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Thioanhydride/isothiocyanate/epoxide ringopening terpolymerisation: sequence selective enchainment of monomer mixtures and switchable catalysis†

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We report a new sequence selective terpolymerisation in which three monomers (butylene oxide (BO) A, PhNCS B and phtalic thioanhydride (PTA) C) are selectively enchained into an $(ABA'C)_n$ sequence. PTA/PhNCS/BO ring-opening terpolymerisation ROTERP can be coupled with $CS₂$ ROTERP to generate tetrapolymers and with εDL ROP in switchable catalysis for blockpolymer synthesis.

Heteroatom-containing polymers have strong potential as degradable replacements for polymers based on saturated aliphatic backbones. $1-3$ Often such polymers are synthesised via the living ring opening polymerisation (ROP) of a heterocycle A to produce $(A)_n$ such as poly(thio)esters, poly(thio)carbonate and poly(thio)ethers. $2,4$ In some cases, the ROP of three or four-membered heterocycles A can be coupled with the insertion of heteroallenes or cyclic anhydrides B to generate alternating copolymers $(AB)_n$ in so called ring-opening copolymerisation (ROCOP).^{5,6} Prominent examples include CO_2 /epoxide ROCOP forming polycarbonates or cyclic anhydride/epoxide ROCOP forming polyesters.⁷⁻⁹ However, sulfur-containing variants also exist and relevant to this study are recent reports on isothiocyanate/epoxide ROCOP forming poly(monothioimidocarbonates), isothiocyanate/thiirane ROCOP forming poly (dithioimidocarbonates), cyclic thioanhydride/epoxide ROCOP forming poly(ester-alt-thioesters).^{10–18} Recently we realised alternating ring-opening terpolymerisation (ROTERP) of ternary monomer mixtures comprising thiophtatlic anhydride (PTA), CS_2 and butylene oxide (BO).¹⁹ Here, a simple lithium catalyst (e.g. lithiumbenzyloxide LiOBn or lithiumhexamethyldisilazide LiHMDS) selectively forms poly(ester-alt-ester-alttrithiocarbonates) in up to 98% selectivity with respect to the

erroneous thioester links from PTA/BO ROCOP. The polymer shows an unusual "head-to-head-alt-tail-to-tail" selectivity, meaning that ester groups sit next to tertiary carbon centres while trithiocarbonates sit next to secondary carbon centres. Our results indicated that $PTA/CS_2/BO$ ROTERP formally derives from Li catalysed $CS₂/BO$ ROCOP reported by Werner and co-workers.¹⁷ Both polymerisations are enabled by a O/S exchange reaction step in which an alkoxide chain end from a ring opened epoxide A isomerizes into a thiolate through incorporation of the oxygen into the adjacent link and this isomerised incorporated epoxide (forming a link akin insertion of a virtual thiirane) has been termed $A'.^{17,20}$ Thus, we hypothesized that other monomer combinations which undergo Li catalysed ROCOP could be suitable for sequence selective ROTERP. Xiong and coworkers very recently reported RNCS/ epoxide ROCOP mediated by a simple LiO'Bu catalyst and accordingly we hypothesised that isothiocyanates could undergo ROTERP (Fig. 1).¹² COMMUNICATION

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> Therefore, we investigated the terpolymerisation of PTA/ PhNCS/BO at different monomer ratios and catalyst loadings at 80 °C with lithiumbenzyloxide (LiOBn) as the catalyst (Table 1). Gratifyingly, we find that mixtures comprising 15 eq. PhNCS and 5 eq. BO with LiOBn loadings of 1–8 mol% per equivalent of PTA form poly(ester-alt-ester-alt-dithioimidocarbonates) in quantitative polymer selectivity. The ¹H NMR spectrum (Fig. 2) of the polymer shows two main aryl resonances corresponding to a symmetrically substituted terephthalate unit (δ = 7.67 and 7.47 ppm) in an approximate 4 : 5 integrative ratio with respect to the NPh aromatic resonances indicating a 1 : 1 ratio of the two aromatic moieties.

> Furthermore, there is one main resonance for the $CHR₃$ protons (δ = 5.23 ppm) in a 2 : 4 integrative ratio and one main signal for the CH₂R₂ protons (δ = 3.80–2.90 ppm) in a 4 : 4 integrative ratio to the aromatic terephtalate signals respectively. The ${}^{13}C_1{}^{1}H$ } NMR spectrum (Fig. 2) reveals that the links formed are primarily dithioimidocarbonates $R-S-C(=NPh)-S-$ R (δ = 159.1 ppm, \approx 31% of all links) and arylesters R–C(=O)– O–R (δ = 166.6 ppm, \approx 64% of all links), alongside minor thio-

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Fig. 1 Comparison of different ROCOPs and ROTERP. [Li] = LiN(SiMe₃) or LiOCH₂Ph (LiOBn).^{12,17,19}

ester R–C(=O)–S–R (δ = 192.7 ppm, \approx 2.5% of all links) and monothioimidocarbonate R-O–C(=NPh)–S–R (δ = 156.0 ppm, ≈2.5% of all links) linkages (95% ROTERP links and 5% ROCOP errors). Due to overlapping $^1\mathrm{H}$ NMR resonances for the respective linkages, the linkage ratios were approximated by integration of the relative integrals of the quaternary carbon resonances which proved to correspond well to the linkage ratios determined by integration of the ¹H NMR spectra in related terpolymers.¹⁹ The thioester and monothioimidocarbonate links represent the links of the parent ROCOP reactions between PTA/BO and PhNCS/BO and are inferred to result from incomplete O/S exchange and insertion of PTA or PhNCS into alkoxide chain ends or from insertion of thiolate chain

ends into PTA as previously shown.¹⁹ 2D NMR spectroscopy (Fig. S3 and S4†) further substantiates that dithioimidocarbonate units are positioned adjacent to $CH₂$ groups while arylesters are connected to the tertiary CHMe groups. Hence, we propose a similar "head-to-head-alt-tail-to-tail" selectivity connectivity as for the previously reported ROTERP involving $CS₂$ in place of PhNCS. Furthermore, no ether links were detected in the polymer, which are a common side products formed in related ROCOPs.^{5,21} The respective resonance ratios remain unchanged after multiple precipitations from DCM : MeOH or THF : pentane confirming that all links are part of the same polymer. Linkage identity could be further substantiated by the ATR-IR spectrum (Fig. 2) showing an arylester $C=O$ stretch at \tilde{v} = 1716 cm⁻¹ as well as a dithioimidocarbonate C=N stretch at \tilde{v} = 1563 cm⁻¹.^{10,19} MALDI-TOF analysis unfortunately only led to decomposition of the materials and no signals could be identified as previously reported for sulfurrich polymers. $22,23$ However the OBn initiator can be identified to be part of the purified polymers (Fig. S5†) suggesting the formation of linear as opposed to cyclic chains. The polymers are colourless amorphous solids (T_g = 30.6 °C, Fig. S8†) with good thermal stability ($T_{d,5\%}$ = 230.1 °C, Fig. S10†). SEC analysis of the obtained materials at catalyst loadings of 1–8 mol% vs. PTA shows that the methodology can yield polymers with molecular weights ranging from $M_n = 2.55$ to 14.00 kg mol⁻¹ ($D = 1.15$ -1.49, Table 1 and Fig. S19†). Attempting lower LiOBn loadings did not result in any polymerisation. Employing Na or K in place of Li resulted in unappreciable turnover supporting that Li acts as a catalyst rather than a spectator countercation for the OBn initiator (Table 1 run #9 and #10). Decreasing the amount of PhNCS in the initial monomer mixture (and therefore increasing the PTA concentration as PhNCS acts as a cosolvent) led to more erroneous thioester links (Table 1 run #6–#8) and a similar amount of monothioimidocarbonate links (ca. 3% of all links). This can be rationalized by kinetic competition between O/S Communication
 $Q_{\text{total}} = \frac{1}{2} \left(\frac{1}{2} \left(\frac{1}{2} \right) \right)$
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^a Copolymerisation at T = 80 °C, LiOBn generated in situ from LiHMDS and BnOH (see ESI†). ^b Determined by comparison of the relative integrals, in the normalised ¹H NMR spectrum (CDCl₃, 25 °C, 400 MHz), of terepthalate CH resonances due to (co/ter)polymer and PTA.
^c Determined by comparison of the relative integrals, in the normalised ¹H NMR spectru due to (co/ter)polymer and cyclic thioimidocarbonate and polyether. ^d Determined by comparison of the relative integrals, in the normalised the
¹³C{¹H} NMR spectrum (CDCl₃, 25 °C) of resonances due to ester relativ measurements conducted in THF, using narrow MW polystyrene standards to calibrate the instrument. JNaHMDS was employed in place of LIHMDS. ^g KHMDS was employed in place of LIHMDS. h CS₂.

rearrangement and propagation from alkoxide intermediates. In these cases, we also observe the onset of PhNCS/BO ROCOP to form poly(monothioimidocarbonates) once all PTA is consumed as confirmed by 1 H (Fig. S6†) and 13 C NMR (Fig. S7†) spectroscopy. This leads to poly(monothioimidocarbonate) blocks forming adjacent to the ROTERP blocks which also explains the higher molecular weights obtained at the same PTA loadings. The new monomer combination is furthermore compatible with PTA/CS₂/BO ROTERP. Tetrapolymerisation of PTA and BO with PhNCS and CS_2 results in poly(ester-alt-esteralt-heterocarbonate) formation in which both RNCS (forming dithioimidocarbonates) and CS_2 (forming trithiocarbonates) are incorporated into the polymer. NMR (Fig. S11–S15†) and FTATR-IR (Fig. S17†) clearly show the presence of trithiocarbonate (δ = 222.8 ppm, \tilde{v} = 1063 cm⁻¹) links in addition to the other links from PTA/PhNCS/BO ROTERP. Corresponding to Table 1 run #11, a polymer comprising 67% arylester, 23% trithiocarbonate, 6% dithioimidocarbonate, 3.5% thioester and 0.5% monothioimidocarbonate links is produced (96% ROTERP links and 4% ROCOP errors). Apparently, PTA/PhNCS/ BO ROTERP follows a similar polymerisation mechanism (Fig. $S31\dot{}$) as the one proposed for PTA/CS₂/BO ROTERP. It involves a central O/S rearrangement step and preferential insertion of PTA into alkoxide chain ends and of the heteroallene into thiolate chain ends at which point a mixture of heteroallenes can be employed which are then both incorporated at this insertion step. Interestingly, although PhNCS and $CS₂$ are employed in an equimolar ratio, $CS₂$ is incorporated preferentially in 77% selectivity. We could confirm this observation by varying the initial PhNCS: CS_2 ratio in which CS_2 was always incorporated to a greater degree than employed in the initial monomer feed (see Fig. S18†). In ROCOP, switchable catalysis has been established as an elegant and valuable tool to synthesise blockpolymers with useful material properties. $24-26$ Here a suitable catalyst first mediates the ROP of for example cyclic esters (e.g. εDL forming PDL) with epoxides present in the mixture until the second ROCOP monomer (e.g. $CO₂$) is

added causing immediate termination of ROP and the onset of (e.g. $CO₂/epoxide forming polycarbonate) ROCOP to form a$ ROCOP block connected to the ROP polymer. We recently showed that ROTERP is also suitable for switchable catalysis in combination with the lithium catalysed ROP of ϵ DL.¹⁹ Having identified a new heteroallene that undergoes ROTERP, we were intrigued whether this monomer combination is also suitable for the construction of blockpolymers via switchable catalysis. Accordingly, we added PhNCS (750 eq. per LiOBn) and PTA (50 eq.) to polymerising εDL (50 eq.) in BO (250 eq.) after 15 min at room temperature which completely and immediately stops the occurrence of εDL ROP. Heating to 80 °C initiates ROTERP and a poly(ester-alt-ester-alt-dithioiminocarbonate) block grows from the PDL-chain-end until the reaction is stopped after 30 min. Under these conditions the ROTERP block consists of 64% arylester links, 31% dithioimidocarbonate links and 2.5% erroneous thioester and monothioimidocarbonate links, respectively (95% ROTERP links and 5% ROCOP errors). Switchable catalysis and block polymer formation were established by various methods: (i) no εDL is consumed after ROTERP starts (Fig. S21†) and the ¹³C{¹H} PDL (δ = 173.2 ppm, Fig. S23†) remains unchanged showing that ROP stops and that no transesterification between blocks occurs; (ii) the number averaged molecular weight shifts from $M_n = 8.26$ ($D =$ 1.11) to 14.57 kg mol⁻¹ ($D = 1.24$, Fig. 3), which shows the growth of existing chains rather than the initiation of new ones; (iii) $3^{31}P$ end group analysis shows the consumption of all PDL end groups (Fig. S30†);²⁷ (iv) the composition of the resulting blockpolymer remains unchanged through multiple precipitations from DCM/MeOH and THF/pentane supporting that the blocks are joint; (vii) DSC analysis exhibits two T_g 's at − 46.5 °C for the ROP block and 35.5 °C for the ROTERP block suggesting microphase separation in the solid-state which could be confirmed by AFM (Fig. 3); 28 (viii) TGA analysis shows a stepwise thermal decomposition profile with two $T_{\text{d,onset}}$ at approximately 235 °C for the ROTERP block and 300 °C for the ROP block (Fig. 3). In conclusion, we have identified a new

Fig. 3 (Left) εDL ROP to ROTERP switchable catalysis sequence. (Right) overlayed SEC traces before and after switch as well as TGA, DSC and 3d image of AFM tapping topography (hight range 4 nm) of the obtained blockpolymer. 3d display of topography is overlaid with a false colour map of the AFM tapping phase which differentiates between the blockpolymer's components.

monomer combination that undergoes lithium catalysed sequence selective terpolymerisation. Mixtures of PTA/PhNCS/ BO forming poly(ester-alt-ester-alt-dithioimidocarbonate)s in up to 95% selectivity with respect to the erroneous links from PhNCS/BO and PTA/BO ROCOP. This has enabled the synthesis of complex tetrapolymers from quarternary monomer mixture or via switchable catalysis. Our results establish ROTERP as a valuable methodology for the synthesis of heteroatom containing blockpolymers.

Conflicts of interest

There are no conflicts of interests.

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