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Tricyclononenes and tricyclononadienes as efficient monomers for controlled ROMP: understanding structure-propagation rate relationships and enabling facile post-polymerization modification

Living ring-opening metathesis polymerization of a library of tricyclic olefins was investigated for comparison to traditional bicyclic ROMP monomers, providing new insights into monomer reactivity and useful new monomers for macromolecular synthesis.

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Tricyclononenes and tricyclononadienes as efficient monomers for controlled ROMP: understanding structure–propagation rate relationships and enabling facile post-polymerization modification†

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Grubbs 3rd-generation (G3) pre-catalyst-initiated ring-opening metathesis polymerization (ROMP) remains an indispensable tool in the polymer chemist's toolbox. Tricyclononenes (TCN) and tricyclononadienes (TCND) represent under-explored classes of monomers for ROMP that have the potential to both advance fundamental knowledge (e.g., structure–polymerization kinetics relationships) and serve as practical tools for the polymer chemist (e.g., post-polymerization functionalization). In this work, a library of TCN and TCND imides, monoesters, and diesters, along with their *exo*-norbornene counterparts, were synthesized to compare their behaviors in G3-initiated ROMP. Real-time ¹H NMR was used to study their polymerization kinetics; propagation rates (k_p) were extracted for each monomer. To understand the relationships between monomer structure and ROMP propagation rates, density functional theory methods were used to calculate a variety of electronic and steric parameters for each monomer. While electronic parameters (e.g., HOMO energy levels) correlated positively with the measured k_p values, steric parameters generally gave improved correlations, which indicates that monomer size and shape are better predictors for k_p than electronic parameters for this data set. Furthermore, the TCND diester—which contains an electron-deficient cyclobutene that is resistant to ROMP—and its polymer p(TCND) are shown to be highly reactive toward DBU-catalyzed conjugate addition reactions with thiols, providing a protecting- and activating-group free strategy for post-polymerization modification.

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Introduction

Ruthenium-initiated ring-opening metathesis polymerization (ROMP) is an invaluable tool in the polymer chemist's repertoire due to the commercial availability of fast-initiating ruthenium initiators and various strained olefin monomers, its wide functional group tolerance, and the ease in which complex polymers may be synthesized in a rapid and controlled manner. Grubbs 3rd-generation pyridine-ligated pre-catalysts (G3) are most frequently used for living ROMP due to their fast initiation rates.¹ Moreover, *exo*-norbornenes (NB) are often

used as monomers due to their fast propagation rates, especially in comparison to their *endo* counterparts.^{2,3} *Exo*-NBs are, however, often more difficult to synthesize and more expensive than their *endo* isomers. Recent detailed studies of the mechanism⁴ and the effects of the *exo*-norbornene monomer structure on the rate of propagation and degree of “livingness” of these polymerizations^{5–8} have led to new insights that may guide the design of next-generation initiator and monomer classes. For example, Matson and coworkers^{5–8} have suggested that the HOMO energy levels of *exo*-NBs correlate with their G3-initiated ROMP propagation rates. Inspired by this work, and seeking synthetically accessible, functional monomers that do not compromise polymerization kinetics and controlled polymerization behavior, we identified tricyclononenes (TCN) and tricyclononadienes (TCND) as monomers ripe for further study (Fig. 1).

TCNs and TCNDs are tricyclic, strained olefins with fused cyclobutane and cyclobutene rings, respectively, connected to the two *exo* positions of the NB skeleton. Compared to other *exo*-substituted NB monomers, which often require high temperatures^{9–11} and/or precious metal catalysts^{8,12} for their

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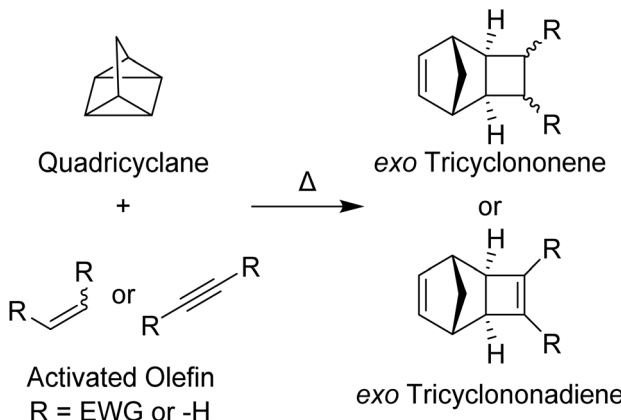


Fig. 1 General syntheses and structures of exo-tricyclononene and exo-tricyclonadiene analogs.

synthesis, *exo*-TCNs and TCNDs can be conveniently assembled by thermal cycloadditions of activated olefins and quadricyclane with perfect *exo* selectivity.^{13,14} While TCN and TCND-based polymers have been synthesized using Mo (Schrock-type initiators)¹⁵ and Ru (Grubbs 1st and 2nd generation initiators) initiated ROMP^{16,17} and studied for applications in gas separations,^{18–25} no studies of their performance under “living” ROMP conditions initiated by G3 or post-polymerization functionalization have been reported. We hypothesized that they may be competitive with commonly used *exo*-NBs in terms of ROMP propagation rates, but that they could offer advantages of facile synthesis and, in some cases, post-polymerization modification. Moreover, by accessing a training set of monomers, we sought to investigate if the HOMO level was primarily responsible for propagation rate as proposed by Matson and coworkers, or whether other molecular features (e.g., steric parameters) of the monomers may also impact performance.

Here, we synthesize a library of TCNs and TCNDs that incorporates a broad functional group scope, including imides, diesters, and monoesters. We assess their structures and properties, investigate their ROMP behavior using G3, and compare the results to analogous *exo*-NB derivatives. We identify trends in the relative propagation rates across these monomers, showing that while their HOMO energies are good predictors of reactivity, readily computed steric parameters (e.g., volume, solvent accessible surface area (SASA), and Sterimol B₅) offer improved structure-reactivity correlations across all *exo*-NB, TCN, and TCND monomer classes. Moreover, we demonstrate that the cyclobutene pendants of p(TCND)s can be excellent substrates for conjugate addition reactions with thiol nucleophiles (i.e., “thiol-Michael-type additions”),^{26–28} offering a straightforward methodology for efficient post-polymerization modification without the need for protecting groups or monomer activation (e.g., activated esters). These studies demonstrate that TCNs and TCNDs are attractive next-generation monomers for ROMP as (a) they are easy to synthesize in high stereochemical purity in as few as two simple steps from commercial starting materials; (b) they display propagation kinetics similar to or faster than many typical *exo*-NBs; and (c)

they offer facile routes to post-polymerization modification that are not possible for common *exo*-NBs. Finally, this study presents the first application of computed steric parameters to correlate monomer structure with propagation rates in ROMP, providing insights into the predictive design of future ROMP monomers.

Results/discussion

TCN and TCND monomer synthesis

Quadricyclane was prepared by photochemical isomerization of norbornadiene in batches of up to 17 g and yields up to >90% (ESI† Section 2). Heating quadricyclane in the presence of maleic anhydride gave TCN *exo*-anhydride **1** as a mixture of *anti* and *syn* diastereomers (67 : 33 *anti* : *syn*) in quantitative yield (Fig. 2a).^{29,30} Crystallization led to enrichment of the *anti*-**1** to 78 : 22 *anti* : *syn* (69% yield); further isolation of pure diastereomers was conducted following a subsequent synthetic step. TCN diester TCN-(CO₂Me)₂ was prepared by acid-catalyzed esterification of **1** with methanol.²⁹ The TCN imides TCN-NBn_{anti}, TCN-NCy, and TCN-NHex were prepared by condensation of **1** with benzylamine, cyclohexylamine, and *n*-hexylamine, respectively, in an analogous fashion to Warrener's procedure for preparing the *N*-methyl TCN imide from **1** and methylamine.³⁰ All of these TCNs were crystalline solids wherein crystallization afforded the *anti*-isomer in a >95 : 5 *anti* : *syn* ratio in each case.

We also sought TCN-NBn_{syn} for comparison to its *anti*-diastereomer (Fig. 2b). While TCN-NBn_{syn} and TCN-NBn_{anti} may potentially be separable by chromatography,³⁰ we identified a kinetic resolution method that leveraged the large difference in imidation rates between the *syn* (slow) and *anti* (fast) amic acids formed upon initial ring-opening opening of **1**, ultimately giving TCN-NBn_{syn} in 4% yield over 2 steps with >98 : 2 d.r. (ESI† Section 2). The relative configurations of TCN-NBn_{syn} and TCN-NBn_{anti} were unambiguously determined by single-crystal X-ray diffraction (Fig. 2b). TCND derivatives TCND-(CO₂Me)₂ and TCND-CO₂Me were synthesized by direct cycloaddition of quadricyclane with dimethylacetylenedicarboxylate and methyl propiolate, respectively (Fig. 2a).^{13,31}

We note that upon storage under ambient atmosphere for extended periods, TCND-(CO₂Me)₂ yellows and a new product is observed *via* ¹H NMR spectroscopy. A small sample of this impurity was isolated chromatographically; its ¹H NMR spectrum contained no olefinic resonances and its high-resolution mass spectrum suggested a mass 16 Da (i.e., 1 oxygen atom) larger than TCND-(CO₂Me)₂. Single-crystal X-ray analysis of the saponified derivative of this impurity revealed its identity to be the *exo*-epoxidized variant of TCND-(CO₂Me)₂ (ESI† Section 11). Thus, we recommend storage of uninhibited TCND-(CO₂Me)₂ under inert atmosphere, as is often done for olefinic monomers, to avoid its oxidative decomposition.

Finally, for comparison of ROMP propagation rates (*vide infra*), *exo*-NB analogs of each TCN and TCND were synthesized following previously reported procedures.^{32–34} All monomers and intermediates were isolated non-chromatographically, and the key intermediates towards TCNs (**1**) and TCNDs



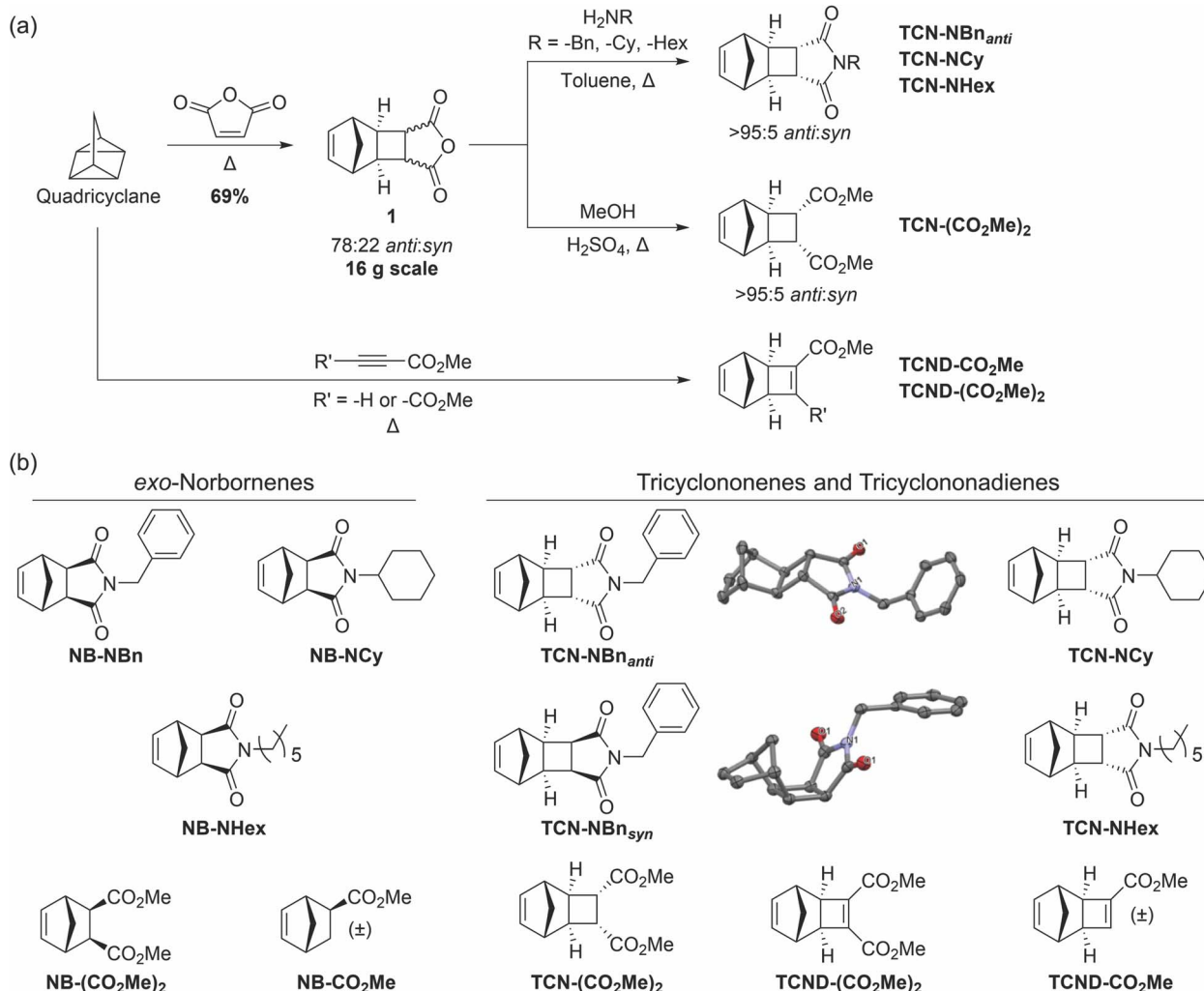


Fig. 2 (a) Synthetic pathways toward TCN and TCND monomers. (b) Library of exo-NBs, TCNs and TCNDs synthesized in this study. The crystal structures of TCN-NBn_{anti} and TCN-NBn_{syn} are shown with ellipsoids drawn at the 50% probability level. Hydrogen atoms are omitted for clarity; black = C; red = O; blue = N.

(quadricyclane) were synthesized on the 16 g and 17 g scales, respectively. Notably, the melting temperatures for all but one (TCND-CO₂Me) of the TCNs and TCNDs synthesized in this study were greater than their *exo*-NB analogs (from 10 °C to >150 °C higher), presumably due to their multicyclic, comparably rigid structures. In particular, all of the TCNs studied here have melting points >40 °C above room temperature, making them generally easy to handle and purify (crystallization).

ROMP of TCN and TCND monomers

Each TCN and TCND described above (200 equiv.) was exposed to G3 (1 equiv.) in CDCl₃ solvent to initiate ROMP. Reactions of TCNs were allowed to proceed for 15 minutes before quenching and precipitating twice from methanol, giving p(TCN)s in good yields (Table 1). Size exclusion chromatography (SEC) revealed low dispersities (D) and number-average molar masses (M_n) close to theoretical values for all p(TCN) samples (Fig. 3a–d).

SEC analysis of p(TCND-(CO₂Me)₂) prepared under the same conditions (*i.e.*, 15 min reaction time) showed a minor high molecular weight shoulder peak at $2 \times M_n$ indicative of cross-linking, as was also reported for Mo-initiated ROMP of this monomer (Fig. 3e).¹⁵ Quenching the ROMP reactions after a shorter time (2 min) using the same batch of TCND-(CO₂Me)₂ or a freshly distilled sample of TCND-(CO₂Me)₂ (discarding the first and last ~10–15% of the distillate) substantially decreased the size of this shoulder peak. Notably, conversion was unaffected by decreasing the reaction time to 2 min, as evidenced by the similar SEC traces, suggesting that this monomer is highly reactive. Moreover, these results indicate that the cyclobutene component of TCND-(CO₂Me)₂ remains largely intact under these ROMP conditions, providing an opportunity for further modification post-polymerization (*vide infra*).

While the synthesis of p(TCND-CO₂Me) appeared to be no different than for the other samples prepared here, the polymer product did not re-dissolve after the first precipitation. We



Table 1 Characterization of p(TCN)s and p(TCND)s prepared by G3-initiated ROMP^a

Polymer	M_n ^b (theo.)	M_n ^c (SEC)	D	Yield	<i>E/Z</i> ratio ^d
p(TCN-NB _n <i>anti</i>)	56.0 kDa	52.2 kDa	1.02	97%	37 : 63
p(TCN-NCy)	54.4 kDa	54.2 kDa	1.02	95%	34 : 66
p(TCN-NHex)	54.8 kDa	52.2 kDa	1.02	92%	37 : 63
p(TCN-(CO ₂ Me) ₂)	47.4 kDa	47.8 kDa	1.02	97%	33 : 67
p(TCND-(CO ₂ Me) ₂)	47.0 kDa	49.5 kDa	1.04	92%	30 : 70
p(TCND-CO ₂ Me)	35.3 kDa	59.3 kDa	1.97	75%	37 : 63

^a Monomer conversion was >98% in each case. ^b Theoretical M_n values were calculated assuming 100% monomer conversion. ^c SEC conditions: DMF (0.025 M LiBr) mobile phase, 60 °C, with differential refractive index and multi-angle light scattering detectors. A 100% mass recovery method was used to calculate M_n . ^d *E/Z* ratio was determined by the relative integrations of the *trans* (downfield) and *cis* (upfield) backbone olefinic protons in the polymers' ¹H NMR spectra.^{35,36}

hypothesized that polymer-associated residual Ru led to cross-linking reactions upon precipitation; the addition of excess pyridine (~2000 equiv. relative to G3) to the quenched ROMP reaction suppressed this cross-linking (ESI† Section 3), enabling NMR and SEC analysis of p(TCND-CO₂Me); however, SEC analysis showed a significant high molecular weight shoulder

(Fig. 3f). Notably, while cyclobutene 1-carboxylic esters, like the cyclobutene fragment of TCND-CO₂Me, cannot homopolymerize via G3-initiated ROMP,³⁷ they are known to slowly form eneonic carbenes that are kinetically stabilized by intramolecular chelation.³⁸ Thus, we propose that ROMP of TCND-CO₂Me occurs first through rapid polymerization of its NB-like alkene followed by slow cross-linking of its cyclobutene pendants, which are more sterically accessible than for TCND-(CO₂Me)₂ above.

¹H NMR spectroscopy was used to characterize the structures of these p(TCN)s and p(TCND)s in more detail (Fig. 4). Resonances associated with each of the monomer protons could be identified in the polymers. Compared to p(*exo*-NB) analogs (ESI† Section 3), the backbone olefins of p(TCN)s and p(TCND)s were slightly skewed toward the *Z* configuration: *E* : *Z* ranged from 48 : 52 to 37 : 63 for p(*exo*-NB)s while p(TCN)s and p(TCND)s ranged from 37 : 63 to 30 : 70 (Table 1). The spectrum of p(TCND-CO₂Me) contained an extra downfield resonance at 6.8 ppm that we attribute to the olefinic proton of the cyclobutene. Otherwise, the spectra generally resembled those of their p(*exo*-NB) counterparts, with the addition of two proton resonances in the carbocyclic region (~1.4 ppm–3.5 ppm) for the p(TCN)s.

We anticipated that TCN and TCND monomers would display controlled polymerization behavior due to their high

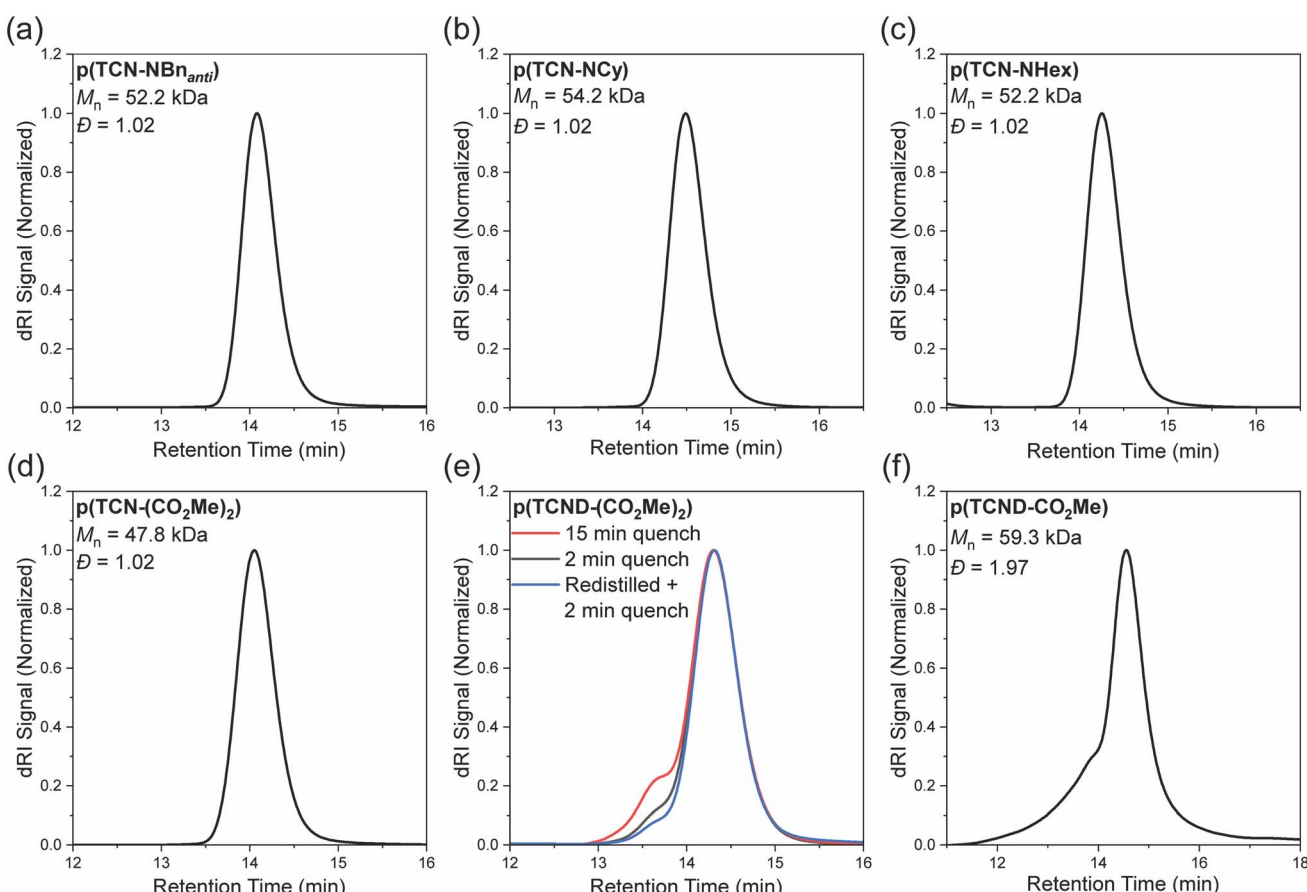


Fig. 3 (a–d) SEC traces of p(TCN)s. (e) Overlaid SEC traces of p(TCND-(CO₂Me)₂) when quenched after 15 min, 2 min, or after carefully distilling the monomer and quenching after 2 min (f) SEC trace of p(TCND-CO₂Me) after working up the polymer in the presence of excess pyridine.



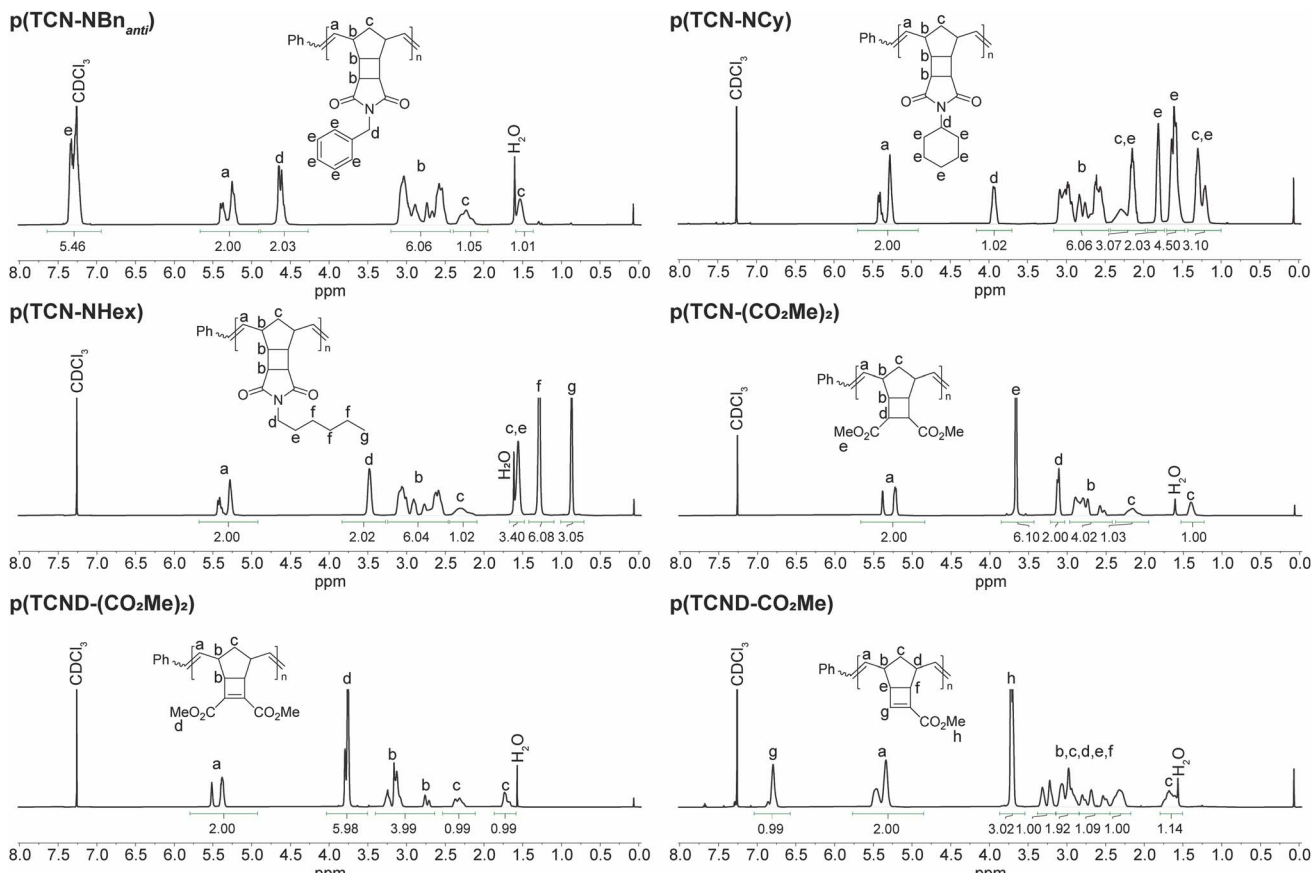


Fig. 4 ^1H NMR spectra for the p(TCN)s and p(TCND)s prepared in this study.

ring strain and low likelihood of backbiting/chain transfer. In support of this hypothesis, $M_{n,\text{SEC}}$ values for **p(TCN-NBn_{anti})** agreed well with the theoretical values and increased linearly as a function of conversion up to a degree of polymerization (DP) of 500 (Fig. 5a). Moreover, a plot of $\ln([TCN-NBn_{anti}]_0/[TCN-$

$NBn_{anti}])$ was linear as a function of time (Fig. 5b), and D remained low throughout the polymerization (Fig. 5a), suggesting minimal irreversible termination or chain transfer. Finally, to demonstrate the active nature of the **p(TCN-NBn_{anti})** Ru-alkylidene chain ends, a block copolymer **p(TCN-NBn_{anti})₁₀₀-**

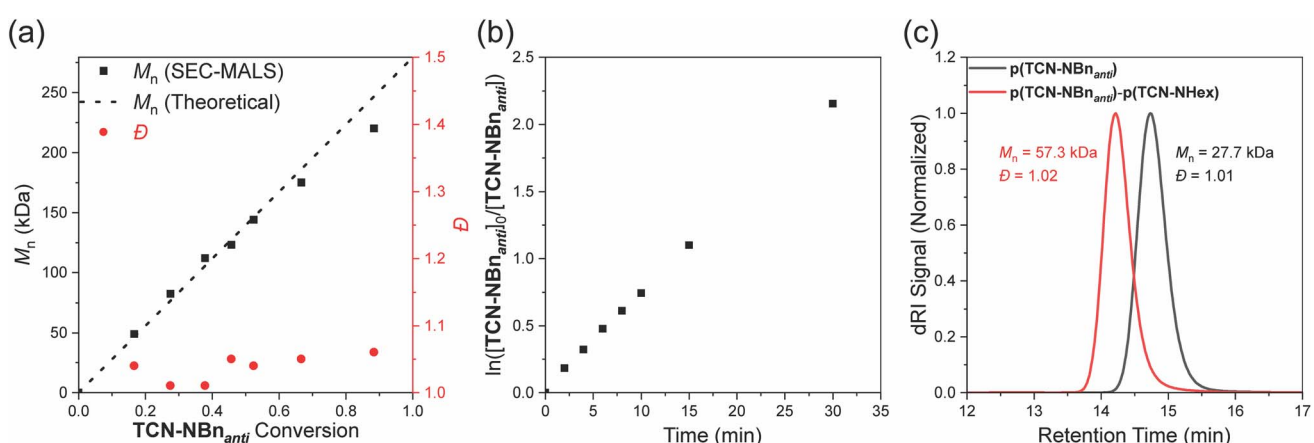


Fig. 5 (a) Plot of $M_{n,\text{SEC}}$ as a function of conversion for the polymerization of TCN-NBn_{anti} with G3. Conditions: 500 equiv. TCN-NBn_{anti}, 5 equiv. pyridine, 1 equiv. G3, 500 mM in CDCl_3 , RT. (b) Plot of $\ln([TCN-NBn_{anti}]/[TCN-NBn_{anti}]_0)$ versus time for the polymerization described in panel a. (c) SEC analysis of the first and second blocks of the block copolymerization of TCN-NBn_{anti} (DP 100) with TCN-NHEx (DP 100). Conditions: 100 equiv. TCN-NBn_{anti}, G3, 100 mM CDCl_3 , RT, then 100 equiv. TCN-NHEx (100 mM soln. in CDCl_3).

b-p(TCN-NHex)₁₀₀ was synthesized *via* sequential monomer addition (Fig. 5c). SEC analysis showed a clear peak shift to shorter retention time upon addition of **TCN-NHex**, suggesting complete chain extension; D for the block copolymer remained low (1.02).

¹H NMR spectroscopic kinetic investigations of TCN and TCND ROMP reactions

Real-time ¹H NMR spectroscopy was used to evaluate the ROMP propagation kinetics for each monomer shown in Fig. 2 in the presence of pyridine (5 equiv. relative to G3; note: pyridine was added to retard the polymerization reactions sufficiently for ¹H NMR analysis). For all monomers except **TCND-CO₂Me**, conversion was measured by comparison of the monomer signal to an added internal standard (mesitylene). The additional downfield olefin resonance of **p(TCND-CO₂Me)** overlapped with that of the internal standard; monomer conversion was measured by relative integration of the monomer signal to the backbone olefin signals in this case. Each polymerization followed pseudo-first order kinetics up to $\geq 95\%$ conversion, indicative of a controlled process (Fig. 6a-f). The propagation rates (k_p), obtained by linear fitting of $\ln([M]_0/[M])$ vs. time plots (Fig. 6, insets, and ESI† Section 4), spanned a factor of ~ 13 ($k_p =$

$0.094\text{--}1.259\text{ min}^{-1}$) for the monomers analyzed in this study (Table 2).

The k_p values for imides **TCN-NBn_{anti}**, **TCN-NBn_{syn}**, **TCN-NCy**, and **TCN-NHex** were within $\sim 20\%$ of each other and were slightly lower than their *exo*-NB imide counterparts. A similar trend was observed for the monoesters, but to a larger extent: k_p for **NB-CO₂Me** was 2.2-fold higher than for **TCND-CO₂Me**. Notably, however, the opposite trend was observed for the diester monomers; k_p for **TCND-(CO₂Me)₂** was greater than **TCN-(CO₂Me)₂**, which was greater than **NB-(CO₂Me)₂**.

Comparing across the imides, diesters, and monoesters, the imides had the lowest k_p values without exception; the fastest imide (**NB-NCy**) had a $\sim 19\%$ lower k_p than the slowest diester/monoester (**NB-(CO₂Me)₂**). This result agrees well with the trends reported for imide *versus* ester anchor groups for both small molecule monomers and macromonomers.⁵⁻⁷ Comparison of the monoesters and diesters shows that while the monoester **TCND-CO₂Me** and diester **TCND-(CO₂Me)₂** propagate at nearly the same rate, **NB-CO₂Me** propagates 5.6 times faster than its diester counterpart **NB-(CO₂Me)₂**. Altogether, these results demonstrate that TCN and TCND monomers possess similar living behavior and G3-initiated ROMP k_p values compared to traditional *exo*-NBs.

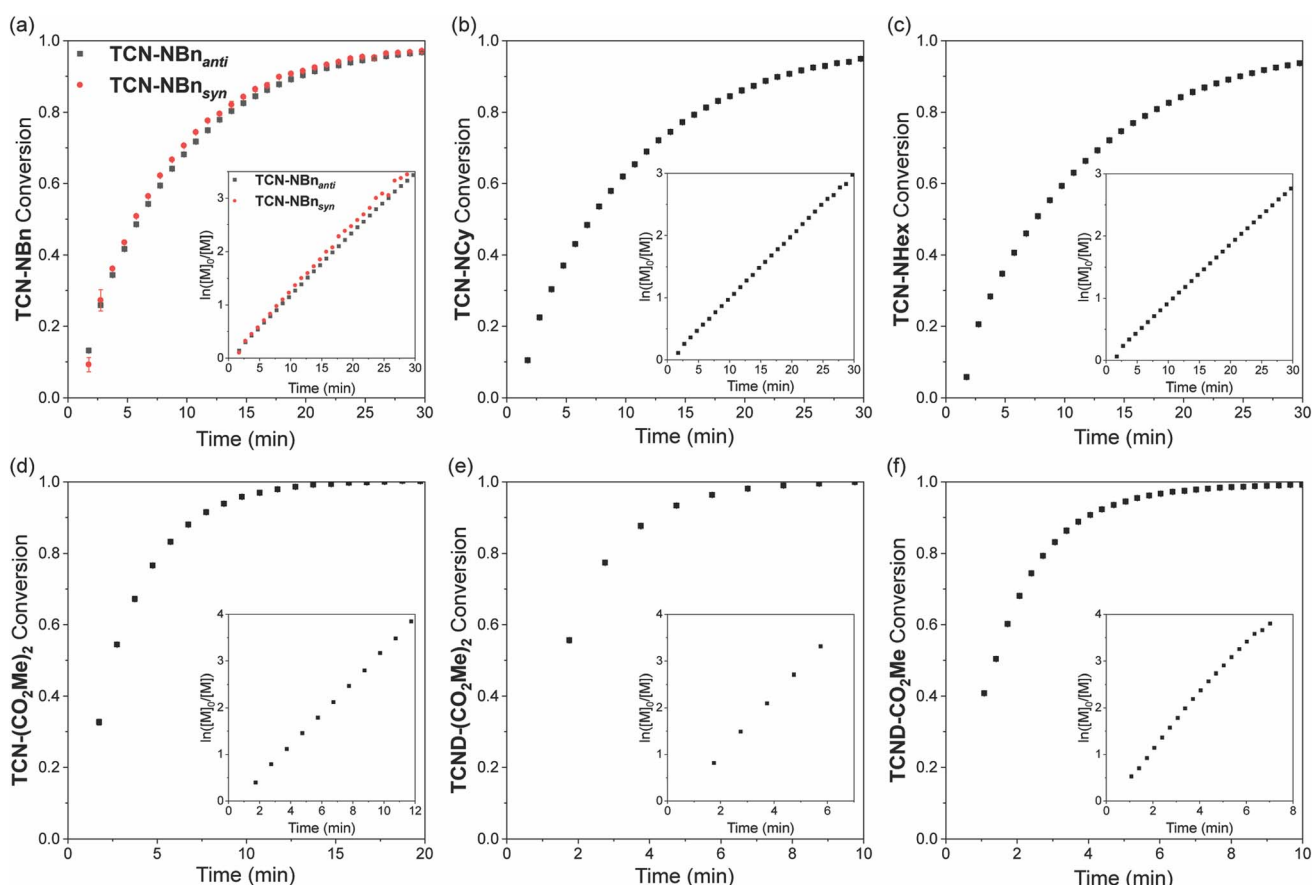


Fig. 6 (a-f) ROMP propagation kinetics of all TCN and TCND monomers in this study. The raw data are presented with an inset plot of $\ln([M]_0/[M])$ versus time. Each was run in triplicate, and error bars are plotted as \pm SD in the raw data plots. Conditions: 95 mM monomer (200 equiv.), 2.4 mM pyridine (5 equiv.), 0.48 mM G3 (1 equiv.) and internal standard (mesitylene) in CDCl_3 at 25 °C.



Table 2 ROMP propagation rates and half-lives of *exo*-NB, TCN, and TCND monomers extracted from the real-time NMR kinetic data^a

Monomer	k_p (min ⁻¹)	$t_{1/2}$ (min)
NB-NBn	0.157 ± 0.001	4.4
TCN-NBn_{anti}	0.117 ± 0.001	5.9
TCN-NBn_{syn}	0.122 ± 0.001	5.7
NB-NCy	0.183 ± 0.001	3.8
TCN-NCy	0.101 ± 0.001	6.9
NB-NHex	0.140 ± 0.001	5.0
TCN-NHex	0.094 ± 0.001	7.4
NB-(CO₂Me)₂	0.225 ± 0.001	3.1
TCN-(CO₂Me)₂	0.342 ± 0.001	2.0
TCND-(CO₂Me)₂	0.622 ± 0.007	1.1
NB-CO₂Me	1.259 ± 0.014	0.6
TCND-CO₂Me	0.564 ± 0.010	1.2

^a Conditions: 95 mM monomer (200 equiv.), 2.4 mM pyridine (5 equiv.), 0.48 mM G3 (1 equiv.) and internal standard (mesitylene) in CDCl₃ at 25 °C.

Density functional theory (DFT) calculations of monomer electronic and steric parameters to predict k_p

Matson and coworkers have proposed that the DFT-calculated HOMO energy levels of *exo*-NB monomers correlate well with their k_p values in G3-initiated ROMP.⁵⁻⁸ To determine if TCNs and TCNDs follow this trend, the HOMO energies for each monomer shown in Fig. 2b were calculated using the M06-2X functional and def2-TZVP basis set (with a conductor-like polarizable continuum model for chloroform) and plotted against the measured $\ln(k_p)$ values (Fig. 7b). The HOMO energies ranged from -183 kcal mol⁻¹ to -194 kcal mol⁻¹, which falls within the range reported for other ROMP monomers (-174 kcal mol⁻¹ to -197 kcal mol⁻¹) calculated with the same functional and basis set.^{5,6,8} In agreement with Matson and coworkers, a positive correlation between k_p values and HOMO energies was observed; however, **NB-CO₂Me** stands out as having a much larger k_p value than its HOMO energy would predict. Notably, **NB-CO₂Me** is the only monosubstituted

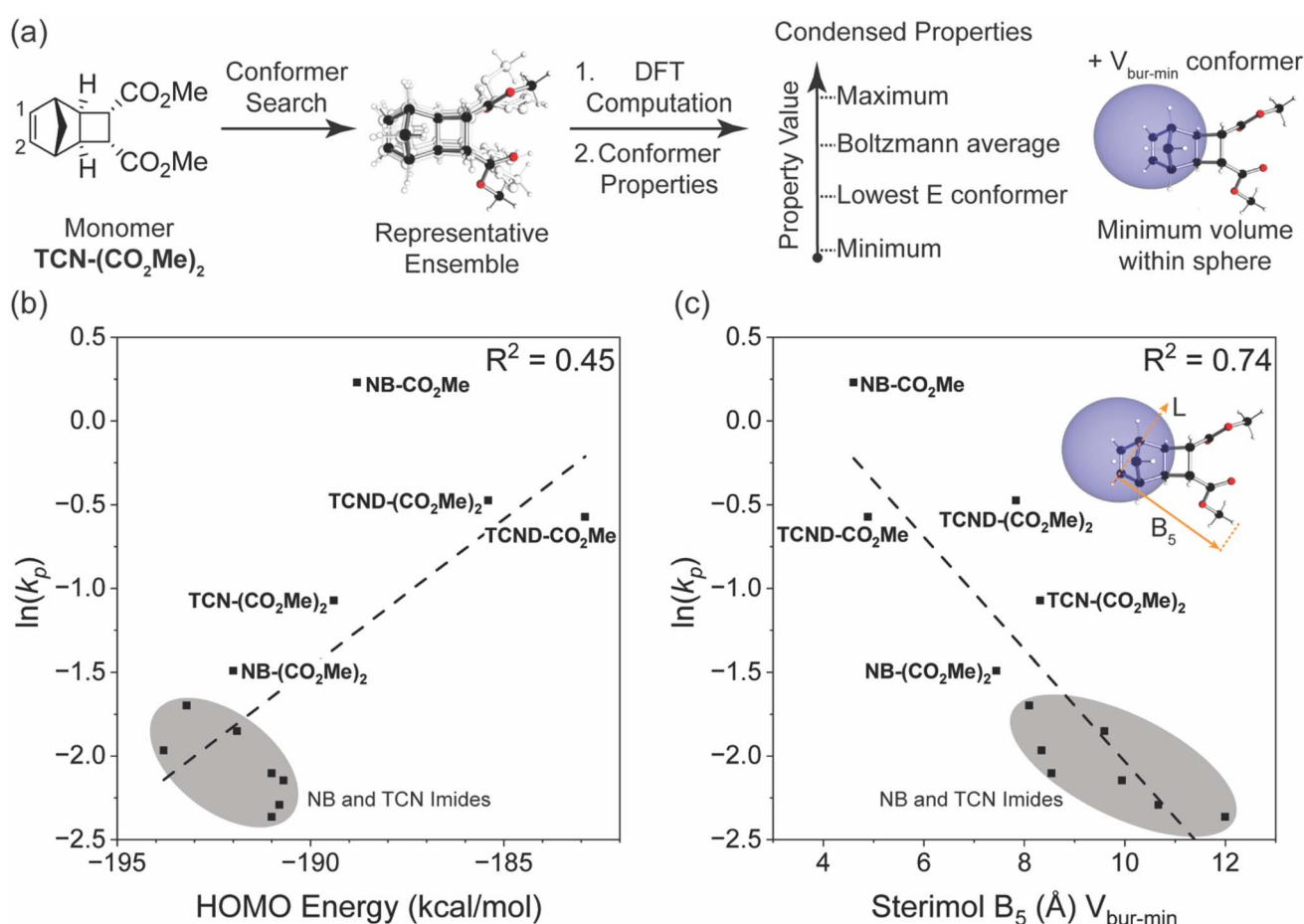


Fig. 7 (a) Modeling workflow for steric parameters. AQME³⁹ (rd-kit⁴⁰) was used to convert monomer SMILES strings into a representative conformer ensemble.⁴⁴ The ensemble was subjected to DFT computation and conformer properties were obtained. For each conformer the maximum, Boltzmann average, lowest energy (*E*) conformer, minimum, property value was obtained. Property values for the $V_{\text{bur-min}}$ conformer at C₁ were obtained as well to further probe steric effects. (b) Plot of measured monomer $\ln(k_p)$ versus their HOMO energy levels calculated from their M06-2X/def2-TZVP optimized geometries. (c) Plot of measured monomer $\ln(k_p)$ versus their calculated Sterimol B₅ length from the minimum buried volume conformer of the adjacent active site olefinic carbon C₂ (B3LYP/def2SVP/M06-2X/def2TZVP).



norbornene (only one of the two *exo* positions is substituted) of the monomers studied here.

Grubbs³ and Matson also found that monosubstituted *exo*-NBs polymerize faster than disubstituted variants; Matson attributed this difference either to the possibility that mono-substituted monomers can form regioisomers where their substituents point away from the ruthenium center (a steric effect),⁷ or to electronic effects (HOMO level and HOMO/LUMO gap).^{5–7} While the chelating ability of NB monomer substituents has also been shown to significantly affect the activation parameters of polymerization and may slow the polymerization rate by ground-state stabilization in certain cases,³ the chelating ability of *exo*-NB monomers seems to have little correlation with their propagation rates.⁵ We note that **NB-CO₂Me** has a ~4-fold larger k_p than **TCN-(CO₂Me)₂** despite having nearly the same calculated HOMO energy (0.6 kcal difference), and that **TCND-(CO₂Me)₂** and **TCND-CO₂Me** both have higher-lying HOMOs (by ~4 and ~6 kcal mol⁻¹, respectively) than **NB-CO₂Me** but significantly lower k_p values. Overall, while HOMO energy is a useful tool for predicting ROMP reactivity for *exo*-NB-like scaffolds, other effects, such as steric effects near the propagation site, may impact the rate as well.

These findings led us to consider alternative readily-computed structural descriptors that may provide better correlations with ROMP monomer performance over a wider structure range, and may then also provide greater predictive power for the design of novel monomers. We began by collecting DFT-derived parameters from an AQME derived conformer ensemble³⁹ using RDKit⁴⁰ for each monomer. From the conformer ensemble, five representative conformers were selected to extract a range of steric and electronic parameters (Fig. 7a).^{41,42} The ensemble conformer properties were condensed to include the lowest energy conformer (low E), minimum (min), maximum (max), and Boltzmann derived parameter values. Additionally, we also extracted condensed properties from the conformer presenting the minimum percent buried volume ($V_{\text{bur-min}}$) parameter value at the active site olefinic carbon (C₁), as we hypothesized that steric effects may be a driver of the observed rate differences. Percent buried

volume probes the proximal steric environment by placing a 3 Å sphere at the site of interest and measuring the resulting overlap.⁴³ The library of parameters was regressed against $\ln(k_p)$ to reveal a range of good correlations with various parameters describing steric effects (molar volume (MV), solvent accessible surface area (SASA) and Sterimol B₅). These correlations all resulted from the $V_{\text{bur-min}}$ conformer values located at the active site olefinic carbon C₁. A representative correlation using Sterimol B₅ is depicted in Fig. 7c. This parameter describes the display of steric information for how the adjacent olefinic carbon C₂ may present itself to the propagating Ru chain end (Fig. 7a). The other two strong correlations, MV and SASA, represent the shape of the monomer while accounting for the local environment defined by the $V_{\text{bur-min}}$ conformer. Notably, the Sterimol B₅ and the MV $V_{\text{bur-min}}$ conformers were found to be collinear, and the same parameters derived from other conformers (*i.e.*, low E , min, and max values) do not perform nearly as well (ESI† Section 13). Overall, these data suggest that the overall size and shape of the monomer has a significant impact on polymerization kinetics.

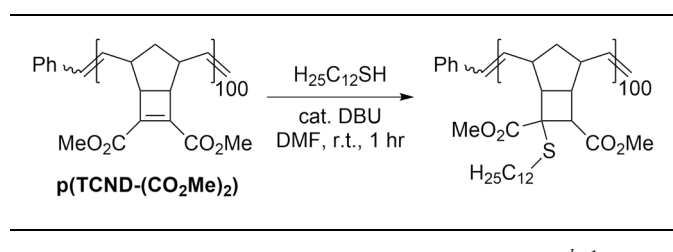
Further derivatization of **TCND-(CO₂Me)₂** and post-polymerization functionalization of **p(TCND-(CO₂Me)₂)**

In addition to their straightforward syntheses and good k_p values, **TCND-(CO₂Me)₂** and **TCND-CO₂Me**, and their corresponding polymers **p(TCND-(CO₂Me)₂)** and **p(TCND-CO₂Me)**, offer unique opportunities for further derivatization and post-polymerization functionalization, respectively, compared to *exo*-NBs. While their cycloaddition reactivity has been explored,^{45–47} we suspected that their strained, electron-deficient cyclobutene substituents would make them good electrophiles for conjugate addition reactions (*e.g.*, “thiolic-Michael” addition). To test this idea, **TCND-(CO₂Me)₂** was exposed to dodecanethiol (1.0 equiv.) in the presence of 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU) (10 mol%) in CDCl₃ for 30 min (Table 3, entry 1); 47% conversion to the conjugate addition product was observed under these conditions, while 0% conversion was observed in the absence of DBU. Polar

Table 3 Conjugate addition of dodecanethiol to various conjugate acceptors

Entry	Conjugate acceptor	LUMO energy (kcal mol ⁻¹)	Solvent	DBU (mol%)	Conversion	Conjugate Acceptors
						$\xrightarrow[0.1 \text{ M}, 25^\circ\text{C}, 30 \text{ min}]{1 \text{ equiv } \text{H}_25\text{C}_{12}\text{SH}, \text{cat. DBU, Solvent}}$
1	TCND-(CO₂Me)₂	-30	CDCl ₃	10%	47%	
2	TCND-(CO₂Me)₂	-30	DMF-d ₇	1%	87%	
3	DDM	-8	CDCl ₃	10%	0%	
4	TCND-CO₂Me	-10	CDCl ₃	10%	3%	



Table 4 Optimization of post-polymerization modification of **p(TCND-(CO₂Me)₂)** with dodecanethiol^a

Entry	H ₂₅ C ₁₂ SH equiv.	DBU equiv.	Conversion ^b (¹ H NMR)
1	5	20 mol%	>95%
2	2	20 mol%	>95%
3	2	5 mol%	>95%
4	1.5	5 mol%	>95%
5	1.1	5 mol%	95%
6	1.1	2.5 mol%	93%
7	1.2	5 mol%	>95%

^a Conditions: **p(TCND-(CO₂Me)₂)** in DMF (50 mg mL⁻¹) at room temperature. Dodecanethiol was added, followed by the addition of DBU as a stock solution in DMF (1:9 v/v DBU:DMF) and was allowed to react for 1 hour. ^b Conversion was assessed by comparing the integration of the backbone olefinic protons (4.8 ppm – 5.7 ppm) versus the integration of the terminal –CH₃ of the dodecanethiol side-chain (0.9 ppm).

solvents are known to accelerate conjugate addition reactions that involve polar mechanisms;⁴⁸ when DMF-d₇ was used as the solvent for this reaction, 87% conversion was achieved in 30 min using only 1 mol% DBU (Table 3, entry 2).

Notably, dimethyl 2,3-dimethylmaleate (DDM), a less-strained electrophile with a similar substitution pattern, did not react under the CDCl₃ conditions described above (Table 3, entry 3), and **TCND-CO₂Me** reached only 3% conversion (Table 3, entry 4). These results correlate with the DFT calculated LUMO energies (M06-2X/def-TZVP) for each molecule (Table 3), suggesting that the rigid structure of **TCND-(CO₂Me)₂** paired with its two electron-withdrawing ester substituents make it a better electrophile for conjugate addition. Meanwhile, as discussed above, these structural features are also key for (1) ensuring a high ROMP *k_p* (by avoiding steric hindrance associated with diesters appended directly to the *exo*-NB) and (2) avoiding undesired cross-linking during ROMP (as observed for **TCND-CO₂Me**), making **TCND-(CO₂Me)₂** a particularly promising monomer for further exploration.

We hypothesized that the electrophilicity of **TCND-(CO₂Me)₂** would not change significantly after ROMP, rendering **p(TCND-(CO₂Me)₂)** a similarly useful substrate for post-polymerization conjugate addition reactions. This hypothesis was supported by DFT calculations (M06-2X/def-TZVP); the LUMO energy of a model substrate designed to mimic the **p(TCND-(CO₂Me)₂)** backbone—the ethenolysis product of **TCND-(CO₂Me)₂**—was very similar (−32 kcal mol^{−1}) to that of **TCND-(CO₂Me)₂** (−30 kcal mol^{−1}). Moreover, nearly quantitative conjugate addition was observed when **p(TCND-(CO₂Me)₂)** was exposed to dodecanethiol (5 equiv.) and DBU (20 mol%) in DMF solvent for 1 h under ambient atmosphere (Table 4, entry 1). The reaction solution became cloudy almost immediately after addition of the DBU, suggesting rapid conjugate addition and limited solubility of the functionalized polymer product in DMF. Decreasing the number of equivalents of thiol to 2 (Table 4,

Table 5 Post-polymerization modification of **p(TCND-(CO₂Me)₂)** with a variety of thiols

Entry	Thiol	Atmosphere	M _n initial	M _n final	Conversion (SEC)	Conversion (¹ H NMR)
1		Air	27.0 kDa	33.4 kDa	33%	45%
2		N ₂	27.0 kDa	39.7 kDa	66%	64%
3		Air	27.0 kDa	32.4 kDa	29%	16%
4		N ₂	27.0 kDa	33.2 kDa	34%	36%
5		Air	27.0 kDa	41.4 kDa	Quant.	N/A
6		N ₂	27.0 kDa	40.0 kDa	Quant.	N/A
7		Air	27.0 kDa	39.4 kDa	Quant.	N/A
8		Air	27.0 kDa	45.4 kDa	Quant.	>95%



entry 2) and lowering the catalyst loading to 5 mol% (Table 4, entry 3) did not affect the conversion. Reduction of the equivalents of thiol to 1.5 (Table 4, entry 4) did not affect conversion either; however, further decreasing the amount of thiol to 1.1 equiv. gave slightly reduced conversion (95%, Table 4, entry 5), which was reduced further with a lower catalyst loading of 2.5 mol% (93%, Table 4, entry 6). Over 95% conversion was achieved with 1.2 equiv. of thiol and 5 mol% DBU (Table 4, entry 7); these conditions were chosen as optimal for further studies.

A variety of thiol nucleophiles was explored using these optimized conditions (Table 5); ^1H NMR and SEC were used to measure conversion of each reaction, with the two methods showing agreement within $\sim 15\%$ in most cases. Incomplete conversion was observed for 4-*tert*-butylthiophenol and 2-naphthalenethiol under ambient (Table 5, entries 1 and 3) or inert (N_2) atmosphere (Table 5, entries 2 and 4), which we attribute to the steric bulk of these thiols. Nearly quantitative functionalization was observed for methyl thioglycolate (Table 5, entries 5 and 6) regardless of atmosphere; however, the SEC peak shape for the product showed distinct broadening and tailing compared to the initial **p(TCND-(CO₂Me)₂)**, which may be due to possible deleterious side-reactions promoted by the relatively acidic α -protons of the α -mercaptopropanoate (ESI† Section 9).^{49,50} By contrast, 1-methyl-3-mercaptopropionate, a β -mercaptopropanoate, underwent high conversion to give a polymer product with no significant change in peak shape (Table 5, entry 7). Finally, hexanethiol also gave high conversion under these conditions (Table 5, entry 8).

Conclusions

This work represents the first detailed study of the polymerization behavior of TCN and TCND monomers in G3-initiated ROMP. A variety of functional (imides, monoesters, and diesters) TCNs, TCNDs, and *exo*-NBs were synthesized and their ROMP propagation rates were measured to span a factor of ~ 13 . To understand the relationship between monomer structure and propagation rate, the monomers were featurized using DFT to extract a variety of electronic and steric parameters and search for correlations with measured propagation rates. While simple electronic parameters (e.g., HOMO levels) positively correlated with monomer k_p , steric parameters that described the overall size and shape of the monomer were more strongly correlated with k_p . **TCND-(CO₂Me)₂**—which contains an electron-deficient cyclobutene—was discovered to be highly active towards DBU-catalyzed conjugate addition of dodecanethiol. This conjugate addition reaction was optimized on **p(TCND-(CO₂Me)₂)**, and a variety of thiol nucleophiles were used to functionalize **p(TCND-(CO₂Me)₂)** in this post-polymerization modification reaction.

Overall, this study revealed that TCNs and TCNDs are a synthetically accessible class of ROMP monomers that do not compromise on activity or controlled polymerization behavior in comparison to other commonly used monomers (e.g. *exo*-NBs). In particular, **TCND-(CO₂Me)₂** is a uniquely promising monomer due to its ease of synthesis (2 steps from commercial materials), fast ROMP kinetics, and its innate activity towards

post-polymerization functionalization with thiols; we envision its future widespread use for functional polymer design. Finally, our experimental and computational study revealed that monomer steric parameters correlate most strongly with their measured propagation rates; these results offer the polymer chemist both the intuitive framework and computational tools for developing future novel ROMP monomers.

Data availability

The datasets supporting this article have been uploaded as part of the ESI.†

Author contributions

Conceptualization: all authors. Data curation: L. K and T. M. formal analysis: L. K. and T. M. funding acquisition: J. J and M. S. investigation: L. K and T. M. methodology: L. K. and T. M. project administration: all authors. Resources: all authors. Software: T. M. and M. S. supervision: J. J. and M. S. validation: L. K. and T. M. visualization: L. K. and T. M. writing – original draft: L. K. and J. J. writing – review and editing: all authors.

Conflicts of interest

There are no conflicts of interest to declare.

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