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Neighboring Group Participation and Internal Catalysis Effects on Exchangeable Covalent Bonds: Application to the Thriving Field of Vitrimer Chemistry

Florian Cuminet, Sylvain Caillol, Éric Dantras, Éric Leclerc, and Vincent Ladmiral*



ABSTRACT: Vitrimers constitute a fascinating class of polymer materials that make the link between the historically opposed 3D networks (thermosets) and linear polymers (thermoplastics). Their chemical resistance, reshaping ability, and unique rheological behavior upon heating make them promising for future applications in industry. However, many vitrimers require the use of high catalyst loadings, which raises concerns for their durability and limits their potential applications. To cope with this issue, internal catalysis and neighboring group participation (NGP) can be used to enhance the reshaping ability of such materials. A few studies report the effect of activating groups on the exchange reactions in vitrimers. Nevertheless, knowledge on this topic remains scarce, although research on vitrimers would greatly benefit from NGP already known in organic chemistry. The present Perspective presents the different types of exchangeable bonds implemented in vitrimers and discusses chemical groups known to have or potentially capable of an enhancing effect on exchange reactions. This analysis is underpinned by a thorough mechanistic discussion of the various exchangeable bonds presented.

I. INTRODUCTION

Since the team of L. Leibler discovered and described the first vitrimer in 2011,¹ these materials have attracted tremendous attention in the polymer and materials communities. This enthusiasm is not surprising given the amazing properties of these materials. Nevertheless, though the definition of this concept seems easy to handle at first glance, caution should be taken when classifying new materials as vitrimers. To understand well what a vitrimer is, the usual classification of polymer materials must be clearly recalled. Historically, polymers have been categorized in two main classes: thermoplastics and thermosets. Thermoplastics consist of entangled linear polymer chains that are free one from another. In contrast, in thermosets, polymer chains are linked by covalent bonds that cannot be cleaved without destroying the network.² This difference at the molecular scale has consequences on the macroscopical behavior of these materials. Most thermoplastics are soluble provided that a

convenient solvent is used (or exists). In thermosets the solvent is not able to set the chains apart, and the material does not dissolve. Besides this different behavior toward solvents, the other main difference lies in the thermal behavior of these materials. In most cases when thermoplastics are heated, the relative movement of the chains becomes easier. Macroscopically this phenomenon is visible when the material flows.² Thermosets cannot flow when heated.^{2,3}

Reversible reactions involving covalent bonds have been known for many decades, and their possible use in polymers had already been inferred in the 1960s.^{4,5} The concept of

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reversible covalent bonds implemented in polymers led to a new class of materials called CANs, short for covalent adaptable networks. In CANs, polymer chains are covalently cross-linked as in thermosets, but these cross-linking bonds are reversibly cleavable, which leads to properties such as malleability in response to an external stimulus such as temperature or light irradiation.⁶ The most iconic reversible reaction used in such materials is probably the Diels–Alder reaction.^{7–13} This reaction is reversible upon heating. When this reaction is used to make cross-linked materials, the crosslinking bonds can break upon heating and re-form at lower temperature, thus leading to reshapability (Figure 1C).⁶ Most



Figure 1. Schematic representations of the mechanisms at work in (A) associative CANs (vitrimers), (B) vitrimer-like dissociative CANS, and (C) dissociative CANs. Vitrimer-like dissociative CANs (B) exhibit a constant K_{eq} over the vitrimer-like temperature regime and a short lifetime of the dissociated state.

CANs designed before the year 2010 were based on the dissociative mechanism whereby a bond has to be cleaved before another bond is formed between the dissociated moieties.

In 2005, Bowman et al. reported the first CAN based on an associative exchange reaction.¹⁴ Allyl sulfides were used as addition—fragmentation chain transfer agents. The exchange proceeded via a photoinduced radical mechanism allowing constant cross-linking density and network connectivity even though the topology changed.¹⁴

Later, in 2011, Leibler et al.¹ described the first thermally triggered associative CAN, based on transesterification as the

exchange reaction. This material exhibited a novel rheological behavior with its viscosity decreasing linearly with increasing temperature. Because this behavior is common in inorganic materials and especially strong glass, this new kind of polymer was coined "vitrimer". On the one hand, vitrimers are often insoluble like thermosets, except for specific cases such as very fast vitrimers for instance.^{15,16} On the other hand, they can be reshaped and recycled by heating without precise control over temperature, like glass. Their singular rheological behavior was attributed to the associative mechanism of the bond exchange. In contrast to the majority of dissociative CANs, because vitrimers keep a constant connectivity regardless of the temperature (even the reshaping temperatures), their viscosity only depends on the rate of the exchange reaction. Therefore, the viscosity and stress-relaxation profiles as a function of temperature follow the Arrhenius law. These behaviors are often considered specific to vitrimers,^{1,17-19} although this point has recently been questioned and is debatable.

Two visions of what a vitrimer is coexist, depending on whether the chemist's or the physicist's point of view is adopted. The chemist's definition focuses on the exchange mechanism. In consequence, only a CAN based on an associative mechanism is a vitrimer (Figure 1A). In contrast, the physicist defines a vitrimer according to the physical properties. For example, Drockenmuller et al.²⁰ described a material based on a dissociative mechanism that exhibits an Arrhenius viscosity dependence with temperature. Such materials were designated as "vitrimer-like".²¹ The hypothesis to explain such vitrimer-like behavior in dissociative CANs lies in the thermodynamic and kinetic of the exchanges. When the equilibrium constant of dissociative bonds is displaced toward the associated state, though the dissociation-association happens, the overall number of bonds and cross-link density is barely reduced. Kinetically, this happens when the association reaction is much faster than the dissociation reaction. The dissociated state is thus transitory, globally the number of bonds hardly changes, and the network integrity remains virtually unchanged (Figure 1B).^{22,23} The Arrhenian rheological behavior can also be observed in a range of temperatures over which the equilibrium constant does not vary much. For instance, Konkolewicz et al. made vitrimer-like dissociative CANs out of anilinium salts. They reported for the adduct formation a ΔH value of -2.8 ± 0.2 kJ mol⁻¹ and compared it with the case of Diels-Alder adduct formation with a ΔH value of -157 ± 14 kJ mol^{-1,21} Because ΔH is very low for their system, the dependence of K_{eq} with temperature is considered negligible. In this case, the ratio of dissociated and associated bonds over this temperature range is constant (Figure 1B), and the network connectivity does not change.^{21,24} With the increasing number of chemical platforms able to undergo exchange discovered, some particular cases of dissociative CANs somewhat blur the boundary with vitrimers.^{20,21,25,26} In addition, networks featuring both permanent and exchangeable cross-links are reported to exhibit vitrimer-like stress-relaxation, but within a certain limit as the permanent cross-links induced a residual stress plateau.²⁷ Such borderline materials indicate that the definition of a vitrimer still remains debatable, depending on the viewpoint adopted.

Vitrimers address the problem of thermoset recyclability and thermoplastic vulnerability toward solvents. Therefore, the interest for these materials is growing in industry, as highlighted in recent reviews.^{22,28,29} Nonetheless, some characteristics may limit their industrialization. They often



Figure 2. Associative exchangeable bonds for vitrimers' design.

Scheme 1. Ester Exchange Mechanisms: (a) Alcoholysis, (b) Acidolysis, and (c) Direct Ester-Ester Exchange



c. direct ester-ester exchange

display high relaxation times and high viscosity during reprocessing, making them unsuitable for the recycling processes usually found for thermoplastics in industry. Furthermore, most vitrimers rely on catalysts and in some cases on high catalyst loadings.¹ The use of catalysts may generate problems such as limited number of recycling/ reshaping cycles due to premature degradation or migration and loss of the catalyst.^{30–32} A solution to this issue is the design of vitrimers that do not rely on catalysis. This can be done by using a large excess of exchanging functional groups^{32–34} or by using exchange reactions that do not require any catalyst.^{18,35–42} Another strategy, inspired by organic chemistry in solutions, relies on chemical functions

near the exchangeable bond to enhance the reaction rate. This effect is called either neighboring group participation (NGP) when the accelerating substituents are at some point covalently bonded to the reaction center during the exchange or internal catalysis for effects at longer distance such as electrostatic interactions, steric effects, or weak bonds.⁴³ Guan et al.⁴⁴ implemented this strategy in boronic ester CANs and enhanced the transesterification rate thanks to neighboring tertiary amines. This effect was later implemented in other kinds of vitrimers featuring ester or silyl ether exchangeable bonds, for instance.^{38,45} In their review, Guerre et al.⁴³ highlighted the few examples of internal catalysis in vitrimers reported so far and the strong potential of this strategy to tune

Scheme 2. Exchange Reactions on Carbonates (a) Transcarbonation and (b) Carbonate Interchange



the properties of vitrimers. More recently, Van Lijsebetten et al.⁴⁶ also emphasized the power of NGP in dynamic covalent chemistry, especially in polymeric materials, in a very clear and useful tutorial review. To proceed further with this concept, a general outline of the activating groups potentially beneficial to vitrimers would give useful insight for further research on this fast rising topic. Nevertheless, caution should be taken when using NGP as this strategy can sometimes drastically modify the exchange mechanism.^{24,43} For instance, transesterification, which usually proceeds via an associative exchange, was reported to follow a dissociative pathway in the presence of specific activating groups.^{47,48}

Activating groups are a very powerful yet complex tool. Because the case of vitrimer-like dissociative CANs is still debated in the vitrimer community, this Perspective focuses on internal catalysis and NGP in associative exchange reactions, to follow the strict definition of vitrimers. The first part is dedicated to the description of the associative exchange reactions implemented in vitrimers, their mechanisms, and some of the pitfalls a vitrimer scientist should be aware of (Figure 2). Then, in light of this mechanistic discussion, the activating groups potentially useful to tune the properties of vitrimers are briefly presented and discussed. Insights into the activating groups proven to be beneficial on small molecules but not yet implemented in vitrimers are also given. For each kind of exchangeable bond the influence on the kinetics of exchange reactions is discussed as well.

II. EXCHANGEABLE BONDS WITH AN ASSOCIATIVE MECHANISM

Esters. Esters are probably one of the oldest known exchangeable bonds, as Friedel and Craft, famous for the eponymous reaction, also discovered in 1865 the exchange reaction occurring between two ester groups.⁴⁹ Several terms such as ester exchange, ester-ester interchange, interesterification or the nowadays most commonly used transesterification are considered as synonymous, though they actually encompass several reactions.⁵⁰ Actually, three reactions are reported to happen in what is commonly considered as the transesterification equilibrium,⁵¹ namely, alcoholysis, acidolysis, and the genuine transesterification. $^{49,52-54}$ Alcoholysis is the main reaction⁵⁵ although the direct ester-ester exchange prevails for esters synthesized from aromatic acids over 300 °C.56 The alcoholysis mechanism proceeds by the attack of the alcohol onto the carbon of the carboxylate. A tetrahedral intermediate is thus formed prior to the elimination of an alkoxy group, leading either to the starting ester or to the exchanged one (Scheme 1a).

The mechanism of acidolysis is similar: the nucleophilic oxygen of the acid attacks the electrophilic carbon of the carbonyl moiety of the ester and forms a tetrahedral

intermediate, prior to the elimination of the attacking acid, or the exchanged one. Elimination of the alkoxy group on the intermediate can also occur. This produces an anhydride intermediate which reacts immediately with the released alcohol (Scheme 1b).^{53,57} The third mechanism is less usual and is described as a concerted interchange (Scheme 1c). This reaction was reported experimentally thanks to kinetics experiment at temperatures over 300 °C.56 Kotliar et al. pointed out that the enthalpy change during this reaction is near zero, suggesting an associative pathway.⁵³ Anyway, these three mechanisms are associative. Catalysis does not seem to change this feature as acid-catalyzed transesterification is reported to proceed through an addition/elimination mechanism.⁵⁸ Additionally, base catalysis only changes the nucleophile from an alcohol to its more reactive alkoxide counterpart.⁵⁹ Transesterification was studied in polymers in the second half of the 20th century, as the exchange reaction is more significant in bulk materials;^{49,52,55,60} then this reaction regained popularity among scientists with the development of biodiesel and biofuels. Recently, the interest in biomass valorization and greener fuels has grown, and catalysts such as *N*-heterocyclic carbenes,⁶¹ calcium oxide,⁶² ZrO_2/SO_4 ,⁵⁹ silicates and clays,^{63,64} or aluminum chloride⁶⁵ were (re)discovered. Metal-based catalysis (such as Sn, Ti, or Mg⁵ for instance) draws particular attention as zinc-based catalysts were recently used in the first reported vitrimer.¹ The mechanism involves a Zn²⁺ species, which coordinates to the ester carbonyl group and activates it with respect to the nucleophilic attack of the alcohol or alkoxide and then proceeds via the pathway previously described.⁶⁶ In summary, all the catalytic mechanisms encountered so far were associative.

Carbonates. Owing to their structural similarity, carbonates and esters are often compared, and their chemistries are considered very close. As in the case of esters, several names are used for the exchange reactions of carbonate. The exchange between an alcohol and a carbonate is called transcarbonation, carbonate transesterification, or carbonate alcoholysis when the exchange occurs on polycarbonate chains and small alcohols (Scheme 2a). Carbonate interchange refers to the exchange between two carbonates (Scheme 2b).

The interest for transcarbonation rose when scientists aimed at synthesizing polycarbonate materials without the use of highly toxic phosgene.⁶⁷ The exchange between diphenyl carbonate and bisphenol A was used as a safer pathway to produce polycarbonates. The mechanism involved was similar to ester transesterification⁶⁸ and was proven quite recently. In the absence of catalyst, there is a prototropy between the phenol and the carbonate. The deprotonated counterpart (phenolate) is the reactive species toward the electrophilic carbon of the carbonate carbonyl moiety. The reaction Scheme 3. (a) Transcarbonation Pathway and (b) Catalytic Effect by Bases and Metal Ions



proceeds through a tetrahedral intermediate before the elimination of a phenolate (from the starting carbonate or from the reactant).^{69,70} However, this exchange is very slow.⁶⁷ Transcarbonation is known to be catalyzed by metal-derived catalysts such as tin salts,^{71–73} zinc acetate,⁷⁴ copper salts,⁷² and even a dimetallic iron–manganese cyanide catalyst.⁷⁵ Bases are also common catalysts, for instance, magnesium hydroxide,⁷⁶ calcium salts,^{77,78} or sodium alkoxides.^{74,79} All the catalyzed mechanisms described are associative,^{69,71,74,76–80} though they are slightly different depending on the type of catalysis employed (Scheme 3).

When a base catalyst is used, the carbonate and the alcohol moiety form a prereaction complex (CX1). Then the reaction proceeds via a tetrahedral transition state (TTS) yielding a postreaction complex (CX2) and ultimately the free exchanged species (Scheme 3b).⁷⁴ The overall reaction follows an "addition–elimination" mechanism. When a Lewis acid catalyst is employed, such as zinc acetate, a cyclic transition state involving a ternary complex between the carbonate, the alcohol, and the catalyst is formed. Then, the reaction proceeds through a concerted mechanism to generate a postreaction complex, following an addition–elimination mechanism.⁷⁴ Direct ester–carbonate interchange has also been reported⁸¹ and described as following an associative pathway as well.

Boronic Esters. Cyclic boronic esters, also known as dioxaborolanes, are synthesized by condensation of boronic acids with 1,2- and 1,3-diols leading to 5- and 6-membered rings, respectively.⁸²⁻⁸⁴ Boronic acids are used in a wide range of reactions in organic chemistry, as catalysts, pronucleophiles (cross-coupling reactions), or reaction intermediates.^{82,85} In carbohydrate chemistry, they are also useful as protective groups for diols and diamines.^{82,85,86} Besides, their ability to bond with diols makes boronic esters particularly useful in biology, biochemistry, or chromatography for sugars and diolbearing compounds purification or carbohydrate sensors, for example.^{85,87-89} In medicine they are also proposed for drug delivery⁹⁰⁻⁹² and glucose sensing for diagnostic purposes.^{93,} The key feature of boronic esters is their reversible formation from diols and boronic acids, resulting in dynamic systems with tunable properties. In biological media, this reactivity results in a dissociative mechanism, as boronic esters in aqueous conditions are prone to hydrolysis yielding boronic acids.^{95,96} The equilibrium between dissociated boronic aciddiol and associated boronic ester (Scheme 4) was exploited to prepare covalently cross-linked self-healing hydrogels, for example.97-99

Over the past few years, growing interest for this functionality has spread in the polymer science community, in particular for the preparation of self-healing polymers.^{100,101}

Scheme 4. Equilibrium between Boronic Acid and 1,2-Diol and Cyclic Boronic Ester

$$R_1 - B \stackrel{OH}{\longrightarrow} + H \stackrel{HO}{\longrightarrow} R_2 \stackrel{-H_2O}{\longrightarrow} R_1 - B \stackrel{O}{\longrightarrow} R_2$$

Interestingly, Sumerlin et al.¹⁰¹ showed that two different mechanisms are involved in the exchange reactions occurring in boronic ester-based cross-linked materials. The exchange occurs either via classical boronic ester dissociation and reesterification or via an associative mechanism similar to transesterification, provided that free diol functions are available in the material. As the dissociative mechanism requires water to cleave the boronic ester, it is assumed that in a completely dry environment only the associative transesterification mechanism occurs.^{85,102} Guan et al. implemented this associative mechanism in the first boronic ester cross-linked vitrimers.⁴⁴ Eventually, Nicolaÿ and Leibler et al.¹⁰³ reported a vitrimer based on a third kind of exchange reaction, namely, the direct metathesis between two boronic ester moieties. Although the exact mechanism of this reaction remains uncertain, the hypothesized transition states and intermediates are expected to have a higher connectivity,¹⁰⁴ as expected for an associative mechanism. Raynaud et al. investigated the mechanism at work in dynamic polymer networks based on pinacol boronates. The experimental and theoretical results corroborate the hypothesis of an associative mechanism as suggested and that the reaction is triggered by adventitious traces of nucleophiles trapped in the medium.¹⁰⁵ Nevertheless, mechanistic studies would be very useful to confirm the associative pathways of boronic ester exchange reactions in anhydrous conditions and to acquire the knowledge required to harness this type of exchange reaction more efficiently.

Sulfur-Containing Bonds. *Disulfides.* For a long time, disulfide bonds have been known to feature an interesting reversible behavior. Historically, this behavior has been used in personal care products. Disulfide bonds naturally occurring in keratin were cleaved by a reducing agent before new bonds were generated by using an oxidant to perform so-called "permanent wave" on hair.¹⁰⁶ This process can be considered as an early chemically reshapable material and was implemented after several decades in reversibly cross-linked polymers.^{107,108} Disulfide bonds are also involved in the rubber vulcanization process. Hence, in 1946, Stern and Tobolsky already discussed possible mechanisms for the reshuffling of these bonds to explain stress—relaxation in these material.¹⁰⁹

associative.¹⁰⁴ Two possible mechanisms have been discussed: (1) a [2 + 2] metathesis mechanism and (2) a radicalmediated [2 + 1] mechanism, where a S–S bond would be homolytically cleaved prior to a radical attack of another S–S bond to yield a three-membered intermediate (Scheme 5).

Scheme 5. Disulfide Exchange [2 + 1] Radical Mechanism



To elucidate which mechanism was involved. in silico studies were carried out and concluded that the radical mechanism was more likely to occur.¹¹⁰⁻¹¹² This conclusion was verified experimentally by model exchange reactions in the presence of radical traps or radical sources.¹¹³ However, when a nucleophilic catalyst is used, two mechanisms occur simultaneously: the radical mechanism described previously and a thiol-mediated exchange due to the formation of thiolate anions. Moreover, UV light can advantageously activate disulfide metathesis. Thus, materials exhibiting self-healing behavior upon 5 min exposure to 320-390 nm UV light were synthesized.¹¹⁴ Although this reaction is convenient to yield self-healing materials,¹¹⁴⁻¹¹⁶ its dissociative mechanism may not be suitable to obtain vitrimer properties. Additionally, disulfide metathesis is not always the only mechanism involved. Hence, depending on the synthetic pathways for disulfide-containing polymers and on the stoichiometry chosen, dangling thiol moieties enable thiol/disulfide exchanges. For this reaction the exchange proceeds through an associative mechanism, whereby a disulfide bond is attacked by a thiolate, yielding a new disulfide bond through a threemembered transition state (Scheme 6).^{111,117} Thus, attention should be paid to stoichiometry when designing a disulfidecontaining vitrimer or self-healing material.

Scheme 6. Thiol–Disulfide Exchange Mechanism



Thioesters. Recently, Bowman et al. reported thioester dynamic networks.¹¹⁸ Without neighboring groups the exchange takes place via a purely associative pathway. Nevertheless, when NGP were present, two competitive mechanisms were discussed: a mechanism involving an equilibrium between the associated thioester form and the dissociated thiol/anhydride form and the associative transthioesterification mechanism requiring free thiol moieties.

Meldrum's Acids. Sulfur-bearing Meldrum's acids were also described as exchangeable cross-linkers in PDMS vitrimers through an associative mechanism involving thiol moieties (Scheme 7). 119

Trialkylsulfonium Salts. In the 1970s, a group at Ghent got interested in poly(propylene sulfide) and the mechanism of its degradation upon incorporation of oxonium salts to introduce sulfonium groups in the material. They reported an exchange reaction between thioethers and trialkylsulfonium likely proceeding through an associative mechanism whereby the sulfur atom of the thioether attacks the α -carbon atom of a sulfonium salt prior to the elimination of the exchanged thioether following an associative pathway (Scheme 8).^{120–123}

Scheme 8. Proposed Mechanism for Thioether–Sulfonium Exchange



The fact that sulfides react with electrophiles (brosylates for instance) and the positively charged sulfonium is a good leaving group comfort the hypothesis of a $S_N 2$ mechanism, while a $S_N 1$ is unlikely considering this pathway would involve a highly unstable primary carbocation. Four decades later, the group of Du Prez studied and updated this forgotten chemistry and implemented it in polythioether networks to prepare catalyst-free vitrimers.⁴⁰ Knowledge on this bond exchange chemistry remains scarce, but the promising opportunities offered in rubber recycling¹²⁴ might favor more in-depth studies.

Silyl Ether Transalkoxylation and Metathesis. Silicones represent an \$11 billion industry, and production of formulated silicones exceeded 2 Mt in 2013.¹²⁵ Silicones are made from silicon, which is also the main component of glass. In the 1950s, polydimethylsiloxanes (PDMS) were reported to exhibit stress-relaxation behavior thanks to a chemical phenomenon of chain exchange, catalyzed by either an acid or a base.^{126,127} The acid-catalyzed mechanism was supposed to involve a protonated oxygen atom on the Si-O-Si bond as an intermediate,¹²⁷ but this mechanism still seems unclear. The base-catalyzed mechanism was supposed to involve the attack of the basic species onto a silicon atom, consequently cleaving a Si-O-Si bond and yielding an anionic ⁻O-Si group.¹² However, the mechanism leading to the exchange between two polymer chains was not fully elucidated. Surprisingly, the selfhealing properties of these materials were somehow forgotten. except for a patent reporting thermally reversible silicone rubbers catalyzed by strong acids, becoming malleable at 150 °C but remaining insoluble and dimensionally stable at room temperature.¹²⁸ The self-healing behavior of polysiloxanes was rediscovered in 2012 by McCarthy et al.¹²⁹ after the concept of vitrimer was first defined. They studied tetramethylammonium

Scheme 7. Mechanism of Meldrum's Acid Exchange with a Thiol as an Example



Scheme 9. Trans-Siloxanation Mechanism¹³⁰

$$R_{1} \xrightarrow{\circ} Si_{0} \xrightarrow{\ominus} + R_{2} \xrightarrow{\circ} Si_{0} \xrightarrow{\circ} Si_{0} \xrightarrow{\circ} R_{3} \implies R_{1} \xrightarrow{\circ} Si_{0} \xrightarrow{\circ} Si_{0} \xrightarrow{\circ} R_{2} + \Theta_{0} \xrightarrow{\circ} Si_{0} \xrightarrow{\circ} R_{3}$$

Scheme 10. Phosphoryl Chloride as a Catalyst Favors the Associative Mechanism of Transcarbamoylation with Phenols¹⁴⁰



Scheme 11. Associative and Dissociative Exchange Pathways for Polythiourethanes^{144,145}



dimethylsilanolate-terminated polysiloxane chains and emphasized that any cross-linked PDMS elastomer could be converted into a self-healable polymer by addition of basic catalysts. The mechanism involved is an associative transsiloxanation. The first step of this mechanism would be the dissociation of the chain-end silanolate-counterion ion pair followed by the attack of the silanolate anion onto a silicon atom and simultaneous cleavage of a Si-O bond (Scheme 9).^{130,131} Thus, the ion pair would behave as a dormant species and the silanolate anion as the active one, enabling chain exchange. Other strong bases can be used to generate living chain ends such as the most effective potassium hydroxide and cesium hydroxide, but a major flaw at high temperature is the formation of volatile cyclic oligomers leading to decreases of connectivity and average molar mass.¹³¹ In 2017, Guan et al. used a silvl ether as the exchangeable moiety in cross-linkers to make polystyrene networks, where the exchange reaction involved free hydroxyl groups without ion pairs.⁴⁵ Finally, the same group reported an exchange reaction in the absence of free hydroxyl moieties and hypothesized a direct silyl ether metathesis mechanism catalyzed by acids, but the mechanism still remains unclear and further investigations are needed to fully understand the behavior of this material.¹³²

Urethanes, Ureas, Thiourethanes, and Hydroxyurethanes. Polyurethanes (PU) are the fifth largest production of industrial polymers with a global production reaching 20 Mt/y.¹³³ The urethane bond, alternatively called carbamate, is usually obtained by reaction between an isocyanate and an alcohol and has been known to be reversible at high temperatures for a long time. Ureas are obtained by reaction between isocyanate and amine, and the direct condensation of ureas leads to biuret functions. In the 1950s, PU materials were shown to exhibit stress–relaxation thanks to urethane cleavage.¹³⁴ Furthermore, relaxation times were shorter when the material included ureas and biuret bonds,¹³⁵ highlighting the increased reactivity of nitrogen in exchange reactions. Catalyst-free vitrimers based on exchangeable urea bonds were also made thanks to bulky groups, which enhances their reactivity toward urea-urea exchange without the need of any catalyst.¹³⁶ Nevertheless, the mechanism involved in this process is uncertain. In the early studies on urethane cleavage, the mechanism was suggested to be dissociative, involving the dissociation of the urethane to its initial isocyanate and alcohol moieties,¹³⁷ as no free hydroxyl groups was available for transurethanization reaction.^{138,139} Even when a free alcohol is available next to a urethane bond, the dissociative pathway occurs most of the time, in some cases along with an associative pathway similar to transesterification (Scheme 1), depending on the system composition and temperature. A low temperature favors the associative mechanism, and reversely, high temperatures favor the dissociative one.¹³⁷ In solution, external catalysts such as phosphoryl chloride were shown to favor the associative mechanism between a urethane and a phenol over the dissociative one (Scheme 10).¹⁴⁰ The direct exchange between two urethane moieties via an associative mechanism has been ruled out.¹³⁹

Ureas seem to be promising bonds to implement in selfhealing materials as the exchange reaction is much faster.¹³⁵ However, the exchange mechanism is similar, with first a dissociation (which rate depends on the steric hindrance of the amine) yielding the starting isocyanate and amine.^{141–143} Thus, based on their apparent dissociative mechanism, polyurethanes and polyureas do not seem to be suitable to obtain vitrimer properties.

Recently, polythiourethanes (PTU) started to be investigated to make self-healing materials. PTU vitrimers catalyzed by dibutyltin dilaurate were synthesized by reaction of a trithiol with different diisocyanates: isophorone diisocyanate, 1,6diisocyanatohexane, or tolylene-2,6-diisocyanate. In these materials, the structure of the isocyanate precursor influenced the relaxation time. Aliphatic isocyanate-derived networks displayed much higher relaxation times at 180 °C than their aromatic analogues. The relaxation times of the isophorone diisocyanate- or 1,6-diisocyanatohexane-derived network were 11.1 and 10.7 min, respectively, whereas it was only 2.9 min for the aromatic tolylene-2,6-diisocyanate-based material. Both associative and dissociative exchange mechanisms were reported to be involved in their reprocessability (Scheme 11).^{144,145}

Compared to traditional PU, the case of polyhydroxyurethanes (PHU) prepared from cyclic carbonates and amines may be different. PHU contain hydroxyurethane bonds, which are carbamate bonds with a hydroxyl group at the β -position (Scheme 12) or γ -position. The synthesis of these materials is advantageously greener compared to PU, as it does not involve the use of isocyanates which are toxic substances.

Scheme 12. Synthesis of Polyhydroxyurethanes (PHU) from (a) Five- and (b) Six-Membered Cyclocarbonates Featuring a Substituent at the β -Position



Fortman et al. studied PHU vitrimers and suggested that transcarbamoylation occurred, albeit slowly, and that decomposition and side reactions occurred more easily at high temperatures.¹⁴⁶ In particular, side reactions were observed at 160 °C and over, whereas the decomposition of the urethane linkage to an isocyanate and an alcohol usually occurs over 250 °C. These observations were highly dependent on the type of PHU: five-membered cyclocarbonate-derived PHU (5-CC PHU, Scheme 12a) were more prone to decomposition via dissociative reactions at 180 °C. 5-CC PHU were shown to decompose back to their 5CC amine precursors (Scheme 13a). Several decomposition reactions were hypothesized: reaction of amine with cyclic carbonates by the irreversible decarboxylative mechanism (Scheme 13b), addition of amines to carbamates to form ureas (Scheme 13c), decarboxylative reaction of free hydroxyl groups with cyclic carbonate to form

ethers (Scheme 13d) or carbonates (Scheme 13e), and oxazolidone formation, to name a few. These irreversible reactions are responsible for the loss of mechanical properties of the reprocessed materials. On the contrary, the dissociative reversion reaction leading to the starting amine and cyclocarbonate was not observed for 6-CC PHU (Scheme 12b), thus limiting the decomposition reactions. This suggests that the associative transcarbamoylation may be responsible for the stress—relaxation behavior observed for these 6-CC PHU materials.^{139,147}

Chen et al. also pointed out that two simultaneous mechanisms were responsible for the network rearrangement of their 5-CC PHU, the associative transcarbamoylation and the dissociative reversible cyclocarbonate aminolysis, yielding back the initial cyclocarbonate and amine.¹⁴⁸ Because two main reactions seem to be accountable for the stress–relaxation behavior of PHU and PTU and because one is associative whereas the other is dissociative, one can rule out this chemistry for associative CANs. However, more research could be done to favor the associative mechanism over the dissociative one, investigating the effects of catalysts, the network morphology, or the monomer structure, for example.

Olefin Metathesis. In the early days of petrochemistry, investigations led on propane cracking revealed that cracking of propylene at 752 °C under 0.2 atm yielded 23-24 mol % of butylene and ethylene.¹⁴⁹ The overall reaction equation is similar to a metathesis. However, this reaction is inexploitable under such conditions that do not allow any control. Olefin metathesis was then developed in the 1960s by chemists working for DuPont and has been extensively studied for both industrial production and research purposes.¹⁵⁰⁻¹⁵² Olefin metathesis requires metal catalysts to occur quantitatively. An in-depth discussion on these catalysts is out of the scope of this article, but a rich literature and very relevant books are available.^{150,152} At first glance, the overall reaction seems to be rather simple and can be summarized as a carbene exchange between two olefins. Nevertheless, the underlying mechanism involved is complex. Yves Chauvin, Richard Schrock, and Robert Grubbs were awarded the Nobel Prize in Chemistry in 2005 for elucidating this mechanism, developing the first

Scheme 13. (a) Reversion of 5-CC PHU to the Original 5-CC and Amine and Hypothesized Decomposition Reactions (b) by Decarboxylation, (c) by Formation of Urea, (d) by Formation of Ether, and (e) by Formation of Carbonate¹⁴⁷



Scheme 14. Olefin Metathesis Mechanism¹⁵⁰

1. Initiation

$$M=CR + R_1 - C=C-R_2 \implies \prod_{R_1-C=C-R_2}^{M=CR} \implies \prod_{R_1-C=C-R_2}^{M-CR} \implies \prod_{R_1-C=C-R_2}^{CR} \implies M=CR_1 + R-C=C-R_2$$

$$\prod_{R_1-C=C-R_2}^{M=CR} \implies \prod_{R_2-C=C-R_1}^{M-CR} \implies \prod_{R_2-R_1-C=C-R_2}^{M-CR} \implies M=CR_1 + R-C=C-R_2$$
2. Propagation

$$M=CR_1 + R_1 - C=C-R_2 \implies \prod_{R_1-C=C-R_2}^{M=CR_1} \implies \prod_{R_1-C=C-R_2}^{M-CR_1} \implies M=CR_1 + R_1 - C=C-R_2$$

$$\prod_{R_1-C=C-R_2}^{M=CR_1} \implies \prod_{R_1-C=C-R_2}^{M-CR_1} \implies \prod_{R_2-C=C-R_1}^{M-CR_1} \implies M=CR_1 + R_1 - C=C-R_2$$

$$M=CR_1 + R_1 - C=C-R_2 \implies \prod_{R_1-C=C-R_2}^{M=CR_1} \implies \prod_{R_2-C=C-R_1}^{M-CR_1} \implies M=CR_1 + R_1 - C=C-R_2$$

$$M=CR_1 + R_1 - C=C-R_2 \implies \prod_{R_1-C=C-R_2}^{M=CR_1} \implies \prod_{R_2-C=C-R_1}^{M-CR_1} \implies M=CR_2 + R_1 - C=C-R_1$$

$$M=CR_2 + R_1 - C=C-R_2 \implies \prod_{R_1-C=C-R_2}^{M-CR_2} \implies \prod_{R_2-C=C-R_1}^{M-CR_2} \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies \prod_{R_1-C=C-R_2}^{M=CR_2} \implies \prod_{R_2-C=C-R_2}^{M-CR_2} \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies \prod_{R_1-C=C-R_2}^{M-CR_2} \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies \prod_{R_1-C=C-R_2}^{M=CR_2} \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies \prod_{R_1-C=C-R_2}^{M-CR_2} \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_1 - C=C-R_2 \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_2 - C=C-R_1 \implies R_2 - C=C-R_2 \implies M=CR_1 + R_2 - C=C-R_2$$

$$M=CR_2 + R_2 - C=C-R_1 \implies R_2 - C=C-R_2 \implies M=CR_2 + R_2 - C=C-R_1$$

$$M=CR_2 + R_2 - C=C-R_1 \implies R_2 - C=C-R_2 \implies R_2 - C=C-R_2 \implies R_2 - C=C-R_1 \implies R_2 - C=C-R_2 \implies R_2 -$$

catalysts, and synthesizing more convenient catalysts, respectively.^{153–155} Their studies deepened the knowledge of this exchange reaction, which was useful to develop new catalysts. Indeed this reaction fully relies on metal-carbene catalysts. The mechanism of olefin metathesis is detailed in Scheme 14.^{150,156–159} All the steps for this reaction are equilibria, so if the reactants and desired exchanged products have a similar enthalpy, the reaction is degenerate. This property is convenient to make exchangeable cross-links in vitrimers. In contrast, when metathesis is used for organic synthesis, aiming at efficient synthesis of a desired exchanged olefin product, the equilibrium can be easily displaced by using a R-CH=CH₂ olefin as the substrate. Thereby, the direct reaction yields the desired product, and easy removal of ethylene from the reaction vessel provides the necessary driving force for the reaction to happen preferentially in the desired direction. An example of polybutadiene vitrimers implementing olefin metathesis as the exchange reaction was reported.^{160,161} These materials were made of cross-linked polybutadiene by reaction of butadiene with benzoyl peroxide. The interest of polybutadiene is the presence of insaturations in the network after cross-linking. These insaturations can undergo olefin metathesis in the presence of a Grubbs catalyst. Therefore, it is possible to make a vitrimer by adding a Grubbs secondgeneration catalyst before the polymerization, but a much more original approach is to make a vitrimer out of a thermoset by swelling it with a solution of the catalyst after the cross-linking process. Such a strategy to make vitrimers has so far been specific to olefin metathesis vitrimers. These vitrimers exhibited self-healing at ambient temperature and below. Selfhealing behavior was also observed for thermoset materials after the catalyst was only applied on the surface where healing was required. When no catalyst was used, no effective self-

healing was observed. In summary, olefin metathesis allows original strategies to make vitrimers, which is not possible with other chemistries. However, the use of a metal catalyst seems unavoidable and could be seen as the major drawback of this exchange reaction.

Imines. Imines result from the condensation of an aldehyde or a ketone and a primary amine to obtain a C=N bond. Imines were investigated in the field of biochemistry, as their reversible covalent bond is known to be involved in enzymatic reactions,¹⁶² and more recently in the field of materials chemistry,^{163–175} with the growing interest in dynamic covalent networks. Actually three exchange reactions are known for imines: a dissociative hydrolysis followed by reaction with another amine; an associative exchange between an imine bond and a free amino moiety called transimination; and a direct associative imine-imine exchange termed imine metathesis.^{166,170} Obviously, only the latter two associative reactions are interesting for associative CANs design. Although dissociative hydrolysis may happen in imine-based vitrimers, if the amount of water in the polymer matrix remains low, this reaction is unlikely to influence significantly the properties of the material. Therefore, the focus here is put on transimination, which requires residual amines, and on imine metathesis, which does not. Transimination has been known to occur through an equilibrated associative pathway for several decades. 176,177 Three successive steps were described. The first and rate-determining step consists in the attack of a free primary amine on the imine carbon atom to form a geminal diamine tetrahedral intermediate.^{162,178–181} Then, two protons transfer from the nitrogen atom of the formerly attacking amino moiety to the nitrogen atom of the former imine moiety.^{162,179,181} Finally the exchanged amino moiety dissociates and leads to the formation of the new imine¹⁶²

Scheme 15. (a) Transimination and (b) Imine Metathesis Mechanisms



Scheme 16. Vinylogous Urethane Transamination Mechanism (a) in Acidic and Neutral Conditions and (b) in Basic Conditions



Scheme 17. Diketoenamine Transamination Catalyzed by p-Toluenesulfonic Acid (p-TsOH)



(Scheme 15a). The mechanism of imine metathesis is less extensively studied but is described as a nucleophilic addition of a neutral imine to a protonated one. The reaction proceeds through a 1,3-diazetidinium intermediate, yielding the exchanged imines or the initial reactants^{179,182} (Scheme 15b). Transimination and imine metathesis have already been implemented in various polymeric materials to make vitrimers directly from polyimines^{163,165,167,169} or by using imine bonds to cross-link polyesters, ^{166,172,173,175} polybuta-dienes, ^{164,168} polydimethylsiloxanes, ¹⁷¹ and polyethers. In the late

Vinylogous Urethanes, Ureas, and Amides. In the late 1970s, the exchange of amine moieties on enaminone structures was reported.¹⁸³ Later in the 1990s, similar exchanges between dialkylamine-derived enaminones and aromatic primary amines were discussed.^{184,185} However, this exchangeable bond was not implemented in polymers until 2014.¹⁸⁶ Moreover, the mechanism involved in this first example was based on a reversible hydrolysis to the amine and carbonyl components, although a possible "component exchange" (amine direct exchange) was evoked by the authors. In this field of chemistry, enaminones are commonly termed vinylogous urethanes, vinylogous ureas, and vinylogous amides depending on their structure. These structures are easily prepared by acetoacetylation of polyols followed by reaction

with a primary amine.¹⁸ In 2015. Denissen et al.¹⁸ reported the first vinylogous urethane vitrimer, exhibiting relaxation times as short as 85 s at 170 °C without the need of any catalyst. The exchange mechanism was later described by the same team¹⁸⁷ and depends on whether the medium is acidic, neutral, or basic. In neutral and acidic conditions, the free primary amine is protonated to form an ammonium, and the ammoniumenamine couple is in equilibrium with the amine-iminium couple. The latter couple can evolve by reaction of the nitrogen atom of the amine with the iminium to form a geminal amineammonium tetrahedral adduct similar to the intermediate described earlier for transimination. Then, dissociation occurs, yielding either the exchanged products or the reactants (Scheme 16a). The amine-iminium pathway written in Scheme 16a (bottom) for a Brønstedt acid can be extended to Lewis acids. Under these conditions, the reaction can proceed through a zwitterionic intermediate.

For instance, Friary et al.¹⁸⁴ studied the transamination of an enaminone bearing alkyl substituents with an aromatic primary amine in solution. The exchanged enaminone was then able to cyclize. When 1 equiv of toluenesulfonic acid was added, the overall yield increased from 29 to 56%. The authors suggested that the acid accelerated the enaminone transamination step (Scheme 17). The catalyst protonates the oxygen atom of the

L TREN	p-TsOH 0, OH	Reference	R (AA/N) ^a	I _g (°C)	p-TsOH	τ (s) ^b	E _a (kJ.mol ⁻¹) ^c
	٥̈́ ()	VU-E _{H+}	1.0	51		80	84.9
	/~	VU-A	0.7	13		29	75.9
		VU-A _{H+}	0.7	17	\checkmark	0.3	60.3
	~YY	VU-NA	1.35	-8		> 46800	-
hexane-1,6-diyl bis(3-o	oxobutanoate)	VU-NA _{H+}	1.35	29	\checkmark	134	75.8

Figure 3. Impact of the stoichiometry and p-TsOH catalysis on vinylogous urethane properties. ^aRatio acetoacetate/amine. ^bRelaxation time at 110 °C. ^cActivation energy (VU-E stands for an equal ratio of amines and acetoacetates, VU-A an excess of acetoacetates, and VU-NA an excess of amines).¹⁹¹

enaminone and generates an iminium tosylate, more electrophilic toward the attack of the amine.

The exchange kinetics can be tuned by using simple additives. Brønsted and Lewis acids such as sulfuric acid, dibultyltin dilaurate (DBTL), or zinc(II) accelerate the reaction. The use of 1 mol % p-toluenesulfonic acid was reported to reduce the time to reach equilibrium from >1 h to <10 min at 100 °C. On the contrary, non-nucleophilic bases such as triazabicyclodecene (TBD) were reported to increase the equilibration time.^{187,188} In a basic environment the exchange reaction happens but is slower and involves a direct Michael-like addition of a neutral amine on a neutral vinylogous urethane (Scheme 16b).^{187,189} These mechanisms are all associative and thus suitable to obtain vitrimer properties. Some vitrimers exhibit a dual temperature response.^{189,190} Interestingly, their dual behavior was explained by possibly concomitant mechanisms a and b (Scheme 16), which are usually independent as they require different pH conditions. The iminium pathway would be dominant at low temperatures whereas the neutral Michael-type pathway would prevail at high temperatures.

Catalysts also have an impact on the behavior of vitrimers. The relaxation time of vinylogous urethane materials was reported to decrease from 10 min at 120 °C in the absence of catalysts to 2 min with 0.5 mol % of p-TsOH.¹⁸⁷ A recent study by Haida and Abetz¹⁹¹ studied the impact of p-TsOH in vinylogous urethane vitrimers with different stoichiometries. The networks were synthesized from hexane-1,6-diylbis(3oxobutanoate) and tris(2-aminoethyl)amine (TREN). When an excess of amine was used, the relaxation time decreased from 29 s without p-TsOH to 0.3 s with 0.05 wt % p-TsOH. Similarly, with an excess of the acetoacetate the relaxation time decreased from >46800 s to only 134 s (Figure 3).¹⁹¹ DBTL and sulfuric acid were shown to be less effective. In the case of sulfuric acid, the sulfate anions probably form salts with the ammonium ions. In consequence, the ammonium species are less available and thus less reactive. Similarly, carboxylate anions are also known to be proton scavengers for ammonium species in nonaqueous solutions, for instance.¹⁸⁷ When DBTL was used, the reaction activation energy dropped from 74 to 45 kJ mol⁻¹, suggesting a different mechanism, probably via the carbonyl moiety activation of the vinylogous urethane. On the contrary, TBD increased the activation energy up to 103 kJ mol⁻¹ because of proton scavenging. Interestingly, Spiesschaert et al.¹⁹² managed to incorporate acidic or basic functionalized fillers in PDMS-vinylogous urethane matrices. When 10 wt % kaolin bearing acidic groups was incorporated, the relaxation time of the resulting material decreased from 5725 to 4650 s at 150 °C, whereas nonfunctional fillers were shown to increase relaxation time. Acid, basic, or neutral alumina fillers were also studied. When the relaxation time for the matrix was 4000 s,

the material containing neutral alumina or basic alumina displayed relaxation times of 4650 and 4500 s, respectively. On the contrary, the material made from acid alumina showed a shorter relaxation time (3600 s), in agreement with the results observed in solution (Figure 4).¹⁹² Other kinds of catalysis were also reported, and not surprisingly an excess of free primary amines in a vinylogous urethane vitrimer accelerates transamination.¹⁸⁹



Figure 4. Effect of different functionalization of alumina fillers on vinylogous urethane vitrimer composites' relaxation time.

Eventually, it is important to highlight the crucial role of the backbone and cross-linking density on the behavior of vitrimers, as already mentioned for imine vitrimers. Spies-schaert et al.¹⁹⁰ recently disclosed in a detailed study that the relaxation rate of various vinylogous urethane-cross-linked polyether vitrimers decreased with increasing cross-linking density. An effect on the activation energy was also reported even if a clear trend was not identified. A dual temperature behavior was reported for polytetrahydrofuran, poly(propylene glycol), and poly(ethylene glycol) backbones, as reported previously in perfluoropolyether.^{189,190} Nonetheless, this effect was not observed on a polypropanediol backbone, emphasizing the influence of both the matrix type and the molar mass of the spacers. The dual mechanism discussed earlier is probably the cause for this dual behavior.¹⁹⁰

Because they are versatile and they allow short reprocessing times, vinylogous urethanes have already been implemented in various polymers networks. Many examples were reported such as polysaccharide hydrogels,¹⁹³ polysiloxanes,^{188,190,192,194,195} polybutadienes,^{190,196} perfluoropolyethers,¹⁸⁹ methyl methacrylate-(2-acetoacetoxy)ethyl methacrylate copolymers,¹⁹⁷ polystyrenes,¹⁹⁸ polydimethylsiloxane (PDMS)–kaolin, PDMS– alumina and PDMS-aminated silica composites,^{188,192} epoxy– amine copolymers,¹⁹⁹ polyethers,¹⁹⁰ and vinylogous urethane polymers bearing photodimerizable anthracene units.²⁰⁰

Diketoenamines. Diketoenamines are known as a Dde protecting group for amines in peptide synthesis.^{201,202} This protecting group is formed by reaction between 2-acetyl-5,5-dimethyl-1,3-cyclohexanedione (also known as 2-acetyldimedone) and primary amines.²⁰³ In 2009, the behavior of Dde was studied in detail on lysine in solid-phase synthesis. Lysine is an interesting amino acid bearing two amino moieties in the α - and ε -positions.²⁰¹ This study reported the ability of ε -NH₂ to attack a Dde-protected ε -NH₂ or α -NH₂, leading to an

Scheme 18. Dde Protection Group Exchange on Amines (a) by Direct Electrophilic Attack and (b) Mediated by Piperidine²⁰¹



Scheme 19. Mechanism of Diketoenamine Exchange



Scheme 20. Example of Synthesis of Diketoenamine Vitrimer; Alternatively, Various Diketones and Mixes of Primary Triamines and Diamines Were Also Used^{204,205}



amine exchange. According to the authors,²⁰¹ the mechanism involves a direct nucleophilic attack of the free amine on the Dde group via a Michael addition, before a retro-Michael addition yielding either the original structure or the amine-exchanged one (Schemes 18a and 19).

Another mechanism involving piperidine as catalyst was also described. Piperidine acts as a mediator to pick up the Dde group from the originally protected amine before attack of another primary amino group. The overall reaction is composed of two successive exchanges: first between the originally protected amine and piperidine and then between the protected piperidine and a free amine (Scheme 18b). In all cases, the mechanism of amine exchange is associative.

This chemistry was implemented in materials synthesis by Christensen et al. to make vitrimers;^{204,205} they reported the synthesis of a network by click polycondensation between β triketones and primary amines (Scheme 20). Triketones were synthesized from polytopic carboxylic acids and 1,3-diketones and the final materials simply by ball-milling of the reactants at room temperature. The materials obtained were reprocessable without requiring a catalyst and followed an Arrhenius behavior. These materials are easily hydrolyzed. While this feature is often considered as a flaw, it can be beneficial for specific applications. In particular, as the materials are fully hydrolyzed in a few hours at room temperature in sulfuric acid (<12 h in 5 M H_2SO_4), the monomers can be easily recovered. Amines are separated from the triketones thanks to an ion-exchange resin, with a recovery above 90%. The recovered monomers can be reused to make new poly(diketoenamines). This feature affords a convenient and efficient chemical recycling by hydrolysis at room temperature, while other hydrolyzable vitrimers require heating and more demanding conditions.

The range of exchange reactions suitable for the design of vitrimers is already large (Figure 2), and new exchange chemistries occurring via associative pathways are still to be implemented in such materials,²⁰⁶ although proving the associative character of an exchange reaction is often challenging and remains uncertain in some cases. In addition, a survey of the activation energies of the associative exchange reactions implemented in vitrimers (Figure 5) shows that these energies can be distributed over relatively large ranges. This is certainly an asset for chemists and materials scientists who want to tune the reshaping ability of vitrimers.



0 10 20 30 40 50 60 70 80 90 100 110 120 130 140 150 160 170 180 190 200 210 220 230 240 250 ${\rm E_{a}}$ (k.J.mol^-1)





Figure 6. Examples of effects to activate bond exchanges.

III. INTERNAL CATALYSIS, STERICS, AND NEIGHBORING GROUPS EFFECTS ON EXCHANGEABLE BONDS

Many parameters influence the properties of vitrimers: the cross-link density, the nature of the network backbone, and obviously the type of exchangeable bonds, for instance. For the exchange reaction to be efficient, many vitrimers rely on external catalysts. These compounds often come with some risks of leaching out of the material or premature aging limiting the recycling abilities of the materials. To address these problems, internal catalysis, steric effects, and neighboring group participation (NGP) are valuable. For a given exchange reaction, well-designed chemical modification of the exchangeable bonds surroundings is a way to accelerate the exchange, without resorting to any external catalyst. Different effects at the molecular scale are gathered in these three concepts (Figure 6). In light of the reaction mechanisms discussed, some relevant activating groups, either already implemented in bulk vitrimer networks or only in solution, are presented below. This should give new insights for further research in such dynamic materials.

Transesterification Enhancement by Neighboring Groups: Influence on the Exchange Mechanism and



Figure 7. Main examples of activating groups activating transesterification. The temperature range²⁵ for stress-relaxation experiments and the range of calculated activation energies²⁴ of the exchange reaction in vitrimers are given.

Vitrimer Properties. A few activating groups have been reported to influence the reactivity of esters toward trans-esterification and/or hydrolysis in solution (Figure 7).

For instance, on phenyl esters, a neighboring hydroxymethyl group on the ortho position increases the transesterification rate. However, the mechanism involves a cyclization after the attack on the ester by the -OH group, leading to a transient lactone (Scheme 21).²⁰⁷

Scheme 21. Catalytic Effect of an Ortho Hydroxymethyl Group on Phenyl Ester Transesterification by Transient Lactone Formation



This chemistry if it were implemented in a CAN would proceed through a dissociative mechanism. A regioselectivity effect is also reported on pyridine triesters as one of the ester groups possesses an ortho hydroxyl group. Interestingly the transesterification reaction was reported to be regioselective on the ortho hydroxy ester, highlighting a reactivity increase (Scheme 22).²⁰⁸ Nevertheless, in this case the regioselectivity

Scheme 22. Regioselective Transesterification: (a) 10 mol % EtNiPr₂, ROH, 60 °C; (b) 5 mol % Sc(OTf)₃, ROH, cat. H_2O , 60 °C²⁰⁸



is due to the hydrogen-bonding ability of the hydroxyl group, and the mechanism is associative, contrary to the previous example.

The same kind of regioselectivity was reported with 3benzamidophthalates. The ester group located ortho to the amide moiety undergoes transesterification whereas that located in the meta position does not (Scheme 23).²⁰⁹

Two mechanisms were discussed depending on the substituent: the first one is the regular attack of the alcohol

Scheme 23. Regioselectivity of 3-Benzamidophtalates Transesterification



on the ester groups activated by hydrogen bonding with the proton of the amide (Scheme 24a), and the second one is the formation of a more reactive 3,1-benzoxazin-4-one intermediate (Scheme 24b).

In contrast, neighboring thioether groups stabilize the ester bond,²¹⁰ hence decreasing its reactivity toward transesterification. Fluorene structures (Figure 8) bearing a methyl carboxylate substituent on the 1-position were studied.

When one ethylthio group was present on the central fivemembered ring, the rate constant of transesterification (using methanol- d_4) was 0.29 times the rate of the unsubstituted fluorene ester. Interestingly, on the fluorene ester bearing two ethylthio groups, the effect was shown to be opposite. This ester was more prone to transesterification. According to the authors of this study,²¹⁰ this phenomenon is likely due to at least two different causes, resulting in opposite effects on the transesterification rate. On one hand, the steric hindrance induced by one ethylthio group would decrease the transesterification rate, as the reactive site is less accessible. On the other hand, ethylthio groups induce a stabilizing electronic effect on the protonated ester intermediate (Scheme 25). The authors hypothesized that the electronic effect were low compared to the steric effect when only one ethylthio substituent was present. On the contrary, when two ethylthio groups were present, the resulting electronic effect is significant and exceeds the steric effect, even though this effect is stronger than in the monosubstituted counterpart.

 β -Ketoesters as well as esters derived from malonates or oxalates, for example, are also more prone to transesterification reactions in solution.²¹¹ Indeed, acetoacetic acid is 10 times stronger than acetic acid, so its conjugated base is more stable and the corresponding ester bond is weaker, leading to easier transesterification.²¹¹ Oxalates' reactivity toward transesterification is explained exclusively by inductive effects. For β -ketoesters and malonates, a stabilized cyclic intermediate was

Scheme 24. Regioselective Transesterification of 3-Benzamidophtalates: (a) Direct Pathway and (b) Pathway with 3,1-Benzoxazin-4-one Intermediate



Figure 8. Effect of substitution by ethylsulfane groups on fluorene esters' transesterification rate.²¹⁰

Scheme 25. Stabilizing Effect of Sulfur on Protonated $\operatorname{Esters}^{210}$



proposed to explain the reactivity of the other aforementioned species, though inductive effects also play a significant role (Scheme 26).

Scheme 26. Proposed Stabilization by Cyclic Intermediate for the Transesterification of β -Ketoesters²¹¹



Recently, diethylmalonate was used as cross-linker in a polyester vitrimer, disclosing the enhancing effect of β -activated esters.²¹²

Phthalate monoesters also displayed enhanced transesterification similarly to the previously discussed *o*-hydroxymethyl phenyl esters,²⁰⁷ as demonstrated by Du Prez et al.⁴⁷ The neighboring carboxylic acid can indeed attack the carbonyl of the ester to yield a cyclic anhydride and an alcohol (Scheme 27a). Recently the same team enhanced this effect by using a β -aminodiol with a dianhydride to make networks (Scheme 27b). The enhancing effect of β -amines had already been described for ester hydrolysis in 1962.²¹³ The tertiary amine at the β -position of the hydroxyl groups induced a dual activation: the neighboring group participation of the dangling -COOH and the internal catalysis effect of the tertiary amine. The stress-relaxation times decreased by a factor 500 compared to the amine-free analogous network.²¹⁴ However, it should be noted that the mechanism was dissociative in these cases, highlighting that great care should be taken when considering neighboring groups participation as they potentially change the bond exchange mechanistic pathway.47,214 The same mechanism was also recently reported with 2,5bis(methoxycarbonyl)benzenesulfonic acid, a benzoester bearing an o-sulfonic acid moiety (Scheme 27c).⁴⁸ These materials showed a faster stress-relaxation than their carboxylic acid analogues.

Amines were also shown to be efficient catalysts for transesterification vitrimers.¹⁷ Triazabicyclodecene (TBD) in particular has often been used. Strategies have been designed to integrate tertiary amine groups into the polymer network. For example, bisphenol A diglycidyl ether (BADGE) was mixed with primary or secondary amines to generate epoxy-terminated tertiary amines oligomers. In a second step, the oligomers were reacted with polyacids to obtain a network bearing covalently bonded tertiary amines (Figure 9a).³⁸ The exchange reaction activation energy (E_a) for this network was 93.6 kJ mol⁻¹, very close to the 80–90 kJ mol⁻¹ range observed for the Zn-catalyzed epoxy vitrimers. Alternatively, triethanolamine was added as coreactant in a BADGE– (methyl nadic anhydride) system for the same purpose (Figure 9b).²¹⁶ The relaxation time at 190 °C decreased from 19460 to

Scheme 27. Dissociative Transesterification Pathway Involving a Cyclization (a) by an o-Carboxylic Acid,⁴⁷ (b) by an o-Carboxylic Acid,⁴⁸ (c) by an o-Sulfonic Acid⁴⁸



Figure 9. Strategies to embed tertiary amines in polyester networks using (a) epoxy-ended tertiary amines (two steps),³⁸ (b) ethanolamine as coreactant,²¹⁶ and (c) a commercial tetraepoxide bearing tertiary amines.²¹⁷

4200 s when the triethanolamine content increased from 5 to 10 mol % (Figure 11).²¹⁶ Recently, catalyst-free transesterification vitrimers were synthesized from a hydrogenated dimerized acid derived from vegetable oil and tetraglycidyl-4.4'-diaminodiphenylmethane (TGDDM), a tetrafunctional epoxide bearing two tertiary amino groups (Figure 9c).²¹⁷ These tertiary amines were shown to act as internal catalysts. The effect was reported to be enhanced by the flexibility induced by the vegetable oil structure. Instead of increasing monomers' flexibility, dangling chains which can plasticize the material were also shown to promote similar enhancement as tertiary amines' internal catalytic effect.²¹⁸ However, a similar internal catalysis was also reported in systems based on shorter, more rigid acids and which did not contain any plasticizer.²¹⁹ These examples of internal catalysis emphasize the role of tertiary amines as promising activating groups to increase the transesterification rate.

Interestingly, a chemistry similar to the first vitrimer¹ featuring β -hydroxyesters pointed out the participation of the hydroxyl moiety in transesterification.^{34,220,221} A six-membered ring with a hydrogen bond between the hydroxyl and the ester groups was hypothesized (Figure 10). The hydrogen bond is assumed to weaken the ester bond as previously discussed for the 3-benzamido group.



Figure 10. Hypothesized activating effect of the β -hydroxyl groups for transesterification. The hydrogen bond (dashed bond) increases the carbonyl C electrophilicity, facilitating the attack of an electrophile such as an alcohol (first step of the transesterification).

Besides, increasing the amount of free hydroxyl moieties was used to prepare catalyst-free transesterification vitrimers.^{29,222} Although this effect cannot be considered as neighboring group activation or internal catalysis, it remains significant in transesterification vitrimers, was often reported in the vitrimer literature, and therefore deserves to be mentioned. The acceleration of the exchange reaction can be explained by the large amount of hydroxyl groups, facilitating the transesterification by a concentration effect.^{33,219} In addition, a catalytic effect is also likely caused by hydrogen bonding, as in the case of oxime–esters (vide infra). Accordingly, the addition of glycerol to an epoxy–anhydride vitrimer matrix increases the concentration of free hydroxyl groups. The concentration effect is clear as the relaxation time of the materials at 180 °C dropped from 204 min to 86 and 66 min for 25, 50, and 75 mol % of glycerol, respectively (Figure 11).³³

The Case of Oxime-Esters. Oxime-ester bonds can be considered as a subgroup of the ester family or as a specific type of bond. The most important difference between the oxime-ester and ester is the nucleophile (an oxime instead of an alcohol). Heat-induced transesterification of the oximeester enabled the synthesis of catalyst-free vitrimers (Scheme 28),²³³ with relaxation times ranging from 5102 to 2725 s at only 100 °C, depending on the free oxime-OH available and their concentrations ranging from 0 to 5 mol % (Figure 11). The acceleration of the exchange reaction is caused mainly by the high concentration of exchangeable bonds. However, part of the enhancing effect can be explained by the higher nucleophilicity of the oxime (exchangeable moiety) compared to hydroxyl groups. This would allow to decrease relaxation temperatures and times. Nevertheless, considering the scarcity of reports dedicated to oxime-ester vitrimers, it would be unwary to compare the relaxation parameters with transesterification vitrimers, as many parameters are varying between the systems reported in the literature, such as the backbone or the cross-linking density, for example.

Carbonate Activation by Electron-Withdrawing Groups. Transcarbonation (Scheme 3) is used to synthesize polycarbonates, but this reaction requires a catalyst (Figure 12). Basic catalysts are probably the most used. Indeed, the generation of a more nucleophilic alkoxide increases the rate of addition onto the electrophilic carbon of the carbonate.^{69,70,74,77–80} Activation of the carbonate with Lewis acids has also been reported but is less common.⁷⁴ Besides, some metal catalysts (such as tin or calcium salts, for instance) that may combine Lewis acid activation of the carbonate^{67,71,73,76–78} and activation of the nucleophilic alco-



Figure 11. Effect of various enhancing strategies on stress-relaxation time: hyperbranched epoxy prepolymers (HBE), network-embedded triethanolamine (TEA), and free oxime hydroxyl group in the oxime-ester network.^{33,216,233} Note that comparison between these three systems should be exerted with caution since structural parameters (backbone, cross-linking density, etc.) differ from one system to another.

Scheme 28. Transesterification of Oxime-Ester Bonds²³³



Figure 12. Examples of activating groups enhancing transcarbonation rate in solution and characteristics of reported transcarbonation vitrimers.²²⁵⁻²²⁸



Figure 13. (a) Structures investigated by Tillett and Wiggins.²²⁴ (b) Effect of the electron-withdrawing group (EWG) position on the exchange rate.²²⁵ (c) Polymerization rate of nitrophenylcarbonates with bisphenol A depending on the substituent position.²²⁶ (d) Effect of CF₃ substituent on the exchange rate.²²⁹

hol^{71,73,75,77–79} have been also used. The formation of metal complexes coordinating both reacting partners increases the electrophilicity of the carbonyl^{67,71,74,77–79} and also has a templating effect that accelerates the reaction. Nevertheless, neighboring group effects on transcarbonation are still relatively unknown (Figure 12).

Inspired by a similar study on esters, Tillett and Wiggins investigated the effect of hydroxy- and methoxy-substituted phenylethylcarbonates on their hydrolysis rate (Figure 13a).²²⁴ An influence on the hydrolysis rate could be extrapolated to transcarbonation reactions due to the strong mechanistic similarities between these reactions. At high pH, the hydrolysis



Figure 14. Polymerization of bis(methylsalicyl)carbonate (BMSC) and diphenylcarbonate (DPC) with bisphenol A (BPA).²²⁷

rates of o-hydroxy-, p-hydroxy-, and o-methoxy-substituted compounds were not significantly different. However, in neutral conditions the o-hydroxy-substituted phenylethylcarbonate was hydrolyzed 10 times faster than the two other compounds. General principles for transcarbonation were given in a very complete review on carbonates in 1996.²²⁵ Generally, the more nucleophilic alcohol displaces the less nucleophilic one, and in the case of equal nucleophilicity, the less volatile alcohol displaces the more volatile one. This rule does not apply to phenols, as dialkylcarbonates react with phenols to make diarylcarbonates. In addition, alcohols bearing bulky groups or electron-withdrawing groups are poorly reactive. On the contrary, when the carbonate contains electron-withdrawing groups, the alcohol exchange is accelerated. For instance, fluorinated dialkylcarbonates and diarylcarbonates bearing -NO2, -CN, or -Cl groups on the ortho position undergo much faster alcohol exchange (Figure 13b). The activation is less noticeable when the substituents are on the para position. Brunelle et al. harnessed this effect with the use of a catalyst. Hence, they successfully polymerized onitrophenylcarbonates with bisphenol A at room temperature and even below, 226 whereas high temperatures (up to 200-300°C) are usually required to prepare polycarbonates.²²⁷ The exchange between nitrophenylcarbonates and phenol was twice faster with the ortho-substituted carbonate than with the parasubstituted homologue (Figure 13c). Similarly, bis-(trifluoroethyl)carbonates were shown to exchange with various alcohols and phenols in refluxing heptane or toluene (at 98 and 111 °C, respectively) in the presence of a base, ^{225,228} and their reactivity was reported to be between the very reactive dimethylcarbonate and the less reactive diphenylcarbonate (Figure 13d).²²

Kamps et al. studied bis(methylsalicyl)carbonate (BMSC, Figure 14) as an activated carbonate to lower the temperature for melt transcarbonation synthesis of polycarbonate.²²⁷ BMSC was polymerized with bisphenol A (BPA), and this polymerization was compared to that of diphenylcarbonate (DPC) and BPA. DPC-BPA polymerization was achieved in 135 min, whereas for the BMSC-BPA system, 40 min was sufficient to reach similar molar masses (Figure 14). This polymerization is composed of two successive substitutions. The rates of the first and second substitutions were reported to be 30 and 9 times higher for the BMSC-BPA system, respectively. Although this study focused on the direct overall two-step reaction, the rate of the reverse reaction for the first substitution was determined for both BMSC and DPC. Indeed, the rate of the first step reverse reaction for DPC was half that of the direct reaction, whereas for BMSC it was 3 decades lower. The overall two-step reaction is an equilibrium. When this equilibrium was reached for the DPC-BPA system, the

main product was the monosubstituted compound, and the same amounts of starting carbonate and disubstituted product were observed. In contrast, for the BMSC–BPA system, the disubstituted carbonate was the main product and the monosubstituted product was also present as byproduct, whereas no starting material remained, suggesting that the equilibrium was more displaced toward the products in this case. Such observation is significant for the design of vitrimers based on exchangeable carbonate bonds.

Groups containing nitrogen seem to exert a significant activating group effect on carbonates. Amine-modified mesoporous silica prepared from amine-terminated silanes showed a catalytic effect on the transcarbonation between ethylene carbonate and methanol to produce dimethylcarbonate.⁸⁰ Thus, an amino neighboring group might have interesting effects on the transcarbonation reaction, too. Examples of vitrimers based on carbonate exchange are very scarce,^{230,231} and they rely on Ti(IV) catalysts; they thus exhibit similar flaws to metal-catalyzed transesterification vitrimers discussed earlier. Smart molecular design based on the knowledge on activating groups effects on transcarbonation might help the development of such transcarbonation vitrimers.

Dioxaborolane Transesterification Acceleration. As discussed previously, boronic esters were mainly used in aqueous media. Therefore, the effects of activating groups on boronic esters were mainly described under such conditions. Although the behavior of an exchangeable bond is very different in solution and in the bulk of a material, especially with a peculiar solvent such as water, it may still be possible to draw general conclusions on the effects of activating groups (Figure 15). In aqueous solution, there is an equilibrium between the dissociated boronic acid—diol couple and the corresponding boronic ester. The formation of a boronic ester



Figure 15. Reported activating groups enhancing dioxaborolanes' exchange rate and range of stress-relaxation experiment temperatures²⁵ and activation energies.²⁴



Figure 16. Synthesis of a network based on boronic esters: (a) poly(ethylene glycol) (PEG) monomers functionalized by boronic acids (blue) and diols (red) and their equilibrium with the boronic ester form; (b) schematic representation of the network.²³⁶

by reaction between a boronic acid and a diol must be performed at a pH higher than the pK_a of the acid.^{232,233} Interestingly, the pK_{a} of boronic acids can easily be tuned by chemical modification. For instance, compared to the pK_{a} value of 8.8 of phenylboronic acid, the pK_a of methylboronic acid is 10.4, whereas the pK_a of the very electron-withdrawing 3-pyridylboronic acid is 4.0.⁸⁴ This shift of the esterification equilibrium gives a glimpse of the possible electronic and internal catalysis effects. An early study showed that arylboronic acids bearing ortho neighboring groups containing oxygen, sulfur, or fluorine do not enhance the esterification or the transesterification.²³⁴ Nevertheless, electron-withdrawing groups decrease the boronic acid pK_{a} , whereas electron-donating groups increase it^{83,84} and destabilize the ester bond. For instance, the boron atom of phenylboronic acids is reported to be highly electron deficient, which reduces the pK_a and facilitates ester formation.^{84,233,235} Recently, benzoxaborole-cross-linked poly(ethylene glycol) (PEG) networks were synthesized out of 4-carboxy-3-fluorophenylboronic acid along with its non-fluorinated counterpart (Figure 16). Four-arm amine-terminated PEG was modified with 4-carboxy-3fluorophenylboronic acid or 4-carboxyphenylboronic acid to afford tetrafunctional boronic acids. The same procedure was applied to D-glucolactone to obtain a tetrafunctional diol. Each tetrafunctional acid was then reacted with the tetrafunctional diol to make a network (Figure 16). This study showed that the resulting partially fluorinated material had a higher dynamic modulus and a faster relaxation time than the nonfluorinated analogue.²³⁶

Nevertheless, the most studied and most promising activating groups so far are nitrogen-containing groups. Aminomethyl-substituted arylboronic acids were shown to be esterified several orders of magnitude faster than non-substituted phenylboronic acids.²³⁷ Their transesterification was also accelerated compared to unsubstituted boronic acids or such acids bearing substituents containing oxygen, sulfur, or fluorine.^{234,238} In solution, the addition of a dimethyl-

aminomethyl group on a phenylboronic ester was shown to accelerate the rate of transesterification by 5 orders of magnitude.⁴⁴ The amino group on the ortho or para position decreases the pK_a of phenylboronic acid²³³ and facilitates the formation of the ester.²³⁹ This is due to electronic substituent effects on the aromatic ring. Even when the amino moiety is not directly attached to the aromatic ring, the acceleration effect is noticeable. In this case, the effect is caused by the formation of B–N dative bonds (a proper example of neighboring group participation, Figure 17a).^{83,85,233,239}

Interestingly, amino activating groups have been implemented in vitrimers.⁴⁴ They demonstrated an accelerating effect on boronic esters transesterification in bulk, which enabled the tuning of the properties of dioxaborolane vitrimers. Indeed, materials containing amino activating groups relaxed stress within 5 min while the reference dioxaborolane material required more than 20 min (Figures 17b and 18).

Other kinds of effects such as steric interactions and ring strain also play a role in the ester stability, as functional groups attached or adjacent to the boron atom on a boronic acid decrease the association constant with diols.^{83,232,240} Steric hindrance is particularly relevant in polymers. Sumerlin et al. reported higher pK_a and lower binding affinity of boronic acidcontaining polymers compared to small molecules. The monomer pK_a was 7.2 whereas the polymer had a pK_a of 8.2. In the polymer, the low mobility of the chains limited the availability of the binding sites.^{87,232} Great care should be taken in the design of the exchangeable bond as the effect obtained can be opposite to the expected one in some cases. For instance, in a recent study, vitrimers based on nitrogencoordinating cyclic boronic diesters were synthesized, with a dative $B \rightarrow N$ bond embedded in the cyclic diester (Figure 19).²⁴¹ This particular structure not only offered an improved resistance to hydrolysis but also lowered the exchange rate compared to other cyclic boronic esters. In this particular case the effect on transesterification is somewhat counterintuitive. In summary, boronic esters in polymers allow the fine-tuning



Figure 17. (a) Example of coordinative B-N bond.²³⁹ (b) Slow exchange cross-linker and (c) fast exchange cross-linker (bottom) studied by Guan et al.44

Relaxation time = 5 min

of materials properties thanks to activating groups⁸⁴ with electronic and/or steric effects. This feature is interesting for vitrimers design. However, the knowledge on these effects in bulk remains scarce and deserves further investigations.²³⁶

Disulfide Metathesis: Effect of Aromatic Substituents. Neighboring group participation (NGP) in the exchange of disulfide bonds has hardly been probed yet, and the data readily available on this topic focus on the dissociative [2 + 1]radical assisted mechanism previously discussed in the Disulfides subsection (Figure 20). However, this general knowledge on disulfide bonds strength is useful to understand how to tune their dynamic behavior. Aromatic disulfides were mainly investigated, as the exchange reaction does not require the use of a catalyst to occur significantly contrary to exchange between alkyl disulfides usually catalyzed by triethylamine or tributyl phosphate, for instance.^{110–113,242,243} The metathesis of alkyl disulfides is a very slow reaction and even considered as not spontaneous in the absence of a catalyst, especially in solid state.¹¹⁵ Adding 6% of triethylamine is, however,

sufficient for the reaction to happen.²⁴⁴ Phosphines are also good catalyst candidates, and 5 mol % tricyclohexylphosphine was reported to accelerate the reaction 240 times in solution at room temperature, whereas the same amount of tri-(dimethylamino)phosphine accelerated it 520 times, although some degradation products were observed in that case.² Nevertheless, a catalyst-free self-healable hydrogel based on alkyl disulfide was reported.²⁴⁶ The network was formed by harnessing the thiol-disulfide exchange reaction of a telechelic thiol-ended PEG with telechelic PEG bearing cyclic fivemembered disulfide (Scheme 29). This network formation did not require any catalyst, as the tension of the five-membered disulfide ring favors its opening and enhances the exchange reaction. This design also allowed self-healing in slightly acidic condition, whereas such behavior is usually observed only in neutral and alkaline mediums, in the presence of nucleophilic thiolates.

Nitrogen has an enhancing effect¹¹⁰ in different kinds of activating groups such as tertiary amines on phenyl 1247 at 111247 at 1248 at 248thiuram disulfides,²⁴⁸ or sulfenamides.² ⁴³ For disulfides, instance aromatic sulfenamide-derived disulfides (Figure 21) exhibited a bond dissociation energy (BDE) of around 35 kcal mol⁻¹, whereas their nitrogen-free counterpart exhibited a BDE around 50 kcal mol^{-1.243} The lower is the BDE, the more radicals are generated, and the easier is the exchange reaction. Thanks to the enhancement of the exchange reaction, some catalyst-free materials based on aromatic disulfides exhibited self-healing at room temperature.¹¹⁶ In those materials bis(4aminophenyl) disulfides were used as dynamic cross-linkers to make poly(urethane-urea) systems.

4-Hydroxyphenyl disulfides are known to undergo faster radical generation than their 4-aminophenyl homologues,¹ but this effect is still poorly documented. 4-Hydroxyphenyl disulfide exchange reaction with diphenyl disulfide reached 60% conversion in only 2 h, whereas 12 h was necessary for the 4-aminophenyl disulfide with diphenyl disulfide under UV radiation. Matxain et al. published a thorough computational reactivity study of the disulfide bonds dissociation energy.¹¹⁰ They pointed out that electron-donating groups lower the disulfide BDE and that electron-withdrawing groups (EWG) increase this BDE. Compared to the 48 kcal mol⁻¹ BDE value of unsubstituted aromatic disulfides, the p-sulfonic acid substituent increased this value to 55.6 kcal mol⁻¹ whereas an amino derivative in the para position decreased the BDE to 41.3 kcal mol^{-1} and even to 30.1 kcal mol^{-1} with the trisubstituted bis(2,4,6-triaminephenyl) disulfide (Figure 22a).



Figure 18. Effect of amine activating groups on the relaxation time of vitrimers with different kinds of exchangeable bonds (NG stands for neighboring group and TEA for triethanolamine).



Figure 19. Nitrogen-coordinating cyclic boronic diester exchange.²⁴¹



Figure 20. Disulfide exchange enhancement by activating groups (activation energy from Jourdain et al.²⁴).

Scheme 29. Hydrogel Polymerization by Thiol–Disulfide Exchange on Strained Five-Membered Ring Disulfides²⁴⁶



Figure 21. Structures of aromatic sulfenamide-derived disulfides and the associated bond dissociation energies (BDE) calculated by DFT.²⁴³

They also reported that dendralene and thiuram disulfides (Figures 22b and 22c) have a low BDE, which may allow self-healing at room temperature and provide a good alternative to aryl disulfides. Though disulfide bonds have been known for a long time, there is still much research needed to understand

how to better tune the exchange reaction in materials, especially in the case of the associative thiol-disulfide reaction about which literature remains scarce.

Internal Catalysis of Silyl Ether Transalkoxylation by Amines, Urethanes, and Ureas. When implementing the



Figure 22. Structures of (a) bis(2,4,6-triaminephenyl) disulfide, (b) dendralene disulfide, and (c) thiuram disulfide.



Figure 23. Examples of activating groups enhancing silyl ether exchange and range of stress-relaxation experiment temperatures²⁵ and activation energies.²⁴

previously discussed ion pair dissociation-triggered mechanism in silyl ethers, the main parameters to tune the exchange reactivity are obviously the nature¹³¹ and concentration¹³⁰ of the ion pairs. The environment of the counterion influences its ability to dissociate and thus the rate of the exchange reaction (Figure 23). For instance, cation complexation by oxygen atoms on neighboring siloxane chains was reported to increase the electrophilic behavior of silicon atoms and the nucleophilic behavior of silanolate moieties, facilitating trans-siloxanation.¹³¹ In 1970, Wright et al. studied the equilibrium between chains and rings in $[R(CH_3)SiO]_r$ structures, showing that for x = 2 or 4 bulkier R led to favored cyclization. However, the opposite effect was shown for $x > 10.^{249}$ Catalysis by triphenylphosphine (TPP) and triazabicyclodecene (TBD) in silicone materials was also reported. TPP led to materials with a temperature of plastic flow around 240 °C whereas the materials containing TBD flowed at 120 °C.²⁵⁰ In 2014, tertiary and secondary amines, urea, and carbamate groups on α - and γ -silanes (Figure 24) were reported to interact with the alkoxy leaving group or the attacking nucleophile, thus increasing the hydrolysis rate of the alkoxy group. This study performed both in solution and in silico showed the significant influence of several activating groups. For instance, secondary amines accelerated the hydrolysis reaction from 5 to 50 times and tertiary amines at least 40 times. Carbamate groups accelerated the reaction from 6 to 80 times, depending on their position and structure.²⁵¹

This result was later successfully implemented in malleable materials with activating amino moieties accelerating the exchange rate of silyl ethers.^{45,252} A copolymer of styrene and 2-((4-vinylbenzyl)oxy)ethan-1-ol was synthesized by free radical copolymerization. Then the resulting copolymer was cross-linked either with N,N'-bis(3-(trimethoxysilyl)propyl)-ethane-1,2-diamine or with 1,10-bis(trichlorosilyl)decane (Scheme 30). Both cross-linked structures were similar, and the cross-linkers used had the same length. The relaxation time for the structure incorporating secondary amines was 260 s at 180 °C, 3 times less than the 779 s needed for the amine-free structure (Scheme 30).⁴⁵ Variations of the silyl ether bulkiness



Figure 24. Structure of (a) α -silanes and (b) γ -silanes. The effects of various -X substituents on the hydrolysis rate of the alkoxy group were studied (c) in basic conditions and (d) in acidic conditions.²⁵¹

Scheme 30. Synthesis of Styrene-Derived Networks Cross-Linked by Silyl Ethers^a



Relaxation time: 779 s (180 °C)

 a Two cross-linkers were used, one bearing secondary amines (top) and the other not (bottom). The reprocessabilities of the obtained materials were compared.⁴⁵



60 kJ.mol⁻¹

Figure 25. Characteristics of polyurethanes, polyhydroxyurethanes, polythiourethanes, and poly(oxime-urethanes): range of stress-relaxation experiments temperatures²⁵ or reprocessing temperatures and activation energies.^{24,253}

and the polymer backbone are suggested to tune the properties of these materials. $^{\rm 132}$

Case of Urethanes and Hydroxyurethanes. As discussed previously, the mechanism involved in polyhydroxyurethane exchange reaction remains uncertain. Hence, the determination of the effects of activating groups on transcarbamoylation is intricate as they were mainly studied for the dissociative decomposition reaction discussed in section I (Figure 25).^{254–256} It should be mentioned that an important activation reported is the oxime-promoted transcarbamoylation, similar to the effect on esters previously discussed. It was implemented in dissociative covalent adaptable networks since the mechanism is purely dissociative in this case.²⁵³

For thiourethanes solvolysis by thiols, secondary amines or alcohols, the suggested reaction pathway was an associative addition/elimination mechanism²⁵⁷ instead of the dissociative elimination/addition commonly accepted. Nonetheless, as previously pointed out, the recent implementation of thiourethane bonds in materials highlighted a dual associative and dissociative mechanism.²⁵⁸ Knowledge on the effects of activating groups on hydroxyurethane-based vitrimers is very limited. Nevertheless, an excess of free exchanging dangling groups, namely alcohol groups for polyurethanes and thiol groups for polythiourethanes, was shown to favor the associative pathway and decrease degradation reactions. Furthermore, this excess of dangling exchanging groups allows a better control over the reprocessing temperature and shorter reprocessing time. 144,258 After 70 min at 140 $^\circ C$ a PU network without free -OH group exhibited 83% recovery of cross-link density in the presence of 1 mol % dibutyltin dilaurate (DBTDL) as catalyst. In contrast, only 15 min at 120 °C was sufficient to reprocess a PU containing 20% of free -OH.²⁵⁸ In contrast, an increase of the relaxation time was observed in the case of thiourethane-based networks (Scheme 11) containing free thiols compared to the thiol-free analogue. This counterintuitive result was explained by a change from a dissociative to a slower associative mechanism.¹⁴⁴ In conclusion, the chemistry of polyhydroxyurethanes is much more complex that it seems at first glance. Dissociative and associative exchange mechanisms compete and are influenced by many parameters such as the availability of exchangeable group, the presence of activating group or internal catalysts, or the temperature. Therefore, much caution should be taken when designing covalent adaptable networks out of polyhydroxyurethanes.

Ureas Exchange. Recently, a polyurethane—urea elastomer was claimed to be a vitrimer.²⁵⁹ The authors first synthesized a PU prepolymer out of poly(oxatetramethylene) glycol (PTMG) and 4,4'-diphenylmethane diisocyanate (MDI). Then this prepolymer was mixed either with 3,5-dimethylth-

Scheme 31. Synthesis of Polyurethane–Urea Self-Healable Elastomers²⁵⁹



io-2,4-toluenediamine (DMTDA) or with 3,3'-dichloro-4,4'diaminodiphenylmethane (MOCA) to form exchangeable biuret groups (N,N'-diarylurea) (Scheme 31). Because the diamines used are primary, each amine can react twice, thus ensuring the formation of a cross-linked materials (primary diamines can be considered as hidden tetrafunctional reactants). The amines used possessed either electron-donating methylthio substituents or electron-withdrawing chlorine atoms. The stress–relaxation time of the methylthio groupcontaining network was 3–4 times shorter than that of the chlorinated analogue, depending on the temperature (100– 120 °C). The relaxation was assigned to urea–urea exchanges, and the mechanism was reported as associative.²⁵⁹ Nevertheless, there is little proof to support this claim.

Transimination Acceleration by Activating Groups. The success of imine-derived vitrimers (Figure 26) may be



Figure 26. Characteristics of transimination: reported range of stress–relaxation experiments temperatures²⁵ and activation energies.²⁶⁰ Two examples of potentially active neighboring groups are given.

explained by the versatility of this linkage but also because these vitrimers can be reprocessed without the need for a catalyst.¹⁶³⁻¹⁷⁴ When used in organic chemistry or biochem-

istry, transimination was reported to be catalyzed by thioureametal complexes (2 orders of magnitude rate acceleration),¹⁷⁷ triflate salts of scandium, terbium, samarium, and other lanthanides (up to 5 orders of magnitude acceleration),^{178,261} zinc bromide,²⁶² iron chloride, copper chloride, and magnesium chloride.²⁶³ Besides, acidic and basic catalyses are known to be efficient as well in aqueous^{175,264} or organic solutions.^{175,265} In particular, interesting examples of internal basic catalysis in solution were reported in the 1980s by using primary amines featuring an internal tertiary amine func-tion.^{264,265} Ciaccia et al.¹⁸⁰ reported a study on aromatic imines in organic solvents at room temperature. Part of this work dealt with the exchange between N-benzylideneanilines and *p*-toluidine (Figure 27). The exchange reaction exhibited low equilibrium constants ($K_{eq} < 10$); thus, the results could have been carefully extrapolated to degenerate system such as those encountered in vitrimers. The authors showed that the rate of the forward transimination reaction was almost unaffected by EWG and EDG on the imine aromatic ring but that the reverse reaction was mildly affected by these substituents. The electron-donating -OCH₃ group slightly increased the reverse reaction rate, whereas electron-withdrawing -CN and -NO2 groups decreased this rate (Figure 27). Similarly, Schaufelberger et al.²⁶² studied the impact of substituents on the exchange of aromatic imines with benzylamines in the presence of zinc bromide as catalyst in acetonitrile. Nevertheless, in this system, two simultaneous reactions were involved: imine isomerization and transimination. Thus, it would be dubious to draw general conclusions on internal catalysis based on this study.

Because the first step of the amine—imine exchange reaction is associative, it occurs at room temperature without catalyst in solvents only with unhindered reactants.¹⁷⁹ This idea underlines the fact that the environment of the exchangeable bond likely plays a major role in the reaction. For instance, the solvent used to make dynamic covalent gels had an influence—

сн, ,	Х	К	k _d (M ⁻¹ s ⁻¹)	k _r (M ⁻¹ s ⁻¹)
Срана Ср	Н	2.6	1.0 × 10 ⁻¹	3.8 × 10 ⁻²
· · · · · · · · · · · · · · · · · · ·	OCH ₃	3.0	1.2×10^{-1}	4.0 × 10 ⁻²
	CN	4.5	1.1 × 10 ⁻¹	2.4 × 10 ⁻²
Ŷ Ŷ	NO ₂	7.2	0.94×10^{-1}	1.3×10^{-2}

Figure 27. Transimination reaction between N-benzylideneanilines and p-toluidine and kinetic parameters studied by Ciaccia et al.¹⁸⁰

acetonitrile allowing faster imine exchange than toluene.²⁶⁶ Additionally, imine metathesis is catalyzed by zirconium, molybdenum, rhenium, and niobium in organic solvents if the temperature reaches $100 \ ^{\circ}C^{179}$ and by hafnium imido complexes at $80 \ ^{\circ}C$ as well.²⁶⁷ An interesting strategy involving minute amounts of primary amines was also reported to accelerate this reaction in solution at room temperature.¹⁷⁹ The effect of primary amines on imine metathesis was investigated in terms of mechanism and thermodynamics. Mechanistically speaking, the overall reaction can be seen as two concomitant transimination reactions (Scheme 32). Thus,

Scheme 32. Imine Metathesis Catalysis by Minute Amounts of Primary Amines $^{179}\,$



it is not *stricto sensu* an imine metathesis, but the overall result is the same. Minute amounts of an imine A bearing an isobutyl group exchange with the minute amount of n-butylamine, leading to the release of minute amounts of the exchanged imine C and isobutylamine. This amine then undergoes transimination with an imine B bearing a n-butyl group. This second step releases the exchanged imine D and n-butylamine able to react with A and so on. The overall reaction is an equilibrium between amines A and B, on one hand, and between C and D, on the other. Thermodynamically speaking, the overall reaction is an imine metathesis.¹⁷⁹ This effect was implemented in imine-cross-linked PEG networks. A slight excess of dangling primary amines enabled self-healing at 50 °C within 5 min.²⁶⁶ Thus, choosing carefully the stoichiometry for the synthesis of imine-derived vitrimers would enable the tuning of their properties.

Recently the effect of hydrogen bonds was also emphasized. Indeed, hydrogen bond donors such as thioureas and squaramides catalyzed both imine metathesis and transimination in solution and their efficiencies depended on their structures.²⁶⁸ Similarly imines derived from salicylaldehyde were reported to be stabilized by hydrogen bonding with -OH or -NHR substituents on the ortho position.²⁶⁹ Once again, this effect could be used in the molecular design of imines to tune the exchangeable bond properties.

Vinylogous Urethanes and Amides, an Efficient Exchange Chemistry without Catalyst. Transamination of vinylogous urethane is a very effective exchange reaction and enables chemists to design easily reprocessable catalyst-free vitrimers (Schemes 16 and 33). This observation might be the cause for the rarity of reports on such effects on vinylogous urethanes. Although the exchange is so efficient that many catalyst-free vitrimers were synthesized, vinylogous urethanebased CANs would still benefit from activating group participation or internal catalysis to lower reprocessing temperatures. Actually, an example of internal catalysis was recently reported in vinylogous urethane-cross-linked epoxy-amine materials.¹⁹⁹ Tri- and difunctional amines were reacted with a bisacetoacetate to prepare a vinylogous urethane bondcontaining hardener for an industrial epoxy resin. Two synthesis methods were compared. On the one hand, the amines, the bisacetoacetate, and the epoxy resin were mixed in one pot. On the other, the amines and the bisacetoacetate reacted in a first step to yield a vinylogous urethane curing agent bearing free primary amines. Then the curing agent was mixed with the epoxy resin in a second step (Figure 28). The difference lies in the water release when amines and the bisacetoacetate reacted together. In the one-pot strategy, this water was trapped in the material, whereas in the two-step strategy the water produced during the first step was removed prior to the second step to yield a water-free material. The materials exhibited a relaxation time at 160 °C of 3.5 s for the one-pot material and of 5.5 s for the two-step synthesis, showing no significant difference between the two preparation procedures. These values must be compared with the 25.5 s relaxation time observed in epoxy-free vinylogous urethanes vitrimers with a similar structure, although having a higher vinylogous urethane bond concentration. Certainly, the network structure influences the relaxation time, and thus direct comparison of these values might be adventurous, but an internal catalytic effect from the dangling hydroxyl groups is likely responsible for the shorter relaxation times.¹⁹⁹ To

Scheme 33. Characteristics of Vinylogous Urethane Transamination: Reported Range of Stress-Relaxation Experiment Temperatures²⁵ and Activation Energies;²⁴ Dangling Hydroxyl Groups Accelerate the Relaxation Time of VU Materials



80-220 °C / E_a 31-129 kJ.mol⁻¹



Figure 28. Two strategies to synthesize VU networks were reported by Spiesschaert et al.¹⁹⁹ The one-pot strategy traps the water released by the reaction. The two-step strategy allows to remove water from the VU curing agent after the first step, allowing water-free network. TREN = tris(2-aminoethyl)amine; BAC = 1,3-bis(aminomethyl)cyclohexane; EG-AA = ethylene glycol-bisacetoacetate.

support this idea, the authors performed model reaction on small molecules. They studied the disappearance of propyl-3-(octylamino)but-2-enoate (octyl VU) with benzylamine in 2hexanol or dodecane at 100 °C. The reaction was faster in 2hexanol, showing the accelerating effect of hydroxyl moieties on the exchange reaction, independently of any matrix effect. As the study of vinylogous urethane vitrimers is popular, numerous tuning possibilities might emerge in the future not only by implementation of internal catalysis in these networks but also by variation of the backbones nature and length.

Hints for Diketoenamine Activation. As previously mentioned, diketoenamines were essentially studied as protecting groups, and there is no extensive knowledge on a potential effect of neighboring groups. Nevertheless, in the work of Augustyns et al.²⁰¹ already discussed earlier, a few interesting trends on reactivity were gathered. The two amino groups on lysine did not have the same reactivity toward the exchange of the Dde group. Unprotected ε -NH₂ groups could exchange with protected ε -NH₂ and α -NH₂ groups, but free α - NH_2 groups could not exchange Dde with ε - NH_2 . This was probably because α -NH₂ groups were too sterically hindered and not nucleophilic enough due to the neighboring electronwithdrawing carboxylic acid. In addition, when 2-acetyldimedone was reacted with polyamine, primary amines were selectively protected.²⁰¹ The use of diketoenamines in vitrimers is just starting to emerge, and there is undoubtedly a lot to discover on this chemistry based on the encouraging proof of concept recently disclosed.^{204,205}

Effect of the Alkylating Agent on Trialkylsulfonium Salt Exchange. As mentioned before, the knowledge on trialkylsulfonium exchange with thioethers is poor except for the proof-of-concept disclosed by Du Prez's group.⁴⁰ Of course, the most obvious parameters to tune the material reprocessability are the nature and quantity of the alkylating agent. Without an alkylating agent, no stress–relaxation was observed in polythioether.⁴⁰ Conversely increasing the concentration of the butyl brosylate alkylating agent from 1 to 10 mol % decreased relaxation time at 150 °C from 75 to 10 min.⁴⁰ The same behavior was observed by using trimethylsulfonium iodide as an alkylating agent.¹²⁴ Once again, there might be room for internal catalysis studies with this kind of chemistry. Indeed, this exchange reaction was studied on small arylalkyl sulfides in solution. If no change in the kinetics was observed with electron-rich aromatic groups, electron-poor aromatic groups slowed the conversion rate. The alkyl substituent bulkiness also played a role. Phenylalkyl sulfides bearing various alkyl groups were transalkylated with methyl iodide. tert-Butylphenyl sulfides reached full conversion in 25 h, whereas the butyl-, 3-pentyl- and 2,2-dimethylpropyl analogues only reached 70, 60, and 8% conversion, respectively. Diphenyl sulfide showed no conversion at all.²⁷⁰ This effect has never been studied in materials but underlines the importance of the exchangeable group and might bring useful insights. These articles also underline the effect of the counteranions which nucleophilicity likely plays a role as seen in the transalkylation of triazolium salts.²⁷¹⁻²

CONCLUSION AND OUTLOOK

Since the invention of vitrimers ten years ago, this class of organic materials has triggered an undeniable enthusiasm in the polymer scientists' community. These materials have not only aroused scientists' curiosity, but their implications for industrial applications are also potentially transformative. Indeed, the rheological behavior of these materials allows a precise control of the viscosity upon heating. This property is central to processing and reprocessing issues. One drawback of these materials is that many vitrimers require catalysts, sometimes at high loadings, which raise concerns for the risks of leaching and premature aging of the materials. Consequently, further work on vitrimers will likely focus on designing catalyst-free vitrimers and on the tuning of the vitrimers' properties, especially the reshaping ability, for the associative systems already known. Parameters such as the cross-linking density, the concentration and availability of exchanging moieties, or the structure and chemical groups of the backbone are already known to influence the behavior of vitrimers. A few studies on the effect of activating groups on the exchange reactions and consequently on materials reshaping are already available. Nonetheless, the use of this strategy to tune the rheologic profile of associative CANs is still in infancy. Beyond the sole tuning of the reshaping parameters, the study of activating group and internal catalysis would be significant progress for understanding how the exchange reactions in vitrimers works, and it would give access to the scientists' community to more intimate knowledge of this innovative class of polymer materials. At the moment knowledge on the phenomena occurring at the molecular scale in vitrimers is still poor. In addition, the mechanistic studies on the exchange mechanisms involved are sometimes only preliminary or even inexistent. Moreover, activating groups can have an influence on the exchange mechanism at work in CANs. They may change an associative exchange into a dissociative one. Mechanistic studies are thus fundamental for the development of CANs. A better comprehension of the phenomena at the molecular scale would indeed allow scientists to design materials with original properties by exploiting such mechanism changes. Broadening the pool of techniques used to characterize CANs would also undoubtedly help to understand their behavior and contribute to a better and more purposeful design. For example, molecular mobility studies using dielectric spectroscopy or dynamic mechanical analysis could enrich knowledge on these fascinating materials.

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Biographies

Florian Cuminet was born in 1996 in Avranches, France. He spent two years at the National Graduate School of Chemistry of Rennes, France (ENSCR), and then three years at the National Graduate School of Chemistry of Montpellier, France (ENSCM), where he was awarded his master's degree in Materials Chemistry in September 2019. He achieved a master's thesis supervised by R. Tavernier on novel biobased benzoxazines at the Institute Charles Gerhardt of Montpellier (ICGM). In 2019, he started a PhD at the ICGM and the Interuniversity Center of Materials Research and Engineering in Toulouse, France (CIRIMAT), under the supervision of Drs. Caillol, Dantras, Ladmiral, and Leclerc. His PhD project, funded by the Chimie Balard Cirimat Carnot Institute, aims at the synthesis and characterization of catalyst-free vitrimers.

Sylvain Caillol is Research Director at CNRS. He was born in 1974. He graduated engineer from the National Graduate School of Chemistry of Montpellier in 1998 and then received his MSc degree in Chemistry from the University of Montpellier. He received his PhD degree in 2001 from the University of Bordeaux. Subsequently he joined Rhodia Company. Later, promoted Department Manager, he headed the Polymer Research Department in the Research Center of Aubervilliers. In 2007, he joined the CNRS at the Institute Charles Gerhardt of the University of Montpellier where he started a new research topic dedicated to green chemistry and biobased polymers. He is coauthor of more than 200 articles, patents, and book chapters. He is Chairman of the Oleochemistry division of European Federation of Lipids. He won the Green Materials Prize in 2018 and 2020 and was awarded Pioneering Investigator by the Royal Society of Chemistry in 2019.

Éric Dantras defended his PhD thesis in 2002, entitled "Molecular mobility analysis of phosphorus containing dendrimers by dynamic and thermostimulated dielectric spectroscopy", under the supervision of Prof. C. Lacabanne. He is now associate professor at the Université Paul Sabatier, Toulouse, France. His research activity is devoted to polymer physics and the analysis of polymer chain dynamic in correlation with their dielectric, mechanical, and thermal properties.

Éric Leclerc received his PhD under the supervision of Dr. P. Mangeney at the Université Pierre et Marie Curie (Paris 6) in 2001. After a postdoctoral stay at the University of Hawaii in the group of Prof. Marcus A. Tius, he was appointed CNRS Research Fellow in 2003 at the INSA & Université de Rouen. In 2011, he moved to the Ecole Nationale Supérieure de Chimie de Montpellier to join the group of Prof. J.-M. Campagne. His research interests mainly focus on the synthesis of fluorinated building blocks and biomolecules, the development of new organo- or metallo-catalyzed reactions, and the application of original small molecules to the development of organic materials.

Vincent Ladmiral graduated from the National Graduate School of Chemistry of Montpellier in 1998 and received his PhD from the University of Warwick in 2006. After postdoctoral fellowships at the University of Kyoto, the University of Sydney, the University of New South Wales, and the University of Sheffield, he was appointed CNRS Research Fellow in 2012 in Institut Charles Gerhardt of Montpellier. His research interests focus on the structure—property relationship of polymers.

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