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Comparison of PGSE NMR and ESI-MS measurements on methylaluminoxane[†]

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PGSE NMR and ESI-MS studies of different grades of hydrolytic methylaluminoxane (MAO) demonstrate a correlation between the average dimensions and anion distribution of MAO. This correlation is revealed through aging and gelation studies of a commercial formulation of MAO. Formulations featuring an anion distribution skewed to higher *m/z* ratios have been observed to contain significantly higher activator contents as measured by ¹H NMR spectroscopy, despite being otherwise very similar. PGSE NMR studies in toluene vs. chlorobenzene media demonstrate that the average hydrodynamic dimensions of MAO are largely unaffected by solvent polarity, although aggregation behavior is somewhat sensitive to solvent polarity. As for catalyst activation and ion-pair speciation, the situation in polar solvents is complicated by the formation of solvated cations (and anions) in chlorobenzene which dramatically lower dimensions. DFT studies of model aluminoxane structures in the size range of MAO featuring a variety of architectures, molar masses and Me₃Al contents reveal a linear correlation between *D*_t as measured by PGSE NMR, and molar mass using simple relationships and estimates of molecular volume if suitable high molecular weight standards are available. There is reasonable agreement in molar mass with available ESI-MS data, recognizing that MAO is not monodisperse.

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Introduction

The structure and properties of methylaluminoxane (MAO)^{1,2} continue to attract interest in the context of olefin polymerization using metallocene catalysts.^{3–5} After nearly 50 years of research, the basic mechanism of action of MAO is clear, yet the structure of the reactive components and activation mechanisms are not fully understood.

Theory suggests that the bulk of MAO consists mainly of relatively unreactive structures built up of tetrahedral O₃AlMe and trigonal O groups in six-membered or larger (MeAlO)_n rings.^{6,7} Reactive structures consist of strained cages, analogous to those adopted by *t*-butylaluminoxanes,⁸ especially

those with vicinal (MeAlO)₂ rings,⁹ extended cages,¹⁰ nanotubes^{11,12} and two-dimensional sheet models¹³ with reactive sites incorporating structural Me₃Al.¹⁴

Recently, a component of this complex mixture has been finally isolated from hydrolytic MAO (h-MAO) and characterized by X-ray crystallography.¹⁵ It has a sheet structure with the formula (MeAlO)₂₆(Me₃Al)₉ (hereinafter 26,9) and a molecular weight of 2156 g mol^{−1} and was used to activate metallocenes.

The average molecular weight and presumably the molecular weight distribution of h-MAO¹⁶ are known to be important in metallocene catalyst activation. Low MW species such as (Me₂AlOAlMe₂)_n (*n* = 3 to 4)¹⁷ are known to be ineffective,¹⁸ while early synthetic methods emphasize the formation of higher MW materials as confirmed through cryoscopic measurements.¹⁹ Cryoscopic measurements of commercial h-MAO (W. R. Grace)²⁰ suggest a (number) average MW (*M*_n) of *ca.* 1100 g mol^{−1} while small angle neutron scattering (SANS) studies of h-MAO (Chemtura/Lanxess) suggest a higher (weight) average MW (*M*_w) of 1803 ± 55 g mol^{−1} with a spread in values between 1400 and 2800 g mol^{−1} depending on the free Me₃Al content (*M*_w = 1803 corresponds to *ca.* 75% of the total Me₃Al content for the sample in question).²¹ The same study suggested a roughly spherical shape for bulk MAO based upon *r*_g = 9.3 and *r*_H = 11.7 Å as measured by SANS and PGSE NMR, respectively. Both estimates were in good agreement with earlier studies of *T*₁ relaxation (*r*_g = 9.7 Å)²² and [Cp₂ZrMe₂AlMe₂][MAO(Me)] (1) ion-pair size (*r*_H = 12–12.5 Å)

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[†]Electronic supplementary information (ESI) available: SI.pdf – [Me₂Al(THF)₂]⁺ and Me₃Al content of MAO samples and representative ¹H NMR spectra. Additional ESI-MS spectra. Electronic energies and thermodynamic quantities for the MAO structures. PGSE NMR signal intensity decay data and non-linear regression analyses. Table S-2.xyz – Coordinates for all new theoretical MAO structures reported. MAO.xlsx – ESI-MS *m/z* ratio vs. intensity (counts) and *m/z* averages of h-MAO and MMAO 12. See DOI: <https://doi.org/10.1039/d4qi02982h>



by PGSE NMR.²³ However, the interpretation of the PGSE NMR data in terms of molar mass (50–60 Al atoms for MAO²¹ vs. 150–200 Al atoms for ion-pair **1**²³) differed significantly between the two studies.

The source and Me_3Al content of MAO are important variables as revealed by PGSE NMR studies on h-MAO (Sigma Aldrich). In particular, dried or chemically modified MAO (DMAO and MAO-BHT) and the ion-pairs formed from Cp_2ZrMe_2 and DMAO had significantly larger dimensions than the h-MAO itself.²⁴ Limiting values of $r_{\text{H}} = 8.6$ and 10.3 \AA for this MAO and ion-pair **1** were significantly lower than estimates from other studies where the sources of h-MAO were different.

PGSE NMR experiments yield translational diffusion coefficients D_{t} in the solvent used but require the use of an internal standard, either deliberately added or the solvent itself, to extract size information using the Stokes–Einstein equation or variations (e.g. eqn (1), where k = Boltzmann constant, T = temperature, c and f_s are the size and shape-dependent factors, η = solution viscosity, and r_{H} = hydrodynamic radius).²⁵ However, the relationship between the hydrodynamic volume and molar mass is never straightforward.

$$D_{\text{t}} = \frac{kT}{cf_s\pi\eta r_{\text{H}}} \quad (1)$$

A more useful approach is to directly relate the measured diffusion coefficients to molar mass, by establishing empirical correlations between D_{t} and species of known molar mass and similar density.²⁶ In the case of lower MW solutes, a relationship incorporating both size and molar mass has been established (eqn (2)),²⁷ and it appears to be applicable to MAO.²⁸

$$\frac{D_1}{D_2} = \left(\frac{2}{1 + r_2/r_1} \right)^2 \left[\frac{1}{2} \left(1 + \frac{m_1}{m_2} \right) \right]^{1/2} \quad (2)$$

Here D_1 and D_2 refer to the diffusion coefficients of the solute and solvent (or standard), while r_i represents the van der Waals radii of these two molecules and m_i represents the corresponding molar masses. It should be noted from the original study establishing the correlation in eqn (2) that the masses of the solutes and solvents were on the same order of magnitude.²⁷ This is generally not true for MAO solutions.

The widespread availability of NMR spectrometers with pulse gradient capability suggests an accessible and powerful approach to MAO or ion-pair speciation,²⁹ particularly if the results could be directly related to absolute molar mass.

Mass spectrometric characterization of MAO dates back to early methods of its synthesis, but only volatile, lower molecular weight components were detected using EI-MS.³⁰ Detection and characterization of high MW ions in h-MAO using ESI-MS³¹ and a donor such as octamethyltrisiloxane (OMTS) in a polar solvent (fluorobenzene) were reported in 2013.³²

This initial study involved the analysis of h-MAO from Sigma Aldrich (Fig. 1a). The Lewis base donor OMTS reacts with MAO according to Scheme 1^{10,14,33} where n, m and $[n, m]^-$

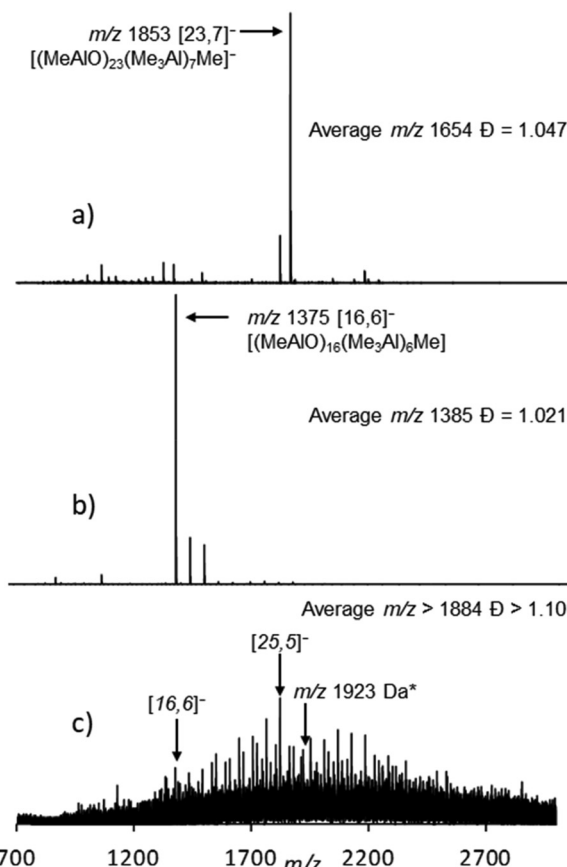


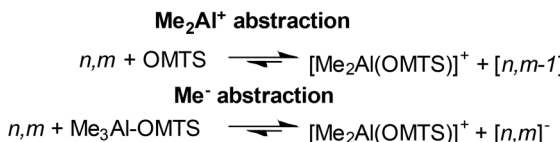
Fig. 1 Negative ion mass spectra of hydrolytic MAO: (a) 10 wt% Sigma Aldrich, (b) 10 wt% W. R. Grace and (c) MMAO 12 (10 wt% Sigma Aldrich) in the presence of 10.0 mol% OMTS in PhF solution with $[\text{Al}] \approx 0.01 \text{ M}$. Assignments are based on MS/MS experiments and nominal mass (a and b) or just nominal mass (c). * The peak with m/z 1923 Da is assigned to the $[(\text{MeAlO})_{24}(n\text{-C}_8\text{H}_{17}\text{AlO})(\text{Me}_3\text{Al})_5\text{Me}]^-$ anion. For details, see ref. 32, 34 and 39.

are shorthand notations for $(\text{MeAlO})_m(\text{Me}_3\text{Al})_n$ and $[(\text{MeAlO})_m(\text{Me}_3\text{Al})_n\text{Me}]^-$, respectively.

Both the cations and anions formed can be separately detected by ESI-MS. The individual anions can be analyzed by MS-MS to determine their formula (i.e. $[n, m]^-$). In the case of Sigma Aldrich MAO, the spectrum was dominated by an anion with m/z 1853 with the formula $[23,7]^-$. We also provide here the number-average m/z ratio $\overline{(m/z)}_n = \sum (m/z)_i I_i / \sum I_i$ and polydispersity $D = \overline{(m/z)}_w / \overline{(m/z)}_n$ of this mixture where $(m/z)_i$ and I_i are taken from the raw MS data files of ion intensity (I) vs. m/z ratio. The average m/z ratio and principal anion present in MAO from Sigma Aldrich were in good agreement with SANS estimates of \bar{M}_w for h-MAO from Chemtura.²¹

More recently, 10 wt% h-MAO from W. R. Grace was analyzed by ESI-MS and the results were quite different.³⁴ At the same additive level (10 mol% OMTS) the Grace material featured significantly lower m/z anions (Fig. 1b) with a significantly lower average m/z ratio, while the appearance of these spectra was very sensitive to the amount of OMTS present. This study also revealed that if h-MAO solutions from





Scheme 1 Ion-pair formation involving MAO and OMTS.

W. R. Grace were stored at room temperature, the anion distribution slowly changed from that shown in Fig. 1b to one similar to that shown in Fig. 1a.

Storage of h-MAO solutions at room temperature is accompanied by gelation and eventual loss of activity (aging).^{1,2,35} The stability to and extent of gelation are dependent on the total Al concentration and the amount of Me₃Al present. With the benefit of hindsight, the material from Sigma Aldrich (which always required filtering to remove gel particles prior to analysis by ESI-MS) had been aged at room temperature prior to and during shipment, as confirmed by this vendor in 2014 for this material sold in North America.³⁶

Much more recently, MMAO 12 (Sigma Aldrich), a 95:5 copolymer of methyl- and *n*-octylaluminoxane,³⁷ was analyzed by low- and high-resolution MS using an inert atmosphere sampling technique.³⁸ Families of ions with *m/z* ratios <1000 Da were detected and they featured hydrolysis, possible pyrolysis, and no evidence for incorporation of *n*-octyl groups.

The same material was also earlier analyzed by ESI-MS using 10 mol% OMTS.³⁹ A broad continuum of anions⁴⁰ with *m/z* ranging from ~1000 to >3000 Da were detected with evidence of *n*-octyl substitution (Fig. 1c) while the corresponding positive ion spectrum showed the [Me₂Al(OMTS)]⁺ and [(*n*-C₈H₁₇)MeAl(OMTS)]⁺ cations in relative amounts (ca. 97:3) consistent with the composition of MMAO 12.

It is evident from the foregoing discussion that a variety of different MAO samples have been analyzed by PGSE NMR, ESI-MS and other techniques (e.g. SANS) which are sensitive to or measure molar mass. Unfortunately, none of these studies involved the analysis of the same sample of MAO by all of these techniques. Also, given the difficulty in relating hydrodynamic dimensions from PGSE NMR directly to molar mass, we wished to establish a linear correlation between *D_t* and MW for MAO as in the work of Zaccaria *et al.* for other materials.

To address these issues we analyzed MAO from W. R. Grace by PGSE NMR spectroscopy as we have extensive experience with the ESI-MS analysis of that material, both 10- and 30 wt% formulations.^{10,34} These formulations have very similar ESI-MS spectra at similar OMTS levels, while different batches also have very similar spectra as long as they are shipped and stored refrigerated (as they were in all of our prior work – see the ESI† for examples).

We were also interested in whether the metallocene ion-pairs formed from this material had similar dimensions as seen in earlier work, and whether their dimensions were sensitive to solvent polarity as the ESI-MS work, unlike previous PGSE NMR studies, was conducted in a polar solvent.

Finally, we have also studied the aging/gelation of MAO by both ESI-MS and PGSE NMR and show that this process can be studied by either technique. We also establish that h-MAO with different anion distributions can have significantly different activator content.

Results and discussion

Relationship between molar mass and *D_t* for MAO

It is evident, based upon both this work and previous literature studies of MAO and the ion-pairs derived from it, that a reliable estimate of molar mass from *D_t* values is needed to relate PGSE NMR experiments to the limited ESI-MS (or MS) data that are available. The linear correlation reported in ref. 26 is not appropriate in the sense that the density of the compounds used to construct it were all $\leq 1 \text{ g mL}^{-1}$ while MAO and the ion-pairs derived from it are expected to have higher densities. For example, the crystalline density of 26,9 is 1.26 g mL^{-1} although toluene is incorporated into the lattice.¹⁵

For the calibration, we selected representative structures for MAO comprising reasonably stable sheets, cages and nanotubes varying in molar mass between 750 and 2000 g mol^{-1} , including two unreported sheet and cage structures corresponding to the neutral compound 23,7 and some 16,4 tube structures located at the M06-2X/TZVP level of theory. We also included the structure of the 26,9 sheet¹⁵ after establishing the correlation reported here. The structures of any unreported aluminoxane models are depicted in Fig. 2. We use the poly-

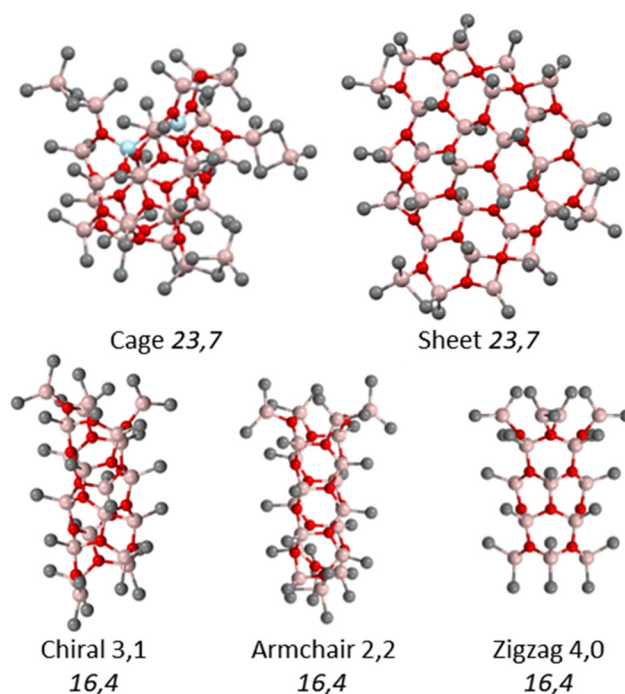


Fig. 2 Structures of some new models of MAO (Al pink, O red, C gray) with H-atoms omitted for clarity. Tetrahedral AlO₄ groups are highlighted in light blue for the cage 23,7 structure. For coordinate files and energies, see ESI.†



hedral octasilsesquioxane (POSS) compounds studied previously by PGSE NMR^{21,26} as standards based on their similar shapes, high molar mass and calculated densities.

The properties of the aluminoxane models, along with those of the POSS standards are summarized in Table 1. Their van der Waals volumes (V_{vdw}) and those of the reference compounds were estimated using the Connolly method⁴¹ with a probe radius of 1.4 Å as implemented in Chem3D.⁴² We chose this convenient approach as the Connolly solvent excluded volumes of the compounds, cholesterol, camphor, neopentane and methane were in very good agreement with the van der Waals volumes previously estimated by the method of Bondi.⁴³ The calculated densities for the aluminoxane models including that for sheet 26,9 vary between 1.55 and 1.72 g mol⁻¹ (Table 1, entries 1–14) while the POSS standards have somewhat lower calculated densities (entries 15–17).

It should be noted that the experimental liquid densities reported for $(i\text{-BuSi})_8\text{O}_{12}$ and $(i\text{-OctSi})_8\text{O}_{12}$ are 1.13 and 1.01 g mol⁻¹.⁴⁴ Our calculated densities are significantly higher as the V_{vdw} value is smaller than molar volume and does not include free volume.⁴³

The best correlation was obtained using the $\text{Ti}[\text{O}_2(i\text{-Bu}_8\text{Si}_8\text{O}_{11})_2]$ standard ($R^2 = 0.99_{85}$, Fig. 3) although good correlations with $R^2 = 0.99_{79}$ or 0.98_{76} were also obtained using $(i\text{-C}_8\text{H}_{17}\text{Si})_8\text{O}_{12}$ ⁴⁵ or $(i\text{-BuSi})_8\text{O}_{11}(\text{OH})_2$, respectively. Only the Ti POSS complex actually lies close to the correlation line for the MAO models due to its similar calculated density.

Going forward we will make use of this correlation both with respect to PGSE NMR studies on MAO from W. R. Grace and discussion of prior literature work where appropriate. Typical experimental errors in the determination of D_t (and thus r_H) are about $\pm 3.0\%$ so one can expect the same error in calculated MW.

As an example, original PGSE NMR data for 30 wt% Chemtura MAO had $D_t = 3.1 \pm 0.1 \times 10^{-10} \text{ m}^2 \text{ s}^{-1}$ at a limiting concentration of 380 mM (see the next section for studies of

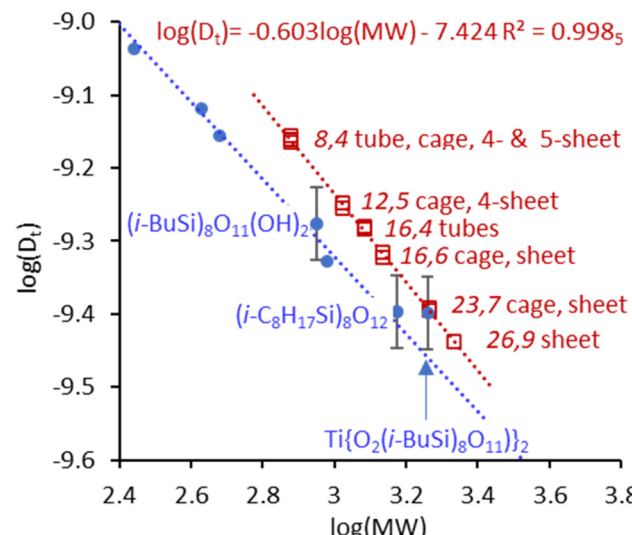


Fig. 3 $\log(D_t)$ vs. $\log(\text{MW})$ for some models of MAO using $\text{Ti}[\text{O}_2(i\text{-Bu}_8\text{Si}_8\text{O}_{11})_2]$ as a reference to construct the red calibration line. Data for POSS compounds reported in ref. 21 and 26 are shown in blue with the correlation established in ref. 26 shown as a dashed blue line.

the effect of MAO concentration). Our correlation gives $\text{MW} = 2865 \pm 92 \text{ g mol}^{-1}$ which is statistically lower but in reasonable agreement with the earlier estimate based on V_H (ca. 3360 g mol⁻¹).²¹ Both estimates are much higher than either the number- or weight-average estimates given in the literature, though the latter depends on free vs. associated Me_3Al content,²¹ while the former is dependent on the amount of low MW materials present including Me_3Al .²⁰

It is important to mention that the average MW determined by PGSE NMR spectroscopy (or other techniques used to study diffusion, such as light scattering in the case of large macro-

Table 1 Properties of models for MAO and POSS standards

Entry	Structure	ΔE^a (kJ mol ⁻¹)	MW (g mol ⁻¹)	V_{vdw}^b (Å ³)	$\rho_{\text{calc.}}^c$ (g mL ⁻¹)	D_t^d
1	8,4 5C-sheet	30.2	753.48	792	1.58	10.5
2	8,4 4C-sheet	48.1	753.48	809	1.55	10.4
3	8,4-cage	24.3	753.48	760	1.65	10.8
4	8,4-tube	0.0	753.48	786	1.59	10.6
5	12,5 sheet	7.3	1057.63	1112	1.58	7.5
6	12,5 cage	0.0	1057.63	1061	1.65	7.8
7	16,4 3,1 tube	18.6	1216.59	1201	1.68	6.8
8	16,4 2,2 tube	55.3	1216.59	1171	1.72	6.9
9	16,4 4,0 tube	93.7	1216.59	1202	1.68	6.8
10	16,6 sheet	9.4	1360.77	1367	1.65	5.9
11	16,6 cage	0.0	1360.77	1439	1.57	5.7
12	23,7 sheet	77.2	1838.96	1895	1.61	4.1
13	23,7 cage	0.0	1838.96	1835	1.66	4.2
14	26,9 sheet ¹⁵		2156	2253	1.59	3.7
15	$(i\text{-BuSi})_8\text{O}_{11}(\text{OH})_2$		891.6	1095	1.35	5.3
16	$(i\text{-C}_8\text{H}_{17}\text{Si})_8\text{O}_{12}$ ⁴⁵		1493.6	1771	1.25	4.0
17 ^e	$(i\text{-C}_8\text{H}_{17}\text{Si})_8\text{O}_{12}$		1332.5	1662	1.33	4.0
18	$\text{Ti}[\text{O}_2(i\text{-Bu}_8\text{Si}_8\text{O}_{11})_2]$		1828.1	1934	1.57	4.0

^a Electronic energy differences at the M06-2X/TZVP level of theory. For complete energies, see Table S1 in the ESI† and for coordinates, see ESI, Table S1.xyz.† ^b van der Waals volume calculated using a probe radius of 1.4 Å and Chem3D. ^c Density calculated from molar mass and V_{vdw} . ^d Experimental or calculated diffusion coefficient ($10^{-10} \text{ m}^2 \text{ s}^{-1}$) estimated from eqn (2) using $\text{Ti}[\text{O}_2(i\text{-Bu}_8\text{Si}_8\text{O}_{11})_2]$ as a reference. ^e Molar mass and volume data from ref. 21.



molecules) is different for polydisperse materials such as synthetic polymers, including MAO. Studies of synthetic polymers have shown that the diffusion average MW (\bar{M}_D) lies between the weight- and Z-average values of the molecular weight distribution but lies closer to \bar{M}_w .⁴⁶ Thus, it is not unexpected that an \bar{M}_D value for commercial MAO is higher than either the number- or weight-average values measured by other techniques.

Characterization of 10 wt% MAO by PGSE NMR spectroscopy

A 10 wt% W. R. Grace sample was analyzed in toluene- d_8 solution by preparing samples upon serial dilution of the commercial material. As in earlier work, MAO aggregates at higher concentrations lead to an increase in the calculated \bar{M}_D . The data obtained are summarized in Table 1, which includes the results obtained earlier for the material from Sigma Aldrich using the same techniques and instrumentation. Fig. 3 shows the changes in \bar{M}_D as a function of [AlMe] concentration as measured by NMR. Within the error of these experiments, the material from W. R. Grace is marginally of higher MW than the Sigma Aldrich h-MAO sample studied previously, although not at the 95% confidence level. The tendency towards aggregation of the two commercial samples is also quite similar. In contrast, DMAO, which features a lower Me_3Al content than either of the commercial samples (ESI, Fig. S1†), shows a much higher tendency to aggregate at the same concentrations, and this feature is responsible for its higher average MW.

Since we were interested in the influence of a polar solvent on these results, we also diluted the W. R. Grace h-MAO sample in chlorobenzene- d_5 . This solvent has a similar dielectric constant to PhF used in the ESI-MS work. In fact, chlorobenzene has been used as a solvent for ESI-MS experiments of this type.⁴⁷ The D_t of this solvent could not be measured due to the overlap of residual ^1H signals with toluene. Instead, the decay of the toluene Me resonances was used to estimate the D_t of toluene in this solvent assuming the viscosity of the mixture was the same as that of pure solvent (0.75 cP). Furthermore, the known $c_f r_H$ properties of toluene and the Chen equation⁴⁸ were used to estimate c_f (f assumed the same for both the solvent and solute) and thus r_H for MAO in this solvent using the decay of the AlMe resonances.

The correlation we developed for \bar{M}_D vs. D_t is not applicable in this more polar solvent. The results suggest that the hydrodynamic dimensions of MAO do not differ significantly in the more polar solvent (Table 2, entries 8–10 vs. 11–13), yet the higher \bar{M}_D values derived from D_t (which are significantly smaller in this solvent compared with toluene) would suggest otherwise. We do note that the slope of V_H vs. [AlMe] is significantly different in the two solvents (ESI†) suggesting that the aggregation tendency of MAO is increased in the more polar solvent.

Characterization of MAO-based ion-pairs by PGSE NMR spectroscopy

Table 3 summarizes the PGSE NMR results for the ion-pairs formed from the 10 wt% W. R. Grace sample and Cp_2ZrMe_2 including those reported earlier using DMAO derived from the Sigma Aldrich material.²⁴ As shown in Scheme 2, both contact

Table 2 Characterization of h-MAO by ^1H PGSE NMR spectroscopy

Sample NMR solvent	Entry	[AlMe] ^a (mM)	D_t ^b ($10^{-10} \text{ m}^2 \text{ s}^{-1}$)	r_H ^b (Å)	\bar{M}_D ^b (g mol ⁻¹)
Sigma Aldrich 10 wt% Benzene- d_6 ^c	1	17.4	4.02	8.57	1860
	2	60.5	3.81	8.99	2040
	3	184.5	3.89	8.82	1970
	4	299.3	3.63	9.41	2200
Sigma Aldrich DMAO Benzene- d_6 ^c	5	10.0	3.58	9.52	2160
	6	55.0	3.47	9.79	2380
	7	127	3.26	10.4	2640
W. R. Grace 10 wt% Toluene- d_8	8	11.0	3.78	9.79	2060
	9	85.5	3.66	10.1	2180
	10	377.0	3.54	10.4	2300
W. R. Grace 10 wt% Chlorobenzene- d_5 ^d	11	31.2	3.30	9.23	—
	12	241.8	3.06	9.90	—
	13	521.3	2.85	10.6	—

^a AlMe concentration calculated from the ^1H NMR spectra. ^b Errors in D_t , r_H and \bar{M}_D are $\pm 3.0\%$. ^c Data from ref. 24. DMAO was prepared by removal of excess Me_3Al in *vacuo*. ^d D_t and r_H were determined as described in the text.

(2) and outer-sphere (1) ion-pairs are formed and are involved in slow exchange on the ^1H NMR chemical shift time scale.

Chem3D Connolly volume calculations on both a cage and sheet model for OSIP **1**⁴⁹ indicate that they have V_{vdW} of 1660 and 1740 Å³ and have comparable densities of $\rho_{\text{calc.}} = 1.70$ and 1.62 g mL⁻¹ with models for MAO itself (Table 1) and so we have calculated \bar{M}_D for these species from their measured D_t .

The results for the inner sphere ion-pair (ISIP) $\text{Cp}_2\text{ZrMe}-\mu-\text{Me-MAO}$ **2** appear largely independent of the source of the MAO in use. The r_H of ISIP **2** correlates with that of the MAO in use and the extent to which MAO aggregates at higher concentrations leads to parallel increases in the apparent size of **2** at higher concentrations as seen in earlier work.²⁴ However, by comparing entry 5 with entry 8 the \bar{M}_D of ISIP **2** is only slightly different at similar concentrations.

Outer-sphere ion-pair (OSIP) $[\text{Cp}_2\text{ZrMe}_2\text{AlMe}_2][\text{Me(}]\text{MAO}\text{)}]$ **1** is much more prone to aggregation at higher concentrations to form ion quadruples and larger aggregates, as are all related ion-pairs of this type.²⁴ The \bar{M}_D of OSIP **1** is also not influenced by the source of the MAO in use (entry 4 vs. 7) though it would appear that the solvated dimensions are slightly different in toluene- d_8 vs. benzene- d_6 solution.

Although the dimensions of MAO appear largely unaffected by solvent polarity (Table 2), the same is not true for ion-pairs formed from MAO and Cp_2ZrMe_2 . In chlorobenzene- d_5 , the dimensions of both ion-pairs are dramatically reduced, even though the size of h-MAO is largely unaffected (Table 3, entries 10–12). The size reduction is much more pronounced for OSIP **1** than for ISIP **2**.

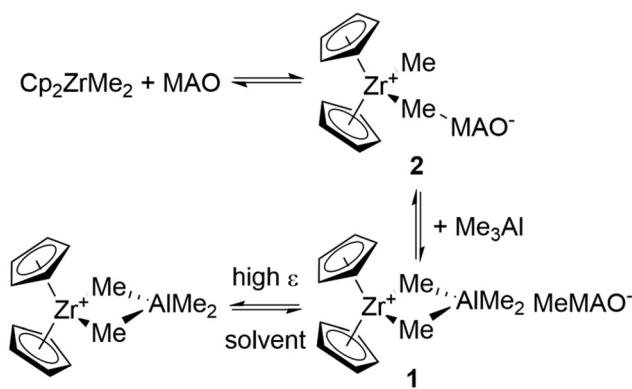
These results can be partially explained with reference to the equilibria shown in Scheme 2. While ion-pairs **1** and **2** are involved in slow exchange on the NMR time scale, diffusion is an even slower process, and the diffusion of each species influences the diffusion of all others related by the equilibria shown. In polar media, one can thus expect that OSIP **1** will be in exchange with solvated cations (and anions).⁵⁰



Table 3 Characterization of ion-pairs by ^1H PGSE NMR spectroscopy

Sample	Conc.	D_t^b	r_{H}^b	\bar{M}_D^b	
Entry	Species	(mM)	($10^{-10} \text{ m}^2 \text{ s}^{-1}$)	(Å)	(g mol^{-1})
Sigma Aldrich DMAO ^a					
1	OSIP 1	0.63	2.89	12.3	3220
2	ISIP 2	5.07	3.04	11.1	2960
3	DMAO	333	2.89	11.6	3220
4	OSIP 1	0.10	3.22	10.3	2690
5	ISIP 2	1.25	3.51	9.7	2330
6	DMAO	84	3.25	10.3	2650
W. R. Grace 10 wt% ^c					
7	OSIP 1	0.12	3.28	11.2	2610
8	ISIP 2	1.23	3.69	10.0	2150
9	MAO	167	3.47	10.6	2380
W. R. Grace 10 wt% ^d					
10	OSIP 1	0.22	4.06	7.69	—
11	ISIP 2	0.88	3.51	8.73	—
12	MAO	122	2.86	10.5	—

^a Excess Me_3Al removed *in vacuo*, samples prepared in benzene- d_6 ref. 24. ^b Errors in D_t , r_{H} and \bar{M}_D are $\pm 3.0\%$. ^c Sample diluted with toluene- d_8 . ^d Sample diluted with chlorobenzene- d_5 ; D_t and r_{H} were determined as described in the text.

**Scheme 2** Equilibria between Cp_2ZrMe_2 , MAO and inner and outer-sphere ion-pairs **2** and **1**.

Since the $[\text{Cp}_2\text{ZrMe}_2\text{AlMe}_2]^+$ cation has a much smaller size, the averaged dimensions of ion-pair **1** will be significantly reduced compared to the situation in toluene. The effect will even be seen for ISIP **2** but to a lesser extent as it is indirectly coupled to the third equilibrium. Alternately, this ion-pair is in direct equilibrium with Cp_2ZrMe_2 and MAO and the larger size of the latter species will counter the effect of the third equilibrium. In other studies,⁵¹ discrete contact ion-pairs like **2** were shown to be in equilibrium with σ -chlorobenzene complexes⁵² so it is possible that this equilibrium may also reduce the average dimensions of ion-pair **2** as well.

Comparison of ESI-MS results with PGSE NMR spectroscopy results

The negative ion ESI-MS spectra of 10 or 30 wt% MAO from W. R. Grace and metallocene complexes are also dominated by the same anion with m/z 1375 and formula $[16,6]^-$.⁵³⁻⁵⁵ In

positive ion mode using Cp_2ZrMe_2 one sees mainly the $[\text{Cp}_2\text{ZrMe}_2\text{AlMe}_2]^+$ cation with m/z 307 at Al : Zr ratios of 100 : 1 or greater.^{53,54} We also note that the average m/z ratio of the anion distribution is sensitive to the Al : Zr ratio.^{54,55} For ratios of 100 : 1 vs. 1000 : 1 the values are 1395 and 1484 Da, respectively (ESI, Fig. S2†). This means the nominal mass of the ion-pair **1** should vary between 1702 and 1791 Da, while that of ISIP **2**, which is not directly detected by ESI-MS, would be between 1630 and 1719 g mol⁻¹.

Obviously, there is a significant discrepancy between the MS results in fluorobenzene and the PGSE NMR results in toluene, well outside the experimental errors of either technique. While it is tempting to convert the D_t values in chlorobenzene- d_5 to \bar{M}_D values, OSIP **1** would end up lower in molar mass than ISIP **2** (1830 vs. 2330 g mol⁻¹), while both would be much lower than that of the MAO in use (3270 g mol⁻¹). This result is certainly not supported by any earlier work and as we have shown the correlation involving MAO does not appear to be applicable in this polar solvent.

The effect of polydispersity should also apply to the PGSE NMR results and thus one would expect \bar{M}_D would be larger than that determined by ESI-MS. However, the ESI-MS results suggest that the polydispersity of the MAO-based anions, even in aged materials (e.g. Fig. 1a vs. Fig. 1b), is very narrow with $D = \bar{M}_w/\bar{M}_n = 1.047$ to 1.021 (see also ESI MAO.xlsx†). This suggests that the \bar{M}_D values for both ion-pairs **1** and **2** should be larger but not much larger than those estimated by ESI-MS. Clearly this is not the case, so we discuss possible reasons for this discrepancy.

The number average m/z ratio (and any higher averages as well as D) calculated from the raw MS data is dependent on all anions being detected with the same sensitivity.⁵⁶ For higher m/z anions, this cannot be true in a general sense given the intrinsic bias of MS towards lower m/z ions.⁴⁰ This leads to a pronounced narrowing of the MWD and a bias towards lower m/z averages as measured by low resolution MS vs. other methods such as GPC.⁵⁶ Even in the case of high-resolution techniques such as FT-ICR MS, the polydispersity of narrow MWD synthetic polymers is still underestimated by mass spectrometry.⁵⁷

If there are also differences in surface activity between homologous ions even more pronounced intensity differences can be observed.⁵⁸ For an example obtained under the same conditions as we use for the analysis of MAO, see ESI Fig. S3† where the sensitivity difference for $[\text{n-Bu}_4\text{N}]^+$ (m/z 242) vs. $[(\text{n-C}_{18}\text{H}_{37})_4\text{N}]^+$ (m/z 1026) is *ca.* 150-fold at equivalent concentrations. If differences of this magnitude also pertain to the aluminoxane anions, particularly those with higher m/z ratios, it is easy to understand why \bar{M}_n , \bar{M}_w , etc. are underestimated using the ESI-MS data.

It is also possible that the higher \bar{M}_D values measured using PGSE NMR reflect weak but reversible aggregation of the ion-pairs with other neutral components of MAO, even at the lowest concentrations examined. These interactions might be simply too weak to be detected in the ESI-MS experiment under the conditions used for the analysis of these mixtures (see the Experimental section).



In any event, the PGSE NMR results for the Grace 10 wt% MAO are in good agreement with the MW of an isolated, neutral component of that mixture,¹⁵ while the NMR estimates of MW reported here for the ion-pairs **1** and **2** seem reasonable given that for MAO itself. In particular, our NMR measurements and the correlation between D_t and \bar{M}_D do not support the conclusion that OSIP **1** is very high in MW or very different from the MAO in use.

Aging of MAO determined by ESI-MS and PGSE NMR

Earlier we obtained ESI-MS and ^1H NMR spectra of a commercial and a proprietary sample of h-MAO with nearly identical Al (13.7 and 13.6 wt%) and similar Me_3Al contents (5.0 and 4.1 wt%), in the presence of OMTS and $\text{THF}-d_8$ donors, respectively. Representative negative ion mass spectra are shown in Fig. S4† while Me_3Al and $[\text{Me}_2\text{Al}(\text{THF})_2]^+$ contents as measured by NMR (Fig. S5†)^{21,59} are summarized in Table S2† as a function of storage time at room temperature.

A 30 wt% commercial formulation ages much faster than a 10 wt% formulation (*vide infra*) and is accompanied by noticeable gel formation after only 2–3 weeks at room temperature. The proprietary sample with higher m/z anions gels faster under these conditions even though the anion distribution is not changing rapidly, a seemingly counter-intuitive result. That the two different anion distributions change at different rates is shown by comparing the average m/z ratio of each mixture *vs.* time (Fig. 5a, open *vs.* closed symbols).

If one recognizes that the proprietary sample has an initial composition (average m/z = 1709 Da) that is similar to the commercial sample at three weeks (average m/z = 1710 Da), the slower rate of aging of the former can be rationalized as being further located along the *same* growth curve (dashed line in Fig. 5a). Presumably the onset of gelation is also dependent on \bar{M}_w ⁶⁰ of the material present and thus more rapid in the material with higher m/z anions.

Although we do not have spectra of these two samples over longer time periods, due to their pronounced tendency to gel, we have included some ESI-MS spectra of the 10 wt% W. R. Grace sample at six weeks and six months (Fig. S6†)³⁴ which shows that this material has an anion distribution that resembles the aged proprietary formulation at longer times.

NMR experiments in $\text{THF}-d_8$ demonstrate that the proprietary formulation with higher m/z anions has *ca.* 50% higher activator content based on the intensity of signals due to the $[\text{Me}_2\text{Al}(\text{THF})_2]^+$ cation. This difference is maintained for the aged samples, as both the sol and gel phases are soluble in excess THF or sufficient amounts of OMTS (*ca.* 2 mol% in PhF).

The only time dependent change seen is the total Me_3Al and toluene content, which decreases in both samples (Fig. S5†), because these two volatile materials evaporate from screw-top vials inside the glovebox over the three-week period of these experiments.

To compare with these results, we monitored the aging of 10 wt% Grace formulation, which is much less prone to gelation, by PGSE NMR. In Table 4 we show D_t and size data at 0 to

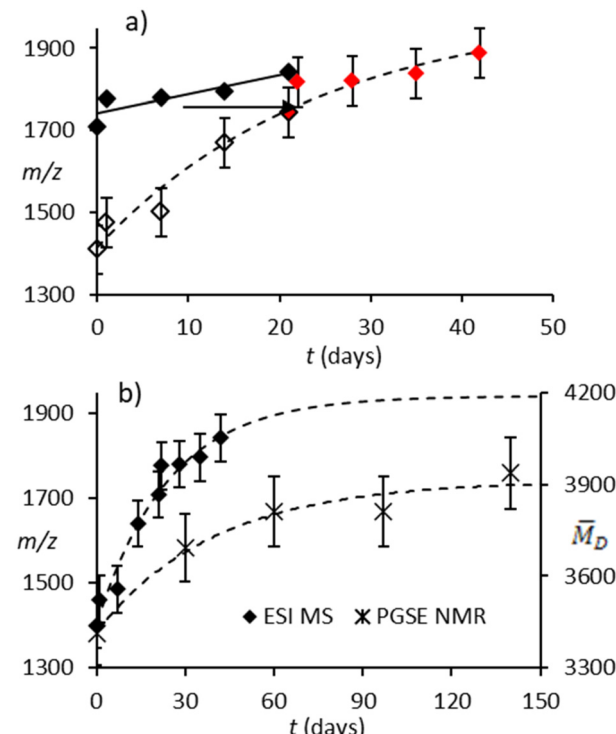


Fig. 5 (a) Change in average m/z ratio of hydrolytic MAO *vs.* time (days). Commercial 30 wt% MAO (open symbols) and a proprietary formulation (black symbols) at 0, 1, 7, 14 and 21 days. The average m/z ratio was calculated from the raw ESI-MS data files of m/z (50–3000 Da) *vs.* intensity (counts). (b) Average m/z ratio (left hand y axis) and diffusion average MW (right hand y axis) *vs.* time for 30- and 10 wt% MAO from Grace. A function of the form $y = a + b(1 - \exp(-ct))$ was fit to each set of data using non-linear regression.

140 days at room temperature with \bar{M}_D values based on the correlation developed. In these experiments, undiluted MAO was employed and so even in unaged materials the dimensions and MW of the MAO are much larger due to reversible aggregation.²⁴ Based on the data in Table 4 (entries 1–4) the increase in \bar{M}_D scales in a linear fashion with $[\text{AlMe}]$ ($R^2 = 0.99$).

Based on the integral ratio of aluminoxane and Me_3Al to toluene in this sample, the amount of soluble aluminoxane decreased by about 20% over 140 days due to gelation. Interestingly, methane formation^{61,62} was not detectable over this period in this (sealed) sample (Fig. S7†) suggesting that gelation occurs without chemical change at room temperature. Indeed, since the gel can be redissolved in a donor solvent, initial gelation must be a process involving non-covalent interactions which makes sense given the sheet structure for an isolated component of MAO. However, it seems plausible that physical gelation may be complicated by chemical cross-linking in samples where water is not completely excluded, for example, if exposed to the atmosphere in a glovebox with ppm background levels of water for long periods of time.

Over the 140-day period, the molar mass of the sol phase increases by about 15% which is lower than the 22% increase in the average m/z ratio for a 30 wt% formulation of the same



Table 4 Monitoring of 10 wt% Grace MAO by PGSE ^1H NMR spectroscopy^a

Entry	Time (days)	[AlMe] (mM)	D_t ^b ($10^{-10} \text{ m}^2 \text{ s}^{-1}$)	r_{H} ^b (Å)	V_{H} ^b (Å 3)	\bar{M}_{D} ^b (g mol $^{-1}$)
1	0	11.0	3.78	9.79	3930	2070
2	0	85.0	3.66	10.09	4300	2180
3	0	377	3.54	10.40	4710	2300
4	0	2020 ^c	2.78 ^d	11.5	6370	3410
5	30	1980	2.65 ^d	12.0	7340	3690
6	60	1920	2.60 ^d	12.3	7750	3810
7	97	1800	2.60 ^d	12.3	7800	3810
8	140	1630	2.55 ^d	12.4	8100	3940

^a Samples diluted with toluene- d_8 except where noted. ^b Errors in D_t , r_{H} and \bar{M}_{D} are $\pm 3.0\%$, $V_{\text{H}} \pm 10.0\%$. ^c Undiluted sample. ^d Using a viscosity of 0.7 cP corresponding to a 10 wt% MAO solution in toluene.

MAO over 21 days. This is to be expected given the difference in concentrations. A comparison of the ESI-MS and PGSE NMR data is shown in Fig. 5b, where the dashed lines correspond to growth curves fit to the experimental data.

It is clear that the same process is being monitored by the two techniques although there is a systematic discrepancy, by more than a factor of two for the 10 wt% solution, between \bar{M}_{D} and the average m/z ratio of these mixtures. Given that the m/z ratio is expected to track the \bar{M}_{n} , while the critical MW for gelation (<3400 g mol $^{-1}$ for the sol phase) will be correlated with \bar{M}_{w} ,⁶⁰ and thus \bar{M}_{D} , the magnitude of the discrepancy, is what one might expect for a polydisperse sample, bearing in mind the concentration-dependent aggregation of MAO in solution.

Conclusions

As outlined here, PGSE NMR and ESI-MS provide reasonably consistent data as to the average molar mass of MAO *viz a viz* the ion-pairs derived from this material and reactive donors. The most pronounced discrepancy is related to outer-sphere ion-pairs, which are known to be prone to self-aggregation²⁴ and may also be susceptible to aggregation with the additional, neutral components of MAO. Future work should examine this hypothesis since it is one that can be experimentally tested, especially given the recent isolation of a reactive component of MAO.

The differences between formulations can be detected using either technique, while we expect that 2D techniques like DOSY²⁸ will provide information on molecular weight distribution, especially when complemented by data from ESI-MS (*i.e.* MMAO 12 *vs.* hydrolytic MAO).

Of the two methods, PGSE NMR is the more practical technique based on the availability of suitable instrumentation and expertise. Also, as it is a non-destructive solution technique, it can be used to study weak aggregation phenomena that are less easily studied in most mass spectrometric experiments.

Finally, we have established a useful, internally consistent, and reasonably precise relationship between the molar mass

of MAO and D_t , now fully consistent with the experimental data which we hope will allow routine characterization of this material by NMR spectroscopy in non-polar solvents going forward.

Experimental section

Methylaluminoxane solution (10 or 30 wt% in toluene [Al] \approx 1.8 or 5.4 M) was obtained from W. R. Grace and stored at -20°C in a glove-box freezer. A sample of anhydrous and degassed THF- d_8 was also provided by W. R. Grace for NMR studies of MAO aging. Octamethyltrisiloxane (OMTS) and Cp_2ZrMe_2 were obtained from Sigma Aldrich and used as received. Fluorobenzene (PhF) was obtained from Oakwood Chemicals Ltd, refluxed and distilled from CaH_2 and stored over activated molecular sieves (4 Å) prior to use.

Aging and activator content of 30 wt% MAO

A solution of 30 wt% MAO in toluene was added to 10 screw top vials containing 0.1 mL (ESI-MS) or 0.25 mL (^1H NMR) of solution. The vials were sealed with Parafilm™ and analyzed after 0, 1-, 7-, 14- and 21-days of storage at room temperature in the glovebox.

For the ESI-MS experiments, the contents of one vial were diluted with 1.0 mL of a solution of OMTS (21.2 mg in 10 mL of PhF). A 0.1 mL aliquot of the homogeneous solution was then diluted by a factor of 10 with dry PhF prior to analysis by ESI-MS: this solution was analyzed using a QTOF Micro spectrometer *via* pumping it at *ca.* 40 $\mu\text{L min}^{-1}$ through PTFE tubing (1/16" o.d., 0.005" i.d.) to the ESI-MS probe and source using a syringe pump. The capillary voltage was set at 2900 V with source and desolvation gas temperatures set at 85 °C and 150 °C, respectively with the desolvation gas flow at 400 L h $^{-1}$.

Raw MS data files of m/z ratio *vs.* ion intensity (counts) were imported into Excel and analyzed to calculate number- and weight-average m/z ratios and dispersity.^{56,57} Some results are deposited as ESI† for a sample of MMAO 12 and 30 wt% h-MAO at the different times analyzed (MAO.xlsx†).

For the NMR experiments, the contents of another vial were diluted with 1.0 mL of THF- d_8 with stirring to mix the contents. After one hour, 0.6 mL of the diluted sample was then transferred to a 5 mm NMR tube equipped with an air-tight Teflon™ valve and analyzed by ^1H NMR spectroscopy at 500 MHz on a Bruker Avance 500 instrument, using a 30° pulse-width and 5 s relaxation delay averaging 64 transients. The spectra were integrated before and after baseline subtraction of the broad MAO resonance as described in the literature^{21,59} to determine the total Al, Me_3Al and $[\text{Me}_2\text{Al}(\text{THF})_2]^+$ contents.

PGSE NMR experiments

^1H PGSE NMR measurements were performed using a double stimulated echo pulse sequence (dstepg3s1d) on a Bruker AVANCE III HD 400 spectrometer equipped with a SmartProbe™ with a Z-gradient coil, at 298 K without spin-



ning. For polydisperse species, the dependence of the resonance intensity (I) on a constant waiting time and on a varied gradient strength (G) is described using the equation

$$I = \sum_i (I_0)_i \exp \left[-(\gamma \delta)^2 (D_t)_i \left(\Delta - \frac{\delta}{3} \right) G^2 \right]$$

where I = intensity of the observed spin echo, $(I_0)_i$ = intensity of the spin echo in the absence of a gradient corresponding to the i -th species, D_t = self-diffusion coefficient of the i -th species, Δ = delay between the midpoints of the gradients, δ = length of the gradient pulse, and γ = magnetogyric ratio. The shape of the gradients was rectangular, their lengths were 4–5 ms, and their strengths G were varied during the experiments. In the present case, we tested both monoexponential ($i = 1$) and biexponential ($i = 2$) treatments by fitting the decays of I versus G with a non-linear regression algorithm and the best model for each sample was chosen by comparing the fitting errors and R values (see the ESI† for the data). All spectra were recorded using 64 K points and a spectral window of 6400 Hz and they were processed with a line broadening of 1.0. Different values of Δ , G , and numbers of transients were used for different samples.

The self-diffusion coefficient D_t , which is directly proportional to the slope (m) of the exponential fitting obtained by plotting I versus G was estimated by evaluating the proportionality constant using a sample of HDO (5%) in D₂O (known diffusion coefficients in the range 274–318 K)⁶³ under identical conditions as for the sample of interest. The solvent was taken as an internal standard in the case of toluene-*d*₈ measurements. In the case of chlorobenzene-*d*₅, the overlap between aromatic signals hampered the precise estimation of D_t and hydrodynamic dimensions were obtained using dissolved toluene as an internal standard. D_t data were treated as described in the literature in order to derive the hydrodynamic dimensions.²⁵ Experimental errors in r_H are $\pm 3.0\%$ and $\pm 10\%$ for V_H .

NMR samples for PGSE NMR measurements were prepared inside a glovebox by successive dilution. After having inserted the sample into the NMR probe, the sample was equilibrated for at least 30 minutes before starting the experiment to achieve optimal thermal stability of the solution. The actual concentration was estimated by integration relative to an external standard. The nominal concentration refers to Al-Me groups and considers both MAO and AlMe₃.

Computational studies

DFT studies of aluminoxane structures employed the M06-2X density functional⁶⁴ and TZVP basis set⁶⁵ for geometry optimization. Gaussian 16⁶⁶ was used for all calculations and stationary points were confirmed as minima using frequency calculations. Electronic energies and thermodynamic quantities for all structures are summarized in Table S1† while coordinate files of structures in Fig. 4 are deposited as the ESI (Table S1.xyz).†

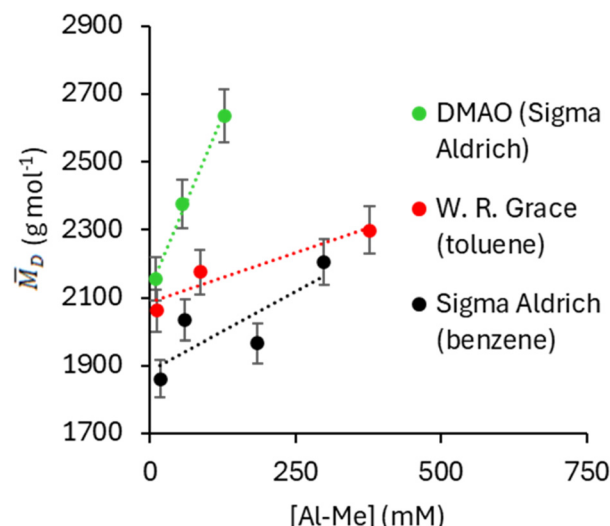


Fig. 4 Diffusion average molecular weight (error bars are $\pm 3.0\%$) vs. AlMe concentration for different MAO samples as measured by PGSE NMR. Data for the Sigma Aldrich samples are from ref. 24.

Author contributions

L. R. carried out PGSE NMR experiments. S. C. carried out ESI-MS and ¹H NMR analyses of commercial and proprietary MAO formulations. S. C. and M. L. conducted quantum chemical calculations. S. C. devised the linear correlation between D_t and molar mass. The manuscript was written through contributions of all authors.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts to declare.

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