A well-defined model system for the chromium-catalyzed selective oligomerization of ethylene†

Wesley H. Monillas, John F. Young, Glenn P. A. Yap and Klaus H. Theopold*

The chromium(I) dinitrogen complex [(i-Pr2Ph)2nacnacCr]2(μ–η2:η2-N2) catalyzes the selective trimerization of ethylene to 1-hexene at ambient pressure and temperature, and in the absence of any cocatalyst. After the conversion of the substrate, the catalyst cleanly converts to another chromium(I) species, namely [(i-Pr2Ph)2nacnacCr]2(μ–η2:η2-C2H4), which is not catalytically active. Binuclear metallacycles containing Cr(II) have been prepared as candidates for catalytically active intermediates; however they are not kinetically competent to explain the catalysis. Turning thus to mononuclear metallacycles featuring Cr(III), a chromacyclopentane, chromacyclopentene and chromacyclopentadiene have been prepared as models of catalytic intermediates. Of these, the latter also catalyzes the trimerization of ethylene. These results support the proposal that selective ethylene oligomerization catalysis involves an interplay between Cr(I) ethylene complexes and mononuclear Cr(III) metallacycles.

Introduction

α-Olefins are valuable precursors for a wide range of chemical processes.1 Owing to the variety of their applications, selective syntheses, e.g. of 1-hexene or 1-octene, are inherently preferable to the statistical (Schulz–Flory) mixtures resulting from the widely practiced ethylene oligomerization reactions based on the Cossee mechanism.2 Originally discovered in the ’60s,3,4 there has been a recent flurry of interest in the selective trimerization of ethylene to 1-hexene, predominantly catalyzed by chromium complexes.5–40 While the differential reactivity of metallacycles of different ring sizes is widely held to be the cause of this unusual selectivity, many questions about the mechanistic details remain; among these are the formal oxidation states and charges of the intermediates, the role of mononuclear versus binuclear catalysts, and the exact nature of the product forming step(s). Most relevant catalysts require activation by cocatalysts (e.g. MAO), which complicate spectroscopic detection or isolation of intermediates.41 There exists thus a need for well-defined compounds that catalyze ethylene trimerization in isolation. This contribution describes such a system.

Results and discussion

Catalysis

Some time ago we reported the synthesis and characterization of several compounds featuring chromium in the rare formal oxidation state +I;42,43 the most reactive of these is the side-on dinitrogen complex [(i-Pr2Ph)nacnacCr]2(μ–η2:η2-N2) (1, (i-Pr2Ph)nacnac = 2,4-pentane-N,N′-bis(2,6-diisopropylphenyl) diketiminate).44 1 reacts cleanly with ethylene to eventually yield [(i-Pr2Ph)nacnacCr]2(μ–η2:η2-C2H4) (2) as the sole organometallic product;43 however, before the latter is formed all available ethylene is oligomerized to 1-hexene and smaller amounts of 1-butene. Thus, structurally characterized 1 is a catalyst – or a catalyst precursor – for the selective trimerization of ethylene, which does not require a cocatalyst (see Scheme 1).

Monitoring (by 1H NMR) the reaction of 1 with an excess of ethylene at ambient temperature in a sealed NMR tube revealed the gradual replacement of the broad and
Table 1 Ethylene trimerization catalyzed by 1

| Run  | Catalyst (mmol) | Solvent (mL) | C2H4 pressure (psi) | Temperature (°C) | Turnover (mmol C6/ mmol cat/h) |
|------|----------------|--------------|---------------------|------------------|-------------------------------|
| 1    | 0.031          | 10           | 15                  | 23               | 0.76                          |
| 2    | 0.031          | 10           | 15                  | 23               | 5.7                           |
| 3    | 0.031          | 10           | 50                  | 23               | 11.4                          |
| 4    | 0.031          | 10           | 60                  | 23               | 26.9                          |
| 5    | 0.093          | 30           | 215                 | 23               | 172.5                         |
| 6    | 0.031          | 10           | 15                  | 40               | 1.2                           |
| 7    | 0.031          | 10           | 15                  | 40               | 8.4                           |

* Description of typical method can be found in the Experimental section.

Turnover for reactions at 40 °C calculated before catalyst poisoning.

Attempts to isolate the spectroscopically observed intermediate formed during the catalytic reaction – e.g. by cooling solutions under ethylene – were not successful, nor were we able to trap this species by the addition of potential ligands (e.g. pyridine) to catalytically active solutions. However, borrowing an experiment used by Mashima et al., 46 we found that addition of a methylene chloride solution of bromine (Br2) to a reaction in progress produced [(i-Pr3Ph)2nacnacCr(μ-Br)2] and 1,4-dibromobutane (identified by retention time in the GC and by GCMS). The formation of the latter at a minimum suggests that the intermediate results from coupling of two ethylene units; it is most likely a metallacycle of some type.

These observations are consistent with the commonly suggested mechanism for the selective trimerization of ethylene by transition metal catalysts (see Scheme 2). According to this proposal, 47 coordination of two molecules of ethylene to each chromium(II), followed by reductive coupling, forms a mononuclear metallacyclopentane containing Cr(II). The latter is a candidate for the yet unidentified species spectroscopically observed during turnover. Insertion of another ethylene into the relatively stable chromacyclopentane yields the homologous chromacycloheptane. Due to its greater conformational flexibility, β-hydrogen elimination and reductive elimination of the resulting alkenyl hydride are kinetically favored over further ring expansion, thus explaining the selectivity for 1-hexene.

The binuclear nature of both 1 and 2 raised the question whether a bimetallic mechanism might be operative. Among the various lines of evidence arguing against such a pathway we mention but two at this juncture. First, the lack of reactivity of 2 with added ethylene would seem to be an obstacle to such a mechanism. Furthermore, we have shown that binuclear chromium alkyl/aryl hydrides, such as might be generated by such a mechanism, are resistant to reductive elimination, 42, 46 militating against such a transformation as the product forming step of the catalytic cycle. However, a concerted 1–5 hydrogen shift has been discussed as an alternative chain transfer step.

Synthesis of potential bimetallic intermediates

Crucial intermediates in a bimetallic reaction mechanism would be dinuclear metallacycles in which carbon chains of various lengths (C4, C6, C8) bridge a dichromium fragment.
Accordingly, we set out to prepare compounds of this type.\textsuperscript{19–51} Alkylation of [[i-Pr\textsubscript{2}Ph\textsubscript{2}nacnacCr(μ-I)]\textsubscript{2}], with 1,6-dilithioalkanes indeed yielded such compounds, after some optimization (see Scheme 3).

The required dilithioalkyl reagents were synthesized using a procedure reported by Negishi.\textsuperscript{52} For example, Li(CH\textsubscript{2})\textsubscript{4}Li was prepared by lithium/halide exchange between 1,4-diodobutane and 4.2 equivalents of t-butyllithium. Although the slight excess of t-butyllithium ensured full conversion, upon quenching with diethyl ether solvent at room temperature, lithium ethoxide (LiOEt) and ethylene are formed.\textsuperscript{53} The LiOEt side product yielded an undesirable chromium ethoxide complex that was observed crystallographically in our initial attempts to alkylate [[i-Pr\textsubscript{2}Ph\textsubscript{2}nacnacCr(μ-I)], with Li(CH\textsubscript{2})\textsubscript{4}Li. Therefore, the lithiation procedure was modified to eliminate the formation of lithium ethoxide. The lithium/iodide exchange reaction was performed at −78 °C and the diethyl ether solvent was removed under vacuum while the solution warmed from −78 °C to 0 °C. The crude product was extracted with pentane and filtered under nitrogen to remove any traces of LiOEt.\textsuperscript{54} Removal of the pentane solvent under vacuum yielded a gummy white precipitate which was dissolved and stored at −30 °C in Et\textsubscript{2}O as a clear yellow solution. The use of the dilithioalkyls generated by this procedure did not produce any chromium complexes containing ethoxide moieties. However, several attempts to synthesize 1,6-dilithiohexane by this route were unsuccessful, presumably due to formation of cyclohexane or polymerization;\textsuperscript{55} accordingly the synthesis of a bimetallacyclooctane complex was eventually abandoned. Reaction of the [[i-Pr\textsubscript{2}Ph\textsubscript{2}nacnacCr(μ-Cl)]\textsubscript{2}]\textsuperscript{56} with Li(CH\textsubscript{2})\textsubscript{4}Li at a variety of temperatures (−78 °C to RT) was always incomplete, giving mixtures of the starting material and the desired bimetallacycle. It was then found that addition of Li(CH\textsubscript{2})\textsubscript{4}Li to iodide [[i-Pr\textsubscript{2}Ph\textsubscript{2}nacnacCr(μ-I)]\textsubscript{2}] at room temperature resulted in complete conversion to the desired product, namely [[i-Pr\textsubscript{2}Ph\textsubscript{2}nacnacCr(μ-CH\textsubscript{2})\textsubscript{2}(CH\textsubscript{2})\textsubscript{2}] (3). The molecular structure of 3 was determined by X-ray diffraction and the result is shown in Fig. 1. The homologous bimetallacycloheptane [[i-Pr\textsubscript{2}Ph\textsubscript{2}nacnacCr(μ-CH\textsubscript{2})\textsubscript{2}(CH\textsubscript{2})\textsubscript{3}] (4) was synthesized in analogous fashion via alkylation of the iodide precursor with 1,5-dilithiopentane (Li(CH\textsubscript{2})\textsubscript{4}Li) at room temperature. The structure of 4, as determined by X-ray diffraction, is depicted in Fig. 2.

The C\textsubscript{3} bimetallacycle 3 crystallized in the monoclinic space group P2\textsubscript{1} with a co-crystallized molecule of hexamethyldisiloxane (solvent of recrystallization). C\textsubscript{4} bimetallacycle 4 crystallized in the triclinic space group P\textsubscript{1} with a co-crystallized molecule of pentane. The molecular structures of 3 and 4 are very similar, except for the incrementally larger ring size of 4.
Both feature terminal methylene groups that bridge the two chromium atoms; in that sense they may be described as a dichromabicyclo[1,1.2]hexane (3) and dichromabicyclo[1.1.3]heptane (4). Extreme electron deficiency – discounting metal–metal bonding, the individual chromium atoms feature 10-valence electron configurations – drive the formation of 3-center/2-electron M–C–M bonds. This structural feature has ample precedent in Cr(u) chemistry; for example, we had earlier prepared compounds of the type [Cp*Cr(μ-R)]2, which show strong similarities in gross structure, metal–metal distances, and magnetism.57 The local coordination environments around the Cr atoms in 3 and 4 deviate from the preferred square planar geometry for four coordinate Cr(u) complexes.58 The Cr2C2-cores of the molecules adopt a butterfly shape and the relevant Cr–C distances exhibit a short/long pattern, rendering this fragment somewhat unsymmetric. The non-coordinated carbon atoms (i.e., C(2), and C(3) of 3 and C(2), C(3), and C(4) of 4) exhibit positional disorder; thus their C–C single bonds appear unrealistically short.59 The Cr–Cr distances in 3 and 4 are 2.5451(15) and 2.5567(8) Å, respectively, and are therefore within the range of weak Cr–Cr bonding. Both compounds have identical e

Reactivity with ethylene

It is difficult to imagine how any bimetallic mechanism for the oligomerization of ethylene catalysed by 1 could circumvent Cr(u) as an intermediate, and 4 is surely representative of the kind of bimetallacycle involved in such a putative mechanism. Accordingly, their reactivity with ethylene and kinetic competency vis-à-vis the catalysis described above is of interest. Preliminary 1H-NMR spectroscopic studies of the reaction of 3 with 1 atm of ethylene in C6D6 indicated the presence of Cr(u) metal–metal bonding, or a superposition of both.61

Entry Dimer Temp (°C) Time (h) Pressure (psi)

| Yield (%) | C4 | C6 | C8 |
|---|---|---|---|
| 4.43 | 11.2 | 0 |
| 16.55 | 36.2 | 8.62 |
| 14.32 | 5.6 | 0 |
| 7.57 | 63.1 | 5.21 |
| 2.10 | 7.55 | 0 |
| 4.35 | 15.9 | 0 |

5a % yield of Cs was less than 1%. Cs was not detected.
undergoing reductive elimination of 1-butene and yielding two Cr(i) fragments. However, analysis by $^1$H-NMR and GC/MS did not show any butadiene; only butene was detected by GC/MS. Since no free butadiene was found, despite the formation of 5, it was hypothesized that another paramagnetic compound formed during the decomposition reaction might be a butadiene complex. In separate $^1$H-NMR experiments in C$_6$D$_6$ or d$_8$-THF, 5 was exposed to 1 atm of butadiene and left to react overnight at room temperature. 5 completely disappeared and resonances for butene and a paramagnetic compound that exactly matched the unassigned $^1$H-NMR resonances noted during the decomposition of 3 were observed (Scheme 4).

In hopes of determining if this new paramagnetic compound was indeed a butadiene complex, 1 was exposed to 1 atm of butadiene in THF; this reaction indeed yielded a monomeric Cr butadiene adduct, namely (i-Pr$_2$Ph)$_2$naacnacCr(1,3-C$_2$H$_4$)(THF) (6). The molecular structure of 6 was determined by X-ray diffraction; its molecular structure is shown in Fig. 3. 6 exhibits square pyramidal geometry and contains a cis-$\eta^1$-bonded butadiene molecule and a coordinated molecule of THF in the apical position (the basal plane is defined by the naacnac ligand and the butadiene). The bond distances of the butadiene’s erstwhile double bonds (C1–C2 and C1A–C2A, 1.378(4) Å) are identical to the formal single bond distance (C2–C2A, 1.379(5)), indicating a modest transfer of electron density to the diene, which is consistent with other reported Cr(η$^1$-1,3-butadiene) complexes. However, the relative Cr–C distances still favour a description as a z-complex rather than as the valence tautomeric metallacyclopentene. The room temperature effective magnetic moment of 6, $\mu_{\text{eff}}$(295) = 3.8(1)$\mu_B$, is consistent with a quartet ($S = 3/2$) ground state. The chromium in 6 has a formal oxidation state of +1, giving it a d$^0$ electronic configuration; 6 apparently adopts an intermediate spin state.

We note that 6 also provides a model for the bonding of two molecules of ethylene to Cr(i); it may be considered an analogue of the proposed L$_2$Cr(C$_2$H$_4$)$_2$ intermediate of ethylene trimerization proceeding by a mononuclear metallacycle mechanism (see Scheme 1). The notion that 3 decomposes to form a butadiene complex was confirmed, since the $^1$H-NMR spectrum of 6 matched the unassigned paramagnetic resonances observed in the decomposition of 3. To summarize, 3 decomposes to equal parts of 5 and 6. This result implies that there are multiple pathways to generate Cr(i) fragments, which may initiate oligomerization catalysis. The observation of butene during thermolysis of 3 is also consistent with the detection of pentene during the reaction of 4 with ethylene, and it suggests that decomposition of these bimetallacycles can lead to catalytically active Cr(i) fragments.

Finally, in a separate $^1$H-NMR experiment in THF-d$_8$, 6 was exposed to 1 atm of ethylene, whereupon it produced 1-hexene and 1-butene (confirmed by GC/MS analysis). Mechanistically, dissociation of butadiene followed by the binding of ethylene would lead to the formation of 1-hexene and then regenerate a Cr(i) fragment that can either continue in trimerization or rebind butadiene. These results suggest that labile monomeric Cr(i) compounds can selectively oligomerize ethylene and it provides some support for a monomeric metallacycle mechanism for ethylene trimerization. This notion will be further explored in the next section.

**Synthesis of mononuclear model compounds**

As the binuclear hypothesis based on Cr(i) olefin complexes and Cr(II) bimetallacycles became discredited [see above], a premium was placed on preparing possible intermediates of a mononuclear mechanism, which would presumably involve Cr(n) metallacycles. Notably, we tried to devise synthetic routes to a mononuclear chromacyclobutane, which might be a credible candidate for the active intermediate observed during catalysis. A severe obstacle to the synthesis of e.g. (i-Pr$_2$Ph)$_2$naacnacCr(C$_2$H$_4$)$_2$ was presented by our finding that the (i-Pr$_2$Ph)$_2$naacnacCr-fragment does not readily support the preparation of trivalent chromium dialkyls of the type (i-Pr$_2$Ph)$_2$naacnacCrR$_2$. Indeed, whereas treatment of (Me$_2$Ph)$_2$naacnac-Cr$^{III}$Cl$_2$(THF)$_2$ with lithium alkyls produces Cr(n) dialkyls, similar treatment of (i-Pr$_2$Ph)$_2$naacnacCr$^{III}$Cl$_2$(THF)$_2$
[(i-Pr₂Ph)₂nacnacrIIICl(μ-Cl)]₂ with such reagents invariably leads to reduction and formation of Cr(II) alkyls of the type (i-Pr₂Ph)₂nacnacrR (R = CH₂SiMe₃) or [(i-Pr₂Ph)₂nacnacr-μ-R]₂ (R = Me). In view of this dichotomy, we chose to initially attempt the synthesis (Me₂Ph)₂nacnacr(CH₂)₄. However, as detailed elsewhere, reaction of (Me₂Ph)₂nacnacrIIICl(THF)₃ with Li(CH₂)₄Li actually yielded the bimetallic molecule structure of (Me₂Ph)₂nacnacr(CH₂)₄(py) (an analogue of 3), presumably due to the greater reducing power of the dilithio reagent. To overcome this problem, we decided to make the chromium(III) ion in the halide precursor more electron rich by substituting it with stronger σ-donors. Gratifyingly, this was met with success. Treatment of (Me₂Ph)₂nacnacrIIICl(THF)₃ with Li(CH₂)₄Li afforded the mononuclear chromacyclobutane (Me₂Ph)₂nacnacr(CH₂)₄(py) (7). The molecular structure of 7 was determined by X-ray diffraction, and the result is shown in Fig. 4.

7 is a rare example of a mononuclear chromacyclopentane. It differs from the putative catalytic intermediate only by virtue of the coordinated pyridine. However, the latter apparently makes all the difference. 7 does not catalyze the trimerization of ethylene, nor does it decompose readily upon heating. We believe that the strongly coordinated pyridine ligand prevents binding and insertion of ethylene as well as decomposition by β-elimination. Attempts to remove the pyridine by adding various Lewis acids (AlMe₃, B(C₆F₅)₃, BF₃) did not yield tractable products, nor did these additives facilitate reactions of 7 with ethylene. In summary, 7 may be a suggestive structural model for a catalytic intermediate, but the strong coordination by pyridine blocks further reactivity (such as binding and insertion of ethylene or even β-hydrogen elimination). Its ancillary ligand also differs from the one which gives rise to catalysis. Accordingly, 7 does not by itself provide convincing evidence for the catalytic activity of mononuclear metallacycles.

We then turned our attention to the preparation of compounds that might serve as models for some of the species proposed to constitute the catalytic cycle (Scheme 1), while employing the same ancillary ligand as the actual catalytic system. First up, we wondered about an analogue of the mononuclear ethylene complex (i-Pr₂Ph)₂nacnacr(C₂H₄), which may enjoy a fleeting existence after the elimination/substitution of the α-olefin product. In the presence of excess ethylene, it would be expected to form (i-Pr₂Ph)₂nacnacr(C₂H₄)₂ to begin another turnover, while lack of substrate presumably leads to irreversible formation of 2. We hypothesized that a stericly more demanding σ-acceptor might stabilize such a 1 : 1 complex, and indeed the reaction of 1 with one equivalent of diphenylacetylenyl chloride produced the mononuclear alkyne adduct (i-Pr₂Ph)₂nacnacr(η¹-C₂H₂) (8). The molecular structure of 8 as determined by X-ray diffraction is shown in Fig. 5. The diphenylacetylene fragment is bound nearly perpendicularly to the plane of the ligand. Both C₁ and C₂ were found to be located at similar distances from the chromium at distances of 1.957(4) Å and 1.963(4) Å, respectively, with a Cr-centroid distance of 1.846(6) Å. The C-C distance of the diphenylacetylene was found to have been elongated by 0.115 Å from free diphenylacetylene to a distance of 1.320(5) Å. The latter distance is the strikingly similar to the C=C double bond distance in stilbene, which measures 1.321(5) Å. The diphenylacetylene fragment in 8 is no longer linear but features an average C-C-Ciso angle of 137.3(4)°. This value is within the range of other structurally characterized chromium η²-diphenylacetylene complexes, which possess a C-C-Ciso angles in the range of 136.3–155.9°. These structural changes in the diphenylacetylene fragment, and the fact that 8 has an effective magnetic moment of 3.9(1)μ₅B, suggests that the chromium center of 8 is in the +III oxidation state, with corresponding twofold reduction of the diphenylacetylene ligand, resulting in a more appropriate description as a metalla-cyclopropene.

If 8 is to be a model of a 1 : 1 ethylene complex, it might be expected to rapidly react with additional ethylene. We already

![Fig. 4](image1.png)  
**Fig. 4** The molecular structure of (Me₂Ph)₂nacnacr(CH₂)₄(py) (7); hydrogen atoms, except those on the alkyl chain have been removed for clarity. Selected interatomic distances [Å] and angles [°]: Cr–C1 2.076(5), Cr–C4 2.120(5), Cr–N1 2.048(4), Cr–N2 2.126(4), Cr–N3 2.138(4), C1–Cr–C4 83.2(2), N1–Cr–N2, 88.99(16).

![Fig. 5](image2.png)  
**Fig. 5** The molecular structure of (i-Pr₂Ph)₂nacnacr(η¹-C₂H₂) (8); hydrogens have been removed for clarity. Selected interatomic distances [Å] and angles [°]: C1–C1, 1.980(2); C1–C7, 1.944(4); C1–C8, 1.970(4); C7–C8, 1.307(5); N1–C1–N1A, 90.78(14); N1–Cr–C7, 130.52(8); N1–Cr–C8, 132.17(8).
have a model of sorts for the ensuing bis(ethylene) complex, in the form of butadiene complex 6, but the covalent linkage of the two olefins in butadiene prevents it from modelling the subsequent reductive coupling step. On this score, 8 delivers. Monitoring the addition of ethylene to a THF-d$_8$ solution of 8 by $^1$H NMR spectroscopy showed formation of a new paramagnetic chromium compound, but no formation of hexene or any other olefin. Isolation and determination of the structure of the new compound (see Fig. 6) showed it to be the five-membered metallacycle (i-Pr$_2$Ph)$_2$NacnacCr(CPh)$_2$(CH$_2$)$_2$ (9), which can the thought of as resulting from the insertion of ethylene into one of the Cr–C bonds of the metallacyclopentene 8, or alternatively as the product of a reductive coupling of a coordinated alkyne and alkene. 9 adopts a square pyramidal geometry with a THF molecule occupying one of the basal sites. The C3–C4 distance of 1.354(4) Å is consistent with a C=C double bond. The C1–C2 distance (1.526(4) Å) and C2–C3 distance (1.508(4) Å) on the other hand are characteristic of C=C-single bonds, consistent with the description of 9 as a metallacyclopentene. In addition, the hydrogen atoms on C1 and C2 can be located in the difference map and their positions refined.

Although the THF molecule coordinated to chromium in 9 might be expected to be substitutionally labile, the compound does not react further with additional ethylene. In particular, it does not catalyse the oligomerization of ethylene in THF (as 1 does). We note the possibility that the large steric demand of the phenyl substituents is responsible for this reluctance, effectively blocking the coordination of additional ethylene. On the other hand, 9 is already the second mononuclear chromacycle (besides 7) that does not produce any higher olefins under ethylene. Fortunately, the situation is about to improve.

Diphenylacetylene is a rather ‘large’ stand-in for ethylene by any reckoning, which in turn led us to exploring smaller alkenes. Accordingly, exposure of a THF solution of 1 to 2-butyne resulted in the rapid formation of a white precipitate. Permitting the reaction to continue yielded a large amount of the white precipitate and effected a slow color change of the solution from brown to green-grown. $^{13}$C NMR spectroscopy proved that the white precipitate was hexamethylbenzene, i.e. the product of a catalytic cyclotrimerization of the alkyne.$^{72,73}$ Removal of the organic product by filtration and standard work-up of the filtrate afforded the five-membered metallacyclic (i-Pr$_2$Ph)$_2$NacnacCr(n$_2$C$_6$Me$_6$)(THF) (10). The molecular structure of 10 has been determined by X-ray diffraction, and is shown in Fig. 7.

The compound is a metallacyclopentadiene (or ‘metallole’), which presumably results from reductive coupling of two alkenes bound to chromium. The compound possesses square pyramidal geometry with one terminus of the butadienediy ligand in the axial position. As expected, the coordinated C$_2$Me$_4$-fragment is planar and exhibits clear bond length alternations, as C1–C2 and C3–C4 were found to be 1.359(4) Å and 1.347(4) Å, respectively, while C2–C3 was found to be 1.485(4) Å. The effective magnetic moment of 10 is 3.8(1)$\mu_B$, consistent with chromium in the +III oxidation state. Compound 10 is the resting state of the cyclotrimerization catalyst, as its exposure to 2-butyne reinitiates the catalysis and yields more hexamethylbenzene.

More to the point, 10 is another mononuclear chromacycle, which (unlike 9) is not encumbered by excessive steric hindrance, and the coordinated THF of which is apparently subject to displacement by additional substrate. Therefore it might be expected to react with ethylene as well. Gratifyingly then, 10 was indeed found to be a catalyst for the selective oligomerization of ethylene. Exposure of a THF-d$_8$ solution of 10 to ethylene yielded 1-hexene, ethylene complex 2, and an unidentified organic compound (presumably 1,2,3,4-tetramethylhexatriene).
Conclusions

We have shown that a well-characterized chromium(1) dinitrogen complex (i.e., 1) catalyzes the selective trimerization of ethylene to 1-hexene in homogeneous THF solution, albeit slowly. To our knowledge, this is the only example of a catalyst for the trimerization of ethylene that unambiguously starts and finishes in the formal oxidation state Cr(I). Since the catalyst functions in the absence of any cocatalysts, which often obscure assignments of structure and oxidation state, this system presents an excellent chance to address fundamental questions regarding a commercially important and mechanistically intriguing catalytic cycle. For example, we have prepared relevant binuclear Cr(II) metallacycles and shown that they are not intermediates in the catalysis. By extrapolation, and absent any unambiguous evidence for a role of binuclear sites in this type of catalysis, we suggest that the alternative mononuclear Cr(III) metallacycles are most likely intermediates in selective oligomerization catalysis. While the independent synthesis of the most likely intermediate in our system – i.e., (i-Pr2Ph)2nacnacCr(CH2)4(THF) – proved elusive, we have prepared several closely related analogues of it. Of these, two proved inactive, but for cogent reasons. However, the mononuclear chromacyclopentadiene (i-Pr2Ph)2nacnacCr(η2-C6Me6)(THF) (10) reacted with ethylene to catalytically produce 1-hexene and enter the same reaction channel as 1. To the extent that these results may prove general, they strongly support the notion that chromium catalysts for selective ethylene oligomerization alternate between mononuclear Cr(I) ethylene complexes and Cr(III) metallacycles. Our results also show that neutral compounds are competent catalysts for the selective oligomerization. However, the activity of this catalyst is rather low by comparison, and we surmise that cationic, more electrophilic intermediates are the mainstays of more active ones.

Experimental

General considerations

All manipulations of compounds were carried out using standard Schlenk, high vacuum line, and glovebox techniques under an atmosphere of N2. Solvents were purchased from Fisher Scientific, degassed, and dried by passing through activated alumina. THF-d8 and C6D6 were purchased from Cambridge Isotopes Laboratory and stored under vacuum over Na/K alloy. All other reagents were purchased from Aldrich or Acrros and dried as appropriate using standard procedures as needed. 23H NMR spectra were taken on a Bruker DRX-400 spectrometer and were referenced to the residual protons of the solvent. FTIR spectra were taken on a Mattson Alpha Centauri or Mattson Genesis Series spectrometers. UV/vis spectra were recorded using a HP 8453 spectrophotometer. Mass spectra were obtained by the University of Delaware Mass Spectrometry Facility. Elemental analyses were performed by Desert Analytics. Room-temperature molar magnetic susceptibilities (μm) in the solid state were determined using a Johnson Matthey magnetic susceptibility balance. They were corrected for diamagnetism using Pascal constants and converted into effective magnetic moments (μeff).

GC/MS measurements for oligomerizations were taken on a HP G1800A GCD. To determine the response factor for the GC 1 mmol of 1-pentene, 1-hexene, 1-heptene, 1-octene, and nonane were placed in a 50 mL volumetric flask and then filled to 50 mL with THF. The flask was shaken and 1 μL was injected into the GC, the average area for three runs was calculated and the ratio of the GC/MS response to the α-olefin to nonane was calculated and plotted versus carbon number. The slope of the linear trendline allowed for the calculation of the RF for the 1-hexene; the RF for nonane was arbitrarily set to 1.0. The raw integrals for the 1-hexene analyses were divided by the RF to obtain the corrected integrals.

Typical oligomerization procedure for reactions <90 psi

A thick walled reactor was heated to 250 °C overnight to remove all traces of moisture and fitted with a cap consisting of an inlet port, gauge, and connections for vacuum and ethylene. The reactor was evacuated and backfilled with ethylene three times. The reactor was charged with 1 atm of ethylene. In the nitrogen dry box the requisite amount of catalyst was dissolved in 10 mL THF; nonane was added as the standard. A syringe containing the aforementioned mixture was removed from the dry-box and the solution mixture was injected into the reactor. The reactor pressure was raised to the necessary pressure and the reaction mixture was stirred. At various times throughout the oligomerization an aliquot was removed from the reactor and injected in the GC.

Typical oligomerization procedure for reaction at 215 psi

In the dry-box, 90 mg (93 mmol) of catalyst was added to a 300 mL Parr reactor followed by 90 mL THF and the nonane standard. Pentane was also added as an additional check. The reactor assembly was brought of the dry-box and the connection line was purged with ethylene for 2 minutes. The reaction was opened to ethylene and the pressure was regulated to 215 psi and stirred for certain time periods. When the set time was reached the ethylene pressure was released, the solution is exposed to air and an aliquot injected into the GC. This procedure was repeated for each time frame.

Typical oligomerization procedure for the reactions of 3 and 4 with ethylene

In a nitrogen-filled dry box the bimetallicacycle was dissolved in 10 mL THF and nonane was added via a micropipette to a glass ampoule. The ampule was removed from the nitrogen dry box and placed onto a high vacuum line. After three freeze–pump–thaw cycles 1 atm of ethylene was backfilled into the vessel. After 24 h the reaction mixture was exposed to air,
filtered with celite, and then 1 μL was injected onto the GC/MS.

Procedure for the reaction done at 250 psi
In a nitrogen-filled dry-box, 180 mg (0.154 mmol) of 4 was added to a 300 mL Parr reactor followed by 30 mL THF and the nonane standard. The reactor assembly was removed out of the dry-box and the connection line was purged with ethylene for 2 minutes. The reactor was opened to ethylene and the pressure was regulated to 250 psi (17 atm) and the solution stirred for 24 h. After 24 h the ethylene pressure was released and reaction mixture was exposed to air, filtered with celite, and then 1 μL was injected onto the GC/MS.

Preparation of 1,4-dilithiobutane Li(CH2)4Li
10 mL of diethyl ether was added to a Schlenk flask in a nitrogen-filled glove box. The flask was then cooled to −78 °C using a dry ice/acetone bath. 4.2 equivalents of 1.7 M n-BuLi in pentane (11.95 mL, 20.33 mmol) were added to the cold ether solution and allowed to stir for 15 minutes. One equivalent of 1,4-diodobutane (1.50 g, 4.84 mmol) was then added via syringe. The reaction was allowed to stir for 1 h at −78 °C. Afterwards, the diethyl ether was removed under vacuum at −78 °C. The reaction was warmed over 1 hour to 0 °C, while maintaining a dynamic vacuum. The resulting white/yellow product was kept under vacuum for an additional hour at 0 °C. The precipitate was then extracted, in an inert atmosphere, with 100 mL of pentane in 50 mL portions. The gummy white precipitate was then dissolved in dry ethyl ether giving a translucent yellow solution which was stored at −30 °C.

Preparation of [μ-tetramethylene][2,4-pentane-N,N'-bis(2,6-diisopropophenyl)ketiminato]dichromium(II), [[i-Pr2Ph]nacnaCn][μ-CH2]2(CH2)4] (3)
To a solution of [[i-Pr2Ph]nacnaCn]2[μ-CH2]2(CH2)4] (0.200 g, 0.168 mmol) in 70 mL pentane, 0.480 M Et2O solution of Li(CH2)2Li (0.385 mL, 0.185 mmol) was added dropwise. The solution rapidly changed color from green to brown. The solution was stirred at room temperature for 3 hours. The solvent was removed under vacuum leaving a crude brown powder. Pentane was then added to the crude reaction mixture and cooled to −30 °C for 1 hour. The mixture was filtered and the pentane solvent was removed under vacuum leaving a crude brown powder. This was recrystallized from hexamethylsiloxane by slow cooling to −30 °C, yielding X-ray quality crystals (0.093 g, 47% yield). 1H-NMR (400 MHz, C6D6): 8.17 (br), 8.0 (br), 7.82 (br), 7.56 (br), 7.03 (br), 4.36 (br), 3.07 (br), 1.85, 1.52, 1.32, 0.22 ((CH3)3Si)2O ppm. IR (KBr): 3057 (w), 2959 (s), 2925 (m), 2867 (s), 1525 (s), 1434 (s), 1311 (s), 1254 (s), 1171 (m), 1098 (m), 1056 (m), 1042 (m), 932 (w), 846 (m), 793 (s), 760 (m) cm⁻¹. UV/Vis (toluene): λmax (e) = 477 (125 M⁻¹ cm⁻¹) nm. μeff = 2.6 (1 mmuCr (294 K). Mp: 95 °C dec. Mass spectrum m/z (%): 469 (100%) (C29H41N2Cr+). Anal. Calcd for C29H41N2O2Cr: C, 69.77; H, 9.14; N, 4.78. Found: C, 69.66; H, 8.42; N, 4.75. The compound decomposes slowly at room temperature.

Preparation of [μ-pentamethylene][2,4-pentane-N,N'-bis(2,6-diisopropophenyl)ketiminato] dichromium(II), [[i-Pr2Ph]nacnaCn][μ-CH2]2(CH2)4] (4)
To a solution of [[i-Pr2Ph]nacnaCn]2[μ-I]2 (0.200 g, 0.168 mmol) in 70 mL pentane, 0.480 M Et2O solution of Li(CH2)2Li (0.385 mL, 0.185 mmol) was added dropwise. The solution rapidly changed color from green to brown. The solution was stirred at room temperature for 3 hours. The solvent was removed under vacuum leaving a crude brown powder. Pentane was then added to the crude reaction mixture and cooled to −30 °C for 1 hour. The mixture was filtered and the pentane solvent was removed under vacuum leaving a crude brown powder. This was recrystallized from hexamethyldisiloxane by slow cooling to −30 °C, yielding X-ray quality crystals (0.093 g, 47% yield). 1H-NMR (400 MHz, C6D6): 8.17 (br), 8.0 (br), 7.82 (br), 7.56 (br), 7.03 (br), 4.36 (br), 3.07 (br), 1.85, 1.52, 1.32, 0.22 ((CH3)3Si)2O ppm. IR (KBr): 3057 (w), 2959 (s), 2925 (m), 2867 (s), 1525 (s), 1434 (s), 1311 (s), 1254 (s), 1171 (m), 1098 (m), 1056 (m), 1042 (m), 932 (w), 846 (m), 793 (s), 760 (m) cm⁻¹. UV/Vis (toluene): λmax (e) = 477 (125 M⁻¹ cm⁻¹) nm. μeff = 2.6 (1 mmuCr (294 K). Mp: 95 °C dec. Mass spectrum m/z (%): 469 (100%) (C29H41N2Cr+). Anal. Calcd for C29H41N2O2Cr: C, 69.77; H, 9.14; N, 4.78. Found: C, 69.66; H, 8.42; N, 4.75. The compound decomposes slowly at room temperature.
Preparation of (i-Pr2Ph)nacnacCr(η2-C5H7)2 (8)

[(LiPrCr)2(μ–η2–η2–N2)] (1) (0.200 g, 0.206 mmol) was dissolved in 20 mL THF and 0.074 mg (0.161 mol) of diphenylacetylene was added and the solution was stirred for 4 hours. The THF was removed and the product was extracted with pentane. Concentration and cooling to −30 °C yielded brown crystals of 8 (0.197 g, 74% yield). 1H NMR (400 MHz, THF-d8): 143.6 (br), 39.9 (br), 20.4 (br), 17.1 (br), 10.0 (br), 6.64 (br), 2.80 (br), −4.55 (br) ppm. IR (KBr): 3056 (w), 2961 (s), 2926 (m), 2867 (m), 1589 (w), 1526 (s), 1489 (m), 1462 (m), 1436 (m), 1383 (s), 1373 (s), 1315 (s), 1257 (m), 1174 (m), 1103 (m), 1056 (m), 1024 (m), 933 (w), 856 (w), 796 (m), 759 (m), 697 (m) cm−1. UV/vis (pentane): λmax (ε) = 390 (4657 M−1 cm−1) nm. μeff (293 K) = 3.9 (1μB). Mp: 211 °C (dec). Mass spectrum m/z (%): 496.3 (70) [M+ − C2Ph2].

Preparation of (i-Pr2Ph)nacnacCr(CPh4)(CH2)2 (9)

(i-Pr2Ph)nacnacCr(η2–C5H7)2 (8) (0.100 g, 0.154 mmol) was dissolved in 20 mL excess ethylene and the solution was stirred for up to 24 hours. The THF was removed and the product was extracted with pentane. Concentration and cooling to −30 °C yielded light brown crystals of 9 (0.140 g, 60% yield). 1H NMR (400 MHz, THF-d8): 29.78 (br), 14.86 (br), 11.01 (br), 7.70 (br), 4.73 (br), −6.66 (br), −13.01 (br) ppm. IR (KBr): 3057 (w), 3016 (w), 2960 (s), 2926 (s), 2866 (m), 1522 (s), 1490 (m), 1472 (m), 1465 (m), 1457 (s), 1437 (s), 1419 (m), 1384 (s), 1374 (s), 1339 (w), 1256 (m), 1172 (w), 1101 (w), 1025 (w), 933 (s), 795 (m), 756 (m), 698 (m), 668 (w) cm−1. UV/vis (pentane): λmax (ε) = 370 (4200 M−1 cm−1), 454 (1195 M−1 cm−1) nm. μeff (293 K) = 3.4 (1μB). Mp: 192 °C (dec). Mass spectrum m/z (%): 469.2 (51) [M+ − C4H14H6 − C2H4O].

Crystallographic structure determinations

A summary of crystal data collection and refinement parameters for compounds 3, 4, 6, 7, 8, 9 and 10 can be found in Table 3. Suitable crystals were selected, mounted with viscous

| Table 3 Crystallographic data and refinement details for 3, 4, 6, 7, 8, 9 and 10* |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Formula         | C35H52CrN5O2S12 | C41H61CrN2O     | C41H52CrN2O     | C41H54CrN2      | C41H54CrN2O     | C41H54CrN2O     |
| Fw              | 1157.76         | 1081.55         | 1593.56         | 73.68           | 647.86          | 649.92          |
| Space group     | P2_1            | P1              | Pna             | P2_1           | Pna             | P2_1/c          |
| Color           | Brown           | Brown           | Green           | Orange         | Brown           | Green           |
| a (Å)           | 13.404(5)       | 13.187(4)       | 11.966(6)       | 11.966(4)      | 11.966(6)       | 11.966(4)       |
| b (Å)           | 14.089(5)       | 13.657(4)       | 14.191(7)       | 14.191(7)      | 14.191(7)       | 14.191(7)       |
| c (Å)           | 18.924(7)       | 18.405(6)       | 16.977(8)       | 16.977(8)      | 16.977(8)       | 16.977(8)       |
| α (°)           | 90              | 90              | 90              | 90             | 90              | 90              |
| β (°)           | 109.268(9)      | 74.798(4)       | 97.191(5)       | 97.191(5)      | 97.191(5)       | 97.191(5)       |
| γ (°)           | 90              | 90              | 90              | 90             | 90              | 90              |
| V (Å³)          | 3374(2)         | 3175.8(17)      | 3135.3(11)      | 3135.3(11)     | 3135.3(11)      | 3135.3(11)      |
| Z, Z'          | 2, 1            | 2, 1            | 4, 1            | 4, 1           | 4, 1            | 4, 1            |
| D(001) (g cm⁻³) | 1.140           | 1.131           | 1.207           | 1.207          | 1.207           | 1.207           |
| β(MO) (°)       | 1.140           | 1.140           | 1.140           | 1.140          | 1.140           | 1.140           |
| Temp. K         | 120(2)          | 120(2)          | 120(2)          | 120(2)         | 120(2)          | 120(2)          |
| Data/params     | 11.546(3)       | 11.546(3)       | 11.546(3)       | 11.546(3)      | 11.546(3)       | 11.546(3)       |
| GOF on F²      | 1.052           | 1.052           | 1.052           | 1.052          | 1.052           | 1.052           |
| R(F²) (%)       | 7.42            | 5.84            | 8.41            | 8.41           | 8.41            | 8.41            |
| Rw(F²)(%)       | 12.77           | 16.20           | 18.15           | 18.15          | 18.15           | 18.15           |
| Flack param     | 0.03(3)         | —               | —               | —              | —               | —               |

* Quantity minimized: Rp(F²) = Σ[w(Fo² – Fc²)]/Σ[w(Fo²)]²/2.
oil and cooled to the data collection temperature. Data were collected on a Bruker APEX CCD diffractometer using graphite monochromated Mo-Kα radiation (λ = 0.71073). Unit cell parameters were obtained from 60 data frames, 0°, from three different sections of the Ewald sphere. No symmetry higher than triclinic was observed for 4 and 10. The systematic absences in the data and the unit cell parameters were consistent to P2₁ and P2₁/m for 3, and Pnma and Pn2₁a (Pna2₁) for 6 and 8, and, uniquely, P2₁/n for 7, and P2₁/c for 9. The centrosymmetric space group options for 4 and 10 (P1), and 6 and 8 (Pnma), and the noncentrosymmetric space group for 3 (P2₁) yielded chemically reasonable and computationally stable results of refinement. The data-sets were treated with SADABS absorption corrections based on redundant multiscan data. The structures were solved using direct methods and refined with full-matrix, least-squares procedures on F². A close inspection of the packing diagram did not reveal any overlooked symmetry for 3 and the Flack parameter refined to 0.03(3) indicating that the true hand of the data was determined. Compound 4 consistently deposits as small, weakly-diffracting crystals resulting in lower-than-ideal 2θ range. Two symmetry unique but chemically identical compound molecules were found in the asymmetric unit of 8. The metallaecyclic carbon atoms and the isopropyl moieties displayed unresolvable disorder causing a larger than usual Ueq range for carbon atoms and associated hydrogen atoms. Compound 6 was located disordered at a mirror plane. Solvent molecules were found cocrySTALLinized in the asymmetric units of 3 (hexamethyldisiloxane), 4 (pentane), 7 (pentane) and 9 (pentane). Disordered pentane molecules of crystallization in 7 with net occupancy of 1.5 per unit-cell, in two void spaces located away from the compound molecule, and two molecules of co-crySTALLinized pentane per compound molecule in 9 were treated as diffused contributions. All non-hydrogen atoms were refined with anisotropic displacement parameters. Hydrogen atoms on metallaecycles were located from the difference map, treated with chemically sensible restraints, and refined with a riding model. All other hydrogen atoms were treated as idealized contributions with 1.2–1.5Ueq of the associated carbon atom. Atomic scattering factors are contained in the SHELXTL 6.12 program library.

Acknowledgements

This work was supported by the U.S. National Science Foundation, through grants CHE-0616375 and CHE-0911081 to KHT.

Notes and references

1 K. Weissermel and H.-J. Arpe, Industrial Organic Chemistry, VCH, Weinheim, 2003.

2 D. Vogt, in Applied Homogeneous Catalysis with Organometallic Compounds, ed. B. Cornils and W. A. Herrmann, VCH, Weinheim, 1996, pp. 245–258.

3 R. M. Manyik, W. E. Walker and T. P. Wilson, Union Carbide Corporation, US, Continuous processes for the production of ethylene polymers and catalysts suitable therefore, US Patent, 3300458, 1967.

4 R. M. Manyik, W. E. Walker and T. P. Wilson, J. Catal., 1977, 47, 197–209.

5 J. T. Dixon, M. J. Green, F. M. Hess and D. H. Morgan, J. Organomet. Chem., 2004, 689, 3641–3668.

6 D. F. Wass, Dalton Trans., 2007, 816–819.

7 T. Agapie, Coord. Chem. Rev., 2011, 255, 861–880.

8 B. R. Aluri, N. Peulecke, S. Peitz, A. Spannenberg, B. H. Muller, S. Schulz, H. J. Drexlner, D. Heller, M. H. Al-Hazmi, F. M. Mosa, A. Wohl, W. Muller and U. Rosenthal, Dalton Trans., 2010, 39, 7911–7920.

9 B. H. Muller, N. Peulecke, S. Peitz, B. R. Aluri, U. Rosenthal, M. H. Al-Hazmi, F. M. Mosa, A. Wohl and W. Muller, Chem.–Eur. J., 2011, 17, 6935–6938.

10 B. H. Muller, N. Peulecke, A. Spannenberg, U. Rosenthal, M. H. Al-Hazmi, R. Schmidt, A. Wohl and W. Muller, Organometallics, 2012, 31, 3695–3699.

11 W. Muller, A. Wohl, S. Peitz, N. Peulecke, B. R. Aluri, B. H. Mueller, D. Heller, U. Rosenthal, M. H. Al-Hazmi and F. M. Mosa, ChemCatChem, 2010, 2, 1130–1142.

12 S. Peitz, B. R. Aluri, N. Peulecke, B. H. Muller, A. Wohl, W. Muller, M. H. Al-Hazmi, F. M. Mosa and U. Rosenthal, Chem.–Eur. J., 2010, 16, 7670–7676.

13 S. Peitz, N. Peulecke, B. R. Aluri, S. Hansen, B. H. Muller, A. Spannenberg, U. Rosenthal, M. H. Al-Hazmi, F. M. Mosa, A. Wohl and W. Muller, Eur. J. Inorg. Chem., 2010, 1167–1171.

14 S. Peitz, N. Peulecke, B. R. Aluri, B. H. Muller, A. Spannenberg, U. Rosenthal, M. H. Al-Hazmi, F. M. Mosa, A. Wohl and W. Muller, Organometallics, 2010, 29, 5263–5268.

15 S. Peitz, N. Peulecke, B. R. Aluri, B. H. Muller, A. Spannenberg, U. Rosenthal, M. H. Al-Hazmi, F. M. Mosa, A. Wohl and W. Muueller, Organometallics, 2011, 30, 2364–2370.

16 S. Peitz, N. Peulecke, B. R. Aluri, B. H. Muller, A. Spannenberg, U. Rosenthal, M. H. Al-Hazmi, K. E. Al-Eidan, A. Wohl and W. Muueller, Organometallics, 2011, 30, 4080–4089.

17 A. Wohl, W. Muller, S. Peitz, N. Peulecke, B. R. Aluri, B. H. Muller, D. Heller, U. Rosenthal, M. H. Al-Hazmi and F. M. Mosa, Chem.–Eur. J., 2010, 16, 7833–7842.

18 S. Licciulli, I. Thapa, K. Albahily, I. Korobkov, S. Gambarotta, R. Duchateau, R. Chevalier and K. Schuh, Angew. Chem., Int. Ed., 2010, 49, 9225–9228.

19 I. Thapa, S. Gambarotta, I. Korobkov, R. Duchateau, S. V. Kulangara and R. Chevalier, Organometallics, 2010, 29, 4080–4089.

20 R. Duchateau, S. Gambarotta, S. Licciulli, K. Albahily, S. V. Kulangara and I. Thapa, Abstr. Pap. Am. Chem. Soc., 2011, 241, 1155-INOR.

21 S. Licciulli, K. Albahily, V. Fomitcheva, I. Korobkov, S. Gambarotta and R. Duchateau, Angew. Chem., Int. Ed., 2011, 50, 2346–2349.
22 K. Albahily, S. Gambarotta and R. Duchateau, Organometallics, 2011, 30, 4655–4664.
23 K. Albahily, V. Fomitcheva, Y. Shaikh, E. Sebastian, S. I. Gorelsky, S. Gambarotta, I. Korobkov and R. Duchateau, Organometallics, 2011, 30, 4201–4210.
24 K. Albahily, S. Liciuilli, S. Gambarotta, I. Korobkov, R. Chevalier, K. Schuhen and R. Duchateau, Organometallics, 2011, 30, 3346–3352.
25 I. Thapa, S. Gambarotta, I. Korobkov, M. Murugesu and P. Budzelaar, Organometallics, 2012, 31, 486–494.
26 Y. Shaikh, K. Albahily, M. Sutcliffe, V. Fomitcheva, S. Gambarotta, I. Korobkov and R. Duchateau, Angew. Chem., Int. Ed., 2012, 51, 1366–1369.
27 K. Albahily, Z. Ahmed, S. Gambarotta, E. Koc, R. Duchateau and I. Korobkov, Organometallics, 2011, 30, 6022–6027.
28 Y. Shaikh, J. Gurnham, K. Albahily, S. Gambarotta and I. Korobkov, Organometallics, 2012, 31, 7427–7433.
29 L. E. McDyre, T. Hamilton, D. M. Murphy, K. J. Cavell, W. F. Gabrielli, M. J. Hanton and D. M. Smith, Dalton Trans., 2010, 39, 7792–7799.
30 A. F. R. Kilpatrick, S. V. Kulatingara, M. G. Cushion, R. Duchateau and P. Mountford, Dalton Trans., 2010, 39, 3653–3664.
31 Y. Qi, Q. Dong, L. Zhong, Z. Liu, P. Y. Qiu, R. H. Cheng, X. L. He, J. V. C. Hamilton and B. P. Liu, Organometallics, 2010, 29, 1588–1602.
32 I. Y. Skobelev, V. N. Panchenko, O. Y. Lyakin, K. P. Bryliakov, V. A. Zakharov and E. P. Talsi, Organometallics, 2010, 29, 2943–2950.
33 T. W. Hey and D. F. Wass, Organometallics, 2010, 29, 3676–3678.
34 S. K. Kim, T. J. Kim, J. H. Chung, T. K. Hahn, S. S. Chae, H. S. Lee, M. Cheong and S. O. Kang, Organometallics, 2010, 29, 5805–5811.
35 Z. Mohamadnia, E. Ahmadi, M. N. Haghighi and H. Salehi-Mobarakeh, Catal. Lett., 2011, 141, 474–480.
36 P. Y. Qiu, H. Cheng, B. P. Liu, B. Tumanskii, R. J. Batrice, M. Botoshansky and M. S. Eisen, Organometallics, 2011, 30, 2144–2148.
37 P. W. N. M. van Leeuwen, N. D. Clement and M. J. L. Tschan, Coord. Chem. Rev., 2011, 255, 1499–1517.
38 E. Ahmadi, Z. Mohamadnia and M. N. Haghhighi, Catal. Lett., 2011, 141, 1191–1198.
39 S. F. Liu, R. Pattaccini and P. Braunstein, Organometallics, 2011, 30, 3549–3558.
40 L. H. Do, J. A. Labinger and J. E. Bercaw, Organometallics, 2012, 31, 5143–5149.
41 T. E. Stennett, M. F. Haddow and D. F. Wass, Organometallics, 2012, 31, 6960–6965.
42 W. H. Monillas, G. P. A. Yap and K. H. Theopold, Angew. Chem., Int. Ed., 2007, 46, 6692–6694.
43 W. H. Monillas, G. P. A. Yap, L. A. MacAdams and K. H. Theopold, J. Am. Chem. Soc., 2007, 129, 8090–8091.
44 W. H. Monillas, G. P. A. Yap and K. H. Theopold, Inorg. Chim. Acta, 2011, 369, 103–119.
45 A. A. Frost and R. G. Pearson, Kinetics and Mechanism, John Wiley & Sons, Inc., New York, 1953.
46 R. Arteaga-Muller, H. Tsurugi, T. Saito, M. Yanagawa, S. Oda and K. Mashima, J. Am. Chem. Soc., 2009, 131, 5370–5371.
47 J. R. Briggs, J. Chem. Soc., Chem. Commun., 1989, 674–675.
48 L. A. MacAdams, G. P. Buffone, C. D. Incarvito, J. A. Golen, A. L. Rheingold and K. H. Theopold, Chem. Commun., 2003, 1164–1165.
49 K. H. Theopold and R. G. Bergman, Organometallics, 1982, 1, 1571–1579.
50 J. R. Moss and L. G. Scott, Coord. Chem. Rev., 1984, 60, 171–190.
51 C. P. Casey and J. D. Audett, Chem. Rev., 1986, 86, 339–352.
52 E. Negishi, D. R. Swanson and C. J. Roussel, J. Org. Chem., 1990, 55, 5404–5409.
53 W. F. Bailey and E. R. Punzalan, J. Org. Chem., 1990, 55, 5404–5406.
54 C. W. Kamienski and D. H. Lewis, J. Org. Chem., 1965, 30, 3498–3504.
55 W. F. Bailey, R. P. Gagnier and J. J. Patricia, J. Org. Chem., 1984, 49, 2098–2107.
56 V. C. Gibson, C. Newton, C. Redshaw, G. A. Solan, A. J. P. White and D. J. Williams, Eur. J. Inorg. Chem., 2001, 1895–1903.
57 R. A. Heintz, R. L. Ostrander, A. L. Rheingold and K. H. Theopold, J. Am. Chem. Soc., 1994, 116, 11387–11396.
58 A. R. Hermes, R. J. Morris and G. S. Girolami, Organometallics, 1988, 7, 2372–2379.
59 P. Müller, R. Herbst-Irmer, A. L. Spek, T. R. Schneider and M. R. Sawaya, Crystal Structure Refinement – A Crystallographers’s Guide to SHELXL, Oxford University Press, Oxford, 2006.
60 R. L. Carlin, Magnetocchemistry, Springer, Berlin, 1986.
61 P. J. Hay, J. C. Thibeault and R. Hoffmann, J. Am. Chem. Soc., 1975, 97, 4884–4899.
62 K. Angermund, P. Betz, A. Dohring, P. W. Jolly, C. Kruger and K. U. Schonfelder, Polyhedron, 1993, 12, 2663–2669.
63 A. Dohring, J. Gohre, P. W. Jolly, B. Kryger, J. Rust and G. P. J. Verhovnik, Organometallics, 2000, 19, 388–402.
64 N. F. Wang, D. J. Wink and J. C. Dewan, Organometallics, 1990, 9, 335–340.
65 D. W. Norman, M. J. Ferguson, R. McDonald and J. M. Stryker, Organometallics, 2006, 25, 2705–2708.
66 C. E. Elschenbroich and A. Salzer, Organometallics, A Concise Introduction, VCH, Weinheim, 1992.
67 L. A. MacAdams, G. P. Buffone, C. D. Incarvito, A. L. Rheingold and K. H. Theopold, J. Am. Chem. Soc., 2005, 127, 1082–1083.
68 K. H. Theopold, L. A. MacAdams, C. Puttmual, G. P. Buffone and A. L. Rheingold, Polyom. Mater. Sci. Eng., 2002, 86, 310.
69 The CC distance of free diphenylacetylene is 1.205(4) Å taken from the following publications retrieved from the CSD: (a) V. D. Samarskaya, R. M. Myasnikova and A. I. Kitaigorodskij, Kristallografija, 1968, 13, 616;
A search of the CSD showed 14 structures of stilbene with an average C≡C distance of 1.321(5) Å (range: 1.288–1.341 Å).

A search of the CSD showed 10 structures with a $\eta^2$-diphenylacetylene fragment with a C–C–C$_{ipso}$ angle of 145.3(19)° (range: 136.3–155.9°). Selected examples:

(a) D. J. Wink, J. R. Fox and N. J. Cooper, *J. Am. Chem. Soc.*, 1985, 107, 5012;
(b) I. M. Barlett, N. G. Connelly, A. G. Orpen and M. J. Quayle, *Chem. Commun.*, 1996, 2583;
(c) A. A. Danopoulos, G. Wilkinson, T. K. N. Sweet and M. B. Hursthouse, *J. Chem. Soc., Dalton Trans.*, 1996, 271;
(d) C. J. Adams, I. M. Bartlett, N. G. Connelly, D. J. Harding, O. D. Hayward, A. J. Martin, M. J. Quayle and P. H. Reiger, *J. Chem. Soc., Dalton Trans.*, 2002, 4281.

Y. Yamamoto, *Curr. Org. Chem.*, 2005, 9, 503–519.

L. S. Zhou, S. Li, K. Kanno and T. Takahashi, *Heterocycles*, 2010, 80, 725–738.

L. E. Bowen, M. F. Haddow, A. G. Orpen and D. F. Wass, *Dalton Trans.*, 2007, 1160–1168.

A. J. Rucklidge, D. S. McGuinness, R. P. Tooze, A. M. Z. Slawin, J. D. A. Pelletier, M. J. Hanton and P. B. Webb, *Organometallics*, 2007, 26, 2782–2787.

A. Jabri, P. Crewdson, S. Gambarotta, I. Korobkov and R. Duchateau, *Organometallics*, 2006, 25, 715–718.

C. Temple, A. Jabri, P. Crewdson, S. Gambarotta, I. Korobkov and R. Duchateau, *Angew. Chem., Int. Ed.*, 2006, 45, 7050–7053.

T. Agapie, J. A. Labinger and J. E. Bercaw, *J. Am. Chem. Soc.*, 2007, 129, 14281–14295.

G. M. Sheldrick, *Acta Crystallogr., Sect. A: Fundam. Crystallogr.*, 2008, 64, 112–122.

H. D. Flack, *Acta Crystallogr., Sect. A: Fundam. Crystallogr.*, 1983, 39, 876–881.

A. L. Spek, *J. Appl. Crystallogr.*, 2003, 36, 7–13.