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1	Thermal aging of acrylic-urethane network: kinetic modeling and end-of-life criteria combined with
2	mechanical properties
3	
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16	
17	Abstract
18	This study addresses the multiscale analysis of acrylic urethane networks (AUN). To establish the

1	kinetic model for predicting AUN oxidation, this study considered the pure thermal oxidation of AUN at
2	160, 180, and 200 °C. Chemical changes were monitored using infrared spectroscopy. These indicated the
3	presence of an imide, presumably generated from the oxidation of CH_2 at the α -position of nitrogen. On the
4	macromolecular and macroscopic scales, oxidation was shown to induce predominant crosslinking, leading
5	to a drop in toughness (i.e., embrittlement). The novel kinetic model of AUN thermal aging was developed
6	from a mechanistic scheme previously established for polyamide 11, by adding some extra paths of
7	thermolytic alkyl radical formation, oxidative N-H bonds decomposition and coupling of aminyl radicals.
8	
9	Keywords: Kinetics, oxidation, macromolecular architecture, acrylic-urethane network
10	
10 11	1. Introduction
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11 12 13 14 15	Acrylic-urethane network (AUN) is a well-known thermoset material used in a wide variety of commercial and technical applications; it is particularly prevalent as a coating for buildings and infrastructure, owing to its excellent flexibility, film-forming properties, and resistance to weathering [1]. The coatings applied to structure surfaces are exposed to several environmental stress factors, such as temperature [2], light [3],

1	Cracking is a major problem for coatings, and it can ultimately determine the lifespan of structures.
2	Cracking phenomena have been widely studied and have attracted attention from coating users in the
3	building and construction communities, because cracking failure can induce a dramatic deterioration of
4	crucial building properties [9,10].
5	For evaluating the resistance to crack propagation, the Essential Work of Fracture (EWF) method [11,12]
6	should be the first choice, which analyzes the fracture properties of notched specimens. As one of the
7	alternative options, it is sometimes evaluated from the area under the stress-strain curve [13] of the classical
8	tensile test, a routine procedure for characterizing mechanical properties in the polymer degradation field
9	[4,14,15]. This loss of ductility is related to various macromolecular characteristics such as internal stress
10	[11], changes in crosslink density [16], fracture energy [17], and changes in sample weight due to the release
11	of small volatile products [18].
12	In real applications of polymer-based materials, oxidation strongly limits the service lifetime, because
13	oxidized materials suffer deteriorations (e.g., embrittlement and yellowing) in their structural properties. In
14	general, oxidation modifies not only the chemical structure but also the macromolecular architecture [19]
15	through chain scission and/or crosslinking [20,21]. Hence, a multi-scale perspective is needed to understand
16	aging behaviors and service lifetimes.
17	AUN oxidation has been studied in earlier papers on photo-aging [22–25]. Oxidation of urethane group

18 was highlighted together with a predominant crosslinking (which increases the glass transition temperature

1	(T_g) and micro-hardness [22]) and a decrease in the free volume and its radius [23]. An elaborated kinetic
2	model for lifetime prediction would represent a valuable tool for AUN users and manufacturers. However,
3	it is difficult to construct a kinetic model for photothermal oxidation from scratch, because photo-oxidation
4	involves complex radical initiation processes determined by the wavelength dependence of the UV-light's
5	effects and the presence of additional physical phenomena such as light attenuation across the material's
6	thickness. In the case of the polypropylene, a kinetic model for photooxidation was developed by
7	completing the model for thermal oxidation [26]. The first aim of this work is an experimental multiscale
8	study of AUN thermal aging since those latter are uncommon in the literature. This study also aims to
9	establish a novel kinetic model for thermal aging (in accordance with Refs. [18] [19][27]) and to investigate
10	the end-of-life criteria related to mechanical resistance.
10 11	the end-of-life criteria related to mechanical resistance.
	the end-of-life criteria related to mechanical resistance. 2. Experimental
11	
11 12	2. Experimental
11 12 13	2. Experimental2.1 Materials
11 12 13 14	 2. Experimental 2.1 Materials The target material was a two-part acrylic-urethane paint (acrylic-polyol and crosslinker) designed for
 11 12 13 14 15 	 2. Experimental 2.1 Materials The target material was a two-part acrylic-urethane paint (acrylic-polyol and crosslinker) designed for commercial use; it consists of a tri-functional network structure cross-linked by hexamethylene

shown in Fig. 1(a), acrylic-polyol is composed of methyl methacrylate (MMA), styrene (St), butyl acrylate
(BuA), and 2-hydroxyethyl methacrylate (HMA). The approximate composition stoichiometry of acrylicpolyol can be inferred from the NMR results, as follows: [MMA]:[St]:[BuA]:[HMA] ~
0.32:0.36:0.16:0.16.

5

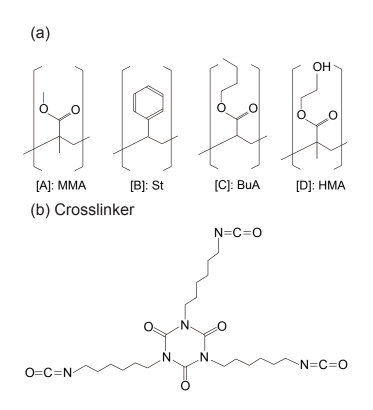


Fig. 1. (a) Structural units of acrylic polyol and (b) chemical structure of crosslinker (HDI trimer).

[(1-column fitting image)].

6

7 Solutions of polyol and crosslinker were mixed in a given polyol/hardener weight ratio of 17/5. The

8 concentrations of OH and NCO in the mixed solution were approximately $[OH]_{mix} \sim 1.30 \, [mol/kg]$ and

1 $[NCO]_{mix} \sim 1.35 \text{ [mol/kg], respectively.}$

2

4 Two-component mixed solutions were cast on a Naflon® sheet (Nichias, Japan) to allow the film sample 5 to be easily peeled off. The samples were hardened under dry conditions (i.e., a desiccator containing silica 6 gel) at 20 °C for a week, after which they were placed in an oven at 105 °C for 2 h. The thicknesses of the 7 film samples were approximately 50–100 μ m (measured by ultrasonic thickness gauge, SDM-3100, 8 SANKO Electronic Laboratory Co. Ltd., Japan). The samples were fully cured, as verified by the total 9 disappearance of the peak at 2275 cm⁻¹ (attributed to the isocyanate group) in the Fourier transform infrared 10 (FTIR) spectra, as described in our previous work [28]. Concerning the so-called "diffusion limited 11 oxidation (DLO)" effect [29,30], Larché, Bussière, and Gardette [22] studied the photo-oxidation of 12 approximately 40-60 μ m acrylic-urethane thin films to investigate the material's photo-oxidation 13 behavior. Accordingly, we assumed that the oxidation behavior of our thin films (50–100 μ m thickness) 14 was not DLO-controlled in this work.

15

16 2.2 Thermal-aging tests

17 Sample films were placed in air-ventilated AP60 ovens (supplied by Systèmes Climatique Service, France)

18 at temperatures of 160–200 $\,^\circ\,$ C.

2 2.3 Characterization 3 2.3.1 Fourier-transform infrared spectroscopy (FTIR) 4 Infrared (IR) spectra were collected in transmission mode using a Frontier spectrophotometer (supplied 5 by Perkin Elmer). The spectra were averaged over 64 scans at a resolution of 4 cm⁻¹ in the range 400-4000 6 cm⁻¹. The concentrations of the oxidation products were calculated using the Lambert–Beer law, expressed 7 as 8 $A = \varepsilon_{\lambda} \times l \times C_{\lambda},$ (1)9

10 where *A* is the absorbance, ε_{λ} [L·mol⁻¹·cm⁻¹] is the molar extinction coefficient, *l* is the sample 11 thickness [cm], and C_{λ} is the concentration of the target chemical species [mol·L⁻¹]. Here, we took the 12 value reported in the literature: $\varepsilon(Imide) = 285 \text{ L} \cdot \text{mol}^{-1} \cdot \text{cm}^{-1}$. [19] 13

14 2.3.2 Classical tensile experiment

15 The mechanical changes induced by the oxidation of the AUN were captured by tensile testing on an

- 16 Instron® 4301 machine. Dumbell samples were obtained by punching the AUN films. The effective zone
- 17 had dimensions of 10 mm (length) and 2 mm (width). The crosshead speed was held constant at 1 mm/min

2	
3	2.3.3 Dynamic Mechanical Analysis (DMA)
4	The dynamic mechanical analysis (DMA) measurements were performed on a Q800 apparatus (TA
5	instruments) driven by a Qseries Explorer in the tensile mode, within a temperature range of 40–140 $^{\circ}$ C
6	using virgin or aged AUN films. The measurement frequency was 1 Hz, and the heating rate was 3 °C/min
7	under a strain of 0.2% with a 0.004 N preload force.
8	
9	2.3.4 Sol-gel analysis
9 10	2.3.4 Sol-gel analysis Swelling experiments were performed using the classical gravimetric method; for this, virgin and aged
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10 11 12	Swelling experiments were performed using the classical gravimetric method; for this, virgin and aged samples were immersed (initial mass m_0) in toluene at room temperature. After reaching the equilibrium swelling states, the samples were removed and lightly wiped with tissue to remove excess solvent from

$$SF = \frac{m_0 - m_{dry}}{m_0}.$$
 (2)

during the test, and a 100 N cell was used.

17 The swelling degree (*Q*) was formulated as described in [23]:

$$Q = \frac{m_s - m_{dry}}{m_{dry}}.$$
(3)

2	
3	Here, Q is the swelling degree in the equilibrium state, and m_s is the weight of the sample after
4	swelling.
5	
6	2.3.5 Pyrolysis gas chromatography-mass spectrometry (GC/MS) analysis
7	A sample of approximately 1 mg was used for the pyrolysis gas chromatography-mass spectrometry
8	(GC/MS) measurements. The sample was placed in a deactivated stainless-steel sample cup and heated in
9	a pyrolyzer at 200 $^\circ$ C for 6 h under a helium atmosphere with a flow rate of 50 mL/min. The resultant
10	volatiles were redirected to a GC/MS system (GCMS-QP2010 SE, Shimadzu, Kyoto, Japan). A proportion
11	of the flow (1 mL/min), reduced by the GC splitter (50:1), was continuously introduced into the MS via a
12	transfer capillary. This capillary was maintained at 280 $^\circ$ C in a GC oven to prevent condensation of less
13	volatile products in the capillary (Ultra ALLOY, 0.25 mm i.d. \times 30 m long, Frontier Lab). For the MS
14	measurements, electro-ionization was performed at 70 eV with an operating mass range of 20–500 m/z and
15	a scan rate of 0.3 s/scan. Observed products were identified using the corresponding mass spectra and the
16	F-search software (Frontier Laboratories Ltd).
17	

1	2.3.6 Differential	Scanning	Calorimetry	(DSC)
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2	Differential scanning calorimetry (DSC) was conducted using a Q10 apparatus (TA Instruments).
3	Approximately 5 mg of each sample was placed in a standard sealed aluminum pan and heated from 0 to
4	200 °C at a heating rate of 10 °C/min. The DSC cell was continuously purged with nitrogen (50 ml/min).
5	The results were analyzed using TA Analysis software.
6	
7	2.3.7 Nuclear Magnetic Resonance (NMR)
8	¹ H-NMR spectra for virgin/aged polyol (linear) components at 200 °C under air and vacuum conditions
9	were collected. The samples dissolved in CDCl3 solution measured at room temperature on a Bruker Avance
10	III HD400 spectrometer (400 MHz) using standard pulse sequences available in the Bruker software.
11	Chemical shifts are expressed relative to SiMe4, which is used as an internal reference.
12	
13	3. Results
14	3.1 Chemical changes
15	Chemical change monitoring should constitute the first step in a mechanistic discussion of thermal aging
16	behavior. In general, the thermal oxidation loop is initiated by thermolysis of the peroxide groups
17	[19,22,27,31]. In addition, homolysis of urethane groups were previously reported under both conditions
18	in heating and UV irradiation [32,33]. We expect the thermal aging at high temperatures (in the range of

1	160-200 °C) can provide insight into both oxidative and non-oxidative degradation chemistry of AUN.
2	Although alkyl radicals (P°) can be formed during thermal oxidation via the hydrogen abstraction of peroxy
3	radicals (POO°), the non-oxidative route of P° formation from thermolytic events of the main functional
4	groups must be investigated in thermal aging cases. Here, we study preliminary thermal aging under
5	vacuum conditions at 200 °C, followed by FTIR (Fig. S1 in the Supplementary Materials). Even in the case
6	of inert thermal aging, thermolytic scission of the N-H bonds was not observed.
7	Next, the time-dependent IR spectra for thermal aging under air are shown in Fig. 2 (a)-(f). The peak
8	absorption correspondences in Fig. 2 were determined from the literature [3,23,34,35] and by comparisons
9	of the FTIR spectra of casted polyol and AUN films (see Fig. S2 in the Supplementary Materials). The
10	bands at 3385 cm ⁻¹ and the shoulder of the C-H region at 2859 cm ⁻¹ are attributed to the N-H stretching
11	vibration in urethane-linkage and C–H bonds at the α -position of an N atom. Absorption in the region
12	1850–1650 cm ⁻¹ region corresponds to C=O vibrations. The vibrational band at 1539 cm ⁻¹ represents the
13	amide II band, which indicates a combination of C–N stretching and N–H in-plane-bending vibrations. First
14	of all, the formation of a left-wing of the C=O band (1800–1750 cm ⁻¹) clearly indicates the progress of
15	oxidation [2,3,19,22,23,36]. This wide band at 1800–1750 cm ⁻¹ is quite complex because of overlapping
16	peaks of oxidized products; however, it may contain a possible indicator of AUN oxidation. In general, it
17	is commonly concluded that the features of urethane oxidation include an N-H group (urethane) loss,
18	consumption of C–H bonds at the α -position of an N atom, development of the band at 1770 cm ⁻¹

1	(attributable for the oxidized product, imide structure), and loss of the amide II band [2,22,23,37]. In
2	addition, a very small peak at 3240 cm ⁻¹ appeared in Fig. 2(a) also corresponds to an N-H bond in the
3	generated imide structure [38]. On the other hand, a formation of the shoulder at 1800–1780 cm ⁻¹ in the
4	severely aged cases possibly implies the oxidation of the acrylic-polyol part (attributed to anhydride and/or
5	lactone) [39,40]. These mechanisms are discussed in detail in the Discussion section (with reference to
6	previous investigations). Furthermore, the decrease of absorbance at 3385 cm ⁻¹ suggests that N-H groups
7	in urethane are destroyed and aminyl radicals are formed [41]. It is well known that the bond dissociation
8	energy (BDE) values for the N-H bond and the C–H bond at the $\alpha\text{-position}$ of the heteroatoms were
9	significantly lower than those for the C-H bond in olefinic materials [31,42]. From observed spectral
10	features in thermal aging, possible structures of alkyl radicals and aminyl radicals can be considered around
11	heteroatoms in this system (i.e., urethane-linkage and isocyanurate ring), as shown in Scheme. 1.

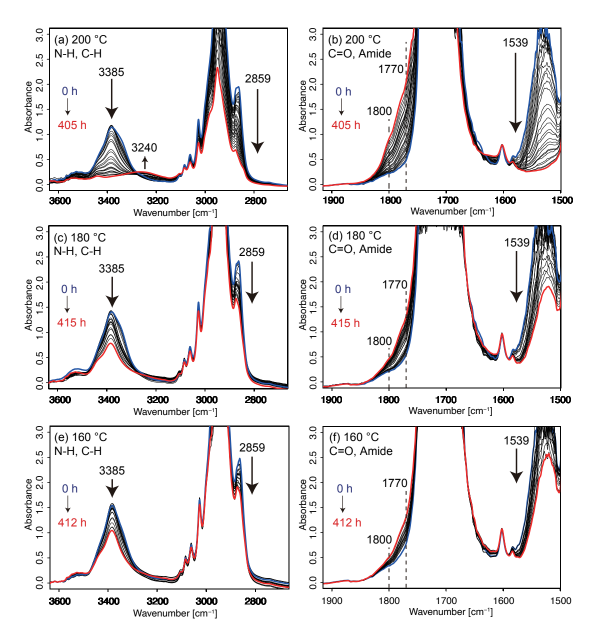


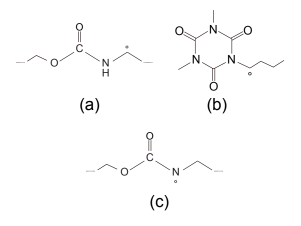
Fig. 2. Time-dependent IR spectra for thermally aging AUN at (a), (b) 200 °C;

(c), (d) 180 °C; and (e), (f) 160 °C under air.

[(Color), (2-column fitting image)].

1

2



Scheme 1. Possible structures of radicals around heteroatoms.

(a), (b): alkyl radicals, (c): aminyl radical

[(1-column fitting image)].

2	To identify volatile compounds, GC/MS measurements were performed. Volatiles were collected from
3	samples heated in the pyrolyzer at 200° C for 6 h under a helium atmosphere. The pyrogram observed
4	for this measurement in the total ion current (TIC) mode is shown in Fig. S3 in the Supplementary
5	Materials. The assigned products listed in Table. 1 indicate the volatilization of the styrenic and MMA
6	units from the acrylic linear component. Note that these fragments were not observed in case of flash
7	pyrolysis at 200° C, adsorbed water and monomer-dissolving solvents could be only detected. In other
8	word, these fragments were not ascribed for unreacted monomers but cleaved product from AUN system.
9	In addition, the BuA unit was not detected in this GC/MS measurement. This may be because the amount
10	of unstable BuA unit was small, or the unit is thermally more stable than the detected the trimer fragments
11	(e.g., St-St-MMA unit, St-MMA-St unit, St-MMA-MMA). Note that no units were detected in the

1	monomeric or dimeric state in the short retention time region on the pyrogram (Fig. S3 in the
2	Supplementary Materials.). It may be that the part of the trimer consisting of MMA and St is thermally
3	unstable and easily ejected, and BuA is possibly not included in the unstable part. Hence, these trimer
4	fragments consisting of MMA and St are potentially responsible for mass loss in the AUN system. This is
5	discussed in the Discussion section (along with the mass-loss behavior).
6	

Table. 1. Main volatiles identified from GC/MS analysis.

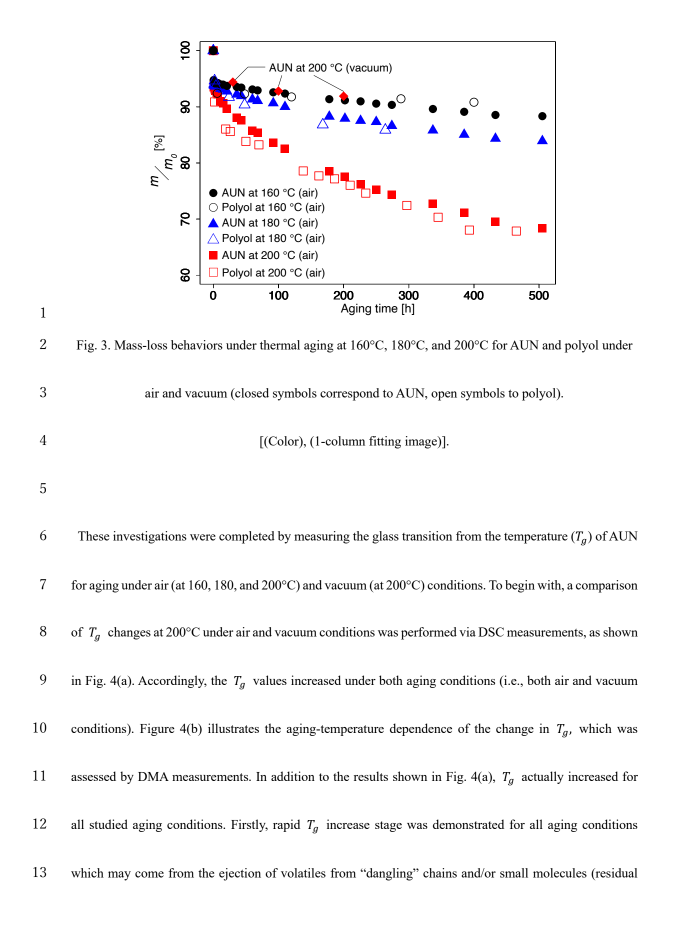
Retention time [min]	Origin	Formula
14.9, 15.1, 16.3	St dimer and MMA	
15.0, 15.2, 15.7	St dimer and MMA	
16.1, 16.2	St and MMA dimer	

8

9 3.2. Macromolecular changes

10 The mass losses obtained for aging under air at several temperatures (160, 180, and 200 °C) are shown in
11 Fig. 3 for the crosslinked AUN and (un-crosslinked) polyol film. Furthermore, we compared AUN

- 1 thermally aged at 200 °C under air and under vacuum.
- 2 The results suggest the following comments:
- 3 Mass-loss curves systematically display a very rapid mass-loss stage, which is typically ascribed to
- 4 absorbed water, and monomer-dissolving solvents [43,44].
- 5 The entire curve exhibits the auto-decelerated shape typical of oxidation processes involving reactive sites
- 6 in the α -position of heteroatoms [2,27].
- 7 For all aging conditions under air (200 °C, 180 °C, and 160 °C), the mass-loss curves are relatively similar
- 8 for linear polyol and AUN. More precisely, the mass loss was slightly higher for the polyol component.
- 9 Expressed otherwise, the reactions responsible for the release of volatiles primarily occur in the acrylic-
- 10 polyol component, which coincides with the detected volatile compounds in the GC/MS measurement.
- 11 The comparison of aging under air and vacuum conditions at 200 °C suggests that thermal degradation
- 12 under vacuum generates only a small number of chain scissions and (later) volatile compounds, as discussed
- 13 below.
- 14
- 15
- 16



1	monomers, absorbed water, and solvents), which coincides with GC/MS and mass-loss results.
2	Subsequently, the T_g values increased regularly which indicates the decrease in molecular chain mobility
3	induced by oxidation [45]. One of the first factors that come to mind is the crosslinking formed by the
4	termination of two alkyl radicals. Another possibility may be the additional hydrogen bonding formation
5	and promoted the steric effect by introducing C=O bonds. In order to argue that the predominant
6	crosslinking in terms of macromolecular architecture, it is not sufficient to only confirm the T_g increase.
7	As a supplemental study, the occurrence of the crosslinking process was tentatively verified by sol-gel
8	analysis of the virgin/aged AUN for aged at 200 °C under air. For the soluble fraction (SF), the unaged
9	AUN had a 7.2 % SF (corresponding to the initial mass-loss as shown in Fig. 3). Then, the SF dropped
10	to almost zero (less than 0.5 %) after even a short time of thermal aging. In addition, a decrease of swelling
11	degrees was demonstrated along with aging process, as shown in Fig. 5. This second result has two possible
12	explanations: (i) the chemical changes occurring in AUN produce a lower affinity with the toluene used as
13	the swelling solvent, or (ii) the crosslink density increases, reducing the swelling degree. Regardless, the
14	most striking result is the absence of SF after thermal aging. If chain scission is favored compared with
15	crosslinking, SF should increase along with the progress of oxidation as in the previous study of AUN
16	oxidation [28]. Therefore, it would be reasonable to consider the "qualitative" predominant crosslinking by
17	combining experimental results (swelling degree, SF , and T_g). In conclusion, the overall experimental
18	results suggest in favor of the ejection of several structural units accompanied by crosslinking. Let us recall

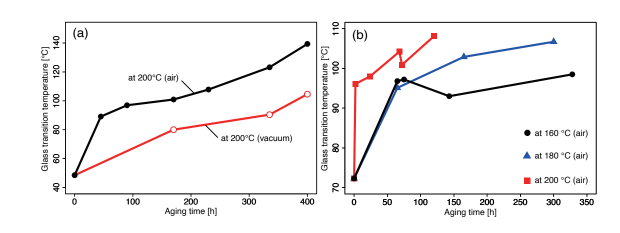
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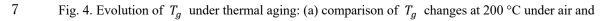
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that in tridmensionnal networks such as AUN system, number of elastically active chains increase by 2 for each crosslinking and decrease by 3 for each chain scission [24]. According to the abovementioned results, it possibly satisfies the 2(crosslinking) - 3(chain scission) > 0, i.e., the concentration in produced crosslinks

4 exceeds 2/3 of chain scission ones.

5



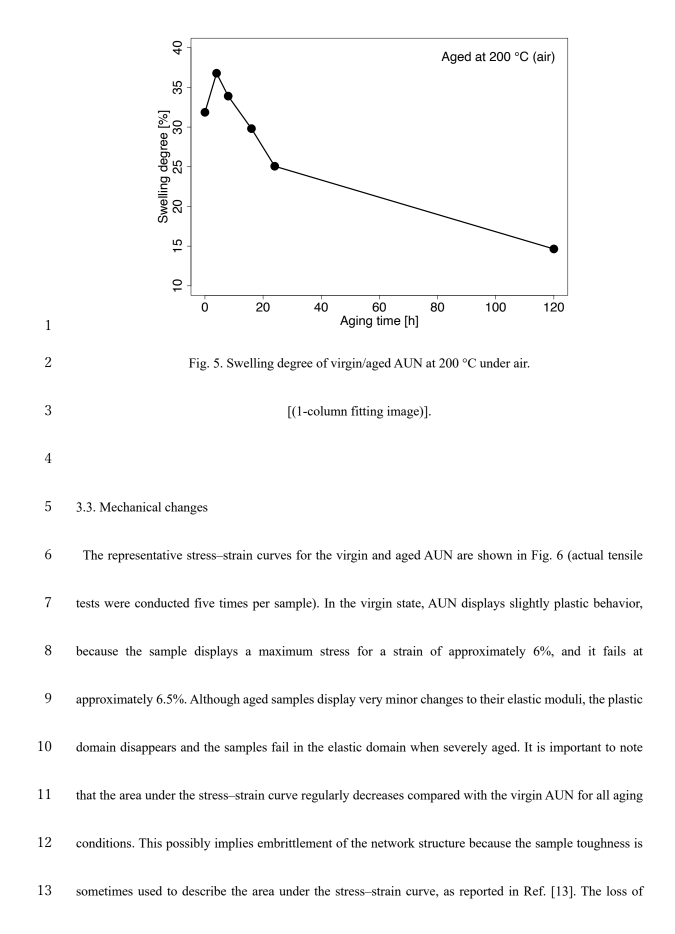


8 vacuum conditions, as assessed by DSC; and (b) temperature dependence of T_g evolution, as assessed by

- 9 DMA.
- 10

6

[(Color), (2-column fitting image)].



toughness is more significant in air aged AUN than aged under vacuum, which possibly originates from the restriction of molecular chain mobility arisen from the oxidation process. It is also important to note that the elastic modulus is not significantly changed for all temperatures under investigation (160, 180, and 200 °C), while it decreases by thermal aging at 200 °C under vacuum. This also relates to the cohesive energy density (CED) associated with the progress of oxidation [46]. These detailed mechanisms will be discussed in the Discussion section.

7

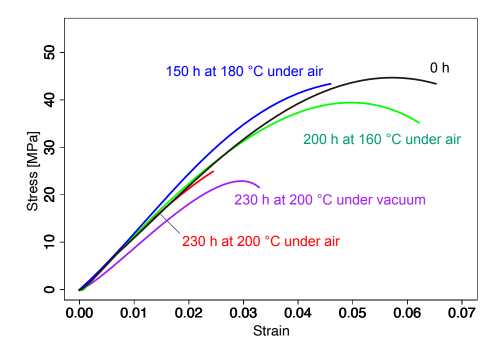


Fig. 6. Representative stress-strain curves for virgin/aged AUN.

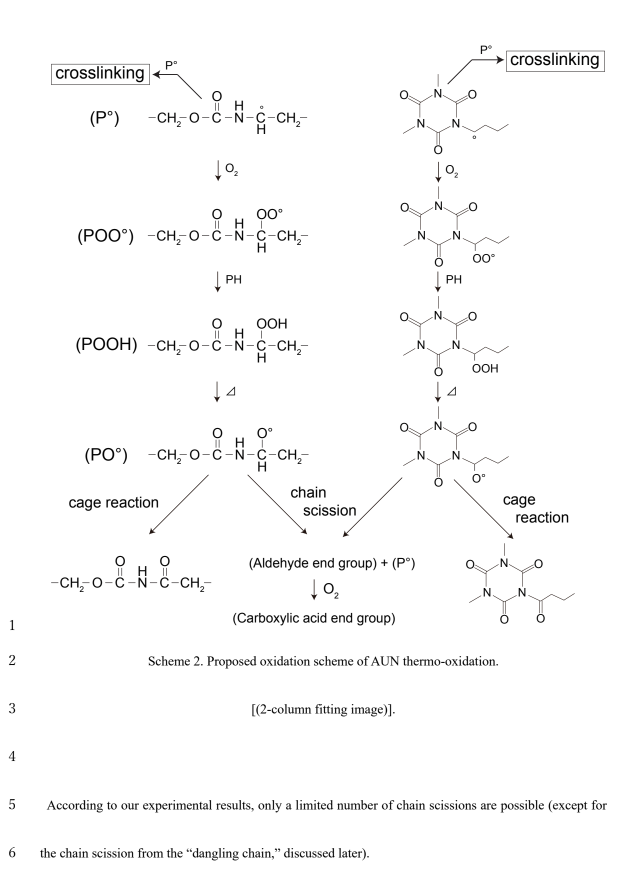
[(Color), (1-column fitting image)].

8

9 **4.** Discussion

1	In this section, l	based on the	experimental	results, v	we provide a	coherent	explanation	of the AUN	thermal
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2	aging mechanism. The main aims of this section are as follows:
3	- To propose a chemical mechanism for the thermal and thermal oxidative aging of AUN.
4	- To discuss the structural properties involved in the mechanical failure of AUN during thermal aging.
5	- To derive a novel kinetic model predicting the appearance of chemical markers relating to chemical aging
6	effect.
7	
8	4.1. Chemical mechanism
9	The main chemical marker associated with thermal oxidative aging is the appearance of carbonyl species
10	(presumably imides), as observed by FTIR spectroscopy (Fig. 2). We stress that this marker is observed
11	only in the presence of oxygen and therefore must correspond to an oxidation product. The most reasonable
12	explanation for this is the oxidation of the two methylene groups in the α -position of heteroatoms in the
13	system (i.e., urethane-linkage and isocyanurate ring). Here, we propose an oxidative degradation path from
14	two reactive sites in the α -position of heteroatoms, as shown in Scheme 2.



7 Expressed otherwise, alkoxy radicals can be converted into "chain" imides without requiring chain

1	scissions, which is consistent with the case of polyamides [19] and epoxies [47]. The chemical structure of
2	"chain" imide in the AUN system corresponds to the N-acylurethane, which has the absorption band in the
3	region of 1780-1765 cm ⁻¹ and at 3240 cm ⁻¹ (broad) [38]. Note that it does not contradict the fact the
4	continuous development of the shoulder at 1770 cm ⁻¹ from the beginning of thermal aging, which could be
5	the important indicator of AUN thermal aging (the absorbance at 1770 cm ⁻¹ is used to estimate the
6	concentration of imide used in the kinetic model). The appearance of a small and broad absorption at 3240
7	cm ⁻¹ is also observed in severely aged AUN at 200 °C. A mechanistic scheme for polyamide 11 is presented
8	in Ref. [19]; there, hydrogen abstraction was reported at the α -position of the amino group in the auto-
9	oxidation process, and this reaction mechanism corresponded well with the photo-oxidation scheme in the
10	acrylic-urethane system, as presented by Larché, Bussière, and Gardette [22]. In this work, the increase in
11	absorbance at 1770 cm ⁻¹ is quite significant, while the development of the peak at 3240 cm ⁻¹ could not be
12	seen as a remarkable peak. This may originate from two reasons: (i) the oxidation of CH2 next to the
13	isocyanurate ring can yield the imide structure without N-H bond (corresponding to the peak at 3240 cm ⁻
14	¹), as shown in Scheme. 2 (on the other hand, the oxidation of CH ₂ next to the urethane group can form the
15	imide structure with N-H bond), (ii) the molar absorptivity of N-H bond in the imide structure is smaller
16	than the N-H bond in pristine urethane (Ref. [48,49]). As another oxidation route in Scheme. 2, carboxylic
17	acid formation via chain scission may be possible, and the bands at 1756 cm ⁻¹ and 1711 cm ⁻¹ in the IR
18	spectra were assigned to isolated carboxylic acid and non-isolated carboxylic acid [19,50]. In addition to

1	these bands, the non-negligible shoulder in our case was also formed in the domain 1800–1780 cm ⁻¹ ,
2	especially in the severely aged AUN sample, as shown in Fig. 2. The development of this domain may
3	imply the accumulation of an oxidation product from acrylic-polyol. Lazzari and Chiantore [39] assumed
4	that the CH_2 in the α -position of the ester group was the reactive site in the oxidation process and could
5	form anhydrides and/or γ -lactones [39,40]. In the IR spectrum, the absorption bands at 1805 cm ⁻¹ and 1780
6	cm ⁻¹ can be attributed to anhydrides and γ -lactones, respectively. However, it is important to note that the
7	BDE for the C–H bond at the α -position of the ester was theoretically estimated to be approximately 380–
8	400–kJ mol ⁻¹ [51,52], significantly larger than the BDE at the α -position of an N atom (approximately 360
9	[kJ·mol ⁻¹]). Therefore, we assumed the radical attack of hydrogen abstraction to primarily occur at the C-
10	H bond in the α -position of an N atom, rather than the CH ₂ in the α -position of the ester group, owing to
11	the difference in BDEs between the two reactive sites. Although the selective attack on the principal
12	oxidation site (i.e., α -position of an N atom) could occur for the pristine AUN, the oxidation of the acrylic-
13	polyol-part could be possibly caused after the depletion of the principal sites. This gradual mechanism may
14	be the reason why oxidation of the polyol-part was suggested in the late stage of degradation. In addition,
15	the decomposition of N-H groups (in urethane) and formation of aminyl radicals were also implied from
16	FTIR spectroscopy (Fig.2). It is also needed to be considered for establishing a kinetic model of AUN
17	thermal aging. According to the previous paper [53], the coupling of those two aminyl radicals can occur;
18	in other words, the following mechanism would be envisaged: $N^{\circ} + N^{\circ} \rightarrow N - N$ (crosslink). According

to the abovementioned discussions, the first attempt at a kinetic model of AUN thermal aging is proposed

2 in Section 4.3.

Next, let us consider the mechanisms of the increase in T_g (Fig. 4) from the theoretical equation governing the T_g change that considers the entropy of network polymers: the DiMarzio theory [65].

$$T_g = \frac{T_{gl}}{1 - K_{DM} nF}.$$
(4)

7 where T_{gl} is the glass transition temperature of the corresponding "virtual" linear polymer, K_{DM} is the 8 DiMarzio's universal constant, n is the crosslink density, F is the flex parameter. The higher is F, the 9 higher is the inertia of rotatable units, and the lower is the chain mobility. Compared with experimental 10 results, the crosslink formation caused by oxidative degradation should contribute to the increase in T_q . It 11 can yield through the coupling of two alkyl radicals, as demonstrated in Scheme. 2. As reported in Ref. 12 [54], such alkyl radicals are created even under an inert atmosphere, possibly explaining why the T_g 13 increased at 200 °C under vacuum conditions. However, the introduction of C=O groups on the molecular 14 chain can also increase T_g through the increase T_{gl} and/or F in Eq. (4). The introduction of C=O groups 15 possibly act as hydrogen bonding sites and enhance steric effects resulting in the increase of cohesive 16 energy and reduction of molecular mobility, which relates with the T_{gl} and F, respectively [45,46]. Fig. 17 4(a) suggests that the increase in T_g under vacuum is somewhat mitigated compared to the thermal aging

1	under air. In addition to oxidative crosslinking, it can also be explained by the elimination of the effect of
2	the introduced C=O groups in vacuum. The initial rapid increase in T_g is also associated with the emission
3	of volatile components generated from the scission of several units (e.g., styrene and MMA) from "dangling"
4	chains of the acrylic linear component, which is consistent with mass loss (Fig. 3) and GC/MS results
5	(Table. 1). Since "dangling" chains can act as similar to "plasticizer" in the system, the loss of "dangling"
6	chains (ejection of several cleaved units from the acrylic linear part) might explain both the increase of T_g
7	and mass loss in aging process. As one of the supporting results of the mechanism, the ¹ H-NMR spectra
8	change for thermally aged polyol (linear prepolymer) was studied (see Fig. S4 in the Supplementary
9	Materials). The results revealed that C=C bonds were not formed in thermally aged uncrosslinked acrylic
10	components at 200 °C under vacuum. If numerous random main chain scissions were present, C=C bonds
11	would be detected. Combining these results with those of the GC/MS analysis, we find support for our
12	claim that the scission and ejection of several structural units are implied by "dangling" chains (i.e., MMA
13	and St units located in chain ends).
14	
15	4.2. End-of-life criteria
16	Here, we discuss the structural properties involved in the mechanical failure of the AUN during thermal
17	aging. From Fig. 6, we can summarize the notable features of these curves as follows:

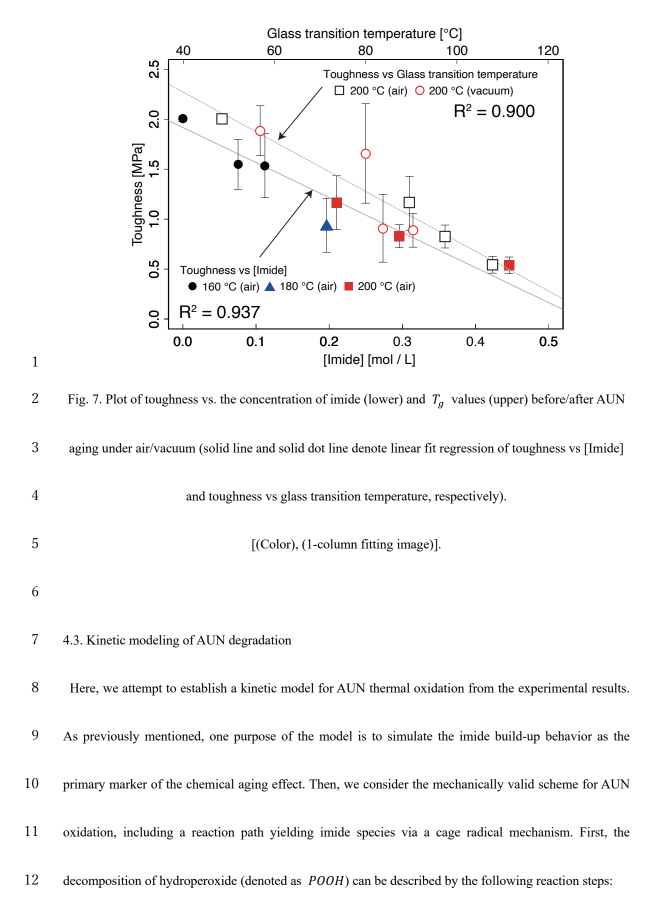
1	- Modulus changes in the elastic region are not significant for all temperatures under investigation (160,
2	180, and 200 °C).
3	- The maximum elongation is decreased in the severely degraded case (i.e., embrittled), and the yield
4	stress is also decreased when the sample is brittle.
5	
6	Note that the tensile test was performed at room temperature, and the analyzed AUN is in the glass state.
7	According to Ref. [55], the elastic modulus of a thermosetting polymer in the glassy state should be
8	proportional to the cohesive energy density (CED). Although the oxidation of CH2 groups at the α -
9	position of an N atom can yield imide structures, we assume the effect on the CED possibly remains limited.
10	As suggested by Van Krevelen (Ref. [46]), the CED can be estimated by the "additive group contribution
11	method":
12	

$$CED = \sqrt{\frac{\sum_{i} E_{coh}^{(i)}}{\sum_{i} V_{m}^{(i)}}}$$
(5)

14 where $E_{coh}^{(i)}$ is the cohesive energy of each group and $V_m^{(i)}$ is the molar volume of each group. The 15 oxidation with the formation of the imide structure can increase both the denominator and the numerator, 16 and their square root contributes to the CED. As a result, the CED of the network structure, which can 17 contribute to the physical properties, may not change significantly by the progress of oxidation. In the

1	present system, the T_g increased, and the modulus did not change significantly along with the thermal
2	oxidation, which the decrease in molecular chain mobility can justify due to crosslinking and/or hydrogen
3	bonding formations. These results are not surprising case. In fact, it has been reported that in the case of an
4	oxidized epoxy system, the change in CED was limited, and the rubbery modulus changed only slightly;
5	however, the T_g increased significantly (Ref. [56]). On the other hand, the decrease of the elastic modulus
6	in thermally aged samples under the inert condition could be explained that the contribution of chain-
7	scission was only occurred without any oxidation. Concerning the second comment on the tensile
8	characteristics, it suggests the significance of toughness as an indicator of the risk of mechanical failure,
9	because toughness loss can be considered as an "embrittlement level." As stated in the Introduction section,
10	toughness could be sometimes assessed from the area beneath the stress-strain curve, as described in [13].
11	In our study, the calculated toughness was markedly decreased by thermal aging for all temperature grades
12	(see Fig. S5 in the Supplementary Materials). Here, we assessed the potential correlation between the
13	toughness loss and the progress of thermal aging. Figure 7 plots toughness with respect to the concentration
14	of the main chemical marker (i.e., imide) and the T_g values. These results present a strong linear
15	correlation between the loss of toughness, accumulation of oxidized species, and T_g increase. As stated in
16	Section 3.2, the increase in T_g should be related to the extent of crosslinking. As shown in Fig. 4(a), the
17	AUN underwent significant crosslinking regardless of whether it was aged under air or vacuum conditions.
18	However, the crosslinking efficiency was much higher when aged under air. Hence, the additional

1	likelihood of chemical and/or physical crosslinking (corresponding to the covalent bonds and inter-chain
2	hydrogen-bonding, respectively) may have arisen from oxidation. In addition, the toughness loss is
3	proportional to both the accumulation of imide and T_g values, as shown in Fig. 7. As a side note, the
4	influence on T_g and toughness from the oxidation of acrylic-part cannot be ruled out, as implied in FTIR
5	spectra (Fig. 2). Suppose we want to decouple the contribution of the urethane-part oxidation from the
6	linear-part oxidation. In that case, comparing oxidized urethane-networks crosslinked with several different
7	polyols (i.e., linear part) will be needed. It should be worth for further study to discuss the impact of liner-
8	part oxidation in a future publication. By the way, it should be noted that the imide accumulation, which
9	indicates the progress of chemical aging, is correlated with the amount of chemical and/or physical
10	crosslinking. Therefore, we can make rough predictions of service lifetime under arbitrary temperatures
11	using a kinetic model of AUN thermal degradation. In the next section, we establish a novel kinetic model
12	for thermal aging.



$$POOH \rightarrow PO^{\circ} + HO^{\circ}$$
,

$$(P0^{\circ} + H0^{\circ})_{cage} \rightarrow \text{imide} + H_20, \qquad (\gamma_1)$$

$$PO^{\circ} + PH \rightarrow \text{end chain imide} + P^{\circ} + \text{chain scission},$$
 (γ_2)

$$PO^{\circ} + PH \rightarrow \text{Hydroxyurethane} (-O - CO - NH - CHOH - CH_2 -) + P^{\circ} (\gamma_3)$$

+ chain scission,

$$H0^{\circ} + PH \rightarrow P^{\circ} + H_20, \qquad (1 - \gamma_1)$$

3 where γ₁, γ₂, and γ₃ are the yields of the corresponding reaction paths; they satisfy γ₁ + γ₂ + γ₃ = 1.
4 Hydroxyurethane should destabilize and be converted into carboxylic acid via chain scission [27]; this is
5 expressed as
6 Hydroxyurethane → Carboxylic acid + chain scission.
7
8 Next, the creation of alkyl radicals and subsequent crosslinking was experimentally indicated, even during
9 thermal aging under vacuum conditions. Hence, an extra path for thermolytic alkyl radical formation must
10 be considered:

1 2 Here, *PH* denotes the polymer substrate (here, C–H bonds at the α -position of an N atom). 3 4 According to Fig. 2, the loss of N-H group was observed for all aging conditions under air (200 °C, 180 °C, 5 and 160 °C). Thus, we introduced the probable paths of oxidative decomposition of N-H bonds [57] as 6 follows: 7 $NH + POO^{\circ} \rightarrow N^{\circ} + POOH$, 8 9 Here, NH and N° denotes the N-H bonds in the urethane-linkage and the aminyl radical (as shown in 10 Scheme. 1(c)), respectively. According to the previous work [53], the bimolecular association of two aminyl 11 radicals can occur and yield a crosslink as follows: 12

 $PH \rightarrow P^{\circ}$.

 $N^{\circ} + N^{\circ} \rightarrow N - N$

13

14 In case of aminyl radicals, hydrogen abstraction and β -scission are rarely occurred as reported by Ogawa

15 and Nomura [58]. The nitroxide radicals (*NO*°) could be formed through the propagation of aminyl radicals,

1likewise stabilizers of the amine family. Although NO° can act as stabilizers by scavenging alkyl radicals,2the stabilizing mechanism is assumed to be negligible because the efficiency of the stabilization effect is3quite limited at high temperature (typically $T \ge 150 \ ^{\circ}C$) [57].4Lastly, the formation of a peroxide-type product (*POOP*) can be assumed in termination processes (i.e.,5coupling of $P^{\circ} + POO^{\circ}$ and $POO^{\circ} + POO^{\circ}$). It has been previously suggested that the decomposition of6*POOP* generates imides and carboxylic acids in equal proportions [19,27]. By summarizing the discussions7so far, the overall mechanism of AUN oxidation is proposed as follows:

$$PH \rightarrow P^{\circ},$$
 (k_t)

$$POOH \rightarrow (\gamma_1 + \gamma_2)$$
 Imide $+ 2(\gamma_2 + \gamma_3)P^\circ + \gamma_3$ Hydroxyurethane $+ \gamma_2 S$, (k_1)

$$P^{\circ} + O_2 \to POO^{\circ}, \tag{k_2}$$

$$POO^{\circ} + PH \rightarrow POOH + P^{\circ},$$
 (k_3)

$$NH + POO^{\circ} \rightarrow N^{\circ} + POOH,$$
 (k_{NH})

$$P^{\circ} + P^{\circ} \to X, \tag{k_4}$$

$$N^{\circ} + N^{\circ} \to X, \tag{k_N}$$

$$P^{\circ} + POO^{\circ} \to POOP, \qquad (k_5)$$

$$POO^{\circ} + POO^{\circ} \to POOP + O_2, \tag{k_6}$$

$$POOP \rightarrow Imide + Hydroxyurethane,$$
 $(k_{1d} \sim k_1)$

Hydroxyurethane \rightarrow Carboxylic acid + *S*, (k_d)

1

2 where S and X denote chain scissions and crosslinking, respectively. k_i is the rate constant for each 3 reaction considered.

4 The formulas of corresponding differential equation system are presented in the Supplementary Materials. 5 We here describe the method for determining the rate constants (k_i) of each reaction process used in the 6 kinetic model on AUN oxidation. It should be noted that the parameter configuration is basically referring 7 to the previously developed kinetic models of polyamide 11 [27] and epoxy/amine [14] systems with similar 8 chemical structures (in terms of the common features of oxidation mechanisms). Firstly, the rate constants 9 concerning propagation steps (k_2, k_3) are fixed from a priori based on literature [14,27,59], since the radical 10 propagation process is known to exhibit common characteristics regardless of the type of polymer (see, e.g., Refs. [18,26,27,31,60]). k_2 is the rate constant of oxygen added to P° ; it is assumed to be significantly 11 rapid in comparison with k_1 and k_3 . Here, k_2 is taken as 10⁶ [L mol⁻¹ s⁻¹] without temperature 12 13 dependence (i.e., $E_2 = 0$), which has been validated in polyamide and epoxy/amine systems [14,27]. In 14 addition, k_3 corresponds to the radical propagation process of hydrogen abstraction by POO° radicals. 15 Conventionally, k_3 values, including its temperature dependence, are determined by Korcek's law [59] 16 according to the dissociation energy values for the C-H bond (BDE) of the considered oxidative site. Hence, 17 we assume that the selective attack of hydrogen abstraction occurred primarily at the C-H bond in the a-

position of an N atom, because the BDE here should be considerably smaller (360 kJ·mol⁻¹) than the olefinic

2 BDE (390–395 kJ·mol⁻¹) [31], expressed as

3

$$k_3 = \exp\left(19.517 - \frac{55000}{RT}\right),\tag{6}$$

4

5 where R is the gas constant (kJ mol⁻¹ K⁻¹) and T is the absolute temperature [K]. Secondly, we assume 6 the radical termination steps $(k_4, k_5, k_6, and k_N)$ are basically performed quickly due to the samples aged 7 at high-temperature environments; hence, they could be assumed to be virtually temperature independent (i.e., $E_4 = E_5 = E_N = 0$). In general, the termination steps should depend on the molecular mobility, and 8 9 previous studies [31] have shown that the POO° + POO° termination has a minimal temperature dependence ($E_6 = 0.10 \, [\text{kJ} \cdot \text{mol}^{-1}]$) because it is expected to have smaller molecular mobility than the other 10 11 termination steps (steric effect). Lastly, the initiation rate constant (k_1) corresponds to the event of POOH decomposition. Since the stability of POOH depends on the chemical nature of target systems, k_1 is 12 13 treated as a fitting parameter in this study. In addition, the rate constant for POOP unimolecular 14 decomposition should be slightly smaller than (but very close to) the value for POOH decomposition 15 found in Sagar's work [61]. Thus, we provisionally assume that $k_{1d} \sim k_1$. The other rate constants (k_t , k_1 , 16 k_{NH} , k_4 , k_N , k_5 , k_6 and k_d) vary from system to system according to several factors (e.g., chemical nature 17 and molecular mobility); these parameters are needed to be obtained by a trial-and-error process. Starting

1 from a set of parameters from existing models [14,27] that have been used successfully, we first determined 2 the main rate constants $(k_1, k_4, k_5, and k_6)$ in the auto-oxidation loop to fit our data (the change in 3 concentration of imide, PH, and NH) and not violate the physically realistic hierarchy ($k_4 > k_5 > k_6 >$ 4 $k_2 \gg k_3$). After these parameters (for the radical initiation, propagation, and termination steps) have been 5 completely fixed, the other minor parameters $(k_t, k_d, k_N, and k_{NH})$ were determined by setting their initial 6 values to 0 and adjusting them manually little by little. In this study, the change in oxygen concentration 7 and the effects of the oxygen profile are neglected because the thin film was subjected to thermal aging 8 (i.e., non-DLO conditions). 9 The set of differential equations were numerically solved using the MatLab® ode23s solver. The 10 following initial conditions were used as input data: $[PH]_0 = 1.0 \text{ [mol } L^{-1}], \ [NH]_0 = 0.5 \text{ [mol } L^{-1}], \ [POOH]_0 = 0.01 \text{ [mol } L^{-1}], \ [O_2] = 4 \times 10^{-4} \text{ [mol } L^{-1}$ 11 $[P^{\circ}]_0 = [POO^{\circ}]_0 = [POOP]_0 = [Imide]_0 = [Carboxylic acid]_0 = [Hydroxyurethane]_0 = X_0 = X_0$ 12 ¹], 13 $S_0 = 0.$ 14 15 The initial concentration of the polymer substrate $[PH]_0$ corresponding to the C-H bond concentration

17 $[NH]_0$ corresponding to the initial concentration of N-H bond in this system can be also estimated in the

in the α -position of an N atom can be calculated from the sample composition, as shown in Section 2.1.

16

18 same manner as $[PH]_0$. Time-dependent [PH] and [NH] (test data) are calculated from the absorbances

1 of corresponding chemical species in FTIR spectra as the relative decrease from the initial concentration of

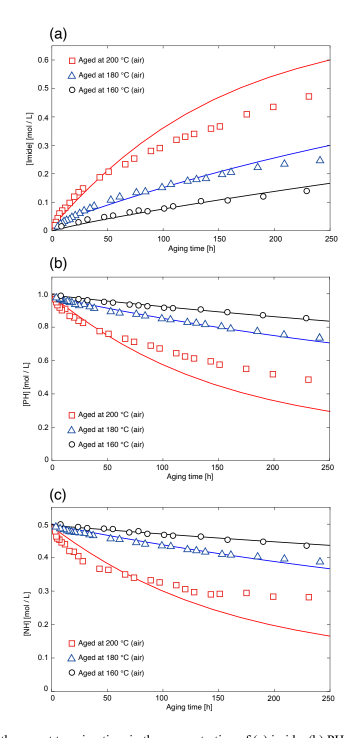
2 $[PH]_0$ and $[NH]_0$.

3

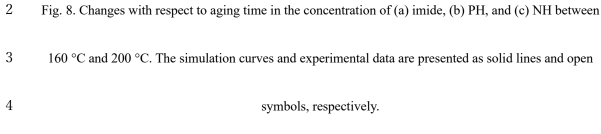
$$[X] = \frac{\operatorname{Abs}(t)}{\operatorname{Abs}(t=0)} [X]_0 \qquad (X = NH \text{ or } PH)$$
(7)

4

5 where Abs(t) is the time-dependent specific absorption in FTIR spectra. To estimate the relative changes 6 in [PH] and [NH], we used the shoulder of the C-H region at 2859 cm⁻¹ and the bands at 3385 cm⁻¹, 7 which are attributed to the C-H bonds at the α-position of the urethane groups and the N-H stretching 8 vibration (in urethane). In addition, $[O_2] = 4 \times 10^{-4}$ suggests a coefficient of O₂ solubility of ~ 10⁻⁸ [mol 9 L⁻¹ Pa⁻¹], by considering Henry's law under the oxygen partial pressure (0.21 bar; ambient air). The order 10 of magnitude of O₂ solubility does not contradict the O₂ solubility of polyurethane rubber reported in Ref. 11 [62]. 12 The output data described the concentration of the following series of reactive species: 13 14 γ_2 and γ_3) are determined by considering compatibility with the experimental results, as shown in Fig. 8. 15 The concentration of imide was estimated using Lambert-Beer's law [Eq. (1)]. The obtained set of rate 16 constants are presented in Table. 2, and γ_1 , γ_2 , and γ_3 are listed in Table 3. 17







[(Color), (1-column fitting image)].

Table 2. Rate constants and activation energies (E_a) used for the simulation and determined by the

2	numerical reverse method at 160 °C.										
	Rate constants	k _t	k_1	<i>k</i> ₂	k ₃	k_{NH}	k_4	k_N	k_5	k_6	k _d
	[mol L ⁻¹ s ⁻¹]	10-10	3 × 10 ⁻⁵	106	69.7	28	9.6 × 10 ¹⁰	10 ¹⁰	5.85×10^{9}	1.6 × 10 ⁹	10-7
	E_a [kJ mol ⁻¹]	155	195	0	55	34.2	0	0	0	3.8	0
3											
4	Table 3	. Yields	of the initi	ation pro	ocess, c	determi	ned by the nu	merical	reverse metho	od.	
			_	γ_1		γ_2	γ_3				
			_	0.78		0.15	0.07				
5			_								
6	The results sugge	st the fo	llowing co	mments	:						
7	- At low conve	ersion (i	.e., [PH] >	0.7 [mo	ol L ⁻¹],	[imide]	< 2.0 [mol I	[⁻¹]), the	validity of th	e simulation	
8	with imide by	uild-up,	PH and NF	I deplet	ion is c	onfirme	ed.				
9	- The simulation	on betw	een 180 °C	and 20	00 °C o	overestii	nated oxidati	ion prog	ress at high co	onversion. It	
10	possibly caus	sed by t	he sensitiv	ity loss	of exp	eriment	al data respo	nsible f	or volatilized	products (as	
11	shown in Fig	. 3) and	/or compen	sated by	y the ox	xidation	of CH ₂ in th	e α-posi	tion of the est	er (so-called	
12	"co-oxidation	n" migh	t coexist) [6	50].							
13	- The simulation	on result	s are sensit	ive to th	$k_1 $ v	values f	or the initiation	on step,	which reflect t	he reactivity	

1	of the AUN system (i.e.	, stability of <i>POOH</i>). We ad	dditionally conducted the sensit	ivity analysis of the
2	simulation results to the	e variation in k_1 values (se	e Fig. S6 in the Supplementary	Materials).
3				
4	Table. 4 compares the	obtained initiation rate cor	stant k_1 , initial <i>POOH</i> conce	entration [POOH] ₀ ,
5	and carbonyl species yi	eld obtained for each POO	<i>H</i> decomposition event for pol	yamide 11 [27] and
6	epoxies [14]. The fitted	k_1 value was lower than t	he other cases, which may refle	ect the difference in
7	<i>POOH</i> stability. The as	ssumed [POOH] ₀ appears	acceptable because it is compa	arable to previously
8	successful models. In th	nis case, the imide yield for	each POOH decomposition e	event was 0.78; that
9	is, 78% of POOH yield	ded imide. This was compar	rable with the case of polyamid	e 11 and lower than
10	that of epoxies.			
11				
12	Table 4. Comparison of	k_1 at 160 °C, [<i>POOH</i>] ₀ , a	nd the carbonyl species yields	for each POOH
13	decomposition event (deno	ted as γ) against the values	obtained in previous works. The second	he k_1 values were
14	extrapolated by the c	corresponding activation end	ergy for cases of polyamide 11	and epoxies.
	Sample	k_1 at 160 °C (s ⁻¹)	$[POOH]_0$ (L mol ⁻¹)	γ
	AUN (present work)	3.00×10^{-5}	1.0×10^{-2}	0.78
	Polyamide 11 [27]	3.76×10^{-3}	3.0×10^{-2}	0.77

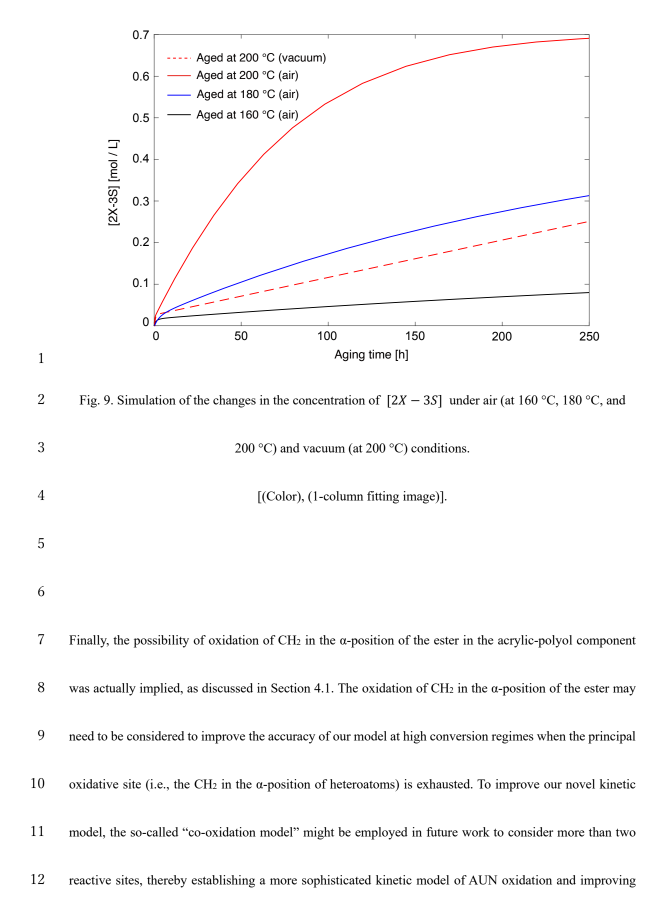
 1.0×10^{-3}

1.00

 1.721×10^{-3}

Epoxies [14]

2	In addition, a simulation of the macromolecular architecture changes is discussed. The experimental
3	characterization of the aged samples implies the predominance of chemical and/or physical crosslinking
4	except for 160 °C, as discussed in Section 3.2. From the experimental results in this study, the number of
5	chain scission is possibly limited, and the thermal oxidation scheme (Scheme. 2) suggests the crosslink
6	formation by coupling alkyl radicals (P°). It is reasonable that the concentration in elastically active chains
7	could be increased consistently under the increase in T_g for thermal aging. According to Refs. [15,63], the
8	time-derivative of the concentration in elastically active chains corresponds to $2X - 3S$. The simulations
9	in this work indicate that $2X - 3S$ remained positive for all aging conditions studied, even under vacuum
10	conditions. In the case of 200 °C aging, $2X - 3S$ was larger under air than under vacuum conditions, as
11	shown in Fig. 9; this is consistent with the evolution of T_g [Fig. 4 (a)]. It was found that $2X > 3S$ for all
12	aging conditions provided $\gamma_2 < 0.27$. Here, the kinetic model was constructed under the (simplified)
13	condition of $2X - 3S > 0$, which is required by the "qualitative" extensive crosslinking inferred from the
14	experimental results. It is required to carry out the continuous and intermittent stress-relaxation test to
15	quantify the rate of crosslinking/scission accurately, submitted by Nichols and Gerlock [64]. Further
16	upgrading the model using this method will be a worth for future works.



1	predictions of crosslinking and chain scission concentration [60]. In addition, extensive crosslinking was
2	reported in the case of AUN "photo-aging" in our previous work [23]. The macromolecular change behavior
3	of AUN exhibits common features between thermal and photo-aging cases. Hence, the possibility of
4	extrapolating the established model for photo-oxidation phenomena is worth further research.
5	
6	5. Conclusion
7	In this paper, the thermal aging of AUN was discussed by comprehensive characterization (FTIR,
8	classical tensile experiment, DMA, sol-gel analysis, DSC, and GC/MS analysis). First, the analysis of
9	chemical modifications arising from the aging process suggested the oxidization of CH_2 in the α -position
10	of an N atom and the volatilization release of MMA and styrenic units from acrylic-polyol, which is
11	responsible for mass loss from the AUN system. Based on the chemical scale discussions, a possible thermal
12	aging scheme including a reaction path yielding "chain" imide via a cage radical mechanism was proposed,
13	in accordance with previous research (i.e., thermo-oxidation of polyamides and AUN photo-oxidation).
14	Another oxidation route corresponds to the conventional closed-loop scheme, which generates carboxylic
15	acids via chain scission.
16	Next, macromolecular changes under thermal aging were studied, primarily from the evolution of the T_g
17	values. In this work, the T_g values were gradually increased for all aging conditions (even at 200 °C

18 under vacuum), which may indicate the predominance of chemical and/or physical crosslinking

1	(corresponding to the covalent bonds and inter-chain hydrogen-bonding, respectively), as confirmed by
2	the absence of a soluble fraction and by the sol-gel analysis. A loss of toughness was also observed
3	alongside the increase in T_g . Toughness can represent an important indicator of the risk of material
4	failure (cracking), because toughness loss might be considered as an "embrittlement level." The
5	toughness loss exhibited a strong linear correlation with the accumulation of the main oxidized product
6	(i.e., imide) and the increase in T_g linked to aging-induced crosslinking. Using these correlations, we
7	might make rough predictions of service lifetime using the kinetic model of the AUN thermal
8	degradation, which can successfully reproduce the build-up of imide.
9	Finally, a first attempt at such a kinetic model of AUN thermal aging was proposed, based on
10	mechanistic discussions derived from the analysis of chemical-scale discussions in the aging process.
11	This was based on a model previously established for polyamide 11, with extra paths of thermolytic alkyl
12	radical formation, oxidative N-H bonds decomposition and coupling of aminyl radicals. The validity of
13	the simulation with imide build-up, PH and NH depletion was confirmed at low conversion. In addition,
14	the predominance of crosslinking during thermal aging could also be reproduced. Our proposing kinetic
15	model is not currently perfect, however, it is possibly acceptable for "mild" thermal aging (at low
16	conversion). As one option for improving our kinetic model, the oxidation of acrylic-polyol may need to
17	be considered. For the unaged system, the probability of oxidation at α -position of an ester group in
18	acrylic-polyol can be estimated to be less than 0.1% at most based on the difference in BDEs between the

1	C–H bond α -position of an N atom and the α -position of an ester group. If the principal oxidation site
2	(PH) is much exhausted, we should envisage the additional oxidation at α -position of an ester group in
3	acrylic-polyol to obtain better prediction even at high conversion. Therefore, proposing a "full" kinetic
4	model of AUN, which takes into account the specificities of linear polymer, would be worthwhile future
5	research, and we believe that a kinetic model for pure thermal aging would help establish a kinetic model
6	of AUN photothermal aging.
7	
8	Acknowledgments
0	Acknowledgments
9	This work was supported by a Grant-in-Aid for JSPS (Japan Society for the Promotion of Science)
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12	Funding sources
13	This work was supported by Grant-in-Aid for JSPS (Japan Society for the Promotion of Science)
14	Fellows (Number 19J20126).
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9			
10	Figu	re Captions	
11	Fig. 1. (a) Structural units of acrylic polyol and (b) chemical structure of crosslinker (HDI trimer).		
12	Fig. 2. Time-dependent IR spectra for thermally aging AUN at (a), (b) 200 °C; (c), (d) 180 °C; and (e), (f)		
13	160 °C under air.		
14	Fig. 3. Mass-loss behaviors under thermal aging at 160 °C, 180 °C, and 200 °C for AUN and polyol unde		
15	air and vacuum (closed symbols correspond to AUN, open symbols to polyol).		
16	Fig. 4. Evolution of T_g under thermal aging: (a) comparison of T_g changes at 200 °C under air and		
17	vacuu	vacuum conditions, as assessed by DSC; and (b) temperature dependence of T_g evolution, as assessed by	
18	DMA.		

- 1 Fig. 5. Swelling degree of virgin/aged AUN at 200 °C under air.
- 2 Fig. 6. Representative stress-strain curves for virgin/aged AUN.
- 3 Fig. 7. Plot of toughness vs. the concentration of imide (lower) and T_g values (upper) before/after AUN
- 4 aging under air/vacuum (solid line and solid dot line denote linear fit regression of toughness vs [Imide]
- 5 and toughness vs glass transition temperature, respectively).
- 6 Fig. 8. Changes with respect to aging time in the concentration of (a) imide, (b) PH, and (c) NH between
- 7 180 °C and 200 °C. The simulation curves and experimental data are presented as solid lines and open
- 8 symbols, respectively.
- 9 Fig. 9. Simulation of the changes in the concentration of [2X 3S] under air (at 160 °C, 180 °C, and
- 10 200 °C) and vacuum (at 200 °C) conditions.