Hydrozirconation–Isomerisation. Reactions of Terminally Functionalised Olefins with Zirconocene Hydrides and General Aspects

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Hydrozirconation of internal aliphatic olefins, with terminal oxygen-, sulfur- or nitrogen-containing functionalities, has been studied. Internally unsaturated ethers, sulfides, a phenylsulfonyl and an ammonium iodide underwent substantial elimination of the functional group, after rearrangement of the zirconium moiety towards the carbon bearing the heteroatoms.

With aromatic olefins, acceptable yields of terminal alkyldienzirconiums (based on deuteration) were usually difficult to achieve. One exception was the reaction of 1-phenyl-1-hexene with ‘Cp*ZrCl [H]Cl’ (Cp* = η5-C5H4Me). Pronounced cleavage of the methyl-sulfur bond occurred in the hydrozirconation of 1-(1-hexenyl)-2-thiomethylbenzene.

Results from deutero- and hydrozirconations of ordinary alkenes indicated dissociation of the metal hydride from the olefins during rearrangement.

Hydrometallation–isomerisation of internal olefins is a way of making unactivated methyl hydrogens accessible to substitution [eqn. (1)]. Among existing procedures, hydroalumination,1–12 hydroboration,13–15 or hydrozirconation1,2 seem to be methods of choice with respect to isomerisation.7 Both hydroalumination and hydroboration require high temperatures for rearrangement to occur, which can cause side reactions.16 For hydroalumination, isomerisation towards the chain terminus is facilitated by the presence of catalysts,17–19 which may also be the case for hydroboration. Irrespective of whether the aluminium or boron approach is used, heteroatoms in the olefinic substrate1,2, Y: eqn. (1)] can decrease the terminal selectivity.17,20–23

\[
\begin{align*}
\text{CH}_3 \cdot \text{(CH}_2)_m \cdot \text{CH} &= \text{CH} \cdot \text{(CH}_2)_n \cdot Y \\
1)^{[\text{M}] \cdot \text{H}} &\rightarrow \text{E} \cdot \text{(CH}_2)_m \cdot \text{n} \cdot \text{n} \cdot \text{n} \cdot \text{Y} \\
2)^{\text{E}^+} 
\end{align*}
\]

Schwartz and coworkers reported the hydrozirconation procedure to be a very facile method of isomerisation under mild conditions [eqn. (2)]. They also showed that the organometallic products (like aluminium and boron alkyls) were useful in organic synthesis. The number of different functionalities that can be introduced by cleavage of the C–Zr bond has increased in recent years.24–27

Since our aim was to use derivatives of unsaturated fatty acids, occurring naturally in large amounts, for the preparation of α,ω-disubfunctionalised compounds (which may also be prepared from internal acetylenes bearing a terminal heteroatom-containing substituent28), we considered hydrozirconation to be an attractive method. Furthermore, we wished to explore the influences of various functionalities on migration of the metal. Studies on aromatic olefins were also undertaken, knowledge of the hydrozirconation of such substrates was very limited at the initial stage of our work.

In a very recent review on hydrozirconation,8 most of our work is not included. We now summarise our...
results, which show some important limitations of hydrozirconation. Finally, a few general comments on hydrozirconation–isomerisation are made.

Results and discussion

The regiochemical outcome of rearrangement was chiefly established by $^2$H NMR spectroscopy after deuteriolysis of the hydrozirconation mixtures. The deuterium content was determined by $^2$H NMR or MS analyses. Reported yields were, if not stated otherwise, determined by GLC in combination with MS.

1. Hydrozirconation of long-chain functionalised aliphatic olefins. We set out to examine derivatives of the readily available oleic and erucic acids. Acids are reduced by Cp$_2$Zr(H)Cl, but can be converted into the corresponding oxazolines, which are inert towards reduction. Attempts to prepare aldehydes from a long-chain internally unsaturated oxazoline through hydrozirconation and subsequent reaction with carbon monoxide were unsuccessful. In contrast, a shorter-chain oxazoline, having a terminal double bond, could be $\alpha$-formylated. The resulting aldehyde was, however, only stable in solution, and therefore we turned to hydroxylation reagents for cleavage of the C–Zr bond.

Hydrozirconation of oleic [1, eqn. (3)] and erucic acid oxazolines gave $\beta$- and $\omega$-hydroxylated compounds after oxidation of intermediate alkylzirconiums. The formation of the $\beta$-OH isomer indicated stability of its zirconium precursor. Stabilisation is presumably due to coordination of oxygen (or nitrogen) to oxophilic zirconium.

When deuterium oxide was used for cleavage of the C–Zr bonds, yet another positional isomer could be observed [x-D; eqn. (4)]. The entire process was also carried out with the oxazoline of hexanoic acid. This control experiment showed no deuterium incorporation. All attempts to oxidize $\alpha$-zirconium intermediates, obtained from short-chain $\alpha$, $\beta$-unsaturated oxazolines, to the corresponding $\alpha$-hydroxy compounds, failed. Clearly, caution must be used when comparing the regiochemical outcome of reactions in which different electrophiles have been utilised. Regioselectivity, with respect to the $\omega$-deuterated isomer, was not improved using THF as the solvent.

The disappearance of unsaturated material was relatively rapid in dioxane (compared with when benzene or toluene were used as solvents) at a slightly elevated temperature ($40^\circ$C). Ether and hexane were not appropriate as solvents. With long-chain substrates, 2 equivalents of the reagent were necessary to achieve complete conversion in dioxane.

Small structural modifications of the protective group cause alterations of the regiochemistry [eqns. (5) and (6)]. Moreover, the position of the double bond influences the composition of deuterated products.

The oxazolines shown in eqn. (7) are mainly hydrogenated upon treatment with [Zr]-H and subsequent deuteriolysis. The reason for this is not understood. Other $\alpha$, $\beta$-unsaturated oxazolines did not show this behaviour under the same conditions. However, at higher temperatures ($\geq 50^\circ$C) hydrogenation, as well as decomposition of the reagent, can become a major problem.

The investigations so far showed that complete regioselectivity was not easily achieved, and called for more experimentation with different functionalities. Our interest was turned to amines, ethers and sulfur-containing compounds derived from oleic acid.

Hydrozirconation results obtained with oleyl amines 2a–c are shown in Fig. 1. As before, use of dioxane...
proven advantageous. It is reasonable to assume that zirconium–heteroatom interactions in intermediate 1- and 2-zirconocenyl species are responsible for the unsatisfactory selectivity. That the proportion of α-zirconium intermediate actually differs significantly when using amine 2a, compared with 2b and 2c, is not certain, since the degree of deuteration differs somewhat in these cases, and by-products (although formed to a limited extent) were obtained.

A considerable amount of cleavage product was afforded by (Z)-9-octadecenyl trimethylammonium iodide [eqn. (8)], under the conditions used for amines.\textsuperscript{39}

When the oleyl ethers 3a–c were subjected to this reaction (20 h, 40–45°C, 3 equivalents of in-situ generated [Zr]–H), migration towards the functional group also resulted in β-elimination (Fig. 2).\textsuperscript{40} Although we never isolated the zirconium alkoxides or 1-octadecene, the occurrence of this type of elimination has been confirmed by others.\textsuperscript{30,41} Neither the saturated analogue of 2d, nor that of 3b, underwent cleavage when treated with the zirconium hydride under identical conditions. With (Z)-9-octadecenyl tert-butylidemethylsilyl ether [i.e., when R in Fig. 2 is –Si(Me)\textsubscript{2}-tert-Bu] elimination also occurred to a certain extent, but in another fashion, producing stearyl alcohol.

It was also of interest to examine the reactions of sulfur-containing substrates, and to compare their reactivity with the ethers. Here too, substantial elimination was observed [eqn. (9)], to give 33–56% of octadecane.\textsuperscript{42}

\[ \text{Y} = \text{–SMe, –SPh, –SO\textsubscript{2}Ph} \]

\[ \begin{align*}
\text{C}_{18}\text{H}_{37}\text{Se} + \text{C}_{18}\text{H}_{37}\text{Y} + \text{unconsumed olefin(s)} \\
\text{1) 3 equiv. Cp}_{2}\text{ZrHCl} \\
\text{2) D}_{2}\text{O} \\
\end{align*} \]

Fig. 2.
Proton donors react with the metal hydride, one equivalent being wasted when oleyl alcohol is subjected to hydridetonation. Only limited cleavage of the functional group was seen here. The reason for this is probably trapping of the zirconium moiety at the γ-position (as indicated by γ-deuteriation) by the formation of a five-membered cyclic intermediate, in which zirconium is connected to oxygen.

Once again it was shown that the apparent regiochemistry was dependent on the electrophile. The o : γ ratio was 62–68: 32–38 and 43–50: 50–57 when deuterium oxide or bromine was used, respectively. Oleyl alcohol and its lithium salt give similar product compositions after hydrozirconation-deuteriolysis.

In summary, hydrozirconation-isomerisation does not seem to be an effective way of producing long-chain, doubly functionalised compounds from derivatives of naturally occurring fatty acids. It may also prove difficult to apply this synthetic scheme to short-chain heteroatom-containing systems other than oxazolines (which also give products of mixed composition[38,39]), unless some variation is introduced, e.g., modification of the reagent. A few attempts were made in this direction (Section III).

It should be mentioned that very long-chain alkenes, on the other hand, are synthetically useful substrates when using the Cp₂Zr(H)Cl reagent (Fig. 3).

**II. Hydrozirconation of aromatic olefins.** We turned our attention to the substitution of alkenes with a terminal aromatic function. Reactions of Cp₂Zr(H)Cl (generated in situ, if not otherwise stated) with substrates 4–9 were initially examined with the aim of preparing zirconium derivatives such as 10. First, 1-phenyl-1-hexene (4) was chosen. Compounds of this type are readily available from various coupling reactions.

Dioxane was used as the solvent throughout the entire study discussed in this section. The reaction of 4 is slow and requires somewhat elevated temperatures. It should be mentioned that all comparisons between rates of reaction of aromatic olefins are based on arylalkane formation (upon quenching) within 1–10 h. As can be seen in eqn. (10), the benzylic isomer (x) predominated almost exclusively at room temperature, which is in agreement with Gibson’s findings[46] (this similar study appeared in the literature during the course of our work[45]). That a major part of the metal is actually directed towards the α-carbon during initial addition at 20–25 °C is indicated by the deuterium distribution in the phenylhexane formed by reaction of 4 with Cp₂Zr(D)Cl (isolated) and subsequent hydrolysis: [β-D/(α-D + β-D)] × 100 = 93 % (compare eqn. (10)).

At 60 °C, good α-selectivity was achieved, but the yield of deuterated phenylhexane decreased owing to competing hydrogenation[45] (thermal decomposition[37,38]).

The unconjugated olefin 5 reacted much faster than 4 at room temperature, but even at 45 °C the terminal selectivity was quite unsatisfactory.

The deuterium distribution with olefin 6 was not appreciably different from that found using 4 under the same conditions (20–45 °C).

Several examples of useful synthetic transformations of the thiophene nucleus are found in the literature.[45] These can in principle be used for preparative purposes after hydrozirconation-functionalisation. We therefore decided to study the reaction of substrates 7–9 with the zirconium hydride. Heteroaromatic olefins 7 and 8 both reacted more slowly than the corresponding phenylhexenes.[45] For 7 and 9 competing hydrogenation was extremely pronounced even at room temperature, resulting in hexyliothiophenes with only 10–30 % and

* That this labelling pattern reflects initial addition requires that the kinetic isotope effect (kH/kD) for β-elimination is of such a magnitude, that the possible β-Zr → α-Zr (or α → β) rearrangement does not lead to pronounced scrambling of D between these positions. For reactions in which other transition-metal alkyls undergo β-H elimination to give alkenes, kH/kD was found to be in the range 2–3, irrespective of whether the β-elimination was considered to be the rate-determining step, or not (Refs. 47 and 48). PhCH(D)CH₄[Zr] and PhCH[Zr][CH₃(D)] (obtained from styrene and [Zr]-D₃), without being in fast equilibrium with each other, both give the corresponding D₀, D₁, and D₃ analogues, i.e., PhCH₃(D)CH₃[Zr] (m = 0–2) and PhCH[Zr][CH₃(D)]ₑₙ (n = 0–3). This was not observed with 1-phenyl-1-propene or 1-phenyl-2-propene (Ref. 49). Nevertheless, the α-D: β-D ratio after deuterozirconation-hydrolysis of 4 should not be affected in any case.
5–10% deuterium, respectively. At higher temperature (45°C), larger amounts of hexyl thiophenes were formed, but with no measurable degree of deuteration. In contrast, 2-{(Z)-3-hexenyl}thiophene (8) can be converted into 2-hexylthiophene with good terminal selectivity [eqn. (11)].

\[
\text{OMe} \quad [Z]-H \quad 1) 2 \text{equiv. } [Z]-H \quad 48 \text{h, } 25-26^\circ C \quad \text{D}_2 \quad 54 \% \text{ yield} \quad (11)
\]

In order to determine the significance of the presence of sulfur, with respect to hydrogenation of hexyl thiophenes (hydrogenation did not depend on added thiophene during the hydrozirconnation of 4), we used olefin 11 as the substrate [eqn. (12)].52 and compared it with the methoxy analogue 6 which reacted without such pronounced reduction.

At 20–25°C the reaction of 11 was very sluggish. When the temperature was raised to 45°C, the reaction took a totally unexpected course [eqn. (12); sometimes other unknown products were also found]. The disulfide 14 was formed on exposure of the deuteriolysed reaction mixture to air. The extent of deuterium incorporation was low in products 12 and 13.

\[
\text{OMe} \quad \text{OMe} \quad [Z]-H \quad 1) 2 \text{ equiv. } [Z]-H \quad 24, 45^\circ C \quad \text{D}_2 \quad (12)
\]

After treatment of 12 [eqn. (12)] with Cp₂Zr(H)Cl (isolated), the aromatic compound was recovered unchanged. Moreover, the addition of thioanisole to solutions of preformed phenylethylzirconiums was not detrimental to the organometallics (based on deuterium content in ethylbenzene). The presence of sulfur in the vicinity of the double bond, as in 11, seems thus crucial, both to induce cleavage of the C–S bond and to hydrogenation.

\[
\begin{align*}
\text{D}_2 & \quad \text{D}_1 & \quad \text{D}_2 \\
22 & \quad 47 & \quad 31 \% (13)
\end{align*}
\]

For compounds 7, 9 and 11, results pointed towards the hydride Cp hydrogen of the reagent being involved in competing hydrogenation and/or H/D exchange.53,52

Next we turned to the aromatic olefin 15.54 It seemed logical that α-functionalised α-phenylpropanes would be obtainable from 15 [eqn. (13)]. The postulated intermediate (16) has shown to give mainly the terminal functionalised product.56 The results obtained with 15 were somewhat unexpected.54 Although propylbenzene was the major constituent of the reaction mixture, the α : ω ratio was higher [α : ω ≈ 4 : 6; eqn. (14)] compared with what had previously been found (α : ω ≈ 1 : 9; 25°C56), using 16 as the substrate. At 39–41°C the isomer ratio was altered slightly (α : ω ≈ 2–3 : 7–8).

\[
\begin{align*}
\text{OMe} & \quad [Z]-H \quad 1) 2 \text{ equiv. } \text{Cp}_2 \text{Zr} \text{HCl} (\alpha) \quad 24, 38-39^\circ C \quad \text{D}_2 \quad (14)
\end{align*}
\]

Some experimental evidence for Cp₂Zr(H)OMe [possibly formed via β-elimination of the ether functionality and subsequent reaction of Cp₂Zr(Cl)OMe with Cp₂Zr(H)Cl] being responsible for the high α : ω ratio [eqn. (14)] was obtained.54

Before leaving this section, we present the results obtained with cinnamyl alcohol (17).54 We reasoned that the relatively high γ-selectivity found using oleyl alcohol (Section 1), combined with the fact that benzyl zirconium derivatives are stable, should lead to the exclusive formation of 18 [eqn. (15)]. Dehydroxylation was unfortunately more pronounced [eqn. (16)] than in the hydrozirconnation of oleyl alcohol.

\[
\begin{align*}
\text{OH} & \quad \text{OH} \quad 1) [Z]-H \quad 2) \text{E}^\circ \quad \text{D} \quad (15)
\end{align*}
\]

\[
\begin{align*}
\text{OH} & \quad \text{OH} \quad 1) 4 \text{ equiv. } [Z]-H \quad 24, 42-45^\circ C \quad \text{D} \quad (16)
\end{align*}
\]

The lactone 19 was prepared in low yield by trapping the benzyl zirconium analogue of 18 with carbon monoxide–iodine.
In summary, the results discussed in this section show further limitations of the synthetic utility of hydrido-zirconation.

III. Experiments with modified zirconocene hydrides. In an effort to enhance the reactivity and improve the regioselectivity with respect to terminal alkylzirconiums, we wanted to make use of a few known zirconocene dihalides for the preparation of modified zirconocene hydrides. 55, 56

 Cp₂Zr(H)Br showed no advantages over the chloride analogue, and neither Cp₂Zr(H)F nor Cp₂⁺Zr(H)Cl (Cp⁺ = η⁵-C₅Me₅) could be obtained in the usual way. 55

"Cp⁺CpZr(H)Cl" (generated in situ) reacted much faster than Cp₂Zr(H)Cl with (Z)- or (E)-5-decene to give 1-deuterodecane after deuteriolysis. 55 The rate enhancement is presumably due to the formation of more soluble zirconium hydride(s). Greater solubility is also believed to cause faster reactions of (Me₅Cp)₂Zr(H)Cl, compared with Cp₂Zr(H)Cl, with 1-hexene or acetophenone. 57

Oleic acid oxazoline (I) (Section I) was converted quite rapidly into the stearil analogue using "Cp⁺CpZr(H)Cl" in toluene at 40°C. No starting material remained after 8 h and 80% of the saturated analogue, having a deuterium content of 89%, was isolated upon quenching (D₂O). 55 The shorter-chain oxazoline shown in eqn. (5) (Section I) was also treated with this modified reagent (~20 h, 40°C, dioxiane). 56 Regardless of the use of different solvents and reaction times, the experiments with these two oxazolines resulted in identical deuterium distribution, but unfortunately the terminal selectivity was even less pronounced than when Cp₂Zr(H)Cl was utilised.

As mentioned in Section II, difficulties were encountered with the aromatic olefin 4 when trying to obtain ω-alkylzirconiums. Even though rate enhancement was observed with (Me₅Cp)₂Zr(H)Cl, improvement of the ω-selectivity was not achieved. "Cp⁺CpZr(H)Cl", on the other hand, showed good terminal selectivity at 45°C [eqn. 17]. 56

IV. Reactions of alkenes with Cp₂Zr(H)Cl. General comments on hydridoconation-isomerisation. The process [eqn. (2)] has been suggested to proceed via insertion of the olefin into the Zr−H bond, followed by rapid eliminations and readditions. 7 The metal in Cp₂Zr(H)Cl formally has the d⁸-configuration, and thus the equilibrium in eqn. (18) is shifted to the right by the energy gained from formation of the new C−H bond. 8 If no stabilisation of the secondary ω-alkylzirconiums can be achieved (cf. previous sections), migration of the metal moiety occurs, most probably owing to the steric requirements of the Cp rings. The intrinsic rate of rearrangement to the terminal carbon has been suggested to be at least as fast as insertion of the starting alkene [(Z)-4-ocetene, benzene, room temp.] into the zirconium–hydrogen bond. 59 This assumption was based on the fact that only starting olefin and the final 1-alkyl compound could be observed when monitoring the reaction by 1H NMR spectroscopy. These observations imply that the metal never dissociates from the hydrocarbon chain, i.e., path (a) in Scheme I. Intramolecular rearrangement [path (a)] has also been indicated in cycloalkylboranes, but it competes with a mechanism corresponding to path (b). 60 In analogy with alkene complexes of d⁸-metals (insofar as they exist as intermediates), there should be no stabilising π-backbonding in the corresponding borane complexes.

In this section we will discuss our observations regarding reactions of alkenes with the zirconium hydride (deuteride), some of which have relevance to Scheme I.

After hydridoconation of long-chain (Z)-olefins, cis–trans isomerisation was often observed. 29, 39, 60 The formation of positional isomers of the starting material with the double bond only one or two carbons away from the original position, was indicated by mass spectral analysis, even after quite a long reaction time. 59 Others too, have found isomerised alkenes using long-chain substrates. 46 In all these cases, reactions were conducted at slightly elevated temperatures (40–50°C) with an excess of reagent, as this gave a faster consumption of olefin. As mentioned above, in a previous study 29 no isomers of the starting alkene were observed during rearrangement.

For a closer look at the process we initially used (Z)-5-decene as a model compound. After reaction in toluene or benzene for 4 h at 40°C, the yield of decane never exceeded 70%. The remaining material consisted of isomeric decenes. The rate of alkane formation was considerably higher in THF. Results of the experiments using toluene as the solvent are summarised in eqn. (19). 51
yield of decane (11% D₁; 4% D₂) was significantly lowered using a less than stoichiometric amount of the metal hydride.

Neither 1-decene nor internally deuteriated decane were detected. The isomeric alkenes formed do not stem from the terminal σ-decyzirconium intermediate (see below), and thus it is obvious that the formation of α-alkylmetal compound in this case is quite slow.

An interesting point is that only 21% of (Z)-5-decene remained using 0.5 equivalents of [Zr]−H. A possible reason could be a competing displacement which has been proposed for alkenes and alkynes [eqn. (20)]. We conclude, however, that this is not an important side-reaction during hydrozirconation [eqn. (19)]; otherwise (Z)-5-decene would not be unchanged, as it actually is (≥97%), after treatment with about one equivalent of the 1-declyzirconium shown in eqn. (21).61

Displacement [eqn. (20)] should result in formation of isomeric decenes from the postulated 5-decylmetal intermediate [compare ordinary hydrozirconation, eqn. (19)]. As can also be seen from eqn. (21), the formation of alkenes from the primary alkylzirconium, via elimination of [Zr]−H, is unimportant (compare Refs. 7 and 8).

The hydrozirconation of (Z)- and (E)-5-decene (benzene, 40°C, 4 h) gave decane with the isotope composition given in eqn. (22).53 The deuterium composition in the remaining olefinic mixture varied in the isomers.

Before making any conclusive remarks, we will present a few results concerning alkenes which can give only secondary alkylmetals.61 Firstly, alkene 20 was hydrozirconated [eqn. (23)]. The formation of 21 was certainly not favoured, independent of conditions. When 20 was treated with Cp₂Zr(DCl), deuterium was found at several carbons in the alkenes recovered after hydrolisis. Secondly, the reactions of cycloalkenes 22 and 23 were studied. Cyclooctene (22) has, like 1-methyl-1-cyclohexene (24), been reported not to give any alkyl compounds in hydrozirconation-functionalisation attempts.44 After deuteriozirconation we recovered 80–100% of these substrates (Table 1). Even if the cycloalkenes 22 and 23 were not consumed, they were deuteriated with total scrambling. The amount of deuterium incorporated was roughly the same as the amount of [Zr]−D used. One might argue that the reason why cyclooctyl- and cycloheptyl-zirconiums are thermodynamically unstable, in contrast with cyclopentyl44 and cyclohexyl65,66 could be transannular interactions in the former intermediates.

Alkene 24 was very unreactive. No evidence for deuterium incorporation in the recovered alkene (80–90%) was obtained.

As would of course be expected, [Zr]−D ([Zr]−H) adds to alkenes 20, 22 and 23 [eqn. (23), Table 1], even if no saturated analogues, or only very small amounts,
<table>
<thead>
<tr>
<th>Deuterorizonated compound</th>
<th>Recovered olefin (%)</th>
<th>Isotope composition (%)</th>
</tr>
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<tbody>
<tr>
<td></td>
<td></td>
<td>$D_0$</td>
</tr>
<tr>
<td>20</td>
<td>97</td>
<td>≈80</td>
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<td></td>
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<td>100</td>
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<tr>
<td>22</td>
<td>90–95</td>
<td>34</td>
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<td>23</td>
<td>80–90</td>
<td>100</td>
</tr>
<tr>
<td>24</td>
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*The cycloolefins were reacted with ≈1 equiv. of $\text{C}_{2}p_{2}\text{Zr}(\text{D})\text{Cl}$ (isolated) for 22–23 h at 40°C using 1,4-dioxane as the solvent. Alkene 20 was deuterorizonated [1.0–1.5 equiv. of isolated $\text{C}_{2}p_{2}\text{Zr}(\text{D})\text{Cl}$] at the same temperature for 4–5 h in toluene. With a longer reaction time (20 h) the $D_1$ and $D_2$ fractions increased considerably, corresponding to 40–50%. Analyses were made after quenching with water. *Mixture of isomers.

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References

   2310.
27. Fryzuk, M. D., Bates, G. S. and Stone, C. J. Org. Chem. 53
   379.
30. Buchwald, S. L., Nielsen, R. B. and Dewan, J. C. Organome-
31. Lappert, M. F., Luong-Thi, N. T. and Milne, C. R. C.
    5 (1986) 1585.
35. Alvähll, J., Gronowitz, S. and Hallberg, A. Chem. Soc. 25
    (1985) 393.
36. Alvähll, J., Gronowitz, S., Hallberg, A. and Svensson, R.
38. Choukroun, R., Gervais, D. and Raoult, Y. Polyhedron 8
    (1989) 1758. Also see Ref. 61 and literature cited there.
41. Erker, G., Schlund, R., Albrecht, M. and Sarter, C.
48. Albismendi, G., Scolaro, L. M., Minitti, D. and Romeo, R.
    Inorg. Chem. 29 (1990) 3467 and literature cited there.
49. Nelson, J. E., Bercaw, J. E. and Labinger, J. A. Organome-
50. See for example, Gronowitz, S., Ed., Heterocyclic Com-
    pounds. Thiophene and its Derivatives, Wiley-Interscience,
    14 (1978) 353.
53. Annby, U. Studies of the Hydrozirconation–Isomerisation
    Chem. 403 (1991) 133.
55. Alvähll, J., Gronowitz, S. and Hallberg, A. Chem. Scr. 28
    (1988) 285. References concerning the preparation of the
    zirconocene dihalides, used as precursors, are found in this
    paper.
    Scand. 44 (1990) 862.
    (1989) 2349. Some references regarding other modified zir-
    concene hydrides are given here (also see Ref. 8, p. 676).
    1729.
    8115.
    6125 and references cited there.
61. Annby, U., Alvähll, J., Gronowitz, S. and Hallberg, A.
62. Swanson, D. R., Nguyen, T., Noda, Y. and Negishi, E.
64. Hart, D. W. Transition Metal Hydrides in Organic Synthesis,
    dissertation, Princeton University 1975, Xerox University
    Microfilms, Ann Arbor, Michigan 48 106.
    Chem. 43 (1972) C 32.
66. Gell, K. I., Posin, B., Schwartz, J. and Williams, G. M.
68. The 1H NMR analyses of the zirconium hydride were made
    according to Carr, D. B. and Schwartz, J. J. Am. Chem.
    Soc. 101 (1979) 3521.

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