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Oxidative $C(sp^3)$ -H Bond Cleavage, C-C and C=C Coupling at a Boron Center with O₂ as Oxidant Mediated by Platinum(II)

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Dimethyl- and diphenylplatinum(II) fragments $Pt^{II}R_2$ (R = Me, Ph) enable facile and efficient oxidative $C(sp^3)$ -H bond cleavage and stepwise C-C and C=C coupling at the boron atom of a coordinated 1,5-cyclooctanediyldi(2-pyridyl)borato ligand with O₂ as the sole oxidant.

The selective oxidation of C-H bonds using dioxygen, an inexpensive and environmentally benign reagent, catalyzed by electrophilic transition metals and pioneered by Hay¹ and Shilov² has attracted recently much attention.³⁻⁷ In turn, organoboron compounds have emerged as valuable nucleophilic partners in various oxidative coupling reactions allowing for the formation of C-C and a variety of C-X bonds.8 In spite of the great practical importance of the transition metal chemistry of organoboranes, little is known about the mechanisms of reactions of organoboron compounds and electrophilic transition metals. In the transition metal - mediated C-C coupling reactions of organoboron compounds it is usually presumed that the boron atom is only involved in the transmetallation of a boron-bound hydrocarbyl to an electrophilic metal whereas the high-valent transition metal center is solely responsible for the product - forming step.^{3,8} Previously, we have characterized the direction and the nature of the migration of the methyl and phenyl groups R between an anionic boron atom of the dimethyl- and diphenyldi(2-pyridyl)borates R₂B(C₅H₄N)₂ (L) and the nearby electrophilic $Pt^{IV}\hat{R'}_2$ fragment (R' = Me, Ph) which can result, in particular, from the oxidation with O2 of the organoplatinum(II) complexes Na[LPt^{II}R'₂] (see examples in Scheme 1).⁹⁻¹¹ In this work we report that the anionic alkylborate 1 present in the Pt^{II} complexes Na[2] - Na[5] (Chart 1) can be involved in a Pt - mediated double oxidative cleavage of the bridgehead alkylborate $C(sp^3)$ -H bonds accompanied by an unprecedented C=C coupling at the *boron* center, with O₂ as the sole oxidant. According to our mechanistic proposal, the C-C/C=C coupling occurs at the boron center.

The new borate ligand and the derived anionic Pt^{II} complexes in the form of their sodium salts, Na[2] - Na[5], were prepared using standard synthetic techniques and fully characterized. The chloride salt of the doubly protonated dipyridylborate, H[1]·HCl, was also characterized by single crystal X-ray diffraction.

The reactivity of the new organoplatinum(II) organoborates

Scheme 1. Oxidatively induced B-to-Pt^{IV} hydrocarbyl transfer.⁹⁻

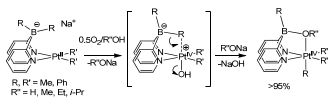
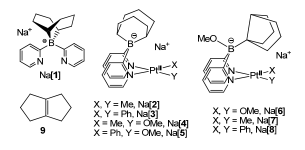
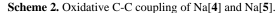


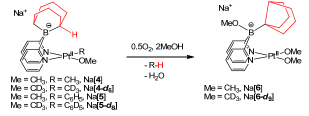
Chart 1. New borate ligand and derived compounds.



Na[2] - Na[5] toward O_2 in methanolic solution was studied next. We anticipated that compounds Na[2] - Na[5] bearing one or two hydrocarbyl groups at the Pt^{II} center will be reactive toward O_2 in methanol.^{9-10,12-14} We also expected that the 9borabicyclo[3.3.1]nonane (9-BBN) fragment present in these complexes will resist the B-to-Pt^{IV} hydrocarbyl migration due to the constraints imposed by its bicyclic structure.

While the platinum-free compound Na[1] dissolved in methanol showed no change under 1 atm O_2 after 4 days at 20 °C, the aerobic oxidation of the hydrocarbyl methoxo Pt^{II} complexes Na[4] and Na[5] in methanol- d_4 was facile and took less than 5 min under the conditions indicated above (Scheme 2). Both reactions led to the formation of a platinum-containing product Na[6] in >96% NMR yield, H₂O and a hydrocarbon, methane or benzene, respectively, identified by means of NMR spectroscopy. The identity of Na[6] was confirmed by the Electro-Spray Ionization Mass-Spectrometry (ESI-MS) and single crystal X-ray diffraction. This characterization revealed that the 1,5-cyclooctanediyl fragment present in Na[4] and





Na[5] is converted to the bicyclo[3.3.0]octyl group (Fig. 1, a) as a result of the cleavage of one of the ligand bridgehead C-H bonds and an intramolecular $C(sp^3)$ - $C(sp^3)$ coupling. The dimethoxoplatinum(II) complex Na[6] is completely inert under 1 atm O₂ in methanol at 20 °C for at least 2 days.

Further exploration of the unusual aerobic C-C coupling reactivity of the Pt^{II} complexes derived from Na[1] involved the diphenylplatinum(II) complex Na[3]. The reaction of Na[3] under 1 atm O₂ in methanol was complete after 10 min at 20 °C and led to a virtually quantitative formation of the bicycloolefin 9, as confirmed by ¹H and ¹³C NMR spectroscopy and GS-MS, dimethoxodi(2-pyridyl)borato¹⁵ diphenylplatinum(IV) hydroxo complex 11 (isolated yield 95%) and two moles of H₂O (Scheme 3). In contrast to the monophenyl complex Na[5], no benzene formed in this reaction. Solid reaction residues were strongly alkaline confirming the formation of sodium methoxide. The identity of 11 was proven by ESI-MS and single crystal X-ray diffraction (Fig. 1b). Hence, a double oxidative cleavage of the ligand 1' bridgehead C-H bonds and an intramolecular oxidative C=C coupling were observed in the reaction of O₂ with the diphenylplatinum(II) complex Na[3] in methanol.[†]

The stepwise character of the oxidative C=C coupling in Scheme 3 was proven in the following experiments. When reaction of Na[**3**] with O_2 was performed under conditions of starvation of oxygen using only 0.5 equiv. O_2 the cleavage of one of the ligand **1**⁻ bridgehead C-H bonds and the selective formation of a C-C coupled product Na[**9**] (94% NMR yield), the diphenylplatinum(II) analog of Na[**7**], was observed.

The identity of Na[9] was confirmed by ¹H NMR spectroscopy and ESI-MS. A small amount of **11** (6% NMR yield) and the equivalent quantity of **9** were also detected, presumably because of a non-ideal control of the reaction stoichiometry. When more oxygen was admitted to the reaction mixture containing Na[8], all of the complex was converted cleanly to **9** and **11**.

A similar reaction of O_2 with the dimethylplatinum(II) complex Na[2] was more challenging to characterize. The use of a low temperature of -60 °C was required to avoid fast protonolysis of Na[2] as well as the protonolysis of the expected intermediate Na[7]. After warming from -60 °C up to 20 °C a solution of the dimethylplatinum(II) complex Na[2] in MeOH while bubbling slowly O_2 through it the formation of **9** and **10** (88% isolated yield), the dimethyl analog of **11**, was

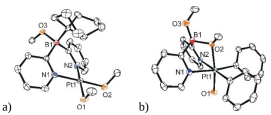


Figure 1. ORTEP drawings (50% probability ellipsoids) of: a) the anionic fragment of Na[6], b) complex **11**. Hydrogen atoms are omitted for clarity.

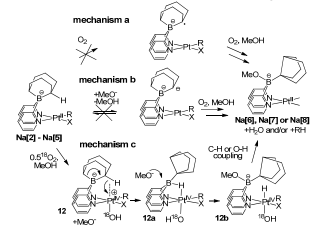
Scheme 3. Aerobic C=C coupling of Na[2] and Na[3].

observed. No methane was detected.

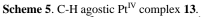
An oxidative C-C coupling of lithium salts containing B,Bdialkyl-9-boratobicyclo[3.3.1]nonane anions to form bicyclo[3.3.0]octylboranes using acyl chlorides as oxidants was reported in mid 1970s but it was never explored mechanistically.¹⁶ To the best of our knowledge, the examples of the platinummediated oxidative C-H cleavage are very rare¹⁷ and the C=C coupling in Scheme 3 is unique.^{8,18} These considerations motivated us to carry out some mechanistic tests. To probe if free alkyl radicals are involved in the oxidative C-H bond cleavage / C-C coupling of Na[2] - Na[5] (Scheme 4, mechanism a), the oxidation with O_2 of two representative compounds, Na[3] and Na[4], was performed in the presence of 2 equiv. of a radical trap TEMPO. These reactions produced no O-alkyl TEMPO derivatives that might have resulted from the recombination of TEMPO and transient free alkyl species. The reactions also were not inhibited. These facts do not support mechanism **a**. In addition, no H/D exchange at the ligand bridgehead carbon atoms in Na[3] dissolved in CD₃OD was observed for at least 3 days so arguing against an initial deprotonation of the ligand bridgehead C-H bonds and subsequent oxidation of the resulting carbanions (Scheme 4, mechanism b).

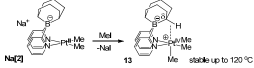
The C-H bond cleavage / C-C coupling mechanism (Scheme 4, mechanism c) involving the formation of a Pt^{IV} center and the hydride abstraction by Pt^{IV} allows to account for the available observations as explained below. First, the oxidation of Pt^{II} hydrocarbyl complexes with O₂ and H₂O₂⁺ in hydroxylic solvents produces Pt^{IV} hydroxo species with an *axial* hydroxo ligand, such as **12**, with the oxygen atom of the hydroxo ligand originating from the oxidant.¹²⁻¹⁴ Indeed, we observed the formation of the ¹⁸O-labeled product **11**-¹⁸O in the reaction of Na[**3**] and ¹⁸O₂ (Scheme 3). Second, the intermediates **12** derived from the complexes Na[**2**] – Na[**5**] transfer the hydride anion from a bridgehead C-H bond of the 9-BBN fragment to the Pt^{IV} center to form Pt^{IV} hydrides **12a** and then **12b**. Third, the hydride ligand at the Pt^{IV} atom of **12b** is oxidatively coupled with either the hydrocarbyl R to produce

Scheme 4. Possible mechanisms of the C-H bond cleavage.



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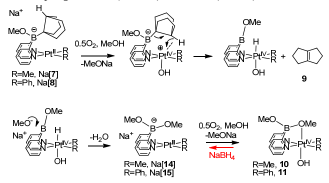
methane or benzene along with Na[6] or with the OH ligand to form H_2O along with the hydrocarbyl Pt^{II} complex Na[7] or Na[8].[‡] Overall, the reactions of Na[4] and Na[5] with O₂ lead to Na[6], H₂O and R-H as the final products (Scheme 4, mechanism c); the dimethoxoplatinum(II) complex Na[6] is too electron-poor to further react with O₂.

In support to the C-to- Pt^{IV} hydride transfer mechanism c, the origin of the protium atom in the hydrocarbon R-H resulting from the oxidation of complexes Na[4] and Na[5] (Scheme 2) was revealed using the partially deuterated compounds $Na[4-d_6]$ and Na[5- d_8]. The oxidation of the labeled complexes in methanol- d_4 leads to the formation of CD₃H or C₆D₅H, respectively, along with Na[6-d₉]. The only source of protium in the reaction mixtures above that could be involved in the formation of CD₃H and C₆D₅H is the 9-BBN fragment. The facile hydride abstraction from the 9-BBN moiety may be due to the rigid structure of 12 that poses the hydrogen atom of one ligand bridgehead C-H bond in the close proximity of the Pt^{IV} center so helping diminish the reaction activation barrier. Some metal complexes derived from 1,5cyclooctanediylbis(1-pyrazolyl)borate, a dipyrazolyl analogue of 1 having the 9-BBN framework, were reported to have an enhanced C-H agostic interactions of their bridgehead C-H bonds with the metal.¹⁹ In support to this hypothesis, a stable trimethylplatinum(IV) complex 13 (Scheme 5) was prepared as a model of 12. Complex 13 was characterized by single crystal X-ray diffraction and found to have a short 1.99 Å agostic $(\delta\text{-}C)H-Pt^{IV}$ bond.§

As opposed to the reaction of Na[4] and Na[5] with O_2 , the reaction of Na[2] and Na[3] with O_2 does not stop at the formation of the C-C coupled product. The dihydrocarbyl platinum(II) complexes Na[7] and Na[8] are much more reducing compared to Na[6] and can be involved in another O_2 activation / hydride abstraction - oxidative C-C coupling / O_2 activation reaction sequence to produce 9 along with 10 and 11, respectively (Scheme 6). In support to the mechanism in Scheme 6, Na[15] was produced *in situ* by reacting the diphenylplatinum(IV) complex 11 with NaBH₄ or NaBH(OMe)₃ in methanol. The solution of Na[15] then reacted with O_2 to form 11 at a fast rate in a quantitative NMR yield.

The observed reactivity suggests that **11** may be a good catalyst for the oxidation with O₂ of some hydride donors. Indeed, the oxidation with O₂ of 0.7M Na[BH(OMe)₃] or NaBH₄ in methanol / NaOMe solutions is catalyzed efficiently by 0.5 mol % **11** with the turnover frequency of ~170 h⁻¹ and ~216 h⁻¹, respectively, at 20 °C.

Scheme 6. Proposed mechanism of the oxidative C-H bond cleavage / C=C coupling of Na[7] (R = Me) and Na[8] (R = Ph).



In summary, we have characterized a rare oxidative Ptmediated $C(sp^3)$ -H bond cleavage accompanied with an unprecedented C=C coupling at the boron center of the 1,5cyclooctanediyldi(2-pyridyl)borato platinum(II) complexes Na[2] – Na[5]. These reactions formally involve two consecutive hydride abstractions from the borate ligand occurring, most likely, via agostic C-H ... Pt^{IV} intermediates. Up to three O₂ activation steps at a single Pt^{II} atom may be involved in these facile transformations. Further study of this novel system might lead to some practical applications utilizing O₂ in catalytic oxidation and C=C coupling of organoboron and similar compounds.

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Notes and references

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^{\dagger} The use of H₂O₂ instead of O₂ in the reaction with Na[**3**] also leads to a fast formation of **9** and **11** (90% NMR yield).

[‡] The methoxo ligand oxygen lone pairs in **12b** (X = OMe vs. X, R = Me or Ph) may favor a faster C-H coupling as a result of their π -donation to the Pt-C antibonding orbital while impeding attacks of bases (e.g., MeO') at the Pt^{IV}-H fragment required for the O-H coupling. § In ¹H NMR spectra (CD₂Cl₂) the agostic proton resonance is at -3.36 ppm with the large coupling constant $J_{Pt-H} = 208$ Hz. The inertness of **13** toward the C-H – to - Pt^{IV} hydride transfer, as compared to **12**, may result from a much stronger *trans*-influence of the CH₃ ligand *trans*- to the agostic C-H bond in **13** *vs.* the OH ligand in **12**, which strongly discourages the hydride migration.

Electronic Supplementary Information (ESI) available: the experimental details. CCDC 932583 - 932586. See DOI: 10.1039/c000000x/

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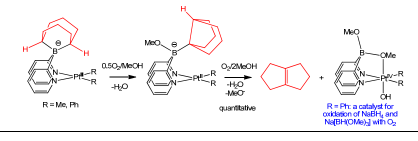
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Platinum(II) 1,5-cyclooctanediyldi(2-pyridyl)borates react with O_2 via $C(sp^3)$ -H bond cleavage and stepwise C-C and C=C coupling of the borate ligand.