

Upgrading Light Hydrocarbons via Tandem Catalysis: A Dual Homogeneous Ta/Ir System for Alkane/Alkene Coupling

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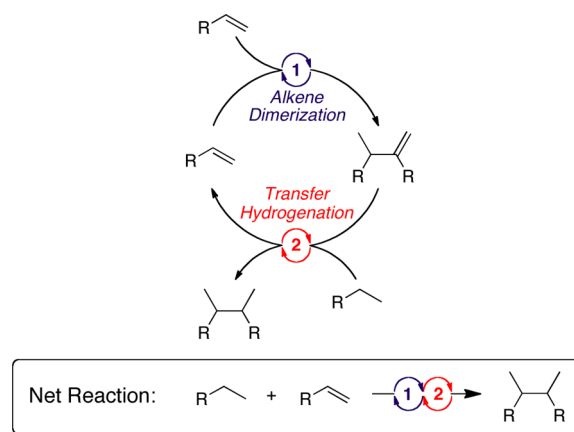
Supporting Information

ABSTRACT: Light alkanes and alkenes are abundant but are underutilized as energy carriers because of their high volatility and low energy density. A tandem catalytic approach for the coupling of alkanes and alkenes has been developed in order to upgrade these light hydrocarbons into heavier fuel molecules. This process involves alkane dehydrogenation by a pincer-ligated iridium complex and alkene dimerization by a $\text{Cp}^*\text{TaCl}_2(\text{alkene})$ catalyst. These two homogeneous catalysts operate with up to 60/30 cooperative turnovers (Ir/Ta) in the dimerization of 1-hexene/*n*-heptane, giving $\text{C}_{13}/\text{C}_{14}$ products in 40% yield. This dual system can also effect the catalytic dimerization of *n*-heptane (neohexene as the H_2 acceptor) with cooperative turnover numbers of 22/3 (Ir/Ta).

Light hydrocarbons, especially those with 4–6 carbons, constitute a significant but underutilized fraction of carbon-based energy carriers.¹ Because of their volatility and low energy density, these materials cannot be used as transportation fuels on a large scale. While pure light alkenes such as ethylene and propylene are extensively used in polymer and chemical synthesis,² mixed alkane/alkene streams, which are unsuitable as monomers or precursors, are abundant byproducts of catalytic cracking³ and Fischer–Tropsch synthesis.⁴ Furthermore, with increased exploitation of natural gas,⁵ heavy oils from bitumen and kerogen,⁶ and lignocellulose,⁷ the proportion of light hydrocarbons in the global energy mix will only increase. One method to turn light hydrocarbons into useful fuels would be to upgrade these feedstocks to higher-molecular-weight compounds. In this vein, alkane metathesis has emerged as a possible technology for upgrading light alkanes via combined alkane dehydrogenation and alkene metathesis; however, such a process tends to afford a statistical distribution of products, with little selectivity for a particular desired weight fraction.^{8–10} While this chemistry may hold promise, a more selective method would simplify purification processes and thus increase the product yield and efficiency.

Herein we describe a complementary approach for upgrading light hydrocarbons based on a tandem alkane dehydrogenation/alkene dimerization sequence. This process takes advantage of the mixed nature of many light byproduct streams by incorporating both alkanes and alkenes as substrates. In an ideal system (shown for a linear alkane and 1-alkene in Scheme 1), one catalyst would dimerize the alkene component of the mixed feedstock to a C_{2n} alkene. Subsequent transfer hydrogenation by

Scheme 1. Idealized Tandem Catalytic Approach toward Alkane/Alkene Upgrading



a second catalyst would convert the alkane component to a 1-alkene while hydrogenating the C_{2n} product to an alkane. The 1-alkene thus formed could then be catalytically dimerized with a second equivalent of 1-alkene, and the cycle would continue; the net reaction would be coupling of the alkane and alkene to give the higher alkane without generation of byproducts. Calculations indicate that such a reaction should be thermodynamically favored below $\sim 250^\circ\text{C}$;¹¹ therefore, the catalysts for dimerization and transfer hydrogenation must operate with appreciable rates at relatively mild temperatures.

The present work represents our initial efforts to develop such an upgrading scheme, involving a dual homogeneous catalytic system in which the tantalum catalyst $\text{Cp}^*\text{TaCl}_2(\text{alkene})$ (1) ($\text{Cp}^* = \text{C}_5\text{Me}_5$) effects alkene dimerization¹² and alkane/alkene transfer hydrogenation is carried out by an iridium catalyst (2).^{8b,10,13} We demonstrate that these two species function in tandem to effect alkane/alkene coupling, although the complete synthetic cycle has not yet been fully achieved.

Often the major challenge in developing a tandem catalytic process is catalyst compatibility, particularly in homogeneous reactions.^{8b,14} In the present case, alkane/alkene transfer hydrogenation requires high temperatures and relatively long reaction times to achieve high conversion. Pincer-ligated iridium catalysts are the current state-of-the-art homogeneous systems for these transformations and have demonstrated applicability in tandem reactions, namely, alkane metathesis with group-6 cocatalysts.^{8b,10}

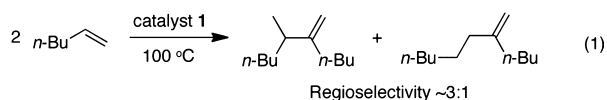
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While these Ir catalysts are reported to be active at temperatures as low as 100 °C,¹³ many alkene dimerization catalysts are temperature-sensitive with short lifetimes.¹⁵ Furthermore, Ir–pincer catalysts are known to be very sensitive to trace impurities, including Lewis acids such as BR₃ (and even N₂!),¹³ whereas many alkene dimerization systems require strong Lewis or Brønsted acid activators, such as methylaluminoxane (MAO) or other group-13 compounds.^{15,16} Finally, deactivation of one or both catalysts by ligand transfer is a concern; for example, Ir–pincer catalysts are known to undergo chloride ligand transfer from Grubbs-type Ru alkene metathesis catalysts, rendering them inactive.^{8b}

In light of these potential complications, we chose the Ta-based alkene dimerization catalyst **1**, developed by Schrock,¹² for our first trials. Cp*TaX₂(alkene) complexes are reported to be “indefinitely active” for the selective dimerization of 1-alkenes to two regioisomers at temperatures up to 100 °C (eq 1); they are inert to internal alkenes and the product 1,1-disubstituted alkenes. Importantly, no cocatalyst or activator is required.



To demonstrate catalytic alkane/alkene coupling, we examined the conversion of a 1-hexene/*n*-heptane mixture in the presence of the Ir transfer hydrogenation catalyst (*t*-Bu₄[POCOP]-Ir(C₂H₄) (**2a**)^{13d} or *t*-Bu₄[PCP]IrH₄ (**2b**)^{13a} and Cp*TaCl₂(C₂H₄) (**1**)^{12c} (Figure 1); the formation of C₁₃ and C₁₄ products here would signal the operation of tandem catalysis.

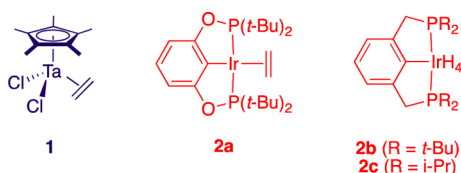
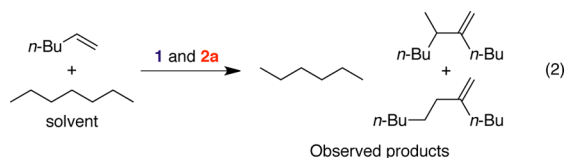


Figure 1. Precatalysts investigated for tandem catalysis.

Heating a solution of [POCOP]Ir catalyst **2a** and **1** in 1-hexene/*n*-heptane (250–1300 mM alkene) under a range of reaction conditions (100–150 °C, 1–48 h, Ta:Ir = 1–3:1) gave no measurable amount of C₁₃ or C₁₄ products; however, it is evident from GC analysis of the resulting hydrocarbon mixture that both catalysts did operate: nearly all of the 1-hexene was consumed, with *n*-hexane and C₁₂ hexene dimers as the major observed products (eq 2 and Figure S5¹¹). *n*-Hexane resulted

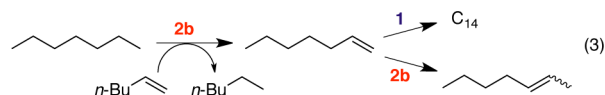


from Ir-catalyzed transfer hydrogenation, while the C₁₂ products formed via Ta-catalyzed 1-hexene dimerization. Hence, deactivation caused by catalyst incompatibility is not responsible for the lack of cooperation. Instead, we hypothesize, as have others,^{8b,17} that catalyst **2a** is not selective for 1-alkene formation, resulting in an overall concentration of 1-heptene that was too low for catalyst **1** to incorporate; internal heptenes, which are not dimerized by catalyst **1**, were therefore the other hydrogen transfer products.¹⁸

In contrast, use of the known terminal-selective^{13c} Ir catalyst **2b** (10 mM) with **1** (16 mM) afforded *n*-hexane, internal hexenes, and C₁₂, C₁₃, and C₁₄ alkenes; the latter two products were formed in a combined yield of 22% (Table 1, entry 1). [The identities of the C₁₂/C₁₃/C₁₄ products were confirmed by comparison to a product mixture resulting from codimerization of 1-hexene and 1-heptene by **1** (Figure S4).¹¹] Thus, this dual catalyst system can incorporate the *n*-alkane solvent into the dimerization cycle, demonstrating the feasibility of the proposed tandem catalytic process. Under these reaction conditions, the catalysts exhibited a “cooperativity”¹⁹ of 35%. The full synthetic cycle of Scheme 1 was not completed, however, as no C₁₂/C₁₃/C₁₄ alkanes were observed; separate control experiments showed that the 1,1-disubstituted alkene isomers afforded by **1** are poor acceptors for hydrogen transfer promoted by **2b**.

To assess the factors that influence the efficiency of this tandem Ta/Ir system, a number of reaction parameters were varied (Table 1). Reducing the catalyst loadings to 8 and 5 mM for **1** and **2b**, respectively, improved the overall turnover number (TON) without sacrificing product yield or cooperativity (entry 2). Decreasing the reaction temperature to 100 °C dramatically improved the cooperativity (63%) with a slight reduction in yield (entry 3), while altering the **1**/**2b** ratio (entries 4 and 5) was detrimental to the cooperativity. Notably, increasing [1-hexene] to 1000 mM merely resulted in the production of more C₁₂ dimer (entry 6); the absolute amounts of C₁₃ and C₁₄ were nearly identical to those obtained with 250 mM 1-hexene (entry 3). Another intriguing observation is that in most cases the amounts of C₁₃ and C₁₄ were very similar. One would expect the C₁₃ codimer to be the major C₇-containing species in view of the large excess of 1-hexene present relative to the amount of 1-heptene generated.

Together, these two observations suggest that most of the tandem catalytic productivity occurs only after high conversion, when [1-hexene] is low. To confirm this, the reaction from entry 3 of Table 1 was monitored over time (Figure 2). Initially, 1-hexene was rapidly consumed and transformed mainly into C₁₂ by homodimerization. The C₁₃ product formed slowly, and the C₁₄ dimer did not form to a significant degree until nearly three half-lives had passed; more than half the amount of C₁₄ was generated after >98% of the 1-hexene was consumed. These results indicate that the relative rates of dimerization and dehydrogenation are best matched at low [1-alkene], in accord with previous observations that high alkene concentrations inhibit transfer hydrogenation.¹³ In addition, competitive alkene isomerization is minimized at low [1-alkene]; isomerization reduces the catalyst cooperativity by converting 1-heptene into internal heptenes that do not undergo dimerization (eq 3).



Importantly, control experiments showed that the initial rates of dimerization and transfer hydrogenation in the tandem catalytic reaction are identical to the catalytic rates exhibited by the corresponding catalysts operating separately (Figure S8).¹¹ This indicates that the two catalysts truly operate independently in solution under these conditions, with no mutual inhibition or deactivation.

The reaction profile in Figure 2 indicates that better C₁₃/C₁₄ yield, catalyst cooperativity, and TONs might be achieved by maintaining a steady, low concentration of 1-hexene.

Table 1. Evaluation of Catalyst Cooperativity in 1-Hexene/*n*-Heptane Coupling by **1** and **2b**^a

1 and 2b
5 hours

Observed products

C₁₂: R = R' = *n*-C₄H₉
C₁₃: R = *n*-C₄H₉, R' = *n*-C₅H₁₁ *or*
R = *n*-C₅H₁₁, R' = *n*-C₄H₉
C₁₄: R = R' = *n*-C₅H₁₁

entry	[1-hexene] ₀ (mM)	[1]/[2b] (mM)	T (°C)	<i>n</i> -hexane (mM) ^b	C ₁₂ (mM) ^b	C ₁₃ /C ₁₄ (mM) ^b	TON for 1 ^c	TON for 2b ^c	coop. (%) ^d
1	250	16/10	125	112	57	14/13	5 (2)	11 (4)	35
2	250	8/5	125	106	58	15/13	11 (4)	21 (8)	39
3	250	8/5	100	52	81	13/10	13 (3)	10 (6)	63
4	250	5/5	100	78	75	13/9	19 (4)	15 (6)	41
5	250	12/5	100	53	85	9/8	8 (1)	11 (5)	47
6	1000	8/5	100	87	432	18/10	58 (4)	17 (8)	45
7	1200 ^e	8/5	98 ^f	329 ^g	293	181/59	67 (30)	66 ^g (60)	91 ^g

■ ACKNOWLEDGMENTS

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- (18) Efforts to quantify internal heptenes by GC were hampered by the large peak for the *n*-heptane reaction solvent, which obscured the peaks for many of these isomers (as determined by coinjection of 2- and 3-heptenes with *n*-heptane).
- (19) Here we define “cooperativity” as the amount of heptene generated by dehydrogenation that is incorporated into C₁₃ or C₁₄ dimers. Since the [1-heptene] produced is equal to the [*n*-hexane] observed, the cooperativity factor is given by $([C_{13}] + 2[C_{14}])/[n\text{-hexane}]$. For the reaction in eq 4, the cooperativity is equal to $([C_{13}] + 2[C_{14}])/[\text{neoheptane}]$.
- (20) The initial [*n*-hexane] was not zero because of activation of precatalyst **2b**. Because the Ir tetrahydride was used, 2 equiv of 1-hexene per equivalent of catalyst was hydrogenated immediately upon mixing; since the catalyst loading was 5 mM, the initial [*n*-hexane] was ~10 mM.