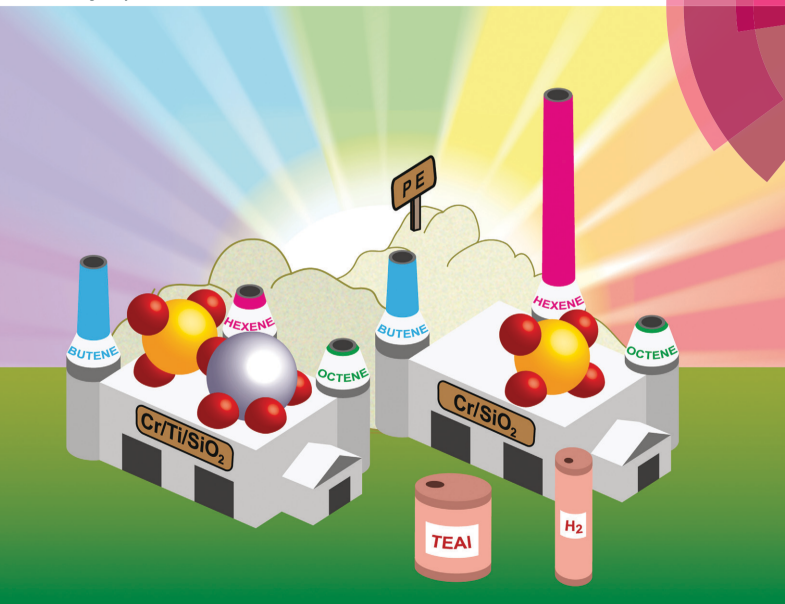
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PAPER

B. M. Weckhuysen et al. Structure–performance relationships of Cr/Ti/SiO $_2$ catalysts modified with TEAl for oligomerisation of ethylene: tuning the selectivity towards 1-hexene

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Introduction

Phillips-type Cr/SiO₂ polymerisation catalysts are responsible for the commercial production of more than one third of all polyethylene sold worldwide.^{1–4} This has attracted a great deal of research attention both in academia and chemical industries since their discovery in the early 1950s by J. P. Hogan and R. L. Banks at Phillips Petroleum Company.^{5,6} Despite the numerous research efforts on the oxidation states of the catalyst,^{7–11} the molecular structure of the active sites,^{2,12–16} and the mechanism of ethylene polymerisation with Phillipstype catalysts,^{17–23} however, these key scientific questions still require further attention.

No univocal consensus has been yet achieved in both academia and chemical industries, especially in studies with polymerisation conditions similar to industrial ones, as many of the research studies involved model systems, temperatures and pressures, which were quite different from the conditions utilised in industrial systems. One of the reasons is the very high sensitivity of the Cr/SiO_2 ethylene polymerisation catalyst to trace amounts of catalytic poisons, *i.e.* H₂O, O₂



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A study on the influence of H₂ and the degree of titanation of a Cr/SiO₂ Phillips polymerisation catalyst on the selective oligomerisation of ethylene induced by pre-contacting the catalyst with triethylaluminium (TEAI) is presented. Ethylene oligomerisation reactions were performed at 373 K and 1 bar, inside a quartz reactor of a specially designed *operando* experimental setup, which allowed examination of the catalysts by UV-vis-NIR diffuse reflectance spectroscopy, while the gas phase was continuously monitored by online mass spectroscopy and subsequently analysed by gas chromatography. With this combination of techniques, it was possible to acquire detailed insight into the different distributions of produced oligomers, which were highly dependent on the catalyst structure. The addition of small amounts of Ti significantly changes the electronic environment of Cr oligomerisation sites by the formation of Cr–O–Ti–O–Si linkages, favouring β -H transfer and increasing the selectivity towards butene at the expense of 1-hexene. Moreover, we show that ethylene oligo-/polymerisation reactions follow at least two different pathways, *i.e.* metallacycle for olefinic species with a broken Schulz-Flory distribution, and Cossee–Arlman for other hydrocarbon species.

> and small oxygenated organic compounds. Furthermore, the fast rate of polymerisation and high productivity of the catalysts make the study of the initial stages of ethylene polymerisation very challenging.

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Certain adjustments of the catalyst furthermore complicate the picture. Cr/SiO_2 ethylene polymerisation catalysts can often be modified by incorporation of titania, yielding a $Cr/Ti/SiO_2$ catalyst with increased activity and capability of producing polyethylene with a lower molecular weight and a broader molecular weight distribution than the pristine catalyst. Titanation can be performed in two manners, either by co-precipitation with silica gel, when titania becomes highly dispersed in the bulk catalyst material,²⁴ or by reaction of a titanium compound with the hydroxyl groups of the support, therefore coating the silica surface with a layer of titania.^{25–28} Consequently, both methods include the formation of Cr–O– Ti–O–Si linkages,²⁹ which change the electronic environment of the active sites and influence the mechanism of ethylene polymerisation.

One of the unique aspects of Phillips-type Cr/SiO_2 catalysts is that they do not require activation by a metal alkyl cocatalyst as compared to Ziegler–Natta and metallocene polymerisation catalysts and therefore these compounds are often excluded, even at industrial production sites.^{30–32} However, when metal alkyls are used in combination with Phillips-type Cr/SiO_2 catalysts, the development of the polymerisation rate

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is commonly accelerated and the induction time is decreased. It is believed that these effects are caused by a more facile reduction of the Cr⁶⁺ species of the activated catalyst to Cr species in a lower oxidation state, which are thought to be responsible for the polymerisation of ethylene.¹ Most of the authors attribute the polymerisation-active sites to Cr²⁺ species of different coordinative unsaturations^{19,33–35} although other species of higher valences, *i.e.* Cr³⁺, Cr⁴⁺ and Cr⁵⁺, should still be taken into consideration,^{8,10,36-38} especially in view of the recent work of the group of Copéret.^{20,21,39} Secondly, metal alkyl co-catalysts can improve the polymerisation rate by alkylation of Cr sites and provide chain initiation similarly to Ziegler-Natta catalysts. However, with Phillips-type catalysts, this chain initiation by a cocatalyst is not necessary. Thirdly, as highly reactive compounds, metal alkyls can remove the poisons still present in the gas feed or in the system, such as the ethylene oxidation products formed during the initiation stage that might remain adsorbed on the Cr sites. Depending on the type of co-catalyst, there is an optimal amount expressed as the metal-to-Cr ratio, which gives the highest catalytic activity. Besides the improvement of the activity and decrease of the induction time, further increase of the co-catalyst amount can deteriorate the polymerisation rate or even destroy the catalyst due to attack on Cr-support bonds. Some of the metal alkyls can act as chain transfer agents, exchanging the alkyl groups with Cr polymerisation sites, and therefore affect the properties of the produced polyethylene, although less than in Ziegler-Natta and metallocene catalysis. Co-catalysts can also influence the polymerisation sites by reacting with Cr or neighbouring groups and directly modifying the polymerisation activity and the type of polyethylene that is produced.

One of the most interesting facts regarding co-catalysts and their interaction with the Cr sites is that the addition of metal alkyl compounds to the reactor containing a Phillipstype catalyst changes the amount of short-chain branching of the produced polyethylene, even if no α -olefin co-monomers were added to the system. This short-chain branching of the polyethylene chain originates from the co-polymerisation of ethylene with the *in situ* produced ethylene α -oligomers on modified polymerisation active sites.^{40,41} In situ branching attracts high interest in the industrial world due to the possibility of reducing or even eliminating the separate feed of α -olefin comonomers, their transportation, storage and purification, therefore simplifying manufacturing procedures and production costs of MDPE and LLDPE. The recent drop in ethylene price of more than \$500 per metric ton from January 2014 to January 2015⁴² decreases the production cost of polyethylene and increases the economic importance of in situ ethylene oligomerisation. Therefore, an accurate description and finding the means to directly influence in situ ethylene oligomerisation, i.e. selectivity, olefin distribution and their yield, proves to be one of the hottest topics in the ethylene polymerisation industry.

Although a lot of progress has been made in the last few decades in resolving these issues by applying advanced characterisation techniques,^{18,40,43-46} no unifying picture has yet been achieved. Here, we present an *operando* UV-vis-NIR diffuse reflectance study (DRS) of a Cr/Ti/SiO₂ catalyst in order to elucidate the TEAl-induced *in situ* ethylene oligomerisation properties.

Experimental

Catalyst preparation

The catalyst samples were provided by Total Research and Technology Feluy, Belgium. Catalyst preparation includes the treatment of two commercially available Cr/SiO₂ pre-catalysts containing either a higher or a lower Cr loading, with the following properties: a pre-catalyst with ~1.0 wt% Cr loading, a surface area of 500 m² g⁻¹, a pore volume of 1.5 cm³ g⁻¹ and a D50 particle size diameter of 72 µm, and a pre-catalyst with ~0.5 wt% Cr loading, a surface area of 318 m² g⁻¹, a pore volume of 1.55 cm³ g⁻¹ and a D50 particle size diameter of 47 µm. The pale white silica pre-catalyst was placed in a fluidised bed reactor under N2 flow and slowly heated to 543 K for dehydration. While still under N₂ flow, in the case of the titanated samples, the pre-catalyst material was surface-titanated, according to the procedure described by Debras et al., to either 2 wt% or 4 wt% of the total Ti loading using titanium isopropoxide (99.999% trace metals basis, Sigma-Aldrich) which was added dropwise to the fluidised bed.²⁵ After flushing under N₂ for 45 min, the gas flow was changed to dry air and the temperature was slowly increased to the desired activation temperature. During the 6 h oxidation step, the colour of the prepared Cr/SiO₂ and Cr/Ti/SiO₂ catalysts changed to intense yellow/orange due to the formation of Cr⁶⁺ species. After cooling down to RT and switching back to N₂ flow, the activated Cr/SiO₂ and Cr/Ti/SiO₂ catalysts could be obtained under inert N2 atmosphere and transferred to an Ar glove box for storage and further use. An overview of the catalysts' properties and activation temperatures is given in Table 1.

Characterisation of the Cr/SiO₂ ethylene polymerisation catalysts

The setup. The prepared catalyst materials were tested in a specially designed setup (Fig. 1), which allows an *operando* UV-vis-NIR DRS study of the catalyst under working conditions and a real-time analysis of the reaction products with on-line Mass Spectrometry (MS) and Gas Chromatography (GC). The setup is suitable for gas phase polymerisation reactions with solid catalysts at 1 bar and temperatures in the range from RT to 473 K.

Ethylene oligo-/polymerisation. Approximately 300 mg of a catalyst sample was loaded into a quartz cell under inert atmosphere and heated to a reaction temperature of 373 K. Before the feed of ethylene, the catalyst was pre-contacted with 0.1 cm³ of 1.3 M (for an Al:Cr ratio of 2:1) triethyl-aluminium co-catalyst (TEAl) in heptanes (~94 wt% TEAl, with ~6 wt% predominately tri-*n*-butylaluminium and less than 0.1 wt% triisobutylaluminium residue, Acros Organics),

Table 1 Overview of the prepared Phillips-type ethylene polymerisation catalysts and support materials, their properties and the positions of the O \rightarrow Cr⁶⁺ charge transfer bands

Sample name	Act. temp. (K)	Cr loading (wt%)	Ti loading (wt%)	Surface area $(m^2 g^{-1})$	Pore volume (cm ³ g ⁻¹)	$O \rightarrow Cr$	⁶⁺ CT (cn	n ⁻¹)
S	1048	0	0	246	1.39		n.a.	
TS	1048	0	4.7	254	1.51		n.a.	
CS	1048	0.52	0	296	1.30	38 500	29 500	21 900
CTS1	1048	0.62	2.2	293	1.44	38100	28700	21700
CTS2	1048	0.56	3.9	277	1.39	37 500	27800	21600
CTS3	923	0.56	4.0	289	1.34	37 200	27 800	21600
CTS4	823	0.55	3.8	294	1.35	36 500	27500	21600
CTS5	1048	0.98	4.7	367	1.13	36 500	27500	21700

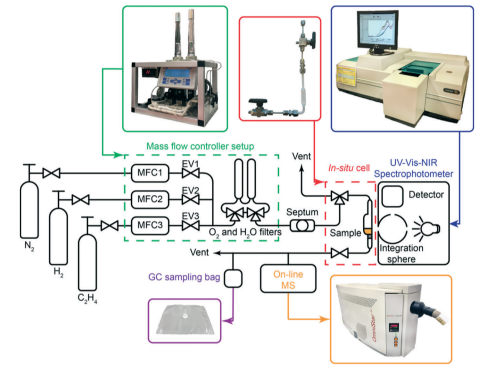


Fig. 1 UV-vis-NIR diffuse reflectance spectroscopy, mass spectrometry and gas chromatography setup developed for the testing of solid catalysts and products obtained in gas phase in reactions at 1 bar and temperatures in the range from RT to 473 K. The *operando* setup includes changeable gas reactant sources, mass flow controller setup (green), septum for the injection of the co-catalyst, 4 cm³ air-tight quartz reactor (red), Varian Cary 500 spectrophotometer (blue), OmniStar mass spectrometer (orange) and gas sampling bag for the GC(-MS) analysis of gas phase (purple). All lines and the reactor are traced and heated to the desired reaction temperature monitored by a couple of thermocouples.

which was injected through a silicon septum into a N_2 stream of 10 cm³ min⁻¹. After the stabilisation of the spectra, a gas reaction mixture consisting of either 45 vol% N_2 , 45 vol% C_2H_4 and 10 vol% H_2 or 50 vol% N_2 and 50 vol% C_2H_4 was fed into the cell for a total flow of 10 cm³ min⁻¹. The gases were provided by Linde Gas with the following purities N_2 (99.999%), H_2 (99.999%) and C_2H_4 (99.95%). Modification of the catalyst with TEAl and polymerisation reaction were simultaneously monitored with *operando* UV-vis-NIR DRS, while the gaseous products were analysed on-line with MS and collected for GC analysis.

UV-vis-NIR diffuse reflectance spectroscopy. UV-vis-NIR DRS measurements were performed with a Varian Cary 500 Scan UV-vis-NIR spectrophotometer equipped with a diffuse reflectance accessory. To obtain the diffuse reflectance spectra of the catalyst materials, a reference spectrum of a Halon

white standard was first measured and automatically subtracted from the actual DRS data using the Cary Win UV Scan Application. All data were acquired in the spectral range of $4000-45\,000 \text{ cm}^{-1}$, with 17 cm⁻¹ spectral resolution and 33 ms data point scan time in the UV-vis region ($12\,500-45\,000 \text{ cm}^{-1}$) and 7.1 cm⁻¹ spectral resolution and 33 ms data point scan time in the NIR region ($4000-12\,500 \text{ cm}^{-1}$). The UV-vis-NIR DRS spectra were corrected for the detector/grating and light source changeover steps at $12\,500 \text{ cm}^{-1}$ and $28\,570 \text{ cm}^{-1}$, respectively, and smoothed in the UV-vis region using a FFT filter with a 0.00147 cut-off frequency. The spectra were repeatedly taken every 120 s.

On-line mass spectrometry. The released gases during the pre-treatment with TEAl and polymerisation of ethylene were analysed by a quadrupole on-line Pfeiffer OmniStar mass spectrometer, which was connected at the exit of the quartz

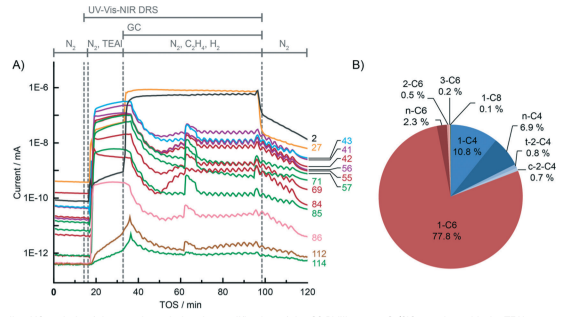


Fig. 2 A) On-line MS analysis of the gas phase during the modification of the CS Phillips-type Cr/SiO_2 catalyst with the TEAl co-catalyst and subsequent oligo-/polymerisation of ethylene using the in-house designed *operando* setup. UV-vis-NIR DRS repeated measurements were started at t = 14 min. At t = 16 min, TEAl was injected into the system *via* evaporation into the N₂ stream. After 17 min, the gas feed was changed to the reaction mixture consisting of 45% N₂, 45% C₂H₄ and 10% H₂, at a total flow rate of 10 cm³ min⁻¹ and the gas phase was sampled for the GC analysis. After 60 min of reaction, the gas feed was changed back to N₂ and UV-vis-NIR DRS and GC sampling stopped. Assignment of the *m/z* is as follows: 2 H₂; 27 ethylene, 1-butene, 1-hexene, hexane, TEAl, butane, heptanes; 43 1-butene, butane, heptanes, octane; 41 1-butene, 1-hexene, hexane, butane, heptanes; 56 1-hexene, hexane, 1-butene; 42, 55, 69, 84 1-hexene, 1-octene; 57 hexane, TEAl, heptanes; 71, 85 TEAl, octane; 71, 85, 114 TEAl; 86 hexane and 112 1-octene. B) GC analysis of the gas phase collected during oligo-/polymerisation of ethylene after the modification of the catalyst with TEAl.

reactor. Ion currents were recorded for 33 different m/z values in the range between 2 and 114, selected to detect and distinguish different produced hydrocarbon species. Data were recorded using the program Quadstar 32-Bit.

Gas chromatography-mass spectrometry. The identification of hydrocarbon species from the gas phase, collected into 1 litre Sigma-Aldrich Tedlar gas-sampling bags during the polymerisation of ethylene, was performed on a QP2010 Shimadzu GC-MS apparatus with a VF-5 ms column by manual injection of 1 cm³ of the gas phase using a 1 cm³ gastight GC syringe. The column flow was set at 1 cm³ min⁻¹ and the column temperature was kept constant at 305 K during the whole analysis. Quantification of the amount of hydrocarbon species in the gas phase was performed on a Varian 430-GC Gas Chromatograph with a FID detector. The same type of column and experimental settings as for the GC-MS experiments were used. The calibration curve was made by the dilution of a reference light hydrocarbon gas mixture, provided by Linde Gas, with N₂.

Results

Integration of UV-vis-NIR diffuse reflectance spectroscopy, mass spectrometry and gas chromatography: an experimental example

For the sake of clarity, the results obtained by the integration of multiple analytical techniques within the specially designed setup, *i.e.* UV-vis-NIR diffuse reflectance spectroscopy (DRS), mass spectrometry (MS) and gas chromatography (GC) (Fig. 1), will be first explained for the ethylene polymerisation reaction using the CS catalyst (Table 1), after which the differences between the catalyst structures and reaction conditions will be discussed in the following sections.

Fig. 2A shows the on-line MS analysis and time evolution of several fragments of the compounds of interest during the pre-run, catalyst modification with TEAl, ethylene oligo-/polymerisation reaction and post-run. The UV-vis-NIR DRS spectra were repeatedly collected during the TEAl-modification stage and polymerisation reaction, while the gas phase was sampled for the GC analysis during the reaction stage. Numerous mass fragments were monitored although only the evolution of the most relevant fragments was presented, based on the GC analysis shown in Fig. 2B.† The injection of TEAl solution in heptanes into the N_2 stream results in a sudden increase of the detector current and saturation of the signal due to the high concentration of the solvents. After 17 min of flushing off the solvent, the gas feed was changed to the reaction mixture consisting of 45 vol% N_2 , 45 vol% C_2H_4

[†] The concentration of the oligomers in the gas phase may be altered by the consumption of the oligomers in the polymer. However, we assume that the oligomer consumption by polymer formation is negligible due to the small amount of PE formed and the low reactivity of α -olefins with the Phillips-type catalyst. For example, Cr/SiO₂ incorporates only 1.85 mol% of 1-hexene at a constant concentration in the reactor.¹

and 10 vol% H₂. The change in the gas composition is testified by a high rise of the signal with m/z values of 2 and 27 due to H₂ and C₂H₄ reactants, respectively. After the start of the reaction, the signals belonging to the fragments of the *in situ* produced 1-hexene slowly rise and remain constant until the feed of the reaction mixture was switched off. Furthermore, the TEAl-modified CS catalyst is able to oligomerise ethylene to 1-butene, although in smaller amounts than 1-hexene. At ~30 min of the reaction, a sudden increase of the signals suggests a temporary production of the oligomers with carbon numbers higher than 6. After 60 min of reaction, the gas feed was switched back to N₂. The signals due to C₂H₄ and H₂ immediately drop due to the abrupt change and continue to decline slowly together with the signals belonging to the oligomerisation products.

The origin of the fragments detected during the on-line MS analysis could be identified and confirmed by the GC-MS and GC analysis of the sampled gas phase (Fig. 2B). In this particular case when the CS catalyst was used, the main oligomer product detected was 1-hexene. Moreover, a small amount of the *in situ* produced saturated and unsaturated ethylene dimers and tetramers was identified. The GC and MS results unambiguously confirm the ethylene oligomerisation properties of the tested catalyst material.

Furthermore, the UV-vis-NIR DRS spectrometer integrated within the setup offered monitoring of the catalyst material during the course of the reaction. Analysis of the UV-vis part is suitable for inspecting the electronic spectra of supported transition metal ions, which allows determination of the oxidation number, coordination state and symmetry of a transition metal ion by studying d–d and charge transfer (CT) transitions.^{47,48} Moreover, the NIR part offers the possibility of examining vibrational properties *i.e.* combination bands and overtones of the vibrational modes of the materials and reaction products. The expected electronic and vibrational bands due to the silica-supported Cr and Ti species (if present inside the catalyst),^{14,16,49,50} and polymer⁵¹ produced therewith are summarised in Table 2.

During the pre-treatment of the CS catalyst with TEAl (Fig. 3A and B), the UV-vis-NIR DRS spectra of the catalyst show a small intensity increase of the bands at 16000 cm⁻¹ and 10 000 $\,cm^{-1}$ due to the $\,^4\!A_{2g}$ \rightarrow $\,^4\!T_{2g}$ and $\,^5\!E_g$ \rightarrow $\,^5\!T_{2g}$ d–d transitions of the formed Cr³⁺_{Oh} and Cr²⁺_{Oh} species. Due to the small amount of modified surface sites, the reduction is more evident after the subtraction of the fresh catalyst spectrum from the final spectrum of the sample after TEAl modification, as shown in Fig. 4. The negative bands at 21 000-22 000 cm⁻¹, 27 000-31 000 cm⁻¹ and 37 000-40 000 cm⁻¹ originating from the decrease of the CT bands of Cr⁶⁺ species are accompanied with the positive bands of the reduced Cr³⁺_{Oh} and Cr²⁺_{Oh} species. The addition of the cocatalyst can be clearly noted in the NIR region of the timeresolved spectra (Fig. 3B). The intensity increase of the bands at 4000-4400 cm⁻¹ is due to the combination modes of stretching and deformation vibrations of CH₂/CH₃ groups originating from the adsorbed alkyl groups of the TEAl and

Table 2 Assignment of the bands in the UV-vis-NIR DRS spectra of the Cr/Ti/SiO₂ Phillips-type polymerisation catalyst and polyethylene (PE)^{14,16,48-50}

Assignment	\tilde{v} (cm ⁻¹)		
$\sigma \rightarrow \sigma^* C - C$	~67 000		
$\pi \rightarrow \pi^* C = C$	52 000-59 000		
$O \rightarrow Ti^{4+} CT$	37 500-40 000		
$O \rightarrow Cr^{6+} CT$	37 600-39 600		
$O \rightarrow Cr^{6+} CT$	27 000-31 000		
$O \rightarrow Cr^{6+} CT$	21 000-22 000		
$Cr^{3+}_{Oh} d-d$	~16000		
$Cr^{2+}_{Oh} d-d$	~10000		
$3v_{as}(CH_2/CH_3)$	8423; 8233		
2v(SiOH)	7325		
2v(TiOH)	7270		
$2v_{as/s}(CH_2/CH_3) + \delta(CH_2/CH_3)$	7189; 7059; 6966; 6482		
$2v_{as/s}(CH_2/CH_3)$	5777; 5668		
$v(H_2O) + \delta(H_2O)$	5260		
$v(SiOH) + \delta(SiOH)$	4525		
$v_{as/s}(CH_2/CH_3) + \delta(CH_2/CH_3)$	4325; 4250		
$v_{\rm as}(\rm CH_2/\rm CH_3) + w(\rm CH_2/\rm CH_3)$	4185		

heptane solvents. At higher wavenumbers, the first overtone of the fundamental symmetric and asymmetric CH_2/CH_3 stretching vibration appears in the 5600–5900 cm⁻¹ region. Furthermore, the first stretching vibration overtone and deformation band can be observed in the 6900–7200 cm⁻¹ region. In the first overtone OH stretching region, consumption of the silanol groups at 7325 cm⁻¹ occurs due to the interaction with adsorbed TEAl and heptanes. After flushing with N₂, the OH functionality of the catalysts is mostly restored, giving almost identical spectra to those obtained before the treatment with TEAl, although a broadening and red shift of the associated absorption band can be noted.

At this stage, ethylene polymerisation reaction with the CS catalyst was started by changing the N₂ flow in the quartz reactor to the gas reactant mixture consisting of 45 vol% ethylene, 45 vol% N2 and 10 vol% H2. Fig. 5A shows the time evolution of the UV-vis-NIR DRS spectra in the UV-vis region during the in situ polymerisation of ethylene using the Cr/ SiO₂ (CS) catalyst. During the reaction, the intensities of the $O \rightarrow Cr^{6+}$ CT bands belonging to mono- and dichromates are decreasing due to the reduction of Cr⁶⁺ species by ethylene with a simultaneous increase of the d-d transition bands of Cr^{3+}_{Oh} and Cr^{2+}_{Oh} species at ~16000 cm⁻¹ and ~10000 cm⁻¹, respectively. Consumption of surface silanol groups is evidenced in the NIR region (Fig. 5B) by the disappearance of the OH stretching overtone and combination bands at ~7325 cm⁻¹ and 4535 cm⁻¹. On the other hand, the formation of the polymerisation product can be monitored in situ by the growing bands originating from the CH₂ and CH₃ of the growing polymer chain and possible shorter oligomers. These bands evolve in the regions of 4000-4400 cm⁻¹, 5600-5900 cm⁻¹ and $6900-7200 \text{ cm}^{-1}$ with the assignation given in Table 2.

From the on-line MS and *operando* UV-vis-NIR DRS data, we can identify two stages of the catalyst reaction. During the first stage, right from the start of the ethylene feed, the catalyst is being reduced. No bands due to the polymer product

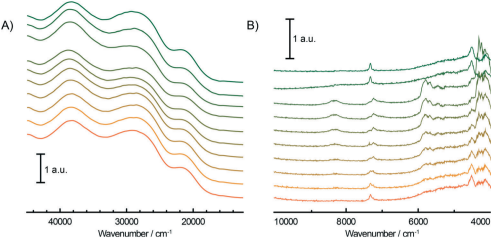


Fig. 3 Time evolution (from green to orange) of the operando UV-vis-NIR DRS spectra of the Cr/SiO₂ Phillips-type catalyst (CS sample) during the addition of TEAl in the UV-vis (A) and the NIR (B) region.

can yet be seen in the NIR region of the spectra. On the other hand, it is possible to observe a rise of the MS signal belonging to the fragments of the produced ethylene oligomers. In this stage, ethylene oligomerisation sites were being created and ethylene oligomerised mainly to 1-hexene. The second stage starts later and includes the production of heavier oligomers and polyethylene, as confirmed by the growing NIR bands of the CH_2 and CH_3 groups. It must be noted that the oligomerisation activity of the catalyst is still preserved during the later stage.

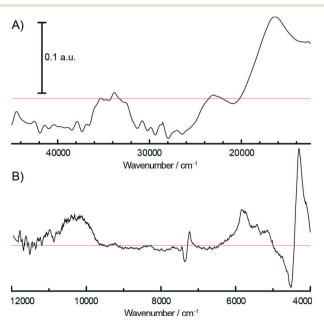


Fig. 4 Difference spectrum of the catalyst modified with TEAl and the fresh Cr/SiO_2 Phillips-type catalyst in the UV-vis (A) and the vis-NIR (B) region showing the reduction of some of the Cr^{6+} species as testified by the negative $O \rightarrow Cr^{6+}$ charge transfer bands and positive bands of the formed Cr^{3+} and Cr^{2+} species at approximately 16 000 and 10 000 cm⁻¹, respectively. Zero is presented with the red line.

UV-vis-NIR diffuse reflectance spectroscopy

In this study we have investigated the influence of Ti and H₂ on the oligomerisation properties of a TEAl-modified Phillips-type ethylene polymerisation catalyst and therefore tested the catalysts with different amounts of titanation (Table 1) and a different content of the gas reactant mixture. As was shown in the previous section, the UV-vis-NIR DRS spectra of the fresh CS catalyst show a highly dehydroxylated catalyst surface, evidenced by a sharp 2ν (SiOH) stretching overtone at ~7325 cm⁻¹ and a combination $v(SiOH) + \delta(SiOH)$ band at ~4525 cm⁻¹. The addition of Ti during the preparation of the catalyst materials introduces a 2v(TiOH) overtone band at 7270 cm⁻¹ as a shoulder of the silanol stretching band, with the intensity increasing with increasing Ti loading. This group increases the acidity of the catalyst surface and appears at lower wavenumbers as expected due to the increased reduced mass of the functional group. Another characteristic of the fresh catalysts is the presence of three O \rightarrow Cr⁶⁺ CT bands of mono- and polychromate species appearing in the UV-vis region. The exact positions of these absorption bands (Table 1) are dependent on the catalyst architecture since any changes in the electronic environment influence the energy levels of Cr. Titanation and an increased amount of Ti from CS to CTS2 cause a shift of the maxima of the UV-vis bands towards lower energy. This is triggered by the formation of Si-O-Ti-O-Cr bonds where Si is replaced with the less electronegative Ti atom, causing a slight shift of the electron cloud towards the O closest to Cr, populating its energy levels and narrowing the $O \rightarrow Cr^{6+}$ CT energy gap, therefore decreasing the energy required for the charge transfer to occur.52

Prior to ethylene polymerisation, all of the tested catalysts were pre-contacted with TEAl. Some of the Cr⁶⁺ sites were reduced to Cr³⁺ and Cr²⁺ species, as testified by a small intensity decrease of the O \rightarrow Cr⁶⁺ CT bands and the appearance of the ${}^{4}A_{2g} \rightarrow {}^{4}T_{2g}$ d–d transition band of Cr³⁺_{Oh} and the ${}^{5}E_{g} \rightarrow {}^{5}T_{2g}$ d–d transition band of Cr²⁺_{Oh}. Furthermore, TEAl also

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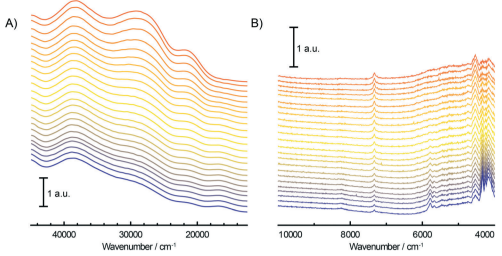


Fig. 5 Time evolution (from orange to blue) of the operando UV-vis-NIR DRS spectra of the fresh Cr/SiO₂ Phillips-type catalyst (CS sample) during the oligo-/polymerisation of ethylene, in the UV-vis (A) and the NIR (B) region.

reacts with the hydroxyl groups of the support, as shown by the broadening and lowering of the intensity of the silanol and titanol group stretching first overtone. However, these changes in the UV-vis-NIR DRS spectra are not substantial, suggesting that TEAI modifies only some of the Cr sites, which were still enough to produce ethylene oligomers.

Fig. 6 shows the evolution of the UV-vis-NIR DRS spectra during ethylene polymerisation using the CS and CTS2 catalysts, with and without H₂, after the modification with TEAl. During ethylene polymerisation, Cr⁶⁺ mono- and dichromate species become significantly reduced by the ethylene reactant to O_h coordinated Cr³⁺ and Cr²⁺ species.⁹ Some of these newformed sites may not only be the active ethylene polymerisation species, but also the products of deactivation of polymerisation sites as a slight shift of the 16000 cm⁻¹ band to higher energies indicates a possible formation of crystalline Cr₂O₃ on the catalyst surface. The influence of titanation on the polymerisation of ethylene can be clearly seen in the time evolution of the operando UV-vis-NIR DRS spectra. By changing the electronic and spatial environments around Cr sites, Ti gives rise to a new type of Cr⁶⁺ species, which prove to be more reducible than the Cr⁶⁺ species of the non-titanated catalyst. This effect is evidenced by a more rapid intensity decrease of the O \rightarrow Cr⁶⁺ CT bands in the case of CTS2 compared to CS. Consequently, easier reducibility causes a faster development of the polymerisation rate, which can be seen by the growth rate of the overtone and combination vibrational bands of the CH_x groups of the growing polyethylene. In the presence of H_2 (Fig. 6B and D), the reduction of Cr is even more pronounced due to the reducing properties of H₂.

The UV-vis absorption bands of the oligo/polymerisation products could not be measured, since they appear outside of the detectable range of the UV-vis-NIR DRS spectrophotometer. Absorption, due to the $\sigma \rightarrow \sigma^*$ transition of saturated hydrocarbons, is expected at ~67000 cm⁻¹. Additional unsaturation of the C bond, in the case of alkene species,

introduces an additional $\pi \rightarrow \pi^*$ transition band in the 52 000–59 000 cm⁻¹ range.⁵³

Gas chromatography analysis of the reaction products

Although *operando* UV-vis-NIR DRS is a spectroscopic technique useful to study both the catalyst and the product during the polymerisation reaction, it cannot discriminate between different types of products due to the overlap of NIR bands appearing at the same wavenumbers. Fig. 7–9 show the GC analysis results of the gas phase sampled during the polymerisation of ethylene using several different catalysts, with varying Ti loading (0, ~2 and ~4 wt% Ti), Cr loading (0, ~0.5 and ~1 wt% Cr) and activation temperature (823, 923 and 1048 K), as summarised in Table 1. Each catalyst was tested both with and without 10 vol% H₂, while the ratio of ethylene and N₂ was kept at 1:1.

For the sake of excluding any ethylene oligomerisation activity of the silica carrier or the titania on the silica carrier,‡ two additional materials without any Cr were tested, *i.e.* wide-pore silica support (S) and 4 wt% titanated silica (TS), both activated using the same procedure as the actual catalyst. During ethylene polymerisations using these materials with H_2 and the TEAl co-catalyst, no substantial ethylene oligomerisation was reported (Fig. 7A and 8A).

For the actual Phillips-type catalysts, it was observed that the concentration of the produced ethylene oligomers greatly depends on the composition of the catalyst. The highest yield of all oligomers was obtained with the CS catalyst containing no Ti, and it decreases with increasing Ti loading. Detailed analysis of the amounts of olefinic and paraffinic fractions

[‡] Supported Ti alkoxides on silica reduced by a metal alkyl co-catalyst are reported as active catalysts for ethylene dimerisation.⁶⁵ In the case of our catalysts, we assume that virtually all alkoxide groups have been burnt off during the calcination step.

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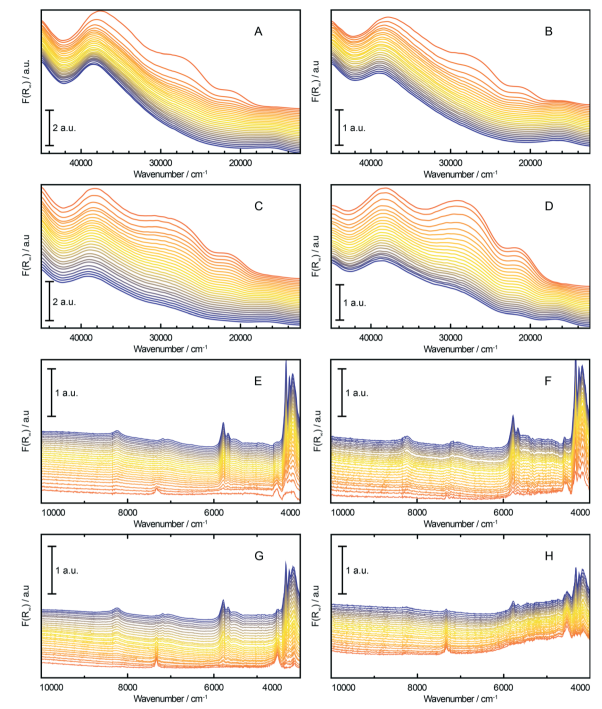


Fig. 6 UV-vis part of the *operando* UV-vis-NIR DRS spectra of CTS2 and CS Phillips catalysts during polymerisation of ethylene without H_2 (A and C) and with H_2 (B and D), respectively, after pre-reacting the catalyst with the TEAl co-catalyst. NIR part of the *operando* UV-vis-NIR DRS spectra of CTS2 and CS Phillips catalysts during polymerisation of ethylene without H_2 (E and G) and with H_2 (F and H) after pre-reacting the catalyst with the TEAl co-catalyst. Time evolution of the spectra is indicated by colour transition from orange to blue with 2 minutes between each successive spectrum.

shows that the yield of olefins significantly drops with increasing Ti loading, while the yield of paraffins is slightly increased. The addition of H_2 to the reaction mixture caused an increase in the overall oligomer yield, with a higher influence on the production of saturated oligomers. Therefore, H_2 might be involved in the termination of polymer chain

growth at the early stages, increasing the concentration of saturated oligomers in the gas phase.

Fig. 7B and 8B show a detailed distribution of the *in situ* produced olefinic and paraffinic hydrocarbons, respectively, normalised to 100%. Firstly, a clear dependence of the ratio of butenes, hexenes and octenes can be observed, depending



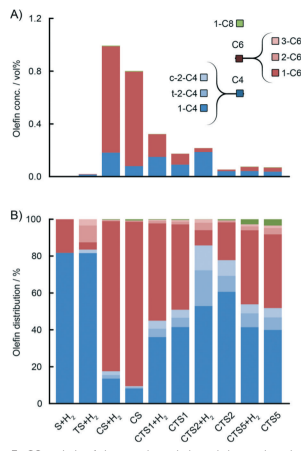


Fig. 7 GC analysis of the gas phase during ethylene polymerisation with various Phillips-type catalysts showing different gas phase concentrations (A) of the *in situ* produced C4 (blue), C6 (red) and C8 (green) olefins depending on the catalyst composition. Olefin distribution normalised to 100% (B) reveals that the titanation of the Phillips-type catalysts has a detrimental effect on ethylene oligomerisation and selectivity towards industrially important 1-hexene fraction.

on the amount of Ti present in the catalyst. The CS system, without any Ti, exhibited the highest selectivity towards ethylene trimerisation, predominantly producing 1-hexene. The addition of Ti caused a further decrease in the overall ethylene trimerisation selectivity as can be noted for the CTS1 catalyst with ~2 wt% Ti. The most pronounced Ti effect is observed for the catalyst with the highest Ti loading of ~ 4 wt% where a strong inhibition of ethylene trimerisation sites diminished the amount of hexenes compared to the higher yield of butenes and octenes. In the reaction with the CTS5 catalyst, which contains double the amount of Cr loading compared to the other examined catalysts, the distribution of olefins matches the distribution produced with the CTS1 catalyst of a similar Cr-to-Ti ratio. The distribution of paraffins shows that the relative ratio of the in situ produced *n*-butane, *n*-hexane and *n*-octane is independent of the Ti loading in the catalyst and that the differences can mainly be attributed to the presence of H₂ in the reactant gas mixture. Overall, a higher Ti loading decreases the olefin-to-paraffin ratio within the ethylene oligomers of all chain lengths.

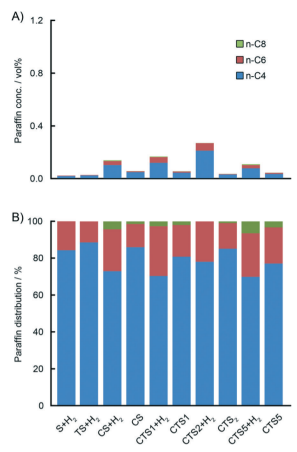


Fig. 8 GC analysis of the gas phase during ethylene polymerisation with various Phillips-type catalysts showing different gas phase concentrations (A) of the *in situ* produced C4 (blue), C6 (red) and C8 (green) paraffins depending on the catalyst composition. Paraffin distribution normalised to 100% (B) reveals that the titanation of the Phillips-type catalysts does not change significantly the ratio of produced oligomers.

A higher activation temperature of the catalyst showed the promoting effect on the *in situ* ethylene oligomerisation as can be seen by the increasing amount of produced oligomers starting from CTS4, over CTS3 to the CTS2 catalyst, activated in dry air at 823 K, 923 K and 1048 K, respectively (Fig. 9). A higher activation temperature improves the dehydroxylation of the catalyst surface, liberating the coordination sphere of Cr producing more reactive Cr sites, which increase the gas phase oligomer yield. A rise in the activation temperature also shifts the olefin distribution towards butenes and promotes the isomerisation and production of linear β - and γ -olefins.

Fig. 10 shows the total distribution of hydrocarbon oligomers, split into olefin and paraffin contributions, which can be further presented as Schulz–Flory (SF) diagrams.† A clear 1-hexene spike can be observed with the CS catalyst with no Ti. An increase of the Ti loading significantly decreases the 1-hexene selectivity. If the products follow the SF distribution, caused by the linear-insertion mechanism, a straight curve can be expected with the slope, which represents the chain-growth probability α , determined by the rates of chain



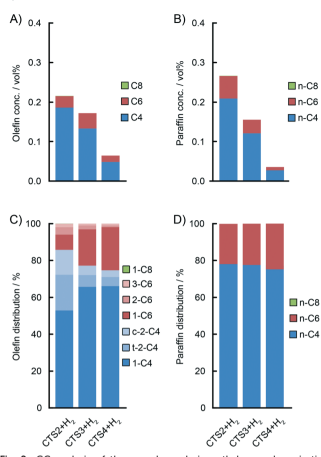


Fig. 9 GC analysis of the gas phase during ethylene polymerisation with CTS2, CTS3 and CTS4 Cr/Ti/SiO₂ Phillips-type catalysts activated at 1048 K, 923 K and 823 K, respectively. Decrease of the activation temperature lowers the yields of the *in situ* produced C4 (blue) and C6 (red) olefins (A) and paraffins (C). A lower activation temperature slightly shifts the distribution of *in situ* produced olefins towards α -olefins, particularly 1-hexene (B), having practically no influence on the distribution of paraffins (D).

growth and termination. The distribution of the *in situ* produced paraffins exhibits an exponential decay with the increase of carbon number, which is reflected in linear SF curves with α values given in Table 3. Addition of 10 vol% H₂ to the reactant mixture increases α and therefore decreases the selectivity to shorter paraffin oligomers, *i.e. n*-butane. On the other hand, the olefinic oligomers do not show a linear trend in the SF plot in the case of the CS catalyst containing no Ti. With increasing Ti loading, due to the decrease in 1-hexene selectivity, the curves become more linear.

Similar trends can be observed with the CTS5 catalyst with the increased Cr loading of ~1 wt% (Fig. 11). A higher Cr loading, while keeping the amount of Ti constant, increases the number of Ti-free Cr sites, which are able to produce selectively ethylene trimers, as illustrated by the nonexponential decay of olefins and the presence of 1-hexene spike. This catalyst sample still contained Ti, which explains the higher amount of ethylene dimers. Nevertheless, the increase in Cr loading appeared to be beneficial for the promotion of ethylene trimerisation.

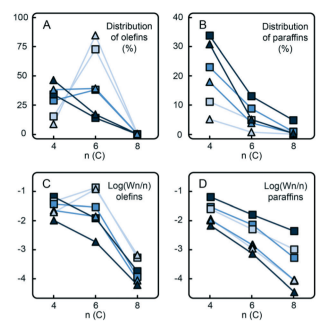


Fig. 10 The increase of Ti loading in the Phillips-type catalyst (from light to dark blue) shifts the selectivity towards C4 at the expense of C6 products in the experiments with (\Box) and without (Δ) hydrogen. A different trend in the distribution of olefinic species (A) compared to paraffinic species (B) is also clearly visible in the Schulz–Flory plots. While paraffinic species show linear curves independent of the Ti loading (D), olefin species exhibit a discrepancy from linearity with increasing Ti loading (C), indicating an oligomerisation mechanism, which does not follow SF distribution.

Table 3 Chain-growth probability factor α for the paraffinic oligomers produced in ethylene oligomerisation reactions

Catalyst name	α , no H ₂	α , with H ₂
CS	0.30	0.45
CTS1	0.33	0.37
CTS2	0.27	0.51
CTS5	0.38	0.46

Discussion

The analyses of the gas phase collected during the ethylene polymerisation show that the titanation of the Cr/SiO_2 Phillips-type catalyst significantly reduces the yield of ethylene oligomers and shifts the selectivity towards butene fractions. A lower Ti-to-Cr loading ratio appeared to be beneficial for the promotion of the trimerisation of ethylene, as the number of Cr sites, which are not inhibited by Ti, is increased and therefore able to selectively produce more ethylene trimers, indicating a change in the oligomerisation pathway.

Currently, the most widely accepted proposed mechanism of ethylene polymerisation is the Cossee–Arlman mechanism, which explains the chain propagation by linear insertion of the ethylene monomer into an existing alkyl group or polymer, bonded to a Cr active site. The termination of the chain occurs *via* β -H elimination leaving a vinyl end group in the released hydrocarbon product. As a result of the mentioned

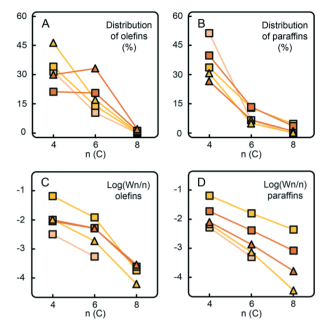
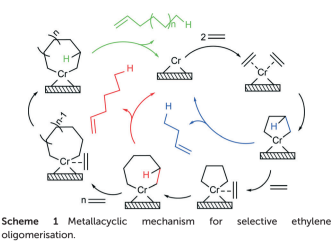


Fig. 11 The increase of Cr loading in the Phillips-type catalyst (from light yellow to orange) increases the selectivity towards C6 products in the experiments with (\Box) and without (Δ) hydrogen. A different trend in the distribution of olefinic species (A) compared to paraffinic species (B), also visible in the Schulz–Flory plots (C and D), causes a discrepancy from linearity with increasing Cr loading. A higher amount of Cr, while keeping the Ti loading constant, increases the amount of ethylene trimerisation sites, which produce higher amounts of 1-hexene.

ethylene insertion mechanism, the products are expected to follow a statistical hydrocarbon distribution, known as the Schulz–Flory (SF) distribution, with only even-membered hydrocarbon chains due to the presence of ethylene as the only monomer. However, the high 1-hexene spike produced with the non-titanated catalyst can be attributed to the complex structure of the catalyst, with different types of active sites and a complex mechanism of oligomerisation, which cannot be explained solely by the Cossee–Arlman mechanism of statistical chain growth.

From the results of our study, it is found that two types of distinct oligomerisation mechanisms occur simultaneously with the extent between each other depending on the precise catalyst compositions and hence the electronic environment of the different active sites. Besides the Cossee-Arlman mechanism of linear insertion, in the case of alkane oligomers and polyethylene, alkene species are predominantly being produced via a metallacyclic mechanism (Scheme 1). Two ethylene molecules are able to coordinate to the vacant coordination sphere of the Cr active site and form a chromacyclopentane ring. Two processes can occur at this stage, β -H transfer and release of 1-butene, or coordination and insertion of another ethylene monomer to form a chromacycloheptane ring. This process can continue even further and form bigger chromacyclic rings, where the release of olefin *via* β -H transfer occurs when the probability of insertion of a new monomer becomes lower than the probability of β -H transfer. The stability of the chromacyclic ring is highly



influenced by the structure of Cr active sites. As revealed in the operando UV-vis-NIR DRS spectra, titanation of the catalyst changes the structure of the Cr oligomerisation sites by forming Si-O-Ti-O-Cr linkages, as testified by the red shifts of $O \rightarrow Cr^{6+}$ CT bands, and introduces free surface titanol groups, which increase the acidity of the catalyst surface. Consequently, an increasing Ti loading decreases the stability of the 5-membered chromacyclopentane ring, favouring β -H transfer and release of 1-butene and therefore reducing the selectivity towards 1-hexene. Alternatively, given the DFT studies of Robinson et al.⁵⁴ and the work of Emrich et al.,⁵⁵ the high 1-butene selectivity in the case of the titanated catalysts can be explained by the Cossee-Arlman mechanism over the oligometrisation sites with a very low α , while 1-hexene is still being produced over trimerisation sites via the metallacyclic mechanism.

Ethylene trimerisation catalysts have been reported for chromium-based homogeneous systems, basically consisting of multidentate ligands reacted with chromium salts and activated with aluminium alkyl co-catalyst, some of them being commercialised.⁵⁶⁻⁵⁹ The isotopic labelling and regiochemistry studies of Agapie et al.^{60,61} on such catalytic systems strongly support the metallacyclic mechanism of the selective ethylene trimerisation and tetramerisation, which might occur through Cr⁺/Cr³⁺ and Cr²⁺/Cr⁴⁺ redox cycles as shown by XAS and EPR.62-64 Similarly, the metallacyclic mechanism was proposed for the silica-supported chromium catalyst discussed in this paper, although it has to be noted that a heterogeneous catalyst is more difficult to define due to the heterogeneity of different surface chromium sites. The advantage of the catalyst material examined in this study is its multifunctionality, i.e. the selective ethylene oligomerisation capabilities of some of the chromium sites and co-/polymerisation activity of other sites, within a single Phillips catalyst.

Conclusions

The oligomerisation properties of a TEAl-modified Phillipstype ethylene polymerisation catalyst were tested in a

specially developed operando setup allowing the relation of data gathered by UV-vis-NIR diffuse reflectance spectroscopy, mass spectrometry and gas chromatography techniques. This combination of methods allowed a real-time study of working catalysts and reaction products. The incorporation of Ti in the Cr/SiO₂ Phillips-type catalysts by the formation of Cr-O-Ti-O-Si linkages alters the electronic environment of Cr ions and changes the oligomerisation capabilities of the catalyst. Consequently, different distributions of the in situ produced ethylene oligomers were revealed, highly dependent on the architecture of the catalyst. Paraffinic oligomers show an exponential decay explained by the Cossee-Arlman linear insertion mechanism. On the contrary, olefinic oligomers in the case of the non-titanated Cr/SiO₂ catalyst exhibit a much more complex distribution, with a higher selectivity towards 1-hexene than what is expected from the linear insertion mechanism. High ethylene trimerisation activity is a consequence of the metallacyclic oligomerisation mechanism. The addition of Ti modifies the trimerisation Cr sites and creates new oligomerisation sites, which favour β-H transfer in the chromacyclopentane intermediate and release of the 1-butene product, therefore decreasing the selectivity towards 1-hexene. These results open new possibilities for both academic and industrial research of the Phillips polymerisation system, as they show that by a proper selection of the catalyst structure it is possible to change the distribution of the in situ produced oligomers and therefore change the amount of α -olefins, which are able to co-polymerise with ethylene to form the desired grade of polyethylene.

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