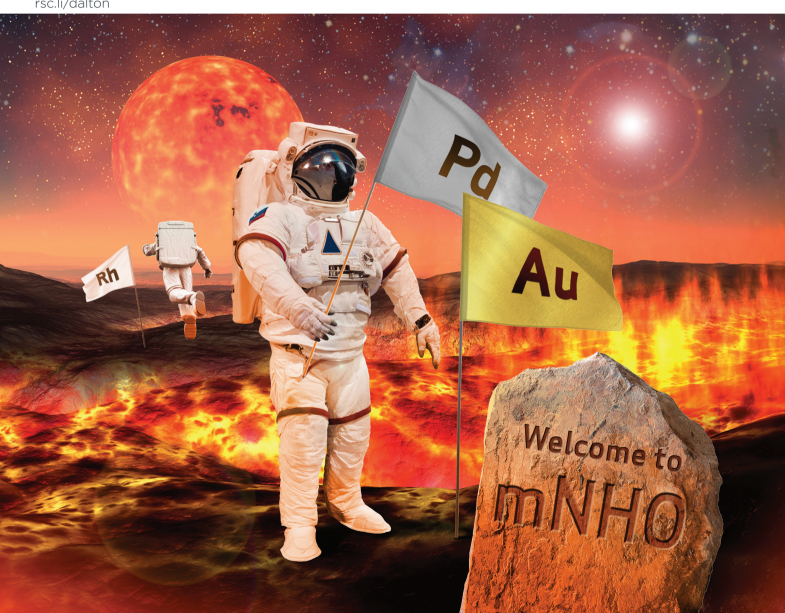
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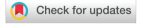
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Towards structurally versatile mesoionic N-heterocyclic olefin ligands and their coordination to palladium, gold, and boron hydride†

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We have developed an efficient and versatile approach for the synthesis of a family of 1,2,3-triazole-based mesoionic N-heterocyclic olefin (mNHO) ligands and investigated their coordination to palladium, gold, and boron hydride experimentally and computationally. We reacted mNHOs obtained through deprotonation of the corresponding methylated and ethylated 1,3,4-triaryl-1,2,3-triazolium salts with [Pd(allyl)Cl]₂ to give the corresponding [Pd(η^3 -allyl)Cl(mNHO)] coordination complexes. ¹³C NMR data revealed the strong σ -donor character of the mNHO ligands, consistent with the calculated bond orders and atom-condensed charges. Furthermore, we also synthesized [AuCl(mNHO)] and a BH₃-mNHO adduct by reacting the triazolium salts with AuCl(SMe₂) and BH₃-THF, respectively. The BH₃-mNHO adduct was tested in the reduction of select aldehydes and ketones to alcohols.

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Introduction

N-Heterocyclic olefins were first prepared in 1993¹ *via* deprotonation of methylated *N,N'*-dimethylimidazolium salts (NHOs, Fig. 1a) and have been identified as a promising new class of electron-rich and carbon-based ligands.^{2–10} Imidazole-based NHO coordination compounds with Mo,¹¹ Rh,^{12–14} Au,^{10,13} W,^{11,15} Ir,^{5,16} Pd,^{8,14} and Ti¹⁷ have been reported. Their utility in catalysis is also documented.^{8,15,16,18}

Subsequently, mesoionic N-heterocyclic olefins (mNHOs, Fig. 1b) were obtained in 2020 by the deprotonation of C5-methylated 1,3-diaryl-1,2,3-triazolium salts. ¹⁹ Unlike NHOs, the ground state structures of mNHOs cannot be described by an uncharged mesomeric Lewis structure, giving them increased basicity and donor abilities (Fig. 1). These triazole-based mNHOs have been investigated as catalysts for hydro-

boration²⁰ and *N*-methylation of primary amines.²¹ Reactions of C4-unsubstituted triazole-based mNHOs with Lewis acids (CO₂, AlMe₃, BH₃·SMe₂ and 9-borabicyclo[3.3.1]nonane),^{22,23} oxygen²⁴ and aryl azides²⁵ have also been described recently. mNHOs recently provided access to diazoolefins and the corresponding copper complexes.^{26–29}

Five rhodium-carbonyl complexes with mNHO ligands have been reported, three of them being triazole-based [Fig. 2a (i)]. ^{3,19,30} To the best of our knowledge, these are the only known coordination compounds with mNHOs apart from the related but peculiar complexes with diazoolefin ligands

Fig. 1 Formation and representative mesomeric structures of: (a) methylated *N,N'*-disubstituted imidazolium salts to form the corresponding imidazole-based N-heterocyclic olefins (NHOs), and (b) C5-methylated 1,3,4-trisubstituted-1,2,3-triazolium salts to form 1,2,3-triazole based mesoionic N-heterocyclic olefins (mNHOs).

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Fig. 2 (a) (i) Previously reported complexes with triazole-based mNHOs Dipp = 2,6-diisopropylphenyl and (ii) triazole-based diazoolefin ligands. (b) This work, featuring: (i) mNHO ligand design, and (ii) mNHO-M coordination compounds that potentially exhibit non-biaryl $C(sp^2)-C(sp^3)$ atropisomerism (providing $R \neq H$) as represented by a Newman projection, viewed along the $C_{exo}-C5$ (also C_{endo}) vector.

[Fig. 2a(ii)]. ²⁹ The studies indicated that mNHOs are more strongly basic compared to traditional NHOs and are much stronger σ -donors than both their NHOs and N-heterocyclic carbene (NHC) counterparts. However, unlike NHCs, they are unable to participate in π -backdonation. ^{2,19}

Thus, to further advance the field and reveal the hitherto unknown peculiarities of this ligand family, we have established novel efficient routes to structurally different 1,2,3-triazole-based mNHOs that can serve as versatile ligands to metals, and investigated their coordination properties towards palladium, gold, and boron hydride. Coordination of a prochiral mNHO (Fig. 2b (i), R \neq H) to a square-planar palladium complex was anticipated to give a novel example of non-biaryl $C(sp^2)$ – $C(sp^3)$ atropisomer (Fig. 2b(ii)), and this was also investigated.

Results and discussion

mNHO ligands

We chose to investigate mNHOs with simple phenyl ring functionalities at the N1 and N3 positions, as shown in **1a** (Fig. 3): the *p*-tolyl substituent at N1 was chosen to enable the compounds under investigation in the reaction mixtures to be easily followed by NMR spectroscopy *via* the distinct resonance of the methyl protons. Instead of employing triazene–alkyne

cycloaddition, which is commonly used to prepare symmetric 1,3-diaryl-1,2,3-triazole based scaffolds, ^{31–33} we chose to utilize copper-catalyzed click chemistry with subsequent N3 arylation as it allows for greater structural diversification. ³⁴

We designed two mNHO pro-ligands 1d and 1g, which potentially offer coordination through either the mNHO or the terminal nitrogen atom of the dimethylamine functionality. While compounds 1a-d can provide monodentate exocyclic carbon-ligation upon deprotonation, the general lack of stability of mNHO-metal compounds compelled us to consider pyridine- and phosphine-functionalized analogues 1e-g and 1h-i, which should enable C^N- and C^P-bidentate coordination, respectively, thus conferring additional stability to the complexes due to the chelating effect. Upon deprotonation and C_{exo} -ligation, the substrates **1a**, **c**, and **e** with a methyl group at the triazole C5 would potentially form conformationally constrained structures with limited rotation around the C_{exo}-C_{endo} vector, leading to an interesting class of monodentate atropochiral ligands. 35 In addition, in the same process, the prochiral substrates 1b, d, and f-i (Fig. 2b(i), R \neq H) would acquire a new stereogenic centre.

The synthesis of C5-substituted triazolium salts **1a–g** commenced from 'click triazoles' **2** which, upon treatment with iodonium salt, gave **1**,3,4-triaryl substituted triazolium triflates **3a** and **b** (Scheme **1a**). Triazolium salts **3a** and **b** were then alkylated at C5 to give **1a–b**, and **e–g**. This approach for the

Fig. 3 The chemical structures of compounds 1a-i initially planned for the synthesis.

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Scheme 1 Two alternative approaches for 1,3,4,5-tetrasubstituted triazolium salts 1 from click triazoles 2: (a) N3-arylation of triazoles 2a,b with diphenyliodonium triflate gives triazolium salts 3a,b, which upon treatment with a base (n-BuLi or KOt-Bu) and electrophile (alkyl iodide or N,N-dimethylmethyleneammonium iodide) affords target compounds 1a-b, and e-g. (b) N3-arylation of 1,4,5-trisubstituted triazole 2a' with bis(4-methoxyphenyl)iodonium hexafluorophosphate to give the target 1c. Percentage yields of isolated pure products are reported.

target compounds enables the synthetic steps to be reversed, e.g., for the synthesis of 1c, starting with C5-methylated triazole 2a'36 could be N3-arylated as shown in Scheme 1b.

For the synthesis of diphenylphosphine functionalized triazolium salt 1h, pent-1-yne (4) was converted to triazole 5 according to the modified literature procedure,³⁷ which was then N3 methylated with Merwein salt (Scheme 2a). For the arylation of 5 with iodonium salt, the phosphorus atom was BH₃-protected to form 6 (Scheme 2b), as it was presumed that the relatively harsh reaction conditions would otherwise lead to undesired P- rather than N-arylation. It turned out, however, that BH₃ protection in 6 was not tolerated by the arylation protocol; in situ phosphine deprotection occurred prior to arylation and we isolated the phosphonium salt 1j instead of the desired product 1i (Scheme 2b).

To obtain the corresponding mNHO derivatives, deprotonation of this set of triazolium salts 1 was attempted using an equimolar amount of potassium bis(trimethylsilyl)amide

(KHMDS, Scheme 3) in dry THF at room temperature. For 1a, this resulted in an immediate colour change, with the reaction mixture turning purple, indicating the deprotonation and formation of mNHO 7a [Scheme 3(a)]. This colour change was consistent with what was previously reported19 indicating the resonance stabilisation of the negatively polarised C_{exo} moiety with the cationic triazole unit. With the other triazolium salts (1b-1g), the reaction mixtures instantly turned purple (7b and 7c), dark blue (7d, 7e, and 7f) or green (7g) [the structures of corresponding mNHOs are shown in Scheme 3(b)]. According to ¹H NMR analyses of the reaction mixtures, the deprotonation of triazolium salts 1a-1g with KHMDS to 7a-7g was quantitative.

Deprotonation of 1a to 7a with KHMDS in THF- d_8 was monitored by ¹H NMR spectroscopy. The spectra indicated the appearance of two doublets at δ = 3.18 and 3.25 ppm corresponding to non-equivalent geminal CexoH2 protons. The chemical shifts were in the range observed for olefinic protons

Scheme 2 Synthesis of compounds 1h and attempted preparation of 1i, forming 1j instead. Percentage yields of isolated pure products are reported.

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Scheme 3 (a) Deprotonation of 1a to mNHO 7a; (b) the structures of the other mNHOs (7b-7g) that were formed.

and in agreement with those reported previously for CexoH2 protons in related compounds.¹⁹ In the case of 7b, obtained by in situ KHMDS deprotonation of 1b, the ¹H NMR spectra showed the appearance of two overlapping quartets at δ = 3.87 and 3.84 ppm (J = 7.4 and 7.3 Hz) in an approximately 1:0.7 ratio that could be assigned to the methine C_{exo}H protons of the two geometric isomers E and Z at the exocyclic double bond (Fig. 4). The resonances for the methyl groups appeared as two overlapping doublets at δ = 1.26 and 1.24 ppm (J = 7.4 and 7.3 Hz). A similar observation of the geometric isomers was made for 7d, 7f and 7g (obtained from 1d, 1f and 1g, respectively). On the other hand, the aromatic regions of the ¹H NMR spectra were in all cases overcrowded with overlapping resonances; those for the methyl groups of the p-tolyls unambiguously showed the presence of the two isomers. All mNHOs 7 were found to be sensitive to moisture, instantly reforming the triazolium salts 1 upon exposure to the atmosphere.

The treatment of 1j with KHMDS resulted in the formation of triphenyl(1,2,3-triazol-4-yl)phosphonium ylide (8j) based on

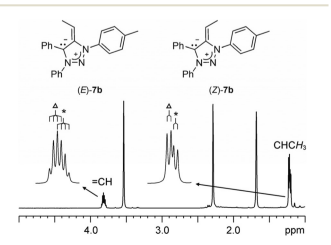


Fig. 4 The selected region of ¹H NMR (THF-d₈) spectrum of in situ generated 7b. ^a and * denote the major and minor isomers, respectively.

the NMR analysis (Scheme 4a, and Fig. S1†). Interestingly, related N-unsubstituted 4-aryl-5-triphenylphosphonium-l,2,3triazole ylides were reported in 1973,³⁸ and since then have not been investigated further.

The treatment of 1h with KHMDS was accompanied by an immediate colour change to dark red, indicating deprotonation (Scheme 4b); however, the NMR spectra of the reaction mixture acquired shortly after the addition of a base revealed decomposition to a complex mixture of unidentified products that likely proceeds through 7h and/or 7h'.

mNHO-palladium coordination compounds

We selected 7a as a model compound to probe coordination abilities toward palladium. Treatment of in situ formed 7a with [Pd(allyl)Cl]₂ in dry THF at room temperature instantly afforded 7aPd (Table 1). The crude product was dissolved in diethyl ether and the solution layered with pentane at −20 °C to form a yellow precipitate, which was isolated by filtration. This yellow solid was stable for several days under a nitrogen atmosphere at room temperature but began to decompose after prolonged storage. Various attempts to prepare crystals of 7aPd suitable for X-ray analysis (including using different solvents, solvent mixtures, temperatures, and crystallisation techniques) were unsuccessful. Thus, 7aPd was characterized by a combination of mass spectrometry (Fig. S2) and NMR spectroscopy (see the ESI†) and its structure was corroborated by density functional theory (DFT) calculations (see below).

High-resolution (HRMS) electrospray ionization mass spectrometry in a positive ion mode (ESI+) revealed a [M - Cl]⁺ ion at m/z 472.0997 (calcd for $C_{25}H_{24}N_3Pd^+$: 472.1005) and was interpreted as the result of in-source collision-induced dissociation of 7aPd (Fig. S2a†). The peak that corresponds to $[(1a-TfO)]^+$ m/z 326.1647 (calcd for $C_{22}H_{20}N_3^+$: 326.1652) was the most intense in the spectrum, likely the result of hydrolysis of 7aPd during the sample preparation.

The structure of 7aPd consists of a neutral square planar complex of Pd(II) in which mNHO monodentately coordinates to Pd through the exocyclic carbon (Cexo) atom. The allyl

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a) KHMDS (1.05 equiv.) THF, rt,
$$N_2$$
, 15 min N_2 , 16 min N_2 , 16 min N_2 , 17 min N_2 , 18 min N_2 , 18 min N_2 , 19 m

Scheme 4 (a) Deprotonation of 1j to 8j, showing the canonical structures of 8j; (b) deprotonation of 1h.

Table 1 Synthesis of Pd complexes 7aPd and 7bPd, and selected NMR data for 1a, 7a, 7aPd and 7bPd

Cmpd.	NMR solvent	$\delta_{ m C1(allyl)}$	$\delta_{ m C2(allyl)}$	$\delta_{\mathrm{C3(allyl)}}$	$\delta_{ m CexoH}$	$\delta_{ m Cexo}$	δ_{C4}	$\delta_{ ext{C5}}$	$\delta_{ m N1}$
1a 7a	$ ext{CDCl}_3$ $ ext{THF-}d_8$				2.51 3.18, 3.25	11.0 46.8	140.9 120.1	140.1 146.1	250 192
7aPd 7bPd	THF THF	63.6 62.4, 63.4	109.9 109.8, 111.4	53.7 56.0, 56.1	$2.01, 2.33$ $2.98, 3.30^a$	-1.6 10.7, 10.9^a	130.2 130.7, 130.7 ^a	161.6 162.2, 162.3 ^a	238

 13 C NMR chemical shifts were extracted from the 1 H $^{-13}$ C gs-HSQC and 1 H $^{-13}$ C gs-HMBC spectra. a Two resonances for the two stereoisomers are given. The first and second figures refer to the upfield and downfield resonances, respectively.

ligand binds to palladium in an η^3 -fashion with its two terminal atoms occupying *cis*-positions and the coordination sphere is completed by a chloride (Fig. 5a), also shown in Table 1.

A square-planar complex may be chiral under certain conditions even if it does not consist of chiral ligand(s).³⁹ The chirality in a generic d⁸ *cis*-M(Me Py)AB₂ platform, shown in Fig. 5b with 2-methylpyridine (a monodentate planar ring with only C_s local symmetry) as one of the ligands, originates from the rotation around the M–N bond that generates the opposite

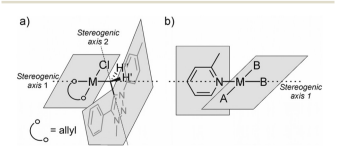


Fig. 5 (a) Graphical representation of planes and possible stereogenic axes in **7aPd**. (b) Axial chirality of a square-planar complex with 2-methylpyridine as an exemplary ligand from the literature.³⁹

enantiomer; this pair of enantiomers can be described in terms of axial chirality. ^{40,41} In addition to pyridine and some other heterocyclic ligands, square-planar metal complexes incorporating carbene ligands with the heterocyclic ring perpendicular to the coordination plane showing axial chirality are well documented. ⁴²

In line with the experimentally deduced structure of 7aPd, DFT calculations demonstrate such an arrangement and configuration to be a stable, ideal d^8 square-planar molecular structure. The computed equilibrium structure, shown in Fig. 6, in addition, reveals the in-plane orientation of the $C_{\rm exo}$ - $C_{\rm endo}$ bond of the mNHO ligand with respect to the plane of the complex. The emerging two conformers, namely $C_{\rm exo}$ - $C_{\rm endo}$ pointing *towards* or *away* from Cl $^-$, differ by 6 kcal mol $^{-1}$ in Gibbs free energy, the former isomer being the more stable. Relaxed scan calculations show that the two isomers are interconnected through a shallow potential energy surface with a rotational maximum at 8 kcal mol $^{-1}$ to the more stable isomer, indicating free rotation along the Pd- $C_{\rm exo}$ bond at room temperature.

In contrast, the rotation along the $C_{\rm exo}$ – $C_{\rm endo}$ bond is sterically hindered: as can be deduced intuitively from the structure shown in Fig. 5a and 6, rotation along this bond leads to the clash of the phenyl and tolyl groups of mNHO with the

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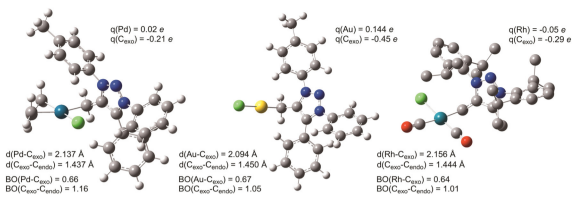


Fig. 6 Equilibrium structures of 7aPd (left), 7aAu (middle) and [RhCl(CO)₂mNHO] (right) obtained at the PBE0/def2-SV(p) level of theory, given together with characteristic structural metrics (metal- C_{exo} and C_{exo} - C_{endo} bond distances), corresponding bond orders and atom condensed charges.

Pd–Cl functionality. Relaxed surface scans imply sharply increasing potential energy surfaces as these groups approach the chloride ligand, triggering the distortion of the structure and causing linearization of the Pd– $C_{\rm exo}$ – $C_{\rm endo}$ angle to avoid close contact with these groups. This leads to unstable wavefunctions and high electronic energies. The hindered rotation creates a stereogenic axis characteristic of the molecular platform, as shown in Fig. 5a, which has implications also for the NMR spectra as discussed below.

A limited rotation around *Stereogenic axis 2* leads to the formation of a pair of enantiomers P and M (Fig. 7a) which, in a non-chiral environment (like in this study), should lead to the observation of one species in the NMR spectra. However, the constitutionally equivalent atoms H' and H'' of the exocyclic methylene group are not symmetry related and should res-

a) $C^{4} \stackrel{H'}{\longrightarrow} H^{0}$ $P^{1} \stackrel{\to} H^{0}$ $P^{1} \stackrel{\to}$

Fig. 7 (a) Viewed along the $C_{\rm endo}-C_{\rm exo}$ vector (Stereogenic axis 2), the triazole ring lies in front of the tetrahedral carbon atom giving stereoisomers P and M. (b) The combined conformationally stable $C_{\rm endo}-C_{\rm exo}$ stereogenic axis (Stereogenic axis 2) and $C({\rm sp}^3)$ stereogenic centre ($C_{\rm exo}$) give rise to two diastereomeric pairs of enantiomers (P,S and M,R; and P,R and M,S) of 7bPd (R = Me, descriptors R,S refer to the absolute configuration at $C_{\rm exo}$).

onate at different chemical shifts in the 1 H NMR spectrum. Indeed, the 1 H spectrum of compound 7aPd shows two doublets at $\delta = 2.33$ ppm (J = 9.6 Hz) and $\delta = 2.01$ ppm (J = 9.6 Hz) belonging to the diastereotopic H' and H" (Table 1, and Fig. S3), while only one set of carbon resonances was observed in the 1 H- 13 C gs-HSQC and 1 H- 13 C gs-HMBC spectra (Fig. S48–S52†).

Now, if a proton of the Cexo methylene group in 7aPd is substituted by another group, say, a methyl group, this introduces a C(sp³) stereocentre into the molecule. Provided that fast rotation is present around Stereogenic axis 1, this results in four stereoisomers, i.e., two diastereomeric pairs of enantiomers (P,S and M,R; and P,R and M,S, Fig. 7b) and, thus, two observable species in the NMR spectra. This was confirmed by the experimental data for 7bPd, the preparation of which is shown in Table 1. Analyses of the ¹H NMR and ¹³C NMR chemical shifts revealed the presence of two species of 7bPd in the THF solution as evident from the two sets of resonances for the allyl ligand and two sets for the mNHO ligand (Fig. S4†). Two sets of ¹³C resonances could also be extracted from the 1H-13C gs-HSQC and 1H-13C gs-HMBC spectra (Fig. S53-S55†). Remarkably, this molecular scaffold with a conformationally stable C(sp2)-C(sp3) stereogenic axis is a novel example of this rare stereogenic element.44

For 7aPd, the ¹³C resonance of C_{exo} was found at δ = -1.6 ppm (Table 1), which is different from those of the previously reported triazole based mNHO-Rh complexes (δ = 7.0-8.3 ppm (ref 19) and 7.9 ppm (ref. 30)) yet similar to that of the imidazole-based-NHO-Pd (δ = -7.0 - (+0.99) ppm). ¹⁴ Increasing the substitution of C_{exo} when going from 7aPd to 7bPd caused a downfield shift of about +11 ppm for the methyl group; in diastereomeric 7bPd, C_{exo} resonated at δ = 10.7 and 10.9 ppm, which is close to the value expected in alkane series (+9 ppm). ⁴⁵

In addition, the η^3 -allyl ligand can adopt an *exo* or *endo* configuration; the designations refer to isomers in which the vectors C_{exo} - C_{endo} and $C2^{\text{ally}}$ -H point in the same or in opposite directions, respectively. Sharp proton peaks of the allyl

ligand in both 7aPd and 7bPd, in conjunction with solution NMR investigations reported in the literature, 46 suggest fast isomerisation^{47,48} between the exo and endo isomers of (n³allyl)Pd and, thus, this dynamic phenomenon has no influ-

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ence on the above-discussed diastereomerism.

Although ¹³C NMR chemical shifts result from a complex interplay between shielding and deshielding effects, 45 themselves stemming from the intramolecular electron flows stimulated by the applied external magnetic field, it has been suggested (on the basis of both experimental⁴⁹ and theoretical investigations⁵⁰) that there is an empirical correlation between the ¹³C NMR shifts and the relative charge (and thus reactivity) at the η^3 -allyl termini in the (η^3 -allyl)palladium complexes. Namely, when the ligands go from pure donors to those with some acceptor character, the 13 C shift at the η^3 -allyl termini trans to the ligand shifts downfield. For example, the η^3 -allyl termini trans to purely donor TMEDA (tetramethylethylenediamine) and chloride were found to resonate at ca. δ = 60-62 ppm whereas, for example, for the corresponding triphenylphosphine ligand with some π -acceptor character, the ¹³C shift can be found at ca. 79 ppm. ⁴⁹ Besides guiding the deduction of a tentative structure, the ¹³C NMR shifts of C3^{allyl} at δ = 53.7 ppm and δ = 56.0 ppm for 7aPd and 7bPd, respectively (Table 1), indicate the strong σ donor character of the mNHO ligands. 19,30 The calculated bond orders and atom-condensed charges, listed in Fig. 6, corroborate this finding; a bond order of 0.66 indeed indicates a strong Pd-Cexo interaction and a Lowdin charge analysis revealed almost identical atomic charges on the allyl terminal carbons trans to mNHO (0.00e) and trans to chloride (-0.03e).

Another interesting feature of 7aPd revealed by calculations is a C_{exo}-C_{endo} bond order of 1.16 that implies a minor but non-negligible π interaction between the triazole ring and the exo carbon, i.e. the reactivity of Cexo's lone-pair and localized negative charge is quenched not only by the interaction with the metal but also by some delocalization to the aromatic heterocycle, which is consistent with the resonance structures of mNHO in Fig. 1. The corresponding π interaction is portrayed clearly by HOMO and HOMO-6 of 7aPd (Fig. S5†).

The allyl ligand in 7aPd and 7bPd appears to stabilize the mNHO-metal bond, probably by the electron-withdrawing effect originating from its π -accepting ability. This may explain why the treatment of 7a and b with non-allyl metal precursors such as [PdCl₂(NCPh)₂] did not give stable products. Nevertheless, even keeping 7aPd or 7bPd dissolved overnight in dry THF under an inert atmosphere was detrimental and led to the formation of a complex mixture of unidentified products as revealed by NMR analysis.

mNHO-gold coordination compounds

Gold is known to be particularly carbophilic and a strong Lewis acid 13,51-53 and, concomitantly, it has been successfully coordinated to NHOs. 10,13 Accordingly, in a similar fashion to the synthesis of the Pd complexes, 7a was prepared in situ from 1a and reacted with chloro(dimethylsulfide)gold(I), AuCl (SMe₂), to give a product that was identified as 7aAu (Table 2)

Table 2 The synthesis of complex 7aAu and 7bAu with selected NMR data

Cmpd.	NMR solvent	$\delta_{ ext{CexoH}}$	$\delta_{ m Cexo}$	$\delta_{ ext{C4}}$	$\delta_{ ext{C5}}$
7aAu 7bAu	THF THF	2.15 2.83,	15.2 15.6,	132 133.0,	153 154.7,
, 5214		2.89^{a}	26.2^{a}	133.2^a	155.4^{a}

¹³C NMR chemical shifts were extracted from the ¹H-¹³C gs-HSQC and ¹H-¹³C gs-HMBC spectra. ^a Two resonances for the two stereoisomers are given. The first and second figures refer to the upfield and downfield resonances, respectively. For the NMR data for 1a, see Table 1.

according to NMR spectroscopy and ESI+ HRMS spectrometry. In the ¹H NMR spectrum (THF solution), the methylene protons of **7aAu** appeared as a singlet at $\delta = 2.15$ ppm. In the carbon NMR spectrum, C_{exo} was found to resonate at δ = 15.2 ppm, which is considerably downfield as compared to the equivalent Cexo in 7aPd (see above). The positive-ion ESI-HRMS revealed the [7aAu-Cl]⁺ ion at m/z 522.1236 (calcd for $C_{22}H_{19}AuN_3^+$ [7aAu-Cl]⁺ 522.1239) and was interpreted as the result of in source collision-induced dissociation of 7aAu (Fig. S6†). In addition, the $[7aAu-Cl + CH_3CN]^+$ ion at m/z at 563.1491 (calcd for $C_{24}H_{22}AuN_4^+$: 563.1505) was found to be the most intense peak. The acetonitrile in this cluster came from the solvent used for the sample preparation. As in the case of 7aPd, the peak that corresponded to the triazolium cation [(1a-TfO)] was also found in the ESI+ HRMS spectrum of 7aAu. As shown in Fig. 6, 7aAu adopted a linear structure, corresponding to a d¹⁰ metal centre, with a rather strong (BO = 0.67) and short (2.09 Å) Au-mNHO bond. Another goldmNHO derivative, complex 7bAu (Table 2), was prepared in the same way as 7aAu. As with the palladium derivatives, the gold complexes 7aAu and 7bAu were stable in the solid state for several days under an inert atmosphere but decomposed overnight under an inert atmosphere in a THF solution.

Phosphine-functionalized triazolium salt coordinated to palladium

The coordination properties of the phosphine-functionalized analogue 1h, which may provide C^P-bidentate coordination, were also investigated. Since the attempts to prepare mNHO 7h were unsuccessful (Scheme 4b), we decided to test a stepwise approach, and compound 1h was first reacted with PdCl₂ in dichloromethane to give a pale yellow bis-phosphine-Pd complex 9hPd (Scheme 5), the structure of which was confirmed by NMR and X-ray diffraction analysis (for crystallographic data, see ESI Fig. S7, S8 and Tables S1, S2†). Treatment of the isolated **9hPd** with KHMDS in the next step

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Scheme 5 Synthesis of the phosphine-functionalized Pd complex 9hPd (for crystallographic data; see the ESI†).

to produce a C^P-bidentate coordination was unsuccessful and resulted in the decomposition of 9hPd and the formation of black palladium. Similar to 1h (vide supra), based on NMR spectra, it appeared that the triazole N3-Me part of the ligand in 9hPd did not survive treatment with a base.

mNHO-boron hydride adduct

In comparison with Pd and Au, mNHO Lewis adducts with trihydridoborane (BH₃) turned out to be exceedingly stable. Compound 10aBH3 was easily prepared starting from 1a through in situ formation of mNHO 7a with subsequent addition of BH3:THF (Scheme 6) and isolated as a light-yellow air-stable solid. Crystals suitable for X-ray diffraction analysis were obtained by layering a dichloromethane solution of the product with hexane at −20 °C (Fig. 8). For crystallographic data, see ESI Fig. S9, S10 and Tables S1 and S3.†

To our knowledge, there are two reports of BH3-mNHO adducts in the literature 20,22 and catalytic activity of mNHOs in

Scheme 6 The synthesis of mNHO-boron hydride adduct 10aBH₃.

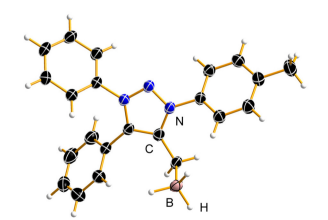


Fig. 8 The asymmetric unit of the 10aBH₃·0.825(CH₂Cl₂) crystal structure; the disordered CH₂Cl₂ molecules were omitted for clarity. Displacement ellipsoids are depicted at the 50% probability level and hydrogen atoms are shown as small spheres of arbitrary radius.

the hydroboration of imines, nitriles and N-heteroarenes with H(Bpin) has been reported.²⁰ Based on theoretical studies, compared with some other N-heterocyclic carbenes (NHCs), N-heterocyclic olefins (NHOs) and mesoionic carbenes (MICs), the mNHO/HBpin adducts possess the most negative charge on H-(Bpin), weakening the B-H bonds and enhancing the hydridic character, which promotes the hydroboration of unsaturated substrates. This prompted us to investigate the reducing ability of 10aBH3 toward selected aldehydes 11 and ketones 13. 3-Phenylpropionaldehyde (11a) was used as a model substrate to briefly optimize the reaction conditions (for more details, please see the ESI†). The scope of the reaction with three benzaldehydes, 11b, c, and d (0.20 M), having different electronic properties (Scheme 7A), was investigated by using 0.5 equiv. of 10aBH3 in acetonitrile as the reaction solvent (Table S4†). Electron-deficient 3-nitrobenzaldehyde (11c) was converted to 3-nitrobenzyl alcohol (12c) in 44% after 4 h, with the starting aldehyde being completely consumed. As expected, the conversion of electron-rich 3,4,5-trimethoxybenzaldehyde (11d) was slower (21% conversion to 12d) under the same reaction conditions.

The reduction of carbonyl compounds by related NHC- and MIC-boranes has been investigated.54-57 Comparison of our results with those previously reported for NHC- and MICboranes⁵⁴⁻⁵⁷ suggests that 10aBH₃ is more active and no additives are needed (both NHC- and MIC-borane reduction of aldehydes require the presence of a Lewis acid such as Sc (OTf)₃ or silica gel) for the reaction to proceed, although the reaction times with 10aBH3 were longer.

Next, the reducing power of 10aBH3 was tested on ketones 13 (Scheme 7B, and Table S5†). Except for strongly activated 4-nitroacetophenone (13a), which led to 76% conversion to 14a under the same reaction conditions as used for the aldehydes, reductions of other substrates by using mNHO-borane 10aBH₃ alone did not occur. The addition of silica gel was beneficial and allowed the reduction of 4-bromoacetophenone (13b) to 1-(4-bromophenyl)ethanol (14b) in 66% after 24 h. For comparison, reduction of 13b with 1 equiv. of NHC-BH3 diMe-Imd-BH₃ gave alcohol 14b in 94% yield after 24 h in DCM.⁵⁴ Control experiments were performed to test the stability of 10aBH₃. A CD₃CN solution of 10aBH₃ with or without silica gel was monitored by ¹H NMR for four days and only negligible changes were observed in the recorded spectra, indicating that BH₃ does not dissociate from mNHO and that the mNHOboron hydride species play an active role in the reaction with the carbonyl compounds.

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Scheme 7 Reduction of (A) aldehydes and (B) ketones. Conversions were determined by ¹H NMR with 1,3,5-trimethoxybenzene as internal standard: a aldehyde 11a (0.029 mmol, 0.058 M in MeCN), 10aBH₃ (1.0, 0.5 or 0.33 equiv.), 3 h, air, room temperature; b aldehyde (0.10 mmol, 0.20 M in MeCN), 10aBH₃ (0.5 equiv.), 24 h, air, room temperature; ^c reaction time was 4 h; ^d ketone (0.10 mmol, 0.20 M in MeCN), 10aBH₃ (0.5 equiv.), 24 h, air, room temperature; e reaction performed in the presence of silica gel (100 mg).

Conclusions

1,4-Disubstituted triazoles obtained by click chemistry were functionalized at the triazole nitrogen atom N3 and at the carbon atom C5 to give the corresponding C5-alkylated triazolium salts. Deprotonation of C5-CexoH was achieved with KHMDS in dry THF at room temperature to give deeply coloured mesoionic N-heterocyclic olefins (mNHO). The colour is due to the resonance stabilisation of the negatively polarised C_{exo} moiety with the cationic triazole unit, which was confirmed by the NMR study through the observation of two geometric isomers indicating C5-Cexo double bond properties. To investigate the coordination properties, selected in situ prepared mHNOs were allowed to react with metal precursors $[Pd(allyl)Cl]_2$ and $AuCl(SMe_2)$. $[Pd(\eta^3-allyl)]$ Cl(mNHO)] and [AuCl(mNHO)] were readily obtained and characterised by multinuclear 1D and 2D NMR spectroscopy and high resolution electrospray ionisation mass spectrometry (ESI+ HRMS). The NMR data showed a strong σdonor character of the mNHO ligands on the metals, which was corroborated by DFT calculations, revealing bond orders of 0.66 and 0.67 for the Pd-mNHO and Au-mNHO bonds, respectively. Besides, a C_{exo}-C_{endo} bond order of 1.1 implies that this bond of the mNHO ligand maintains a non-negligible double bond character and, thus, π -interaction with the triazole core. The $[Pd(\eta^3-allyl)Cl(mNHO)]$ complex with C5-ethylated mNHO also exhibited a rare conformationally stable $C(sp^2)$ - $C(sp^3)$ stereogenic axis, which was scrutinized experimentally and computationally. Noteworthily, these are the first known palladium and gold coordination complexes of mNHOs.

A BH₃-mNHO adduct was prepared and its structure was confirmed by NMR and X-ray single-crystal analysis. The reactivity of BH₃-mNHO in the reduction of aldehydes and ketones to the corresponding primary and secondary alcohols was investigated. It was found to be more active than the

related NHC- and MIC-boranes as no Lewis acid additives were required for the reduction of aldehydes to proceed.

Author contributions

TŽ: investigation, methodology, validation, visualization, writing - original draft, and writing - review & editing; BP: investigation, methodology, resources, visualization, and writing - review & editing; ML: investigation, resources, visualization, and writing - review & editing; DU: investigation, visualization, and writing - original draft; RDJv-V: writing - review & editing; JK: conceptualization, data curation, funding acquisition, project administration, resources, supervision, visualization, writing - original draft, and writing - review & editing.

Conflicts of interest

There are no conflicts of interest to declare.

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