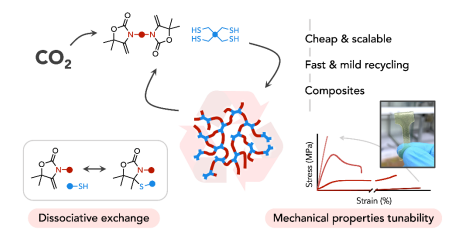
Covalent adaptable networks through dynamic *N,S*-acetal chemistry: toward recyclable CO₂-based thermosets

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Abstract



Finding new chemistry platforms for easily recyclable polymers has become a key challenge to face up environmental concerns and the growing plastics demand. Here, we report a dynamic chemistry between CO₂-sourced alkylidene oxazolidones and thiols, delivering circular non-isocyanate polyurethane networks embedding *N*,*S*-acetal bonds. The production of oxazolidone monomers from CO₂ is facile and scalable starting from cheap reagents. Their copolymerization with a polythiol occurs under mild

conditions in the presence of a catalytic amount of acid to furnish polymer networks. The polymer structure is easily tuned by virtue of monomer design, translating in a wide panel of mechanical properties similar to commodity plastics, ranging from PDMS-like elastomers (with Young's modulus (Ε) of 2.9 MPa and elongation at break (ε_{break}) of 159%) to polystyrene-like rigid plastics (with E = 2400 MPa, ε_{break} = 3%). The highly dissociative nature of the N,S-acetal bonds is demonstrated and exploited to offer three different recycling scenarios to the thermosets: (1) the mechanical recycling by compression molding, extrusion or injection molding – with multiple recycling (at least 10 times) without any material properties deterioration, (2) the chemical recycling through depolymerization, followed by repolymerization, also applicable to composites, and (3) the upcycling of two different oxazolidone-based thermosets into a single one with distinct properties. This work highlights a new facile and scalable chemical platform for designing highly dynamic polymer networks containing elusive oxazolidone motifs. The versatility of this chemistry shows great potential for the preparation of materials (including composites) of tunable structure and properties, with multiple end-of-life scenarios.

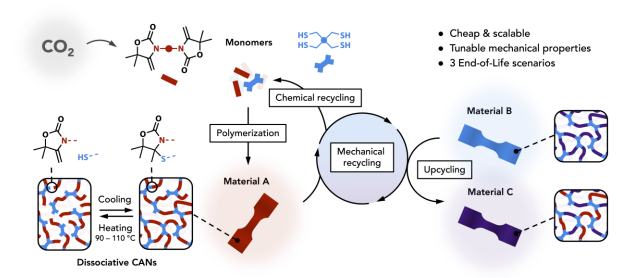
Introduction

Plastics are ubiquitous in our daily life on account of their vast panel of applications within most economical sectors. The global plastic use accounted for 460 million tons in 2019 and, following worldwide trends, is expected to triple by 2060¹. Regrettably, plastic waste became one of the greatest environmental threats causing irreversible damages to many ecosystems on the planet. Replacing them is not a feasible option as there are no viable alternatives so far which can match their outstanding features: low weight, affordability, versatility, and durability. A worldwide effort is devoted to re-think the plastics design through valorization of waste²,³ or by utilizing biobased and/or CO₂-sourced molecules⁴-6 to circumvent fully fossil-based chemistry. However, their recycling must be considered as importantly as their production to tackle environmental concerns³-10. Thus, appropriate solutions need to be developed in both the design and the recyclability of these materials.

In the recent years, covalent adaptable networks (CANs) have emerged as a new class of polymers whose properties lie between thermoplastics and thermosets 11-15. Despite their cross-linked 'fixed' structure, recycling is enabled through exchange reactions between dynamic bonds under a certain trigger, most generally heat 16. Many works already took advantage of bond dynamics to reprocess conventionally unrecyclable thermoset materials, notably via smart insertion of dynamic disulfide, boronic ester, acetal or Diels-Alder adducts bonds within the polymer matrix 17-21. However, the reshaping process is most often limited by the high cross-link density and associated

viscosity even at high temperature, restraining their processing to compression molding^{22,23}. Most examples of CANs reprocessed using a larger arsenal of techniques are designed with lower cross-link densities²⁴ through long oligomeric or polymeric chains within their structure to reach lower viscosities at high temperature^{20,25–28}.

CANs can be categorized in two main categories depending on their bond exchange mechanism²⁹. Associative CANs are characterized by a constant cross-link density during reprocessing as bonds associate before breaking, rendering their fast relaxation challenging. To address this issue, smart molecular engineering through additional functional groups and catalysts¹⁶ was successfully implemented to extrude associative CANs imparted with enough short relaxation times²². In contrast, dissociative CANs contain bonds that follow a stepwise elimination and addition mechanism, resulting in a loss of cross-link density as a function of temperature.³⁰



Scheme 1. Illustration of the alkylidene oxazolidone – thiol chemistry, and overview of the three recycling scenarios of *N*,*S*-acetal oxazolidone-based dissociative CAN materials (mechanical, chemical and upcycling).

However, the addition of large amount of catalyst or the insertion of specific groups in the matrix are generally needed to shift enough the equilibrium toward the dissociated state to endow the material with enough low viscosity to be reprocessed like thermoplastics^{24,31}. Another challenge associated with CANs is the access to milder reprocessing conditions. Most dynamic networks need long exposures to high

temperatures and pressures to be reshaped, typically above 150 °C from some minutes to several hours^{30,32–34}. Such harsh conditions might generate unwanted side reactions,^{14,32,35–37} resulting in material properties alteration during multiple reshaping cycles. Fast reprocessing of CANs in mild conditions is therefore of key interest to facilitate recycling. Lastly, finding new chemistries of high potential in terms of recyclability, cost, scalability, and tunability toward materials with modular thermomechanical properties became a new pillar in the search for a more sustainable and affordable future for plastics.

In the last years, many works were devoted to design sustainable polymers containing degradable acetal linkages^{38,39}. Recent contributions related to CANs have also shown that acetals and thioacetal derivatives undergo stress relaxation through bond exchange at elevated temperatures, rendering the acetal chemistry a promising platform for making greener materials^{31,40–42}.

In this work, we introduce a new highly dynamic chemistry based on N,S-acetals and its application in CANs. Unlike conventional synthetic approaches, N,S-acetals are synthesized by a facile cationic thiol-ene reaction between thiol and exovinylene oxazolidone obtained from CO₂. Their utility is highlighted for the edition and reutilization of thermoset poly(oxazolidone)s, a type of emerging isocyanate-free polyurethanes with remarkable thermal and chemical stability⁴³⁻⁴⁵. Mechanistic and kinetic insights of the N,S-acetal formation and its dynamics were first illustrated on model compounds. Then, diverse CO₂-sourced exovinylene bisoxazolidone monomers were prepared and copolymerized with a polythiol at room temperature to provide thermosets featuring a wide diversity of mechanical properties ranging from PDMS-like elastomers to rigid polystyrene-like ones. The N,S-acetal bond in the thermosets was proven to be highly dynamic by means of rheology through stress relaxation and temperature sweep cycling experiments. This unique bond dynamics was then exploited for developing three different recycling scenarios: (i) the facile and rapid mechanical reprocessing by compression molding, extrusion or injection molding of densely cross-linked bulk material, (ii) the closed-loop chemical recycling through depolymerization into the offspring monomers, followed by their repolymerization, and (iii) an upcycling strategy by repurposing two different oxazolidone-based thermosets into a single one with new thermo-mechanical properties (Scheme 1). The utility of this new dynamic chemistry is exemplified by the facile manufacturing of a recyclable structural composite.

Overall, this work provides a new tool for the facile preparation of *N*,*S*-acetal compounds and recyclable polymer networks from CO₂. To date, there is no example of oxazolidone-based networks with intrinsic recyclability, and recent contributions on poly(oxazolidone)s were focused on linear polymer synthesis^{46–50}.

Results and discussion

Inspired by the "click" reaction of enamides with thiols in acidic conditions to provide N, S-acetals⁵¹, we aimed to investigate whether the enecarbamate structure within alkylidene oxazolidones **2** exhibits similar reactivity (Figure 1a). Our group has previously confirmed the viability of this chemistry for post-functionalizing linear polymers containing alkylidene oxazolidone moieties⁴⁹. However, a comprehensive understanding of the reaction at the molecular level remains unexplored. Intrigued by the potential of this reaction to deliver novel N, S-acetal oxazolidone scaffolds and the presumed dynamic nature of the N, S-acetal bond, we conducted initial studies at the model molecule level before implementing this chemistry to macromolecular engineering.

Model reactions

Alkylidene oxazolidones $\mathbf{2}$ can be obtained via the spontaneous and quantitative reaction between primary amines and CO₂-sourced α -alkylidene cyclic carbonates α CC. This reaction yields hydroxyoxazolidones $\mathbf{1}$, which are subsequently dehydrated into the desired alkylidene oxazolidones $\mathbf{2}$ (Figure 1a). Previous works have introduced this last step^{48,52}, but existing approaches led to a mixture of $\mathbf{1}$ and $\mathbf{2}$. To address this limitation, we developed a novel optimized protocol for the synthesis of $\mathbf{2}$ at high yields, ensuring both quantitative reaction and potential scalability.

In the frame of this study, we started from a cheap and abundantly available propargyl alcohol, 2-methylbut-3-yn-2-ol, that produced the corresponding α -alkylidene cyclic carbonate αCC by the room temperature reaction with CO_2 in the presence of a low catalyst loading (Ag₂CO₃/PPh₃; 1 mol%). After 14 h of reaction at 40 bar, 126 g of αCC was obtained in a single batch with a high isolated yield (76 %). The quantitative dehydration of 1 was then realized with methanesulfonic acid (MSA, 0.1 eq.) in refluxing continuously dried acetonitrile (Figure S1), and 2 was collected with a high isolated yield of 77-83% after purification. For the purpose of this work, two compounds were easily produced from two structurally different primary amines, i.e. propylamine (product 2a) and benzylamine (product 2b) (Figure 1b).

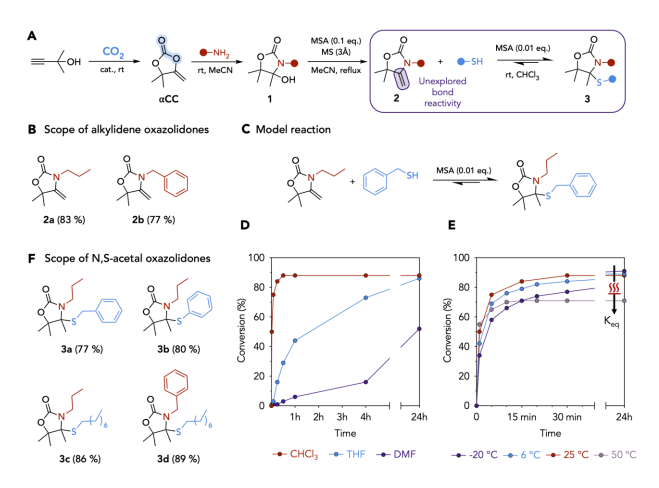


Figure 1. Overview of the synthetic pathway toward N,S-acetal oxazolidones and model reactions. (A) Carboxylation of 2-methylbut-3-yn-2-ol into reactive αCC , followed by its aminolysis into hydroxyoxazolidone 1. Dehydration can be triggered to yield alkylidene oxazolidone 2. N,S-acetal oxazolidones 3 are then obtained by the cationic addition of the thiol onto 2. (B) Scope of synthesized alkylidene oxazolidones 2 (isolated yields in brackets). (C) Model reaction between 2a and benzyl mercaptan to study the effect of (D) solvent and (E) temperature on the reaction kinetics. (F) Scope of synthesized alkylidene oxazolidones 2 (isolated yields in brackets).

With compounds **2** in our hands, we initiated our investigation by subjecting them to reaction with equimolar contents of thiols to yield *N*,*S*-acetal oxazolidones **3**. To assess how solvent and temperature influenced the reaction, we conducted reaction kinetics using the model oxazolidone **2a** and a commercial thiol, benzyl mercaptan, in the presence of MSA as acid catalyst (0.01 eq. compared to the oxazolidone), which has shown to efficiently catalyze the cationic thiol-ene reaction⁵¹ (Figure 1d, Figures S2-S5).

Chloroform was the ideal solvent for the reaction, which ended after nearly 15 minutes with the formation of **3a** (Figure 1e). Importantly, the reaction never reached completion and a conversion of 88 % at equilibrium was obtained at 25 °C, in contrast to previously studied enamides which quantitatively provided the product⁵¹. Solvents with a higher Lewis base character⁵³ significantly decreased the reaction rate. In THF, a similar conversion required over 4h of reaction and, in DMF, the rection remained uncomplete after two days. Our findings indicate that solvents with a higher base nature tend to act as proton traps, affecting the alkylidene protonation efficiency, in accordance with the previous work of Kloxin et al⁵¹.

Conversions were lower at higher temperature as the dissociated state was entropically favored (71 % at 50 °C) but were slightly increased at a lower temperature (89 % at 6 °C and 91 % at -20 °C) at the cost of longer reaction times (Figure 1f). This suggested that the conversion was dictated by the thermodynamic equilibrium between the associated and the dissociated state, the equilibrium constant being affected by temperature changes.

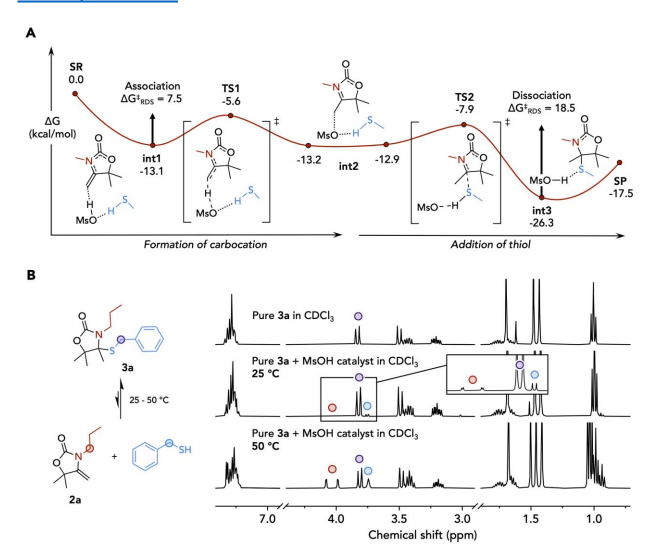


Figure 2. Energetics of the reaction pathway modelled by DFT and dissociative nature of the *N*,*S*-acetal oxazolidone moiety. (A) Gibbs-Free Energy profile with intermediates and transition states structures for the reaction of a model alkylidene oxazolidone with a model thiol. Vertical arrows indicate the energy barriers for both the association and the dissociation reactions at their rate-determining step (RDS). (B) Dissociation experiment of 3a in the presence of MSA in deuterated chloroform (CDCl₃) analyzed by ¹H-NMR spectroscopy at different temperatures.

We probed the reactivity of **2a** with various thiols, i.e. octanethiol, benzyl mercaptan and thiophenol (Figure S6a). Octanethiol behaved similarly to benzyl mercaptan, with a slight increase of the reaction rate, certainly due to a decrease in steric hindrance around the thiol group. Thiophenol reacted rapidly, however its conversion remained low at

equilibrium (73 %). Using the more hindered oxazolidone **2b** with octanethiol did not seem to significantly affect the reaction rate (Figure S6b).

The potential of this reaction can be leveraged by choosing ideal conditions (25 °C, excess of thiol) to synthesize novel *N*,*S*-acetal oxazolidone compounds **3a-d** (Figure 1c) with high isolated yields (up to 89 %). The full characterization of these molecules by NMR spectroscopy and HRMS can be found in Supporting Information (Figures S7-S14). This chemistry therefore opens an avenue to the facile construction of a vast library of compounds whose oxazolidone motifs are of high interest in many fields of chemistry (pharmacology, agriculture, synthetic intermediates, etc.). ^{54–57}

To have a better insight on the mechanism of this reaction, we performed density functional theory (DFT) calculations at the ω B97-XD/6-311++G(d,p) level of theory with a CPCM solvation model (ε =4.7113, chloroform) (Figure 2a). For simplicity's sake, the system was modelled with methyl groups as substituents for the N and S atoms, mimicking aliphatic chains used in the experimental model reactions. As it might have a drastic impact on calculations, MSA was used as catalyst in the system. Our results revealed that the reaction is performed in two separate steps. The starting reactants complex is very stable compared to separate reactants with a stabilization energy of 13.1 kcal·mol⁻¹. This is enabled by hydrogen bonding interactions between the different species and more especially due to the acidic proton of MSA in interaction with the negatively charged atom of the alkene (Figure S17). This interaction creates a high degree of delocalization in the oxazolidone ring thanks to the presence of the electronrich nitrogen group in the vicinity of both the alkene and the carbonyl groups. The first step of the reaction, which is the rate-determining step (RDS) of the forward reaction, has a low activation barrier of 7.5 kcal·mol⁻¹ explaining the experimentally observed high reaction rate. In this first step, the acidic proton of MSA is transferred to the terminal side of the alkene to form a stabilized carbocation, which is of very similar energy than the reactants complex int1 (difference of 0.1 kcal·mol-1). The high stabilization of this intermediate int2 can be explained by the formation of a new stable pi bond between the nitrogen and the carbocation, resulting in a decrease of bond length from 1.373 to 1.292 Å (- 0.081 Å). The N-C(O) bond readily increases its length of 0.073 Å as the nitrogen atom shares less electrons with the carbonyl group (Table S1). In a second time, the thiol adds onto the electrophilic carbon of the ring with a very low barrier of 5.0 kcal·mol-1, thus breaking the pi bond created in the first step. In a concerted manner, the hydrogen of the thiol already in interaction with the catalyst is then completely transferred to form a stabilized complex of the product and the regenerated MSA int3, whose energy is -26.3 kcal·mol⁻¹. This enhanced product stability renders the reverse reaction less energetically favorable though the reverse barrier stays rather low with an activation energy of 18.5 kcal·mol⁻¹. These differences in energy support why the

equilibrium was shifted towards the product under ambient reaction conditions, with only slight reversibility occurring at room temperature.

To further support that dissociation occurred under ambient conditions, to pure oxazolidone $\bf 3a$ was added by MSA (0.01 eq.) in CDCl₃ and the mixture was analyzed by ¹H-NMR spectroscopy (Figure 2b). Besides *N*,*S*-acetal oxazolidone $\bf 3a$, we observed the formation of the alkylidene oxazolidone $\bf 2a$ and the thiol in low amounts (\approx 3 mol%). Without MSA, only compound $\bf 3a$ was observed. This result provided strong evidence that dissociation slightly occurred in the presence of MSA at room temperature. When increasing the temperature to 50 °C, a great effect on dissociation was observed with the formation of $\bf 2a$ and the thiol in large amount (\approx 25 mol%).

The exchange kinetics was monitored at different temperatures using ¹H-NMR spectroscopy by reacting 3a with a fivefold excess of octanethiol in the presence of MSA (0.01 eq. vs 3a) (Figures 3a, S18). The reaction was monitored from 7 to 50 °C, showing slow exchange kinetics at 7 °C and strong acceleration of the reaction rate constant k at 50 °C, passing from 0.5×10⁻³ to 23×10⁻³ s⁻¹. The exchange reaction therefore showed a great temperature dependance. An Arrhenius plot was built from the different reaction rate constants k at different temperatures and an activation energy of 15.5 kcal.mol⁻¹ was extracted for the bond exchange (Figure 3b-c), lying in the same range of S,Sacetal or vinylogous urethane bonds^{42,58} (Table S2). This energy is also of same order as the dissociation energy determined by DFT (18.5 kcal.mol⁻¹). It must be noted that even in the presence of a high excess of thiol, the alkylidene oxazolidone 2a could still be observed in traces amount by NMR during the reaction kinetics, further proving the dissociative mechanism (Figure S18). Overall, the alkylidene oxazolidone - thiol dynamic chemistry showed activity at room temperature with strong increase in bond dynamics when the temperature was increased, highlighting a high reversible character in mild conditions.

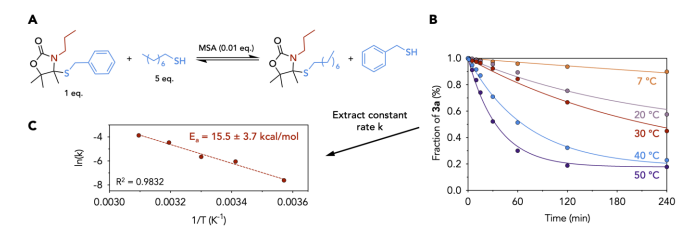


Figure 3. Exchange reaction kinetics at the model compound level. (A) Model reaction for the exchange kinetic study. (B) Kinetics of exchange at several temperatures (from 7 to 50 °C). (C) Arrhenius plot from the extracted reaction rate constants at the different temperatures of reaction.

Thermoset preparation and characterization

Taking advantage of the highly reversible character of this reaction, we explored its potential for the design of covalent adaptable networks based on N,S-acetal bonds. We exploited the versatile and facile α -alkylidene cyclic carbonate chemistry to design bifunctional alkylidene oxazolidone monomers of different structures (Figure 4a). Four compounds 2c-2f were synthesized from different commercially available diamines: mxvlvlenediamine 2,2'-(Ethylenedioxy)bis(ethylamine) (XDA). hexamethylenediamine (HMDA) and Priamine 1075 (PRI). This scope of diamines was chosen to access structurally diverse monomers, which are expected to impart networks with diamine-dependent properties. The synthesis of all four monomers was achieved by reacting CO₂-sourced αCC with a diamine in acetonitrile at r.T. Then, tandem dehydration was triggered without isolation of the bis(hydroxyoxazolidone) intermediate using the previously described dehydration setup. All substrates were easily purified by a chromatography-free process in high isolated yield (up to 84 %), rendering their synthesis facile, cheap, and up-scalable (see supporting information for details and a full characterization of these molecules, Figures S20-S27).

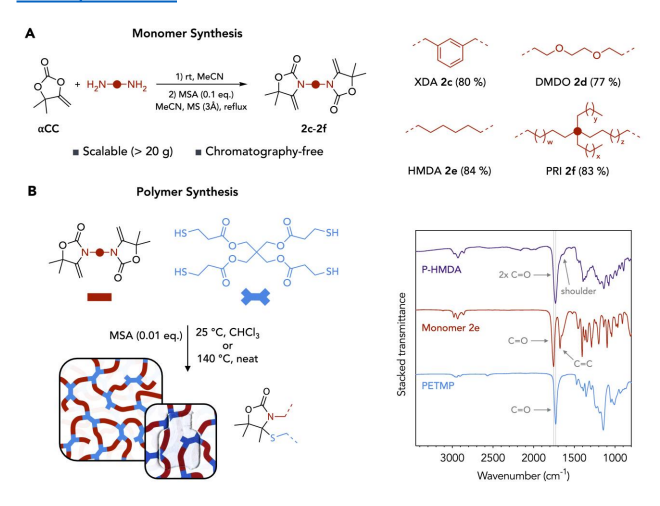


Figure 4. Monomer and polymer syntheses. (A) Tandem synthesis of bis(alkylidene oxazolidone)s 2c-2f from α CC and diamines (isolated yields in brackets). (B) Polymerization of a bis(alkylidene oxazolidone) with PETMP into a cross-linked material. A specimen of P-XDA is displayed to highlight its high transparency. ATR-IR spectra of P-HMDA and its respective monomers.

Different polymers were designed from the oxazolidone monomers **2c-f** and a 4-arm thiol, pentaerythritol tetrakis(3-mercaptopropionate) (PETMP) (Figure 4b). For all these syntheses, an equimolar ratio between the oxazolidone and the thiol functions was used, and the reactions were catalyzed by MSA (0.01 eq. *vs* oxazolidone monomer). The polymerization proceeded either in solution or in bulk (Figure S28). When achieved in solution, the mixture of the monomers in chloroform (2 M concentration of reactive functions) was stirred at r.T. until the formation of a gel (between 10 and 60 minutes depending on the monomer). The gel was then recovered and dried under vacuum at 70 °C followed by slow cooling to yield the pure material. Another option, which might be more easily adopted for large-scale production, was the bulk polymerization by stirring the components in the melt at 140 °C for 5 minutes followed by slow cooling to directly

yield the pure material. In the two processes, the polymer networks were transparent with a special emphasis for P-XDA showing no coloration (Figure 4b).

Successful polymerization was confirmed via infrared (FT-IR) spectroscopy (Figure 4b). The networks were characterized by an intense elongation at 1732 cm⁻¹ contributing for the stretching of the carbonyl groups of both the oxazolidone and ester groups. A very low intensity signal around 1620 cm⁻¹ was indicative of some residual alkylidene function (Figures S29-30). This is in line with the previous model reaction study that revealed a non-quantitative conversion of the reactants. Nevertheless, high gel contents were obtained for all polymers in THF (from 87 to 99 %), attesting for the crosslinked nature of the materials (Table 1). A good swelling (between 140 and 314%) was observed for all polymers in THF (Table 1).

The thermal properties of the polymers were then studied by thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC) (Table 1, Figures S31-38). All polymers displayed a temperature at 10 % of degradation ($T_{deg,10\%}$) around 300 °C except for P-PRI characterized by a higher value of 366 °C. The highest T_g was recorded for the aromatic P-XDA ($T_g = 60$ °C) with a more rigid backbone. On the other side, P-PRI picturing more chain mobility was characterized by the lowest T_g (8 °C). P-DMDO and P-HMDA had intermediate T_g s of 51 and 44 °C, respectively.

Table 1. Thermal properties, swelling ratios, gel contents and mechanical properties of the polymers.

Polymer	T _{deg,10%} ^a (%)	Tg ^b (°C)	SR _{THF} ^c (%)	GC _{THF} ^d (%)	Young modulus ^e (MPa)	Yield Strength ^e (MPa)	Stress at breake (MPa)	Strain at break ^e (%)
P-XDA	296	60	196 ±	90.4 ± 0.6				
P-DMDO	303	51	140 ±	99.0 ± 0.3	2438 ± 38	f	46.5 ± 1.2	2.7 ± 0.1
P-HMDA	282	44	215 ± 5	94.8 ± 0.4	1778 ± 116	f	12.6 ± 1.4	0.7 ± 0.1
P-PRI	366	8	287 ± 21	86.6 ± 1.3	2.9 ± 0.1	f	2.2 ± 0.1	159 ±
P-(HMDA75)- (PRI25)	289	44	228 ±	93.8 ± 0.2	1851 ± 41	f	34.0 ± 1.2	2.1 ± 0.1
P-(HMDA50)- (PRI50)	291	37	205 ± 6	91.2 ± 0.2	655 ± 72	8.6 ± 0.3	13.0 ± 0.3	81 ± 0.3
P-(XDA75)- (PRI25)	289	62	314 ±	86.1 ± 0.1				
P-(XDA50)- (PRI50)	310	43	292 ± 12	93.7 ± 0.4	1238 ± 45	29.9 ± 1.0	20.8 ± 0.8	10 ± 1

^aTemperature at 10% of degradation determined by TGA. ^bGlass transition temperature determined from the reversing curve of the first heating rate by modulated DSC. ^cSwelling ratio (SR) after immersion of the polymer for 24h in solvent (THF). ^dGel content (GC) after immersion of the polymer for 24h in solvent (THF). ^eMechanical properties evaluated by tensile tests. ^fMaterials experienced a brittle fracture.

To extend the scope of materials and modulate their thermo-mechanical properties, four additional polymers were synthesized by mixing pre-defined ratios of XDA or HMDA with PRI. The new polymers were labeled P-(A X)-(B Y) to denote their content in X wt% of monomer A and Y wt% of monomer B. Their temperature of degradation was quite similar, around 300 °C. We expected intermediate $T_{\rm g}$ for these polymers. Surprisingly,

P(HMDA75)-(PRI25) was characterized by the same T_g as P-HMDA (44 °C), and P(XDA75)-(PRI25) was characterized by a similar T_g as P-XDA (62 °C and 60 °C, respectively). By increasing the PRI content to 50 wt% within the materials, their T_g dropped while remaining around 40 °C. Overall, the thermal properties of the networks were judiciously adjusted through the choice of the diamine chemical structure.

Next, the mechanical properties of the various polyoxazolidones thermosets were determined through tensile tests on specimens obtained by compression molding of the samples (Figure 5a). Unfortunately, P-XDA and P-(XDA75)-(PRI25) samples could not be tested due to their extreme brittleness. P-HMDA was a brittle material too, breaking at very low strain (0.7 %). Despite its similar T_g lying between P-XDA and P-HMDA, P-DMDO showed excellent mechanical properties with a Young modulus of 2400 MPa and a yield strength of 46 MPa, with a brittle fracture occurring at low strain (2.7 %). The extra toughness of this polymer compared to P-HMDA was attributed to the higher degree of mobility provided by the presence of the ether bonds within the polymer microstructure. On the other hand, P-PRI demonstrated elastomeric behavior, with a relatively low modulus of 2.9 MPa and an elongation at break of 159%. Interestingly, when PRI was added to the formulation of brittle P-XDA and P-HMDA, the resulting polymer's mechanical behavior was significantly altered. Specifically, P-(HMDA75)-(PRI25) exhibited a modulus of 1850 MPa and a brittle fracture, while increasing the PRI content in P(HMDA50)-(PRI50) decreased the modulus to 655 MPa and resulted in high plastic deformation with an elongation at break of 81%. P(XDA50)-(PRI50) exhibited a modulus of 1240 MPa and plastic deformation.

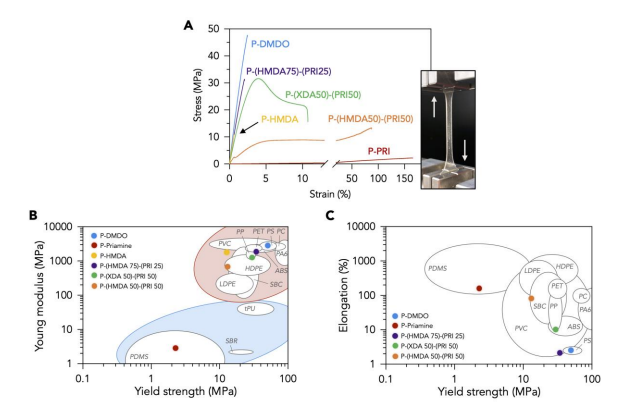


Figure 5. Mechanical properties of the polymers determined by tensile tests and comparison to commodity polymers. (A) Stress – strain curves of all polymers displaying a large range of properties. (B) Ashby plot showing Young modulus and yield strength compared to commodity polymers. (C) Similar Ashby plot showing elongation and yield strength. Commodity polymers mechanical properties were found in the literature⁵⁹.

To compare the properties of the synthesized polymers to commodity polymers, we plotted Ashby diagrams⁶⁰ based on three critical parameters: the Young's modulus, the yield strength, and the elongation at break (Figure 5b-c). These plots revealed a wide diversity in the mechanical properties of the synthesized polymers, with some of them exhibiting properties comparable to some commodity polymers. For example, P-DMDO had similar properties to polystyrene (PS), with a modulus, strength, and elongation within the literature values. Furthermore, P-PRI was located in the PDMS area of the diagram, while P(XDA50)-(PRI50) was situated in the polypropylene (PP) zone, albeit with low elongation. Despite P(HMDA75)-(PRI25) and P(HMDA50)-(PRI50) exhibiting intriguing properties, they did not meet those of the reported commodity polymers.

These results illustrate the huge potential of our new polymers to furnish materials of tunable mechanical properties by simply adapting the alkylidene oxazolidone monomer structure. Moreover, this study only represents a limited overview of the myriad of possibilities through monomer design and formulation to develop a wide range of

materials with on-demand modular properties to fit with the specification of diverse applications.

Dynamic behavior of the networks

The dynamic behavior of the networks was then characterized by rheology and exemplified for P-HMDA in Figure 6. First, the temperature-dependence on the storage (G') and loss modulus (G") was assessed by a temperature sweep experiment (Figure 6a). Values below T_g could not be determined by rheology as the sample was hard, resulting in a poor surface contact between the rheometer plateau and the specimen 61 . They were obtained by dynamic mechanical thermal analysis (DMTA) from rt to $80~^{\circ}$ C, showing a drop in storage modulus at the T_g of the polymer (Figure S39). Above T_g , the sample showed a higher value of G' than G", indicating that the material was in a crosslinked rubber state. Importantly, G' showed a constant decreasing trend, attesting for the progressive loss of cross-linking density with increasing temperature. This result was in line with the dissociative nature of the bonds as observed in the model compounds study. The curves tend to cross above $80~^{\circ}$ C where enough dissociation occurred to decrosslink the material and enable flow. The crossover point between the two curves represents the gel-to-sol transition and was denoted $T_{gel-to-sol}$ for the different polymers (Table 2).

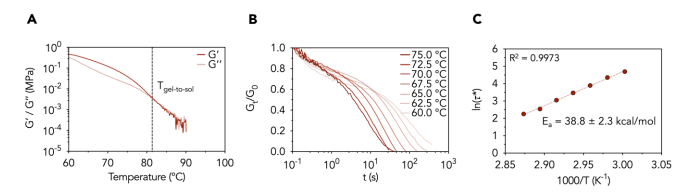


Figure 6. Rheological experiments conducted on P-HMDA. (A) Temperature sweep. (B) Stress relaxation at different temperatures whose extracted characteristic relaxation times could fit the (C) Arrhenius law to obtain an energy of activation.

Similar trends were observed for P-DMDO and P-PRI, each of them showing different T_{gel-to-sol} transition temperatures (Figure S40-S42). P-DMDO presented a higher transition of 91 °C and P-PRI a lower one of 73 °C. As P-PRI was characterized by a low T_g, rheological data were recorded from room temperature. Interestingly, the value of G' was slowly decreasing from 30 to 55 °C, indicating a slight loss of cross-link density at

low temperature. An abrupt drop was observed starting around 60 °C until the gel-to-sol transition at 72.6 °C.

Table 2. Gel-to-sol transition and flow activation energy for the polymers studied by rheology.

Polymer	T _{gel-to-sol} ^a (°C)	E _{flow} ^b (kcal·mol ⁻¹)
P-XDA	0	c
P-DMDO	91.3	26.4 ± 3.0
P-HMDA	81.4	38.8 ± 2.3
P-PRI	72.6	34.5 ± 2.8

^a Gel-to-sol transition temperature as determined by temperature sweep rheology. ^b Flow activation energy (E_{flow}) as determined by stress relaxation measurements in rheology. ^c Could not be determined.

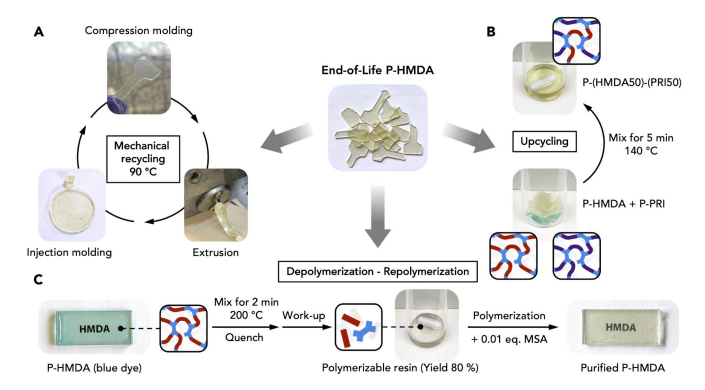
Stress-relaxation experiments were achieved before the flow transition threshold in the viscoelastic region of the networks. As observed in Figure 6b for P-HMDA, the material readily relaxed the applied stress through bond dynamics. The normalized stress relaxation curves have shown to follow a three-element Maxwell model, revealing the presence of three distinct relaxation modes during the material topology reorganization. However, most studies on CANs revealed that single-element Maxwell or stretched exponential models were generally accurately fitting the relaxation experiments data¹². On the other side, multi-element Maxwell could be effectively used to describe materials with kinetically different exchange processes^{62,63}. At each temperature were extracted three characteristic relaxation times (τ_1 , τ_2 and τ_3). τ_1 and τ_2 showed to be independent from temperature changes and were both of relatively short timescale (0.1-0.7 s for τ_1 and 0.7-15 s for τ_2) (Figures S40-S42; Table S3). Thus, we hypothesized these relaxation modes to be attributed to chain rearrangement at stress exposition⁶⁴. Although these two first relaxation modes with short timescales may not interfere in the network reprocessability, the use of a three-element Maxwell model is critical to correctly fit and extract reliable data. The last relaxation mode of characteristic relaxation time τ_3 was characterized by a strong temperature dependence and may be attributed to bond dynamics within the network. The different characteristic relaxation times τ_3 were plotted against temperature in a typical Arrhenius plot and an excellent linear correlation was observed, thus enabling the determination of a flow energy activation Ea,flow of 38.8 kcal·mol⁻¹ (Figure 6c). Stress relaxation experiments on P-DMDO and P-PRI showed

that the polymer matrix and thereby, the monomer structure, had a strong influence on the viscoelastic behavior of the polymer (Table 2). P-DMDO was characterized by a lower E_{a.flow} of 26.4 kcal·mol⁻¹, indicating a less pronounced sensitivity of the viscosity to temperature 14,65 . P-DMDO was indeed imparted with a shorter relaxation time τ_3 at 60 °C (88 s) but higher at 75 °C (15 s). This result was in line with the temperature sweep experiment displaying a higher T_{gel-to-sol} transition for P-DMDO than P-HMDA despite their close T_g. On the other side, P-PRI was characterized by a lower but yet similar E_{a,flow} as P-HMDA, of 34.5 kcal·mol⁻¹. The early gel-to-sol transition observed in P-PRI was rationalized by an increased chain mobility. However, matrix effects on CANs are yet still not fully understood and many parameters are thought to play a role such as polarity changes⁶⁴, solvation effects, cross-linking density and degree of chain flexibility⁶⁶. Overall, monomer structure seems to have critical influence on the viscoelastic properties of N,S-acetal oxazolidone dynamic networks and a better rationalization of the experimental results might be needed to further understand all the structural effects. This study is however out of scope of this paper. The determined flow energy activation for the different studied polymers are in the same range of reported S,S-thioacetal dynamic networks reported by the group of Du Prez⁴² (Table S2).

Overall, this rheological study suggests a high reprocessing ability for these materials which are able to relax extremely rapidly (relaxation times in the order of seconds) in their rubber state and can even flow as thermoplastics above their gel-to-sol transition temperature where the cross-linking integrity is lost.

Recycling of the networks

While most studies on CANs materials mainly display one or two recycling routes, this work exploited the highly reversible character of the alkylidene oxazolidone-thiol chemistry to provide three different end-of-life scenarios for the materials (Scheme 2). Multiplying the recycling possibilities for new emerging plastics is of great interest as the recycling approach might be selected depending on the market needs and the state of the material after use.



Scheme 2. End-of-life scenarios for P-HMDA. (A) Mechanical recycling using compression molding, extrusion, and injection molding. (B) Upcycling of P-HMDA and P-PRI toward P-(HMDA50)-(PRI50). (C) Chemical recycling through depolymerization and re-polymerization of contaminated P-HMDA.

Mechanical recycling

The most straightforward approach is the mechanical recycling (Scheme 2a). Although conventional networks cannot be reshaped, stress can be dissipated in CANs by virtue of bond dynamics and reshaping is therefore made possible. As highlighted by rheological investigations, the polymers reached very low complex viscosities η^{\star} (in the range of 10 to 40 Pa·s at 90 °C) (Figure S43) and flowed at temperatures above the gelto-sol transition. With this in mind, we first reprocessed P-HMDA by compression molding at 90 °C (in the flowing state) for 10 min under mild pressure (1 ton metric). Upon cooling, the material was recovered from the molds. We also successfully subjected P-HMDA to extrusion at high rate (150 rpm) or injection molding at 90 °C. Reshaping the network was thus possible through multiple industrially relevant techniques in mild conditions and in a short timeframe while most studied CANs require long exposition to high pressure and/or temperature $^{30,32-34}$. In all cases, T_g as well as IR spectrum of the reprocessed polymer were identical to the starting material, suggesting no degradation (Figure S44).

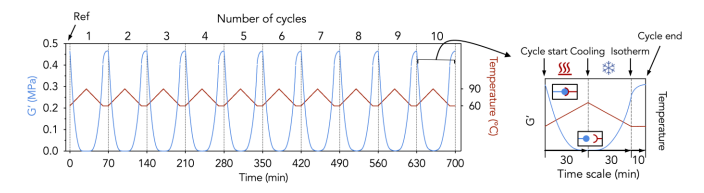


Figure 7. A temperature sweep cycling test was performed on P-HMDA at heating and cooling rates of 1 °C/min between 60 and 90 °C. The storage modulus G' was totally recovered (at 100%) at the end of each cycle.

We performed on-line storage modulus measurement as a function of time and temperature over up to 10 cycles (Figure 7). The sample, heated from 60 to 90 °C and cooled down to 60 °C at each cycle, displayed total recovery of the storage modulus. It must be noted that after cooling at 1 °C/min, around 85 % of the storage modulus was recovered. Additional 10 minutes at 60 °C allowed for a total recovery of G' before beginning a new cycle, highlighting that slow cooling was critical to recover the starting material properties. This experiment evidenced that the network underwent total disassembly to a flowing state at 90 °C by cycling above and below the gel-to-sol transition temperature with a total and reproducible recovery of the storage modulus, i.e. of the cross-link density within the material, without any hysteresis. This suggested that these materials might undergo a significant number of fast reprocessing cycles at mild temperatures, well below the degradation of the polymer, thus avoiding material degradation that might happen in CANs exposed to high pressure and temperatures for a prolonged period of time¹⁴. Tensile tests of reprocessed samples after fracture were performed and showed a good recovery of the Young's modulus which was very similar to the original material after two fracture-reprocessing cycles (Figure S45). However, both stress and strain at break were not fully recovered, as already observed for many healed materials containing dynamic linkages^{35,67–70}.

Upcycling

Recently, it was demonstrated that upcycling of CANs was possible through bond exchange between two distinct materials into a divergent one^{71,72}. Dichtel et al. notably demonstrated that two soft and hard polyurethane networks could be mixed enough efficiently to provide a new material by extrusion at 200 °C. An akin strategy can be used in our case by mixing two different *N*,*S*-acetal oxazolidone dissociative CANs. Upon thermal treatment above the gel-to-sol transition, bond rupture enabled the

material to flow, delivering back alkylidene oxazolidone and thiol functionalities and, upon cooling, random re-association of the bonds was expected to provide a hybrid network consisting of mixed structures (Scheme 2b). This was illustrated by efficiently mixing identical contents of two different materials, P-HMDA and P-PRI, at 140 °C for 5 min under solvent-free conditions using a magnetic stirrer, and then to slowly cool down the mixture. DSC and IR analyses evidenced identical T_g and IR spectra than the previously synthesized P(HMDA50)-(PRI50), suggesting that the mixture of the two separate networks were converted into the mixed one (Figure S46). This approach enabled to easily upcycle the end-of-life networks into new ones exhibiting distinct properties.

Chemical recycling

The chemical recycling is the third considered end-of-life option (Scheme 2c). This approach, especially the one that considers the depolymerization into the constitutive monomers, is highly desirable as it generally allows to recover the monomers from impurities such as dyes and other additives^{8,73}.

Acidic hydrolysis of acetal linkages is a common and attractive way to promote degradation and recycling of polymers in mild conditions^{31,41,74}. Thus, the presence of the *N*,*S*-acetal linkages could render poly(oxazolidone)s scaffolds sensitive to acidic aqueous environments. However, *N*,*S*-acetals derivatives might be more resistant than oxygen-containing *S*,*O*- and *O*,*O*-acetals⁵¹. Low swelling ratios were observed in water after 24h for all synthesized polymers, the highest value being 2% for P-DMDO. Lowering the pH using acidic aqueous solutions of H₂SO₄ at different concentrations (0.1, 1.0 and 5.0 M) did not lead to any degradation with gel contents as high as 99% and IR spectra showed no difference compared to the pristine polymer (Figure S47), attesting for the resistance of the polymers to acidic aqueous environment.

However, the versatility of the *N*,*S*-acetal oxazolidone chemistry and its high dissociative character at high temperatures allowed us to find another approach toward a recovery of the constitutive monomers. We synthesized P-HMDA containing a blue dye (methylene blue) as a model additive. In order to shift the equilibrium toward the monomers and thus to favor depolymerization, we heated the network at 200 °C for 2 minutes only. We then quenched the medium with a catalytic amount of triethylamine (3 eq. per equivalent of MSA) to avoid the MSA-catalyzed reverse reaction, quantitatively yielding a mixture of constitutive monomers and quenched catalyst (Figure S48). The mixture was purified by filtration through activated charcoal to remove the dye followed by a simple liquid extraction with water to remove triethylamine and MSA. After evaporation of the solvent, a colorless viscous resin was obtained. ¹H-NMR analysis of the resin revealed the presence of the two starting monomers in equimolar amount that were recovered at 80

% yield (Figure S48). By adding MSA to this mixture and following the synthetic procedure, P-HMDA was recovered with identical DSC and IR results as the starting product (Figure S49).

Composite preparation and recycling

Thermosets are largely used in composites to furnish high performance light structural materials used for instance in wind turbine blades, aeronautics, automotive, etc. ⁷⁵ However, the recycling of composites is an important issue due to the difficulty to separate the filler from the crosslinked polymer matrix. For the proof of concept to illustrate the potential of our technology in composite preparation and in their facile recycling, we prepared a composite composed of flax as the filler and P-DMDO as the network matrix. For that purpose, P-DMDO was hot pressed on flax under 3 bars of pressure at 100 °C for 15 minutes (Figure 8). This straightforward process was aided by the very low viscosity of the polymer above its gel-to-sol transition temperature, rendering the filler impregnation a facile task. As expected, the composite tensile strength was strongly increased compared to the lone polymer (170 vs 50 MPa) as well as the Young modulus (5700 vs 2400 MPa) (Figure 8). The strain at break passed from 2.7 % to 3.5 %.

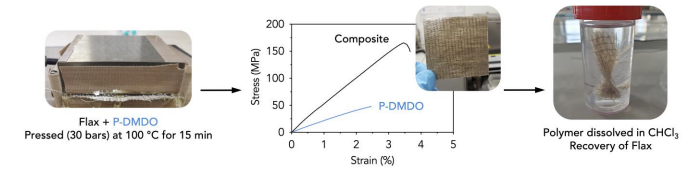


Figure 8. Composite preparation from flax and P-DMDO. Grinded P-DMDO was pressed with Flax to provide a composite material with excellent mechanical properties. Polymer-free flax could be recovered by dissociation of the polymer matrix in CHCl₃ (catalyzed by MSA within the polymer matrix).

To separate the fibers from the crosslinked matrix, we exploited the dynamics of the N,S-acetal type oxazolidone linkages. Based on our model reaction reported in Figure 2b, the N,S-acetal linkages slowly dissociated in chloroform. By immersing the composite (containing MSA catalyst in the polymer matrix) in a large volume of chloroform to displace the equilibrium towards the dissociated state, P-DMDO was completely dissolved after one day at room temperature. While dissolution in organic solvent is generally unwanted for polymer networks and composites that should be

solvent resistant, *N*,*S*-acetal based polyoxazolidone networks were only soluble in a large amount of this specific solvent. This allowed the recovery of the filler when needed (Figure S50). Under identical conditions, no dissolution was noted in THF nor DMF, consistently with the model reactions that demonstrated extremely slow kinetics of reaction in these solvents. An absence of dissolution was also noted in other solvents, therefore proving dissolution of these polymers is selective (Table S4).

Conclusion

In this work, the alkylidene oxazolidone – thiol chemistry that reversibly delivered N,Sacetal type oxazolidone scaffolds was studied at the model reaction level, attesting for fast kinetics in mild conditions with a great dynamic character. The synthesis of bifunctional alkylidene oxazolidone monomers was reported to be easy, affordable, and scalable starting from a readily available propargyl alcohol coupled with CO2 and a primary diamine. Different polymer networks were synthesized from structurally divergent monomers and a commercially available tetrathiol in the presence of a catalytic amount of methane sulfonic acid (MSA). Networks with a wide range of mechanical properties were prepared, ranging from PDMS-like elastomers to rigid polystyrene-like ones. The highly dynamic character of the covalent adaptable networks was highlighted by rheological study reflecting observations made at the model level. High dissociation rates can be accessible in relatively mild conditions, thus enabling fast reprocessing with thermoplastic-like techniques such as extrusion and injection molding without promoting any side reactions and material alteration. Through this high dissociative exchange capability, two divergent materials were also mixed and upcycled by fast dissociation and random re-association events to furnish a new scaffold with distinct properties. At high temperatures, total dissociation was obtained in minutes timeframe, vielding back the initial monomers that could be repolymerized into the same network. Lastly, this chemistry showed potential to be easily applied to composite materials with recycling possibilities. Further work will be devoted to diversify thermal and mechanical properties of the networks with the objective to broaden the application field of this very promising family of easily recyclable polymer networks. Overall, this study shows great potential for constructing a wide range of circular materials that might find applications across the numerous domains covered by polyurethanes, and well beyond as the dynamic linkages might be introduced in other thermosets for improved recyclability.

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Notes

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