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Discrete and Stereospecific Oligomers Prepared by Sequential and Alternating Single Unit Monomer Insertion

Zixuan Huang,^{*a*} Benjamin B. Noble,^{*b*} Nathaniel Corrigan,^{*a*} Yingying Chu,^{*a*} Kotaro Satoh,^{*c*} Donald S. Thomas,^{*d*} Craig J. Hawker,^{*e*} Graeme Moad,^{*f*} Masami Kamigaito,^{*c*} Michelle L. Coote,^{*b*} Cyrille Boyer,^{*a*} and Jiangtao Xu^{*a*}*

^{*a*}Centre for Advanced Macromolecular Design and Australian Centre for NanoMedicine, School of Chemical Engineering, UNSW Sydney, NSW 2052, Australia

^bARC Centre of Excellence for Electromaterials Science, Research School of Chemistry, Australian National University, Canberra, ACT 2601, Australia

^cDepartment of Molecular and Macromolecular Chemistry, Graduate School of Engineering, Nagoya University, Furo-cho, Chikusa-ku, Nagoya 464-8603, Japan

^dNuclear Magnetic Resonance Facility, Mark Wainwright Analytical Centre, UNSW Sydney, NSW 2052, Australia

^eMaterials Research Laboratory and Departments of Materials, Chemistry and Biochemistry, University of California, Santa Barbara, California 93106, United States

^fCSIRO Manufacturing Bag 10, Clayton South, VIC 3169, Australia

ABSTRACT

Natural biopolymers, such as DNA and proteins, have uniform microstructures with defined molecular weight, precise monomer sequence and stereoregularity along the polymer main chain that affords them unique biological functions. To reproduce such structurally perfect polymers and understand the mechanism of specific functions through chemical approaches, researchers have proposed using synthetic polymers as an alternative due to their broad chemical diversity and relatively simple manipulation. Herein, we report a new methodology to prepare sequence-controlled and stereospecific oligomers using alternating radical chain growth and sequential photoinduced RAFT single unit monomer insertion (Photo-RAFT SUMI). Two families of cyclic monomers, the indenes and the *N*-substituted maleimides, can be alternatively inserted into RAFT agents, one unit at a time, allowing monomer sequence to be controlled through sequential and alternating monomer addition. Importantly, the stereochemistry of cyclic monomer insertion into the RAFT agents is found to be *trans*- selective along the main chains due to steric hindrance from the repeating monomer units. All investigated cyclic monomers provide such trans- selectivity, but analogous acyclic monomers give a mixed *cis*- and *trans*- insertion.

INTRODUCTION

Natural polymers, such as nucleic acids and proteins, are the basic structural components of life, which participate in the construction of genes, cells, and tissues. These bioorganic macromolecules possess precise monomer sequences and monomer chirality in their chemical structures which contributes to tuning interand intracellular interaction, defining functional interfaces, and storing chemical or genetic information.¹⁻⁴ The monomer sequence along these polymer chains defines the primary structure of macromolecules and the position of specific functionalities, while the uniform chirality of monomer units is responsible for the chemical configuration of molecular segments which will principally contribute to the helical formation and spatial arrangement of macromolecular assemblies. For most naturally-occurring polymers, helical structures are essential for exerting their sophisticated functions in nature.⁵⁻⁷

Reproducing the chemical structures of these functional biomacromolecules and mimicking their natural processes to understand the underlying mechanisms has been a great challenge for more than half a century. There has been limited success, for instance, scientists have duplicated these structurally perfect polymers from their natural monomers (nucleotides and amino acids) using chemical approaches, such as solid-phase peptide synthesis and oligonucleotide synthesis, to create new macromolecules for a range of biomedical and pharmacological applications.⁸⁻¹⁰ However, problems still remain in these systems with a limited monomer and reaction setup scope restricting the large scale production and innovation of diverse polymer structures. Additionally, such synthetically assembled polymers based on natural monomers have the issue of in vitro and in vivo biostability, which limits the biomedical applicability due to adverse enzymatic degradation.¹¹ Taking these concerns into account, researchers have proposed using synthetic polymers sequence and chirality as an alternative, due to their broad chemical diversity, relatively simple manipulation, and improved biostability.¹² Most importantly, such polymers have the potential to open new opportunities for the development of advanced materials for data storage, ¹³⁻¹⁶ catalysis¹⁷⁻¹⁸ and drug delivery.¹⁹⁻²¹

Various strategies have been proposed for regulating monomer sequence and chain length of synthetic polymers, including condensation chain polymerization,²² alternating radical polymerization,²³⁻²⁹ anionic polymerization,³⁰⁻³¹ ring-opening or cross metathesis polymerization,³²⁻³⁷ atom transfer radical addition (ATRA),^{24-25, 38-40} iterative exponential growth,⁴¹⁻⁴⁶ single unit monomer insertion (SUMI)^{12, 47-53} and various organic coupling, condensation or "click" addition reactions.^{19, 54-68} Separation techniques, which include automated flash chromatography⁶⁹⁻⁷¹ and preparative recycling size exclusion chromatography,^{51, 72} have also been developed to isolate discrete oligomers from synthetically derived oligomer mixtures. Recent developments have been summarized in comprehensive review articles.⁷³⁻⁷⁶

 Compared to sequence control, stereoselectivity in the polymerization of non-natural monomers is far more challenging and has received relatively little attention. Since the first discovery of synthetic isotactic polypropylene in 1955,⁷⁷ the development of polymerization techniques capable of controlling the stereochemistry of vinyl polymers has generated considerable interest in polymer science, as the stereoregularity significantly affects the physical and chemical properties of synthetic polymers.^{7, 78-80} To mimic the natural precision inherent in most biopolymers, both sequence- and stereo-control are vital, which requires monodisperse or uniform synthetic polymers. In recent elegant reports by Johnson's group,^{43-44, 81} exponential chain growth of chiral monomers (R and S configurations) can be incorporated into discrete polymers with different block sequences using "click" chemistry. Hawker and Qiao's work employed column chromatography to separate discrete and monodisperse isotactic and syndiotactic poly(methyl methacrylate) (PMMA) with defined monomer units, which were used to investigate the triple helix formation of such polymers.⁷¹

Recently, our group prepared discrete oligomers with monodisperse character and defined monomer sequences by SUMI into thiocarbonylthio compounds in high yield (up to 95% isolated yield).^{49, 52, 82} The synthesis of these discrete oligomers was based on the principle of selective photoactivation of SUMI adducts, in which the increased C-S bond dissociation energy of the SUMI adduct was used to suppress reactivation and multiple monomer insertions, which would reduce product purity and yield. As such, a major challenge is the selection of appropriate monomer/chain transfer agents, and catalytic systems/light wavelength combinations that meet these activation selectivity criteria. As a result, the monomer pool for subsequent SUMI is reduced, which increases the difficulty in preparing longer oligomers with control over the oligomer structures (**Figure 1a**).

In this article, these previous SUMI procedures have been significantly improved to allow for preparation of discrete oligomers with multiple functionalities and theoretically "infinite" chain growth through a sequential and alternating process (**Figure 1b**). This new protocol relies on alternating SUMI in which two families of monomers, namely, indene (Ind) and maleimide, are able to be alternatively inserted into a suitable RAFT agent and show little or no propensity for homopolymerization. The reaction kinetics for one pair of alternating reactions are much easier to study in this process, while repetition of the alternating process provides a facile method for increasing chain length and maintaining specificity. The selected monomer families can be adapted to carry diverse functionality, in particular, a plethora of maleimide monomers are available through substitution at the nitrogen position. There are a range of other monomer families available that could expand the pool of potential monomers.⁸³ As a result, the currently disclosed strategy allows the monomer sequence and position of functionality to be tuned along the polymer chain. Additionally, compared with other existing methods, our approach possesses the advantages of high ACS Paragon Plus Environment.

conversion and isolated yield for each step, simple purification, and easy functionalization, while reserving the merits of photo-RAFT such as the use of low energy sources, mild reaction conditions, functional group tolerance, which allows the potential to easily access large scale production.

Most significantly, these cyclic monomers insert in a stereospecific manner to form *trans*- linkages along the polymer backbone (**Figure 1b**), even though the SUMI process is a radical chain growth mechanism, which generally has less stereoselectivity. This is the first report of investigating stereochemistry of discrete polymers prepared by SUMI processes. This SUMI method provides the opportunity to engineer stereochemistry of every monomer through organic reactions during each cycle of insertion at a molecular level. The discrete oligomers formed herein can be expanded to access a library of functional building blocks with designed molecular chirality, enabling high order polymers to be prepared with sequence-specific positioning of functionalities and chirality, and thus to rationally design interaction capabilities by mimicking biological systems.



Figure 1. (a) Discrete oligomers synthesized in our previous work and (b) discrete and stereospecific oligomers synthesized in this work by photo-RAFT SUMI. X denotes functional substitution of the maleic or maleimide monomer unit.

In this article, we investigate the mechanism of sequential and alternating photo-RAFT SUMI and introduce the general criteria for screening monomers, RAFT agents and reaction conditions. Next, two model pentamers with alternating Ind/maleimide monomers with diverse functionalities, were synthesized to showcase the efficiency, versatility and robustness of this method. Interestingly, these cyclic monomers have specific stereochemistry when inserted into the polymer chain, which gives only *trans-* configurations due to steric hindrance, regardless of the reaction conditions, monomer families and functionalities. In the last part of this article, we discuss the stereospecificity of the cyclic monomers inserted into various RAFT agents with chiral and non-chiral R (leaving) groups. 1D/2D NMR, XRD and modeling simulations are employed to verify the stereoconfigurations with acyclic monomers being investigated for comparison.

RESULTS AND DISCUSSION

Scheme 1. Mechanism for photo-RAFT SUMI process and essential conditions for effective SUMI. k_d : decomposition rate constant of initial RAFT agent under light; k_{-d} : recombination rate constant of initial RAFT agent; k_{add} : addition rate constant of monomer to radical; k_p : propagation rate constant of monomer to its SUMI adduct radical; k_{tr} : chain transfer rate constant of SUMI adduct radical to initial RAFT agent; k_{-tr} : chain transfer rate constant of R group radical (R·) from initial RAFT agent to SUMI adduct macro-RAFT agent; k_{tr} ': chain transfer rate constant of SUMI adduct radical to SUMI adduct macro-RAFT decomposition rate constant of SUMI adduct macro-RAFT agent; k_{-d} ': recombination rate constant of SUMI adduct macro-RAFT agent. For clarity, the PET-RAFT technique, which involves electron transfer in the presence of a photoredox catalyst, was not depicted in the scheme.

The reaction mechanism for light activated RAFT SUMI is proposed in **Scheme 1.** It can be considered as a development of the pioneering work of Zard and coworkers who described photo-RAFT SUMI of less-activated vinyl monomers into xanthates.^{47, 84-85} RAFT SUMI has also been extensively explored by the CSIRO group and used as a process for synthesizing functional RAFT agents.^{50, 53, 86-87} The initial RAFT agent (ZCS₂R) is activated under light to generate an R group radical ($R \cdot$) and thiocarbonylthio anion or radical (ZCS₂·); this process can occur in the presence or absence of photocatalyst, depending on the employed RAFT agents and the irradiation wavelength. The R· radical can react with monomer to form an adduct radical, with the generation of the desired SUMI product occurring through coupling with thiocarbonylthio anion or radical species, or alternatively, through an addition/fragmentation chain transfer ACS Paragon Plus Environment

process with dormant RAFT species. The latter pathway is accompanied by the generation of another R· radical through fragmentation of the RAFT radical intermediate species.

From the standpoint of the proposed mechanism, the monomer and RAFT agent selection plays a critical role in determining the success of SUMI. The ideal monomers for achieving single unit monomer addition are those which cannot undergo radical homopolymerization (propagation rate constant, $k_{\rm p} \sim 0$) but give alternating copolymerization with other monomers under a range of conditions, and with high monomer addition rate constants (k_{add}). Therefore, α,β -disubstituted and conjugated vinyl monomers, such as N-substituted maleimide (Scheme 2a) and Ind (Scheme 2b), are ideal candidates due to their superior capability for alternating copolymerization with high reaction rates. Moreover, most maleimide and Ind monomers have also been reported to be non-homopolymerizable or low k_p through radical processes, which limits the formation of homopolymer side products.⁸³ In addition, maleimide monomers are electron accepting, while Ind monomers are electron donating, leading to formation of a charge transfer complex and allowing for alternating copolymerization to occur with high reaction rates,⁸⁸ although there has been debate of the mechanism for on alternating copolymerization in the literatures.⁸⁹⁻⁹⁰ Maleic anhydride (MAn, Scheme 2a) presents similar reaction behavior as maleimides, however, there is a high potential for hydrolysis and thus ring opening during the reaction. Meanwhile, although maleate, fumarate and trans- β -methylstyrene (*Trans*-MeSt, Scheme 2b) have low $k_p (\sim 0)$,⁸³ their reaction yield for SUMI into RAFT agents are very low (SI, see Section 2.3), suggesting addition rates of these monomers are not as high as maleimides and Ind, and thus they are not ideal monomers for the SUMI process used in this study.

Another critical component in RAFT mediated SUMI reaction protocols is the selection of the chain transfer agent. The high decomposition rate constant (k_d) of RAFT agent (ZCS₂R) was essential to ensure rapid generation of the initiating radical; the k_d value is directly related to the C-S bond dissociation energy of RAFT agent. Employing the PET-RAFT (photoinduced electron/energy transfer - RAFT) technique⁹¹⁻⁹² and a great variety of photoredox catalysts, most RAFT agents can be activated under light with high k_d . For instance, trithiocarbonate can be easily activated to generate radicals with high efficiency in the presence of zinc tetraphenylporphyrin (ZnTPP) under low energy red light.⁹³ Direct evidence of high k_d and k_{add} is the rapid consumption of RAFT agents (ZCS₂R) observed in these SUMI reactions.

Scheme 2. Various α,β -di-substituted vinyl monomers (electron acceptor (a) and electron donor monomers (b)) investigated in this study. All monomers are employed for the investigation of stereochemistry of SUMI adducts. Four monomers (PMI, HEMI, CEMI and Ind) are employed for the synthesis of discrete pentamers.

The reaction of the carbon centered R group radical from the starting RAFT agent with the disubstituted monomers can also be manipulated through RAFT agent selection. For instance, *n*-butyl benzyl trithiocarbonate (BBTC, **Scheme 3**) with an electron donating benzyl group (R group) can react with electron acceptor monomers such as malemides, maleates and fumarates, but not electron donor monomers such as Ind and *Trans*-MeSt. The trithiocarbonate (CPDTC, **Scheme 3**) with an electron accepting cyano-isopropyl group (R group) reacts in the reverse way. Meanwhile, high monomer/RAFT ratios (> 1/1) may be necessary in specific cases to increase the rate of insertion. The excess monomer does not lead to the formation of undesired higher oligomers (dimers, trimers, etc.) through multiple insertions as the monomer will not homopolymerize (*vide supra*).

The other essential condition for effective SUMI processes is a high recombination rate constant (k_{-d}, k_{-d}) [ZCS₂·] >> k_p [Monomer] ~ 0), or high chain transfer rate constant $(k_{tr} > k_{-tr}, k_{tr}$ [ZCS₂R] >> k_p [Monomer] ~ 0); this will ensure the effective formation of the SUMI adduct and prevent radical accumulation and undesirable termination reactions. Other factors such as a low decomposition rate constant (k_d) for the SUMI adduct are beneficial for the reaction efficiency and product purity but are not essential. Although it is difficult to determine the rate constants for these SUMI processes experimentally, qualitative estimates are helpful for understanding and optimizing such reactions.

Scheme 3. Various RAFT agents with diverse R (leaving) groups investigated in this study. For the full names of these compounds refer to Supporting Information. All RAFT agents are employed for the synthesis of SUMI adducts shown in Scheme 5. Only BBTC is employed for the synthesis of discrete pentamers. The asterisk (*) indicates the chiral carbon.

2. Sequence control of discrete oligomers using sequential and alternating SUMI

The initial RAFT agent for the synthesis of discrete oligomers was screened from the selection shown in **Scheme 3**. Dithiobenzoates, CPDB and CPADB, are poor choices for sequential SUMI processes as their mono-adducts cannot be reactivated ($k_d \sim 0$) by common photocatalysts used for PET-RAFT technique.⁴⁹ All trithiocarbonates are suitable for SUMI reactions, but BBTC was selected as a model RAFT agent because it has the simplest chemical structure (benzyl R group and *n*-butylthiol Z group), which simplifies NMR analysis compared to other RAFT agents. For example, the chiral R group in CDTPA leads to doubled diastereomers in the final product and complicates NMR analysis. BBTC was synthesized using an analogous procedure to BSTP, which has been reported in previous literature.^{91,94} Following silica column purification, nearly 60 g of BBTC RAFT agent was obtained; with the lower polarity of BBTC compared to BSTP reducing the difficulty of the column purification process. Furthermore, while other trithiocarbonates (CPDTC, CDTPA and DDMAT, **Scheme 3**) are commercially available, they are still relatively expensive.

Scheme 4. Synthetic chemistry of two model discrete pentamers, BBTC-M-I-M-I-M and BBTC-M-I-HM-I-CM.

Two model pentamers, BBTC-PMI-Ind-PMI (denoted as BBTC-M-I-M) with alternating *N*-phenylmaleimide (PMI) and Ind, and BBTC-PMI-Ind-HEMI-Ind-CEMI (denoted as BBTC-M-I-HM-I-CM) (Scheme 4) with diverse functionalities, were synthesized to evaluate the reaction conditions and kinetic efficiency. There are two key SUMI steps which are repeated during the synthesis of long oligomers: (1) the insertion of Ind to BBTC-M to generate BBTC-M-I, and (2) the insertion of maleimide into BBTC-M-I to generate BBTC-M-I-M. The reaction kinetics of both steps were comprehensively investigated using online monitoring techniques. The SUMI adducts were fully characterized by NMR, GPC and ESI-MS. The structural verification of long oligomers (> 5 monomer units) is challenging through NMR due to substantial signal overlap; as such, only GPC and ESI-MS were used for characterization of the pentamers.

2.1 Synthesis of BBTC-M mono-adduct in Step A

We started the SUMI process by activating BBTC through the PET-RAFT technique in the presence of PMI. Following the design principles outlined in Section 1, a suitable electron donor-acceptor pair between the donating BBTC R group (benzyl) and maleimide acceptor was formed, leading to effective SUMI of PMI into BBTC. The reaction was performed in the presence of ZnTPP as photocatalyst under red light,

using the molar ratio of [PMI]:[BBTC]:[ZnTPP] =1/1/0.005. The kinetic profile of this reaction was determined by monitoring the conversion of BBTC against time using online ¹H NMR spectroscopy using deuterated DMSO as reaction solvent (**Figure 2a**). The reaction proceeded quickly within the first 2 h and slowed down to reach 65% after 10 h of irradiation. With the consumption of BBTC, the concentration of BBTC-M increased and reached an equilibrium stage. The conversion of RAFT agent could not reach 100% since the benzyl radical in BBTC is a poorer leaving group than the maleimidyl radical of the newly formed SUMI adduct (BBTC-M). The light activation and addition/fragmentation process of BBTC-M competes with that of BBTC, i.e., k_d [BBTC] competes with k_d [BBTC-M], while k_{tr} [BBTC] competes with k_{dr} [BBTC-M]. The decomposition rate of BBTC-M is higher than that of BBTC ($k_d > k_d$) due to the lower C-S bond dissociation energy of BBTC-M compared with BBTC. The chain transfer rate k_{tr} is higher than k_{tr} in this SUMI, which suggests fragmentation to produce SUMI product of BBTC-M (SI, Scheme S1) is unfavorable.

The stacked ¹H NMR spectra (SI, **Figure S3**) at different time points (0, 1, and 9 h) clearly showed the SUMI process proceeded smoothly without obvious side reactions, such as RAFT agent degradation by irreversible termination and dimer formation (BBTC-M-M). The crude mono-adduct was purified by column chromatography to give an isolated yield of 56%. Neat ¹H and ¹³C NMR spectra (SI, **Figures S4** and **S5**) clearly confirmed the chemical structure and high purity of the mono-adduct. ESI-MS (**Figure 3c** and SI, **Figure S6**) further confirmed the isolated product is BBTC-M. Surprisingly, a higher [PMI]/[BBTC] ratio such as 5/1 could not improve the yield and adversely increased the risk of multiple PMI insertion (SI, **Figures S7** and **S8**).

Figure 2. Kinetic study of sequential PET-RAFT SUMI processes from BBTC to BBTC-M-I-M monitored by online ¹H NMR analysis (DMSO- d_6 , 400 MHz). Conversions of RAFT agents *versus* irradiation time: BBTC (**a**), BBTC-M (**b**) and BBTC-M-I (**c**); Stacked ¹H NMR spectra indicated the SUMI processes: BBTC-M to BBTC-M-I (**d**) and BBTC-M-I to BBTC-M-I-M (**e**). The scheme on the right depicts the corresponding chemical structures of the growing SUMI products examined in this kinetic study.

2.2 Synthesis of BBTC-M-I dimer in Step B

The insertion of Ind to BBTC-M to prepare BBTC-M-I followed a similar process to Step A, with identical reaction conditions employed, except for a higher monomer/RAFT ratio ([Ind]:[BBTC-M]:[ZnTPP] = 5/1/0.005). An increased monomer/RAFT ratio was used due to the relatively low addition rate of Ind to maleimidyl radicals, compared to that of PMI to benzyl radicals. In this step, the proposed reaction pathway is similar with Step A ($k_d' > k_d$ and $k_{-tr} > k_{tr}$), as shown in the SI, Scheme S2. The slightly higher stability of

Ind radicals compared to maleimidyl radicals causes $k_d' > k_d$, and also leads to favorable fragmentation of the intermediate to generate the Ind radical and not the SUMI product (BBTC-M-I). Therefore, the high monomer/RAFT ratio improves the RAFT agent conversion and reaction yield by pushing the reaction towards the desired SUMI product (BBTC-M-I), although it also increases the probability of dimer formation.

Figure 3. GPC traces and ESI-MS spectra for all SUMI adducts. GPC traces for the successive synthesized oligomers, from BBTC to BBTC-M-I-M-I-M (a) and from BBTC to BBTC-M-I-HM-I-CM (b), and their corresponding ESI-MS spectra (c) and (d). The full spectra for these SUMI adducts can be found in Supporting Information.

The reaction kinetics showed the conversion of macro-RAFT agent, BBTC-M, reached 100% after 20 h of irradiation (**Figure 2b**). ¹H NMR showed the reaction proceeded smoothly with the increased doublet peak signals at δ 5.42 ppm and δ 6.05 ppm assigned to the two diastereomers of the terminal –CH– proton (proton 3) in BBTC-M-I (**Figure 2d**). The peak signals at δ 5.28 ppm and δ 3.65 ppm assigned to the protons *1* and *2* dropped to 10 % as the reaction time increased from 0 h to 12 h. After 20 h of irradiation, a signal peak (*m*/*z* = 684.20) in ESI-MS was observed, which suggested the presence of double Ind insertion (BBTC-M-I-I) ionized with Na⁺ (SI, **Figure S9b**). The double insertion is attributed to the low chain transfer rate (*k*_{-tr} > *k*_{tr}), causing an increased probability of multiple monomer insertion after long irradiation times in

 the presence of excess monomer. Fortunately, a shorter reaction time (< 15 h) effectively suppressed the formation of this Ind double insertion adduct (ESI-MS, SI, **Figure S9a**). As a result, the RAFT conversion in Step B had to be limited to 85~95%. However, it is not necessary to push macro-RAFT conversion to 100%; the unreacted BBTC-M will be unable to participate in the subsequent maleimide insertion reaction in Step C, as the low maleimide homopolymerization rate ($k_p \sim 0$) suppresses the formation of the dimeric BBTC-M-M adduct. More importantly, this unreacted BBTC-M can be easily removed by column chromatography to afford pure BBTC-M-I-M trimer in Step C (*vide infra*). The ¹H NMR spectrum for the crude product after excess monomer (Ind) removal clearly demonstrated the presence of a mixture of BBTC-M-I and BBTC-M (SI, **Figure S10**).

2.3 Synthesis of BBTC-M-I-M trimer in Step C

This step involves the highly efficient insertion of PMI into BBTC-M-I dimer to form the BBTC-M-I-M trimer, which differentiates the insertion of PMI into BBTC in Step A due to a different R group (benzyl radical *versus* Ind radical). Ind radicals are better leaving groups than both benzyl and maleimidyl radicals, and thus the insertion of PMI into BBTC-M-I reached high conversions (95% isolated yield). The high chain transfer rate ($k_{tr} < k_{tr}$) made the intermediate species preferentially generate the trimer SUMI adduct rather than the RAFT dimer and trimer adduct radical (SI, Scheme S3). The high decomposition rate ($k_d' < k_d$) also allowed for fast conversion of the RAFT agent (BBTC-M-I), which was reflected in the kinetic results (Figure 2c). Nearly full conversion of the initial RAFT agent (BBTC-M-I) was observed after 15 h of irradiation, with a molar ratio of [PMI]/[BBTC-M-I] = 1/1. The ¹H NMR spectra (Figure 2e) at different time points of the reaction (0, 1, and 9 h) clearly indicated the gradual consumption of BBTC-M-I dimer by the decrease of the peak signals at δ 5.42 and δ 6.05 ppm assigned to the proton 3 of terminal –CH- in the dimer BBTC-M-I, and generation of BBTC-M-I-M trimer by the increase of peak signals at δ 5.44, δ 5.38 and δ 5.0 ppm, assigned to the proton 4 of terminal –CH– in the trimer BBTC-M-I-M. As anticipated, the unreacted BBTC-M mono-adduct (the peak signals for protons 1 and 2 in BBTC-M at 0 h) remained intact during the reaction, which demonstrates that the presence of this species (from the previous insertion step) does not affect the production of the target trimer. The unreacted BBTC-M can simply be removed by conventional column chromatography. The $R_{\rm f}$ values in silica gel thin layer chromatography (TLC) for the impurity, BBTC-M, and the product, BBTC-M-I-M, are very different ($R_{\rm f} = 0.7$ for BBTC-M and 0.3 for BBTC-M-I-M using 5/1 (v/v) hexane/ethyl acetate as the eluent), due to the increased polarity and molecular mass of BBTC-M-I-M. The ¹H NMR spectra also indicated that other side reactions (multiple maleimide insertion or RAFT agent degradation) were negligible.

After purification by column chromatography, the structure of the isolated BBTC-M-I-M trimer was ACS Paragon Plus Environment

verified by ¹H and ¹³C NMR, HSQC, and HMBC (SI, Figure S11, S12, S13 and S14). The ¹H NMR spectrum is complicated due to the presence of several diastereomers for this compound. Furthermore, the ¹H NMR spectrum measured in CDCl₃ (SI, Figure S11) was different to that in DMSO- d_6 (Figure 2e). Particularly, the terminal -CH- proton shows different chemical shifts and profiles, which gives three peaks from δ 4.9~5.5 ppm in DMSO- d_6 and two peaks from δ 4.6~4.9 ppm in CDCl₃. Moreover, the peaks from δ 4.6~4.9 ppm in CDCl₃ are not all signals of terminal -CH- proton in these molecules, which was demonstrated by the low integration values of the peaks. The other part of the signal for this proton is overlapped in the area from δ 3.95~4.15 ppm, which was demonstrated by the analysis of ¹³C NMR (SI, Figure S12), high resolution ¹H-¹³C HSOC (SI, Figure S13), and ¹H-¹³C HMBC (SI, Figure S14). HSOC confirmed the rough assignment of every proton in the molecule and HMBC confirmed the peak location of the terminal -CH- proton. To further confirm the structure of the synthetic compound, GPC and ESI-MS analysis was performed. The GPC trace for this product showed a unimodal and symmetric peak, and a clear shift in molecular weight compared to BBTC-M (Figure 3a). ESI-MS confirmed the molar mass of the product to be 741.19, which matched well with the target compound ionized with Na⁺ (Figure 3c and SI. Figure S15). A more detailed discussion for the structural verification of this compound can be found in Supporting Information.

2.4 Synthesis of BBTC-M-I-M-I tetramer in Step D and BBTC-M-I-M pentamer in Step E

The tetramer and pentamer were successfully synthesized and isolated using the same procedures as those for dimer and trimer synthesis under identical conditions in high yield (85% isolated yields for two reaction steps). A conventional silica column was employed to remove excess monomer and trace amounts of impurities, however, difficulties in purification arose due to reduced discrepancy in both molecular weights and polarity between two neighboring SUMI products. The final purified products were confirmed by ¹H NMR (SI, **Figure S16** for BBTC-M-I-M-I tetramer and **Figure S18** for BBTC-M-I-M pentamer), GPC (**Figure 3a**) and ESI-MS (**Figure 3c** and SI, **Figure S17** for BBTC-M-I-M-I tetramer and **Figure S19** for BBTC-M-I-M pentamer).

According to the reaction conditions and efficiency, the trithiocarbonate end group in each step can always be activated easily in the co-monomer system. The SUMI process described in this work can therefore be repeated numerous times with the biggest challenge being separation of the product adducts from the reaction mixture. However, the use of a support resin similar to those used in solid-phase peptide synthesis could potentially increase the commercial viability of this SUMI process.

To demonstrate the end-group fidelity and the capability of the pentamer as a functional building block

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for constructing complex polymer architectures, the synthesized BBTC-M-I-M pentamer was employed as macro-RAFT agent for the polymerization of methyl acrylate (MA) to prepare diblock copolymer ((BBTC-M-I-M)-b-poly(MA)) using the PET-RAFT technique with ZnTPP as the photocatalyst. The $\ln([M]_0/[M]_1)$ derived from monomer conversion linearly increased with the exposure time to light (Figure 4a), indicating a constant radical concentration in the system. The kinetic plots of M_n versus monomer conversion, $\ln([M]_0/[M]_1)$ versus exposure time and molecular weight distribution were investigated. The linear increase of $M_{n,GPC}$ versus monomer conversion (Figure 4b) demonstrated that this macro-RAFT agent was able to mediate the controlled radical polymerization of MA. The experimental molecular weights at different monomer conversions were slightly higher than the theoretical values calibrated by polystyrene standards, which was attributed to the molecular weight of macro-RAFT agents. Regardless, the uniform and symmetric molecular weight distributions showed no signs of tailing and the clear and complete shift in molecular weight indicated the remarkable living character and end-group fidelity of macro-RAFT agents (Figure 4c). Moreover, the dispersity $(M_w/M_n$ in Figure 4b) was as low as 1.08 during polymerization, illustrating that the pentamer is an excellent chain transfer agent. Additionally, "ON"/"OFF" experiments revealed that the polymerization process can be started and stopped by switching the light "ON" and "OFF" (Figure 4d); similar temporal control has been demonstrated previously in other PET-RAFT polymerization systems.^{49, 93, 95}

Figure 4. Online Fourier transform near-infrared (FTNIR) measurement for kinetic study of PET-RAFT polymerization of methyl acrylate (MA), using pentamer BBTC-M-I-M-I-M as macro-RAFT agent and

ZnTPP as photocatalyst, with the molar ratio of [MA]:[macro-RAFT]:[ZnTPP] = 300:1:0.01 (~33 ppm catalyst with respect to monomer concentration): (a) $\ln([M]_0/[M]_t)$ versus exposure time; (b) M_n and M_w/M_n versus conversion; (c) molecular weight distributions at different time points, and (d) monomer conversion versus time in the presence (ON) or in the absence (OFF) of light, red arrows represent the four sampling time points for GPC analysis.

2.5 BBTC-M-I-HM trimer and BBTC-M-I-HM-I-CM pentamer

A range of other functional trimers and pentamers were prepared by following the same synthetic protocol as those for BBTC-M-I-M trimer and BBTC-M-I-M pentamer, except that the maleimide monomers were changed to HEMI and CEMI (Scheme 2) for the third and fifth monomer insertion instead of PMI (Scheme 4). The purification procedures were also similar, using conventional silica gel column and gradient elution, except that the composition of solvents was a little different due to increased polarity of the new SUMI compounds. The product for each step was fully characterized by ¹H NMR (SI, Figure S20, S22 and S23) with GPC (Figure 3b) and ESI-MS (Figure 3d and SI, Figure S21 and S24) of the trimer and pentamer providing solid evidence of successful production in high purity. Significantly, the isolated yields were 95% and 86% for the trimer and pentamer, respectively.

3. Stereospecificity of synthetic discrete oligomers

The properties of natural and synthetic polymers are determined in large part by their geometrical and stereochemical structure, i.e., degree of regularity of the substituents along macromolecular chain. In general, high regularity imparts strength, high glass transition temperatures and hardness, together with restricted solubility in solvents. Symmetry and regularity in macromolecular structures also enhance micro-crystallinity, which is a key property for many applications of polymeric materials. Designing and tuning micro-crystallinity of polymers at the molecular level is therefore a fundamental challenge for synthetic chemists.

Scheme 5. (a) Photo-RAFT SUMI of cyclic monomers into various RAFT agents generating SUMI adducts with all-trans configuration; (b) Photo-RAFT SUMI of acyclic monomers to RAFT agent (BBTC) generating SUMI adducts with mixed *cis*- and *trans*- configurations.

Having demonstrated the ability to control the sequence and chain length of synthetic oligomers, our attention was turned to stereochemistry. As discussed previously in the characterization of synthetic oligomers (Section 2), full structural determination of compounds with more than three monomer units is difficult by NMR spectroscopy. Conveniently, it is not necessary to determine the stereoconfiguration for the full length of final oligomer, as the oligomer is composed of repeating alternative monomer units of N-substituted maleimide and Ind; if we can determine the stereoregularity of one repeating unit, it will be the same for all repeating units with increasing chain length. Initially, the stereochemistry of maleimide and

Ind independently inserted into various RAFT agents with chiral and non-chiral R groups was studied (**Scheme 5**). Maleic anhydride and three acyclic monomers, DEM, DMF and *Trans*-MeSt, were also investigated for comparison.

3.1 Stereochemistry of N-substituted maleimide unit in discrete oligomers

In the chain growth polymerization of vinyl monomers, each propagation step results in the formation of either one (α -substituted and α, α -disubstituted monomers) or two (α, β -disubstituted) stereogenic centers with four different substituents for each. For α, β -disubstituted monomers, both backbone atoms are stereogenic with each possessing either *R* or *S* configuration. For the α, β -disubstituted cyclic vinyl monomers, four potential backbone configurations (*R*, *R*), (*R*, *S*), (*S*, *R*) and (*S*, *S*) exist; a pair of *cis*enantiomers and a second pair of *trans*- enantiomers. With increasing degree of polymerization, the number of possible diastereomers rises exponentially and higher order stereochemical classification (e.g. *threo* and *erytheo*) becomes necessary.

Figure 5. ¹H NMR spectra (400 MHz, CDCl₃) (δ 2.5~5.5 ppm) to demonstrate the *trans*- configuration of maleimide unit in purified (a) BBTC-M and (b) CPDTC-I-M.

During the SUMI of PMI into trithiocarbonate of BBTC, the fragment radicals could theoretically add to the maleimide double bond in both *cis-* and *trans-* fashions. Therefore, the mono-adduct of BBTC-M gives two diastereomers of *cis-* and *trans-* configurations in five-member ring units with their corresponding enantiomers of mirror-image molecules (**Scheme 5a**). The *cis-* and *trans-* diastereomers have different stereochemical structures and molecular environments for the corresponding protons. Therefore, ¹H NMR analysis, particularly the chemical shifts and coupling constants (*J* values) of specific protons, allows *cis-* and *trans-* configurations to be distinguished. However, the ¹H NMR spectrum (**Figure 5a** and SI, **Figure S4**) of BBTC-M showed only one set of diastereotopic signals, implying formation of either a *cis-* or *trans-* configuration exclusively. The signal at δ 4.6 ppm was assigned to the proton of –CH– (5) giving a doublet, ACS Paragon Plus Environment

 which is coupled with the proton of -CH- (6) and showed the same coupling constant of $J_{H5,H6} = 7.2$ Hz. The other configuration (either *cis-* or *trans-*) is supposed to show another doublet signal for the proton of -CH- (5), but was not observed.

The stereochemistry of N-substituted maleimides has interested researchers since 1960s in attempting to use homo- and co-polymerization to prepare polymers with helix-sense chirality.96-97 The anionic homopolymerization of maleimides provided all-trans configuration at low (-78 °C) to room temperature with anionic initiators.⁹⁸ Radical homopolymerization of maleimides at elevated temperature (100 °C) with the assistance of chiral solvent or catalyst also showed trans- stereo-configuration.⁹⁹ The radical copolymerization of maleimides and styrene or propylene provided a helical structure, with the maleimide monomers most likely possessing an all trans- configuration.⁹⁶ In contrast, the electron-donor-acceptor interaction of styrene and cyclic citraconic anhydride (a maleic derivative) resulted in a mixture of *cis*- and trans- configurations.¹⁰⁰ Additionally, the single monomer radical addition of PMI to a xanthate gave a trans- configuration due to the steric hindrance. 47-48, 85 However, the chemical structure of the products was not comprehensively confirmed. Due to the challenge of structural characterization of complex polymers, the stereochemical outcome of maleimide polymerization is thus arguable. There are no effective tools for polymers with large numbers of repeating monomer units, and as noted above previous experiments have delivered conflicting results. In contrast, SUMI provides an excellent tool to synthesize discrete oligomers with exact molecular weights, making it possible to confirm the stereoconfiguration and reaction mechanism, using widely used analytical equipment (NMR and XRD, etc). Most importantly, this tool can be applied to any monomers on the condition that they can be polymerized by iterative SUMI. In this study, the SUMI adducts of α . β -disubstituted vinvl monomers were synthesized and their stereochemistry was investigated for the first time. Particularly, this approach is able to assess the stereospecific configuration of our synthetic discrete oligomers in Section 2.

To elucidate the *cis-* or *trans-* configuration of the product of SUMI of PMI into trithiocarbonates, we performed single crystal X-ray diffraction (XRD) analysis and computational simulations. Single crystal XRD is a powerful technique that provides accurate structural information on small organic compounds, including the location of atoms, bond lengths, bond angles and stereoconfiguration. The sample for XRD requires a crystalline compound, therefore, crystallization of mono-adduct compound, BSTP-M (**Scheme 5a**, ¹H NMR shown in SI, **Figure S25**) was carefully performed; BSTP-M was selected rather than BBTC-M due to the carboxylic acid moiety in BSTP, which facilitates more efficient crystallization. Fine needle-like crystals were obtained, suggesting that BSTP-M has good crystallinity and thus structural stereoregularity, as crystallinity is often regarded as external manifestation of a regular arrangement at the molecular level. XRD analysis of the obtained crystal perfectly matched with the expected molecular structure of BSTP-M ACS Paragon Plus Environment

and unambiguously demonstrates that PMI insertion results in a *trans*- configuration (Figure 6a).

In agreement with the XRD results, computational modelling indicates that the *trans*-BSTP-M diastereomer is around 12.4 kJ/mol more stable than the corresponding *cis*-isomer (**Figure 6b**). This Gibbs-Free energy corresponds to a thermodynamic selectivity of over 99% for the *trans*- diastereomer (at room temperature), indicating that the PMI insertion into BSTP is highly stereospecific.

Figure 6. Single crystal XRD of BSTP-M (a) and modeling results showing the most stable conformations of the *trans-* and *cis-* BSTP-M diastereomers as well as their respective solution-phase Gibbs-Free Energies (b).

To illustrate the broad applicability of this strategy, *N*-substituted maleimide monomer, HM, and MAn were also used to prepare the corresponding SUMI adducts BBTC-HM and BBTC-MAn (Scheme 5a). The ¹H NMR spectroscopy (SI, Figure S30 and S31) revealed the same stereospecific preference for *trans*-configurations of HM and MAn units in the adducts, suggesting cyclic maleimide and maleic monomers have the same stereogenic preference regardless of their substitutions.

The RAFT agents of both BBTC and BSTP have non-chiral R groups. To investigate RAFT agents with chiral R groups, the insertion of PMI into CPDTC-I was performed and the resulting product analyzed by NMR (Scheme 5a). Careful analysis of the NMR results can be used to elucidate the stereochemical configuration of the PMI unit in the adduct (Figure 5b and SI, Figure S34), indicating the same *trans*-configuration of the PMI unit in the adduct. There are two series (a and b) of proton signals which are assigned to the two diastereomers of CPDTC-I-M, each containing a pair of enantiomers as verified by 2D NMR analysis (SI, Figure S35-S37). If both *trans*- or *cis*- configurations for PMI exist, there should be four diastereomers which are attributed to each configuration (*trans*- or *cis*-) with two different connection types for the Ind R group (*threo* and *erytheo*). Thus, if all four diastereomers are generated, four series of proton

signals would be anticipated. However, only two series signals were observed in the ¹H NMR spectrum, which are assigned to two diastereomers of the *trans*- configurations shown in **Figure 5b**.

It is worth noting that the two chemical shifts of the same protons in the adduct, such as -CH-S-(5), were were highly sensitive to the adduct stereochemistry. The chemical shift for one diastereoisomer is δ 6.1 ppm and δ 5.2 ppm for the other, which is attributed to the aromatic ring current effect (SI, **Scheme S4**). This proton has a very different chemical environment in each of the two *trans*- diastereomers; in one isomer (*threo*) it is deshielded by the phenyl ring of the adjacent Ind unit and thus leads to higher chemical shift.

3.2 Stereochemistry of Ind unit in discrete oligomers

Figure 7. ¹H NMR spectra (400 MHz, CDCl₃) (δ 2.5~6.4 ppm) to demonstrate the *trans*- configuration of Ind in purified (a) CPDB-I and (b) CPADB-I.

As one of the alternating monomer units in these synthetic and discrete oligomers, Ind is different from maleimide due to its asymmetrical structure. Interestingly, there are not many previous reports for its polymerization stereochemistry.¹⁰¹⁻¹⁰³ Five RAFT agents with electron accepting R groups (**Scheme 3**) were therefore investigated for SUMI of Ind; three with non-chiral R groups and two with chiral R groups.

The characterization of these SUMI adducts was carried out by 1D and 2D NMR. For instance, the ¹H NMR of purified CPDB-I (**Figure 7a** and full spectrum shown in SI, **Figure S28**) was obtained to confirm the *trans*- configuration of the inserted cyclic Ind monomer with similar proton signals as that for BBTC-M being observed (**Figure 5a** and SI, **Figure S4**). Computational modelling also indicated that the *trans*-CPDB-I diastereomer is around 12.5 kJ/mol more stable than the corresponding *cis*-isomer (**Figure S29**). Other SUMI adducts including CPDTC-I (SI, **Figure S26**) and DDMAT-I (SI, **Figure S27**) had the same stereospecific preference for *trans*- configurations as demonstrated by ¹H NMR results. In the case of chiral ACS Paragon Plus Environment

R group such as CPADB-I, the ¹H NMR spectrum (Figure 7b and full spectrum shown in SI, Figure S32) showed a more complicated signal pattern than the non-chiral R group of CPDB-I due to two diastereomers derived from the chiral carbon of the R group, in which two sets of signals were very closely positioned. For instance, the signal for proton 1 in Figure 7b (inset) at $\delta 6.02 \sim 6.15$ ppm leads to two doublets centered at δ 6.06 ppm and 6.09 ppm, which belong to two different diastereomers. The CDTPA-I SUMI mono adduct had the same ¹H NMR profile (SI, **Figure S33**) as CPADB-I, attributed to the same chiral R group.

Computational modelling further confirmed the *trans*- configuration of Ind unit in the synthetic oligomers with the potential energy surface (PES) for the insertion of PMI into CPDTC-I being examined. In this reaction, the relative free-energy of four intermediate diastereometric configurations (*threo-cis*, erytheo-cis, threo-trans and erytheo-trans, SI, Figure S38) was compared to determine the most favorable reaction pathways. The modelled conditions are identical to the experimental ones (solution-phase in DMSO, at 25 °C). The results clearly showed that both *trans*- insertion reactions are more kinetically favorable (> 10 kJ/mol) than either *cis*- insertion pathway, suggesting that the Ind unit also prefers *trans*-insertion as for the PMI unit.

3.3 Stereochemistry of acyclic monomer unit in discrete oligomers

In the contrast to the behaviors observed for cyclic monomers, the SUMI of acyclic monomers (Scheme 2) into RAFT agents did not show stereospecificity due to the lack of cyclic structure and a free rotating single bond along the backbone. For instance, the ¹H NMR spectrum of BBTC-DEM (SI, Figure S41) showed two series of proton signals (a and b) which belong to two diastereomers, each of them with a corresponding pair of enantiomers. The signals for the same protons in each diastereoisomer presented different chemical shifts (0.12 ppm difference for the proton 5a and 5b of -CH-S-, 0.3 ppm for the proton 6a and 6b of -CH-CH-S-) and splitting patterns (doublet for the proton 7a and two quartets for proton 7b). 2D NMR (¹H-¹H COSY, ¹H-¹³C HSOC and ¹H-¹³C HMBC) was employed to confirm the structure and our conclusion. It is interesting to note that the integrals for two proton signals are different, which suggested that one of the diastereomers formed preferentially over the other during the SUMI process, presumably due to a difference in steric hindrance.

To summarize the above results, the stereospecificity in the SUMI process relies on the monomer structure; the cyclic monomer prefers *trans*- insertion regardless of the chirality of R groups, but the acyclic monomer shows a mixture corresponding to both *cis*- and *trans*- configurations. This can be attributed to steric hindrance and the restricted rotation of cyclic monomers during radical addition (or recombination step). In contrast, radical addition (or recombination) with acyclic monomers was not stereospecific due to free rotation around the single bond of the C-C backbone. ACS Paragon Plus Environment

 This stereochemical preference was independent of temperature and solvents. High temperature will not change the *trans*- opening of cyclic monomers (SI, **Figure S39**). Low polarity solvents, such as cyclohexane, was also shown to have no effect on the stereospecific insertion of cyclic monomer into RAFT agents (SI, **Figure S40**). Similarly, the photocatalysts investigated in this work do not induce stereospecificity, because the *trans*- configuration is formed in both the presence and absence of photocatalysts. The synthetic conditions used to synthesize the SUMI adducts varied depending on the RAFT agents and monomers employed. For instance, BBTC-M and CPDTC-I were prepared through two distinct photocatalytic processes. The synthesis of BBTC-M, described in the first section employed the PET-RAFT technique to activate the RAFT agent BBTC to generate benzyl radical using a photocatalyst ZnTPP under red LED light, followed by PMI monomer addition to form the desired adduct. CPDTC-I was synthesized by direct photolysis of the RAFT agent CPDTC under blue LED light, and subsequent single Ind addition. This protocol is similar to that used for single styrene insertion into a trithiocarbonate, which was described in a previous report using the photo-RAFT SUMI approach.⁵²

CONCLUSION

In this contribution, we report a robust and highly efficient method of sequential and alternating single unit monomer insertion for the preparation of discrete oligomers with sequence control and stereospecific configuration (*trans-*) of inserted cyclic monomers. Two families of cyclic monomers, indene and *N*-substituted maleimide, were inserted into RAFT agents alternatively one unit at a time in high reaction yields, which effectively controlled the monomer sequences. The structures were confirmed by an online NMR kinetic study, GPC and ESI-MS. Moreover, for each investigated cyclic monomers, stereospecific *trans*-insertion was observed due to steric hindrance from the growing oligomer chain. The stereochemistry of these oligomer adducts was verified by 1D and 2D NMR and XRD and explained by quantum-chemical modelling. This methodology provides a versatile approach to synthesize discrete polymers with precise sequence and location of functionalities, while also illustrating a powerful synthetic pathway to model oligomers for the investigation of absolute structure (e.g. stereochemistry) of synthetic polymers. The research to expand the monomer pool (i.e. other non-homopolymerizable monomer families⁸³ and functional indenes^{104, 105}) for this advanced method is currently under investigation.

ASSOCIATED CONTENT

Supporting information

Experimental details, NMR spectra and all raw computational data (Figures S1~S47, Tables S1, and ACS Paragon Plus Environment

Schemes S1~S5). This material is available free of charge via the Internet at http://pubs.acs.org.

Notes

The authors declare no competing financial interest.

AUTHOR INFORMATION

Corresponding authors

michelle.coote@anu.edu.au (M.C.) cbover@unsw.edu.au (C.B.)

j.xu@unsw.edu.au (J.X.)

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