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Improved access to polythioesters by heterobimetallic aluminium catalysis

Bimetallic Al(III) catalysis exhibits improved rates, and greater monomer tolerance and linkage selectivity for thioanhydride/epoxide copolymerisation than Cr(III), resulting in the synthesis of novel co- and ter-polymers as well as high Mn materials with functional vinyl substituents.





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Improved access to polythioesters by heterobimetallic aluminium catalysis†

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Bimetallic Al(III) catalysis mediates thioanhydride/epoxide copolymerisation at greatly improved rates and monomer tolerance than analogous Cr(III) catalysis. Moving to sulfurated monomers furthermore generally improves rates and selectivites.

Heteroatom containing polymers are more susceptible to degradation than current commodity polymers comprising an all-carbon backbone, due to the presence of reactive centres in the polymer main-chain.¹⁻⁴ In this regard, moving from oxygenated polymers like polyester and polycarbonates to sulfur containing analogues imparts additional desirable thermal and optical properties onto the polymers while further enhancing their degradability.5-14 In this regard the ring-opening copolymerisation (ROCOP) of cyclic thioanhydrides with epoxides results in poly(ester-thioesters) from alternating enchainment of the two monomers. 15-23 Bicomponent catalysts featuring a metal or main group based Lewis acid in combination with soluble Lewis bases or onium salts (which have also been combined within the same scaffold) can mediate this process, yet these suffer from distinct challenges: as many rely on the use of toxic and expensive chromium or $[(Ph_3P)_2N]^+Cl^-$; are not air and moisture tolerant; and are synthetically nontrivial to access. Tackling some of these problems, our group established the use of methoxy-substituted bisphenoxyimine based chromium catalysts in the ROCOP of phthalic thioanhydride (PTA) or CS2 with cyclohexene oxide (CHO) to yield sulfurated polymers.²⁴⁻²⁶ These could solubilise alkali metal acetates AMOAc (AM = Li, Na, K, Rb, Cs) by means of coordination to an all-oxygen donor site in the ligand sufficiently, to form heterobimetallic complexes that also form in situ during catalysis, and these then facilitate PTA/CHO or CS2/CHO

volatiles in vacuum, and the adduct formation could be con-

firmed by ¹H NMR spectroscopy (see ESI,† Fig. S6). Due to the

greater practicality of this method, we employed all following

ROCOP. Viewed differently these complexes coordinatively

enable the use of cheap AMOAc salts as cocatalysts in place of onium salts. Our previous work on Cr(III) catalysts revealed that

both ligand modifications and alkali metal choice affect activity

and selectivity and that there are parallelling trends regarding

this for CS₂/CHO and CO₂/CHO ROCOP. Unfortunately though,

these complexes still relied on the use of toxic and expensive

Chromium, which is undesirable as catalysts traces in the

produced polymers rule out many applications (as e.g. in a

biological context). Hence, we explore in this report the use of

Fig. 1 Outline of the current work; R=H (CHO), CH=CH₂(vCHO).

the Al(III) variants featuring an earth abundant and non-toxic metal and not only find these to exhibit improved rates but also offer insight into how monomer sulfuration affects catalysis while improving the monomer scope to yield unexplored co and terpolymers from thioanhydrides (Fig. 1). We initially prepared L^{Cy}AlOAc, i.e. the Al(III) version of our previous Cr(III) catalyst *via* the literature reported procedure.^{27,28} Testing L^{Cy}AlOAc in PTA/CHO ROCOP at a loading of 1 eq. L^{Cy}AlOAc: 250 eq. PTA: 1000 eq. CHO, with one equivalent of added RbOAc for in situ complexation, results in 97% PTA conversion in 25 min at 80 °C. The same activity was found when the preformed adduct was employed. To do this, L^{Cy}AlOAc and RbOAc were combined in methanol, followed by removal of

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Cat:L*AIOAc + M(OAc)_x This contribution: Al improves rate and (Li/Na/K/Rb/Cs)OAc R-group tolerance (Mg/Ca/Sr/Ba)OAc2 Sulfur improves rate and selectivity

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catalysis with preformed adducts. We followed the monomer consumption via aliquot analysis by ¹H NMR which showed a maximum TOF of 638 h⁻¹ for the initial linear increase of the PTA conversion *versus* time plots. In comparison, the previously investigated Cr(III) version showed a maximum TOF of 50 h⁻¹ meaning that, moving to non-toxic and abundant Al(III) improves activity by at least one order of magnitude.‡ The formed polymer shows an apparent $M_{\rm p}$ of 11.3 kg mol⁻¹ with a polydispersity of D = 1.4 (see ESI,† Fig. S37). The GPC curve is bimodal indicating two co-occuring initiation processes which are explored in more detail below. No (thio)ether links are formed indicating alternating insertion of CHO and PTA which would result in a 1:1 distribution of thioester: ester links, i.e. 50% thioester. Yet, only 45% thioester links are formed, and this can be rationalised by an oxygen enrichment mechanism (see ESI,† Scheme S2) producing thiirane by-products which could be identified in the crude reaction mixture (see ESI,† Fig. S11). Note that LCyAl itself without RbOAc also catalyses PTA/CHO ROCOP, albeit with the formation of (thio)ether links from epoxide or thiirane homopropagation as well as a low TOF $< 20 h^{-1}$, furthermore supporting the cooperative catalysis between LCyAlOAc and cocatalytic RbOAc.²⁹ As we previously found the ligand backbone on the bisphenoxyimine spectator ligand to influence the catalysis we proceeded to investigate the phenylene variant L^{Ph}AlOAc, the ethylene variant L^{en}AlOAc and the dimethylpropylene variant LproAlOAc in PTA/CHO ROCOP under the same conditions. Whilst all aliphatic variants exhibited the same activity (see ESI,† Fig. S10), the aromatic LPhAlOAc shows a reduced maximum TOF of 11 h⁻¹ and also forms (thio)ether links. This contrasts the behaviour of the Cr(III) case for which the aromatic backbone showed similar reactivity to other aliphatic ones. Furthermore, the Al(III) complexes were not all equally stable. L^{pro}AlOAc was found to decompose in methanol solution under ambient conditions (ESI,† Fig. S5), while LCyAlOAc, LenA-IOAc and L^{Ph}AlOAc remained stable. Hence, we proceeded with L^{Cy}AlOAc to investigate how the choice of acetate salt affects the catalytic activity.

We employed the series of alkali metal acetates AMOAc (AM = Li, Na, K, Cs), alkaline earth metal acetates AEM(OAc)₂ (AEM = Mg, Ca, Sr, Ba) as well as La(OAc)₃. For all cases, but

LiOAc and Mg(OAc)2, adduct formation could be confirmed via ¹H NMR, (see ESI,† Fig. S7-S9) suggesting that the lighter metals do not seem to be a good size match for the oxygenated binding pocket of L^{Cy}AlOAc. Again, kinetic studies were carried out through ¹H NMR aliquot analysis following the consumption of PTA. It should be noted that although trying to minimise systematic errors associated with the aliquoting method, the precise value of the maximum TOF presented in Table 1 should viewed with caution! Yet our results do allow us to draw conclusions on how metal acetate choice comparatively affects PTA/CHO and PA/CHO ROCOP, which will be discussed in the following. Surprisingly, all added acetate salts led to an improvement of the PTA/CHO ROCOP activity compared to stand-alone L^{Cy}Al yielding poly(ester-thioesters) without (thio)ether links with 39-47% thioester links. The alkali metals Cs, Rb and K are the most active exhibiting very similar kinetic traces (Fig. 2) while Li and Na are substantially less active. The alkaline earth metals Mg, Ca and Ba show similar performances to Li and Na, while an initiation period is observed in the case of Ba. La shows activity ranging between the best alkali and alkaline earth metals while giving the best thioester content of 47%. Next, we were wondering how this compared to the all-oxygen case, as one open question in ROCOP catalysis is, how catalyst selection criteria change when moving down the periodic table from O to S. Therefore PA/CHO ROCOP was investigated under identical conditions with all catalyst combinations and analysed by ¹H NMR aliquot analysis. As can be seen, from the kinetic results presented in Table 1, PTA conversion in PTA/CHO ROCOP is faster than PA conversion in PA/ CHO ROCOP for almost all the catalysts, and we attribute this to the increased nucleophilicity of thiocarboxylate intermediates formed by PTA insertion, compared to carboxylate intermediates from PA insertion. Most interestingly though for L^{Cy}AlOAc with LiOAc, Ca(OAc)₂, Sr(OAc)₂, Ba(OAc)₂ and La(OAc)₃ ether links from CHO homopropagation are formed during the polymerisation process. This is most extreme with Mg(OAc)2, where only polyether and no polyester is formed. Ether formation is associated with broadened polydispersities, in that, while, LCyAlOAc with RbOAc yields a perfect polyester with $M_n = 6.7 \text{ kg mol}^{-1}$ and D = 1.2, the polyether formed with

Table 1 Comparison of PTA/CHO and PA/CHO ROCOP with different M(OAc), with L^{Cy}AlOAc

Alkali metal	PTA conv. ^a [%]	Time [h]	Max. $TOF^b[h^{-1}]$	Ester: thioester: ether ^c	PA conv. ^a [%]	Time [h]	Max. $TOF^b[h^{-1}]$	Ester: ether
Li(ı)	89	6	52	61:39:0	26	4	19	9:91
Na(ı)	94	6	54	59:41:0	94	10	43	100:0
K(ı)	94	0.75	491	55:45:0	98	3	79	100:0
Rb(ı)	99	0.42	638	55:45:0	99	2.5	107	100:0
Cs(I)	98	1.25	248	56:44:0	98	2.5	96	100:0
Mg(II)	91	8	39	56:44:0	0	2	0	0:100
Са(п)	97	5.5	39	59:41:0	53	13	9	26:74
Sr(II)	83	5.5	35	56:44:0	54	4.5	31	20:80
Ва(п)	85	10	25	54:46:0	26	10	8	67:33
La(III)	98	2.5	105	53:47:0	22	10	2	13:87

ROCOPs conducted with 1 eq. L^{Cy}AlOAc: 1 eq. M(OAc)_s: 250 eq. P(T)A: 1000 eq. CHO at 80 °C. ^a Relative integral of aromatic resonances from residual PTA/PA/CHO *versus* polymer in the normalised ¹H NMR spectrum of final aliquot. ^b TOF (turnover frequency) determined by linear fitting of PTA/PA conversion *vs.* time. Data points are excluded once conversion *versus* time plots deviate from linearity due to depleting monomer concentration. ^c Relative integral of ester, thioester vs. ether resonances in normalised ¹H NMR spectrum of the crude polymer.

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100 PTA conversion [%] 60 Na Κ Rb Cs Mg Ca 20 Sr Ba lα 100 200 300 400 500 600

Fig. 2 PTA/CHO ROCOP conversion versus time plots corresponding to Table 1. For PA/CHO ROCOP see ESI,† Fig. S25.

Time [min]

 $Mg(OAc)_2$ shows a $M_n = 18.8$ kg mol^{-1} and D = 1.9 and the poly(ether-ester) with Ca(OAc)₂ a $M_n = 7.0 \text{ kg mol}^{-1}$ and D = 2.0(see Fig. S37, ESI†). In contrast, the same catalysts for PTA/CHO ROCOP produce no ether links whatsoever and this implies that PTA incorporation might occur more facilely than PA incorporation. To investigate this, we performed PTA/PA/CHO terpolymerisation employing L^{Cy}AlOAc with RbOAc at a loading of 1 eq. Cat.: 250 eq. PTA: 250 eq. PA: 1000 eq. CHO, but observed preferential PA incorporation by ¹H aliquot analysis (see ESI,† Fig. S36). Only once PA is consumed PTA/CHO ROCOP occurs to eventually form a polymer with a $M_n = 11.1 \text{ kg mol}^{-1}$ (D = 1.6) after full PA and 64% PTA consumption. Hence, we infer that the linkage selectivity improvement when moving from PA to PTA ROCOP cannot stem from more facile PTA over PA insertion, but much rather the suppression of epoxide homopropagation by the sulfurated species, which needs to be the subject of further investigation.

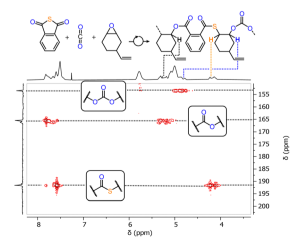
Having established that moving from Cr(III) to Al(III) accelerates heterobimetallic PTA/CHO ROCOP, we wondered whether this also enables the use of epoxides with functional vinyl substituents, as the chromium analogues cannot achieve their copolymerisation with PTA at all (see ESI,† Fig. S38). Hence, we attempted the ROCOP of 4-vinyl cyclohexene oxide (vCHO) with PTA employing L^{Cy}AlOAc with RbOAc at a loading of 1 Cat.: 250 PTA: 1000 vCHO at 80 °C, which resulted in 99% PTA conversion after 45 min to yield a poly(ester-thioester) with $M_{\rm n} = 15.8 \text{ kg mol}^{-1} (D = 1.4) \text{ (Table 2 run #1)}$. This represents an approximately 40% improvement of $M_{\rm n}$ over the analogous PTA/CHO ROCOP. Employment of vCHO rather than CHO to improve molecular weights has been used before in PA ROCOP with related catalysts.30 As vCHO also apparently results in improved molecular weights in PTA ROCOP, we thereafter decreased the catalyst loading to investigate the maximum obtainable molecular weights employing L^{Cy}AlOAc with RbOAc. Reducing the loading to 1 eq. Cat.: 1000 eq. PTA: 2000 eq. vCHO yields a copolymer with $M_n = 45.6 \text{ kg mol}^{-1} (D = 1.6)$ at 80% PTA conversion while moving to 1 eq. Cat.: 2000 eq. PTA: 4000 eq. vCHO further improves the $M_{\rm n}$ to 64.6 kg mol⁻¹ (D = 1.7) at 92%

Table 2 PTA/vCHO ROCOP employing L^{Cy}AlOAc with RbOAc

Cat:PTA:vCHO	Time (h)	PTA conv. a (%)	$M_{\rm n}^{\ b} \left({\rm kg \ mol}^{-1} \right)$	D^b
1:250:1000	0.75	>99	15.8	1.4
1:1000:2000	3	80	45.6	1.6
1:2000:4000	16	92	64.5	1.7
1:4000:6000	24	70	67.2	1.8

T = 80 °C. ^a Relative integral in the normalised ¹H NMR spectrum of aromatic resonances from residual PTA versus polymer. ^b Determined by GPC (gel permeation chromatography) measurements conducted in THF, using narrow polystyrene standards to calibrate the instrument.

PTA conversion (Table 1 run #2 and #3); a further reduction of loading only resulted in slight improvement of M_n which can also be attributed to the decreased maximum PTA conversion of 70% (Table 1 run #4). Although the molecular weights range within the highest achieved for PTA/epoxide ROCOP, they are somewhat lower than the theoretical weights of $M_{n,theo.}$ = 36.1- $265.3 \text{ kg mol}^{-1} (M_{\text{n.GPC}} = 15.8-67.2 \text{ kg mol}^{-1})$. We attribute this to side reactions and protic diol impurities in the vCHO monomer operating as bifunctional initiators that act alongside acetate initiation by the catalyst coligands. This also helps explain the bimodal molecular weight distributions observed as reported previously. Seeking to confirm this notion we deliberately added 20 eq. of a diol chain transfer agent in the form of 1,4-benzendimethanol to a PTA/vCHO ROCOP conducted at 1 eq. Cat.: 250 eq. PTA: 1000 eq. vCHO. This yields a copolymer with a narrowed monomodal molecular weight distribution with a M_n of 3.7 kg mol⁻¹ (D = 1.2) at ca. one third of the molecular weight than without diol chain transfer agent (see ESI,† Fig. S43). This supports that protic impurities are a primary factor for the deviation of molecular weights and broadened bimodal distributions. Nevertheless PTA/vCHO ROCOP could be combined with other ROCOP processes such as CO2/vCHO ROCOP producing some polycarbonate links in PTA/CO₂/vCHO ring-opening terpolymerisation (ROTERP, see ESI,† Section S5 and Fig. 3). Terpolymerisation at a loading of 1 Cat.: 300 PTA: 300 vCHO under 4 bar CO₂ pressure at 80 °C for 1.75 h yields a terpolymer with $M_n = 10.8 \text{ kg mol}^{-1}$ (D = 1.3) at



¹H-¹³C HMBC spectrum (CDCl₃) of PTA/CO₂/vCHO terpolymer.

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58% PTA conversion in which approximately 6% of links are carbonates that result from CO2 insertion and 94% of links are poly(thioester-esters) that result from PTA insertion.§ Increasing the CO_2 pressure to 40 bar yields a terpolymer with M_n = 11.8 kg mol⁻¹ (D = 1.4) at 80% PTA conversion in which approximately 16% of links resulting from CO₂ insertion.

Finally, we sought to demonstrate the utility of the vinyl substituents on the vCHO copolymers that the aluminium catalysis enabled us to incorporate. Hence vulcanisation with elemental sulfur, a waste product of the petrochemical industry, was attempted to crosslink the vinyl functional groups, an underexplored method in the context of ROCOP copolymers (see ESI,† Section S6). Therefore, a powdered blend of 90 wt% vCHO/PTA copolymer (Table 2 run #3) and 10 wt% sulfur was hot processed at 200 °C for 20 minutes and a vulcanized polymer film was obtained as a red, brittle sheet. FTIR analysis shows the stark reduction of the vinylene double bonds upon vulcanisation. In contrast to the parent linear polymer which shows good solubility in organic solvents like DCM, no solubility (95% gel fraction) is observed for the vulcanised material. Furthermore, while the parent linear thioester could be degraded by aminolysis in methanolic ammonia, 26 no degradation was observed for the vulcanised materials further substantiating crosslinking.

In conclusion, we report that moving from chromium to aluminium based heterobimetallic catalysts leads to a drastic improvement of reaction rate in thioanhydride/epoxide ROCOP. Compared to the performance of aluminium catalysis in the alloxygenated anhydride/epoxide ROCOP a general improvement of rate and linkage selectivity was also found. This allowed for the introduction of functional vinyl substituents, the synthesis of high $M_{\rm n}$ materials and novel thioanhydride based terpolymers. Our report highlights clear performance advantages of catalysts featuring sustainable metals in the synthesis of sulfurated polymer which could inspire future catalyst development.

Conflicts of interest

There are no conflicts to declare.

Notes and references

- ‡ For Cr(III), polymerisation were conducted at 100 °C, though at these temperature, we found the Al(III) catalyst to be too fast for the TOF to be determined by the same experimental technique implying that one order of magnitude improvement is the lower limit of the performance
- § This particularly loading was chosen to avoid CO₂/CHO ROCOP after PTA consumption allowing the determination of the relative CO₂ incorporation during the ROTERP process.

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