Fig. 3d). The fluorescence quantum yields (PLQY) for complexes 3-7 in CH₃OH were 5.00%, 10.33%, 8.68%, 4.66%, and 12.55%, respectively (Fig. 3c). Among these analogues, the methyl orange anion stabilized complex 7 exhibited the highest PLQY in the polar CH₃OH. Interestingly, the PLQY for all these complexes were dropped sharply in less polar THF and complex 7 presented the lowest value (Fig. 3e), which was roughly consistent with their emission intensities. For better comparison, we also measured the PLQY for these clusters in (CH₃COCH₃, CHCl₃ and CH₃CN) and similar moderate values were obtained (Fig. S10[†]). It was worth noting that all these complexes exhibited higher PLQY in polar CH₃OH or CH₃CN than less polar solvents (CH₃-COCH₃, CHCl₃ and THF). This result could be attributed by the complex strength of solvent polarity and the possible hydrogen bonding interactions between the negative ions and protic CH₃OH.

Besides the PLQY, the emission intensity of complex 7 also varied significantly when the solvents polarity changes with the excitation wavelength set at 250 nm (Fig. 3f). This result could be attributable to that the polarity of solvents could significantly impact the energies of π - π * and n- π * transitions, therefore, the fluorescence intensity of substances may also change as the polarity of the solvent varies.³⁶ Basically, the excited states possessed greater polarity and was more beneficial from increasing solvent polarity.38-40 Generally, complex 7 conformed to the rule of fluorescence intensity enhancement with increasing solvent polarity and presented apparently stronger emission in CH₃OH. Meanwhile, the fluorescence wavelengths also presented red shift when the polarity of organic solvents were increased. This clear difference was also probably caused by the interactions of hydrogen bonds with hydroxyl in CH₃OH.³⁷ Overall, the CIE coordinates of complex 7 changed from blue to teal with the increase of solvent polarity (Fig. 3g). Besides 7, the luminescence properties for complexes 3-6 were also examined in different solvents, and obvious varying for emission Intensities and wavelengths were observed, probably caused by the introduction of different groups on the anionic aryl sulfonates (see ESI, Fig. S5[†]).

In addition, the color comparisons for methyl orangesodium, methyl orange-silver (2d), complex 3 or 7 in DMF were presented (Fig. 3h). Different with the commercial sodium analogue, the synthesized silver salt 2d showed a deeper red color as solid but decreased solubility in water or DMF, resulting in yellow color in solutions. Comparing complexes 3 and 7, we found that the methyl orange anion introduced complex 7 showed more pronounced red color, which was probably caused by the color enhancement effect from the dye-based anionic fragment.⁴¹

X-ray photoelectron spectroscopy (XPS)

In order to know the bonding forms and valence states for these aryl sulfonate anions stabilized new aromatic triangular tripalladium complexes 3-7, we further examined them by XPS. Taking complex 7 as an example, the observed binding energies for their involved elements: Pd, C, O, S, Cl, P, N, and F presented to be completely consistent with atoms in analogous structures in literature49 and definitely manifested their existence by comparing with established values (Fig. 4a). In addition, the high-resolution XPS spectrum for C 1s in complex 7 presented five characteristic peaks at 284.00, 284.35, 284.60, 286.75, and 290.90 eV (Fig. 4b), which were ascribed to the sp^2 carbon, sp^3 carbon, C-P, C-Cl, and C-F, respectively.42,43 Among them, the presence of bonding energies for sp³ hybridized carbons which belonged to the two methyl groups on the methyl orange anion illustrated the intimate paring between the sulfonate anion and the tri-palladium cation.

Theoretically, the Pd^{2+} , Pd^{n+} (0 < n < 2) and Pd^{0} species have relatively fixed and different binding energies according to literature.⁴⁴ As expected, the XPS for Pd 3d in complex 7 showed two peaks at 336.55 and 342.00 eV, which were certainly assigned to $Pd^{n+} 3d_{5/2}$ and $Pd^{n+} 3d_{3/2}$, respectively (Fig. 4c). The found Pd $3d_{5/2}$ bond energies for cyclic tri-palladium core $[Pd_3]^+$ was much lower than $Pd^{2+} 3d_{5/2}$ (about 337.60 eV) and significantly higher than that of $Pd^{0} 3d_{5/2}$ in species like nanoparticles (about 335.00 eV).^{45,46} Similarly, the Pd $3d_{3/2}$ bond energies for metallic core $[Pd_3]^+$ was perfectly appeared between the $Pd^{2+} 3d_{3/2}$ (about 342.91 eV) and $Pd^{0} 3d_{3/2}$ (about 341.75 eV).^{47,48} So,

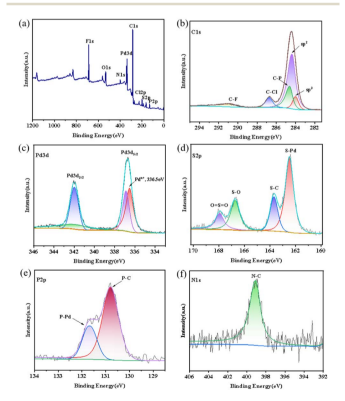


Fig. 4 (a) XPS survey spectrum for complex 7, high-resolution for (b) C 1s, (c) Pd 3d, (d) S 2p, (e) N 1s, (f) P 2p spectra in complex 7.

our recorded XPS data certainly proved the unified palladium valence state (+4/3) in these tri-palladium cation $[Pd_3]^+$ and their aromaticity characterized by definite cyclic electron delocalization.⁴⁹

In addition, the high-resolution XPS for S2p in complex 7 presented four peaks at 162.50, 163.70, 166.70, and 167.95 eV, responding to the S–Pd, S–C, S–O, and O=S=O, respectively (Fig. 4d). As expected, the XPS spectrum for N 1s showed only one peak at 399.10 eV (Fig. 4e), representing the unique N–C bonding in the anionic fragment. Clearly, the XPS spectrum for P2p showed two peaks at 130.8 and 131.70 eV, assignable to the P–C and P–Pd bonding modes, respectively (Fig. 4f).⁵⁰ Overall, the above measured electron binding energies by XPS are perfectly in line with the high resolution mass and single crystal X-ray diffraction results, definitely revealed the chemical compositions, bonding forms, and valence states for elements in these aryl sulfonates stabilized aromatic triangular tripalladium complexes.

Conclusions

In this work, we synthesized a series of new aromatic triangular tri-palladium complexes 3-7 stabilized by substituted aryl sulfonate anions, and investigated their structures and properties. Besides the methyl orange anion, several functional groups (methyl, hydroxy, chloro, vinyl) were successfully introduced to the benzenesulfonates. Firstly, the cationic and anionic modes for these complexes by ESI-MS were fully consistent with calculated values and confirmed the structural integrity. Secondly, the single crystal X-ray diffraction confirmed that complex 7 possessed a D_{3h} symmetric metallic core and became less C3 symmetry compared to its hexafluoroantimonate analogue due to its increased steric effect and weaker σ -donating effect from the negative methyl orange ion. While, the three palladium atoms were still coplanar and the palladium triangles were almost equilateral, especially, the short Pd-Pd bond lengths were supportive for metal-metal bonding. Thirdly, the absorption and fluorescence properties for these complexes were examined in different organic solvents, and presented apparently stronger absorption and emissions in CH₃OH, caused by the interactions of hydrogen bonds and solvent polarity. Finally, the measured XPS for Pd 3d in complex 7 showed two peaks at 336.55 and 342.00 eV, which were assigned to $Pd^{n+} 3d_{5/2}$ and $Pd^{n+} 3d_{3/2}$, respectively. This recorded XPS data certainly proved the unified palladium valence state (+4/3) in these ArSO3⁻ paired aromatic tripalladium complexes.

Overall, these experimental results are consistent with the formulation and cognition that the metallic core $[Pd_3]^+$ in these complexes are aromatic framework. The milder coordination effect between the aryl sulfonates anions and the tri-palladium cations endows their paired complexes with good stability and stronger Lewis-alkalinity. Therefore, the introduction of aryl sulfonates to these new triangular aromatic palladium complexes could potentially realize their practical applications in the fields of coordination chemistry, catalytic reactions, and functionalization for macromolecular materials.

Author contributions

MW, ZH performed the synthetic chemistry and measured the NMR, HRMS, and luminescence data. MW collected and solved the XRD data. MW and MC collected and analysed the XPS data. YW and MW conceived the idea. All authors wrote/edited the manuscript.

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgements

We thank for National Natural Science Foundation of China (Grant No. 21901097); Natural Science Foundation of Shandong province (Grant No. ZR2023MB075); Science and Technology Foundation of Liaocheng University (Grant No. 318051728); Introduction and Cultivation Project for Young Innovative Talents in Shandong Provincial Colleges and Universities (Innovation Team of Functional Organometallic Materials Presided by Prof. Yanlan Wang).

Notes and references

- 1 A. I. Boldyrev and L.-S. Wang, *Chem. Rev.*, 2005, **105**, 3716–3757.
- 2 J. M. Mercero, A. I. Boldyrev, G. Merino and J. M. Ugalde, *Chem. Soc. Rev.*, 2015, 44, 6519–6534.
- 3 N. V. Tkachenko, I. A. Popov, M. Kulichenko, N. Fedik, Z.-M. Sun, A. Muñoz-Castro and A. I. Boldyrev, *Eur. J. Inorg. Chem.*, 2021, **41**, 4239–4250.
- 4 F. Bigi, D. Cauzzi, N. Della Ca, M. Malacria, R. Maggi, E. Motti, Y. Wang and G. Maestri, *ACS Org. Inorg. Au*, 2022, 2, 373–385.
- 5 S. Blanchard, L. Fensterbank, G. Gontard, E. Lacôte,
 G. Maestri and M. Malacria, *Angew. Chem., Int. Ed.*, 2014, 126, 2018–2022.
- 6 Y. Wang, P.-A. Deyris, T. Caneque, F. Blanchard, Y. Li, F. Bigi, R. Maggi, S. Blanchard, G. Maestri and M. Malacria, *Chem. -Eur. J.*, 2015, **21**, 12271–12274.
- 7 M. Kulichenko, N. Fedik, A. Monfredini, A. Muñoz-Castro, D. Balestri, A. I. Boldyrev and G. Maestri, *Chem. Sci.*, 2021, 12, 477–486.
- 8 F. Bigi, G. Cera, R. Maggi, Y. Wang, M. Malacria and G. Maestri, *J. Phys. Chem. A*, 2021, **125**, 10035–10043.
- 9 I. A. Popov, A. A. Starikova, D. V. Steglenko and A. I. Boldyrev, *Chem. Eur. J.*, 2018, **24**, 292–305.
- 10 X. R. You and H. J. Zhai, Chin. J. Chem., 2019, 37, 126-130.
- 11 P. von R. Schleyer and R. B. King, *Chem.-Eur. J.*, 2007, 13, 978–984.
- 12 Y. Wang, A. Monfredini, P.-A. Deyris, F. Blanchard, E. Derat, G. Maestri and M. Malacria, *Chem. Sci.*, 2017, **8**, 7394–7402.

- 13 P.-A. Deyris, T. Caneque, Y. Wang, P. Retailleau, F. Bigi, R. Maggi, G. Maestri and M. Malacria, *ChemCatChem*, 2015, 7, 3266–3269.
- 14 A. Monfredini, V. Santacroce, L. Marchiò, R. Maggi, F. Bigi,
 G. Maestri and M. Malacria, ACS Sustain. Chem. Eng., 2017,
 5, 8205–8212.
- 15 M. Lanzi, T. Cañeque, L. Marchiò, R. Maggi, F. Bigi, M. Malacria and G. Maestri, ACS Catal., 2018, 8, 144–147.
- 16 X. Li, J. Li, X. Wang, L. Wu, Y. Wang, G. Maestri, M. Malacria and X. Liu, *Dalton Trans.*, 2021, **50**, 11834–11842.
- 17 F. Fu, J. Xiang, H. Cheng, L. Cheng, H. Chong, S. Wang and Y. Li, *ACS Catal.*, 2017, 7, 1860–1867.
- 18 C. J. Diehl, T. Scattolin, U. Englert and F. Schoenebeck, *Angew. Chem., Int. Ed.*, 2019, **58**, 211–215.
- 19 C. Cecchini, M. Lanzi, G. Cera, M. Malacria and G. Maestri, *Synthesis*, 2019, **51**, 1216–1224.
- 20 A. Serafino, N. Camedda, M. Lanzi, N. Della Ca, G. Cera and G. Maestri, *J. Org. Chem.*, 2021, 86, 15433–15452.
- X. Wang, L. Sun, M. Wang, G. Maestri, M. Malacria, X. Liu,
 Y. Wang and L. Wu, *Eur. J. Org Chem.*, 2022, 10, e20220.
- N. W. J. Scott, M. J. Ford, N. Jeddi, A. Eyles, L. Simon,
 A. C. Whitwood, T. Tanner, C. E. Willans and
 I. J. S. Fairlamb, *J. Am. Chem. Soc.*, 2021, 143, 9682–9693.
- 23 J. Ren, P. C. Lan, M. Chen, W. Zhang and S. Ma, *Organometallics*, 2019, **38**, 3460–3465.
- 24 S. S. Deshmukh, S. R. Gaikwad, N. R. Mote, M. Manod, R. G. Gonnade and S. H. Chikkali, *ACS Omega*, 2019, 4, 9502–9511.
- 25 M. Zabransky, I. Císarova and P. Štěpnička, *Organometallics*, 2018, 37, 1615–1626.
- 26 N. Li, J. Wang, X. Zhang, R. Qiu, X. Wang, J. Chen, S. F. Yin and X. Xu, *Dalton Trans.*, 2014, **43**, 11696–11708.
- A. Monfredini, V. Santacroce, P.-A. Deyris, R. Maggi, F. Bigi,
 G. Maestri and M. Malacria, *Dalton Trans.*, 2016, 45, 15786– 15790.
- 28 A. Fedorchuk, Y. Slyvka, V. Kinzhybalo, T. Lis and M. Myskiv, *Inorg. Chim. Acta*, 2019, **484**, 79–86.
- 29 H. Liu, H. Wu, J. Ma, S. Song, J. Yang, Y. Liu and Z. Su, *Inorg. Chem.*, 2007, **46**, 7299–7311.
- 30 R. Şuteu, A. Toma, M. Mehring and A. Silvestru, J. Organomet. Chem., 2020, 920, 121343.
- 31 A. Toma, C. Raţ, O. Pavel, C. Hardacre, T. Rüffer, H. Lang, M. Mehring, A. Silvestru and V. Pârvulescub, *Catal. Sci. Technol.*, 2017, 7, 5343–5353.
- 32 L. M. Mirica and J. R. Khusnutdinova, *Coord. Chem. Rev.*, 2013, **257**, 299–314.
- 33 J. Li, X. Li, L. Sun, X. Wang, L. Yuan, L. Wu, X. Liu and Y. Wang, *Eur. J. Inorg. Chem.*, 2021, 40, 4230–4237.
- 34 J. R. Shakirova, E. V. Grachova, V. V. Gurzhiy, I. O. Koshevoy, A. S. Melnikov, O. V. Sizova, S. P. Tunik and A. Laguna, *Dalton Trans.*, 2012, 41, 2941–2949.
- 35 G. J. Zhao, B. H. Northrop, K. L. Han and P. J. Stang, *J. Phys. Chem. A*, 2010, **114**, 9007–9013.
- 36 M. Liu, H. Hao, G. Liao, C. Li, K. Liu, N. Wang, Q. Niu and X. Yin, *Adv. Opt. Mater.*, 2023, 2202918.
- 37 M. Fan, Y. Tang, C. Li, B. Chen, T. Wang, P. Zhou and X. Cui, J. Phys. Chem. B, 2023, 127, 3187–3196.

- 38 G. Singh, Pawan, S. Sharma, Mohit, P. Satija, Diksha, Priyanka, Y. Thakur and A. Kaur, *Inorg. Chem.*, 2022, **61**, 12043–12061.
- 39 A. Barak, N. Dhiman, F. Sturm, F. Rauch, Y. Lakshmanna,
 K. Findlay, A. Beeby, T. Marder and S. Umapathy, *J. Phys. Chem. C*, 2023, **127**, 5855–5865.
- 40 R. Hamze, J. Peltier, D. Sylvinson, M. Jung, J. Cardenas, R. Haiges, M. Soleilhavoup, R. Jazzar, P. Djurovich, G. Bertrand and M. Thompson, *Science*, 2019, 363, 601–606.
- 41 H. Danjo, K. Asai, T. Tanaka, D. Ono, M. Kawahata and S. Iwatsuki, *Chem. Commun.*, 2022, **58**, 2196–2199.
- 42 Y. Liu, Y. Zhou, J. Li, Q. Wang, Q. Qin, W. Zhang, H. Asakura, N. Yan and J. Wang, *Appl. Catal.*, *B*, 2017, **209**, 679–688.
- 43 Y. Liu, C. Zou, K. Wang, Z. Bian, S. Jiang, Y. Zhou and J. Wang, *Mol. Catal.*, 2021, **505**, 111487.

- 44 K. Wang, W. Cui, Z. Bian, Y. Liu, S. Jiang, Y. Zhou and J. Wang, *Appl. Catal.*, *B*, 2021, **281**, 119425.
- 45 Y. Lou, J. Ma, W. Hu, Q. Dai, L. Wang, W. Zhan, Y. Guo, X. Cao, Y. Guo, P. Hu and G. Lu, *ACS Catal.*, 2016, 6, 8127–8139.
- 46 Y. Zhang, Q. Sun, G. Guo, Y. Cheng, X. Zhang, H. Ji and X. He, *Chem. Eng. J.*, 2023, **451**, 138867.
- 47 W. Shi, G. Xu, X. Han, Y. Wang, Z. Liu, S. Xue, N. Sun, X. Shi, Y. Yu and H. He, *J. Environ. Sci.*, 2023, **126**, 333–347.
- 48 A. Ibrahim, U. B. Memon, S. P. Duttagupta, R. K. Singh Raman and A. Sarkar, *J. Electron. Mater.*, 2023, **52**, 446–462.
- 49 Y. Yun, H. Sheng, J. Yu, L. Bao, Y. Du, F. Xu, H. Yu, P. Li and M. Zhu, *Adv. Synth. Catal.*, 2018, **360**, 4731–4743.
- 50 Z. Jiang, S. Guo and T. Fang, *ACS Appl. Energy Mater.*, 2019, 2, 7233–7243.