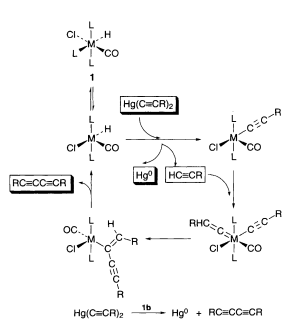
Transalkynylation and catalytic demercuration of bis(alkynyl)mercurials: two alternative mechanisms

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The complexes $[RhCl(CO)(PPh_3)_2]$ and $[MHCl(CO)(PPh_3)_3]$ (M = Ru, Os) catalyse the demercuration of $[Hg(C\equiv CR)_2]$ (R = Ph, C₆H₄Me-4, Buⁿ) to provide buta-1,3-diynes, however the reactions proceed by quite different mechanisms as indicated by (*i*) the isolation of $[Ru\{C(C\equiv CR)=CHR\}Cl(CO)(PPh_3)_2]$ and its reintroduction into the catalytic cycle and (*ii*) the isolation of the iridium complex $[IrCl(C\equiv CR)(HgC\equiv CR)-(CO)(PPh_3)_2]$.

The coupling of terminal alkynes by copper salts has long been established and remains the preferred route for the formation symmetrical buta-1,3-diynes.¹ Bis(alkynyl)mercurials, [Hg(C=CR)₂] emerged early as convenient crystalline derivatives for the characterisation of terminal alkynes² although their deployment as alkynyl transfer reagents has not been extensively studied. We have described unsuccessful³ attempts to use these reagents for the mild transfer of alkynyl groups to transition metals, in particular ruthenium.⁴ Our unsuccessful efforts led to the development of a convenient route to 3-en-1-yn-3-yl complexes of ruthenium, $[Ru{C(= CHR)C=CR}]$ - $Cl(CO)(PPh_3)_2$] [R = Ph 2a, C₆H₄Me 2b, Buⁿ 2c] via the reaction of [RuHCl(CO)(PPh₃)₃] 1a with [Hg(C=CR)₂] (Scheme 1)³ however we find that this route fails for the corresponding osmium complex [OsHCl(CO)(PPh₃)₃] 1b. Heating 1b with 1 equiv. of [Hg(C=CPh)₂] in refluxing toluene leads as in the case of 1a to the deposition of elemental mercury, however the only organometallic complex isolated is unreacted 1b. Indeed if 1b is treated under similar conditions with a



Scheme 1 Catalytic demercuration of bis(alkynyl)mercurials by 1b (L = PPh₃, R = Ph, C₆H₄Me, Buⁿ)

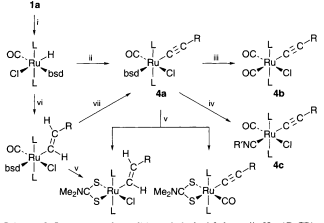
tenfold excess of [Hg(C=CPh)2] the only isolated species (after 2 min) are 1b, elemental mercury and diphenylbutadiyne. Thus the complex 1b catalyses the extrusion of mercury from $[Hg(C \equiv CPh)_2]$. Heating $[Hg(C \equiv CPh)_2]$ alone in refluxing toluene fails to produce mercury deposition thereby confirming the complicity of 1b in the reaction. The mechanism for this reaction would appear to reflect, at least initially, that for the formation of 1a. In the osmium case, however the complex $[Os{C(=CHPh)C=CPh}Cl(CO)(PPh_3)_2]$ 2d is unstable under the reaction conditions, proceeding via β-Os-H elimination to provide 'OsHCl(CO)(PPh₃)₂' for re-entry into the catalytic manifold (Scheme 1). These observations led us to investigate the reaction of $[Hg(C=CPh)_2]$ with 5 mol% of 2a which led after 1-2 min to the formation of PhC=CC=CPh and recovery of 2a. Furthermore similar treatment of 2a with $[Hg(C \equiv CC_6H_4Me)_2]$ provided MeC₆H₄C=CC=CC₆H₄Me and **2b**, thereby verifying the proposed mechanism.

In the initial preparation of 1a, the facile reaction of the putative intermediate complex $[Ru(C=CR)Cl(CO)(PPh_3)_n]$ (n = 2, 3, ?) with terminal alkynes is presumed to be due to the lability of one of the bulky triphenylphosphine ligands. Accordingly, the reactions of the complex [RuClH-(CO)(bsd)(PPh_3)_2] **3** (bsd = 2,1,3-benzoselenadiazole)⁵ with $Hg(C \equiv CR)_2$ (R = Buⁿ, C₆H₄Me-4, Ph) were investigated, assuming that bsd would compete more effectively with alkyne than PPh₃. The complexes [Ru(C=CR)Cl(CO)(bsd)(PPh₃)₂] 4a were obtained in good yield under surprisingly mild conditions (CH₂Cl₂, 25 °C, 30 min) accompanied by the formation of 1 equiv. of free alkyne and elemental mercury. Similar results are obtained with the hydride complexes [RuClH(CO)(PPh₃)₂L'] (L' = pyridine, dimethylpyrazole) however the products are colourless or pale yellow. The bsd ligand is apparently sufficiently strongly bound to prevent reaction with liberated RC=CH, but labile enough to be replaced by sterically modest π -acidic ligands 'L' (CO, CNR) to provide stable octahedral acetylide complexes $[Ru(C=CR)Cl(CO)(PPh_3)_2L]$ (L = CO 4b; $\pm CNC_6H_3Me_2-2,6$ 4c), one example of which (4b, R = C₆H₄Me-4) has been crystallographically characterised.§ The dicarbonyl derivatives could also be prepared by the reaction of [Ru(CH=CHPh)Cl(bsd)(CO)(PPh₃)₂] with an excess of RC=CH in refluxing ethanol, followed by addition of CO (Scheme 2). Reaction of $[Ru(C \equiv CC_6H_4Me-4)Cl(bsd)(CO)(PPh_3)_2]$ with Na(S₂CNMe₂) in CH₂Cl₂-EtOH leads to the formation of $[Ru(C \equiv CC_6H_4Me-4)(S_2CNMe_2)(CO)(PPh_3)_2]$ along with a small amount of a second compound identified as [Ru(CH= CHC₆H₄Me-4)(S₂CNMe₂)(CO)(PPh₃)₂] which could be prepared independently by the reaction of $Na(S_2CNMe_2)$ with [Ru(CH=CHC₆H₄Me-4)Cl(bsd)(CO)(PPh₃)₂].⁵

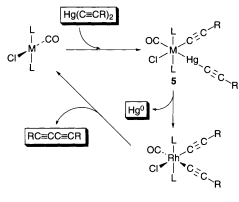
It is perhaps surprising that **4a** fails to react cleanly with electrophiles at the carbon β to ruthenium (to form vinylidenes) in contrast to the copious precedent for acetylides of divalent ruthenium.⁴ Rather, reaction with HCl leads to cleavage of the acetylide and formation of [RuCl₂(CO)₂(PPh₃)₂] suggesting protonation at the ruthenium–carbon bond or ruthenium itself followed by reductive elimination. We note that all successful acetylide to vinylidene conversions on divalent ruthenium have

so far involved complexes with donor ligands (rather than π -acids) coordinated *trans* to the acetylide.

Collman and Kang have reported the oxidative addition of bis(phenylethynyl)mercury to Vaska's complex to provide the complex [Ir(C=CPh)(HgC=CPh)Cl(CO)(PPh_3)_2] **5a**.⁶ We considered that such a strategy might provide access to bis(alkynyl) complexes of rhodium following thermal extrusion of mercury from the presumably more labile Rh–Hg–C linkage, given that we have observed **5a** to be both thermally and photochemically labile. The reaction of [RhCl(CO)(PPh_3)_2] with [Hg(C=CPh)_2] does not however *ultimately* provide [Rh(C=CPh)(HgC=CPh)Cl(CO)(PPh_3)_2] **5b**. Rather, following deposition of elemental mercury, only unreacted [RhCl(CO)(PPh_3)_2] is obtained in addition to diphenylbutadiyne. Furthermore we find that the complex [Rh(CO)(PPh_3)_2 [H_2B(bta)_2]] (bta = benzotriazolyl)⁷



Scheme 2 Reagents and conditions: i, bsd, thf, heat; ii, Hg (C=CR)₂, CH₂Cl₂, 25 °C; iii, CO, CH₂Cl₂, 25 °C; iv, CNR', CH₂Cl₂, 25 °C; v, NaS₂CNMe₂·2H₂O, CH₂Cl₂-EtOH; vi, HC=CR, thf, heat; vii, HC=CR, EtOH, heat (L = PPh₃, R = C₆H₄Me; R' = C₆H₃Me₂)



 $Hg(C\equiv CR)_2 \longrightarrow Hg^0 + RC\equiv CC\equiv CR$

Scheme 3 Catalytic demercuration of bis(alkynyl)mercurials by [RhCl(CO)(PPh_3)_2] (L = PPh_3, R = Ph, C_6H_4Me, Bu^n)

also behaves in a similar manner. The mechanism for this catalytic demercuration is therefore presumably as shown in Scheme 3 and is clearly distinct from that in Scheme 1.

To conclude, two catalytic cycles for the demercuration of bis(alkynyl)mercurials have been demonstrated. Furthermore novel hydride/acetylide metathesis has been shown to be effected under very mild conditions by transalkynylation. The convenience of preparation of alkynyl mercurials² makes these routes to both butadiynes and σ -acetylides attractive.

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Footnotes

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\$ Selected data for 4b: Yield (1.0 mmol scale) 0.77 g (93%). IR (Nujol): 1986, 2048 (v_{CO}), 2111 (v_{C=C}). NMR (CDCl₃, 25 °C): ¹H δ 2.27 (s, 3 H, Me), 6.74, 6.90 [(AB)₂, 4 H, C₆H₄, J(AB) 7.9 Hz]; ¹³C{¹H} δ 194.6 [t, RuCO, J(PC) 11.6 Hz], 192.0 [t, RuCO, J(PC) 8.0 Hz], 115.5 (s, RuC=C], 106.8 [t, RuC=C, J(PC) 19.7 Hz], 21.3 [CH₃]; ³¹P{¹H} δ 23.1. FAB MS m/z832 [M]+, 804 [M – CO]+, 797 [M – CI]+, 717 [M – CCR]+, 689 [M – CCR - CO]+, [RuCO(PPh₃)₂]+, 625 [Ru(PPh₃)₂]+, 363 [RuPPh₃]+. § Crystal data for 4b: $C_{47}H_{37}ClO_2P_2Ru \cdot 0.75CHCl_3$, M = 921.8, monoclinic, space group $P2_1/n$, a = 10.216(2), b = 37.395(8), c = 23.007(3) Å, $\beta = 102.29(1)^\circ$, U = 8588(3) Å³, Z = 8 (two crystallographically independent molecules in the asymmetric unit), $D_c = 1.43 \text{ g cm}^{-3}$, μ (Mo- $K\alpha$ = 0.68 mm⁻¹, F(000) = 3756. 15 118 Independent reflections (Mo-K α radiation; graphite monochromator, ω -scans). $R_1 = 0.068$, $wR_2 = 0.127$ for 8663 independent observed reflections $[|F_o| > 4\sigma(|F_o|), 2\theta < 50^\circ]$ and 879 parameters. Whilst the analysis unambiguously defined the identity and stereochemistry of the complex, minor structural anomalies in the bond lengths precluded detailed discussion. Atomic coordinates, bond lengths and angles, and thermal parameters have been deposited at the Cambridge Crystallographic Data Centre (CCDC). See Information for Authors, Issue No. 1. Any request to the CCDC for this material should quote the full literature citation and the reference number 182/25.

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