Synthesis and investigation of chiral poly(2,4-disubstituted-2-oxazoline) based triblock copolymers, their self-assembly and formulation with chiral and achiral drugs

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Keywords: poly(2-oxazoline), stereoactive polymers, optically active, curcumin, paclitaxel, ibuprofen

Abstract

Considering the largely chiral nature of biological systems, there is interest in chiral drug delivery systems. Here, we investigate for the first time polymer micelles based on poly(2-oxazoline)s (POx) ABA-type triblock copolymers with chiral and racemic hydrophobic blocks for the formulation of chiral and achiral drugs. Specifically, poly(2-ethyl-4-ethyl-2-oxazoline) (pEtEtOx) and poly(2-propyl-4-methyl-2-oxazoline) (pPrMeOx) were used as hydrophobic block B and poly(2-methyl-2-oxazoline) (pMeOx) as hydrophilic block A. Using these triblock copolymers, nanoformulations of curcumin (CUR), paclitaxel (PTX) as well as chiral (R and S) and racemic ibuprofen were prepared. For CUR and PTX, the maximum drug loading dependent significantly on the structure of the hydrophobic repeat units, but not the chirality. In contrast, the maximum drug loading with chiral/racemic ibuprofen was neither affected by the polymer structure nor by chirality, but minor effects were observed with respect to the size and size distribution of the drug loaded micelles.
Introduction

Chirality is an essential property for many biologically relevant molecules, including bio(macro)molecules like sugars, amino acids and their polymers (proteins, polysaccharides, DNA/RNA). Despite the seemingly minute structural difference, enantiomers of drugs can exhibit significant differences in their biological activity such as pharmacology, toxicology, pharmacokinetics, pharmacodynamics, etc.¹ For instance, for the analgesic drug ibuprofen, its S-enantiomer has a higher efficacy than its stereoisomer.² In the worst case, a stereoisomer can produce undesired or toxic effects. A notorious example is thalidomide, which was first marketed in 1957 as a racemic mixture, but due to severe teratogenic effects (phocomelia, amelia) caused by its S-enantiomer, it was withdrawn from the market in the 1960.³ Therefore, the isolation of therapeutically active enantiomers is of utmost importance. Chiral resolution can be achieved, inter alia, by chiral chromatography, in which a chiral compound is immobilized on the surface of the stationary phase.² Accordingly, it may appear logical to utilize chiral drug delivery systems to also preferentially interact/solubilize a drug enantiomer of interest. Therefore, synthetic stereoactive polymers, in which repeating units feature chiral centers have attracted some attention. Recent studies on poly(lactide) (PLA),³ poly(glutamic acid)⁴ and poly(leucine) based block copolymers⁵ have reported the effect of polymer stereoregularity on the physicochemical and functional properties of their self-assembled nanostructures.⁶ Feng et al. investigated micelles based on methoxy-poly(ethylene glycol)-b-poly(l-lactide) micelles (mPEG-b-PLLA, L-micelles) and mPEG-b-poly(d-lactide) micelles (mPEG-b-PDLA, D-micelles) to solubilize the glycosylated antibiotic nocathiacin I (containing multiple chiral centres) and other chiral compounds (containing D- or L-sugars).⁷ They observed that the nocathiacin I loaded D-micelles exhibited better loading efficiency and smaller particle size than that of L-micelles. Also, for micelles loaded with other chiral compounds, D- and L-micelles showed a marked difference in particle size, even though the loading efficiency between D- and L-micelles were not significantly different. Hu et al. loaded insulin in stereo multiblock copoly(lactide)s (smb-PLAs) with different stereoregularity.⁸ They found that smb-PLAs
with a high stereoregularity show much higher insulin loading efficiency than the atactic PLA. However, the effect how polymer stereoregular blocks change the properties such as micellar size, micellar thermodynamic stability, drug loading and cell interaction are generally not broadly investigated and understood, in particular beyond poly(lactide).

Poly(2-oxazoline) (POx) are a family of polymers of the larger group of polymers classified as pseudo-polypeptides. In the past two decades, POx have gained increasing interest for a wide range of applications, especially in the biomedical field, e.g., drug, protein and gene delivery, tissue engineering, regenerative medicine, 3D bioprinting and biofabrication, etc. The 2-oxazoline monomers with different substituents in the 2-position can be tailor-made by relative easy and straightforward synthesis, and numerous monomers have been used in cationic ring-opening polymerization to obtain the corresponding poly(2-alkyl/aryl-2-oxazoline)s with narrow molar mass distributions and a wide range of physico-chemical properties. Among them, the hydrophilic POx, poly(2-methyl-2-oxazoline) (pMeOx) and poly(2-ethyl-2-oxazoline) (pEtOx) have been studied intensively as they exhibit stealth/protein repellent effects. They are regularly discussed as one of the potential alternatives to PEG. pMeOx and pEtOx have been used widely as hydrophilic polymer component, e.g., in polymer nanoparticles and micelles (in combination with other hydrophobic POx e.g., with 2-butyl, 2-nonyl or 2-isopropyl-2-oxazoline), polymer-peptide conjugates, polymer-protein conjugates, lipopolymers for liposome stabilization, polymer-drug conjugates, as well as antimicrobial polymers. While the toolbox of 2-substituted 2-oxazolines and their corresponding polymers has been expanding rapidly, however, work on monomers and polymers with substituents in the 4- and 5-position in the 2-oxazoline monomers is quite limited. The proof-of-principle that such 2,4-disubstituted POx are accessible was provided by Saegusa et al., who synthesized optically active poly(ethylenimine) derivatives by ring-opening polymerization of 4-substituted-2-oxazoline and 4,5-disubstituted-2-oxazoline for the first time. Much later Schubert, Hoogenboom and co-workers reported the synthesis and properties of chiral poly(2-R-4-
ethyl-2-oxazoline)s ($R = \text{ethyl, butyl, octyl, nonyl, undecyl}$),\textsuperscript{35-38} and further discussed the self-assembly of chiral amphiphilic block copolymers composed of a hydrophilic block of pEtOx and a hydrophobic block of poly(R-2-butyl-4-ethyl-2-oxazoline) ($p^R_{\text{BuEtOx}}$) or racemic $p^{RS}_{\text{BuEtOx}}$.\textsuperscript{39} They found that varying the hydrophobic/hydrophilic ratio in the copolymers could control the type of self-assembled structures from spherical and cylindrical micelles to sheets and vesicles. Unfortunately, in this contribution, no direct comparison of chiral and racemic polymers of the same composition was provided.\textsuperscript{39} Jordan et al. investigated the influence of chirality on the lower critical solution temperature (LCST) behavior of water soluble poly(2-alkyl-4-methyl-2-oxazoline)s (alkyl: methyl, ethyl).\textsuperscript{40} Introduction of chirality via the alkyl substituents in the main chain of poly(2,4-disubstituted-2-oxazoline)s allows for the formation of secondary structure in aqueous and non-aqueous environments as well as in bulk.\textsuperscript{37, 40} In a patent, Schmidt and Bott proposed a possible application of poly(4(S)-4-ethyl-2-phenyl-2-oxazoline) in the separation of enantiomeric mixtures of D,L-2-chloro-4-methyl-phenoxy-propionic acid methylester.\textsuperscript{41}

As many drugs, including hydrophobic ones, are chiral, it is interesting to study the effect of chirality on POx based drug formulations. While in recent years, significant structure property relationships regarding hydrophobic drug formulations using a large variety of amphiphilic POx have been studied,\textsuperscript{42-45} a drug formulation based on main chain chiral POx has not been studied before. Aiming to improve our understanding of stereoregular polymers as drug carriers in general and to enlarge the toolbox of POx based drug formulations in particular, we have synthesized two series of chiral POx based ABA amphiphilic triblock copolymers. We explored their potential application for drug formulation and their selectivity and affinity for particular drug enantiomers. Specifically, ABA triblock copolymers were synthesized via living cationic ring-opening polymerization (LCROP), comprising pMeOx as hydrophilic blocks A and chiral poly((R)-2-ethyl-4-ethyl-2-oxazoline) ($p^R_{\text{EtEtOx}}$), poly((S)-2-ethyl-4-ethyl-2-oxazoline) ($p^{S}\text{EtEtOx}$) and racemic poly((RS)-2-ethyl-4-ethyl-2-oxazoline) ($p^{RS}_{\text{EtEtOx}}$) as well as poly((R)-2-propyl-4-methyl-2-oxazoline) ($p^R_{\text{PrMeOx}}$), $p^{S}\text{PrMeOx}$ and $p^{RS}_{\text{PrMeOx}}$ as hydrophobic block B. Their aqueous solubility, optical
activity, thermal properties and drug loading with respect to chirality were investigated. The corresponding hydrophobic homopolymers were also synthesized and investigated to help understand the properties of triblock copolymers. Curcumin (CUR) and paclitaxel (PTX) were used as models of common hydrophobic drugs, whereas R-ibuprofen (R-IBU), S-IBU, RS-IBU were used as model compounds for chiral drugs and a racemic drug mixture.

Scheme 1. (a) Chemical structures of the A-B-A triblock copolymers used in this study, where hydrophilic blocking A are poly(2-methyl-2-oxazoline) (pMeOx) and the hydrophobic block B is poly(2-ethyl-4-ethyl-2-oxazoline) (pEtEtOx, pEtEtOx, pEtEtOx) or poly(2-propyl-4-methyl-2-oxazoline) (pPrMeOx, pPrMeOx, pPrMeOx). (b) Chemical structure of the model drugs used in this study, paclitaxel, curcumin and ibuprofen. (c) Schematic representation of the formulation procedure (thin film method).
Experimental Section

Materials.

All substances for the preparation of the polymers were purchased from Sigma-Aldrich (Steinheim, Germany) or Acros (Geel, Belgium) and were used as received unless otherwise stated. D-Alaninol (purity 98%), (S)-(+-)-2-Amino-1-propanol (purity 98%), (R)-(+-)-2-Amino-1-butanol (purity 98%), (S)-(+-)-2-Amino-1-butanol (purity 98%) were purchased from abcr (Karlsruhe, Germany). DL-2-Amino-1-propanol (purity 98%) and DL-2-Amino-1-butanol (purity 98%) were purchased from TCI (Eschborn, Germany). Curcumin (CUR) powder from Curcuma longa (Turmeric) was purchased from Sigma-Aldrich (curcumin = 79%; demethoxycurcumin = 17%, bisdemethoxycurcumin = 4%; determined by HPLC analysis). Paclitaxel (PTX) was purchased from LC Laboratories (Woburn, MA, USA). (R)-(+-)-Ibuprofen (R-IBU) (98.5%) was purchased from MedChemExpress (distributor Hycultec, Beutelsbach, Germany). (S)-(+-)-Ibuprofen (S-IBU) (99%) and racemic Ibuprofen (RS-IBU) (pharmaceutical secondary standard; certified reference material) was purchased from Sigma-Aldrich. Deuterated solvents for NMR analysis were obtained from Deutero GmbH (Kastellaun, Germany).

The monomers (R)-2-ethyl-4-ethyl-2-oxazoline (REtOx), (S)-2-ethyl-4-ethyl-2-oxazoline (SEtOx), (RS)-2-ethyl-4-ethyl-2-oxazoline (RSOx), (R)-2-propyl-4-methyl-2-oxazoline (RPrMeOx), (S)-2-propyl-4-methyl-2-oxazoline (SPrMeOx) and (RS)-2-propyl-4-methyl-2-oxazoline (RSPrMeOx) were prepared following the procedure by Witte and Seeliger et al.11-12 For monomer synthesis and characterization, see Fig. S1-4 in SI. The substances used for polymerization, specifically methyl trifluoromethylsulfonate (MeOTf), 2-methyl-2-oxazoline (MeOx), REtOx, SEtOx, SRSOx, RPrMeOx, SPrMeOx, RSPrMeOx and sulfolane were refluxed over CaH2, distilled and stored under argon.

Polymer synthesis. The polymerization and workup procedures were carried out similar to Lübtow et al. described previously.43 The initiator MeOTf was added to a dried and argon flushed flask and dissolved in
the respective volume of sulfolane, followed by monomer addition. Subsequently, the reaction mixture was heated to 100 °C or 130 °C (according to the type of monomers, see SI). Reaction progress was controlled by $^1$H-NMR spectroscopy. After complete consumption of monomer, additional monomer was added in case of block copolymer synthesis, or termination was carried out by addition of 1-Boc-piperazine (PipBoc) at 50 °C. Subsequently, K$_2$CO$_3$ was added, and the mixture was stirred at 50 °C. The crude product was purified by dialysis. For polymer synthesis details and characterization, see Fig. S5-7 and Fig. S11-12 in SI.

**Nuclear magnetic resonance spectroscopy (NMR).** NMR spectra were measured with a Fourier 300 ($^1$H; 300 MHz), Bruker Biospin (Rheinstetten, Germany) at 298 K. All chemical shifts are given in ppm. The spectra were calibrated to the signals of residual protonated solvent signals (e.g., CDCl$_3$: 7.26 ppm).

**Gel Permeation Chromatography (GPC).** GPC of polymers was performed on an Agilent 1260 Infinity System, Polymer Standard Service (Mainz, Germany) with hexafluoroisopropanol (HFIP) containing 3 g/L potassium trifluoroacetate; precolumn: 50 mm × 8 mm PSS PFG linear M (particle size 7 μm); main column: 8×300 mm AppliChrom ABOA HFIP-P350 (pore size 0.1-1000 kDa). The columns were kept at 40 °C and flow rate was 0.3 mL/ min. Prior to each measurement, samples were dissolved in HFIP/potassium trifluoroacetate and filtered through 0.2 μm PTFE filters, Roth (Karlsruhe, Germany). Conventional calibration was performed with PEG standards (0.1-1000 kg/mol) and data was processed with Win-GPC software.

GPC of pPrMeOx homopolymers was also performed on an alternative GPC system, Malvern GPCMax system (Malvern, UK) with a VE 3580 RI detector, PSS Polymer Standard Service (Mainz, Germany); two Malvern LC4000L column: 300×8 mm (exclusion limit: 400 kDa). Chloroform was used as the eluent with a 100 μL sample volume injection. The columns were kept at 35 °C and flow rate was 1 mL/min. Prior to each measurement, samples were dissolved in chloroform and filtered through 0.2 μm PTFE filters, Roth.
(Karlsruhe, Germany). Conventional calibration was performed with polystyrene standards (1.2-40 kDa) and data was processed with OmniSEC software.

**Thermogravimetric Analysis (TGA).** A TG 209 F1 IRIS, NETZSCH (Selb, Germany) was used for thermal analysis. The samples (5-10 mg) were added into aluminium oxide crucibles (NETZSCH) and heated under synthetic air from 30 °C to 900 °C (10 °C/min) while detecting the mass loss.

**Differential Scanning Calorimetry (DSC).** DSC was performed on a DSC 204 F1 Phoenix, NETZSCH under N₂-atmosphere (20.0 mL/min). The samples were placed in aluminium pans with pierced crimped-on lids and heated from 30 °C to 190 °C and subsequently cooled to -50 °C (10 °C/min). The heating/cooling cycle was repeated two additional times from -50 °C to 190 °C (10 °C/min).

**X-ray diffraction (XRD).** XRD measurements were performed on a D8 Advance diffractometer with DaVinci design (Bruker AXS, Karlsruhe, Germany). The following measurement parameters were applied: a 2θ range of 5-60°, a step size of 0.02° 2θ, an integration time of 2 s, copper Kα radiation, generator settings of 20 kV and 5 mA and a 0.344° divergence slit. The data was exported by software DIFFRAC.EVA (Bruker AXS, Karlsruhe, Germany).

**Fluorescence Spectroscopy-Critical Micelle Concentration (CMC).** CMC of triblock copolymers was determined using fluorescence probe pyrene. Pyrene solutions (24 µM, 5.0 mg/L in acetone) were added to glass vials and followed by acetone removal by a gentle stream of argon. Afterwards, various amounts of aqueous polymer stock solutions were added, and the solutions were diluted with water (Millipore) to yield a final pyrene concentration of 5×10⁻⁷ M. The samples were shaken gently for 30 min and stored overnight at ambient temperature (≈20 °C) under the exclusion of light. Fluorescence measurements were performed in a FP-8300, Jasco from 360 nm to 400 nm (λₑₓ = 333 nm) at 25 °C with 10x10 mm fluorescence cuvettes.
The fluorescence spectrum of pyrene shows five characteristic vibronic bands around 360-400 nm. The ratio of the fluorescence intensities of the first and third vibronic bands of pyrene ($I_1:I_3$ ratio) increases characteristically with increasing polarity of the probe environment. The CMC was determined as the concentration at which the fitted $I_1:I_3$ ratio decreased to 90% of its initial value.

**Circular dichroism (CD) characterisation.** CD spectra were measured in methanol or water solutions (0.1 g/L polymer concentration) with a JASCO J-810 circular dichroism spectrometer (JASCO International Co., Ltd., Tokyo, Japan). The following scanning conditions were used: 200 nm/min scanning rate; 1 nm bandwidth; 0.5 nm data pitch; 1 s response time; and 3 accumulations. Samples were measured in a 1 mm path length quartz cuvette (110-QS, Hellma Analytics).

**Drug Formulation.** Drug-loaded polymer micelles were prepared using the thin film hydration method. Ethanolic polymer (20 g/L), curcumin (5 g/L), paclitaxel (5 g/L), R-IBU (5 g/L), S-IBU (5 g/L) or RS-IBU (5 g/L) stock solutions were mixed in desired ratio. After complete removal of the solvent at 50 °C under a mild stream of argon, the films were further dried in vacuum ($\leq$0.2 mbar) for at least 30 min. Subsequently, preheated (37 °C) H$_2$O (Millipore) was added to obtain final polymer (10 g/L) and desired drug concentrations. To ensure complete solubilization, the solutions were shaken at 55 °C for 15 min, at 1250 rpm with a Thermomixer comfort (Eppendorf AG, Hamburg, Germany). Non-solubilized drug was removed by centrifugation for 5 min at 9000 rpm (relative centrifugal force (rcf) 7788 g) by a MIKRO 185 (Hettich, Tuttlingen, Germany). The solubilization experiments were performed with three individually prepared samples and results are presented as means ± standard deviation (SD).

The loading capacity (LC) and loading efficiency (LE) were calculated using the following equations:

$$LC = \frac{m_{drug}}{m_{drug} + m_{polymer}} \times 100\%$$
$LE = \frac{m_{\text{drug}}}{m_{\text{drug,added}}} \times 100\%$

Where $m_{\text{drug}}$ and $m_{\text{polymer}}$ are the weight amounts of the solubilized drug and polymer excipient in solution and $m_{\text{drug,added}}$ is the weight amount of the drug initially added to the dispersion. No loss of polymer during micelles preparation was assumed.

**UV-Vis spectroscopy.** CUR quantification was performed by UV-Vis absorption on a BioTek Eon Microplate Spectrophotometer, Thermo Fisher Scientific (Waltham, MA) using a calibration curve obtained with known amounts of CUR, dissolved in ethanol. Samples were prepared in Rotilabo F-Type 96 well plates, Carl Roth GmbH & Co. KG (Karlsruhe, Germany) at a constant volume of 200 µL. Spectra were recorded from 300 to 700 nm at 25 °C. Curcumin absorption was detected at 430 nm. Prior to UV-Vis absorption measurements, the aqueous formulations were appropriately diluted with ethanol to give a final absorbance between 0.3 and 2.5 (diluted at least 1/20 (v/v)).

**High Performance Liquid Chromatography (HPLC) analysis.** HPLC analysis was carried out using a prominence LC-20A modular HPLC system (Shimadzu, Duisburg, Germany) equipped with a system controller CBM-20A, a solvent delivery unit LC-20 AT (double plunger), an on-line degassing unit DGU-20A, an auto-sampler SIL-20AC, a photo-diode array detector SPD-M20A, a column oven CTO-20AC, and a refractive index detector RID-20A. As stationary phase, a ZORBAX Eclipse Plus, Agilent (Santa Clara, CA, USA) C18 column (4.6 x 100 mm; 3.5µm50mm x 4 mm) was used. The volume of samples injected was 20 µL and elution was performed using a mobile phase of H₂O and acetonitrile (ACN) containing 0.05% trifluoroacetic acid (TFA) at 40 °C and a flow rate of 1 mL/min.

Quantification of paclitaxel (PTX) and ibuprofen (IBU) was performed at 220 nm. For PTX, within the first 10 min, the proportion of ACN was increased from 40% to 60%. Solvent proportion was kept constant for 5 min prior to decrease it to initial proportion of 40% ACN within 0.5 min. For IBU, the proportion of
ACN was increased from 40% to 60% ACN within the first 10 min, afterwards was increased to 80% ACN in 0.1 min and kept constant for 1.9 min, and finally was decreased to initial proportion of 40% ACN in 0.1 min. The retention times were 8.2 min for PTX and 9.5 min for IBU.

**Dynamic Light Scattering (DLS).** Triblock copolymer aqueous solutions, CUR or IBU loaded formulations were prepared with PBS (pH 7.4) and measured on Zetasizer Nano ZSP from Malvern, (Malvern Instruments, Worcestershire, UK) in disposable cuvettes (UV cuvettes semi micro, BRAND GmbH, Wertheim, Germany) at ambient temperatures (≈25 °C). Data was analysed by using Zetasizer software 7.11. All the samples were measured after filtration using 0.45 µm PVDF syringe filter (Rotilabo, Karlsruhe). The filtered samples were further diluted with PBS and measured again to exclude variation due to dilution effect. The data obtained are the average of three measurements.

**Long-term stability studies.** For long-term stability studies, formulated IBU was stored at ambient conditions (≈25 °C). The samples were collected at day 0, 1, 8, 20, 30 and day 60. Before the determination of the drug loading by HPLC, all samples were centrifuged for 5 min at 9000 rpm with a MIKRO 185 (Hettich, Tuttlingen, Germany). Long-term stabilization experiments were performed with three individually prepared samples and results are presented as means ± SD, quantification was carried out as described previously.

**Statistical analysis.**

Statistical significance was calculated by Student’s t-test. Differences with a value of p < 0.05 were considered statistically significant.
Results and discussion

Synthesis and characterization of homopolymers

The EtEtOx and PrMeOx series monomers were prepared following the procedure by Witte and Seeliger et al. 11-12. Nitrile and chiral or achiral alkanolamine were heated for 2-6 d under the catalysis of zinc acetate dihydrate to produce corresponding monomers. Detailed description of monomer synthesis and characterization can be found in SI, Fig. S1-4.

Since synthesis of poly(2-propyl-4-methyl-2-oxazoline) (pPrMeOx) and its characteristics were not reported before, the respective homopolymers were synthesized and characterized first. Knowledge of the homopolymerization and homopolymer properties are also important to support the synthesis of corresponding polymer amphiphiles and their characterization. Also, this allows a direct comparison with its isomer pEtEtOx. Accordingly, the homopolymers p\textsuperscript{6}EtEtOx, p\textsuperscript{5}EtEtOx, p\textsuperscript{6}EtEtOx, p\textsuperscript{6}PrMeOx, p\textsuperscript{5}PrMeOx and p\textsuperscript{6s}PrMeOx were prepared by LCROP. Using a [M]\textsubscript{0}/[I]\textsubscript{0} = 20, the complete polymerization of monomers p\textsuperscript{6}EtEtOx, p\textsuperscript{5}EtEtOx and p\textsuperscript{6s}EtEtOx was achieved only after about 45 h at 130 °C. In comparison, the monomer consumption for p\textsuperscript{6}PrMeOx, p\textsuperscript{5}PrMeOx and p\textsuperscript{6s}PrMeOx was complete after only approximately 24 h at 130 °C, which can be attributed to decreased steric hinderance due to the smaller Me substituent at the 4-position affecting the nucleophilic attack of monomer at the 5-position. We tested the solubility of homopolymers in different solvents used in following experiment. All these homopolymers are well soluble in HFIP (≥ 5 g/L) and excellently soluble in chloroform, methanol and ethanol (all ≥ 200 g/L), but poorly soluble in water (< 0.5 g/L in water). The poor water solubility is roughly in line with the well-known side chain size dependence of POx solubility.\textsuperscript{51} Interestingly, when the saturated aqueous solutions of the homopolymers equilibrated at 4 °C were brought to room temperature, the previously clear solutions turned turbid. Surprisingly, the concentration of saturated aqueous solutions (4 °C) was 8-10 g/L (Table 1), much higher than that at room temperature. This observation shows that these homopolymers
exhibit a thermoresponsive behaviour (LCST-type) in water. The LCST behaviour is well known for POx and POzi with C3 side chains\textsuperscript{52}, but, to the best of our knowledge, has not been described for higher substitution in published research.

The homopolymers were further characterized by \textsuperscript{1}H-NMR and GPC, the results of which is detailed in the supporting information (Fig. S5-9) and Table 1. The polymerization was terminated with N-Boc-piperazine (PipBoc), wherein the Boc moiety yields a sharp and intense singlet in \textsuperscript{1}H-NMR spectrum (Fig. S6-7), facilitating end-group analysis. Based on this, p\textsuperscript{8}EtEtOx, p\textsuperscript{5}EtEtOx and p\textsuperscript{8}EtEtOx have a degree of polymerization (DP) of 21, 23, 21, respectively, while p\textsuperscript{8}PrMeOx, p\textsuperscript{5}PrMeOx and p\textsuperscript{8}PrMeOx have a DP of 22, 22 and 23, respectively. Both of pEtEtOx and pPrMeOx series are reasonably close to the targeted DP of 20 (from monomer to initiator ratio [M]:[I]), but it must be noted that all polymer signals are very broad and overlapping, making an accurate end-group analysis difficult. The dispersity as determined by GPC (Table 1, Fig. S8 a and b) was very low with values ranging from Đ = 1.09-1.16. The molar mass obtained from GPC is considerably smaller than from end group analysis by \textsuperscript{1}H-NMR, which can be attributed to a different solution behaviour in the eluent compared to the utilized PEG-standards used for calibration in GPC. Besides, the pPrMeOx series was also characterized using another GPC system (chloroform as eluent), to elucidate the effects of eluent and calibration standard (Fig. S8c).

To investigate the thermal stability of homopolymers, TGA was performed in the temperature range of 30 °C to 900 °C (Fig. S9). Around 220 °C, 3-4 % mass loss was observed in all homopolymers, which is consistent with the Boc weight percent of homopolymers.\textsuperscript{53} Onset temperature of major mass loss T\textsubscript{d} of all homopolymers is around 350 °C, which is consistent with widely reported thermal stability of POx-based polymers.\textsuperscript{44, 54}

DSC measurements were performed from -50 °C to 190 °C in order to determine the thermal transitions of the homopolymers. In the given temperature range, no melting temperature (T\textsubscript{m}) of homopolymers
was observed (Fig. 1), which may be connected to the rather short chain length. The glass transition temperature \((T_g)\) of pEtEtOx series homopolymer, \(p^6\)EtEtOx, \(p^5\)EtEtOx and \(p^{55}\)EtEtOx, are very similar (around 80 °C), i.e., the different chirality of pEtEtOx series homopolymers, as expected, does not affect chain segment mobility (Table 2). Similarly, in the pPrMeOx series, of the \(T_g\) values (around 55 °C) for \(p^6\)PrMeOx, \(p^5\)PrMeOx and \(p^{55}\)PrMeOx are essentially identical. Clearly, the \(T_g\) of pPrMeOx series is lower than that of pEtEtOx, which signifies higher chain segment mobility for pPrMeOx than pEtEtOx. There has been some research on the thermal properties of various POx and poly(2-alkyl-2-oxazines) (POzi) (Table 2). Two conclusions were drawn in this previous work: the \(T_g\) of POx decreased linearly with increasing carbon number in the side chain (from 1 to 5 carbon atoms), while linear POzi have lower \(T_g\) than the POx with same side-chain, which is attributable to the additional methylene unit in the main chain. Previously reported \(p^6\)BuEtOx \(([M]/[I]=60, T_g=52 °C)\) has two methylene groups more in side-chain than pEtEtOx and a \(T_g\) about 30 °C lower than the \(T_g\) of \(p^6\)EtEtOx \((T_g=82 °C)\), which shows that with increasing number of carbon atoms in the side-chain the \(T_g\) also decreases for main-chain branched POx. Besides, the \(T_g\) of pEtEtOx is 20 °C higher than that of pEtOx, pPrMeOx has a 15 °C higher \(T_g\) than poly(2-propyl-2-oxazoline) (pPrOx), and the \(T_g\) of pEtEtOx is about 30 °C higher than the \(T_g\) of pPrMeOx. It should be noted, that for the 2-substituted POx, polymers with higher DP values were investigated, suggesting that at similar DPs, the difference would be even more pronounced. In addition, poly(2-ethyl-4-methyl-2-oxazoline) (pEtMeOx) has a reported \(T_g\) of 75 -80 °C, which is basically identical with value found here for pEtEtOx. It is apparent that the presence of an additional methylene group at the polymer backbone branch, compared to the amide side chain, significantly impedes the macromolecular segment mobility. Also, increasing the length (from C1 to C2) of the side chain branching directly from the polymer backbone does not decrease the \(T_g\), which stands in contrast to the amide side chain. However, it would be interesting to see how the \(T_g\) evolves for longer side chains (≥ C3) branching from the main chain.
Table 1: Physicochemical characterization of synthesized homopolymers including the yield, number average molecular weight $M_n$, dispersity $D$ and the extrapolated onset temperature of major mass loss $T_d$.

| Polymer     | Yield [%] | $M_n$ a) | $M_n$ b) | $M_n$ c) | $D$ c) | $T_d$ d) | Solubility e) |
|-------------|-----------|----------|----------|----------|--------|---------|---------------|
| p^REtEtOx  | 69.3      | 2.7      | 2.9      | 1.3*     | 1.09*  | 362     | 7.8           |
| p^SEtEtOx  | 90.5      | 2.7      | 3.1      | 1.0*     | 1.16*  | 345     | 7.8           |
| p^RSEtEtOx | 57.7      | 2.8      | 2.9      | 1.2*     | 1.13*  | 361     | 9.4           |
| p^RPrMeOx  | 76.0      | 2.8      | 3.0      | 1.3*     | 1.10*  | 357     | 9.3           |
| p^SPrMeOx  | 86.6      | 2.8      | 3.0      | 1.3*     | 1.09*  | 344     | 8.9           |
| p^RSPrMeOx | 78.3      | 2.8      | 3.1      | 1.0*     | 1.11*  | 351     | 9.9           |

a) theoretical molar mass from $[M]_d/[I]_0$; b) as obtained by $^1$H-NMR (CDCl$_3$; 300 MHz) evaluated as mean of all relevant integral ratios; c) as obtained by GPC (*eluent: HFIP, calibrated with PEG standards; ‡ eluent: chloroform, calibrated with polystyrene); d) Extrapolated onset temperature of major mass loss (TGA); solubility in DI water at 4 °C in g L$^{-1}$.

Table 2: Glass transition temperatures $T_g$ of polymers reported here and compared with literature values for similar polymers.

| R= | R’= | DP | $T_g$ [°C] |
|----|-----|----|-------------|
| Me | H   | 60 | ≈80 $^{56}$ |
| Me | H   | 35 | 73 $^{54}$  |
| Me | Me  | 10 | 90 $^{40}$  |
| Et | H   | 60 | ≈60 $^{56}$ |
| Et | Me  | 25 | 75-80 $^{40}$ |
| Et | $^R$Et | 21 | 82 $^a$ |
| Et | $^S$Et | 23 | 80 $^a$ |
| Et | $^RS$Et | 21 | 79 $^a$ |
| Pr | H   | 60 | ≈40 $^{56}$ |
| Pr | $^R$Me | 22 | 55 $^a$ |
| Pr | $^S$Me | 22 | 55 $^a$ |
| Pr | $^RS$Me | 23 | 56 $^a$ |
| CH$_3$(CH$_2$)$_3$(Bu) | H | 60 | ≈25 $^{55}$ |
| CH$_3$(CH$_2$)$_4$(Pent) | H | 60 | ≈5 $^{56}$ |

a) this work, mean $T_g$ obtained from second and third heating curve (DSC)
Bloksma et al. investigated the enantiopure polymer p⁸BuEtOx, p⁵BuEtOx and racemic p⁸⁵BuEtOx via X-ray diffraction (XRD). The enantiopure polymers were found to be semicrystalline, while the racemic polymer was amorphous. However, pEtEtOx and pPrMeOx with short side-chain and main-chain branches have not been investigated using XRD before. Therefore, XRD measurements were performed on the pEtEtOx and pPrMeOx series at room temperature. All the homopolymers showed broad bands (Fig. S10), indicating that pEtEtOx and pPrMeOx series homopolymers are indeed amorphous, which is consistent with the absence of a T_m in the DSC measurement. The two broad bands in the XRD are in a similar position as was found previously for p⁸⁵BuEtOx. Generally speaking, poly(2-n-alkyl-2-oxazoline)s with 4 or more carbon atoms in side-chain are found to be semicrystalline, while the POx with 1-3 carbon atoms in side-chain are amorphous, albeit crystallization can has been reported for those when aqueous solutions are kept above their respective LCST. Our data suggests, that the additional carbon atoms branching off
of the main chain do not promote the formation of crystalline domains in case of POx with short side-chains, although we cannot rule out that under specific conditions, crystallization might occur.

CD spectroscopy is one of the few spectroscopic techniques that is applied to analyse the secondary structure of biopolymers and synthetic polymers. In 1992, Oh et al. carried out a molecular mechanics calculations for pMeMeOx with DP=20 and a corresponding tetramer. The calculated structures were defined by a left-handed helices containing 14 residues/3 turns with an identity period of 17.8 Å. Since, several kinds of chiral POx have been shown to form a secondary structures in solution, including p^R^EtEtOx. This flexible secondary structure is akin to the better-known polyproline type II helix. Accordingly, CD measurements of pEtEtOx and pPrMeOx series homopolymers were performed between 200 and 255 nm in methanol solution (0.1 g/L) at 25 °C. A positive Cotton effect (CE) with maximum values between 210 and 220 nm was observed for the S-homopolymers while a negative CE was observed for the R-homopolymers. This is in the wavelength range of n-π* transition of the amide chromophore (Fig. 1 c and d). The opposite CE and almost symmetrical CD spectra of R- and S- homopolymers indicate that they have a similar helical conformation as proposed by Oh et al., but with opposite handedness. The racemic polymer and the 1/1 (w/w) mixtures of two corresponding chiral homopolymers did not show CE. Besides, the CE maximum values of chiral pPrMeOx were markedly higher than that of chiral pEtEtOx, suggesting that the secondary structure formation of chiral pPrMeOx is more favorable than that of chiral pEtEtOx in methanol.

**Synthesis and characterization of ABA triblock copolymers**

After successful homopolymers preparation, we expanded our synthetic library to the ABA triblock copolymers comprising pMeOx as hydrophilic blocks A, and p^R^EtEtOx, p^S^EtEtOx, p^R^S^EtEtOx, p^R^PrMeOx, p^S^PrMeOx and p^R^S^PrMeOx as hydrophobic blocks B. As the hydrophilic blocks A are common in all polymers,
they were labelled according to their hydrophobic blocks A-p^6EtEtOx-A, A-p^5EtEtOx-A, A-p^6EtEtOx-A, A-
p^6PrMeOx-A, A-p^6PrMeOx-A and A-p^6PrMeOx-A, respectively. All triblock copolymers were characterized by \(^1\)H-NMR spectroscopy and GPC (Fig. S11-13 and Table 3). All the polymers exhibited excellent solubility in water and ethanol (solubility > 200 g/L).

Same as in the case of the homopolymers, the terminal Boc moiety was used for end group analysis by \(^1\)H-NMR spectroscopy (Fig. S11-S12). NMR spectra revealed a good synthetic control. The determined DP for block A are close to 35 and for block B are close to 20, which corresponds to the respective \([M]_0/([I]_0 values. After dialysis and lyophilisation, the triblock copolymers were analysed by GPC (Fig. S13). GPC elugrams of all copolymers exhibited a narrow molar mass distribution with a reasonably low dispersity (D <1.2). Thermally, the ABA triblock copolymers were slightly more stable than their respective homopolymers, the extrapolated onset temperature of major mass loss was at \(T_o >370\) °C (compared to \(T_d\) of the homopolymers =350 °C) (Fig. S14, Table 3). Again, a minor weight loss step corresponding the loss of the Boc group is observed at around 220 °C. In comparison to the homopolymers, this first mass loss step is less pronounced because of the lower relative weight percentage.

The \(T_g\) values for the various triblock copolymers of the A-pEtEtOx-A and A-pPrMeOx-A series are ≈76 °C and ≈71 °C, respectively. No melting transition was observed in the temperature range of -50 to 190 °C. As in the case of the homopolymers, no significant differences were observed for the different stereoisomers within A-pEtEtOx-A and A-pPrMeOx-A series. Compared to the corresponding homopolymers, the triblock copolymers containing pEtEtOx have a lower \(T_g\) while the triblock copolymers comprising pPrMeOx blocks have higher \(T_g\). Obviously, no microphase separation occurs in this system under the investigated conditions, leading to one \(T_g\) value in between the \(T_g\) values of pMeOx and pEtEtOx or pPrMeOx, respectively. Similar to the situation in the homopolymers, the polymers in the A-pEtEtOx-A and A-pPrMeOx-A series exhibited a higher \(T_g\) than their structural isomers A-pPrOzi-A (\(T_g\approx50\) °C) and A-
pBuOx-A (\(T_g\approx62\) °C).\(^{47}\)
Table 3. Physicochemical characterization of the triblock copolymers including the yield, molecular weight $M_n$, dispersity $Đ$, glass transition temperature $T_g$ and the extrapolated onset temperature of major mass loss $T_d$, critical micelle concentration CMC, and hydrodynamic diameter $D_h$.

| polymer       | Yield [%] | $M_n$ a) [kg mol$^{-1}$] | $M_n$ b) | $M_n$ c) | $Đ$ c) | $T_g$ d) [°C] | $T_d$ e) | CMC f) [M·10$^{-4}$] | CMC f) [g·L$^{-1}$] | $D_h$ g) [nm] |
|---------------|-----------|--------------------------|----------|----------|--------|-------------|---------|--------------------|------------------|-------------|
| A-p$^R$EtEtOx-A | 79.5      | 8.7                      | 8.7      | 4.4      | 1.13   | 76          | 377     | 3.2                | 2.7              | 4.9±1.0     |
| A-p$^S$EtEtOx-A | 62.5      | 8.7                      | 8.8      | 4.2      | 1.17   | 76          | 377     | 3.4                | 3.0              | 4.8±1.1     |
| A-p$^R$S$^R$EtEtOx-A | 76.5      | 8.7                      | 8.7      | 4.2      | 1.16   | 77          | 378     | 3.6                | 3.2              | 4.5±1.2     |
| A-p$^R$PrMeOx-A | 70.9      | 8.7                      | 8.8      | 4.2      | 1.15   | 71          | 374     | 4.1                | 3.6              | 4.6±1.2     |
| A-p$^R$S$^R$PrMeOx-A | 69.3      | 8.7                      | 9.1      | 4.1      | 1.15   | 71          | 373     | 1.2                | 1.1              | 4.7±1.2     |
| A-p$^R$S$^S$PrMeOx-A | 81.9      | 8.7                      | 8.8      | 4.2      | 1.18   | 73          | 380     | 2.2                | 1.9              | 4.9±1.4     |

a) According to $[M]_0/[I]_0$; b) Obtained by $^1$H-NMR (CDCl$_3$; 300 MHz) end-group analysis evaluated as mean of all relevant signals; c) Obtained by GPC (eluent: HFIP, calibrated with PEG standards); d) Mean $T_g$ obtained from second and third heating curve (DSC); e) Onset temperature of major mass loss (TGA); f) Obtained by pyrene assay, measured at 25 °C; g) Mean $D_h$ obtained by DLS from the size distribution by volume at 25 °C (polymer concentration 10 g/L in PBS, day 0).

The A-pEtEtOx-A and A-pPrMeOx-A series were also characterized with XRD. No crystalline peaks were observed in diffractograms (Fig. S15), confirming the amorphous character of all A-pEtEtOx-A and A-pPrMeOx-A series polymers. The chiroptical properties of the triblock copolymers were also investigated by CD in methanol (0.1 g/L) at 25 °C (Fig. 2 c and d). Clearly, the CD spectra of chiral triblock copolymers show a pronounced CE. The optically inactive pMeOx blocks do not prevent the secondary structures induced by the chiral block. In addition, the CE of chiral triblock copolymers in CD also retains in aqueous solution at 25 °C and 50 °C (Fig. S16). In contrast, as expected the racemic triblock copolymers and the 1/1 (w/w) mixtures of two corresponding chiral triblock copolymers did not show CE either in methanol or aqueous solution.
Figure 2. DSC heat flow of the third heating cycle of triblock copolymers (a) A-pEtEtOx-A and (b) A-pPrMeOx-A series. The samples were heated 3 times and cooled two times from -50 °C to 190 °C (10 °C/min). CD spectra of (c) A-pEtEtOx-A and (d) A-pPrMeOx-A series in methanol at 25 °C (polymer concentration was 0.1 g/L).

Formulation studies

PTX is a commonly applied chemotherapeutic agent in the treatment of various cancers, such as lung, ovarian, and breast cancers. CUR is a natural yellow orange dye derived from Curcuma longa. Because it reportedly has a plethora of biological effects such as affecting the expression of inflammatory cytokines, adhesion molecules, enzymes, the activity of several transcription factors and their signalling pathways et al., CUR is considered by many as a potential treatment in cancer, atherosclerosis, neurodegenerative disease, hepatic disorders, diabetes, psoriasis, autoimmune diseases and so on. However, its chemical instability in aqueous media also prompted a very critical discussion, labelling it as a pan assay interference compound (PAIN) or invalid metabolic panacea (IMP). While we acknowledge the importance of these issues, we also think that specifically these issues make CUR an interesting model compound for formulation studies.
Both compounds are poorly water soluble; the solubility of PTX is about 0.4 – 4 μg/mL\(^{68}\) while the solubility of CUR is in the range of 1 – 10 μg/mL\(^{69}\) depending on the polymorph. This poor solubility is one of the major problems for both compounds. Accordingly, both compounds have seen extensive efforts to improve their apparent solubility and thus, bioavailability.\(^{70-71}\) Among the plethora of drug delivery systems investigated for formulation of PTX and CUR, some POx and POzi based formulations stand out for their extraordinary high drug loading capacity and overall solubilisation for PTX and CUR.\(^{43,47,72-74}\) While CUR is achiral, PTX is chiral with multiple chiral centers but, to the best of our knowledge, does not have a known enantiomer. The formulations were prepared by the thin film hydration method. Briefly, ethanolic solutions of the polymer and drug were mixed in desired ratios, followed by ethanol removal. The resulting thin film was dissolved by adding water (Millipore). In the resulting solution, the polymer concentration was kept at 10 g/L, while increasing the drug concentration from 1 g/L to 10 g/L in each series. The actual drug concentration achieved in the aqueous phase was assessed using HPLC or UV spectroscopy using a microplate reader after removal of non-solubilized drug, if any, by centrifugation.

In addition to the triblock copolymers with chiral and racemic hydrophobic blocks, we also investigated 1/1 (w/w) mixtures of the chiral triblock copolymers for drug formulations. These mixtures are designated as \(^{69}\)A-pEtEtOx-A and \(^{69}\)A-pPrMeOx-A, respectively. The optical appearance of formulations is shown in Fig. S17. The centrifuged formulations of CUR-loaded various chiral/racemic A-pEtEtOx-A and \(^{69}\)A-pEtEtOx-A appeared homogenous and transparent up to 4 g/L CUR feed. In contrast, a minor precipitate was observed at 6 g/L CUR feed, but the supernatant was still transparent. Interestingly, when the CUR feed increased to 8 g/L and 10 g/L, the formulation separated to three layers: a small amount of precipitate at the bottom of the tube, an opaque layer in middle making up the majority of the sample and a thin transparent layer on top (Fig. S17 a). Also extended centrifugation for 5 min (rcf= 7788 g) did not sediment the opaque layer. This seems different from a gel-like agglomerate or coacervate which was reported in the formulation of A-poly(2-(3-ethylheptyl)-2-oxazoline)-A (A-pEtHepOx-A) and CUR.\(^{44}\) This behaviour
would be interesting to understand in more detail but this is outside the scope of the present contribution.

Here, the thin transparent layer was sampled for the measurement of CUR concentration. CUR-loaded A-pPrMeOx-A and M-A-pPrMeOx-A formulations were also transparent and homogenous at 1-4 g/L CUR feed. In contrast, at 6 g/L, 8 g/L and 10 g/L CUR feed, there was a significant amount of precipitate with a transparent supernatant observed, with the notable exception the formulation with A-pRSPrMeOx-A at 10 g/L CUR feed. The appearance of A-pRSPrMeOx-A/CUR=10/10 (g/L) (feeding ratio) was similar to the A-pEtEtOx-A formulations at 10 g/L CUR feed, and different to A-pRSPrMeOx-A, A-pPrMeOx-A and M-A-pPrMeOx-A formulations at 10 g/L CUR feed.

With increasing CUR or PTX feed, the solubilized drug amount increased until it reached the maximum LC (Fig. 3 a-e and Fig. S19). At the same CUR or PTX feed, A-pRSEtEtOx-A, A-pRSEtEtOx-A, A-pEtEtOx-A and physical mixtures M-A-pEtEtOx-A solubilized similar amount of CUR (Fig. 3 a) or PTX (Fig. 3 c), respectively. The maximum CUR and PTX LC was found to be 39-40 wt % (6.3-6.5 g/L) and 17-20 wt % (2.0-2.4 g/L), respectively (Fig. 3 e). Similarly, there is relatively little difference in solubilization for CUR (Fig. 3 b) and PTX (Fig. 3 d) in the pPrMeOx series. The maximum CUR LC is 28-29 wt % (3.6-4.0 g/L), while the maximum PTX LC is 32-34 wt % (4.6-5.0 g/L) (Fig. 3 e). Based on these results, it seems that the chirality of the hydrophobic block in ABA triblock copolymers has no obvious effect in solubilizing CUR and PTX. There is only one major difference observed for A-pRSPrMeOx-A at a CUR feed of 10 g/L (Fig. 3 b). This particular formulation was prepared two additional times (total 5 times) confirming the extraordinarily high solubilization in this particular case. As mentioned before, the visual appearance of A-pRSPrMeOx-A/CUR=10/10 (g/L) was more similar to the A-pEtEtOx-A formulations at 10 g/L CUR feed.

Although no significant differences were observed within each polymer series, the two different platforms based on pEtEtOx and pPrMeOx obviously show a different pattern of drug loading for CUR and PTX, respectively. The polymers containing pEtEtOx tend to load more CUR than PTX, while the polymers containing pPrMeOx solubilize more PTX than CUR. In previous work of Lübtow et al., it was shown that
even different positioning of a methylene group between the polymer side chain to the polymer main chain (comparing A-pPrOzi-A and A-pBuOx-A) can lead to a surprisingly specific drug loading for CUR and PTX, respectively. Here, the difference in positioning is between the amide side chain to the backbone branch, but also leads to a similarly specific drug loading of A-pEtEtOx-A and A-pPrMeOx-A for CUR and PTX, respectively. Specifically, comparing to literature with the different isomers, the order for maximum LC for CUR is A-pPrOzi-A (≈54 wt %) > A-pEtEtOx-A (≈40 wt %) > A-pPrMeOx-A (≈29 wt %) ≥ A-pBuOx-A (≈24 wt %). In contrast, for PTX, the order for maximum LC is A-pBuOx-A (≈48 %) > A-pPrMeOx-A (≈32 %) > A-pPrOzi-A (≈25 %) > A-pEtEtOx-A (≈18 %).

Figure 3. Solubilized CUR concentrations of CUR formulated (a) A-pEtEtOx-A and (b) A-pPrMeOx-A series in dependence of the CUR feed concentration. Solubilized PTX concentrations of PTX formulated (c) A-pEtEtOx-A and (d) A-pPrMeOx-A series in dependence of the CUR feed concentration. The maximum drug loading capacity of triblock copolymers and mixed polymers, for (e) CUR (orange) and PTX (green), and (f)
R-IBU (red), S-IBU (blue) and RS-IBU (purple). Polymer feed was 10 g/L. The data is given as means ± SD (n = 3, with the exception of A-pRSPrMeOx-A/CUR=10/10 (g/L) which is n = 5)

IBU is a non-steroidal anti-inflammatory drug (NSAID), possesses a single stereogenic carbon atom which gives rise to two enantiomers, S- and R-IBU. This drug is commercially available as racemate, even though S-IBU is more potent than R-IBU as inhibitor of cyclo-oxygenase I. IBU is practically insoluble in water (about 21 mg/L) and has therefore been used as a model drug to prepare formulations. Yang et al. encapsulated IBU into micelles of a copolymer brush poly(poly(vinylidene fluoride-co-vinylidene fluoride) (PVDF-co-VDF)-b-poly(ethylene glycol) methyl ether methacrylate) (P(PLAMA-co-MAA)-b-PPEGMA) and achieved a LC of 20.5 wt %. Lehto et al. absorbed IBU into siliceous mesoporous material TUD-1 and reported a LC of 19.6 wt % after washing. In addition, there are a few reports on IBU loaded POx-based hydrogels.

IBU is commercially available as its chiral and racemic forms and due to its low aqueous solubility, we chose it as a model drug to study solubilization in our novel chiral and achiral POx. The series of A-pEtEtOx-A, A-pPrMeOx-A triblock copolymers were used to solubilize R-IBU, S-IBU and RS-IBU.

The overview over maximum IBU LC in the resulting formulations is shown in Fig. 3f and Table S2. Interestingly, both A-pEtEtOx-A and A-pPrMeOx-A series have a similar and relatively high IBU LC exceeding 30 wt %. In terms of maximum IBU LC of A-pEtEtOx-A series formulations, the highest LC is A-pRtEtEtOx-A/R-IBU (37 wt %), the lowest LC is A-pRtEtEtOx-A/S-IBU (32 wt %). The formulation in the A-pPrMeOx-A series with highest LC is A-pRtPrMeOx-A/R-IBU (36 wt %), the lowest LC values are observed for A-pRtPrMeOx-A/RS-IBU and M-A-pPrMeOx-A/S-IBU (31 wt %). The maximum LC of the rest formulations vary within these limits without any notable regularity. In contrast to CUR or PTX, there is no clear and significant difference for IBU solubilization between the A-pEtEtOx-A and A-pPrMeOx-A series, even though overall the A-pEtEtOx-A series seems to have somewhat higher LC values. In order to view the LC
of different IBUs loaded in one of our triblock copolymers, the LC data was arranged in one coordinate system (Fig. S18 a-f). Up to 6 g/L IBU feed no significant differences were observed between R, S and RS-IBU for any triblock copolymer. At 8 g/L, the LC values for R-IBU and RS-IBU are similar for the same triblock copolymer, but the LC values of S-IBU trails behind for several polymers, especially in A-pRPrMeOx-A and A-pRPrMeOx-A. Coincidentally, the copolymer A-pRPrMeOx-A (Fig. S18 c) and A-pRPrMeOx-A (Fig. S18 f) have the maximum LC of R and RS-IBU at drug feed 8 g/L, while the others have their maximum LC at drug feed 6 g/L. It is rather unexpected and remains unexplained that the two racemic polymers show the highest deviations. In contrast at 10 g/L, again no clear or systematic difference as observed.

Figure 4. Solubilized (a) R-IBU, (b) S-IBU and (c) RS-IBU concentrations of IBU formulated A-pRPrMeOx-A (red), A-pRPrMeOx-A (blue), A-pRPrMeOx-A (purple) and M-A-pRPrMeOx-A (green) in dependence of the IBU feed concentration. Solubilized (d) R-IBU, (e) S-IBU and (f) RS-IBU concentrations of IBU formulated A-pRPrMeOx-A (light red), A-pRPrMeOx-A (light blue), A-pRPrMeOx-A (light purple) and M-A-pRPrMeOx-A (light green) in dependence of the IBU feed concentration. Polymer feed was 10 g/L. Data is given as means ± SD (n = 3). * p<0.05; ** p<0.01; *** p<0.001.
Besides, formulations for the different A-pEtEtOx-A (Fig. 4a-c) or A-pPrMeOx-A formulations (Fig. 4d-f) with different IBUs are compared. Again, up to 4 g/L drug feed, no real difference was observed. However, with increasing IBU feed to 6 g/L, the drug loading of A-pRS\textsuperscript{Et}EtOx-A was always lowest while drug loading for A-p\textsuperscript{Et}EtOx-A was highest in all three cases (Fig. 4a-c). The drug loading for MA-pEtEtOx-A and A-p\textsuperscript{Et}EtOx-A were basically the same and intermediate. At 8 g/L the picture looks much different, with the A-pRS\textsuperscript{Et}EtOx-A showing highest LC values for R and RS-IBU.

Also, for the A-pPrMeOx-A, no significant difference is observed at low feed. Different from the A-pEtEtOx-A series, A-pPrMeOx-A series did not show a regularity of drug loading between each other at 6 g/L IBU feed. Similar to the situation at 8 g/L for A-pEtEtOx-A, the A-pRS\textsuperscript{Pr}MeOx-A shows highest LC values for R and RS-IBU. To some degree, this indicates that the solubilization of IBU is affected more by the chirality of pEtEtOx hydrophobic block than of the pPrMeOx hydrophobic block. Besides, when the IBU feed increased to 8 g/L, the copolymers containing p\textsuperscript{Et}EtOx or p\textsuperscript{Pr}MeOx hydrophobic blocks showed only low drug loading. Apparently, the S-isomer “disadvantage” discussed before for S-IBU also appears in A-p\textsuperscript{Et}EtOx-A and A-p\textsuperscript{Pr}MeOx-A, but we cannot rationally explain this at the moment.

**DLS analysis**

The hydrodynamic diameter (D\textsubscript{h}) of unloaded, CUR loaded and IBU loaded polymer micelles were analysed by DLS. Interesting to note in this context, as determined by pyrene assay, both A-pEtEtOx-A and A-pPrMeOx-A series polymers exhibit a rather high CMC (1.1-3.6 g/L, i.e. 1.2-4.1 \times 10^{-4} M), compared to their isomers A-pBuOx-A (8 mg/L, 1× 10^{-6} M).\textsuperscript{45} The triblock copolymers were dissolved in PBS (polymer concentration 10 g/L) , filtered (0.45 µm) and measured by DLS at 25 °C. The DLS profiles of the micelles were bi- or multimodal with broad size distribution, indicating the formation of heterogeneous particle populations (Fig. S19). Comparing the size distribution by intensity, volume and number (Fig. S20), it
becomes clear that for $^\text{MA}$-pEtEtOx-A and $^\text{MA}$-pPrMeOx-A mainly small self-assemblies (presumably micelles) of $D_h \approx 5$ nm along with very few, much larger particles (apparent $D_h \leq 250$ nm) are present. However, from these simple DLS experiments, we only obtain apparent hydrodynamic sizes based on the assumption of spherical shape. It is however not unlikely that these self-assemblies indeed have a different morphology. The other triblock copolymer solutions had similar multimodal distribution as $^\text{MA}$-pEtEtOx-A and $^\text{MA}$-pPrMeOx-A. Important to note, while they had similar size distribution when freshly prepared, three sets of polymer solutions turned turbid after several days: $^\text{MA}$-pPrMeOx-A solutions (10 g/L) were turbid at day 7, while $^\text{A}$-p$^\text{R}$PrMeOx-A and $^\text{A}$-p$^\text{S}$PrMeOx-A solutions (10 g/L) showed a slight turbidity after one month. On the contrary, $^\text{A}$-p$^\text{R}$PrMeOx-A and all $^\text{A}$-pEtEtOx-A series remained transparent and retained the same particle size distribution after one month (Fig. S21 a). In order to investigate if the turbidity of $^\text{MA}$-pPrMeOx-A, $^\text{A}$-p$^\text{R}$PrMeOx-A and $^\text{A}$-p$^\text{S}$PrMeOx-A DLS samples was caused by crystallization, XRD measurements were performed again after freeze-drying the corresponding polymers from cloudy water (Millipore) suspension, but no signals suggesting crystalline domains were found (Fig. S22). Besides, the turbid DLS samples did not revert to transparent after storage at 4 °C for several days. This indicates that the turbidity of the triblock copolymer solutions is caused by self-assembly, and the thermoresponsive behavior of the hydrophobic block appears not to be sufficient to revert the self-assembly.

The micelles size of CUR loaded $^\text{A}$-pEtEtOx-A and $^\text{A}$-pPrMeOx-A series formulations were also analysed by DLS. All the formulations (polymer/CUR =10/2 g/L) self-assemble to form micelles with essentially the same size ($D_h \approx 25$ nm, PDI < 0.11; Table S3, Fig. 5a and b). Clearly, CUR loaded micelles exhibited a more uniform size distribution compared to the blank micelles. Dilution by 1/2 and 1/10 (v/v) samples (to 5 and 1 g/L) resulted in no change in size and distribution (data not shown). At the same time, all the CUR loaded formulations were quite stable. The size and size distribution of formulations were observed no significant change at day 7, except that of $^\text{MA}$-pPrMeOx-A/CUR (Fig. S23 a and b). Here, scattering intensity is
dominated by a narrow distribution of larger particles ($D_h \approx 200$ nm) after 7 days’ storage, even though in terms of volume or number the distribution at $D_h = 25$ nm remains dominant (Fig. S23 c). All the formulations remained optically clear after one month, including $^\text{MA}$-pPrMeOx-A/CUR (Fig. S21 b).

Similarly, S-IBU loaded A-pEtEtOx-A and A-pPrMeOx-A series formulations (polymer/S-IBU =10/2 (g/L)) were also studied with respect to their size distribution and dispersion stability. After thin-film hydration and filtration (0.45 µm), the maximum of the size distribution for both series of formulations lies between 10 and 20 nm (PDI<0.23, Table S4, Fig. 5c and d) with A-p$^\text{p}$EtEtOx-A, $^\text{MA}$-pEtEtOx-A and $^\text{MA}$-pPrMeOx-A showing somewhat broader distributions. The higher PDI values are primarily due to a minor population at larger sizes, in particular for A-p$^\text{p}$EtEtOx-A. Obviously, the formulations of the chiral compound S-IBU are quite similar but appear less uniform compared to the curcumin formulations, however, no clear trend between the different chiralities can be seen at day 0. After 7 days’ storage, the formulations remain transparent and their size and size distribution appear more uniform, except $^\text{MA}$-pPrMeOx-A/S-IBU (Fig. S24 a and b). This formulation turned turbid similar to the plain $^\text{MA}$-pPrMeOx-A/CUR formulation. Also, after one month, the other S-IBU loaded formulations remained transparent.
Figure 5. The size distribution by intensity of (a) A-pEtEtOx-A/CUR, (b) A-pPrMeOx-A/CUR, (c) A-pEtEtOx-A/S-IBU and (d) A-pPrMeOx-A/S-IBU formulation 10/2 (g/L) in PBS at day 0. The samples were measured at 25 °C after filtration using 0.45 µm PVDF syringe filter.

Long-term stability studies of the formulations.

In order to investigate the potential shelf-life of the formulations (A-pEtEtOx-A and A-pPrMeOx-A series), the freshly prepared CUR, PTX and IBU aqueous formulations were stored at ambient conditions containing the initial precipitate after thin-film hydration (if any). The samples were collected at day 0, 1, 8, 20, 30 and 60 with centrifugation before each collection to sediment precipitate (if any). The soluble drug remaining in the supernatant was then quantified.

In each polymer series (A-pEtEtOx-A and A-pPrMeOx-A), the stability of most formulations follows the certain pattern. Exemplarily, CUR loaded M-pEtEtOx-A and M-pPrMeOx-A formulations are shown in Fig. 6a, b. The formulations with a CUR feed of 1-8 g/L were relatively stable up to 24 h, less than 1 wt % loss in the LC was observed in both of M-pEtEtOx-A and M-pPrMeOx-A formulations. In the case of M-pEtEtOx-A formulations (Fig. 6a), up to 4 g/L CUR feed, no reduction in drug loading was observed even after 60 days. In contrast, at 6 g/L CUR feed and above, a gradual but moderate decrease in LC was observed. Overall, the A-pPrMeOx-A formulations series behaves different compared to A-pEtEtOx-A series, but within the series all formulations behave very similar, with the notable exception of A-pRSPrMeOx-A at 10 g/L CUR feed (vide infra). Up to a CUR feed of 6 g/L, all formulations are very stable for 60 days (Fig. 6b). However, at CUR feed of 8 and 10 g/L, the concentration of CUR found in the supernatant increased gradually and quite significantly. For instance, at 8 g/L CUR feed M-pPrMeOx-A formulation, a 10-fold increase in the drug loading (0.16 ± 0.04 g/L (day 0) to 1.66 ± 0.45 g/L (day 60)) was observed. Previously, the similar phenomenon was also reported for POx/POzi micelles with moderately hydrophobic block, such as in the CUR-loaded A-pBuOx-A formulation (CUR feed ≥5 g/L)\(^4\) and CUR-loaded
A-EtHepOx-A formulation (CUR feed 2-10 g/L). It is speculated that an initially formed drug/polymer coacervate redissolves over time, probably via an internal reorganization in the polymer drug self-assembly.\textsuperscript{44, 82} Such time-dependent change in the self-assembly of nanoformulations and its effect on the biodistribution and pharmacological performance has been recently reported by Kabanov.\textsuperscript{82}

The formulation of A-p\textsuperscript{R5}PrMeOx-A/CUR at 10 g/L CUR feed showed a very different behaviour, exhibiting a rather high drug loading at day 0 (23 wt %, Fig. 3 b), followed by an initial small decrease and a subsequent stronger increase leading to an LC = 32 wt % at day 60 (Fig. S25). At this point, we cannot explain this behaviour satisfactorily.

The long-term stability of PTX-loaded \textsuperscript{MA}pEtEtOx-A and \textsuperscript{MA}pPrMeOx-A formulations was also studied. The maximum PTX-loaded \textsuperscript{MA}pEtEtOx-A formulation (2 g/L PTX feed) was quite stable up to 24 h (Fig. 6c). Afterwards, the PTX LC dropped rapidly from 14 wt % (day 1) to 6 wt % (day 8). At PTX feed of 4 g/L, a minor loss (i.e., 2 wt %) in the LC was observed after 24 h, then the LC dramatically dropped to 1 wt % (day 8). At day 20, less than 3 wt % PTX was left in supernatant of all \textsuperscript{MA}pEtEtOx-A/PTX formulations. In comparison, the PTX-loaded \textsuperscript{MA}pPrMeOx-A formulations were relatively stable at 4 g/L PTX feed, as no LC reduction was observed at 24 h (Fig. 6d). While at the 6 g/L PTX feed, \textsuperscript{MA}pPrMeOx-A formulation showed 3 wt % LC loss at 24 h. However, the PTX-loaded \textsuperscript{MA}pPrMeOx-A formulation was also not stable for more days. After 8 days' storage, the LC of 4 g/L PTX feed decreased from 26 wt % (day 1) to 20 wt % (day 8), and the 6 g/L PTX feed decreased from 24 wt % (day 1) to 3 wt % (day 8). The PTX LC was less than 3 wt % in all \textsuperscript{MA}pPrMeOx-A/PTX formulations at day 20.
Figure 6. Long term stability of CUR-loaded (a) physical mixture $^M$A-pEtEtOx-A and (b) physical mixture $^M$A-pPrMeOx-A formulation in dependence of CUR feed concentration (polymer feed 10 g/L, CUR feed 1-10 g/L, 0-60 d). Long term stability of PTX-loaded (c) physical mixture $^M$A-pEtEtOx-A and (d) physical mixture $^M$A-pPrMeOx-A formulation in dependence of PTX feed concentration (polymer feed 10 g/L, PTX feed 1-10 g/L, 0-20 d). Data is given as means ± SD (n = 3).

Generally, the chirality of copolymers does not appear to affect the long-term stability of IBU loaded formulation, with the notable exception of few A-p$^{RS}$EtEtOx-A and A-p$^{RS}$PrMeOx-A formulations (vide infra). Exemplarily, the long-term stability of $^M$A-pEtEtOx-A/RS-IBU and $^M$A-pPrMeOx-A/RS-IBU formulations are used to discuss the general behaviour of A-pEtEtOx-A and A-PrMeOx-A series formulation, respectively (Fig. 7a, b). Up to 6 g/L RS-IBU feed, the LC of $^M$A-pEtEtOx-A/RS-IBU decreased slowly in 30 days. In contrast, at 8 and 10 g/L RS-IBU feed, the LC dropped significantly around day 20 (Fig. 7a). Comparing to A-pEtEtOx-A series, the IBU loaded A-pPrMeOx-A formulations appeared to be relatively more stable formulation. The LC of $^M$A-pPrMeOx-A/ RS-IBU decreased evenly and relatively slowly over 60 days, irrespective of the RS-IBU feed (Fig. 7b). At day 60, only 7 wt % LC loss was observed at 6 g/L RS-IBU feed.
As mentioned in drug loading part, A-pRS\textsuperscript{EtEtOx}-A and A-pRS\textsuperscript{PrMeOx}-A have the maximum LC of R- and RS-IBU at a drug feed of 8 g/L. Accordingly, the long-term stability of RS-IBU loaded A-pRS\textsuperscript{EtEtOx}-A and A-pRS\textsuperscript{PrMeOx}-A formulations are shown for these special cases (Fig. 7 c and d). Similar to M\textsuperscript{A}-pEtEtOx-A/RS-IBU, up to 6 g/L RS-IBU feed, the LC of A-pRS\textsuperscript{EtEtOx}-A/RS-IBU decreased gradually over 30 days period. However, at 8 g/L RS-IBU feed, formulations were not very stable, as within 24 hours, a 6 wt % LC loss was observed (Fig. 7 c). Finally, the LC of 8 g/L RS-IBU feed A-pRS\textsuperscript{EtEtOx}-A/RS-IBU formulation dropped to the same level of M\textsuperscript{A}-pEtEtOx-A/RS-IBU after 60 days. Similarly, the A-pRS\textsuperscript{PrMeOx}-A/RS-IBU formulations were relatively stable for 60 days at 6 g/L RS-IBU feed (Fig. 7 d). In contrast, at 8 g/L RS-IBU feed, stability was compromised.

![Figure 7. Long term stability of RS-IBU-loaded (a) A-pRS\textsuperscript{EtEtOx}-A, (b) physical mixture M\textsuperscript{A}-pEtEtOx-A, (c) A-pRS\textsuperscript{PrMeOx}-A and (d) physical mixture M\textsuperscript{A}-pPrMeOx-A formulation in dependence of RS-IBU feed concentration (polymer feed 10 g/L, RS-IBU feed 1-10 g/L). Data is given as means ± SD (n = 3).](image-url)
Conclusion

In summary, we have successfully synthesized a series of homopolymers of chiral and racemic poly(2,4-disubstituted-2-oxazoline)s, namely $p^8$EtEtOx, $p^8$EtOx, $p^8$EtEtOx, $p^8$PrMeOx, $p^8$PrMeOx and $p^8$PrMeOx via LCROP. Subsequently, novel ABA triblock copolymers were synthesized using these chiral and racemic hydrophobic $p^8$EtEtOx, $p^8$EtOx, $p^8$EtEtOx, $p^8$PrMeOx, $p^8$PrMeOx and $p^8$PrMeOx as block B and hydrophilic pMeOx as block A. The polymers were extensively characterized by $^1$H-NMR spectroscopy, GPC, TGA, DSC and CD-spectroscopy. Attributable to the steric hindrance caused by the methyl/ethyl group on the polymer backbone, both pEtEtOx and pPrMeOx series show less chain flexibility than pEtOx and pPrOx, respectively. Additionally, the results from TGA and DSC indicate that the thermal properties of $p^8$EtEtOx, $p^8$EtOx and $p^8$EtEtOx are similar, and that of $p^8$PrMeOx, $p^8$PrMeOx and $p^8$PrMeOx as well. The homopolymers $p^8$EtEtOx, $p^8$EtOx, $p^8$PrMeOx and $p^8$PrMeOx maintain their chirality, confirming that no racemization occurs during LCROP.

The homopolymers and triblock copolymers were studied by CD spectroscopy in solution. The CD spectra of all chiral polymers showed notable CE in methanol, and CD spectra of chiral triblock copolymers retained clear CE in water as well. It indicates that chiral polymers (including chiral block) form secondary structure in solution. Both A-pEtEtOx-A and A-pPrMeOx-A series copolymers formed rather heterogeneous self-assemblies in aqueous solution.

CUR and PTX loaded formulations were prepared through thin film method, to investigate the chirality influence of ABA triblock copolymers on the solubilization of common model drugs. However, the difference of LC between chiral, racemic triblock copolymers and the 1/1 (w/w) mixtures of two corresponding chiral copolymers is not significant for either CUR or PTX in most cases. However, A-pEtEtOx-A and A-pPrMeOx-A exhibit specific drug loading for CUR and PTX similar to the isomeric pair A-pBuOx-A and A-pPrOzi-A shown previously. It indicates that shifting a methylene group from the N-
substituted side chain to the backbone branch can also lead to specific drug loading. The chirality influence of ABA copolymers on loading chiral and racemic ibuprofen (R-IBU, S-IBU and RS-IBU) is not very significant up to 6 g/L drug feed. At higher drug feed (8 g/L and above), the chirality influence of polymers and IBU appears to affect the LC, as the LC of A-p^[EtEtOx]-A and A-p^[PrMeOx]-A are noticeably lower than their R- and RS- isomers. The origins for the observed, albeit small differences for chiral POx require further investigations, in particular the influence on interactions with biological systems will be a matter for future investigations.

**Supporting information**

NMR spectra; GPC elugrams; TGA; XRD patterns; CD spectra (in water solution); graph of drug solubility; data of maximum LC and LE; DLS; optical appearance of selected formulations; long term stability of A-p^[RSPrMeOx]-A/CUR (10/10 g/L).

**Acknowledgments**

This work was supported by the Deutsche Forschungsgemeinschaft, project # 398461692 (awarded to R.L.). Moreover, M. Y., C. H. and M. S. H. are grateful to the China Scholarship Council (CSC) and higher education commission of Pakistan-German academic exchange services (HEC-DAAD Pakistan), respectively, for a doctoral fellowship.

We also thank the Department for Functional Materials in Medicine and Dentistry (Julius-Maximilians-University of Würzburg) for instrument access and Prof. Paul D. Dalton and Prof. Dirk Kurth for valuable discussions.
Conflict of interest

The authors declare no conflicts of interest.

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