Preparation of new diazene complexes of ruthenium and osmium †

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The hydrazine complexes $\text{[Ru(NH₂NH₂)L₅](BPh₄)₂}$ **1**, **8** or $\text{[Ru(NH₂NH₂)L'L₄](BPh₄)₂}$ **2** [L = P(OEt)₃, L' = PPh(OEt)₂] were prepared by allowing the corresponding hydride species $\text{RuHL}_3\text{BPh}_4$ or $\text{RuHL}_4\text{BPh}_4$ to react first with HBF₄ and then with hydrazine. Oxidation of these hydrazine complexes or $[Os(NH₂NH₂)L₅](BPh₄)₂$ with Pb(OAc)₄ at -30 °C led to the corresponding stable and isolable 1,2-diazene complexes $[M(NH=NH)L_s](BPh₄)₂$ **3**, **5** (M = Ru, Os) or $[Ru(NH=NH)L'L_4]$ (BPh₄)₂ 4. The phenyldiazene derivatives $[M(PhN=NH)L_5]$ (BPh₄)₂ 6, 7 (M = Ru, Os) were also prepared by treating the hydride [MHL**5**]BPh**4** species with phenyldiazonium tetrafluoroborate. The aquo-complex, $[Ru(H_2O)\{P(OEt)_3\}$ ₅](BPh₄)₂ **8**, was obtained by substitution of the NH=NH ligand in $[Ru(NH=NH)L_5]$ (BPh₄)₂ and was characterised by X-ray crystal structure determination. Oxidation reactions of the bis(hydrazine) complexes $[Ru(NH_2NH_2)_2L_4]$ $(BPh_4)_2$ or $[Ru(CH_3NHNH_2)_2L_4]$ $(BPh_4)_2$ [L = P(OEt)₃] with Pb(OAc)₄ were reinvestigated and were found to give the diazene complexes $\text{[Ru(NH=NH)}, L_{\text{A}}\text{[BPh}_4), 9, \text{[Ru(CH}_3N=NH)}, L_{\text{A}}\text{[BPh}_4), 11 \text{ or } \text{[Ru(CH}_3N=N)$ NH)(CH**3**NHNH**2**)L**4**](BPh**4**)**² 10**.

Introduction

Diazene NH=NH is a very reactive molecule in the free state, and disproportionates¹ at -150 °C into N₂ and NH₂NH₂. Nevertheless, it is an important molecule of possible relevance as a metal-bound intermediate in inorganic and bioinorganic N_2 fixation processes,² and useful as a reagent in stereoselective *cis*-hydrogenation of unsaturated organic compounds.**³** Stabilisation of NH=NH in standard conditions has only been achieved in a few cases by coordination to transition metals, mainly in bimetallic complexes containing a μ -NH=NH ligand.⁴ Stable derivatives containing monodentate diazene have also been reported.**⁵**

We are interested in the chemistry of partially reduced dinitrogen ligands, and have previously reported⁶ the synthesis of both methyldiazene and aryldiazene complexes of ruthenium and osmium, obtained by oxidation of the related hydrazine derivatives. However, no evidence of formation of stable complexes containing coordinated 1,2-diazene was obtained. We have therefore extended those studies, searching for an appropriate metal fragment stabilising the diazene NH=NH molecule by coordination, and results are reported here. Reinvestigation of our previous reports **⁶***a***,***^c* on oxidation of bis(hydrazine) $[M(NH_2NH_2)_2L_4]^2$ ⁺ (M = Ru, Os) derivatives, which allows bis(1,2-diazene) complexes to be prepared, is also described here.

Experimental

All synthetic work was carried out in an inert atmosphere using standard Schlenk techniques or a Vacuum Atmosphere drybox. Once isolated, the complexes were found to be relatively stable in air, but were stored in an inert atmosphere at -25 °C.

All solvents were dried over appropriate drying agents, degassed on a vacuum line and distilled into vacuum-tight storage flasks. The phosphite P(OEt)₃ (Aldrich) was purified by distillation under nitrogen; PPh(OEt)₂ was prepared by the method of Rabinowitz and Pellon.**⁷** Diazonium salts were obtained in the usual way.**⁸** Hydrazine NH**2**NH**2** was prepared by decomposition of hydrazine cyanurate (Fluka) following the reported method.**⁹** RuCl**3**3H**2**O salt was a ChemPur product and (NH**4**)**2**OsCl**6** was purchased from Johnson Matthey; both salts were used as received. High-grade (99.99%) lead(IV) acetate was purchased from Aldrich. Other reagents were purchased from commercial sources in the highest available purity and used as received. Infrared spectra were recorded on a Nicolet Magna 750 FT-IR spectrophotometer. NMR spectra (**1** H, **³¹**P) were obtained on a Bruker AC200 spectrometer at temperatures varying between -90 and $+30$ °C, unless otherwise noted. **¹** H spectra are referred to internal tetramethylsilane. $3^{31}P{^1H}$ chemical shifts are reported with respect to 85% H**3**PO**4**, with downfield shifts considered positive. The conductivity of 10^{-3} mol dm⁻³ solutions of the complexes in $CH₃NO₂$ at 25 °C was measured with a Radiometer CDM 83 instrument. **CH1**_HPh(DF), 1987
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Preparation of complexes

Hydrides RuH_2L_4 [L = P(OEt)₃, PPh(OEt)₂] and [OsH{P(O- Et ₃}₅]BPh₄ were prepared following previous methods.^{6*c*},10</sup>

 $[\text{RuHL}_5]\text{BPh}_4$ $[\text{L} = \text{P}(\text{OEt})_3$ and $\text{PPh}(\text{OEt})_2]$. Hydrides [RuHL**5**]BPh**4** were prepared by a modification of the method previously reported.**¹¹** An equimolar amount of the appropriate phosphite (0.56 mmol) was added to a solution of $RuH₂L₄$ (0.56 mmol) in 5 cm**³** of ethanol and the mixture cooled to -196 °C. A slight excess of HBF₄·Et₂O (0.57 mmol, 82 μ L) was then added and the reaction mixture, brought to room temperature, stirred for about 2 h. The addition of an excess of NaBPh₄ (1.2 mmol, 0.41 g) caused the separation of a white solid, which was filtered and crystallised from CH₂Cl₂ and ethanol; yield ≥ 80%. (For [RuH{P(OEt)**3**}**5**]BPh**4** Found: C, 51.64; H, 7.85.

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[†] Electronic supplementary information (ESI) available: observed and calculated ${}^{31}P\{{}^{1}H\}$ NMR spectra of compounds $[Ru(NH_2NH_2)-]$ ${PPh(OEt)_2}$ ${P(OEt)_3}$ $d{PBH_4}$ ₂ 2 (Fig. S1) and [Ru(NH=NH){PPh- $(OEt)_2$ } ${P(OEt)_3}$ ₄](BPh₄)₂ **4** (Fig. S2). See http://www.rsc.org/ suppdata/dt/b2/b202888n/

mol⁻¹. IR (KBr) v/cm^{-1} : 1915 (m) $v(Ru-H)$. δ _H [(CD₃)₂CO, 25 C]: 7.40–6.78 (m, 20 H, Ph), 4.11 (m, 30 H, CH**2**), 1.31, 1.30 $(t, 45 H, CH_3)$, -9.25 to -10.03 (m, 1 H, H⁻). δ_P [(CD₃)₂CO, 25 ^oC]: A₄B spin system, δ_A 142.0, δ_B 140.4, J_{AB} = 46.6 Hz).

 $\textbf{[RuH}\{\textbf{PPh}(\textbf{OEt})_2\}\{\textbf{P}(\textbf{OEt})_3\}_4\{\textbf{BPh}_4\textbf{.} \text{A slight excess of HBF}_4\textbf{.}\}$ Et₂O (0.42 mmol, 60 μ L) was added to a solution of RuH₂- ${P(OEt)_3}_4$ (0.4 mmol, 0.31 g) in 10 cm³ of C₂H₅OH and the reaction mixture was brought to about -30 °C and stirred for 30 min. A slight excess of $PPh(OEt)$ ₂ (0.42 mmol, 83 μ L) was added and the resulting solution brought to room temperature. After 1 h of stirring, an excess of NaBPh₄ (0.8 mmol, 0.27 g) was added and the white solid obtained filtered and crystallised from CH₂Cl₂ and ethanol; yield $\geq 80\%$. (Found: C, 54.34; H, 7.45. C₅₈H₉₆BO₁₄P₅Ru requires C, 54.25; H, 7.53%. Λ_M = 57.5 S cm² mol⁻¹. IR (KBr) v/cm^{-1} : 1936 (w) $v(Ru-H)$. δ _H (CD₂Cl₂, 25 C): 8.01–6.70 (m, 25 H, Ph), 4.20–3.50 (m, 28 H, CH**2**), 1.26 $(m, 42 \text{ H}, \text{ CH}_3)$, $-8.81 \text{ to } -9.70 \text{ (m, 1 H, H⁻)}$. $\delta_{\text{P}} (\text{CD}_2 \text{Cl}_2)$, 25 °C): ABC₂M spin system, $\delta_{\rm M}$ 166.4, $\delta_{\rm A}$ 141.0, $\delta_{\rm B}$ 140.1, δ _C 137.6, J_{AM} = -435.7, J_{BM} = 56.0, J_{CM} = 31.6, J_{AB} = 67.7, J_{AC} = 38.3, $J_{BC} = 55.4$ Hz).

 $[\text{Ru(NH,NH₂)}$ $[\text{P(OEt)}₃$ $]\underline{\text{O(BPh_4)}}, 1.$ An excess of $HBF_4 \cdot Et_2O$ (0.4 mmol, 57 µL) was added to a solution of hydride $[RuHL_{5}]$ BPh_4 (0.125 g, 0.1 mmol) in 10 cm³ of CH_2Cl_2 cooled to -196 °C and placed under a hydrogen atmosphere. The reaction mixture was brought to room temperature, stirred for 2 h and then, after replacing the H**2** atmosphere with argon, an excess of $NH₂NH₂$ (1 mmol, 31 μ L) was added. The resulting solution was stirred for 20 h and then the solvent was removed under reduced pressure. The oil obtained was triturated with ethanol containing an excess of $NABPh_4$ (0.2 mmol, 68 mg) giving a white solid, which was filtered and crystallised from CH₂Cl₂ and ethanol; yield ≥ 60%. (Found: C, 58.35; H, 7.58; N, 1.72. C**78**H**119**B**2**N**2**O**15**P**5**Ru requires C, 58.47; H, 7.49; N, 1.75%. $A_M = 114.9$ S cm² mol⁻¹).

 $\textbf{[Ru(NH₂NH₂)}$ $\textbf{[PPh(OEt)₂]}$ $\textbf{[P(OEt)₃}$ $\textbf{[ABPh₄)}$ 2. This compound was prepared exactly like the related pentakis(phosphite) complex 1 by reacting the hydride $\text{RuH}\{\text{PPh}(\text{OEt})_2\}\{\text{P-}$ $(OEt)_{3}$ ¹₄]BPh₄ first with HBF₄·Et₂O and then with NH₂NH₂; yield ≥ 65%. (Found: C, 60.39; H, 7.45; N, 1.76. C**82**H**119**B**2**- $N_2O_{14}P_5$ Ru requires C, 60.26; H, 7.34; N, 1.71%. Λ_M = 115.1 S $\text{cm}^2 \text{ mol}^{-1}$).

 $\textbf{[Ru(NH}_2NH_2)_2\{\textbf{PPh}(\textbf{OE}t)_2\}_4\textbf{]}(\textbf{BPh}_4)_2.$ This complex was obtained in an attempt to prepare the pentakis(phosphite) $[Ru(NH_2NH_2)\{PPh(OEt)_2\}$ ₅](BPh₄)₂ following the method reported for the related compounds **1** and **2**. In this case, the known**⁶***^a* bis(hydrazine) derivative was obtained as the only isolated product, with a yield of about 75%. (Found: C, 66.12; H, 6.95; N, 3.45. C**88**H**108**B**2**N**4**O**8**P**4**Ru requires C, 66.21; H, 6.82; N, 3.51%. $A_M = 114.7$ S cm² mol⁻¹).

 $[Os(NH₂NH₂)L₅](BPh₄)₂ [L = P(OEt)₃]$. This complex was prepared following the method previously reported.**⁶***^c*

 $[Ru(NH=NH){P(OEt)}_{3}$ ₅ $]$ $(BPh_{4})_{2}$ 3 and $[Os(NH=NH){P-C}$ $(OEt)_{3}$ **[BPh₄**)₂ 5. A sample of the appropriate hydrazine complex [M(NH**2**NH**2**)L**5**](BPh**4**)**2** (0.16 mmol) was placed in a three-necked 25 cm³ round-bottomed flask fitted with a solidaddition sidearm containing an equimolar amount of Pb(OAc)₄ (0.16 mmol, 71 mg). Dichloromethane (10 cm³) was added, the solution cooled to -30 °C and $Pb(OAc)₄$ added portionwise over 20–30 min to the cold stirring solution. The solution was then brought to 0° C, stirred for 10 min and the solvent removed under reduced pressure. The oil obtained was treated at 0° C with ethanol (2 cm^3) containing an excess of NaBPh**4** (0.2 mmol, 68 mg). A white solid slowly separated out, which was filtered and crystallised by dissolving in $CH₂Cl₂$ and, after filtration and concentration, by fast precipitation with ethanol; yield ≥ 60%. (Found: C, 58.70; H, 7.47; N, 1.70. C**78**H**117**B**2**N**2**O**15**P**5**Ru **3** requires C, 58.54; H, 7.37; N, 1.75%. $A_M = 118.4$ S cm² mol⁻¹. Found: C, 55.38; H, 7.04; N, 1.75. C**78**H**117**B**2**N**2**O**15**OsP**⁵ 5** requires C, 55.45; H, 6.98; N, 1.66%. $A_M = 120.7$ S cm² mol⁻¹).

 $\textbf{[Ru(NH=NH)\{PPh(OEt)_2\}\{P(OEt)_3\}_4\textbf{]} (BPh_4)_2}$ 4. This complex was prepared exactly like the related species **3** and **5** by oxidation with Pb(OAc)**4** of the starting hydrazine compound **2**; yield ≥ 45%. (Found: C, 60.16; H, 7.30; N, 1.61. C**82**H**117**B**2**- $N_2O_{14}P_5$ Ru requires C, 60.33; H, 7.22; N, 1.72%. Λ_M = 116.9 S $\text{cm}^2 \text{ mol}^{-1}$).

 $[Ru(PhN=NH){P(OEt)}_3{}$ ^{$[OBPh_4]_2$} 6 and $[Os(PhN=NH){P-}$ $(OEt)_{3}$ ₅ $(OBH₄)_{2}$ 7. In a three-necked 25 cm³ round-bottomed flask were placed solid samples of the appropriate hydrides [MHL₅]BPh₄ (0.1 mmol) and an excess of phenyldiazonium tetrafluoroborate $[PhN_2]BF_4$ (0.5 mmol, 96 mg) and the mixture was cooled to -196 °C. Acetone (10 cm³) was slowly added and the reaction mixture, brought to room temperature, stirred for 4 h. The solvent was removed under reduced pressure giving a brown oil which was treated with ethanol (5 cm**³**) containing an excess of $NaBPh₄$ (0.2 mmol, 68 mg). A yellow solid slowly separated out, which was filtered and crystallised from CH₂Cl₂ and ethanol; yield $\geq 75\%$. (Found: C, 60.02; H, 7.35; N, 1.60. C**84**H**121**B**2**N**2**O**15**P**5**Ru **6** requires C, 60.18; H, 7.27; N, 1.67%. $A_M = 120.3$ S cm² mol⁻¹. Found: C, 57.01; H, 6.99; N, 1.67. C**84**H**121**B**2**N**2**O**15**OsP**⁵ 7** requires C, 57.14; H, 6.91; N, 1.59%. $A_M = 117.8$ S cm² mol⁻¹).

 $\textbf{[Ru(H₂O)(P(OEt)₃)}$ **(BPh₄)₂ 8.** This complex was obtained in an attempt at crystallisation of the diazene derivative $[Ru(NH=NH)\{P(OEt)_3\}$ ₅ $[(BPh_4)_2$ 3. In fact, by slow cooling to -25 °C of a saturated solution of 3, prepared by treating the solid sample (150 mg) with ethanol (8 cm**³**) and enough CH**2**Cl**²** to obtain a saturated solution at room temperature, white microcrystals of the aquo-complex **8** were obtained. These crystals were also suitable for X-ray analysis; yield $\geq 80\%$. (Found: C, 59.11; H, 7.49. C**78**H**117**B**2**O**16**P**5**Ru requires C, 58.98; H, 7.42%. $A_M = 118.2$ S cm² mol⁻¹).

 $[Ru(CH_3N=NH)(CH_3NHNH_2){P(OEt)_3}$ ¹⁰and $[Ru-I_3]$ $(CH_3N=NH)_2$ { $P(OEt)_3$ }₄**]**(BPh_4)₂ 11. These complexes were obtained by oxidation of the [Ru(CH**3**NHNH**2**)**2**{P(OEt)**3**}**4**]- $(BPh_4)_2$ complex^{6*a*} with Pb(OAc)₄ in CH₂Cl₂ at -30 °C following the method used for **3** and **5**. In this case, by reacting 0.2 mmol (0.300 g) of $\text{[Ru(CH_3NHNH_2)_2\{P(OEt)_3\}_4\text{][BPh}_4)_2}$ with 0.4 mmol (0.177 g) of Pb $(OAc)_4$, a mixture of both compounds **10** and **11** was obtained. Their separation involves the slow cooling to -25 °C of a saturated solution of the mixture prepared at room temperature using as solvent ethanol and enough CH₂Cl₂ to dissolve the solid. The first solid obtained was the mixed-ligand compound **10** in about 25% yield, while from the mother liquor, after repeated crystallisation, the bis- (methyldiazene) **11** was separated in about 10% yield. (Found: C, 59.31; H, 7.48; N, 3.66. C**74**H**110**B**2**N**4**O**12**P**4**Ru **10** requires C, 59.48; H, 7.42; N, 3.75%. $\Lambda_M = 118.5$ S cm² mol⁻¹. Found: C, 59.42; H, 7.41; N, 3.63. C**74**H**108**B**2**N**4**O**12**P**4**Ru **11** requires C, 59.56; H, 7.29; N, 3.75%. $A_M = 108.9$ S cm² mol⁻¹).

Oxidation of $\text{[Ru(NH}_2\text{NH}_2), L_4\text{]}(\text{BPh}_4), \text{[L = P(OEt)}_3 \text{ and}$ $PPh(OEt)_{2}$ **]** and $[Os(NH_{2}NH_{2})_{2}$ { $P(OEt)_{3}$ }₄**]**(BPh_{4})₂

Also the oxidation of the bis(hydrazine) complexes **⁶***a***,***^c* was carried out with $Pb(OAc)₄$ at -30 °C, following the method used for the other hydrazine complexes 1 , 2 , $[Os(NH₂-)]$ $NH_2)L_5$ [BPh₄)₂ and [Ru(CH₃NHNH₂)₂L₄](BPh₄)₂. A typical experiment involves the addition of solid $Pb(OAc)₄$ (0.4 mmol,

Table 1 Crystal data and structure refinement for $\text{Ru}(H_2O)\$? $(OEt)_{3}$ ₅ $[(BPh_{4})_{2}$ ⁸

Empirical formula Formula weight Temperature/K Wavelength/Å Crystal system Space group alĂ <i>hl</i> Ă	$C_{78}H_{117}B_{2}O_{16}P_{5}Ru$ 1588.26 293(2) 0.71069 Triclinic P ₁ 14.763(5) 16.613(5)
β /°	18.646(5) 82.05(5)
Volume/Å ³	4529(2)
Z	\mathfrak{D}
Absorption coefficient/mm ^{-1}	0.318
F(000)	1684
Reflections collected	26755
Independent reflections	18872 [<i>R</i> (int) = 0.0555]
Data/restraints/parameters	18872/6/845
Final R indices $[I > 2\sigma(I)]$	$R1 = 0.0948$, $wR2 = 0.2383$
R indices (all data)	$R1 = 0.1649$, $wR2 = 0.2968$

0.177 g) to a cooled solution $(-30 °C)$ of the appropriate bis(hydrazine) complex (0.2 mmol) in CH₂Cl₂. Removal of the solvent at the end of the reaction gave an oil, which was triturated at 0° C with ethanol (2 cm^3) containing an excess of NaBPh**4** (0.4 mmol, 136 mg). The white solid that slowly separated out was filtered and dried under vacuum. The samples contained a mixture of bis(diazene) $[M(NH=NH)₂$ - L_4 $(BPh_4)_2$ and acetate $[M(\kappa^2-O_2CCH_3)L_4]BPh_4$ complexes, which cannot be separated in pure form. However, in the case of $\text{[Ru(NH=NH)}, \text{[P(OEt)}, \text{]}$ ₄ $\text{[BPh_4)}, 9$, the sample contains the bis(diazene) as the major product (about 60–70%), while in the other cases only little amounts $(5-10\%)$ of bis(diazene) are present in the reaction product. For the mixture containing $[Ru(NH=NH)_2\{PPh(OEt)_2\}$ ₄](BPh₄)₂ δ_H (CD₂Cl₂, 20 °C): 16.23, 15.62 (dm, 2 H, NH=NH; ${}^{3}J_{\text{HH}} = 32.0$ Hz). For the mixture containing $[Os(NH=NH)_2{P(OEt)_3}$ ₄ $](BPh_4)_2$ δ_H (CD_2Cl_2) , 20 °C): 16.63, 16.31 (dm, 2 H, NH=NH).

Crystallography

Crystals were air sensitive, a white irregular prism single crystal was sealed in a glass capillary along with some drops of solution. X-Ray diffraction data were collected on a Bruker-Siemens SMART AXS 1000 equipped with a CCD detector, using graphite monochromated Mo-K α radiation ($\lambda = 0.71069$) Å). Data collection details are: crystal to detector distance $=$ 5.0 cm, hemisphere mode, time per frame = 30 s, oscillation $\Delta \omega = 0.300^{\circ}$. Data reduction was performed up to $\theta = 28^{\circ}$ by the SAINT package¹² and data were corrected for absorption effects by the SADABS**¹³** procedure. Data collection and refinement results are summarised in Table 1. The crystal suffered a severe merohedral twinning expressed by a mirror plane perpendicular to *b*, which simulated a 2/*m* Laue symmetry, whilst the actual structure is described in $P\bar{1}$, with a and β close to 90°. The phase problem was solved by direct methods¹⁴ and the structure was refined by full matrix least squares on all F^2 by taking into account the twinning as implemented in Shelxl 97,**15** using the WinGX package.**16** The refinement without considering the twinning was unsuccessful. Anisotropic displacement parameters were refined for all nonhydrogen atoms, while hydrogen atoms were introduced in calculated positions, except for those belonging to the water, located on the Fourier maps. Phenyls were treated as rigid bodies. Use of the Cambridge Crystallographic Database **¹⁷** facilities was made for structure discussion. Meaningless electron density residues were left around the metal in the final map.

CCDC reference number 182442.

See http://www.rsc.org/suppdata/dt/b2/b202888n/ for crystallographic data in CIF or other electronic format.

Results and discussion

Hydrazine complexes of ruthenium $\text{[Ru(NH, NH)}\text{]}$ Et)₃} $\{P(OEt)_{3}\}\{P(OEt)_{3}\}$ **1** and $[Ru(NH_{2}NH_{2})\{PPh(OEt)_{2}\}$ $\{P(OEt)_{3}\}$ (BPh_4) , 2 were prepared by reacting the hydride $[RuHL]$ -L**4**]BPh**⁴** first with an excess of fluoroboric acid and then with hydrazine, as shown in Scheme 1.

$$
[RuHL'L_{4}]^{+}\xrightarrow{exc. HBF_{4}} \begin{bmatrix} [Ru(\eta^{2}-H_{2})L'L_{4}]^{2+} \\ \xrightarrow{exc. NH_{2}NH_{2}} \end{bmatrix} \longrightarrow
$$

\n
$$
= H_{2} \begin{bmatrix} [Ru(\eta^{2}-H_{2})L'L_{4}]^{2+} \\ 1.2 \end{bmatrix}
$$

Scheme 1 $L' = L = P(OEt)$, **1**; $L' = PPh(OEt)$, $L = P(OEt)$, **2**.

Reactions of the pentakis(phosphite)hydride $[RuHL'L_4]^+$ with an excess of HBF_4 gives the $\eta^2 - H_2$ dihydrogen intermediate $[M(\eta^2 - H_2)L'L_4]^2$ ⁺ (by ¹H NMR), which reacts with NH₂NH₂ to give final hydrazine complexes **1** or **2**. In contrast, the reaction of the hydride $\text{[RuH}\{\text{PPh}(\text{OEt})_2\}$ ₅ IBPh_4 , first with HBF_4 and then with hydrazine, in all conditions afforded the known^{6*a*} bis(hydrazine) complex $\text{[Ru(NH}_2\text{NH}_2)_2\{\text{PPh(OEt)}_2\}_4\}^{2+}$, as shown in Scheme 2.

In the pentakis(phosphite) $[Ru(\eta^2 - H_2)L_5]^{2+}$ intermediate, hydrazine replaces not only the labile η**²** -H**2** ligand, but also one phosphite, giving the bis(hydrazine) species. Attempts to prepare other pentakis(phosphite) hydrazine complexes different from 1 and 2 failed, because the reaction of $\text{[RuHL/L}_4]^+$ first with a Brønsted acid and then with an excess of CH₃NHNH₂ or C**6**H**5**NHNH**2** only gave mixtures of solid products not containing the hydrazine ligand. It seems that only the $NH₂NH₂$ species can bind to the metal centre in the ML^{'L}₄ fragment to give stable and isolable derivatives.

Hydrazine complexes **1** and **2** were isolated as white [BPh**4**] salts, stable in air and in solutions of polar organic solvents, in which they behave as 2 : 1 electrolytes. Analytical and spectroscopic data (Table 2) support the proposed formulation. The presence of the hydrazine ligand was supported by infrared spectroscopy, which showed the characteristic νNH mediumintensity bands of NH_2NH_2 between 3387 and 3266 cm⁻¹. Further support for the presence of $NH₂NH₂$ came from ¹H NMR spectra, which showed two broad signals at 4.37 and 3.36 ppm (**1**) or at 4.41 and 3.46 ppm (**2**), due to the two NH**²** proton resonances. Integration measurements and homodecoupling experiments confirmed the proposed assignment. In the temperature range between $+30$ and -80 °C, the ³¹P{¹H} NMR spectrum of $[M(NH_2NH_2)L_5]^2$ ⁺ 1 appeared as an AB₄ multiplet, fitting the proposed formulation (geometry **I**, Fig. 1). In contrast, the spectra of $\text{[Ru(NH}_2\text{NH}_2)\{\text{PPh}(\text{OEt})_2\}\{\text{P-}$ (OEt)**3**}**4**](BPh**4**)**² 2** showed a rather complicated pattern, simulated as a set of two systems of types AB₄ and ABC₂M, partly overlapping, and with the parameters listed in Table 2. This result may be interpreted as due to two geometries in solution of the type shown in Fig. 1, with the hydrazine and $PPh(OEt)_{2}$ ligands in mutually *trans* (**II**) or *cis* (**III**) positions, respectively.

Pentakis(phosphite)hydrazine complexes of both ruthenium **1**, **2** and osmium^{6*c*} $[Os(NH₂NH₂)L₅](BPh₄)$ ₂ reacted with $Pb(OAc)₄$ at low temperature (-30 °C) with the selective oxidation of the $NH₂NH₂$ ligand to give 1,2-diazene complexes $[M(NH=NH)L₅]²⁺$ (M = Ru 3, Os 5) and $[Ru(NH=NH)\{PPh-$

^a In KBr pellets. ^b In CD₂Cl, at 25 °C, unless otherwise noted. ^c Phenyl proton resonances are omitted. ^d Positive shift downfield from 85% H₃PQ₄. ϵ In $(CD_3)_2CO$. The sample contains the acetate complex $\left[\text{Ru}(\kappa^2-O_2CCH_3)\right]\left\{\text{P(OEt)}\right\}\hat{}_{34}\left\{\text{BPh}_4\right\}$

 $(OEt)_2$ ₂²⁺ **4**, which were isolated as white solids and characterised (Scheme 3).

It was essential for the successful synthesis of diazene derivatives 3–5 to carry out reactions at low temperature $(-30\degree C)$ and to use high-purity Pb(OAc)**4** (99.99%) in equimolar amounts. Otherwise, mixtures of products not containing the 1,2-diazene complex were obtained. In particular, the use of common Pb(OAc)₄ (95%) did not yield pure samples of $3-5$, but only mixtures containing small amounts $(10-15%)$ of the diazene complex.

All NH=NH derivatives 3–5 are stable as solids, but slowly decompose in solution, losing the diazene ligand. An attempt to crystallise **3** in a mixture of ethanol and dichloromethane as solvent gave aquo-complex [Ru(H₂O)(P(OEt)₃) ₅ [BPh₄) ² **8**, which was isolated and characterised. The H₂O molecule probably comes from the traces of water always present in the common "anhydrous" solvents. In the solution the presence of hydrazine was also detected and confirms the decomposition of free NH=NH.

Diazene complexes **3**–**5** are 2 : 1 electrolytes, and their analytical and spectroscopic data (Table 2) support the proposed formulation. In the low-field region of the **¹** H NMR spectrum of complex $[Ru(NH=NH)L₅]²⁺$ 3, two multiplets (Fig. 2) at

 ${}^{1}H$ NMR

Fig. 2 Observed (bottom) and calculated (top) **¹** H NMR spectra, in the diazene region, of $\text{[Ru(NH=NH)\{P(OEt)_3\}}$ **5** $\text{[(BPh_4)_2 3, in CD_2Cl_2}$ at 25 °C. The simulated spectrum was obtained using the parameters reported in Table 2.

16.76 and 15.36 ppm appeared, which were attributed to the two NH protons of a monohapto NH=NH ligand. This pattern is due to coupling with **³¹**P nuclei (AB**4** multiplet), as confirmed by simulation of the spectrum using an AB_4XY model (X = HA, Y = HB), with the parameters listed in Table 2. The ${}^{3}J_{\text{HH}}$ value of 32.5 Hz (J_{XY}) also suggested ^{4*d*,5} a probable *trans*-NH= NH geometry. The related osmium complex $[Os(NH=NH)L₅]²⁺$ **5**, in the low-field region of the proton spectra, also showed the characteristic A**4**BXY multiplet of the diazene ligands. In the temperature range between -80 and $+30$ °C, the ³¹P{¹H} NMR spectra of both complexes 3 and 5 appeared as AB_4 or A_4B multiplets, fitting geometry **IV** (Fig. 3).

The presence of the diazene ligand was also confirmed in the mixed-phosphite derivative $[Ru(NH=NH)\{PPh(OEt)\}$ $(OEt)_{3}$ ²₄](BPh₄)₂ 4, the ¹H NMR spectrum of which showed the characteristic NH multiplet between 17 and 15 ppm (Table 2). In the temperature range between -80 and $+30$ °C, the ³¹P{¹H} NMR spectrum appeared as a complicated pattern, simulated

as a set of one AB**4** and one ABC**2**M multiplets, partially overlapping. On this basis, and taking into account the **³¹**P spectra of hydrazine precursor **2**, we propose the existence of two isomers of the type shown in Fig. 4 for our complex **4**, containing the NH=NH and PPh (OEt) ₂ ligands in mutually *cis* (V) or *trans* (**VI**) positions, respectively.

Fig. 4 $L = P(OEt)_{3}$, $L' = PPh(OEt)_{2}$.

Complexes containing the monodentate 1,2-diazene ligand are very rare,**⁵** and only one example **⁵***^c* of isolable species has been reported for ruthenium and osmium metal centres. Using the pentakis(phosphite) $ML₅$ (or $ML'L₄$) fragment stabilises the NH=NH molecule through coordination, affording stable and isolable species for both $Ru(II)$ and $Os(II)$.

Unfortunately, no other $[M(RN=NH)L₅]²⁺$ diazene complexes besides **3** and **5** could be prepared by oxidation of coordinated hydrazine, owing to the absence of the corresponding precursors. In our hands, hydrazines RNHNH₂ different from $NH₂NH₂$ could not be coordinated to the $ML₅$ fragment to give stable species. However, taking into account that aryldiazene can also be prepared by insertion of aryldiazonium cations into a M–H bond, we treated the $[MHL_5]^+$ $(M = Ru, Os)$ hydrides with the phenyldiazonium cation PhN_2 ⁺ BF_4 ⁻ and observed the formation of aryldiazene derivatives [M(PhN $NH)L₅$ ²⁺ (M = Ru 6, Os 7) which were isolated as $BPh₄⁻$ salts and characterised (Scheme 4).

Scheme 4 $M = Ru 6$, Os 7; $L = P(OEt)$ ₃.

The formation of complexes **6** and **7** may indicate that the steric requirement of aryldiazene PhN=NH is lower than that of arylhydrazine PhNHNH**2**, whose coordination on the ML**⁵** fragment did not take place. Aryldiazene [M(PhN=NH)L₅]- $(BPh₄)₂$ derivatives **6** and **7** are stable as solids and in solutions of polar organic solvents, in which they behave as 2 : 1 electrolytes. The presence of the diazene ligand was confirmed by the **1** H NMR spectra, which displayed the characteristic highfrequency NH proton resonances at 13.72 (**6**) and 14.15 (**7**) ppm. In the temperature range between -90 and $+30$ °C, the **³¹**P{**¹** H} NMR spectrum appeared as an AB**4** multiplet, fitting the proposed formulation.

Table 3 Selected bond lengths (A) and angles $(°)$ for $\text{[Ru(H, O)/P(O Et$)₃}₅](BPh₄)₂ **8**, with s.u.s in parentheses

Ru – $O16$ $Ru-P4$ $Ru-P1$	2.219(7) 2.238(3) 2.348(3)	$Ru-P3$ $Ru-P2$ $Ru-P5$	2.352(3) 2.356(3) 2.372(3)
$O16 - Ru - P4$ $O16 - Ru - Pl$ $P4 - Ru - P1$ $O16 - Ru - P3$ $P4 - Ru - P3$ $P1-Ru-P3$ $O16 - Ru - P2$ $P4 - Ru - P2$	178.3(2) 89.9(2) 91.1(1) 80.7(2) 98.0(1) 91.81(9) 87.0(2) 94.3(1)	$P1 - Ru - P2$ $P3-Ru-P2$ $O16 - Ru - P5$ $P4-Ru-P5$ $P1-Ru-P5$ $P3-Ru-P5$ $P2-Ru-P5$	88.2(1) 167.7(1) 86.8(2) 92.1(1) 176.6(1) 88.7(1) 90.6(1)

Scheme 5 $L = P(OEt)$ ³, $M = Ru$.

Although aryldiazenes **6** and **7** were stable in solution, related 1,2-diazene complexes **3**–**5** slowly decomposed giving aquocomplex $[M(H_2O)L_5]^2$ ⁺, probably by substitution of the labile NH=NH ligand with traces of water present in the solvent. In the case of ruthenium, compound [Ru(H**2**O)L**5**](BPh**4**)**² 8** (Scheme 5) was isolated as white microcrystals and characterised spectroscopically (Table 2) and crystallographically (Table 1). The presence of H_2O as a ligand in the complex was indicated by the proton NMR signal at 2.65 ppm, and the ${}^{31}P{^1H}$ spectra confirmed the ML₅ fragment.

The structure and atom labelling of $[Ru(H_2O){P(OEt)_3}_5]^2$ ⁺ is shown in Fig. 5, and the coordination geometry is summarised in Table 3. The metal coordination is octahedral, with ligand

Fig. 5 Perspective view of the solid-state molecular structure of cation $[Ru(H_2O)\{P(OEt)_3\}^2$ ⁺ $\mathbf{8}^{2+}$. Anisotropic displacement parameters are at the 50% probability level. Ethoxy groups are omitted for clarity.

at P4 located *trans* to the water molecule, the remaining phosphites being sterically equivalent to each other. The steric crowding of the bulky phosphite ligands is partly relieved by pushing them towards the water position, thus reducing the width of the *cis* O–Ru–P bond angles, which range between $80.7(2)$ and $89.9(2)$ °. The presence of the water ligand also affects the Ru–P4 bond *trans* to it [2.238(3) Å], which is far shorter than the remaining *cis* ones [2.348(3)–2.372(3) Å]. The large *trans* influence of the unique $P(OEt)$ ³ ligand is evidenced by the fact that the Ru–O bond is among the longest found in Ru^{2+} water complexes (*e.g.* $Ru-O = 2.139$ Å in the hexa-aquo cation**¹⁸**). There are no intermolecular interactions among non-hydrogen atoms below 3.55 Å in the structure.

Results from oxidation reactions of hydrazine complexes [M- $(NH_2NH_2)L_5]^2$ ⁺ using the new high-purity Pb(OAc)₄ (99.99%) prompted us to reinvestigate the oxidation of the bis(hydrazine) complexes $[M(NH_2NH_2)_2L_4]^2$ ⁺, that were previously studied by us,^{6*a*,*c*} using low-purity Pb(OAc)₄ (95%) as oxidising agent. These earlier reactions did not give diazene complexes, but only acetate derivative $[M(\kappa^2-O_2CCH_3)L_4]^+$. The reaction was studied in CH_2Cl_2 at -30 °C using high-purity $Pb(OAc)_4$ in a Ru : Pb ratio 1 : 2, and showed that, in the case of [Ru(NH**2**- $NH₂$ ${}_{2}$ {P(OEt)₃}₄]²⁺ it does give a white solid containing a mixture of bis(diazene) $\text{[Ru(NH=NH),L₄](BPh₄)}$, 9 and acetate $[Ru(\kappa^2-O_2CCH_3)L_4]BPh_4$ derivatives, as shown in Scheme 6.

Scheme 6 $L = P(OEt)$ ₃.

From this mixture, we attempted to separate the diazene complex by fractional crystallisation at low temperature, but the instability in solution of $\left[\text{Ru(NH=NH)}_{2}L_{4}\right]^{2+}$ on the one hand, and the comparable solubility of the two complexes on the other, always gave samples containing more products. However, the **¹** H NMR spectra of the mixture strongly supports the formation of bis(diazene) **9**, showing two doublets of multiplets at 16.71 and 15.83 ppm, attributed to the two NH protons of the two diazene ligands. The ${}^{3}J_{\text{HH}}$ value of 32 Hz also suggests^{4*d*,5} a *trans*-NH=NH geometry. The ³¹P{¹H} NMR spectra of the oxidation product also shows two A_2B_2 multiplets, due to acetate compound $6a$ [Ru(κ^2 -O₂CCH₃)L₄]⁺ and bis(diazene) complex **9**, fitting the proposed formulation. The A**2**B**2** pattern also suggests the mutually *cis* positions of the two diazene ligands, as in geometry **VIII**, Scheme 6.

Oxidation of the related $\text{[Ru(NH, NH_2), \{PPh(OEt),\}_a\}.$ $(BPh_4)_2$ and $[Os(NH_2NH_2)_2{P(OEt)_3}_4](BPh_4)_2$ derivatives afforded white solids, with **¹** H NMR spectra showing diazene signals, probably attributable to bis(diazene) $[M(NH=NH)₂-]$ L_4 [BPh₄)₂ (M = Ru, Os) complexes. In these cases, however, the predominant species is acetate compound $[M(\kappa^2-O_2CCH_3)L_4]^+$, with the diazene complex present only in a small amount (from 5 to 10%).

These studies on the oxidation of bis(hydrazine) complexes with high-purity Pb(OAc)₄ (99.99%) show that 1,2-diazene complexes can also be formed in these cases, but their stabilisation on the ML**4** fragment is restricted, as compared with related $[M(NH=NH)L₅]²⁺$ derivatives, and does not allow separation of the samples in pure form. However, although substitution of the NH=NH ligand by the acetate ion also takes place using high-purity Pb(OAc)**4**, a solid sample containing the coordinated 1,2-diazene (**9**) as the predominant species was isolated. In addition, these results confirm our previous hypothesis^{6a,*c*} on the formation of a diazene complex in the oxidation of $[M(NH_2NH_2)_2L_4]^2$ ⁺ cations, and highlight the importance of the purity of Pb(OAc)**4** in determining the nature of the final oxidation products.

On this basis, we extended the oxidation reaction to methylhydrazine complexes [Ru(CH**3**NHNH**2**)**2**L**4**] **²** and found that, using high-purity Pb(OAc)₄ (99.99%), the reaction gives a mixture of bis(methyldiazene) $[Ru(CH_3N=NH)_2L_4]^{2+}$ 11 and methyldiazene–methylhydrazine [Ru(CH₃N=NH)(CH₃NH- $NH_2)L_4$ ²⁺ 10 complexes, which were isolated as BPh_4^- salts and characterised (Scheme 7).

Complexes **10** and **11** were separated by fractional crystallisation into two pale yellow solids, stable in air and in solutions of polar organic solvents, in which they behave as 2 : 1 electrolytes. Analytical and spectroscopic data (Table 2) support the proposed formulation. The **¹** H NMR spectrum of bis(methyl-

diazene) complex **11** shows a slightly broad multiplet at 13.58 ppm, characteristic of the diazene ligand. Beside the signals of phosphite and BPh**4** ion, the spectrum also contains, a singlet at 3.81 ppm, due to the methyl–proton resonance of the methyldiazene group. In the temperature range between -80 and +30 °C, the ³¹P{¹H} NMR spectra appear as an A_2B_2 multiplet, fitting the presence of two diazene ligands in a mutual *cis* position (**IX**).

In mixed-ligand complex $\text{[Ru(CH_3N=NH)(CH_3NHNH_2)}$ -L**4**](BPh**4**)**² 10**, the **¹** H NMR spectra show the presence of both nitrogenous ligands. The broad multiplet at 13.74 ppm of the NH proton and the singlet at 3.76 ppm of the methyl resonances confirm the presence of the $CH₃N=NH$ ligand, whereas the broad signals at 4.64 and 3.45 ppm, due to $NH₂$ and $NH₂$ resonances respectively, and the doublet at 2.65 ppm are diagnostic of the hydrazine group. The ABC₂ multiplet present in the **³¹**P{**¹** H} spectra also suggests mutual *cis* position of the hydrazine and diazene ligands in the complex (**X**).

The RuL**4** fragment is therefore able to stabilise both 1,2 diazene and methyldiazene by coordination, affording the related bis(diazene) species, which are very stable, in the case of $CH₃N=NH$. Partial substitution was observed with the NH=NH ligand, affording a mixture of diazene (9) and acetate [Ru(κ**²** -O**2**CCH**3**)L**4**] derivatives. However, isolation of solid samples containing diazene complexes of ruthenium was only achieved by using high-purity Pb(OAc)**4** (99.99%), and the absence in previous studies⁶ of any evidence of these diazene species is probably due to the fact that standard-grade $Pb(OAc)₄$ (95%) was used, which leads to decomposition of diazene derivatives (see ESI).

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