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Computational investigations of click-derived 1,2,3-triazoles as keystone ligands for complexation with transition metals: a review†

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In recent years, metal complexes of organo 1,2,3-triazole click-derived ligands have attracted significant attention as catalysts in many chemical transformations and also as biological and pharmaceutical active agents. Regarding the important applications of these metal–organo 1,2,3-triazole-based complexes, in this review, we focused on the recently reported investigations of the structural, electronic, and spectroscopic aspects of the complexation process in transition metal complexes of 1,2,3-triazole-based click ligands. In line with this, the coordination properties of these triazole-based click ligands with transition metals were studied *via* several quantum chemistry calculations. Moreover, considering the complexation process, we have presented comparative discussions between the computational results and the available experimental data.

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1. Introduction

1,3-Dipolar cycloaddition is a chemical reaction between a 1,3-dipole and a dipolarophile to afford a five-membered ring. Mechanistic study and the synthetic application of this

important reaction were investigated initially *via* Rolf Huisgen's study reported in the 1960s.^{1,2}

Although this reaction is often referred to as a 1,3-dipolar cycloaddition reaction, it is worth noting that the specific reaction is actually between an organic azide and an alkyne to construct a common and useful heterocyclic system of 1,2,3-triazoles, as initially realized by Rolf Huisgen^{1,2} (Scheme 1). It was the American chemist K. Barry Sharpless who referred to this cycloaddition as “the cream of the crop” of click chemistry. Click chemistry reactions are usually high yielding and wide in scope and also generate only by-products that can be removed

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† Dedicated to the unique essence of a great teacher, Professor Majid M. Heravi, who is the realization of simorgh in the form of a scientist.



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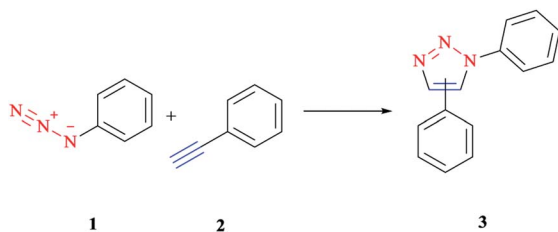
Iran. Her research interests focus on computational organic and organometallic chemistry and statistical thermodynamics. She has published more than 31 papers in reputed journals and has been recently serving as the guest editor of the thematic issue in *Current Organic Chemistry*.



Fatemeh Ebrahimpour-Malmir was born in Tehran, Iran in 1990. She received her B.Sc. degree in Applied Chemistry from Sharif University of Technology, Tehran, Iran (2014) and her M.Sc. degree in Organic Chemistry at Alzahra University, Tehran, Iran (2016) under the supervision of Prof. Majid M. Heravi and the advison of Dr Tayebeh Hosseinnejad. Her researches are concentrated on

the catalyzed click reactions from the combined experimental and computational viewpoints.





Scheme 1 Synthesis of a mixture of 1,4- and 1,5-triazoles via an azide-alkyne cycloaddition reaction.

easily with no chromatography required. Click reactions are regioselective and stereospecific, simple to conduct and can be also performed in easily removable or benign solvents.³

An outstanding variant of the Huisgen 1,3-dipolar cycloaddition is the copper(i)-catalyzed cycloaddition click reaction, in which organic azides and terminal alkynes are reacted to afford 1,4-regioisomers of 1,2,3-triazoles as the only products under mild reaction conditions and in a much shorter reaction time.

After the discovery of the regioselective copper(i)-catalyzed azide-alkyne cycloaddition (CuAAC) reaction, so-called click reaction,^{4,5} the synthesis of click ligands (the archetypal example of click chemistry) attracted significant attention due to its reliability, simplicity, and mild reaction conditions.

It should be mentioned that the mild-functional-group-tolerant click synthesis of 1,2,3-triazole species led to their widespread applications as building blocks throughout synthetic chemistry. The application of the CuAAC click reaction in the synthesis of 1,2,3-triazoles as a stable linkage between chemical or biological components and a coordinating agent with metal ions and also their potential in catalytic systems were assessed, subsequently, increasing their impact far beyond the initial purpose of click chemistry.^{6–10}

As a matter of fact, part of the special interest and focus on this type of reaction is due to the significant and diverse biological activities shown by 1,2,3-triazoles.^{11–15} These compounds can act as rigid linking units and can mimic the atom placement and electronic properties of a peptide bond, but with no hydrolytic cleavage effect. It is important to note that 1,2,3-

triazoles have the following structural and electronic features, which have a significant high impact on their biological pharmaceutical applications. The dipole moment of triazoles is much stronger than that of an amide bond, which leads to increasing the hydrogen bond donor and acceptor properties of triazoles, subsequently enhancing the peptide bond mimicry.^{15,16}

Due to the aforementioned potential of 1,2,3-triazoles to mimic the peptide bond role, several biological activities have been screened for 1,2,3-triazoles, including anti-HIV activity,¹⁷ selective β_3 adrenergic receptor inhibition,¹⁸ anti-bacterial activity,¹⁹ and potent anti-histamine activity.²⁰ It is important to note that 1,2,3-triazole click compounds have anticancer potency, which can be subsequently increased upon complexation with transition metal ions.^{21–23}

On the other hand, based on the solid-phase click chemistry, switching from the solution-phase to solid-phase chemistry provides a playground for the application of triazoles to act as linkers to attach other functionalities to solid-phase resins²⁴ as well as being able to functionalize metal surfaces for modification of their properties²⁵ and to enable prompt adhesion.²⁶

Recently, the promise of metal complexes of organo 1,2,3-triazole-based ligands has been demonstrated in the catalysis of some chemical transformations,^{27,28} such as hydroformylation and intramolecular hydroamination.^{29–36}

It was shown that the catalytic efficiency of these complexes essentially depends on the appropriate electron-donating ability of atoms in 1,2,3-triazole moieties. Precisely speaking, the 1,2,3-triazole unit in the click ligand has an N-donor coordinating role to play through two types of nitrogen atoms: N(2) and N(3).³⁷ From a mechanistic viewpoint, such a set of donor groups may stabilize intermediate species *via* coordination and increase reactivity and can also reduce the energy of transition states by partial dissociation to consequently improve the catalytic efficiency of the complex.

To the best of our knowledge, a few research studies have reported on the synthesis of 1,2,3-triazole organo-based ligands and the applications of their metal complexes as catalysts.^{38–42}

In addition to their catalysis applications, some transition metal complexes of 1,2,3-triazole-based ligands have attracted interest for their luminescent properties and consequent applications as radiopharmaceutical agents in bioimaging techniques.^{43–45}

Envisaging the importance of such complexes and their several applications, the structural, electronic, and spectroscopic features of these metal complexes with 1,2,3-triazole-based ligands have been widely investigated from experimental and computational viewpoints.

In line with this, several spectroscopic techniques have been used to assess the structural and electronic properties of these complexes. On the other hand, the veracity of computational methods was ascertained by comparison of the calculated results with experimental observations. However, the calculated geometrical parameters of these complexes could be compared with the available X-ray crystal structure and also the frequency calculations can be attributed to FT-Raman spectra. It should be mentioned that Raman spectroscopy can probe the nature of



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electronic transitions and also provide an insight into the structural character of the donor and acceptor MOs.⁴⁶ Clearly speaking, the resonance effect enhances the Raman spectra bands corresponding to the structural changes associated with the photoexcitation.^{47–52} So, the pattern of band enhancements can be combined and interpreted with a computational analysis of the vibration normal modes.

On the other hand, the HOMO (highest occupied molecular orbital) and LUMO (lowest unoccupied molecular orbital) energy gap of a complex can be correlated to its chemical reactivity and this information can be applied to interpret the catalytic role of these transition metal complexes of 1,2,3-triazole-based ligands.

In this route, the structural and electronic aspects of coordination in a series of mono and multinuclear transition metal complexes containing nitrogen-donating triazole ligands were investigated computationally through structural optimization calculations *via* density functional theory (DFT)^{53,54} and ligand field analysis.⁵⁵ Moreover, the conformational fluxionality of flexible triazole-based click ligands in coordination with metal ion centers was assessed in the formation of *trans* or *anti-trans* conformers of six-membered metallacycles, and the isomerization mechanism was also studied to obtain the activation and entropy of the reactions *via* DFT computations.^{56,57}

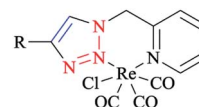
Moreover, the structural aspects of complexation between 1,2,3-triazole-based organochalcogen ligands and metal ions were studied *via* DFT computations and crystallographical methods and showed a disposition of donor atoms around the metal center with several coordination modes through the triazole skeleton.^{38–41}

The functionalization effect on the structural and photo-physical properties of the complexation of triazole-based ligands with transition metal ions have been studied and reviewed.^{58–62}

In this respect, the relation between the energy adsorption band of the electronic spectra and metal-to-ligand charge transfer (MLCT) for all the complexes were modeled *via* DFT approaches.^{61,63–65}

The coordination and electronic features of functionalized triazaphosphole click ligands and their substitution pattern were studied in comparison with triazole-based click ligands using quantum chemical computations, and the results were verified by crystallographical data.⁶⁶

We recently investigated the synthesis and characterization of various heterogeneous nanocatalysts and their applications in various organic reactions, especially in the regioselective synthesis of 1,2,3-triazoles *via* a click reaction.^{67–75} We also recently focused on combining experimental and



Scheme 3 *fac*-Re(CO)₃Cl·2-(1-R-1H-1,2,3-triazol-4-yl)pyridine (6) complexes.

computational studies to provide a deeper insight into this important reaction. We reviewed the mechanistical aspects of the 1,3-dipolar Huisgen cycloaddition reactions and the regio-selective behavior of metal-catalyzed click reactions using computational chemistry.^{5,76}

Armed with these experiences, herein, we show that mild one-pot click methods can be used to readily and rapidly synthesize a family of functionalized multi-dentate 1,2,3-triazole ligand scaffolds, containing electrochemically, photo-chemically, and biologically active functional groups in a reliable yield. Overall, in this review, we present the experimental and computational features of the synthesis and properties of functionalized click ligands and their complexes with transition metal cations. The structural, photochemical, and biological properties of the complexes will be discussed *via* the spectroscopic and computational analysis.

2. 1,2,3-Triazole-containing chelators: click synthetic vision

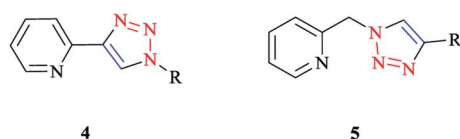
Click ligands are generally prepared using the functional-group-tolerant Cu(I)-catalyzed 1,3-dipolar cycloaddition of organic azides with terminal alkynes (CuAAC reaction),^{10,77,78} which leads to the rapid generation of a wide range of functionalized ligands in association with the mild reaction conditions.

The 2-(1-R-1H-1,2,3-triazol-4-yl)pyridine **4** and [(4-R-1H-1,2,3-triazol-1-yl)methyl]pyridine **5** (Scheme 2) click ligands have emerged as readily modified bipy and phen surrogates.^{8,79,80}

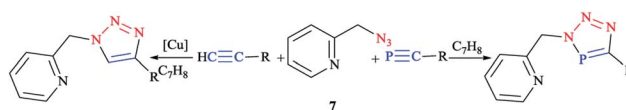
These bidentate click ligands form rhenium(I) complexes with a 1 : 1 metal : ligand ratio. The complexes of these ligands have wide applications in electronic^{81,82} and luminescent^{29,83–89} materials, catalysts,^{90–92} and in metallo-pharmaceuticals.^{93–95}

Moreover, with the goal of the development of novel rhenium(I) complexes as luminescent and pharmaceutical agents, rhenium(I) complexes from mono-functionalized 2-pyridyl-1,2,3-triazole derivatives (pyta) were synthesized as alternative ligands to 2,2'-bipyridines.⁶³ Nevertheless, only a few of them possessed a tethering group for further coupling to a biomolecule.^{63,96}

In short, the synthesis of novel pyta derivative grafted onto a bioactive pharmacophore was reported by considering the advantages of trendy click chemistry. Moreover, the

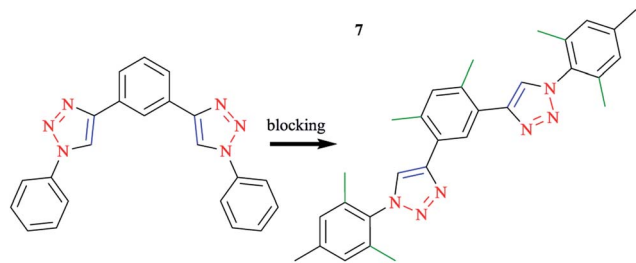


Scheme 2 Bidentate N^N chelators: 2-(1-R-1H-1,2,3-triazol-4-yl)pyridines (**4**) and [(4-R-1H-1,2,3-triazol-1-yl)methyl]pyridines (**5**).



Scheme 4 Syntheses of 2-pyridylmethyl-functionalized triazoles (left) and 2-pyridylmethyl-functionalized triazaphospholes (right).





Scheme 5 Optimization process, alternative coordination modes (left), and desired coordination mode (right).

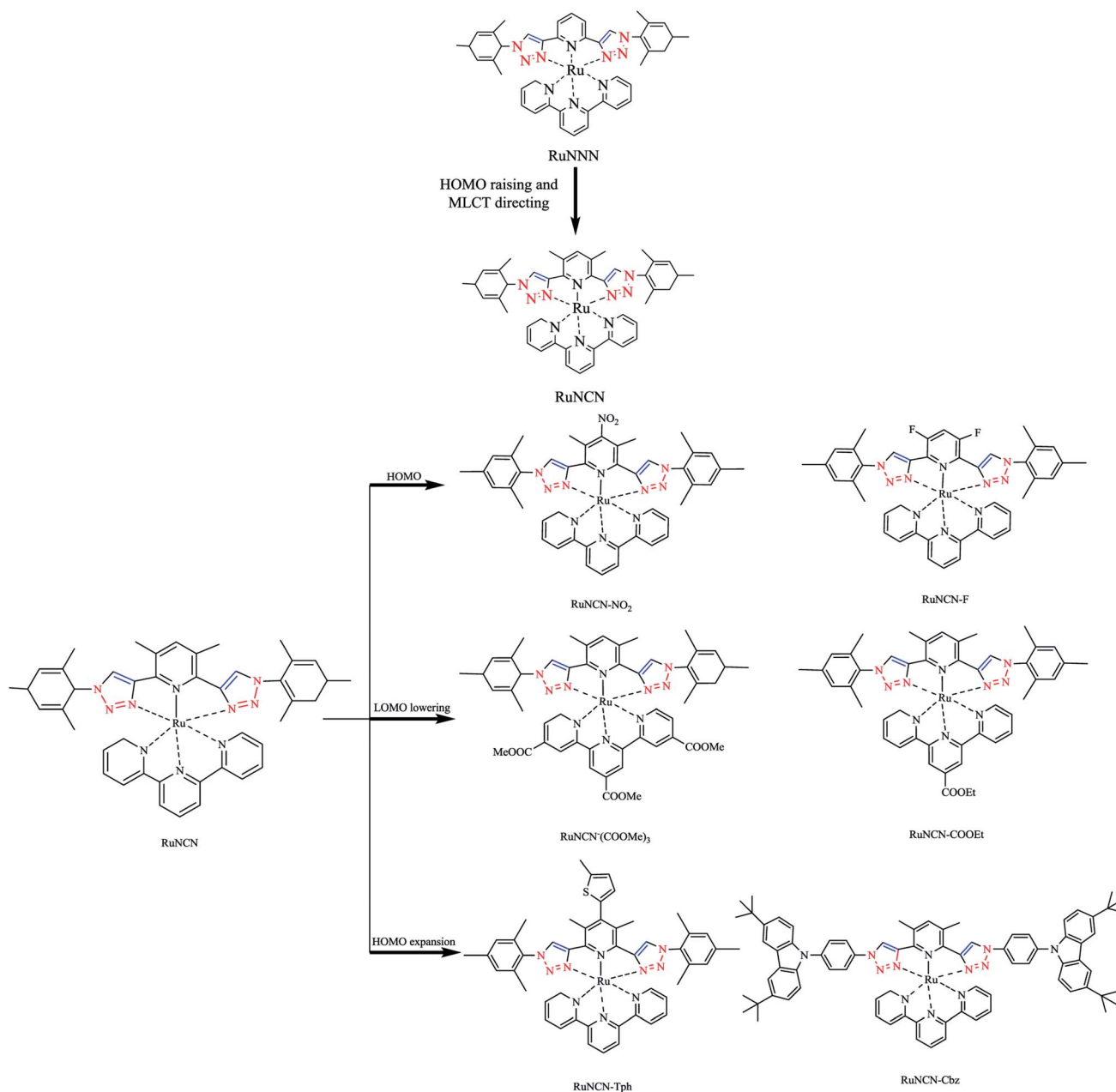
corresponding rhenium complexes were synthesized and characterized in the solid and solution states.^{63,97}

In another research, a family of electronically tuned *fac*- $\text{Re}(\text{CO})_3\text{Cl} \cdot 2\text{-(1-}R\text{-1}H\text{-1,2,3-triazol-4-yl)pyridine}$ **6** complexes (Scheme 3) was generated using efficient click reactions.

These safe one-pot CuAAC click methodologies produced a series of electronically modified 2-pyridyl-1,2,3-triazole ligand architectures, which were readily converted into the corresponding family of *fac*- $[\text{Re}(\text{CO})_3\text{Cl}]$ complexes.

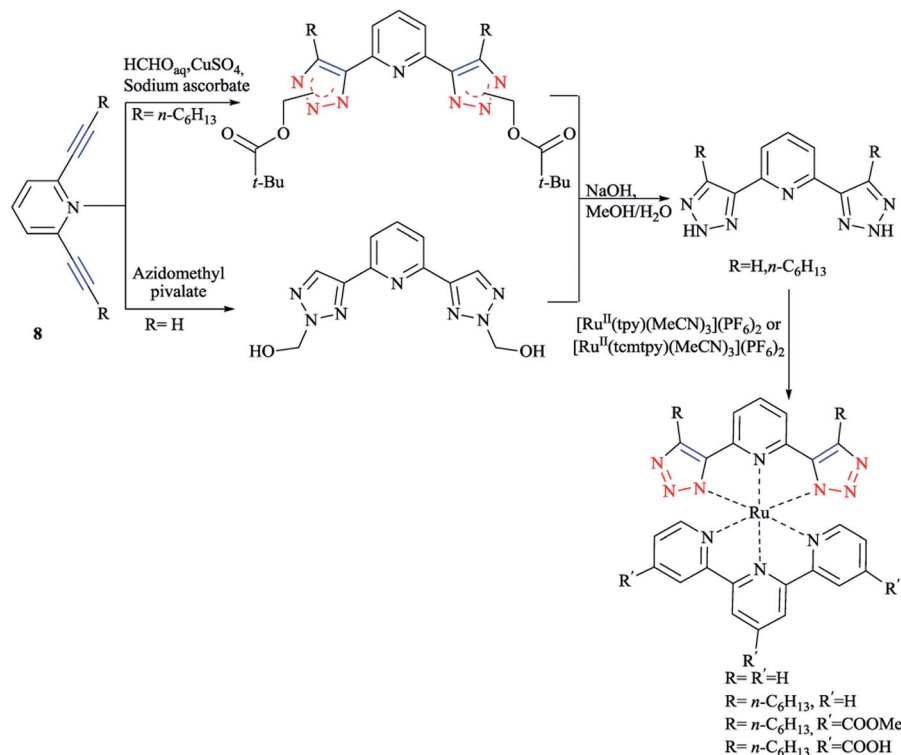
Elemental analysis demonstrated that the isolated powders were pure and showed that the bidentate ligands generated rhenium(i) complexes in a 1 : 1 metal/ligand ratio.⁶⁴

The coordination chemistry of analogous donor-functionalized triazoles has been widely investigated over the



Scheme 6 Synthesis procedures of ruthenium(II) complexes.



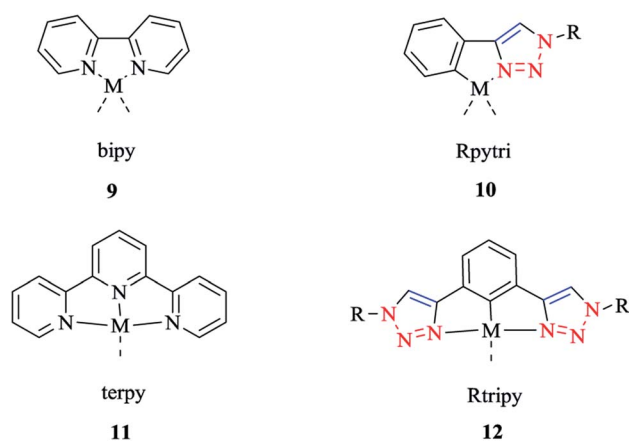


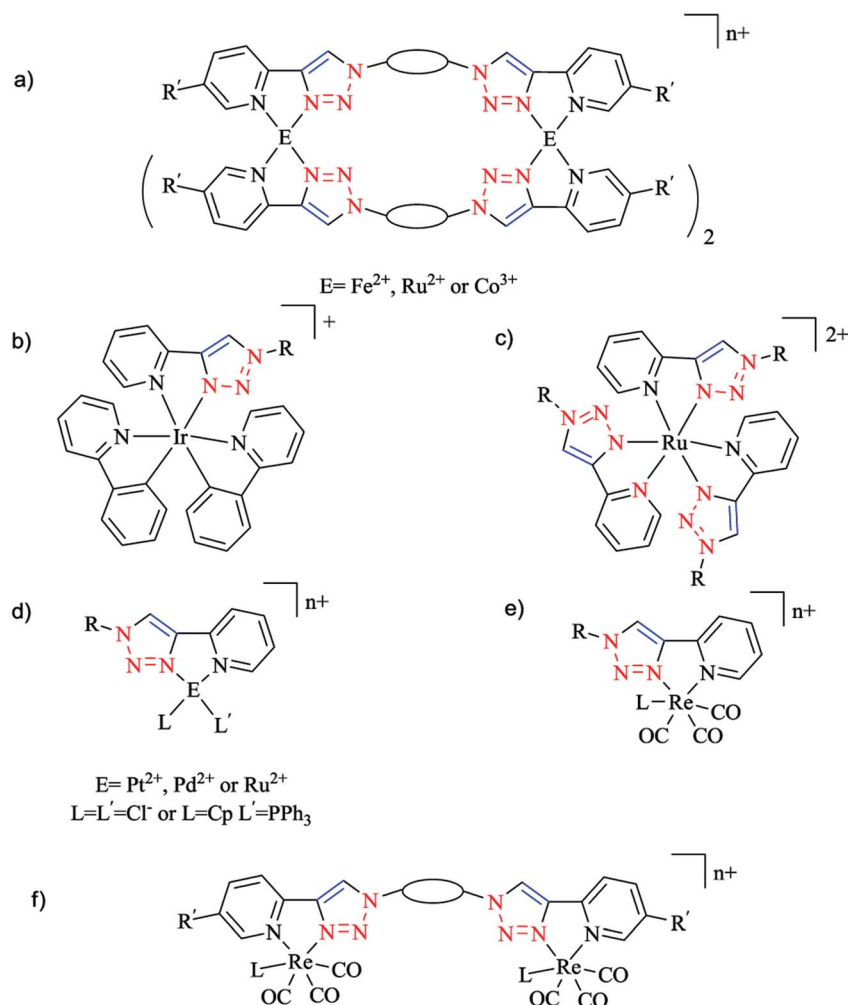
Scheme 7 Synthesis of bis(tridentate)ruthenium(II) complexes.

past decades,^{8,59,79,80,82,94,98,99} whereas the chelating behavior of donor-functionalized 3*H*-1,2,3,4-triazaphosphole derivatives is still practically uncharted.^{100–102} Clearly speaking, the click reaction between donor-functionalized azides and phosphalkynes enables access to polydentate ligands, so that we can adjust the stereoelectronic properties based on the additional phosphorus heteroatom and the nature of the substituents. However, few investigations have been performed on the influence of an additional heteroatom on the properties of triazole derivatives. So far, only one example of such a species was assessed and characterized crystallographically, namely a tridentate ligand, tripodaltris (triazaphosphole), coordinated toward a Pt(0) center *via* P dentate, as reported by Jones *et al.*¹⁰³ In the next research studies, the click reaction between 2-(azidomethyl)pyridine¹⁰⁴ **7** with *t*Bu-C≡P¹⁰⁵ and TMS-C≡P¹⁰⁶ (trimethylsilyl is denoted as TMS) was reported according to the standard literature processes. Furthermore, the reaction between equimolar amounts of the azide and *tert*-butylphospha-ethyne was carried out at room temperature in toluene *via* a click methodology. Interestingly, it has been also reported that trimethylsilylphospha-ethyne reacts straightforwardly with 2-(azidomethyl)pyridine **7** and proceeds to give the TMS-substituted triazaphosphole derivatives (Scheme 4).⁶⁶

Click-derived^{3,107} ligands have been introduced as analogs of polypyridyl ligands, in particular of terpyridine (tpy),^{96,108–111} and successfully used to produce tridentate cyclometalating polypyridyl ligands, namely 1,3-dipyridylbenzene. In this respect, aryl azides react with diethynylbenzene building blocks, in good yields using a standard click procedure.^{112–114} With the purpose

of blocking alternatives, methyl groups were placed at strategic positions when possible and reasonable.¹¹⁵ Therefore, *o*-xylene was selected as the central ring and mesityl species were selected for the clicked-on functionalities (Scheme 5). It should be mentioned that mesityl was kept as the reference moiety in all the other experiments to discuss the important influences of substituents directly attached to the cyclometalating phenyl ring or the opposing ligand (Scheme 6).

Scheme 8 Structures of the 2,2'-bipyridine (bipy) (**9**) metal complex, 2-pyridyl-1,2,3-triazole (Rpytri) (**10**) analog, 2,2':6',2''-terpyridine (terpy) (**11**) metal complex, and 2,6-pyridyl-bis(1,2,3-triazole) (Rtripy) (**12**) analog.



Scheme 9 Structures of biologically active click Rpytri complexes.

In this context, a new and systematic series of click-derived tridentate cyclometalated Ru(II) complexes was considered to assess their potential for dye-sensitized solar-cell applications.⁸⁵ These combined theoretical photophysical studies as well as electrochemical experiments led to a consistent and emergent explanatory picture of the new dyes.¹¹⁶

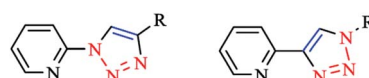
In another research, the synthesis of 2,6-bis(1,2,3-triazol-4-yl)pyridine **8** ligands were performed *via* the Cu(I)-catalyzed cycloaddition reaction of hydroxyl methylazide followed by a base-induced cleavage of formaldehyde. It could also be synthesized *via* a thermal azide-alkyne cycloaddition with azidomethyl pivalate and an internal alkyne. In this azide-alkyne cycloaddition, a mixture of adducts was produced and after cleavage and reprotonation, and free NH-triazole was obtained.⁷

As illustrated in Scheme 7, the corresponding ruthenium(II) complexes were synthesized in good yields using $[\text{Ru}^{\text{II}}(\text{tpy})(\text{MeCN})_3](\text{PF}_6)_2$ ($\text{tpy} = 2,2':6',2''\text{-terpyridine}$) or $[\text{Ru}^{\text{II}}(\text{tcmtpy})(\text{MeCN})_3](\text{PF}_6)_2$ ($\text{tcmtpy} = 4,4',4''\text{-tricarboxymethyl-2,2':6',2''-terpyridine}$) as precursors.¹¹⁶ The solubilities of the charge-neutral complexes were measured and it was shown that

the alkyl chains increases the solubility, allowing the investigation of the photophysical and electrochemical properties (*vide infra*).¹¹⁷

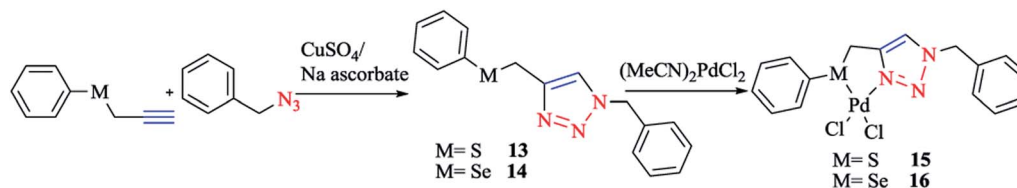
The synthesis and biological activity of Fe, Ru, Co, Pt, and Pd complexes of 2-pyridyl-1,2,3-triazole (Rpytri) **10** bidentate and 2,6-pyridylbis(1,2,3-triazole) (Rtripy) **12** tridentate click ligands (Scheme 8) were investigated. It is important to mention that Rpytri and Rtripy click ligands are commonly considered as analogs of the 2,2'-bipyridine (bipy) **9** and 2,2':6',2''-terpyridine (terpy) **11** ligand systems (Scheme 8).^{8,79}

In this vein, the CuAAC click reaction was exploited to synthesize a series of iron(II),¹¹⁸ ruthenium(II),¹¹⁹ and cobalt(III)¹²⁰ bis-2-pyridyl-1,2,3-triazole (bis-Rpytri) ligands (Scheme 9). It was demonstrated that iron(II) bis-Rpytri complexes have no biological activity and are rapidly decomposed under biological



Scheme 10 Structures of inverse-Rpytri (left) and regular-Rpytri (right) isomers.





Scheme 11 Synthesis of palladium(II) complexes based on the 1,2,3-triazole organosulfur/selenium ligand.

conditions, while the more inert ruthenium(II) and cobalt(III) bis-Rpytri complexes showed biological stability. The same authors also produced a family of mono- and di-*fac*-rhenium(I) tricarbonyl complexes using the Rpytri and bis-Rpytri ligands with aliphatic and aromatic substituents and linkers (Schemes 9e and f). These showed modest biological activity, attributed to the low charge on the rhenium centers as well as their molecular shape.¹²¹

Moreover, Henderson and coworkers synthesized complexes of iridium(III) with Rpytri ligands including aromatic substituents and assessed their biological potency (Scheme 9b).²³

In another work, Aldrich-Wright and coworkers generated a set of six heteroleptic platinum(II) complexes bearing chiral diaminocyclohexane and either aliphatic or aromatic Rpytri ligands (Scheme 9d).¹²²

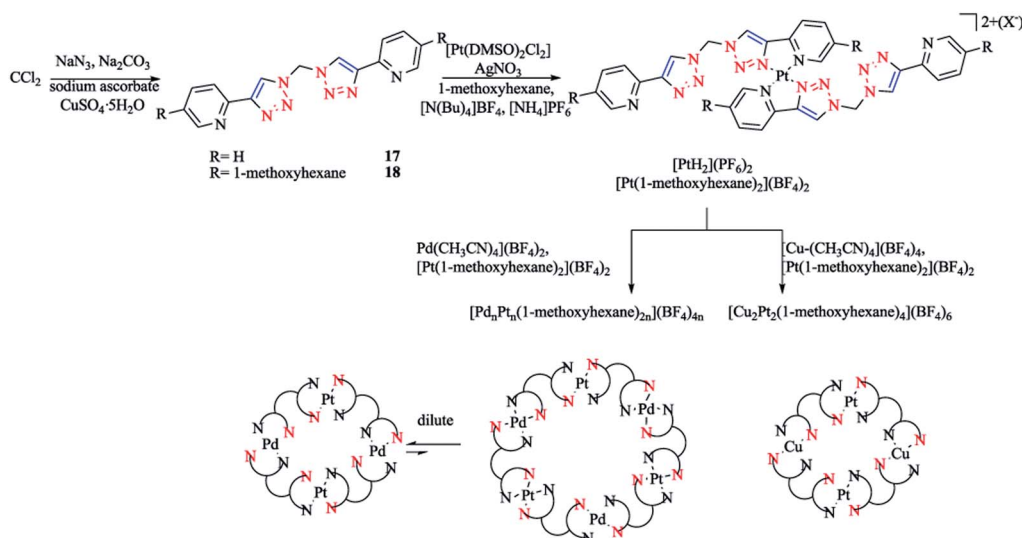
Pyridyl-substituted $[Pd(Rpytri)Cl_2]$ complexes were also synthesized by Sangtrirutnugul and coworkers. The regular Rpytri complex demonstrated a significant catalytic activity in the Suzuki reaction under aerobic conditions without catalyst degradation, indicating it was more resilient than the inverse Rpytri complex (Scheme 10).^{22,123}

According to the procedure illustrated in Scheme 11, complexes **15** and **16** were synthesized as yellow solids by reacting $[(MeCN)_2PdCl_2]$ with **13** and **14**, respectively, in a 1 : 1 molar ratio, in THF solution at room temperature, leading to 90% and 85% yields for complexes **15** and **16**, respectively. The two air- and moisture-insensitive complexes **15** and **16** were moderately soluble in $CHCl_3$, CH_2Cl_2 , and CH_3OH , but had

good solubility in CH_3CN and DMSO. **13** and **14** and their Pd complexes **15** and **16** were characterized with 1H , $^{13}C\{^1H\}$ and $^{77}Se\{^1H\}$ NMR, HR-MS, and IR spectra, and the results were found to be in good confirmation with those reported earlier.^{37,124}

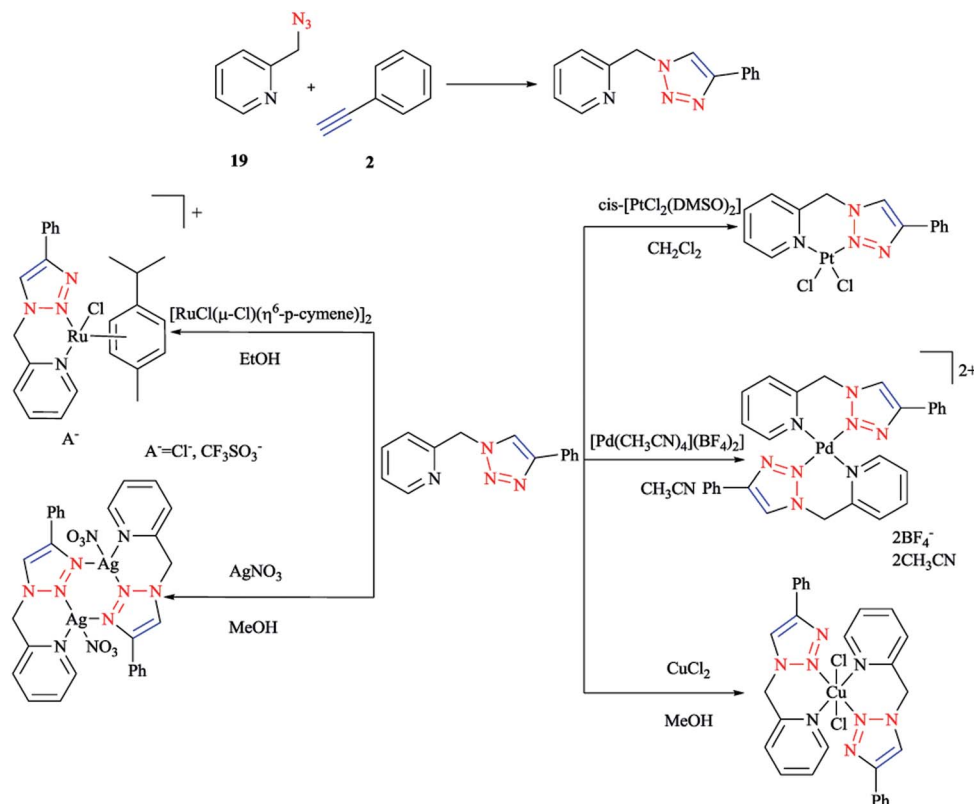
In another work, $[Pd(Rpytri)_2]^{2+}$ complexes were synthesized from the ligands in a head-to-tail arrangement avoiding steric clashes and leading to favorable interligand hydrogen bonding.^{56,125} Furthermore, related platinum Rpytri compounds are known, including $[Pt(Rpytri)_2Cl_2]$ complexes,^{82,125–129} heteroleptic $[Pt(Rpytri)_2(L)_2]^{2+}$ complexes, where L = phenylpyridine^{130,131} or diaminocyclohexane,¹²² and bis-pyridyl-triazolato complexes.¹³² Two bis-2-pyridyl-1,2,3-triazole ligands (L) with methylene spacers were also synthesized, which had the shortest possible “regular” pyridyl-triazole linkage, with the fewest possible degrees of freedom of any “hinged” bis-pyridyl-1,2,3-triazole. Then, these ligands were used to produce bis-pyridyl-triazole $[Pt(L)_2]^{4+}$ complexes, which were assembled exclusively in a head-to-tail conformation. These Pt(II) metal-ligands had vacant peripheral binding sites that could then be employed in the generation of heterometallic complexes. Mixed Pd(II)/Pt(II) macrocycles were synthesized in a concentration-dependent manner, while a mixed metal Cu(I)/Pt(II) was prepared independent of the concentration (Scheme 12).

While a wide range of bispyridyltriazole ligands^{133–135} are known, the previously unreported methylene-linked ligands



Scheme 12 Synthesis of heterometallic $[M_nPt_n(L)_{2n}]^{x+}$ macrocycles based on dichloromethane-derivative bis-2-pyridyl-1,2,3-triazole ligands.





Scheme 13 Syntheses of 1-(2-picolyl)-4-phenyl-1H-1,2,3-triazole ligand and some of the complexes investigated.

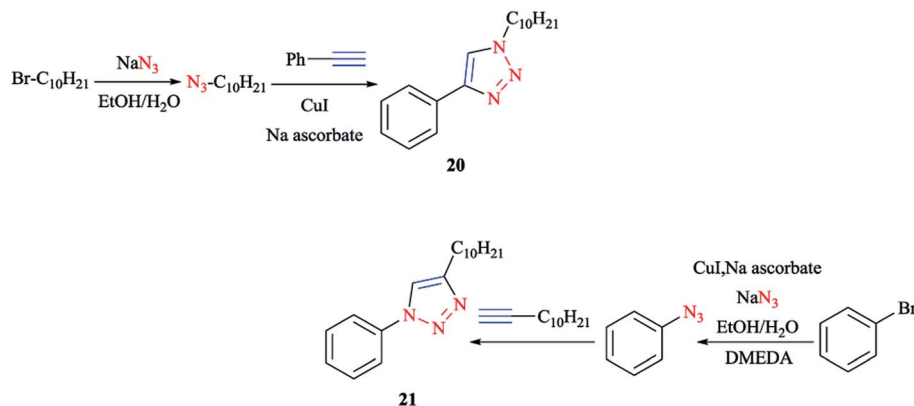
were targeted, to reduce the number of degrees of freedom in the ligand architecture.^{136–138}

The 1,1-bis(2-pyridyl-1,2,3-triazol-4-yl)methane **17** ligand and 1,1-bis(4-(5-(hexyloxy)pyridin-2-yl)-1H-1,2,3-triazol-1-yl)methane **18** ligand were synthesized through a CuAAC click reaction from sodium azide and dichloromethane (DCM) in a microwave in dimethyl formamide (DMF) up to 110 °C in the presence of carbonate, leading to the explosive diazide *in situ*, which then reacts with 2-ethynylpyridine or 2-(2-(trimethylsilyl)ethynyl)-5-(hexyloxy)pyridine.¹³⁹

In the following (Scheme 13), we display the synthesis of a click ligand through a CuAAC one-pot reaction between phenylacetylene

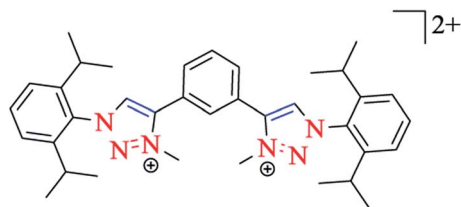
2 and 2-picolylazide **19** and also the coordination of a click ligand to Pt^{II} , Pd^{II} , Cu^{II} , Ru^{II} , and Ag^{I} using $\text{cis-}[\text{PtCl}_2(\text{DMSO})_2]$, $[\text{Pd}(\text{CH}_3\text{CN})_4](\text{BF}_4)_2$, CuCl_2 , $[\text{RuCl}(\mu\text{-Cl})(\eta^6\text{-p-cymene})]_2$, and AgNO_3 . In the case of $[\text{Pd}(\text{CH}_3\text{CN})_4](\text{BF}_4)_2$ and CuCl_2 , bis-bidentate complexes were obtained, respectively. Synthesis of these two compounds was repeated with a ligand–metal precursor molar ratio of 2 : 1. Other stable products crystallized out from the reaction mixtures and were then isolated by filtration in a 60–82% yield.⁹⁹

The synthesis of cyclometalating click ligands was investigated in a highly efficient one-pot consecutive process. In this regard, the acetylene derivative was reacted with the organic azide.^{140,141} It is important to note that there was no necessity for



Scheme 14 Syntheses of 1- and 4-decyl-4-phenyl-1H-[1,2,3]triazole ligands.





Scheme 15 Structure of the bis-monodentate 1,2,3-triazole-5-ylidene (22) ligand.

any further copper(i) catalyst after the formation of the aryl azide for the click reaction. So, 1-decyl-4-phenyl-1*H*-[1,2,3]triazole **20** and 4-decyl-1-phenyl-1*H*-[1,2,3] triazole **21** were produced as the potential ligands for the formation of iridium(III) complexes in high yields (Scheme 14).¹⁴⁴

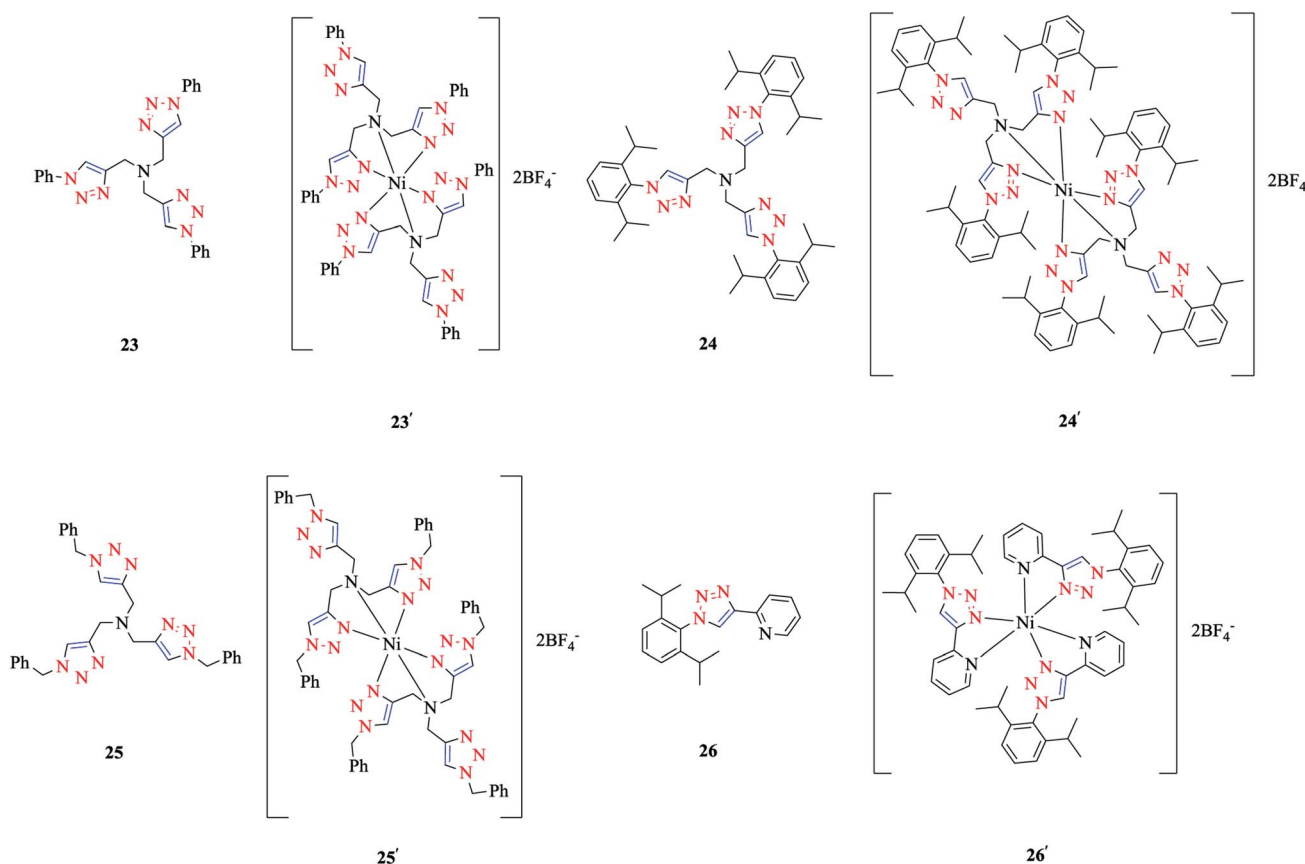
Metallo-macrocycles can be assembled by arranging bis-1,2,3-triazol-5-ylidene **22** carbene ligands in conjunction with linear Ag(i) or Au(i) ions. Crudden and coworkers produced a Ag(i) macrocycle from a bis-1,3(1-aryl-1,2,3-triazol-4-yl) benzene ligand (Scheme 15).¹⁴² An equilibrium mixture of mononuclear $[AgL]^+$ and macrocyclic $[Ag_2L_2]^+$ complexes were synthesized by Frutos and Tóree by applying a 1,4-xylyl linked steroid substituted bis-1,2,3-triazol-5-ylidene ligand. In order to expand the range of Au(i) 1,2,3-triazol-5-ylidene catalysts, Crowley and coworkers synthesized a di-Au(i) macrocycle with a 1,3-phenyl linked bis-1,2,3-triazol-5-ylidene ligand.¹⁴³

Furthermore, White and Beer produced four bis-monodentate ligands with a 1,3-(5-*tert*-butyl)phenyl core.¹⁴⁴ The triazole rings bound *via* N1 or C4 atoms and C4-connected ligands gave cluster-like compounds consisting of four Ag(i) cations and four ligands, while three triazole nitrogens were coordinated to each Ag(i) center and the fourth coordination bond for each metal ion was to water.

Building on their previous Ag(i) metallocycle work, Crowley and coworkers exploited the functional group tolerance of the CuAAC “click” reaction to synthesize a structurally diverse family of bis-triazole ligands. Reaction of a suitable dialkyne with two equivalents of azide gave a range of bis-monodentate ligands with the two triazole groups separated by a linker (either propyl, 1,3-phenyl or 1,4-phenyl units), and substituted with a wide array of benzyl, aryl, and alkyl groups.^{21,145,146}

3. Computational study of electronic energy properties in transition metal complexes of triazole-based ligands

The coordination of Ni(II) with the tripodal ligands tpta (tris [(1-phenyl-1*H*-1,2,3-triazol-4-yl)-methyl]amine) **23**, tbta (tris [(1-benzyl-1*H*-1,2,3-triazol-4-yl)methyl]-amine) **24**, and tdta (tris [(1-(2,6-diisopropyl-phenyl)-1*H*-1,2,3-triazol-4-yl)methyl]amine) **25**, and the bidentate ligand pyta (1-(2,6-diisopropylphenyl)-4-(2-pyridyl)-1,2,3-triazole) **26** led to the $[Ni(tpta)_2](BF_4)_2$ **23'**,



Scheme 16 tpta, tbta, tdta, and pyta ligands and their corresponding Ni(II) complexes.



[Ni(tbta)₂](BF₄)₂ **24'**, [Ni(tdta)₂](BF₄)₂ **25'**, and [Ni(pyta)₃](BF₄)₂ **26'** complexes (Scheme 16).

In order to obtain a better insight into their absorption spectra, quantum chemistry calculations at the NEVPT2 level of theory^{147–154} were performed. Based on the NEVPT2 calculated values, the electronic transitions in the absorption spectra of complexes **23'**, **24'**, and **26'** were assigned (while corresponding calculations for complex **25'** could not be done due to its large size). The energies of the spin-allowed d–d transitions of complexes **23'**, **24'**, and **26'** were calculated comparatively based on their X-ray and DFT-geometry-optimized structures.

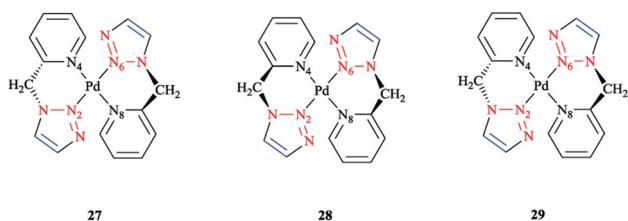
The experimental absorption spectrum of **23'** in CH₃CN was compared with the NEVPT2 calculated values. All the three main bands were adjusted well with three Gaussian distributions, indicate the three-times-three possible transitions, with the sum of these distributions in reliable agreement with the experimental spectrum.

The absorption spectra of **23'**, **24'**, and **26'**, which were calculated at the SA-CASSCF/NEVPT2 level of theory in acetonitrile included 24 excited states (9 triplets and 15 singlets, which with the triplet ground state resulted in a total of 45 microstates of d⁸). All the triplet states of the absorption spectrum could be attributed to the spin-allowed transitions, while the spin-forbidden transitions were not elucidated.

It should be noted that the excited singlet states can be ascertained by second-order perturbation theory (PT2) and make a minor contribution. In particular, when comparing SA-CASSCF¹⁵⁵ with NEVPT2 (ref. 156) approaches, since the dynamic electron correlation is added by the NEVPT2 method, the calculated values of the excited triplet states increase.

Consequently, the zero-field splitting (zfs) parameters *D* and *E* are decreased by almost 50%, while zfs qualitatively is inversely proportional to the excited state energies, which can be ascribed as the covalency of the system. The final *D* values calculated by quasi degenerate perturbation theory (QDPT)¹⁵⁷ were 2.94 and 3.79 cm^{−1} for **23'** and **24'** complexes, respectively, which were in reliable agreement with the values obtained by high-frequency and high-field electron paramagnetic resonance HFEP. ¹⁵⁸

Comparison of QDPT with PT2 approaches revealed that QDPT yielded more accurate results due to the fact that QDPT considers all contributions to the zfs, but this approach does not allow discussing the single contribution of excited states to the total zfs in comparison with PT2.



Scheme 17 Schematic structures of *anti-trans*-[Pd(L)₂]²⁺, *syn-trans*-[Pd(L)₂]²⁺, and *anti-cis*-[Pd(L)₂]²⁺ isomers based on the 4-phenyl-1-(2-picolyl)-1,2,3-triazole ligand (L).

For complex **26'**, the angular overlap model (AOM) parameters¹⁵⁹ were derived *via* a least-squares fit to the V matrices (with a rather small standard deviation) by using the exact structural data of complex **26'** obtained from the X-ray and DFT-optimized structure.

A comparison of the calculated results for the two ligands revealed that Ni–N bonding was dominated by Ni–N^{ta} and Ni–N^{py} σ-antibonding interactions. On the other hand, the parameters *e*_{πs}^{py} = −131 cm^{−1} and *e*_{πs}^{ta} = −276 cm^{−1} (attributed to the out-of-plane π-bonding) had very small and negative values.

In another research study, the interconversion between two boat conformers in a palladium(II) complex of the flexible click chelator (Pd{(py)₂CHNHR}Cl₂)¹⁶⁰ was assessed computationally. The rate constants for the interconversion between the two conformers were calculated. The calculated values for the activation enthalpy and activation entropy (from the Eyring plot) were 13.0 ± 0.4 kcal mol^{−1} and 2.7 ± 1.6 cal mol^{−1} K^{−1}, respectively, which were in reliable agreement with other literature results for boat-to-boat inversions in some similar palladacycles.¹⁶¹ It should be noted that the low activation entropy can be attributed to a nondissociative isomerization mechanism.

The species involved in equilibrium, including *syn*- and *anti*-conformers of the *trans* isomer, namely **27** and **28**, respectively, and the structure with an *anti-cis* arrangement **29** (Scheme 17) were investigated *via* DFT calculations. The calculated geometrical data clearly indicated that the change in the square-planar core was not considerable when going from the *anti* (**27**) to the *syn* (**29**) conformer; except the Pd–N₂ bond, which was slightly elongated to 0.014 Å.

Considering the calculated values of the electronic energy as well as the gas phase free energy, isomer **28** was found to be more stable than **27** by about 1 kcal mol^{−1}, thus predicting the equilibrium was shifted in the direction of **28**.

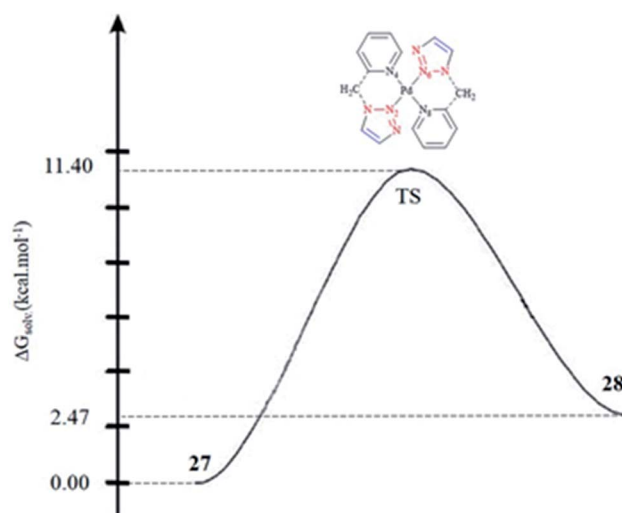


Fig. 1 One-step mechanism for the isomerization of **27** into **28**.



Moreover, the calculated value of steric repulsion, DE_{ster} , (which was defined as the sum of the Pauli repulsion and electrostatic interaction *via* the Ziegler–Rauk decomposition framework^{162,163}) was smaller between ligands in **28** ($DE_{\text{ster}} = -48.68 \text{ kcal mol}^{-1}$) than in **27** ($DE_{\text{ster}} = -50.36 \text{ kcal mol}^{-1}$), while the calculated value of the orbital interaction (DE_{oi}) was maintained at about $463 \text{ kcal mol}^{-1}$. This feature could be due to the slightly shorter N4–N6 distance in **27** ($3.016 \text{ \AA}$) than in **28** ($3.062 \text{ \AA}$).

To assess the kinetics of the isomerization procedure theoretically, we focused on the determination of the transition state (TS) connecting **27** and **28** *via* a one-step mechanism, as illustrated in Fig. 1. It was shown that the calculated activation barrier values and the entropy contributions were in a good agreement with the experimental data.⁵⁷

It was shown that in the following Pd complexes, the calculated electron densities at Se, Pd, and N(3) of the triazole ring were higher in complex **16** than in complex **15**, mainly due to the strong metal–ligand interaction in complex **16** and its subsequent higher stability and lower reactivity relative to complex **15**.^{37,56,126} Moreover, in these complexes, there was a good agreement between the experimentally observed and calculated bond distances and angles. However, some of calculated Pd–M bond distances ($M = \text{S/Se}$) deviated from the corresponding observed ones.¹²⁴

In another research study, the geometric and electronic structures of ruthenium complexes with bidentate ligands, including bis pyridyl and bis triazole ligands (denoted as bpy and

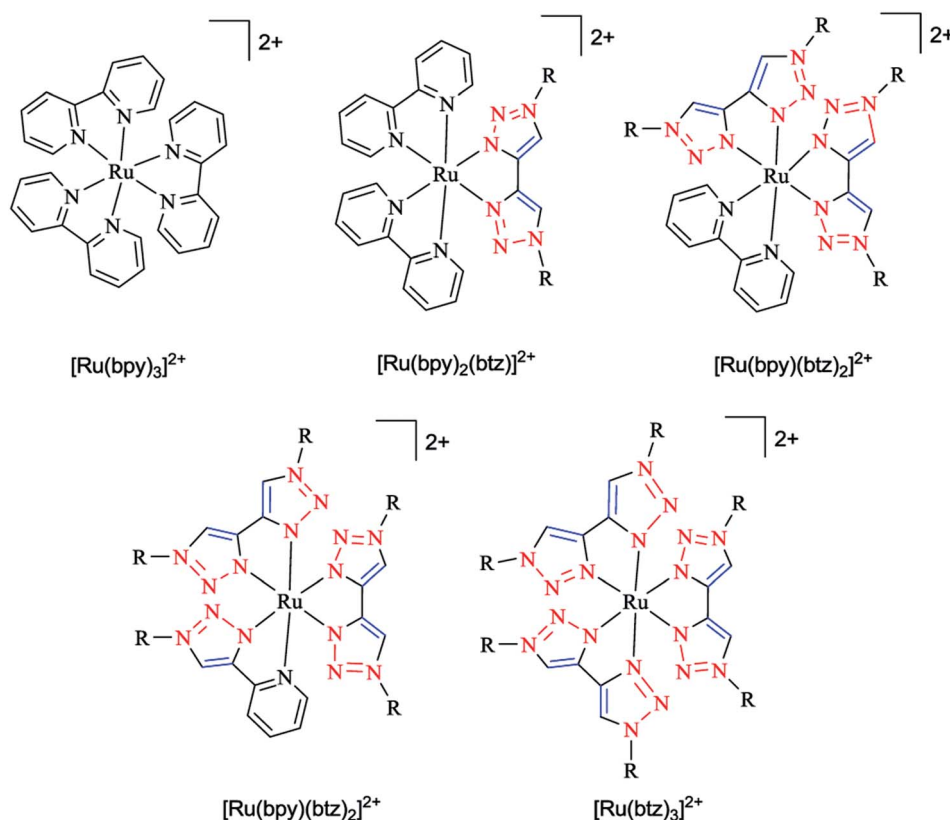
btz ligands, respectively), were investigated computationally (Scheme 18). In the first step, the optimized geometry of the complexes were obtained and then the localization of the d_{z^2} -like and $d_{x^2-y^2}$ -like antibonding $d\sigma^*$ orbitals were assessed to recognize the participating Ru–N bonds. In the next step, a pair of stimulated starting geometries was designed by elongation of the main participating Ru–N bonds and were then optimized with a spin multiplicity of 3.

It was shown that the Ru–N bond length increases due to the population of d_{z^2} -like $d\sigma^*$ orbitals. Moreover, the $d_{x^2-y^2}$ -like $d\sigma^*$ orbital population leads to the distortion of all three ligands. It was also demonstrated that a metal–ligand orbital overlap could be deducted based on the elongation of both the Ru–N bonds of one chelate ligand participating in the $d\sigma^*$ orbital, showing the stabilizing electronic effect.¹⁶⁴

4. Structural properties and frontier molecular orbital analysis in the complexation reactions

4.1 Ni(II) and Zn(II) complexes

The coordination behaviors of the complexes of Ni(II) with tripodal and bidentate ligands were investigated using X-ray diffraction studies and computational methods, specifically $[\text{Ni}(\text{tbta})_2](\text{BF}_4)_2$ **24'** as a tripodal ligand and $[\text{Ni}(\text{pyta})_3](\text{BF}_4)_2$ **26'** as a bidentate ligand. In the $[\text{Ni}(\text{tbta})_2](\text{BF}_4)_2$ **24'** complex, the Ni(II) center was coordinated by four triazole (N^{ta}) and two *trans*



Scheme 18 Structures of Ru(II) tris(bidentate) complexes (in experimental studies $R = \text{benzyl}$, to limit the computational expense in the calculations. R is simplified to methyl).



amine (N^{am}) nitrogen donors, in a distorted-octahedral space, while in $[Ni(pyta)_3](BF_4)_2$ **26'**, $Ni(II)$ was coordinated by pyridine (N^{py}) and triazole (N^{ta}) N-donors in a meridional arrangement.

As could be deduced from the crystal structure of $[Ni(tbta)_2](BF_4)_2$ **24'**, the nickel center was coordinated by each tbta ligand *via* the nitrogen atoms of two 1,2,3-triazole rings and the central amine nitrogen donors, while the third triazole ring of the ligand remained uncoordinated. The three nitrogen donors of triazole rings are located at the equatorial coordination position around the nickel center. It should be mentioned that the amine nitrogen atoms are poorer donors than the triazole nitrogen atoms, and this property leads to the larger $Ni-N^{am}$ distances in comparison with the $Ni-N^{ta}$ distances. Furthermore, it was shown that the average values of the axial-equatorial angle, denoted by α , and distortion angle, denoted by $\delta\alpha$, were significantly distorted from 90° . In the next step, the structural properties of the complexes were reproduced by using DFT geometry optimization in conjunction with a polarized continuum model (COSMO),¹⁶⁵ which showed a good agreement with the X-ray results.⁵⁵

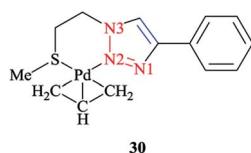
Petitjean and coworkers investigated systematically the synthesis and structural properties of $Fe(II)$, $Ni(II)$, and $Zn(II)$ ions with 1,2,3-triazol-4-yl pyridine chelators. They presented the spacer and substitution pattern for the complexation and assessed the various factors affecting the self-assembly procedure and also the stereochemical features of these architectures.¹³⁵

In another work, it was reported that the reaction of bis-(2-pyridyl-1,2,3-triazole) ligands including 1,3-propyl, 1,4-xylyl, or 2,6-dimethyl naphthalene spacers with $Fe(II)$ or $Ni(II)$ ions led to the production of M_2L_3 cylinders. It should be mentioned that by using several functional groups, a large series of bis-(2-pyridyl-1,2,3-triazole) ligands were synthesized *via* the CuAAC procedure and were then studied in the reaction with $Fe(II)$, $Ni(II)$, and $Zn(II)$ ions.^{135,166}

In the case of $Zn(II)$ complexes, the synthesis and structure of the $[2 + 2]$ macrocycle of $Zn(II)$ ions with the bis-(4-(2-pyridylthiomethyl)triazolyl) ligand including a pentyl linker were assessed. The zinc(II) ions were coordinated to two N3 triazole nitrogen atoms and two chloride anions in a tetrahedral coordination geometry. It was also demonstrated that the ethyl- and butyl-linked ligands formed $[(ZnCl_2)_2L]$ dinuclear complexes so that each metal ion was coordinated to both a triazole and chelating pyridine.^{21,167}

4.2 Pd(II) and Pt(II) complexes

The calculated optimized geometry for the $[Pd(\eta^3-C_3H_5)(1-(2-(methylthio)ethyl)-4-phenyl-1H-1,2,3-triazole)]^+$ (Scheme 19)

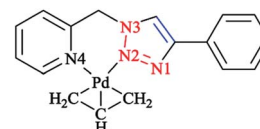


Scheme 19 Structure of the $[Pd(\eta^3-C_3H_5)(1-(2-(methylthio)ethyl)-4-phenyl-1H-1,2,3-triazole)]^+$ complex (**30**).

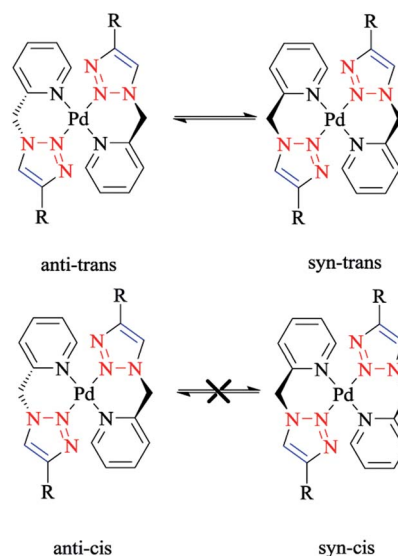
complex **30** was assessed computationally. The computations were performed at the M06 level of theory, including with the C-PCM solvation model and in the presence of acetone as a solvent, and the results highlighted the binding geometry adopted by the ligand.

Then, a comparison was made between the calculated results and the single-crystal data for the tetrafluoroborate salt of the cationic complex, which demonstrated a reliable confirmation. Moreover, based on the X-ray data, it could be seen that palladium was coordinated to N2 and the sulfur atoms and consequently formed a six-membered ring with the metal center.

The phenyl substituent and the triazole were coplanar, while the angle between N2, Pd, and S was around 95° . Another important structural feature of the complex was that the C-allyl-terminal *trans* to S and *trans* to triazole-N2 deviated from planarity specified by Pd, S, and N2, by 0.180 Å and 0.095 Å, respectively. It is important to mention that there was a similar coordination behavior between the triazole species in $[Pd(\eta^3-C_3H_5)(1-(2-(methylthio)ethyl)-4-phenyl-1H-1,2,3-triazole)]^+$ **30** with the pyridine-triazole $Pd(II)$ allyl complex (Scheme 20).^{91,168} Strictly speaking, coordination of the S-donor moiety of (1-(2-(methylthio)ethyl)-4-phenyl-1H-1,2,3-triazole) to the $[Pd(\eta^3-C_3H_5)]^+$ can be evaluated with other neutral bidentate ligands consisting of thioether groups and can generate six-membered rings with the metal center.^{169–173} This coordination feature was assessed



Scheme 20 Structure of the $[Pd(\eta^3-C_3H_5)(2-((4-phenyl-1H-1,2,3-triazol-1-yl)methyl)-pyridine)]^+$ complex.



Scheme 21 The solid-state structures under investigation by their molecular drawing ($R = Ph$).



computationally in a number of complexes through adopting starting geometries where the coordination of the triazole fragment occurred through the N1 nitrogen atom. The calculated results demonstrated that, in all cases, the obtained optimized structure was similar to the X-ray structural data, where a Pd–N2 bond was clearly formed.¹⁷⁴

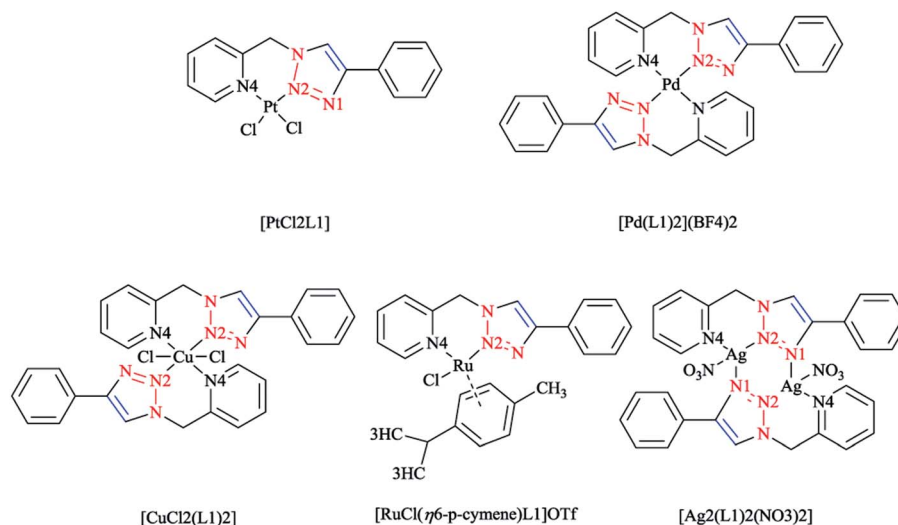
In another research study, experimental and theoretical DFT studies on the structural properties of $[\text{Pd}(\text{L})_2](\text{BF}_4)$ CH_3CN ($\text{L} = 4\text{-phenyl-1-(2-picolyl)-1,2,3-triazole}$) as a bischolate complex showed that the central square-planar palladium(II) cation was coordinated by two L substrates in *trans* position, each through pyridine and triazole N2 nitrogen atoms, generating two six-membered metallacycles. More precisely, *anti-trans*- $[\text{Pd}(\text{L})_2]^{2+}$ and *syn-trans*- $[\text{Pd}(\text{L})_2]^{2+}$ were produced as two boat-like conformations, in which the picolyl methylene carbons were *anti* or *syn*, respectively, depending on the palladium coordination plane, and could be easily converted to each other in solution (Scheme 21). It should be mentioned that $[\text{Pd}(\text{L})_2](\text{BF}_4)\text{CH}_3\text{CN}$ was synthesized from 4-phenyl-1-(2-picolyl)-1,2,3-triazole (L) and $[\text{Pd}(\text{CH}_3\text{CN})_4](\text{BF}_4)_2$ in CH_3CN .⁹⁹

Frontier molecular orbital analysis was also performed for palladium(II) complexes of sulfur–selenium 1,2,3-triazole-based ligands using DFT calculations to rationalize the nature of bonding within these complexes and their reactivity. In both the complexes, the HOMOs were mainly located over Pd and Cl atoms and included the interaction of a d orbital of Pd(II) and a p orbital of chlorine atoms. More precisely, in the HOMO-1 of both complexes' metal d orbitals, p orbitals of a chalcogen donor atom, chlorine atoms, and one of the nitrogen of the triazole ring overlap with each other. Since the high reactivity of a complex may be partially related to a low value for the HOMO–LUMO energy gap, among the two complexes, complex **15**, which had the lower HOMO–LUMO energy gap, was supposed to be a better catalyst than **16**, as was observed experimentally.¹²⁴

In another research study, the chelation properties of some mononuclear bis-bidentate triazole-based complexes (as illustrated in Scheme 22) were reported *via* X-ray crystallographical analysis. The X-ray structures of Pt and Pd complexes showed a square-planar coordination environment of the central Pt and Pd. The coordination sphere around the platinum atom consisted of two nitrogen atoms of the ligand and two chloride ions in the *cis* position, while in the palladium mononuclear bis-bidentate complex, the palladium ion is *trans*-chelated by two ligands. There was a close agreement between the bond distances in the coordination sphere of platinum in its complex with those found in the X-ray crystal structure of other square-planar complexes of Pt(II).¹⁷⁵ In the crystal geometry of the palladium complex, it was shown that the diagonal bonds are nonequivalent, so that N2–Pd is shorter than N4–Pd. In a copper complex, the metal ion is in a distorted-octahedral coordination environment with two additional chloride ions involved, and the ligands are located in a *trans* configuration comparable to that of the palladium complex. This displacement of two axial N2 atoms from the basal plane formed by two pyridine N4 atoms and two chloride ions leads to a pyramidal distortion of the octahedral geometry. In another reported ionic complex of $[\text{Cu}(\text{L})_2(\text{CH}_3\text{CN})-(\text{ClO}_4)](\text{ClO}_4)$,¹⁷⁶ the copper cation displays the square-planar geometry of four nitrogen atoms with CH_3CN and ClO_4^- ligands at the axial positions.

In the crystal structure of the ruthenium complex, a three-legged piano-stool geometry with the metal center coordinated by *p*-cymene in an η^6 fashion, a chloride ion, and ligand L1 was observed. The distances between Ru(II) and the nitrogen atoms of L1 were not equivalent, similarly to Pt and Pd complexes.

The X-ray geometry of the Ag complex showed two silver centers bridged by two L1 molecules. Both metal ions in the structure were coordinated by two nitrogen atoms from one L1 ligand (N4 from the pyridine and N2 from the triazole part) and another triazole N1 atom from the second L1 ligand in the



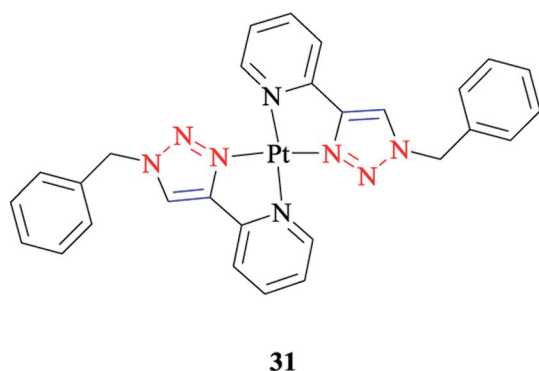
Scheme 22 Structures of $[\text{PtCl}_2\text{L1}]$, $[\text{Pd}(\text{L1})_2](\text{BF}_4)_2$, $[\text{CuCl}_2(\text{L1})_2]$, $[\text{RuCl}(\eta^6\text{-p-cymene})\text{L1}]\text{OTf}$ and $[\text{Ag}_2(\text{L1})_2(\text{NO}_3)_2]$ complexes based on 1-(2-picolyl)-4-phenyl-1H-1,2,3-triazole ligand (L1).



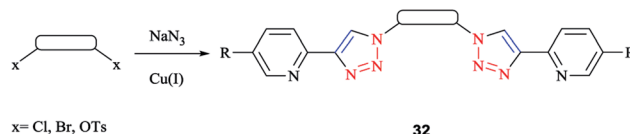
complex. This showed a distorted tetrahedral coordination consisting of the oxygen atom of a nitrate anion coordinated to both silver atoms in a monodentate mode. Alternatively, in the crystal structure of the Ag complex, each silver atom can be displaced from the plane of the coordinating nitrogen atoms and then coordination of the silver atoms can also be assumed to involve a distorted trigonal planar. In the aromatic L1, it was observed that the angle between the plane of the pyridine ring and the plane of the triazole ring was 87.7° and a distortion was found when ligand was bound to the metal center. Coordination and structural assessments of traditional tetrahedral complexes of Ag(I) and Cu(I) with 1,2,3-triazol-4-yl-pyridine click ligands revealed that these complexes can be used as appropriate motifs for the self-assembly of supramolecular architectures.¹⁷⁷

For computational investigation of this series of complexes, in accordance with our elucidations, the complexes with central atoms of copper, palladium, and platinum, $[\text{ML}_n(\text{NH}_3)_2]^{2+}$ ($\text{M} = \text{Cu}, \text{Pd}, \text{Pt}$), were considered with a coordination number of 4, while silver complexes $[\text{AgL}_n\text{NH}_3]^+$ were modeled with a coordination number 1. It is important to notice that the expected X-ray square-planar ($\text{M} = \text{Pd}, \text{Pt}$) and trigonal-planar ($\text{M} = \text{Ag}$) geometries were obtained with a singlet ground state *via* DFT computations. The natural bond orbital (NBO) analysis affirmed the donation of the in-plane lone pairs of the appropriate nitrogen atoms of the ligand and two ammonia to the empty s and d orbitals of the metal, which decreased the charge localized on the metal. On the other hand, since in $[\text{AgL}_n\text{NH}_3]^+$ the d orbitals are filled, electron donation occurs only to the 5s orbital of the metal, which leads to a weaker metal-ligand interaction. It should be mentioned that these calculated results are in agreement with the observed smallest stabilization energy for $[\text{AgL}_n\text{NH}_3]^+$ complexes.

Frontier molecular orbital analysis demonstrated that the d_{z^2} orbital of the metal (d_{z^2} is σ interacting in the D_{4h} symmetry) is combined with the high-lying orbitals of the triazole ring that can be observed in this series of complexes. It was also deduced that due to these interactions, the metal d electrons delocalize more efficiently to the π system of the ligand and so increase the aromaticity and also the stability of the complexes with the



Scheme 23 Structure of the $[\text{Pt}(\text{Bnpytri})_2]^{2+}$ (31) complex.



Scheme 24 Formation of the bis-bidentate pyridyl-triazole ligand (32).

ligand. This stabilization effect was confirmed by the second-order perturbation theory analysis of the Kohn–Sham matrix.⁹⁹

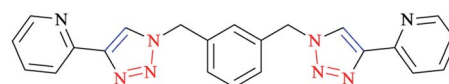
The geometrical and energetic properties of $[\text{Pt}(\text{Bnpytri})_2]^{2+}$ 31 complexes (where Bnpytri = 2-(1-benzyl-1H-1,2,3-triazol-4-yl)pyridine and its schematic structure are illustrated in Scheme 23), and the corresponding palladium(II) analogs were assessed computationally through DFT calculations at the B3LYP/LanL2DZ/6-31G(d) level of theory and using dimethyl sulfoxide (DMSO) as the solvent. The calculated results of $[\text{Pt}(\text{Bnpytri})_2]^{2+}$ 26 complexes revealed that the head-to-tail configuration was significantly favored from a thermodynamic point of view, as was obtained similarly for the corresponding palladium(II) analog.^{56,125} Under special conditions that allow the formation of the thermodynamic product, the head-to-tail configuration would be the dominant form for the Pt(II) complexes.

Moreover, for tetramer complexes, $[\text{Pd}_2\text{Pt}_2(\text{L})_4]^{8+}$ and also $[\text{Cu}_2\text{Pt}_2(\text{L})_4]^{6+}$ (Scheme 12), DFT geometry optimization was performed at the B3LYP/LanL2DZ/6-31G(d) level of theory in the presence of nitromethane as the solvent. The $[\text{Pd}_2\text{Pt}_2(\text{L})_4]^{8+}$ tetramer complex showed the expected box-like geometry, while optimization for the hexamer (with hexyloxy chains) gave the expected geometry. It was also extracted that applying a labile metal with a tetrahedral geometry might have less scope for the production of larger macrocycles, in addition to closing the Pt(II) centers and stabilizing the tetramer through platinum–platinum and π -stacking interactions.

In the case of di-magnetic Cu(I) and Ag(I) $[\text{M}(\text{Rpytri})_2]^+$ complexes, the calculated results showed that there was a tendency to adopt a head-to-tail square-planar geometry with bispyridyl-triazoles,^{129,133} contrary to the primary preference of Ag(I) for a tetrahedral geometry over a square-planar geometry.¹⁷⁸

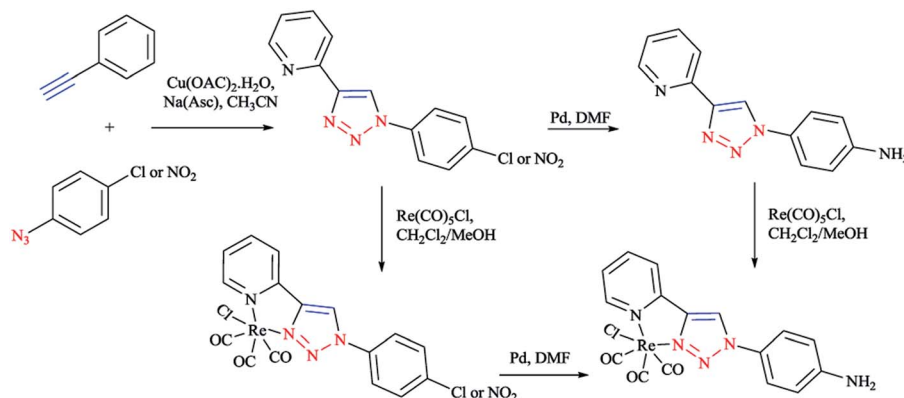
The crystal structure and DFT calculated structural data for $[\text{Cu}_2\text{Pt}_2(\text{L})_4]^{6+}$ complexes demonstrated their tetrahedral geometry,^{129,177,179} but only using the asymmetric ligands with orthogonal binding sites in combination with *cis*-protecting directional bonding strategies at the Pt(II) centers.^{139,180,181}

A family of bis-(2-pyridyl-1,2,3-triazole) ligands 32 (Scheme 24) were synthesized by employing *in situ* azide formation conditions and then by methylene-bridging of the bis-(2-pyridyl-



Scheme 25 Structure of the bis-2-pyridyl-1,2,3-triazole ligand (33).





Scheme 26 Production of bidentate 2-pyridyl-1,2,3-triazole ruthenium complexes.

1,2,3-triazole) ligands **32** to produce larger heterometallic macrocycles.

$[\text{Pt}(\text{L})_2]^{2+}$ metallo-ligands showed a square-planar structure with 2-pyridyl-1,2,3-triazole **33** units adopting a head-to-tail arrangement with vacant peripheral binding sites.

In this vein, tetracopper(i) macrocycles were prepared using the bis-(2-pyridyl-1,2,3-triazole)-1,2-xylylene ligand so that each pair of copper(i) ions was bridged by two iodides.

Furthermore, $[\text{Cu}_2\text{L}_2]^{4+}$ macrocycles were generated from Cu(II) ions and bis-2-pyridyl-1,2,3-triazole ligands *via* embedding a *m*-xylene (1,3-) or 2,7-dimethylnaphthalene spacer unit^{21,182} (Scheme 25).

4.3 Re(I) complexes

In Scheme 26, the production of $\{\text{Re}(\text{CO})_3\text{Cl}\}$ bidentate 2-pyridyl-1,2,3-triazole coordination compounds was illustrated *via* the reaction of the cycloadducts with the commercially available $[\text{Re}(\text{CO})_5\text{Cl}]$.

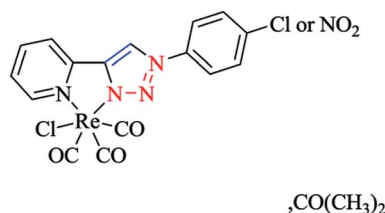
X-ray diffraction analysis was also performed to identify the structural properties of the aforementioned rhenium complexes and the results compared with the DFT-optimized geometries (*via* C-PCM computations in methanol solution). In the optimized structures of the complexes, the rhenium ion attains the usual distorted-octahedral coordination. It should be noted that the obtained bond lengths and angles were in the usual range of tricarbonyl rhenium complexes based on bidentate 2-pyridyl-1,2,3-triazole derivatives⁵⁹ (Scheme 27).

Polcar and coworkers deeply investigated the complexes of Re(I) centers coordinated with a series of functionalized 1,2,3-triazole ligands in an octahedral geometry as noninvasive

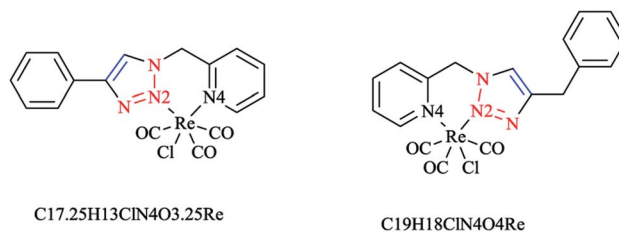
imaging probes. In this vein, a series of pyridyl-1,2,3-triazole and quinoliny-1,2,3-triazole ligands bearing various substituted alkyl side chains were designed and synthesized using click procedures. Furthermore, their photophysical properties were characterized and it was shown that the luminescent properties are enhanced in complexes bearing long alkyl chains.⁴⁴

In some rhenium complexes, the inverse click ligands are generally coordinated to the rhenium(I) center in a bidentate fashion through the pyridyl and triazole nitrogen atoms. In this context, the Re(I) complexes of the [(4-*R*-1*H*-1,2,3-triazol-1-yl)methyl] pyridine ligand series (Scheme 28) were investigated structurally through X-ray crystallography, which showed that all of them were crystallized in the triclinic space group.

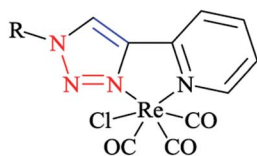
As expected, the rhenium metal center forms a distorted-octahedral coordination environment and is coordinated to a chloride and three carbonyl ligands in a facial array. Moreover, due to the presence of a methylene spacer between the triazole and pyridyl groups, a six-membered chelate ring is generated and consequently, the bite angles of the ligands vary little, ranging from 81.98(8) to 83.5(3)°. Chelation of the ligand



Scheme 27 Structure of the $[(\text{C}_{13}\text{H}_9\text{N}_5\text{O}_2)\text{ReCO}_3\text{Cl (or NO}_2)] \cdot \text{CH}_3 \cdot \text{COCH}_3$ complex.



Scheme 28 Structures of the $\text{C}_{17.25}\text{H}_{13}\text{ClN}_4\text{O}_{3.25}\text{Re}$, $\text{C}_{19}\text{H}_{18}\text{ClN}_4\text{O}_4\text{Re}$, and $\text{C}_{22}\text{H}_{22}\text{ClN}_8\text{O}_6\text{Re}$ complexes.

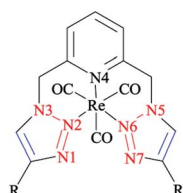


R = toluene, 1-ethyl-4-methoxybenzene, methyl acetate, 1-(2-methoxyphenyl)-4-propylpiperazine

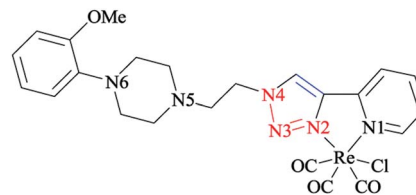
Scheme 29 Structure of the [(N-N)Re(CO)₃Cl] regular five-membered rhenium click complexes.

to the metal center leads to the formation of six-membered rings in boat conformations, with the fold angles between the pyridyl and triazole rings ranging from 111.09 to 114.22°. It has been shown that the triazole ring substituent effect is not considerable on the metric properties of rhenium complexes and so it can be deduced that side-arm substitution leads to little electronic tuning of the ligand. Furthermore, in these [(4-R-1H-1,2,3-triazol-1-yl)methyl]pyridine **5** set of ligands, it was found that the N^{py}-Re bond length value (2.198(5)–2.218(9) Å) is bigger than the N_{trz}-Re bonds (2.155(2)–2.180(7) Å) that occur in other metal complexes of this ligand set. Further, it was assessed experimentally and computationally^{56,81,126} that metal complexes of the related regular 2-(1-R-1H-1,2,3-triazol-4-yl)pyridine **4** ligands were more stable than those with the [(4-R-1H-1,2,3-triazol-1-yl)methyl]pyridine **5** inverse click ligands. Clearly speaking, regardless of whether the triazole is coordinated to the Re center, the obtained values of the Re-N_{2trz} and Re-N_{2trz}-N_{4py} bond lengths and angles in the entire series of regular five-membered rhenium click complexes^{63,64,97,183,184} (Schemes 27 and 29) were very near to those of inverse six-membered rhenium click complexes, and this feature has been also studied and confirmed in the Pd(II) complexes of these ligands.^{56,95}

It was observed in the crystal structure of these complexes that double-stranded supramolecular polymers are formed *via* the many C-H...Cl hydrogen bonds and CH-π or face-to-face π-stacking interactions.^{185–188} It was also observed that the chloride of one complex bonds to the adjacent complex through the acidic methylene CH (Cl...H = 2.638(2) Å, Cl...C = 3.569(7) Å), leading to the formation of a single-stranded linear polymer. A face-to-face π-stacking interaction (centroid-centroid distance of 3.736 Å) occurs between the phenyl group and 1,2,3-triazole units on adjacent complexes to produce the double-stranded structure. In addition, CH-π interactions between the benzyl



Scheme 30 Structure of the 2,6-bis(4-substituted-1,2,3-triazol-1-yl)methylpyridine complexes (R = Ph, C₃H₇).



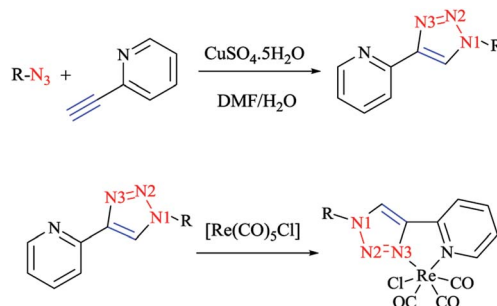
Scheme 31 Structure of the 4-piperazinyl-2-pyridyl-1,2,3-triazole rhenium(I) complex.

substituent methylene hydrogens and the phenyl unit of the adjacent complex were found with CH-centroid distances of 3.032 and 3.122 Å. More interestingly, in the caffeine-substituted complex, the individual complexes are folded into a U-shaped conformation and two complexes interact through C-H...O hydrogen bonds between a carbonyl of the caffeine moiety and an acidic CH of the pyridyl ring, which consequently leads to a supramolecular cage that is filled with two acetone molecules as the solvent. These face-to-face π-stacking interactions result in the formation of a slipped stacked polymer with a centroid-centroid distance of 3.381 Å between the caffeine groups.

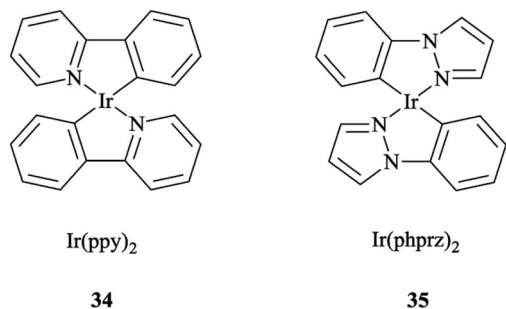
The structures of 4-*n*-propyl- and 4-phenyl-substituted *fac*-Re(CO)₃ complexes coordinated with the tridentate click ligand (2,6-bis(4-substituted-1,2,3-triazol-1-yl)methyl)pyridine (Scheme 30) were also investigated *via* X-ray crystallography and DFT methods. Based on the X-ray and DFT computational results, it was shown that both complexes had the expected facial coordination geometry about distorted-octahedral rhenium(I) centers. More concisely, the Re(I) cations were coordinated to three carbonyl donors and to the pyridyl and triazolyl nitrogen atoms of the ligands.⁵⁸

In another research study, the complex of rhenium(I) with functionalized 2-pyridyl-1,2,3-triazole derivatives (pyta)-based ligands was crystallized by slow evaporation in dichloromethane/methanol solution and then investigated using the X-ray diffraction method.

A comparative study between the structural features of rhenium(I) complexes of the related pyta derivatives^{82,133} exhibited an anti-arrangement in them adopted by the N(1) and N(2) atoms of the triazole and pyridine rings, respectively. In the 1,2,3-triazole ring, the N=N bond length of the 1,2,3-triazole



Scheme 32 Syntheses of [Re(CO)₃Cl] with bidentate pyridyl-1,2,3-triazole click ligands and [(pytri-R)Re(CO)₃Cl] complexes.

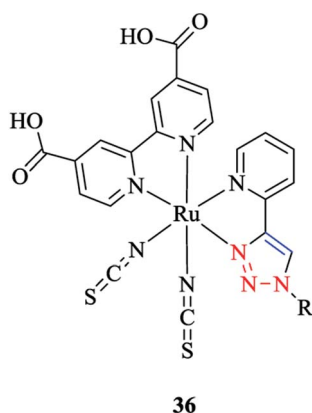


Scheme 33 Structures of the Ir(ppy)₂ (34) and Ir(phprz)₂ (35) complexes.

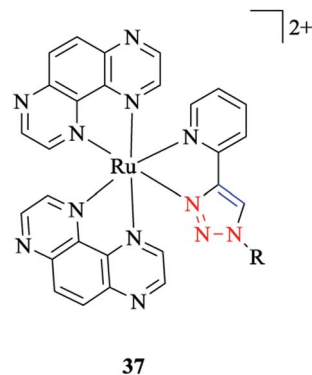
ring was decreased and became shorter than the adjacent N–C and N–N bonds. Moreover, a network of intra-layer hydrogen bonds between the ligands and lattice waters was observed that led to the crystal cohesion. Consequently, the lattice water molecule forms O–H...N bonds with one nitrogen of the piperaziny moiety ($d(\text{O–H}\cdots\text{N}(5)) = 3.013 \text{ \AA}$, $\text{O–H–N}(5) = 173.08^\circ$) (Scheme 31). It was observed that the ethylene bridge leads to a large distance between the (2-methoxyphenyl)piperazine group with the chelating part of the molecule. So, the components near the coordination sphere do not interact through coordination, hydrogen-bonding interaction, or by folding of the pendant arm toward the metal center, which was previously reported in other flexible spacers by Santos' group.¹⁸⁹ The structural results show that the interaction of the metal center with the bioactive fragments is not considerable and so this suggested a suitable affinity for the serotonin receptors.⁶³

Time-dependent DFT (TD-DFT) calculations¹⁹⁰ on Re(I) complexes of readily functionalized bidentate pyridyl-1,2,3-triazole click ligands (Scheme 32) demonstrated the MLCT nature of the HOMO $\rightarrow$ LUMO transitions (the blue-shift of Re $\rightarrow$ bpy MLCT, which lies at 633 nm).

It was also found that the emission from Re $\rightarrow$ pytri-R MLCT (where R = benzyl) is blue-shifted at 538 nm in CH₃CN. It should be mentioned that the electronic role of pytri-R in communication between the differing emitting centers has also



Scheme 34 Structure of a ruthenium complex with the (pytri-R)(bpy(CO₂H)₂)(CNS)₂ ligand.



Scheme 35 Structure of a ruthenium complex with the (TAP)₂(pytri-R) ligand.

been studied systematically.⁸⁸ It was reported that Ir(ppy)₂ 34 and Ir(phprz)₂ 35 (where phprz is 1-phenylpyrazole) (Scheme 33) fragments could bind to pytri-R and were then connected *via* the (N1) sp² triazole nitrogen to produce dimer and trimer structures. Furthermore, it was shown that the modulation of the linkers through the N1 sp² triazole did not affect the photo-physical properties of the individual Ir units.

The coordination behavior of ruthenium ion with (pytri-R)(bpy(CO₂H)₂)(CNS)₂ as a pytri-R based ligand was investigated *via* DFT analysis⁸⁵ (Scheme 34).

It should be mentioned that these complexes are widely used in dye-sensitized solar cells. DFT computations showed that the LUMO and LUMO+1 of [Ru(pytri-R)(bpy(CO₂H)₂)(CNS)₂] 36 were based on the bpy ligand. Moreover, a systematic study of the substituent effects on the ruthenium complexes with substituted pytri-R ligands was performed by Happ *et al.*⁸⁷ In this respect, the pytri-R ligand was substituted with ethynylphenyl groups at either the 5-pyridyl or N1 triazole positions, and it was shown that substitution of the 5-pyridyl position had a significant effect on the emission properties and increased the emission wavelength from 602 nm with the 1-ethynyl-4-methoxybenzene substituent to 674 nm with the 1-ethynyl-4-nitrobenzene substituent. Interestingly, the [Ru(pytri-R)(bpy(CO₂H)₂)(CNS)₂] 36 complex substituted by 1-ethynyl-4-nitrobenzene at N1 triazole nitrogen showed the emission wavelength at 610 nm. This behavior revealed that the linkage of the 1-ethynyl-4-nitrobenzene substituent with the triazole insulated the electron-withdrawing power of this substituent from the Ru center.

In another research study, the properties of [Ru(TAP)₂(pytri-R)]²⁺ 37 complexes with a DNA active moiety (where TAP = 1,4,5,8 tetraazaphenanthracene) were also assessed (Scheme 35).¹⁹¹



Scheme 36 Structure of the Re(CO)₃Cl complex series with 2-(1-R-1H-1,2,3-triazol-4-yl)pyridine.



It was demonstrated that the d and p orbitals of ruthenium are stabilized at 50 mV relative to L = 1,10-phenanthroline, which could be attributed to a poorer overlap between the $d\pi$ and π^* (pytri-R) molecular orbitals. Moreover, it was investigated and shown that alkaline and selected first row transition metals have no considerable effect on varying levels of emission intensity and no tuning of the emission frequency.¹⁹² On the other hand, the N1 substituent does not shift the emission spectrum for [(pytri-R)Re(CO)₃Cl] complexes. For instance, by using R = (CH₂)₂CO₂H and (2-methoxyphenyl) piperazine as substituents in the [(pytri-R)Re(CO)₃Cl] complex, the emission maxima were seen to lie at 526 nm and 522 nm in DMF/MeOH solution, respectively.^{63,183} DFT calculations on the molecular orbital properties of these systems demonstrated a modest contribution from the N1 substituent in the LUMOs.

Moreover, a series of electronically tuned *fac*-Re(CO)₃Cl·2-(1-R-1*H*-1,2,3-triazol-4-yl)pyridine complexes were prepared (Scheme 36). In this systematic study, the substituent effect on the electronic structure of a series of *fac*-[(pytri-R)Re(CO)₃Cl] complexes was examined. The substituents were chosen so that they could couple conjugatively with the triazole and could insulate it *via* a CH₂ group from such communication. The nature of the lowest energy transition resonance was studied *via* Raman spectroscopy in conjunction with DFT calculations. The lowest excited state properties were also established using time-resolved emission spectroscopy. Then, the calculated values of a number of bond lengths and angles were compared with X-ray crystallographical data, and it was seen that there was a reliable agreement with the experimental data.

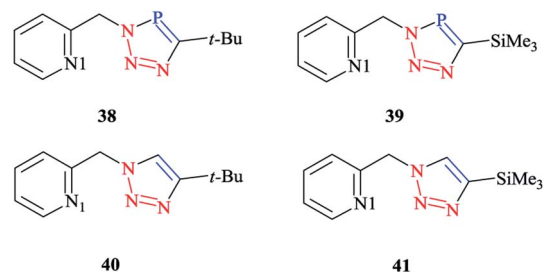
A comparison of the calculated Raman wavenumbers and intensity values with the corresponding experimental data showed a mean absolute deviation (MAD) of <10 cm⁻¹ for bands between 400 and 1650 cm⁻¹ that had more than 20% of the maximum band intensity.

The electronic absorption behavior of all the complex series was investigated. The lowest energy transition at approximately 330 nm was not changed by substitution, regardless of whether was unconjugated or conjugated with the triazole ring.

The band at 330 nm could be assigned to a metal-ligand charge transfer (MLCT)⁹⁷ as was confirmed by the TD-DFT data. The higher energy MLCT transitions (at 290 nm) and additional transitions associated with the ligand (at 275 nm) were obtained *via* TD-DFT computations. It is important to note that DFT methods systematically overestimate the conjugative effects and so, the effect of the conjugated group was predicted computationally to be stronger than the experimental values by up to 0.4 eV.^{193,194}

Moreover, it could be observed that conjugated and non-conjugated ligands affect the nature of the MLCT band at 330 nm. Precisely speaking, the acceptor orbitals are localized solely on the triazole species for the unconjugated complexes, while in the case of conjugated complexes, they are spread over the R substituent in addition to the triazole moiety.

This different behavior can be attributed to the overweighing of the conjugation interactions associated with DFT methods, which can be examined through measuring the resonance Raman spectroscopy.



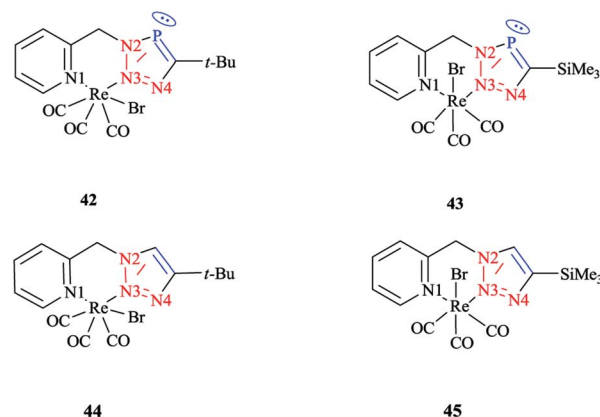
Scheme 37 Structures of click-derived chelating phosphorus- and nitrogen-containing heterocycles.

It was shown that different substitution on the triazole ring resulted in different Raman spectra; for instance NO₂-substituted complexes showed a distinguished band at 1349 cm⁻¹ attributed to NO₂ functionality.^{195,196}

A significant aspect was observed in the resonance Raman spectra, which was the identical spectra for the complexes with similar pytri species (bands at 1285, 1571, 1587, and 1625 cm⁻¹). It should be stated that DFT normal mode calculations also produced these bands and showed that they are based on the pytri moiety. Small differences in the Raman spectra of the conjugated systems were observed. It should be noticed that the Raman spectra showed pytri modes, which were also observed for non-conjugated systems. Two bands were observed due to the stretching of the R-triazole moiety and an enhancement of the NO₂ group^{195,196} was present, which was in agreement with the acceptor molecular orbital nature of a nitro functional group.

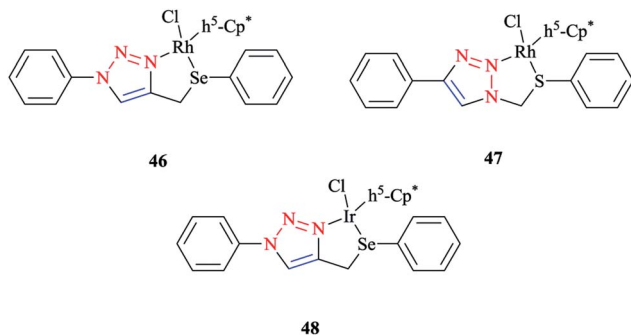
From the computational viewpoints, the DFT calculated values revealed that the LUMO was nitrobenzene based, and also the TD-DFT computations for the lowest energy transitions demonstrated no or only a minor LUMO character. The calculated results were in confirmation with the resonance Raman data, which indicated an enhancement of nitro vibration, reflecting the role of the nitro group on the electronic transition.⁶⁴

In order to compare the stereoelectronic properties of triazoles with triazaphospholes, Re(I) complexes containing these



Scheme 38 Syntheses of Re(I) complexes 35–38.





Scheme 39 Structures of Rh(III)/Ir(III) complexes with 1,2,3-triazole-based organochalcogen ligands.

chelating ligands were assessed (Scheme 37). The coordination behavior of the Re(I) cation with the least nucleophilic nitrogen atom N3 in complexes **38–41** is depicted in (Scheme 38).

The crystallographic properties for two complexes of $[L_2Re(CO)_3Br]$ were comparatively characterized. The crystal structure revealed that the Re(I) cation is coordinated to chelating ligands *via* the N(3) atom of the triazaphosphole species, rather than *via* the phosphorus atom. Since, coordination through the N(3) atom of the triazaphosphole species has not been observed for triazaphospholes before, it can be claimed that this behavior was enforced by the chelating effect of the ligand.

As was observed in the crystal geometry of compounds **42** and **43**, the nitrogen atoms N(3) of the triazole units were coordinated to the Re(I) centers in the facial geometry of complexes **44** and **45**, $[(40)Re(CO)_3Br]$ and $[(41)Re(CO)_3Br]$. Moreover, we can claim that replacing the phosphorus atom in **42** and **43** by an isoelectronic C–H group in **44** and **45** had no considerable effect on the bite angle N(1)–Re(I)–N(3).

DFT calculations at the B3LYP level of theory⁵⁴ were performed on compound **42** ($R = t\text{-Bu}$) to understand the electronic structure of the triazaphospholes. A large coefficient of π symmetry at the phosphorus atom was obtained on the LUMO of low-coordinate phosphorus heterocycle compounds, which showed the π -accepting nature of these systems. In HOMO-1,

a large value of the coefficient of σ symmetry was obtained at the nitrogen atoms N4 and N3 of the heterocycle ring and HOMO-7 clearly indicated the π -donor properties of triazaphospholes.

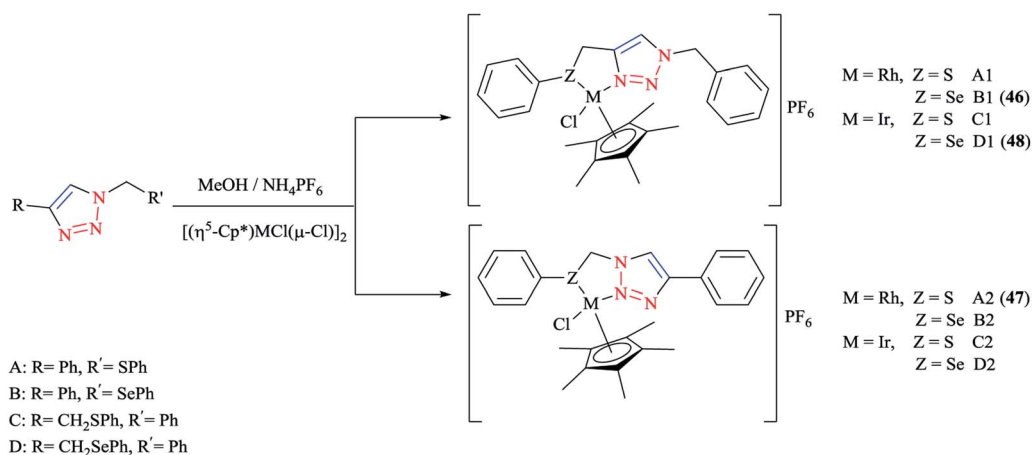
It is important to mention that transition metal cation centers coordinate preferentially to the nitrogen atom N3 in triazoles, but in the ligands of **38** and **39** this nitrogen atom cannot coordinate *via* σ bonds due to the chelating effects. Furthermore, in the HOMO-4 orbital, similar large coefficients of σ symmetry at both the phosphorus atom and nitrogen atom N3 appeared. More importantly, the coordination of a transition metal center with the pyridyl nitrogen and the nitrogen N3 or with the pyridyl nitrogen and the phosphorus *via* a σ coordination mode led to a small energetic difference, and so both coordination modes could be possible.⁶⁶

4.4 Rh(III) and Ir(III) complexes

Some 1,2,3-triazole-based organochalcogen ligands and their Rh(III)/Ir(III) complexes (Schemes 39 and 40) were synthesized and characterized using elemental analysis, multinuclear NMR, IR, X-ray crystallography, and mass spectral data.

To synthesize the complexes, the chloro bridge cleavage of $[(\eta^5\text{-Cp}^*)RhCl(\mu\text{-Cl})]_2/[(\eta^5\text{-Cp}^*)IrCl(\mu\text{-Cl})]_2$ occurred, followed by the reaction with 1-benzyl-4-((phenyl-thio/phenylseleno)methyl)-1H-1,2,3-triazole (A/B) or 4-phenyl-1-(phenylthio/phenylseleno)methyl-1H-1,2,3-triazole (C/D) at room temperature. Consequently, rhodium(III) (A1, B1, A2, and B2) and iridium(III) (C1, D1, C2, and D2) half-sandwich complexes were generated and the process was facilitated by anion exchange with NH_4PF_6 .

The nitrogen of triazole five-membered ring is coordinated to the metal center in all of the complexes. Almost similar X-ray molecular structures were obtained for all the rhodium and iridium complexes, which demonstrated an overall three-legged piano-stool conformation. More clearly, a pseudo-octahedral half-sandwich “piano-stool” disposition of donor atoms is formed around the Rh/Ir metal center. The centroid of the $\eta^5\text{-Cp}^*$ ring occupies nearly the center of a triangular face of an octahedron. The coordination sphere is occupied with



Scheme 40 Syntheses of A–B ligands and A1–D1 and A2–D2 complexes.



a chlorine donor atom and nitrogen and chalcogen atoms, which form a chelate ring with the metal center.

In comparison with complex **46**, the Rh–Se bond length of the half-sandwich complex of Rh(III) with the ligand 2-(phenylselenomethyl)pyridine was 2.487 Å (ref. 197) and for the complex $[\eta^5\text{-Cp}^*\text{RhCl}\{\eta^2\text{-(SePPh}_2)_2\text{N}\}]$ was 2.5266 Å,¹⁹⁸ which were larger than those obtained for the half-sandwich complex of Rh(III) with the ligand 1,2-dicarba-*closo*-dodeca-borane-1,2-dichalcogen.¹⁹⁹

In complex **47**, the Rh–S bond distance was 2.432 Å and was also greater than the corresponding value for the complexes $[\eta^5\text{-Cp}^*\text{RhCl}(1,1'-(1,2\text{-ethanediy})\text{bis}(3\text{-methylimidazole-2-thione}))]\text{Cl}$ at 2.3967 Å (ref. 200) and $[\eta^5\text{-Cp}^*\text{RhCl}\{2\text{-(phenylthiomethyl)pyridine}\}]\text{PF}_6$ at 2.383 Å (ref. 197) and $[\eta^5\text{-Cp}^*\text{RhCl}\{\eta^2\text{-S,P-Ph}_2\text{P(S)NHPPH}_2\}]\text{BF}_4$ at 2.404 Å.²⁰¹ Furthermore, in complex **47**, the Rh–N bond distance was 2.091 Å, which was in reliable agreement with those of $[\eta^5\text{-Cp}^*\text{Rh}(2,6\text{-(mesityl)}_2\text{C}_6\text{H}_3\text{S})(\text{bpy})][\text{B}(3,5\text{-(CF}_3)_2\text{C}_6\text{H}_3)_4]$.²⁰² The obtained bond length values for Rh–C(Cp* centroid) in complexes **46** and **47** were in the normal range, *i.e.*, 1.784–1.786 Å.²⁰¹

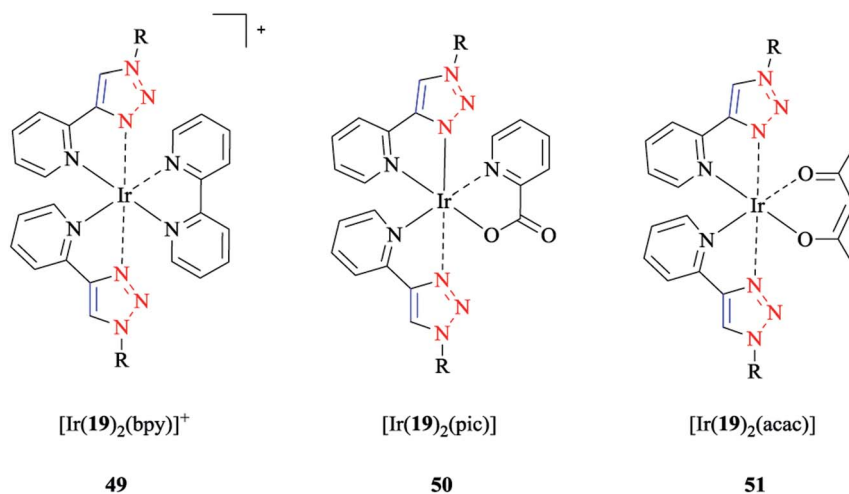
In the case of complex **48**, the Ir–Se bond distance value was 2.4822 Å, which was longer than the values obtained for the complex $[\eta^5\text{-Cp}^*\text{Ir}\{\text{Se}_2\text{C}_2(\text{CO}_2\text{Me})_2\}]\text{Cl}$ (2.3494 and 2.3520 Å)²⁰³ and near to the distances reported for $[(\eta^5\text{-Cp}^*)\text{Ir}(\{2\text{-arylselenomethyl} \} \text{pyridine})\text{Cl}][\text{PF}_6]$ at 2.453 Å,¹⁹⁷ $[\eta^5\text{-Cp}^*\text{IrCl}(\mu\text{-SeCOC}_6\text{H}_5)(\kappa^2\text{-SeCOC}_6\text{H}_4)\text{Ir}(\eta^5\text{-Cp}^*)]$ at 2.445–2.495 Å,²⁰⁴ and $[\eta^5\text{-Cp}^*\text{Ir}(\mu^3\text{-Se})_2(\text{PtTol}\text{-}(\text{PPh}_3)_2)]$ at 2.416–2.422 Å.²⁰⁵ Moreover, the Ir–N bond distance was 2.094 Å, which was very close to the Ir–N bond length in $[(\eta^5\text{-Cp}^*)\text{Ir}(2\text{-arylthiopyridyl})\text{pyridineCl}][\text{PF}_6]$ at 2.099 Å (ref. 197) and shorter than those of Ir complexes of the pyridine-triazole ligand at 2.137 Å (ref. 29) and 2-(1-benzyl-1*H*-1,2,3-triazol-4-yl)pyridine at 2.151–2.159 Å.²⁰⁶ The Ir–C ($\eta^5\text{-Cp}^*\text{centroid}$) bond length in complex **48** was 1.791 Å, which was consistent with the Ir–C bond distance in $[(\eta^5\text{-Cp}^*)\text{Ir}(\{2\text{-arylselenomethyl} \} \text{pyridine})\text{Cl}][\text{PF}_6]$ at 1.781 Å and shorter than the reported value for the complex $[(\eta^5\text{-Cp}^*)\text{Ir}(\text{phpy})\text{Cl}]$ at 1.863 Å.²⁰⁷

More interestingly, in complexes **46–48**, C–H...F secondary interactions and consequent chain-like structures have been found.

Theoretical analysis of the frontier molecular orbitals and charge distribution of complexes A1–D1 and A2–D2 were carried out through DFT approaches. Computational results on N(1)-coordinated complexes showed that the HOMOs were essentially located over the metal center and the π orbital of the Cp* ring. Clearly speaking, the HOMO orbitals consist of d orbital of the metal interacting with the π orbital of the Cp* ring and p orbitals of N(1) in the triazole ring and chalcogen atom, while in the HOMOs of N(2)-coordinated complexes, the π orbital of 4-phenyl-1,2,3-triazole interacts with the metal d orbital and π orbital of the Cp* ring and the p orbital of the chlorine atom. The HOMO–LUMO energy gap was also investigated to interpret the chemical reactivity of the complexes. The calculated results indicated that the HOMO–LUMO energy gaps of complexes A1, B1, A2, and B2 were smaller than those of C1, D1, C2, and D2.

It should be stated that the obtained result was in confirmation with the higher reactivity of Rh complexes in comparison to Ir complexes. Moreover, it can be predicted that among the complexes A1, B1, A2, and B2, complexes A2 and B2 could be expected to be more catalytically active than A1 and B1 complexes, and within the Ir complexes, the reactivity of complex D2 should be more than the other three Ir complexes. It should be mentioned that the obtained DFT results were in reliable agreement with the elucidations in terms of the catalytic activities of the various complexes.

From the electronic viewpoint, the coordination strength can be assumed to be a significant parameter in the stability of a complex. In complexes A1–D1, the calculated values of electron density at the coordinated N(1) of the triazole ring were quite high and so these complexes are more stable electronically, while for the complexes where N(2) of the triazole ring is involved in the coordination, the calculated electron density on the N(2) atom and consequently the stability of the complex was reduced. In conclusion, the electronic stability of complexes involving N(1) in coordination was greater than in those



Scheme 41 Structures of the $[\text{Ir}(\mathbf{19})_2(\text{bpy})]^+$ (**49**), $[\text{Ir}(\mathbf{19})_2(\text{pic})]$ (**50**), and $[\text{Ir}(\mathbf{19})_2(\text{acac})]$ (**51**) complexes based on 1-decyl-4-phenyl-1*H*-[1,2,3]triazole ligand (**19**).



involving N(2) and so their reactivity is less.^{37,56,126,208} It should be noted that the crystallographical structural data for complexes **46**, **47**, and **48** were in good agreement with the calculated values.

It is important to note that all DFT computations were performed using the B3LYP functional. In the case of Ir and Rh, effective core potential (LANL2DZ) calculations^{209–212} were applied for core electrons, while for other elements, the 6-311G** basis set were used.

The initial geometry coordinates were directly specified as a single-crystal X-ray structure. In order to establish the stationary points as the minima, frequency calculations were carried out at the end of the geometry optimization process and the molecular orbital properties were calculated with respect to the optimized coordinates using the same level of theory and the same basis set.²¹³

The frontier molecular orbital analysis of iridium(III) complexes with phenyl-1*H*-[1,2,3]triazoles showed that for all the complexes, the HOMO corresponded to the 5d_{xy}-orbital, while the LUMO was a π^* -orbital. In the case of [Ir(**19**)₂(bpy)]⁺ **49**, a π -orbital of the cyclometalating ligand was intercalated within the occupied 5d-orbitals. Interestingly, this antibonding orbital was located on the ancillary ligand in the case of [Ir(**19**)₂(bpy)]⁺ **49** and [Ir(**19**)₂(pic)] **50**, but was on the cyclometalating ligand in the case of [Ir(**19**)₂(acac)] **51** (schematic structures of the iridium(III) complexes are displayed in Scheme 41).

Since the strong ligand field effect of the phenyl anion ligand increased the gap between the t_{2g} and e_g orbitals, the e_g orbitals of Ir(III) were strongly destabilized in these complexes. The calculations demonstrated that the substituents on the cyclometalating ligands as well as on the ancillary ligand destabilized the HOMO and stabilized the LUMO, and so decreased the HOMO–LUMO energy gap.

The calculated values of the LUMO levels and the theoretical band gaps E_g showed particularly an overestimation compared to the corresponding experimental data. It should be mentioned that the relative change in the complex series was in

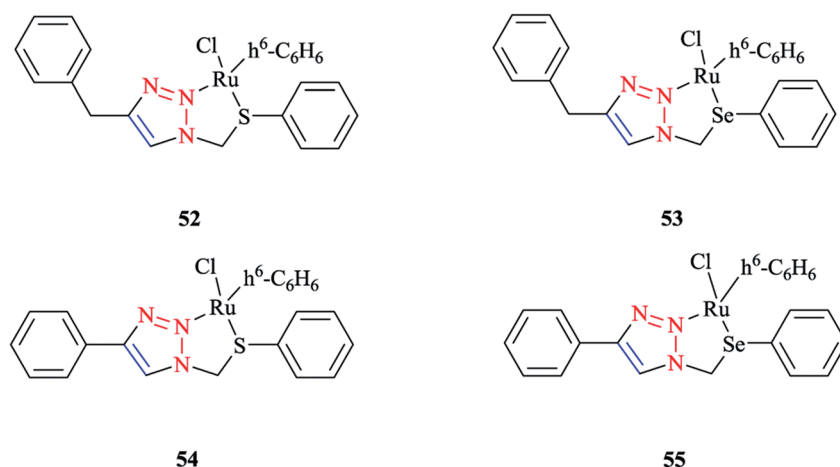
agreement with the LUMO energies and the HOMO–LUMO energy gaps ($E_g^{\text{acac}} > E_g^{\text{pic}} > E_g^{\text{bpy}}$).

The calculated results for [Ir(**19**)₂(acac)] **51** could be used to evaluate the influence of the cyclometalating ligand on the HOMO–LUMO gap. Apparently, 1,2,3-triazole ligands increase the energy gap in comparison with the ppy ligand-based complex, which can be attributed to the stabilization of HOMO and destabilization of LUMO, and so can be caused by the increased π -acceptor ability of the 1,2,3-triazole moiety. Moreover, back-bonding effects for receiving electron density in the π^* -orbitals of the ligand-centered π -system stabilized the occupied t_{2g} orbitals of the iridium complexes. On the other hand, it also destabilized the unoccupied π^* -orbitals of the ligands including the LUMO level.

Computational frontier orbitals assessment on the effect of the ancillary ligand (bpy, acac, pic) led to the following results: first, a larger band gap was obtained for the neutral compounds in comparison to the cationic [Ir(**19**)₂(bpy)]⁺ **49** complex and Ir(III) complexes. More precisely, the HOMO energies were stabilized in the cationic compound in comparison to in the neutral complexes. However, the LUMO was considerably more stabilized than the HOMO, and so the HOMO–LUMO energy gap was reduced. Second, it was shown that the major differences between the neutral compounds (*i.e.*, [Ir(**19**)₂(pic)] **50** and [Ir(**19**)₂(acac)] **51**) were mainly due to the LUMO energies rather than the HOMO energies through a comparison between the neutral compounds (*i.e.*, [Ir(**19**)₂(pic)] **50** and [Ir(**19**)₂(acac)] **51**). As mentioned earlier, both LUMOs showed a π^* -character, but were located on the cyclometalating and on the ancillary ligand, respectively. For the Ir [Ir(**19**)₂(pic)] **50** complex, the LUMO (π_{anc}^*) on the ancillary ligand was stabilized with respect to the π^* and orbital in [Ir(**19**)₂(acac)] **51**, which had higher energy values than the π^* -orbital of the cyclometalating ligands.¹¹⁴

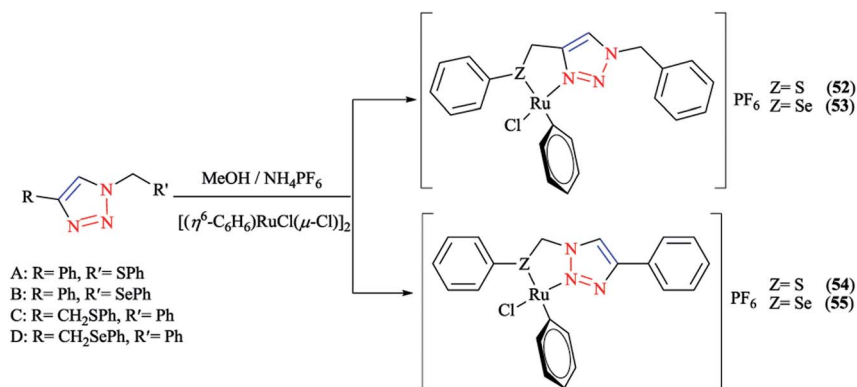
4.5 Ru(II) complexes

Next, we report the crystallographical assessments of some ruthenium complexes of click-generated 1,2,3-triazole-based organosulfur–selenium ligands (Schemes 42 and 43).



Scheme 42 Structures of Ru(II) complexes with 1,2,3-triazole-based organosulfur/selenium ligands.



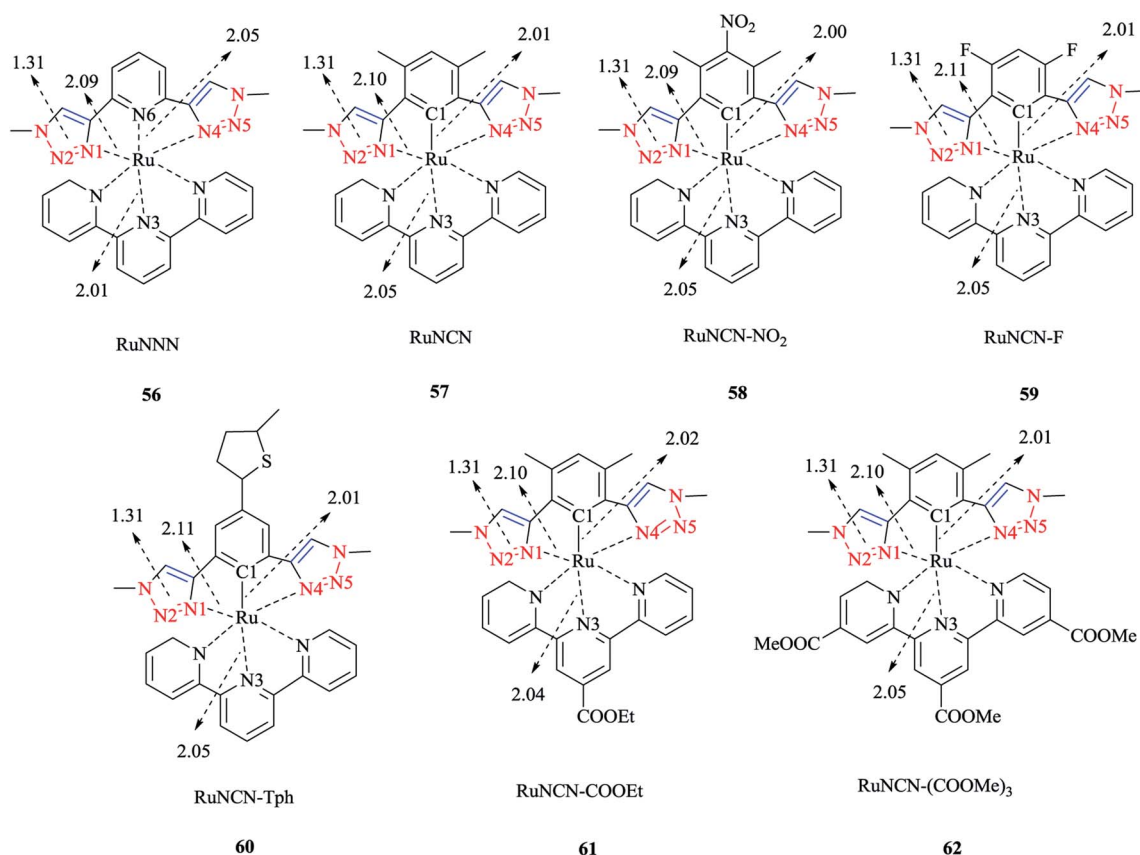


Scheme 43 Syntheses of A–D ligands and complexes 52–55.

A pseudo-octahedral half-sandwich “piano-stool” disposition of donor atoms occurs around the Ru metal center in complexes 52–55. More precisely, one face of an octahedron in all four complexes was occupied with a benzene ring and a chelate ring is formed with nitrogen and chalcogen atoms around the metal center, while a chlorine donor atom completes the coordination sphere. Overall, a three-legged piano-stool conformation was observed in the X-ray crystal structure.

The Ru–S bond lengths were obtained as 2.3847 Å in complex 52 and 2.3893 Å in complex 54, which are consistent with each other.

On the other hand, the Ru–Se bond distance in complex 53 was obtained as 2.497 Å, which was longer than that of complex 55 at 2.4859 Å, but was similar with the values for [(η⁶-C₆H₆)RuCl(N-{2-(phenylseleno)ethyl}pyrrolidine)]⁺ at 2.480 Å and for [(η⁶-C₆H₆)Ru(2-MeSC₆H₄CH=NCH₂CH₂SeC₆H₅)]²⁺ at 2.4848 Å. However, the Ru–Se bond length for [RuCp(CO)(C=CPh)(μ-Se)ZrCp₂] at 2.494 Å was in confirmation with that of complex 53. In complexes 52–55, the Ru–N bond lengths varied in a narrow range (2.072–2.101 Å) and were similar with the reported Ru–N bond distance for [Ru(2,6-bis(1-benzyl-1,2,3-triazol-4-yl)pyridine)₂](PF₆)₂ at 2.050 Å (ref. 214) and [(η⁶-C₆H₆)Ru(2-MeSC₆H₄CH=N-CH₂CH₂SC₆H₅)]²⁺ at 2.073 Å. In



Scheme 44 Structures of Ru(II) complexes with some selected calculated bond distances (Å).



addition, the Ru–C bond lengths of 2.154–2.22 Å and the C–Ru–C bond angles of complexes **52–55** were in the normal range.^{215–217} The PF₆ groups were involved in C–H...F secondary interactions in all the complexes **52–55**.

From the computational viewpoint, the cases of Ru–Cl, Ru–N, and Ru–C (centroid) showed an excellent agreement between the DFT calculated and experimental bond distances values. Furthermore, the X-ray values of the bond angles were very close to the calculated ones.

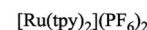
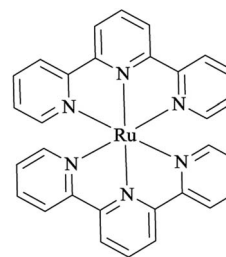
In the next step, DFT calculations were performed to assess qualitatively the optimized geometries, frontier orbitals, and charge distribution of complexes **52–55**. In complexes **52** and **53**, the HOMOs are positioned primarily over the Ru and Cl atoms and include the orbital of Ru(II) interacting with a π orbital of the η^6 -benzene ring, a p orbital of the triazole nitrogen, and a chalcogen atom. In complexes **54** and **55**, the HOMOs constitute a π orbital of 4-phenyl-1,2,3-triazole and a d orbital of the metal. However, the d orbital of Ru(II) interacts with the π orbital of the η^6 -benzene ring, the p orbital of triazole nitrogen, and a chalcogen donor atom in HOMO-1.

DFT calculations indicated that the HOMO–LUMO energy gaps in complexes **54** and **55** were lower than those of **52** and **53**, which corresponds to the higher reactivity of complexes **54** and **55**.

It is important to note that in complexes **54** and **55**, there is N(2) coordinating moiety, which can be attributed to the low energy gaps. On the other hand, the calculated electron densities of the coordinated N(1) triazole ring in complexes **52** and **53** were higher than the calculated value for N(2) in complexes **54** and **55**. This electronic feature can be interpreted as due to a strong coordination between N(1) of triazole and Ru in complexes **52** and **53**, which leads to greater electronic stability in comparison with **54** and **55**.³⁷

In another work, the crystal structure of Ru(II) complexes with NNN, NCN, NCN–NO₂, NCN–F, and NCN–Tph ligands were determined *via* the X-ray diffraction method and some considerable changes were observed attributed to the packing effects, such as a strong distortion within RuNCN–F.

In the RuNCN–Tph complex **60**, the thiophene-phenyl torsion angle was obtained as 30.78°, which confirmed the calculated results. This feature led to the partial extension of conjugation into the thiophene ring. It was shown that when a dative Ru–N bond of the poly-pyridyl-type complex RuNNN was replaced with a covalent organometallic Ru–C1 bond in RuNCN complex **57**, the bond distance was reduced from 2.02 to



63

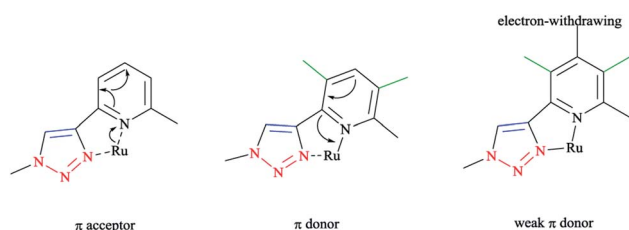
Scheme 46 Structure of [Ru(tpy)₂](PF₆)₂ complex.

1.98 Å. This behavior could be mainly attributed to s donation and additional p donation as well as by electrostatic interactions with the anionic, aromatic carbon donor. Moreover, the triazole N–Ru bond length was increased due to the decrease in the orbital overlap by the small bite angle, while, a trans influence occurs and the opposed Ru–N bond distance is increased to 2.01 Å based on the considerable electron-donation ability of the carbanion. More interestingly, the outer pyridine N–Ru bonds were shortened, which could be ascribed to the p back donation from the electron-rich Ru(II) metal center (Schemes 44 and 45).

Further, the bond length of the N–N double bond in the triazole ring was increased due to the increasing p back donation into the π^* orbitals. By the substitution of NO₂ as an electron-withdrawing group, it was indicated that the cyclometalation process was affected less than by installing a CN-withdrawing moiety in the RuNCN complex **57**.

Another important structural feature is the dihedral angle with the central phenyl ring (as illustrated in Scheme 44), which has a 51.88 X-ray determined value and 51.68 calculated value and can lead to a strong decrease in the π -donation ability of the cyclometalating carbanion in the *para* position, whereby consequently, the Ru–C1 bond is elongated to 2.00 Å.

In the case of the RuNCN–F complex **59**, it was found that substitution of the fluorine atom as a strong s-accepting group with a moderate p-donation character, in the *meta* position, has a not considerable effect on the carbanion and so results in a very short Ru–C1 bond of 1.98 Å. Moreover, it should be mentioned that a decrease in the bond length of the triazole N2–N3 double bond arises from the fluoro and nitro substituent effect, which causes a reduction of the p back electron-donation ability of the carbanion. Strictly speaking, the fluorine substituent essentially affects the σ donation and indirectly reduces the energy of the p system as an inductive effect. This occurs while the NO₂ group polarizes the p system additionally and so lowers its p-donation character *via* a mesomeric effect. It was also demonstrated for the RuNCN–F complex **59** that the HOMO is stabilized and located on Ru(II) metal center and also covers the fluoro-substituted cyclometalating phenyl ring. This occurs while an additional LUMO stabilization occurs in RuNCN–NO₂ complex **58** since the LUMO is located on the opposed tpy



Scheme 45 Schematic of the electronic features of the cyclometalation and an electron-withdrawing group.



ligand and so this can be related to the effect of the aromatic p system and d orbitals of Ru(II).¹¹⁶

In another report, the electronic properties of ruthenium(II) complexes with 2,6-bis(1,2,3-triazol-4-yl)pyridine 7 ligands were assessed using DFT calculations. The calculations showed that the highest occupied molecular orbital (HOMO) of the complexes included a metal d orbital and π orbitals located on the triazolate rings, which were in confirmation with the electron-rich π system of the anionic ring.^{218–220} Based on the electron repulsion between the anionic ligand and the metal center, the HOMO is strongly destabilized in comparison to polypyridyl complexes such as [Ru(tpy)₂](PF₆)₂ **63** (Scheme 46). For complexes with terpyridine ligands, the lowest unoccupied molecular orbital (LUMO) is mainly composed of π^* orbitals of the ligand. Considering the electron-withdrawing groups, the HOMO and, in particular, the LUMO are stabilized due to the electron-withdrawing effect, leading to a significantly smaller HOMO–LUMO gap. A similar effect on the frontier orbitals energies was observed using –COOH anchoring groups. In contrast, for a deprotonated complex, the HOMO and LUMO of the complex are destabilized and the HOMO–LUMO gap slightly increased, which is attributed to the electron-donating effect of the –COO-groups.

There was good confirmation between the calculated electronic excitations and the experimental UV-vis absorption spectrum. The lowest energy absorption was a pure HOMO–LUMO transition, which could be assigned to a metal-to-ligand charge transfer (MLCT) with some ligand-to-ligand charge-transfer (LLCT) character.¹¹⁷

5. Conclusions

In summary, 1,2,3-triazole-based ligands synthesized *via* CuAAC click reaction are introduced as diverse biological and pharmaceutical compounds and also their complexes with transition metals were widely applied as efficient catalysts in chemical reactions and reported here. The coordination properties of Ni, Pd, Pt, Re, Rh, Ru, and Ir transition metal complexes with a series of triazole-based click ligands were investigated *via* computational chemistry approaches by the calculation of their structural properties, electronic energies, and by frontier molecular orbital analysis. More clearly, in the aforementioned metal complexes of triazoles- and triazaphospholes-based ligands, their X-ray crystal structure, FT-IR and NMR spectroscopic features, metal-to-ligand charge transfer, *trans* and *cis* chelation modes, and emission properties were computationally assessed *via* comparison with the experimental data, which showed reliable agreement.

Conflicts of interest

There are no conflicts to declare.

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