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Pierre-Luc Durand, Antoine Brège, Guillaume Chollet, Etienne Grau, Henri Cramail. A Simple and Efficient Approach toward Photosensitive Biobased Aliphatic Polycarbonate Materials. ACS Macro Letters, Washington, D.C: American Chemical Society, 2018, 7 (2), pp.250-254. 10.1021/acsmacro-lett.8b00003. hal-01917959

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A simple and efficient approach towards photo-sensitive biobased aliphatic polycarbonate materials

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ABSTRACT: Fatty acids were used as precursors for the synthesis of photo-sensitive polycarbonate materials. In order to avoid multi-step reactions, a simple and straightforward methodology toward the synthesis of photo-sensitive monomers has been developed. Hence, a fatty acid-based cyclic carbonate bearing an unsaturation was synthesized and subsequently polymerized in a controlled manner (Đ=1,07) by organo-catalyzed ring-opening polymerization (ROP). A thio-cinnamate derivative was then readily synthesized *via* a one-pot reaction and grafted onto the polycarbonate backbone by thiol-ene reaction. The content of photo-responsive cinnamoyl moiety grafted on the polycarbonate was tunable with the reaction time. Such functionalized polycarbonates could be cross-linked (by UV irradiation at 365 nm) and de-cross-linked (irradiated at 254nm) and exhibit versatile properties ranging from rather tough materials to elastomeric networks with respect to the content of the photo-sensitive cinnamoyl moiety grafted on the polymer.

Aliphatic polycarbonates are well known for their specific features such as low T_g, resistance towards hydrolysis, biocompatibility and biodegradability.1 Their synthesis can be achieved via different routes but such polymers are mainly synthesized through ring-opening polymerization (ROP) of cyclic carbonate monomers.² Under suitable conditions, the polymer chain length, dispersity and microstructure as well as the nature of end-groups can be controlled. The advantage of designing functional polycarbonates over the traditional PTMC stems from the modulation of their physico-chemical properties for specific needs, thereby broadening and improving their performance characteristics. Functional polycarbonates can be synthesized either upon direct polymerization of functional monomer or upon chemical modification postpolymerization.^{3,4} Moreover, because of environmental concerns and also in view of searching novel functionalities, the use of building blocks from renewable resources such as vegetable oils is of great interest.⁵ In this purpose and adapted from the work of Venkataraman and coll.,⁶ fatty acid-based cyclic carbonates were synthesized as precursors of original aliphatic bio-based polycarbonates.

In addition, polycarbonate networks exhibiting elastomeric properties are desirable thanks to a large number of applications in the biomedical area, especially in the emerging field of soft-tissue engineering or drug delivery.⁷⁻¹² Thus, several research groups have investigated various cross-linking methods affording polycarbonate materials.¹³⁻¹⁷ However, the main drawback of such crosslinked materials is their inability to be reshaped or recycled. As a consequence, new self-healing materials involving reversible cross-linking reactions have been developed.¹⁸⁻²¹ Such materials could be of interest in many fields such as protective coatings, biomedical applications, piping and electronics.¹⁹

Among all stimuli that can be employed to activate a reversible transformation, light is one of the most used.^{22,23} In the midst of photosensitive moieties, the photo-reactive cinnamate group dimerization, occurring through the [2+2] photochemical cyclo-addition reaction, was implemented for polymerization reactions and for cross-linking purposes.²⁴ Using the photo-sensitive cinnamate group, the reversibility of the cross-linking was, for instance, exploited for the synthesis of self-healing polymers²⁵⁻²⁷ and tunable shape-memory materials.²⁸ However, a challenge in the field of self-healing polymers is their ease of synthesis. The photo-sensitive moieties are mainly brought through copolymerization of the functional monomer, the latter needing several synthetic steps.²⁴⁻²⁶

Herein, a simple and efficient post-functionalization method involving a thiol-ene coupling is developed towards photo-sensitive aliphatic polycarbonate networks. After synthesizing a bio-based aliphatic polycarbonate bearing pendant unsaturations, several cinnamatecontaining polycarbonates were easily prepared and reversibly cross-linked through UV irradiation. The influence of the cinnamate content on the network properties was also studied.



i: TBD, 80°C, 7h; *ii*: ethyl chloroformate, triethylamine, THF, RT, overnight; *iii*: DBU/TUS, BnOH, DCM, RT, 4h; *iv*: toluene, reflux, 2h; *v*: v₇₀, DCM, 40°C.

The fatty-acid based cyclic carbonate was obtained in two steps. First is the formation of the amide functionalized **NH-Und-1,3-diol** by coupling the methyl undecenoate and 2-amino-1,3-propanediol. This amidation is performed in bulk at 80°C, under nitrogen to remove the methanol by-product, in the presence of 1,5,7triazabicyclo[4.4.0]dec-5-ene (TBD) as an organo-catalyst (Yield: 70%., see Figs. S1-S4). The second step is the intramolecular cyclization giving the corresponding 6membered ring cyclic carbonate **NH-Und-6CC** by using ethylchloroformate in THF in the presence of triethylamine as a base (Yield: 55%., see Figs. S5-S8).

The polymerization of **NH-Und-6CC** was carried out in DCM ($[M]_o = 2.o M$) at room temperature with benzyl alcohol (BnOH) as initiator and $[M]_o/[BnOH]_o = 5o$ (**Erreur ! Source du renvoi introuvable.**). DBU was chosen as catalyst associated with the Schreiner thiourea (TUS) as co-catalyst with a ratio of $[M]_o/[BnOH]_o/[DBU]_o/[TUS]_o = 50/1/1/1$. P(**NH-Und-6CC**) was successfully obtained with yield up to 90% (DP= 5o; D=1.07, see Fig S9).

The **cinnamate-SH** moiety was first synthesized through the direct and simple reaction of cinnamic acid chloride with the 2-mercaptoethanol as illustrated in **Erreur** ! **Source du renvoi introuvable.**. This reaction was performed in stoichiometric conditions with complete conversion and very high (yield: 95%, see Figs. Sto-St3). The next step towards the design of photo-responsive polycarbonate material is the grafting of **Cinnamoyl-SH** on P(**NH-Und-6CC**). As above-mentioned, the thiol-ene coupling reaction was used to link these two entities. Nevertheless, since **Cinnamoyl-SH** is sensitive to UV irradiation, a radical initiator which decomposes thermally (V70) was used to initiate the thiol-ene reaction as depicted in **Erreur** ! **Source du renvoi introuvable.**.

Syntheses of polycarbonates bearing different amount of pendant cinnamate moieties were carried out (Table 1). The cinnamate content was controlled by the reaction time, see Fig. S15. As expected, an increase of the polymer molar mass values associated to a decrease of the T_g (23°C to -20°C) was observed with the cinnamoyl content grafted on the polymer, see Fig. S16. Such a decrease is ex-

plained by the flexibility introduced by the sulphur atom to the polymer chains.

Polycarbonate networks were obtained by UV crosslinking using the reactivity of the cinnamate moieties. With UV light at $\lambda = 365$ nm, the cinnamate group switches from the trans to the cis conformation. When two cis cinnamate groups react together, they undergo a [2+2] cyclo-addition reaction leading to the formation of a cyclobutane ring (**Erreur** ! **Source du renvoi introuvable.**). This reaction was used to cross-link all cinnamate-functionalized cx-P(**NH-Und-6CC**).

The curing kinetics was studied using UV–vis spectrophotometry by monitoring the absorption maximum at 280 nm, which relates to the double bonds adjacent to carbonyl group of the cinnamate functionality (Figure 1-(a)). As seen in the UV–vis spectrum, a decrease in intensity of the peaks at 280 nm was observed within 6 h, meaning that almost 100% of the cinnamate groups have undergone the [2+2] cyclo-addition reaction. The fast initial absorbance decrease is characteristic of the polymer cross-linking through irradiation because the concentration of cinnamate groups is higher in the initial stages. As the cross-linking increases, concentration and mobility of the chains decreases leading to a plateau.

Scheme 2. Photo-reversible cross-linking of a cinnamate containing bio-based polycarbonate



Subsequently, upon exposing the cross-linked polycarbonate to 254 nm light irradiation, the de-cross-linking of the polycarbonate material occurs.

Polymer	Reaction time (min)	Cinnamoyl content ^a (mol. %)	$M_n{}^b(g.mol{}^{-1})$	[Ð] ^b	T _g (°C) ^c
P(NH-Und-6CC)	0	0	5 700	1.07	23
c ₁₀ -P(NH-Und-6CC)	0.5	10	5 800	1.10	11
c ₃₀ -P(NH-Und-6CC)	2	30	5 900	1.18	5
c ₅₅ -P(NH-Und-6CC)	6	55	6 500	1.25	-1
c ₇₀ -P(NH-Und-6CC)	15	70	8 500	1.29	-12
c100-P(NH-Und-6CC)	30	100	9 400	1.31	-20

Table 1. Molecular	characterization (of cinnamate-c	ontaining pol	vcarbonates
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^a: Determined by ¹H NMR; ^b: Determined by SEC in THF (PS Std); ^c: Determined by DSC at 10°C.min⁻¹ from the second cycle

The de-cross-linking reaction is relatively fast but the absorbance doesn't reach its former value before cross-linking indicating that the de-cross-linking is not complete probably due to the poor power of the UV lamp at this wavelength. In order to have a total decross-linking, an appropriate UV lamp must be used, allowing a proper irradiation below 300 nm. Nevertheless, Figure 1 demonstrates the reversible feature of the photo-mediated cross-linked polycarbonate.



Figure 1. a) Cross-linking kinetic under 365 nm and b) decross-linking kinetic under 254 nm in solution (DCM)

All functional polycarbonates were cross-linked by the [2+2] photochemical cyclo-addition of cinnamoyl units to form cross-linked polycarbonates (CL-cx-P(NH-Und-6CC)). First, to evaluate the network cross-linking density, swelling tests have been performed. It can be noted from Table 2 that very low gel content (7%) was observed when only 10 mol.% of cinnamoyl units were grafted to P(NH-Und-6CC). The poor photoreactivity in a solidstate coated film explains this low gel content. However, the gel content increases significantly when the pendent cinammoyl content increases in the polymer to almost reach 100%. At the same time, the swelling ratio decreases when the cross-linking increases. This feature shows that the cross-linking density can be controlled by adjusting the polycarbonate cinnamoyl content. After crosslinking, thermal and mechanical chacterizations of these networks were investigated by differential scanning calorimetry (DSC, Figs. S16-17), thermogravimetry (TGA, Fig. S18), tensile tests experiments (Fig. S19) and dynamic mechanical analysis (Fig. S20-21). All the results are summarized in Table 2. Such cross-linked materials demonstrated slight increase in Tg with an increase of crosslinking density. More interestingly, mechanical properties of the networks strongly depend on the cross-linking density. Indeed, CL-c10-P(NH-Und-6CC) and CL-c30-P(NH-Und-6CC) are transparent and flexible materials at room temperature as shown in Figure 2. However, when high cinnamovl content was used, the materials became more rigid. Consequently, the fully functionalized crosslinked polymer was very brittle. In addition, the influence of the cross-linking density on the polycarbonate mechanical properties can be clearly seen. CL-c10-P(NH-Und-6CC) displays very low Young modulus calculated from the initial slope of the stress-strain curve (1.3 MPa) while CL-c100-P(NH-Und-6CC) exhibits a Young modulus 1 000 times higher. In between these values, the higher the cross-linking density, the higher the Young modulus. Max stress at break values follows the same trend. However, the elongation at break follows opposite tendency. Therefore, the mechanical properties of the polycarbonate networks are significantly affected by the cinnamoyl content grafted on P(NH-Und-6CC).

The elastic modulus measured above T_g by DMA exhibited an increase up to 9.5 MPa for the most highly crosslinked polycarbonate. Moreover, in agreement with the results obtained by DSC, we observe that the T_{alpha} increases with the increase of the cross-linking density.



Figure 2. Pictures of flexible CL-c₃₀-P(NH-Und-6CC)

In conclusion, several cinnamate-containing polycarbonates were synthesized *via* the simple and controllable thiol-ene reaction between P(NH-Und-6CC) and thiofunctionalized cinnamate moiety. Photo-reversible polycarbonate networks were prepared thanks to the photoinduced [2+2] cyclo-addition reaction between two cinnamoyl groups.

Table 2. Summary of mechanical properties exhibiting by photo-mediated cross-linked polycarbonates

Network	T _g network (°C)ª	Modulus (MPa) ^b	Elongation at break (%) ^b	Max stress (MPa) ^b	Gel content (%)	Swelling ratio (%)
CL-c ₁₀ -P(NH-Und-6CC)	8	1.3 ± 0.5	156 ± 15	0.5 ± 0.2	7	69
CL-c ₃₀ -P(NH-Und-6CC)	11	39 ± 7	49 ± 8	2.4 ± 0.7	64	153
CL-c ₅₅ -P(NH-Und-6CC)	13	110 ± 21	15 ± 4	6 ± 1	85	62
CL-c ₇₀ -P(NH-Und-6CC)	24	622 ± 50	6 ± 3	29 ± 5	99	48
CL-c100-P(NH-Und-6CC)	17	1266 ± 55	3 ± 1	35 ± 5	95	57

^a: Determined by DSC at 10°C.min⁻¹ from the second cycle; ^b: Calculated from tensile tests

Among the advantages of this synthetic approach, is the use of the cinnamate moiety to control the cross-linking density in time and space *via* UV light without requiring any initiators or chemical cross-linkers that can remain in the final polymer. Thanks to the reversibility of the crosslinking reaction (proved by UV-Vis spectrophotometry), such materials could be reshaped or recycled with appropriate UV lamp.

ASSIOCATED CONTENT

Supporting Information. Figures, experimental procedures, full characterization data and copies of relevant NMR (functional monomers and polymers).

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ACKNOWLEDGMENTS

This work was performed, in partnership with the SAS PIVERT, within the frame of the French Institute for the Energy Transition (Institut pour la Transition Energétique - ITE) P.I.V.E.R.T. (www.institut-pivert.com) selected as an Investment for the Future ("Investissements d'Avenir"). This work was supported, as part of the Investments for the Future, by the French Government under the reference ANR-001-01. The authors thank Equipex Xyloforest ANR-10-EQPX-16. The financial support from the CPER CAMPUSB project funded by the French state and the Région Nouvelle Aquitaine is gratefully acknowledged

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Supporting Information

A simple and efficient approach towards photo-sensitive bio-based

aliphatic polycarbonate materials

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Experimental Section

Materials.

Ethyl chloroformate (97%), 2-mercaptoethanol (98%), benzyl alcohol (BnOH), benzoic acid, 1,5,7-triazabicyclodec-5ene (TBD, 98%), sodium sulfate (Na₂SO₄), magnesium sulfate (MgSO₄), 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU) and 1,3-bis[3,5-bis(trifluoromethyl)phenyl]thiourea (Schreiner TU, >98.0%) were obtained from Sigma-Aldrich. Triethylamine (Et₃N, 99%) was purchased from Alfa Aesar. Methyl 10-undecenoate was obtained from Nu-Chek Prep, Inc. Cinnamoyl chloride (predominantly trans, 98%) was purchased from TCI Europe. 2,2'-Azobis(4-methoxy-2,4dimethylvaleronitrile) (V70) was obtained from Wako.

ITERG kindly provided NH-Und-1,3-diol.

All products and solvents (reagent grade) were used as received except otherwise mentioned. The solvents were of reagent grade quality and were purified wherever necessary according to the methods reported in the literature. Flash chromatography was performed on a Grace Reveleris apparatus, employing silica cartridges from Grace. Cyclohexane: ethyl acetate and dichloromethane: methanol gradients were used as eluents depending on the products. The detection was performed through ELSD and UV detectors at 254 nm and 280 nm.

Characterization.

¹H and ¹³C-NMR spectra were recorded on Bruker Avance 400 spectrometer (400.20 MHz or 400.33 MHz and 100.63 MHz for ¹H and ¹³C, respectively) by using CDCl₃ as a solvent at room temperature, except otherwise mentioned. Two-dimensional analyses such as ¹H-¹H COSY (COrrelation SpectroscopY), ¹H-¹³C HSQC (Heteronuclear Single Quantum Spectroscopy) and were also performed. Size Exclusion Chromatography (SEC) analyses were performed in THF (25°C) on a PL GPC50 and with four TSK columns: HXL-L (guard column), G4000HXL (particles of 5 mm, pore size of 200A, and exclusion limit of 400000 g/mol), G3000HXL (particles of 5 mm, pore size of 75A, and exclusion limit of 60000 g/mol), G2000HXL (particles of 5 mm, pore size of 20 A, and exclusion limit of 10000 g/mol) at an elution rate of 1 mL/min. The elution times of the filtered samples were monitored using UV and RI detectors and SEC were calibrated using polystyrene standards. Differential scanning calorimetry (DSC) thermograms were measured using a DSC Q100 apparatus from TA instruments. For each sample, two cycles from -80 to 100°C at 10°C.min⁻¹ were performed and then the glass transition and melting temperatures were calculated from the second heating run. Thermogravimetric (TGA) analyses were performed on TGA-Q50 system from TA instruments at a heating rate of 10 °C.min⁻¹ under nitrogen atmosphere from room temperature to 600°C. Dynamic mechanical analyses (DMA) were performed on RSA 3 (TA instrument). The sample temperature was modulated from -50 °C to 150 °C, depending on the sample at a heating rate of 4° C.min⁻¹. The measurements were performed in a tension mode at a frequency of 1 Hz, an initial static force of 0.1 N and a strain sweep of 0.04%. Photo-crosslinking were performed using a UV lamp HAMAMATSU equipped with a LC8 lamp (full power of 4000 mW.cm⁻¹) and an A9616-03 filter transmitting in the range 280-400 nm, avoiding the heating of the mixture reaction. The samples were placed at a distance of 9 cm of the UV lamp

Synthesis of NH-Und-1,3-diol

Methyl 10-undecenoate (1 equiv.), was mixed with 2-amino-1,3-propanediol (1.3 equiv.). TBD (0.05 equiv.) was added to the reaction mixture and heated up at 80°C under a nitrogen flow to remove methanol formed. The reaction mixture was heated for 3h. Depending on the matrix, the reaction mixture might become solid indicating the end of the reaction. The product is then extracted with CHCl₃ and washed 3 times with water. Brine is added in case of emulsion. The organic phase was dried over Na₂SO₄ and the solvent was removed under vacuum to yield **NH-Und-1,3-diol**. The diol was obtained as white crystals. Yield: 70%. ¹H NMR (400 MHz, CDCl₃) δ (ppm): 6.22 (s, NH), 5.81 (m, 1H), 4.93 (m, 2H), 3.96 (m, 1H), 3.84 (m, 2H), 3.78 (m, 2H), 2.72 (2 OH), 2.24 (t, 2H), 2.04 (q, 2H), 1.65 (m, 2H), 1.38-1.25 (m, 10H). ¹³C NMR (100 MHz, CDCl₃) δ (ppm): 174.39 (CH₂-CO-NH), 139.33 (CH₂=CH-CH₂), 114.31 (CH₂=CH-CH₂), 63.76 (CH-(CH₂-OH)₂), 52.49 (CH-(CH₂-OH)₂), 36.94 (CH₂-CO-NH), 33.92 (CH₂=CH-CH₂), 29.44-29.02 (CH₂), 25.84 (CH₂-CH₂-CO-NH).

Synthesis of NH-Und-6CC

In a 500 mL round bottom flask at 0°C equipped with magnetic stirrer, **NH-Und-1,3-diol** (1.0 equiv.) and ethyl chloroformate (4.0 equiv.) were dissolved in 300 mL THF. To the cold reaction mixture, triethylamine (4.0 equiv.) was added dropwise over 10 min. The reaction was allowed to proceed in ice-cold conditions for about 1 hour and then allowed to proceed at room temperature overnight. The precipitated solids were filtered off and the volatiles were removed to result in crude product, which was subjected to flash column chromatography, using a gradient of DCM (100 %) to DCM (95 %) and methanol (5 %) solvent mixture, followed by the removal of volatiles to result in white solid **NH-Und-6CC** as the functional monomer. Yield: 55%. ¹H NMR (400 MHz, CDCl₃) δ (ppm): 7.63 (s, NH), 5.76 (m, 1H), 4.93 (m, 2H), 4.54-4.40 (m, 5H), 2.22 (t, 2H), 1.99 (q, 2H), 1.60 (t, 2H), 1.25 (m, 10H). ¹³C NMR (100 MHz, CDCl₃) δ (ppm): 174.33 (CH₂-CO-NH), 148.66 (OCOO), 139.19 (CH₂=CH-CH₂), 114.21 (CH₂=CH-CH₂), 71.45 (CH-(CH₂-OCOO)₂), 40.94 (CH-(CH₂-OCOO)₂), 36.14 (CH₂-CO-NH), 33.81 (CH₂=CH-CH₂), 29.34-28.93 (CH₂), 25.57 (CH₂-CH₂-CO-NH).

Polymerization of NH-Und-6CC

All polymerizations were performed under inert atmosphere (nitrogen) using standard Schlenk, vacuum line, and glovebox techniques. In a 15 mL schlenk flask containing a magnetic stirrer, in glove box, **NH-Und-6CC** (50 equiv.), BnOH (2.0 equiv.) and Schreiner TU (1.0 equiv.) were dissolved in dry DCM ([**NH-Und-6CC**] = 2 mol.L⁻¹). To this solution, DBU (1.0 equiv.) was added to initiate polymerization. The reaction mixture was allowed to stir at room temperature. After 5h, the reaction was quenched by the addition of benzoic acid (2 equiv.) and purified by precipitation in cold methanol to yield P(**NH-Und-6CC**). The polymer was obtained as white solid. Yield: 91%. ¹H NMR (400 MHz, CDCl₃) δ (ppm): 5.80 (m, 1H), 4.93 (d, 2H), 4.64-3.93 (m, 5H), 2.24 (m, 2H), 2.01 (q, 2H), 1.62 (m, 2H), 1.28 (m, 10H). SEC (THF, RI): Mn= 5 700 g.mol⁻¹, D=1.07.

Synthesis of SH-Cinnamoyl

In a round-bottom flask equipped with magnetic stirrer, cinnamoyl chloride (20 g, 120 mmol) was dissolved in toluene (300 mL). 2-mercaptoethanol (9.3 g, 120 mmol, 1 equiv.) was added and the mixture was stirred under reflux for 2 hours. The solvent was removed on rotary evaporator yielding to **SH-Cinnamoyl** as viscous liquid without further purification. Yield= 90%. ¹H NMR (CDCl₃, 400 MHz) δ (ppm): 7.73 (d, 1H), 7.53 (m, 2H), 7.38 (m, 3H), 6.47 (d, 1H), 4.33 (t, 2H), 2.82 (q, 2H), 1.55 (t, SH). ¹³C NMR (100 MHz, CDCl₃) δ (ppm): 167.45 (CH-<u>C</u>OO-CH₂), 146.56 (Ph-<u>C</u>H=CH), 133.19-128.31 (Ph), 118.21 (CH=<u>C</u>H-COO), 66.95 (CH-COO-<u>C</u>H₂), 23.74 (CH₂-<u>C</u>H₂-SH)

Grafting of SH-Cinnamoyl on P(NH-Und-6CC)

In a round-bottom flask equipped with magnetic stirrer, P(NH-Und-6CC) (1 equiv.), SH-Cinnamoyl (3 equiv.) and V70 (1 mol.%) were dissolved in DCM (1 mol.L-1). The reaction was allowed to proceed at 40°C during the appropriate period of time. The solvent was removed to result in c_x -P(NH-Und-6CC) which was purified by dialysis in methanol (1L for 1g of polymer, 48h, and methanol was changed after 24h). Grafting of cinnamoyl moieties were calculated by 1H NMR (Figure S14) according to Erreur ! Source du renvoi introuvable..

$$\% Cinnamoyl = \frac{\int H_{11}}{\int H_7 + \int H_{11}}$$

Equation 1: Formula used for the calculation of cinnamoyl percentage grafted after the thiol-ene reaction using ¹H NMR integrations of characteristic protons (H₁₁ and H₇) in CDCl₃

Photo-cross-linking of cx-P(NH-Und-6CC)

The polymer c_x -P(**NH-Und-6CC**) was dissolved in chloroform (CHCl₃) at 1g.mL⁻¹ in a vial. After evaporation of the solvent, the films were exposed to UV light at a distance of 9 cm, wavelength $\lambda = 365$ nm, for 12 h on each side to form a network structure CL- c_x -P(**NH-Und-6CC**) and to ensure uniform curing.



Figure S1. ¹H NMR spectrum of NH-Und-1,3-diol in CDCl₃



Figure S2. ¹³C NMR spectrum of NH-Und-1,3-diol in CDCl₃



Figure S3. ¹H-¹H COSY spectrum of NH-Und-1,3-diol in CDCl₃



Figure S4. ¹H-¹³C HSQC spectrum of NH-Und-1,3-diol in CDCl₃



Figure S5. ¹H NMR spectrum of NH-Und-6CC in CDCl₃



Figure S6. ¹³C NMR spectrum of NH-Und-6CC in CDCl₃



Figure S7. ¹H-¹H COSY spectrum of NH-Und-6CC in CDCl₃



Figure S8. ¹H-¹³C HSQC spectrum of NH-Und-6CC in CDCl₃



Figure S9. ¹H NMR spectrum of P(NH-Und-6CC) in CDCl₃



Figure S10. ¹H NMR spectrum of SH-Cinnamoyl in CDCl₃



Figure S11. ¹³C NMR spectrum of SH-Cinnamoyl in CDCl₃



Figure S12. ¹H-¹H COSY spectrum of SH-Cinnamoyl in CDCl₃



Figure S13. ¹H-¹³C HSQC spectrum of SH-Cinnamoyl in CDCl₃



Figure S14. ¹H NMR spectrum of cinnamoyl-containing polycarbonate in THF. Example with 30 mol.% of cinnamoyl content grafted on P(**NH-Und-6CC**)



Figure S15. SEC traces of cinnamoyl-containing polymers in THF (Polystyrene standards)



Figure S16. Second heating scans of DSC measurement of cinnamoyl-containing polycarbonates



Figure S17. Second heating scans of DSC measurement of cross-linked cinnamoyl-containing polycarbonates



Figure S18: TGA analysis of cross-linked cinnamoyl-containing polycarbonates



Figure S19. Stress–strain curves at room temperature of the polycarbonate networks prepared with the [2+2] photochemical cyclo-addition of pendant cinnamoyl moieties



Figure S20. Thermo-mechanical experiments showing the tensile storage modulus (E') measured using DMA at an oscillation frequency of 1 Hz



Figure S 21: Thermo-mechanical experiments showing Tan Delta measured using DMA at an oscillation frequency of 1 Hz