Allowing the Direct Interaction of N-Aryl α-Diimines with a High Valent Metal Chloride: One-Pot WCl₆-Promoted Formation of Quinoxalinium Salts

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Abstract. The full potential of a high valent metal chloride as both a chlorinating and an oxidative agent was explored by allowing WCl₆

10 to react with N-(2,6-diisopropylphenyl) α -diimines, in CH₂Cl₂ at room temperature. These a-diimines underwent unprecedented conversion to quinoxalinium cations via intramolecular C-N coupling.

 α -Diimines (also called 1,4-diaza-1,3-dienes = DAD) constitute a 15 fascinating class of compounds whose steric and electronic properties may be finely tuned by varying the substituents on the N=CC=N skeleton.¹ Due to their versatility and robustness, α diimines have been widely employed as ancillary chelating ligands in coordination chemistry,² assisting a variety of ²⁰ stoichiometric and catalytic transformations.³ α -Diimines are prone to undergoing a stepwise reduction (up to two electrons reduction),^{1c,4} and for this reason they are considered as redox non-innocent ligands when bound to a metal centre,^{2b,5} possibly determining some ambiguity in the assignment of the oxidation

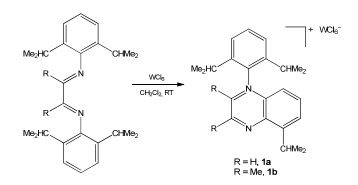
- $_{25}$ states.⁶ On the other hand, the oxidation chemistry of α -diimines still remains, surprisingly, an almost unknown field. As a matter of fact, the electro-oxidation behaviour of non coordinated α diimines has not been clearly assessed hitherto.⁷ Besides, the reactivity studies with high valent metal compounds (oxidation
- $_{30}$ state of the metal \geq 4), that are potential, strong oxidizing agents, are exceedingly rare in the literature.4,8 With reference to homoleptic, high valent transition metal chlorides, the reactivity of NbCl₅,⁹ TaCl₅,¹⁰ MoCl₅ ¹¹ and WCl₆ ¹² with α -diimines has aroused a recent and notable consideration. However, all the 35 described reactions have been performed with prior reduction, by

one or two electrons, of the metal centre. This strategy quenches the activation ability of the strongly acidic metal chlorides, 13,14,15 so to straightforwardly afford coordination compounds with intact, redox active a-diimine ligands. A similar outcome has

(including Scheme SI1), X-ray crystallography, electrochemical studies (including Table SI3 and Figure SI1), computational studies (including Scheme SI2 and Figures SI2-SI6), NMR spectra (Figures SI7-SI12). CCDC 1551416 (1a·CH₂Cl₂) and 1551419 (2b) contain the supplementary crystallographic data. These data can be obtained free of charge from The Cambridge Crystallographic Data centre via www.ccdc.cam.ac.uk/data_request/cif.

⁴⁰ been achieved by allowing TiX₄ (X = Cl, Br) 2d,16 and MCl₅ (M = Nb, Ta)^{16a,17} to react with dilithium(1,4-diaza-1,3-dienyl) salts.

Herein, we describe for the first time the reactions of two α diimines with a high valent metal chloride, in a weakly coordinating solvent (dichloromethane) and in the absence of 45 further reactants (e.g., reductants). Thus the 1:1 molar reaction of N,N'-bis(2,6-diisopropylphenyl)-1,4-diaza-1,3-WCL with butadiene (DAD^{Dip}) afforded the quinoxalinium salt [{2,6- $C_6H_3(CHMe_2)_2$ N(CH)₂NCC(CHMe₂)(CH)₃C][WCl₆], 1a, in 60% yield (Scheme 1).¹⁸ Compound 1a was isolated and then 50 characterized by spectroscopic methods and single crystal X-ray diffraction (see Figure 1, and Table SI1 in the Supporting Information). The analogous 1b was obtained in 45% yield from WCl_6 and 2,3-bis(2,6-diisopropylphenylimino) butane (Me-DAD^{Dip}), see Scheme 1.



60 Scheme 1. α-diimine to quinoxalinium conversion promoted by WCl6.

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[†] Supporting Information contain details of experimental procedures

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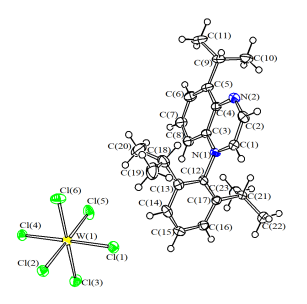


Figure 1. View of the structure of **1a** with key atoms labeled. Displacement ellipsoids are at the 50% probability level. Only one of the ⁶⁵ two independent cation-ion pair present within the unit cell is represented.

- Significant amounts of isopropyl chloride were clearly recognized by NMR/GC-MS analyses on the liquids distilled from the WCl₆/DAD^{Dip}/CD₂Cl₂ and WCl₆/Me-DAD^{Dip}/CD₂Cl₂ reaction 70 systems, respectively. By using the same method, we detected limited amounts of alkyl halides as produced from the combination of WCl₆ with some other *N*-aryl α-diimines (see Experimental for details), suggesting that the activation process observed for DAD^{Dip} and Me-DAD^{Dip} might hold some general 75 character.
- The crystal structure of **1a** consists in an ionic packing of $[WCl_6]^$ anions ¹⁹ and $[\{2,6-C_6H_3(CHMe_2)_2\}N(CH)_2NCC(CHMe_2)$ $(CH)_3C]^+$ cations. The bonding parameters of the quinoxalinium core do not significantly differ from those previously reported for
- core do not significantly differ from those previously reported for ⁸⁰ N-alkyl-quinoxalium cations.²⁰ Quinoxalinium salts have aroused interest as possible chemosensors,²¹ in synthetic organic chemistry ²² and for their pharmaceutical ²³ and photophysical properties.²⁴ They are usually generated via the condensation of α -diketones with 1,2-phenylenediamines.^{24a,25a-b,26} A restricted
- ⁸⁵ number of N-aryl quinoxalinium salts have been synthesized to date, 25b,27 and the crystallographic characterization of **1a** provides the first X-ray structure of a N-aryl-quinoxalium cation ever reported. It should be remarked here that the known activation reactions of α -diimines are rare, 5c,28 being usually limited to C–C
- ⁹⁰ and C–Cl coupling additions to the imine carbons.^{9,10,28,29,30} A tricyclic dipyrrolopyrrole was obtained by FeCl₂-promoted oligomerization of a N-aryl α -diimine, accompanied by the loss of half of one DAD fragment.³¹ The cyclization reaction presented herein has more affinity with the chemistry of 2,3-
- ⁹⁵ bis(diphenylphosphino)-1,4-diazadienes, undergoing a peripheral ring closure upon coordination of the phosphine groups to group 10 metal halides.³²
- Several experiments, including electrochemical analyses and spectroscopic studies, and DFT calculations allowed to elucidate 100 some mechanistic aspects of the WCl₆ promoted cyclization
- reaction of DAD^{Dip}. First, the voltammetric investigation of a series of α -diimines (see
- Chart 1 in the SI and Figure 2 for DAD^{Dip}) at a Pt disk electrode was performed in CH_2Cl_2 . The examined compounds underwent,
- ¹⁰⁵ in addition to the well established one-electron reduction, an oxidation process at potentials ranging between 1.43 and 1.70

V.³³ These values would allow, in principle, a chemical oxidation by WCl₆ in CH₂Cl₂ ($E^{\circ'}$ ca. 1.6 V vs Ag/AgCl/KCl).³⁴

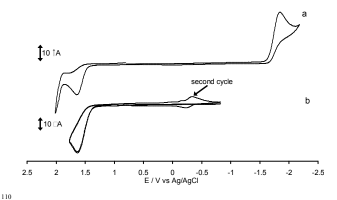
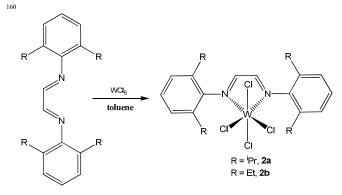


Figure 2. Cyclic voltammograms of DAD^{Dip} recorded at a Pt electrode in CH₂Cl₂ solution containing [$^{n}Bu_{4}N$][PF₆] 0.2 M. Scan rate 0.1 V s⁻¹. a) Cyclic voltammetry obtained starting the scan toward negative potentials from +0.8 V; b) 2 cycles voltammetry between -0.8 and 1.8 V.

- Notwithstanding, variable temperature EPR analysis of a WCl₆/DAD^{Dip} mixture in CH₂Cl₂ did not allow to detect any trace of [DAD^{Dip}]⁺. Otherwise, the electrochemical oxidation of DAD^{Dip} to [DAD^{Dip}]⁺ did not lead to final formation of the quinoxalinium cation contained in **1a** (see comparison of UV spectra in Figure SI1). Accordingly, the reaction in CH₂Cl₂ of DAD^{Dip} with NOBF₄, i.e. a typical oxidizing agent whose power is comparable to that of WCl₆,³⁴ did not give the quinoxalinium derivative among a complicated mixture of products. These facts ¹²⁵ suggest that the synthesis of **1a**, albeit containing a W(V) anion, cannot be the consequence of a simple electron transfer from DAD^{Dip} to W(VI). In agreement with these experimental observations, key intermediates along a plausible reaction pathway to **1a** were DFT calculated (see Scheme SI2 and Figures ¹³⁰ SI2-SI6).
- On theoretical grounds, it appears that the initial interaction WCl₆-DAD^{Dip} results in a favourable Cl-attack to one of the two C=N moieties (Figures SI2-SI3). Experimentally, the 1:1 reaction in CD₂Cl₂ of DAD^{Dip} with PCl₅, *i.e.* a typical chlorinating 135 substance employed in organic synthesis, ^{35a} selectively proceeded with H/Cl substitution at one imine group to give [{2,6- $C_6H_3(CHMe_{2)_2}N=CHC(Cl)=N\{2,6-C_6H_3(CHMe_{2)_2}\}]$ as the stable product (Scheme SI1).³⁵ Following chlorination of the organic substrate, one-electron transfer from the newly formed 140 ligand to the metal centre is likely to occur. This step is concomitant with the migration of one chloride from the metal to isopropyl-substituted carbon atom. The cyclization step should then take place favoured by the presence of both an activated aromatic ring and a formally anionic nitrogen atom. The release 145 of isopropyl chloride (see above) is likely to represent the driving force for the entire process. The radical derived from isopropyl elimination (Figure SI6) could be oxidized to the final quinoxalinium cation either by WCl₅ ($\Delta G = -7.5 \text{ kcal mol}^{-1}$) or more easily by still unreacted WCl₆ ($\Delta G = -40.8 \text{ kcal mol}^{-1}$).
- ¹⁵⁰ The W(IV) coordination complex WCl₄(DAD^{Dip}), **2a**, ^{12a} was identified as a side-product in the synthesis of **1a**. The solvent polarity revealed to be crucial in regulating the **1a/2a** ratio (see SI, page S13). In fact, when toluene was employed as the reaction medium in the place of dichloromethane, **2a** became the major ¹⁵⁵ product (60% yield, Scheme 2). It is noticeable that this method may provide an alternative route to W^{IV}Cl₄(DAD) coordination complexes, in the absence of a Cl-abstracting agent. ^{12a} We found

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that the reaction of WCl₆ with DAD^{Det} in toluene afforded the W(IV) complex **2b** in 40% yield (Scheme 2).



Scheme 2. Synthesis of W(IV) coordination complexes from WCl₆ and α -diimines in toluene.

The new compound **2b** was characterized by analytical and spectroscopic techniques, and by X-ray diffraction (Figure 3, Table 1). The structure of **2b** resembles that previously reported for **2a**.^{12a}

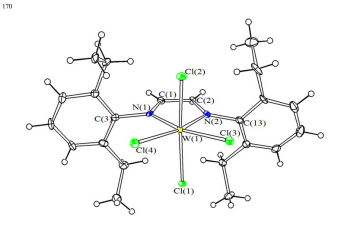


Figure 3. View of the structure of **2b** with key atoms labeled. Displacement ellipsoids are at the 50% probability level.

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W(1)-Cl(1)	2.351(3)	W(1)-Cl(2)	2.349(3)
W(1)-Cl(3)	2.296(3)	W(1)-Cl(4)	2.250(3)
W(1)-N(1)	2.032(10)	W(1)-N(2)	2.054(9)
N(1)-C(1)	1.344(15)	N(2)-C(2)	1.351(15)
C(1)-C(2)	1.381(17)		
Cl(1)-W(1)-Cl(2)	169.60(11)	Cl(3)-W(1)-N(1)	161.7(3)
Cl(4)-W(1)-N(2)	160.9(3)	Cl(3)-W(1)-Cl(4)	110.88(12)
N(1)-W(1)-N(2)	73.7(4)	W(1)-N(1)-C1	120.1(8)
N(1)-C(1)-C(2)	113.4(11)	C(1)-C(2)-N(2)	113.3(10)
C(2)-N(2)-W(1)	118.9(8)		

Table 1. Selected bond distances (Å) and angles (°) for 2b.

In summary, the unusual, *direct interaction* of N-aryl α-diimines, ¹⁸⁰ i.e. widely employed robust ligands in coordination chemistry, with a high valent metal chloride has provided the unprecedented one-pot conversion to quinoxalinium species, *via* C_{aryl}–C_{alkyl} bond activation and subsequent intramolecular C–N coupling. Based on experimental and computational outcomes, the most probable ¹⁸⁵ reaction pathway exploits the rather unique synergic properties offered by WCl₆,¹³ acting as both a powerful chlorinating species ^{13,36} and a strong oxidant.^{19,34,37} We are currently working to develop this concept in the direction to achieve unconventional transformations of suitable organic substrates.

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