In this paper we present the aggregational motifs of the widely used alkali-metal cyclopentadienides (CpLi, CpNa, CpK, CpRb, CpCs) in THF-\(d_8\) solution estimated by ECC-DOSY NMR spectroscopy. They form monomeric contact ion pairs (CIPs) in THF-\(d_8\) solution, whereas in NH\(_3\) solvent-separated ion pairs (SSIPs) are observed. The applicability of ECC-DOSY is further advanced by introducing ECC-MW estimation software.

Alkali metal cyclopentadienide (CpM) derivatives are among the most widely used starting materials in organometallic synthesis.\(^1\) They can be used to synthesise a variety of different sandwich or half-sandwich d-block organometallics through transmetallation or salt elimination, which in turn can be used e.g. as polymerisation catalysts.\(^2\) Ferrocene, which was synthesised by Kealy and Pauson in 1951 is often described as the first metallocene, but alkali metal cyclopentadienides CpMs were indeed synthesised more than 50 years earlier (Thiele, 1900).\(^5,6\) The solubility of CpM compounds has to be described as poor in hydrocarbons as well as in ethers. THF is the only solvent that provides reasonable concentrations in solution. High melting points and low volatilities also underline their salt-like character.\(^7,8\) Crystal structure analysis shows that the donor free compounds form polymeric chains that are linear for [CpLi]\(_x\) and [CpNa]\(_x\) but bent for all higher homologues (for CpRb there are ambivalent structures, Scheme 1).\(^9\)–\(^11\) Various solvation states account for various excerpts from the solid state structures\(^12\) like for example [Li(NH\(_3\))\(_3\)][Cp]\(^-\),\(^13\) [(TMEDA)LiCp],\(^14\) [(H\(_2\)N)\(_2\)NaCp],\(^15\) [H\(_3\)N\(_2\)]LiCpLiCp,\(^15\) [Ph\(_2\)P][Cp\(_x\)Li],\(^16\) or [Ph\(_2\)P][Cp\(_x\)Cs].\(^17\)

Most of the crystal structures have to be classified as contact ion pairs (CIP), but there are a few examples of solvent-separated ion pairs (SSIPs) preferentially generated by the use of ammonia or crown ethers.\(^11,16\) Aggregation of organometallic molecules in solution determines not only their reactivity but can also shed light on reaction mechanisms, and hence provide a handle to improve e.g. yields and/or selectivities. Though crystal structure analysis can elucidate the molecular bonding situation,\(^15\) in many cases it differs quite substantially from solution structures, which in turn can vary in different solvents considerably. Therefore, the absolute size of molecules in solution has been of interest for a long time. In 1992 Johnson and Morris introduced an NMR spectroscopic method to tackle this problem: DOSY (Diffusion Ordered Spectroscopy).\(^19\) This technique correlates chemical shift information with the self-diffusion coefficient \(D\) of the compound. Li and Williard \textit{et al.} pioneered DOSY NMR spectroscopy for small reactive organometallic molecules by addition of at least three internal references to one NMR sample to get an internal calibration curve (ICC).\(^20\) In the last decades many methods have been proposed to link diffusion coefficients either to molecule sizes or molecular weights (MWs).\(^21\)–\(^25\) Recently, our group developed and published an advanced method.\(^26\) We established power law based external calibration curves (ECCs) together with normalised diffusion coefficients to estimate the MWs of organometallic compounds via DOSY in solution with much improved accuracy compared to previous approaches.\(^27\)–\(^30\) We were also able to extend the scope of this method towards other commonly used solvents \(e.g.\) DMSO-\(d_6.\)\(^31\) In the current...
study samples were prepared by addition of internal references to Cpm in order to normalise the diffusion coefficients of these analyses (log $D_{x,\text{norm}}$). This way MWs can be estimated independently of NMR spectrometer properties and differences in temperature or viscosity en route to estimate MWs for the solution structures of Cpli, CpNa, CpK, CpRb and CpCs in THF- $d_8$. CpLi was also analysed in ammonia solution.

In THF the $^1$H- and $^13$C-NMR spectra of all alkali metal cyclopentadienides show only a single resonance (see ESIT). The signal tends to be shifted towards higher field when descending Group 1. $^7$Li and $^{133}$Cs NMR resonances for Cpli and CpCs, respectively, were found to be in accordance with literature. The $^2$Na NMR resonance ($-28$ to $-31$ ppm) is shifted towards lower field compared to solid state MAS NMR findings for CpNa and [CpNa-THF] and can be shifted even lower upon cooling. The same was observed for the $^7$Li NMR signal, which shifts and also broadens upon cooling. In 1990 it has been reported by Paquette et al. that Cpli undergoes a fast exchange process between monomeric and a “sandwich” dimeric species, which results in the splitting of the $^7$Li signal at lower temperatures, but they could not “tell whether the monomer–dimer equilibrium of ‘Cpli’ is shifted to either side at room temperature”. The neat lithocene anion [Cp$_2$Li]$^-$ is characterised triply in the solid state. We recorded spectra in THF solution down to $-100^\circ$C and could not observe any splitting of the $^7$Li NMR signal and at $-105^\circ$C CpLi precipitates. In addition, DOSY-NMR-spectroscopy was used to estimate MWs with ECCs. As shown before, molecular shapes are quite important to accurately interpret diffusion data. We found that for most organometallic compounds the dissipated spheres and ellipsoids (DSE) calibration curve is the most appropriate, especially for lithiated compounds. Therefore, all MWs (MW$_{\text{det}}$) of Cpm in THF were estimated via the DSE-ECC, if not stated otherwise. It seems also mandatory to calculate molecular densities (MD$_{w}$) to foresee whether or not proposed structures might evoke deviations in the ECC-MW estimation. For all herein proposed aggregates, however, such errors due to higher MD$_{w}$ can be excluded (see ESIT). For a more straightforward procedure we developed ECC-MW estimation software, in which all current and up-coming ECC-MW estimation techniques and calibration curves as well as different references can be selected and applied. With this software estimated MW$_{\text{det}}$ in THF were compared to MWs of likely monomeric [CpM-THF] and dimeric aggregates ([CpM]$_2$-THF$_x$) with $x = 0$–4, M = Li, Na, K, Rb, Cs (see Table 1 or ESIT). The results fit best for monomeric aggregates with different quantities of coordinated THF per alkali metal (except for CpCs, see Table 1 and Scheme 2), hence we propose this to be the most populated species for Cpm in THF solution. Just considering the MW$_{\text{det}}$ of dimeric aggregates, degraded by most coordinated THF they would also be an option (see ESIT). Metalloocene-like aggregates of e.g. [Cp$_2$M-THF]$_x$ were not considered because they could not be differentiated from [CpM-THF]$_{x+1}$, because MW(THF) $\approx$ MW(Cp).

For Cpli the predominant monomeric aggregate seems to be coordinated by only two THF molecules. This molecule is stable over a large temperature range. If normalised log $D_{x,\text{norm}}$ are compared, there is almost a perfect fit for all temperatures: log $D_{x,\text{norm}}$(CpLi, +50) = $-8.886$; log $D_{x,\text{norm}}$(CpLi, +25 $^\circ$C) = $-8.901$; log $D_{x,\text{norm}}$(CpLi, $-50^\circ$C) = $-8.905$; log $D_{x,\text{norm}}$(CpLi, $-80^\circ$C) = $-8.909$; $\Delta$log $D_{x,\text{norm}}$(CpLi, +50/−$80^\circ$C) = 0.023.

The formation of CIPs for Cpli in THF can also be confirmed. This behaviour can be deduced from the same log $D_{x,\text{norm}}$ values from the $^7$Li- and $^1$H-DOSY NMR spectra (see Fig. 1 and ESIT). Unfortunately, no signal was observable in the $^7$Li- and $^1$H-HOEYS NMR experiment to confirm this. For CpNa and CpK coordination by 3 THF molecules is preferred with a slightly bigger MW$_{\text{diff}}$ for CpK. After cooling CpNa to $-50^\circ$C the same result could be obtained (log $D_{x,\text{norm}}$(CpNa, +25 $^\circ$C) = $-8.978$); log $D_{x,\text{norm}}$(CpNa, $-50^\circ$C) = $-8.971$; $\Delta$log $D_{x,\text{norm}}$(CpNa, +25/−$50^\circ$C) = 0.007. It was previously proposed that the mono-solvated [CpNa-THF] is a possible aggregate in the solid state, which we could not confirm in solution. Cooling led to precipitation of Cpli, CpRb and CpCs, therefore no further insight could be gained for thermal dependence. CpRb seems to be coordinated by only 2 THF molecules at $25^\circ$C. In the literature a coordination polymer of CpRb chains is known, which crystallised with two THF molecules attached to the metal. This provides plausibility to the deaggregation in solution by breaking one of the two Cp-Rb bonds. Supposedly this is due to the fading donor capacity of

| Table 1 | ECC$_{\text{DSE}}$ was used to determine MW$_{\text{det}}$ of Cpm in THF-$d_8$, except for CpCs, for which ECC$_{\text{DSE}}$ was used. The accuracy of the DSE ECC is in the range of $\pm 9$%. Standard deviations calculated from error propagation are listed in parenthesis. All results in this table are from measurements at 25 $^\circ$C |
|---|---|---|---|---|
| Cpli | 218(11) | 1 | 32 | 65 |
| CpNa | 295(16) | 21 | 3 | 27 |
| CpK | 335(18) | 26 | 4 | 17 |
| CpRb | 294(16) | 0 | 25 | 49 |
| CpCs | 1855(198) | — | — | — |

![Scheme 2](image-url)
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THF descending Group 1. This was e.g. calculated for alkali metal/ammonia binding energies ($\Delta E[M^+\cdot NH_3]$); 170.3 (Li), 117.2 (Na), 82.0 (K), 71.1 (Rb), 61.9 (Cs) kJ mol$^{-1}$. We recently observed the same trend in hexamethyldisilazides crystallized from liquid ammonia. The same could be valid for $\Delta E[M^+\cdot$THF$]$ as the heavier metal Cp compounds tend to form coordination polymers instead of taking on more coordinated solvent molecules. Rb seems to be the borderline case between not adding polymers instead of taking on more coordinated solvent molecules from liquid ammonia. TTh ese same could be valid for $\Delta E[M^+\cdot$THF$]$ as the heavier metal Cp compounds tend to form coordination polymers instead of taking on more coordinated solvent molecules. Rb seems to be the borderline case between not adding polymers instead of taking on more coordinated solvent molecules. Prosenc could show the formation of a caesocene-tripledecker. In THF solution they form monomers and could analyse the donor solvent coordination in alkali metal compounds. THF- is formed because coordination polymer bonds are more important to caesium than to rubidium. In 1996 Harder and Prosenc could show the formation of a caesocene-tripledecker. From this and the estimated MW, we envisage a penta- [CpCs]$^5$ (MW,merge) = $-8\%$ and hexamic [CpCs]$^6$ (MW,merge) = $12\%$), whereas other motifs with different amounts of THF or Cp are feasible. Still, it needs to be emphasised that current ECCs are not optimised for aggregates that are heavy since reference compounds do not cover MWs > 600 g mol$^{-1}$ yet.

Furthermore, we measured a sample of CpLi in ammonia solution (see ESI†), where the formation of SSIPs was expected, and saw a change in chemical shift for the $^7$Li NMR spectroscopic signal towards $-0.59$ ppm. Moreover, a coupling between ammonia and lithium in a $^7$Li-$^1$H-HOESY experiment could be observed, which confirms the formation of SSIPs.

In conclusion, with the new ECC-DOSY NMR method we could analyse the donor solvent coordination in alkali metal cyclopentadienides. In THF solution they form monomers and quite surprisingly the lithium and rubidium derivatives only coordinate two solvent molecules while sodium and potassium accommodate three. Presumably the caesium derivative forms a pentamer with two THF molecules coordinated to each metal. We found SSIPs for CpLi in ammonia solution. In addition, we introduced our new ECC-MW estimation software available for anyone to use from our website.

All NMR experiments were recorded on either a Bruker Avance 400 spectrometer equipped with an observer broadband probe with $z$-axis gradient coil with maximum gradient strength of 57 G cm$^{-1}$ or Bruker Ascend 400 spectrometer equipped with an inverse broadband probe with $z$-axis gradient coil with maximum gradient strength of 51 G cm$^{-1}$. All spectra were acquired in 5 mm NMR tubes. Sample spinning was deactivated during measurements. All DOSY spectra were recorded using the standard Bruker dstebpg38 pulse sequence with zero spoil gradients with convection compensation. The diffusion time was $\Delta = 0.1$ s. The duration of the magnetic field pulse gradients $\delta/2$ was adjusted for every compound in a range of 1–3 ms (2 to 7 ms for $^7$Li). The delay for gradient recovery was 0.2 ms and the eddy current delay 5 ms. For each DOSY-NMR experiment, a series of 16 spectra on 32 K data points was collected. The pulse gradients were incremented from 2 to 98% of the maximum gradient strength in a linear ramp with a total experiment time of 51 min. The temperature was set and controlled at 298 K with an air flow of 400 l h$^{-1}$ in order to avoid any temperature fluctuations due to sample heating during the magnetic field pulse gradients if not stated otherwise. After Fourier transformation and baseline correction, the diffusion dimension was processed with the Topspin 3.1 software. Diffusion coefficients were calculated by exponential fits with the $T_1/T_2$ software of Topspin. All samples have been prepared inside a glove box. THF-$d_8$ is stored over 4 Å molecular sieves under argon. All samples were prepared using 15 mM solutions of analyte and internal reference (2,2,3,3-tetramethylbutane (TMB) or 1,2,3,4-tetraphenylcyclohexane (TPhN)). For NMR measurements in NH$_3$, gaseous NH$_3$ was introduced into the NMR tube for one minute at 78 °C; afterwards 0.1 mL of toluene-$d_8$ was added for referencing and after sealing the NMR tube spectra were recorded at ambient temperature. Diffusion coefficients of compounds in THF-$d_8$ were normalised either with the fixed TPhN signal of $\log D_{THF}(TPhN) = -9.1054$ or the fixed TMB signal of $\log D_{THF}(TMB) = -8.7749$.

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Notes and references

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The fixed diffusion coefficient \( \log D_{\text{fix}} \) of TPhN or TMB was estimated by using the middle \( \log D \) value of multiple DOSY measurements of 15 mM solutions at 25 °C.

\[ MW_{\text{det}} = \frac{\text{MW}_{\text{calc}}}{\text{MW}_{\text{det}}} \times 100\% \]

where \( MW_{\text{det}} \) is the ECC-determined MW of the analyte and \( MW_{\text{calc}} \) the calculated MW.


A term coined “molar van-der-Waals density” can be calculated with MDW = \( \sum V_W \), where \( \sum V_W \) is the sum of the van-der-Waals volumes of all atoms of a specific molecule. This MDW can be used to account for errors in ECC-MW-calculation, when using compounds incorporating heavier elements. For detailed informations see ES†.


