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## 1 THERMAL OXIDATION OF DIALLYLBISPHENOL A - BISMALEIMIDE NETWORKS. COMONOMER EFFECT

# 2 AND KINETIC MODELLING

- 3
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# 11 ABSTRACT

12 Pure BMI resin based on 4,4'-bismaleimidodiphenylmethane (BMI) and its copolymers with various quantities of 2,2'-diallylbisphenol A (BMI-DABPA 2-1, BMI-DABPA 1-1, BMI-DABPA 1-2) were thermally 13 14 aged. Ageing at 160°C was monitored by FTIR and showed the greatest reactivity of diallyl groups which 15 accelerate the degradation of BMI-DABPA copolymers. It was confirmed by a gravimetric study of 16 ageing under nitrogen or oxygen at 350 and 375°C. A co-oxidation kinetic model was proposed for 17 describing the oxidation of BMI and BMI-DABPA systems. The effect of the main kinetic parameters was 18 discussed from a parametric study. The main rate constants were estimated from the fitting of 19 experimental curves and comparable gravimetric data from literature.

20

## 21 KEYWORDS

22 Bismaleimide resins, 2,2'-Diallylbisphenol A, thermal oxidation, mass loss, kinetic modelling

23

## 24 1. INTRODUCTION

BMI resins were first obtained from homopolymerization of bismaleimide monomers, the most common one being the 4,4'-bismaleimidodiphenylmethane (Figure 1a). Polymerization can be achieved even in absence of peroxide or imidazole catalysts  $[^{1},^{2},^{3}]$  and gives networks with high glass transition (300°C or more)  $[3,^{4}]$  and interesting mechanical properties (specific strength at break) but displaying a poor toughness (below 1 MPa.m<sup>1/2</sup>)  $[4,^{5}]$ . Many strategies were implemented for improving this latter such as (i) increasing the distance between crosslink nodes or (ii) adding comonomers such as
for example 2,2'-diallylbisphenol A (DABPA – Figure 1b). This is the case of existing commercial systems
such Matrimid 5292 [<sup>6</sup>]. For example, a Allyl + BMI « copolymer » [<sup>7</sup>] displays stress at break and impact
resistance are respectively around 145 MPa and 12 kJ.m<sup>-2</sup>, which is much higher than data given for the

34 « homopolymer » BMI (40-80 MPa for stress at break [4] and impact strength around 8 kJ.m<sup>-2</sup> [<sup>8</sup>]).

Such materials are usually employed in demanding applications where thermal stability is crucial. Previous works address the case of various homopolymers where the non-reactive group between maleimide groups was changed [<sup>9</sup>,<sup>10</sup>,<sup>11</sup>]. However, there is, to the best of our knowledge, very few studies [<sup>12</sup>,<sup>13</sup>] dealing with the effect of comonomers and in particular the effect of the BMI/diallyl stoichiometric ratio on thermal stability.

Degradation of pure BMI was mostly investigated in non-isothermal conditions by means of classical
 TGA tests [9,10,11]. However, isothermal ageing studies exist but are scarce [9,<sup>14</sup>,<sup>15</sup>,<sup>16</sup>], which limits the
 understanding of the degradation mechanism.

43 Ageing experiments must in general be used for predicting lifetime in service conditions. For that purpose, it is well established that a kinetic model must be used. Kinetic parameters must correspond 44 45 to elementary processes involved in degradation, and obey Arrhenius law. The simulation of 46 experimental results allows an estimation of the values of the parameters used in kinetic models in 47 accelerated conditions. The extrapolation of those values at service temperatures allows to predict the 48 degradation rate at those conditions. A « proof of concept » of the predictive capability of kinetic model was published by Colin et al [<sup>17</sup>] who successfully simulated the mass loss kinetics of a BMI based 49 50 material. However, the proposed kinetic model corresponds to the simple case where there is only one 51 sort of reactive unit in the material, which is clearly not the case in the case of BMI-DABPA networks.

52

53 In this paper, we will hence synthesize and degrade samples with various alkyl/maleimide 54 stoichiometry's in order to:

- investigate the effect of comonomers on stability for isothermal ageing under nitrogen, or underoxygen by performing isothermal tests,

57 - characterize the chemical trackers of oxidation in link with the BMI-DABPA formulation,

- discuss on the effect of DABPA and propose a first kinetic model of degradation for pure BMI and BMI DABPA copolymers. We will focus on its capability to simulate mass loss since this latter is linked to
 shrinkage and latter cracking [<sup>18</sup>,<sup>19</sup>].

# 62 2. EXPERIMENTAL

63

## 64 2.1. Materials

65

66 Materials were synthesized using two comonomers given in Figure 1: 4,4'-67 Bismaleimidodiphenylmethane (CAS: 13676-54-5 - ref 227463-100G supplied by Sigma Aldrich) and 68 2,2'-Diallylbisphenol A (CAS: 1745-89-7 - ref 413526-100ML supplied by Sigma Aldrich).



71 Figure 1. Structure of 4,4'-Bismaleimidodiphenylmethane (a) and 2,2',diallylbisphenol A (b).

72

73 Materials were obtained from the following curing cycles:

- for BMI films: Thickening of reactive mixture during 10 min at 170°C, then 10 min at 175°C after which

samples were pressed using a laboratory press (240 bars without use of spacer) during 1 h at 180°C + 1
h à 190°C + 2 h à 200°C. The obtained films have a thickness about 40 μm.

- for BMI-DABPA films: Thickening of reactive mixture during 30 min at 150°C + 20 min at 180°C after
which samples were pressed using a laboratory press (240 bars without use of spacer)1 h at 180°C + 1 h
at 190°C + 2 h at 200°C. The obtained films have a thickness about 20 μm. Copolymers are detailed in
Table 1.

81

# 82 2.2. Exposure conditions

83 Samples were aged in ventilated ovens (AP60, SCS, France) under atmospheric air at 160°C. Ageing was

84 monitored by FTIR and elemental analysis.

85 Ageing under 100% N<sub>2</sub> or 100% O<sub>2</sub> atmosphere was performed in situ in TGA cell.

#### 87 2.3. Characterization methods

88

#### 89 2.3.1. ThermoGravimetric Analysis

TGA measurements were performed using a Q500 apparatus driven by QSeries Explorer (TA Instruments). After a 20°C min<sup>-1</sup> heating, isothermal degradation was performed at a constant temperature equal to 350 and 375°C either under 100%  $N_2$  or 100%  $O_2$  atmosphere supplied by a continuous 50 ml.min<sup>-1</sup> gas flow.

94

#### 95 2.3.2. Fourier Transform InfraRed spectroscopy

96 FTIR spectroscopy in transmission mode was carried out using a Frontier spectrophotometer
97 (PerkinElmer) by averaging 8 scans at 1 cm<sup>-1</sup> resolution between 400 and 4000 cm<sup>-1</sup>.

98

#### 99 2.3.3. Elemental Analysis

100 Carbon and hydrogen measurements were performed on a homemade Carbon / Hydrogen elemental 101 micro-analyzer (Institut des Sciences Analytiques, UMR5280 CNRS, Villeurbanne). About 1.5 mg of sample weighted in silver cups (3.2 × 6, Säntis Analytical AG, Teufen, Switzerland) dropped into a unit 102 103 combustion in a flow of 50 ml.min<sup>-1</sup> of pure oxygen. The combustion system is equipped by two 104 furnaces held at 1050°C for the upper one and 850°c for the lower and contain a vertical glass 105 combustion tube, half filled with 11 cm of CuO oxidation catalyst (Copper oxide wire 0.7mm, Säntis 106 Analytical) and silver wool (Silberwolle Carl ROTH GmbH, Karlsruhe, Germany) used as halogens trap. 107 Carbon and hydrogen have been respectively turned into carbon dioxide ( $CO_2$ ) and water ( $H_2O$ ). The 108 measurements of both molecules have been done using a CO<sub>2</sub>/H<sub>2</sub>O non-dispersive infrared detector 109 (Rosemount NGA 2000). A calibration has been daily established by measuring standards references 110 (molecules pure at 99.9%). Then up to 8 samples were injected. To be sure of the lack of calibrationdrift, 2 standards reference were introduced and measured as sample, every eight analyses. The 111 112 uncertainty of measurements has been determined at +/-0.10 for H measurement and +/- 0.30 for carbon measurement. Reference standards used for calibration and stability controls are high purity 113 compounds purchased from Sigma Aldrich corp. (Saint-Louis, Missouri, USA): Methionine (C: 40.25 %, 114 H: 7.43 %), Glycine (C: 36.36%, H: 6.10%), Taurine (C: 19.20%, H: 5.20%), Cystine (C: 29.99%, H: 5.03%), 115 Valine (C: 51.26, H: 9.46%), Tryptophan (C: 64.69%, H: 5.92%). 116

#### 118 <u>3. RESULTS</u>

119

#### 120 3.1. Thermal stability

121 The thermal stability of systems was first investigated by studying in situ degradation of BMI and BMI-122 DABPA networks using TGA either under inert atmosphere or under pure oxygen at elevated 123 temperatures (350 and 375°C) as shown in Figure 2. The following conclusions can be drawn:

BMI can be considered as relatively stable under inert atmosphere (less than 1% mass loss)
consistently with literature data [9] but the addition of DABPA comonomer promotes thermal
degradation.

127 - The presence of oxygen accelerates the degradation rate.

In presence of oxygen, the DABPA also accelerates degradation compared to pure BMI. However,
differences between BMI-DABPA 2-1, BMI-DABPA 1-1 and BMI-DABPA 1-2 under oxygen are weaker
than for degradations under nitrogen. BMI-DABPA 1-2 always is the less stable material for ageing
under inert atmosphere, but this is not necessarily the case under oxygen. A definitive explanation
would need to finely characterize the structure of copolymers with high DABPA content (presence of
homopolymerized DABPA sequences) and the stabilizing effect of phenolic groups, which is out of
reach using basic laboratory tools used in this work.







ratio					
theoretical	%C	0.704	0.738	0.757	0.776
	%O	0.179	0.156	0.144	0.131
	%Н	0.039	0.051	0.057	0.064
experimental	%С	0.693	0.699	0.713	0.702
	%O	0.188	0.159	0.187	0.205

		%Н	0.039	0.048	0.059	0.056					
150											
151	Table 1. Mass composition of the reactive mixture and elemental analysis of BMI and BMI-DABPA										
152	networks.										
153											
154	During thermal ageing, all materials display an increase in their oxygen content (illustrated as the ratio										
155	of oxygen concentration over carbon one $[^{20}]$ ). Despite some outliers, data also confirm that BMI-										
156	DABPA copolyme	ers oxidize	faster than BMI h	nomopolymer.							





159

Figure 3. Changes in elemental analysis.

160

161 In order to go further in the understanding of chemical structure and its changes, FTIR spectra analysis 162 were performed for unaged and aged samples. Spectra for unaged samples in several regions are given

163 in Figure 4.

164 As expected from their theoretical structure given in Scheme 1, BMI homopolymers are mainly

165 characterized by the methylene groups at 2930 cm<sup>-1</sup> (Figure 4a) and the presence of residual double

166 bonds concentration (Figure 4d).

167 The main following differences due to DABPA comonomer can be observed:

168 - the stretching of hydroxyl bond at around 3400 cm<sup>-1</sup> (Figure 4a)

169 - the stretching of C-H in methyl groups CH<sub>3</sub> at 2970 cm<sup>-1</sup> (Figure 4b)

170 - the C-O stretching of phenol groups at 1250 cm<sup>-1</sup> (Figure 4c)

171 - the out of plane bending of alkenes in the 600 – 1000 cm<sup>-1</sup> (Figure 4d). Here, absorbances of allyl at 172 989 and 914 cm<sup>-1</sup> [13] are not detected, meanwhile 832 ( $\delta_{C=C}$  in BMI) and 690 cm<sup>-1</sup> ( $\delta_{C=C+H}$  cis in BMI) 173 absorbances still seem present. It would suggest that networks are not fully polymerized [1]. We 174 however did not try to optimize curing degree, keeping in mind that post-curing must occur in situ 175 during the heating ramp prior to 350 and 375°C isothermal ageing for the experiments reported in 176 Figure 2.

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- 181
- 182





Figure 4. InfraRed spectra of BMI homopolymer and BMI-DABPA copolymers in the region of
 hydroxyls (a), C-H stretching (b), fingerprint (c) and C=C double bond bending region (d).

After thermal ageing at 160°C, the following differences (discussed in the following of this paper) were
systematically observed (Figures 5 and 6):

192 - an increase in the carbonyl region, more specifically in the 1600 to 1700 cm<sup>-1</sup> wavenumber range,

193 which could correspond to the appearance of benzophenone functions, or any other kinds of

194 conjugated ketones, consistently with previous results [16,<sup>21</sup>].

195 - the disappearance of aromatic =C-H stretching (above 3000 cm<sup>-1</sup>) which may be due to volatile loss

196 (see above) or isomerization. Actually, the appearance of a new absorbance at 930 cm<sup>-1</sup> accompanied

197 by the increase of a sharp band at  $1610 \text{ cm}^{-1}$ , could correspond to the appearance of new >C=C< double

198 bonds (see also Supplementary information SI-1).







Figure 6. FTIR spectra in the C-H stretching region for BMI (a), BMI-DABPA 2-1 (b), BMI-DABPA 1-1 (c) and BMI-DABPA 1-2 (d) thermally oxidized at 160°C under air.

Finally, our results unambiguously show that the use of DABPA as comonomer is detrimental to thermal stability of BMI based resins, either under nitrogen or in presence of oxygen.

216

218 4. DISCUSSION

219

The first aim of this section is to propose an explanation of the chemical changes in link with the initial structure of networks, and try to explain why diallylbisphenol A lowers thermal stability of BMI-DABPA copolymers. The second aim is to propose a reasonable kinetic model based on the reactivity of each comonomer so as to describe the reactivity of the family of BMI-DABPA networks.

224

#### 225 4.1. Structural changes observed in BMI-DABPA

226

Let us start by recalling the theoretical structure of BMI and BMI-DABPA 2-1 copolymers (Scheme 1). In
BMI homopolymers, the polymerization occurs along the double bonds (even in absence of radical
initiator), as depicted in Scheme 1. In BMI-DABPA copolymers, polymerization occurs in two stages: a
Michael addition of alkene (hold by diallylbisphenol A) followed by a Diels Alder reaction with another
bismaleimide group. The ideal stoichiometry is here one diallyl group for two maleimides ones.

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- 235
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237

238

239





(b)

248

246

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Scheme 1. Curing mechanism in BMI (a) and BMI-DABPA (b).

249

Since only BMI and BMI-DABPA 2-1 correspond to well defined structures, we will focus on those two
networks in the following, keeping in mind that thermal stability of BMI-DABPA 2-1, BMI-DABPA-1-1
and BMI-DABPA 1-2 seem somewhat close.

To better understand the differences in terms of "structure-stability relationships", some representative units of BMI homopolymer and BMI-DABPA 2-1 groups were analyzed using a recently developed Machine Learning tool (Alfabet) allowing the determination of Bond Dissociation Energies,
 and the possible « Achylle Heel » of each kind of polymer. Such an approach was recently successfully
 tested on epoxies [<sup>22</sup>]. Using SMILES codes given in Supplementary Information SI-2, some BDE
 corresponding to weak groups are given in Figure 7.





In the case of copolymer, it seems clear that the allylic C-H are easily oxidized into various kinds of
ketones. In the case of copolymers, it is hence not surprising that the carbonyl region displays a new
signal located between 1750 et 1800 cm<sup>-1</sup>.



Scheme 3. Oxidation of allylic sites in BMI-DABPA.

287 288

The qualitative comparison of oxidized spectra for BMI homopolymer and BMI-DABPA copolymers suggests that copolymers are less stable than homopolymers. This is well in line with elemental analysis results. Dealing now with mass loss curves, it seems that oxidation generates more volatile compounds in the case of copolymers. We will now try to derive a kinetic model for describing the degradation kinetic and in particular mass loss curves of materials.

294

#### 295 4.2. Kinetic modelling

296

We based on the general mechanistic scheme used for describing the oxidation of several polymers. This latter was in particular implemented for describing the oxidation of bismaleimide networks by Colin et al [17]. In its initial formulation, the model assumed that maleimides were the only reactive sites which is not the case here since there are strong discrepancies between BMI and BMI-DABPA 2-1. Instead of using a simple model with « genuine » constants for BMI and BMI-DABPA systems, we decided to develop a cooxidation kinetic model [<sup>23</sup>,<sup>24</sup>] where reactive sites for BMI (subscript 1) and DABPA (subscript 2) react independently or by crossed reactions and where the reactivity of BMI- 304 DABPA copolymers is ruled by rates constants specific to each site. Moreover, our results (in particular 305 of mass loss data under inert atmosphere) showed the existence of a « pure » thermolytical process 306 that we simply modeled by reaction (0) generating two alkyl radicals and (4) corresponding to their 307 termination by coupling. The mechanistic scheme is written as follows:

308

309	Initiation	$P_1H \rightarrow P_1^{o} + \gamma_{01}V_{01}$	$k_{01}$
310	Initiation	$P_{2}H\toP_{2}^{\circ}+\gamma_{02}V_{02}$	<b>k</b> <sub>02</sub>
311	Initiation	$P_1OOH + \gamma P_1H + (1-\gamma)P_2H \rightarrow 2(1-\gamma_{K1}).\gamma P_1^{\circ} + 2(1-\gamma_{K1}).(1-\gamma)P_2^{\circ} H_2O + \gamma_1V_1$	<b>k</b> <sub>11</sub>
312	Initiation	$P_2OOH + \gamma P_1H + (1-\gamma)P_2H \rightarrow 2(1-\gamma_{K2}).\gamma P_1^{\circ} + 2(1-\gamma_{K2}).(1-\gamma)P_2^{\circ} H_2O + \gamma_2V_2$	<b>k</b> <sub>12</sub>
313	Propagation	$P_1^{\circ} + O_2 \rightarrow P_1 OO^{\circ}$	<b>k</b> <sub>21</sub>
314	Propagation	$P_2^{\circ} + O_2 \rightarrow P_2OO^{\circ}$	<b>k</b> <sub>22</sub>
315	Propagation	$P_1OO^{\circ} + P_1H \rightarrow P_1OOH + P_1^{\circ}$	<b>k</b> <sub>311</sub>
316	Propagation	$P_1OO^{\circ} + P_2H \rightarrow P_1OOH + P_2^{\circ}$	<b>k</b> <sub>312</sub>
317	Propagation	$P_2OO^{\circ} + P_1H \rightarrow P_2OOH + P_1^{\circ}$	<b>k</b> <sub>321</sub>
318	Propagation	$P_2OO^{\circ} + P_2H \rightarrow P_2OOH + P_2^{\circ}$	<b>k</b> <sub>322</sub>
319	Termination	$P_1^{o} + P_1^{o} \rightarrow \text{inactive products}$	<b>k</b> 411
320	Termination	$P_1^{o} + P_2^{o} \rightarrow inactive products$	<b>k</b> <sub>412</sub>
321	Termination	$P_2^{o} + P_2^{o} \rightarrow inactive products$	<b>k</b> <sub>422</sub>
322	Termination	$P_1^{o} + P_1OO^{o} \rightarrow inactive products$	k <sub>511</sub>
323	Termination	$P_1^{o} + P_2OO^{o} \rightarrow inactive products$	k <sub>512</sub>
324	Termination	$P_2^{\circ} + P_1OO^{\circ} \rightarrow inactive \text{ products}$	<b>k</b> <sub>521</sub>
325	Termination	$P_2^{\circ} + P_2OO^{\circ} \rightarrow inactive products$	<b>k</b> <sub>522</sub>
326	Termination	$P_1OO^\circ + P_1OO^\circ \rightarrow inactive \ products + O_2$	<b>k</b> <sub>611</sub>
327	Termination	$P_1OO^\circ + P_2OO^\circ \rightarrow \text{inactive products} + O_2$	<b>k</b> <sub>612</sub>
328	Termination	$P_2OO^\circ + P_2OO^\circ \rightarrow inactive \ products + O_2$	<b>k</b> <sub>622</sub>

329 Meaning of symbols are detailed in Table 2.

Symbol	Meaning
$P_1H$ , $P_2H$	Unreacted susbtrate
$P_1^{o}, P_2^{o}$	Alkyl radical
P <sub>1</sub> OO°, P <sub>2</sub> OO°	Peroxy radical
P <sub>1</sub> OOH, P <sub>2</sub> OOH	Hydroperoxide
V <sub>01</sub> , V <sub>02</sub>	Volatiles genenerated by thermolysis
[] <sub>01</sub> , [] <sub>02</sub>	Yield for volatile generation from substrate decomposition
V <sub>1</sub> V <sub>2</sub>	Volatiles generated by thermal oxidation
$[]_1 \bigcup [g_2]$	Yield for volatile generation from oxidation
$\upsilon_{01}M_{v01}, \upsilon_{02}M_{v02}$	Contribution to mass loss from substrate decomposition
$\upsilon_{01}M_{v01}, \upsilon_{02}M_{v02}$	Contribution to mass loss from oxidation
У	Probability of $P_1H$ or $P_2H$ attack by any HO° or $P_iO^\circ$ radical
<b>Ук1, Ук</b> 2	Yield for cage reaction $P_iO^o + HO^o \to inactive\ product\ (carbonyl)$
ki, kij	Rate constant
E <sub>i</sub> , E <sub>ij</sub>	Activation energy

332

# Table 2. Signification of kinetic parameters.

333

The balance equation for initiation are justified in Supplementary Information SI-3. This scheme leads to the differential system given in Supplementary Information SI-4. It allows mass loss to be predicted from :

337

$$\frac{d \frac{m}{m_0}}{dt} = \frac{32}{\rho} \left( k_{21} [P_1^{\circ}] [O_2] + k_{22} [P_2^{\circ}] [O_2] - k_{611} [P_1 OO^{\circ}]^2 - k_{612} [P_1 OO^{\circ}] [P_2 OO^{\circ}] - k_{622} [P_2 OO^{\circ}]^2 \right) - \frac{(18 + v_1 M_{v_1} P_1 OO^{\circ})}{\rho}$$

339

340 System was solved using ODE23s from Matlab<sup>®</sup> using the following initial conditions at t = 0,  $[P_1^o]_0 =$ 341  $[P_2^o]_0 = [P_1OO^o]_0 = [P_2OO^o]_0$ 

342

- 343 The initial substrate concentration was calculated as follows:
- $344\quad \ \ \, \mathbb{O}$  For pure BMI, each monomer holds a diaryl methylene group so that:

345 
$$[P_1H]_0 = \frac{\rho}{M_1} = 3.6 \, mol \, l^{-1}$$

346  $M_1$  being the molar mass of BMI (358 g mol<sup>-1</sup>)

347 ② For BMI-DABPA 2-1, concentration in BMI comonomer and diallylbisphenol A is weighted by the
348 weight ratio of BMI in the mixture considering that each diallylbisphenol A holds 4 reactive sites:

349 
$$\left[P_{1}H\right]_{0} = \frac{\rho}{M_{1}} \cdot \frac{2M_{1}}{2M_{1} + M_{2}} = 2.5 \, mol \, l^{-1}$$

350 
$$[P_2H]_0 = 4 \frac{\rho}{M_2} \cdot \frac{M_2}{2M_1 + M_2} = 5 \, mol \, l^{-1}$$

351 M<sub>2</sub> being the molar mass of DABPA (308 g mol<sup>-1</sup>)

352 The total initial concentration in hydroperoxide was arbitrarily fixed equal to 0.001 mol l<sup>-1</sup> so that:

353 - For pure BMI:

$$[P_1OOH]_0 = 0.001 \, mol \, l^{-1}$$

355 - For BMI-DABPA 2-1 copolymer:

356 
$$[P_1OOH]_0 = 0.00035 \, mol \, l^{-1}$$

357 
$$[P_2OOH]_0 = 0.00065 \, mol \, l^{-1}$$

358

359 The oxygen concentration was fixed equal to 0.04 mol l<sup>-1</sup> consistently with [<sup>25</sup>].

360

The main complexity is the high number of parameters in the kinetic model. Many of them were fixed in order to have the less possible number of adjustable parameters. Here, we implemented the following approach:

364 - Rate constants were first determined for pure BMI and then for BMI-DABPA 2-1.

- Under nitrogen, only  $k_{0i}$  and  $k_{4ij}$  are needed to fit the curves of mass loss. Those values were then utilized for simulating degradation under oxyen without change. - The rate constants k<sub>01</sub>, k<sub>02</sub> and the yields for volatile release were adjusted from the parametric study
presented in Figure 2 by fitting the curves under inert atmosphere. Figure 8a illustrates a parametric
study for the k<sub>0</sub> value.

- According to the analysis of oxidation mechanisms presented in the previous section, reactive sites were identified because of their bond dissociation energies. It allows the estimation of the rate constants values for the POO<sup>o</sup> + PH  $\rightarrow$  POOH + P<sup>o</sup> values from the semi-empirical values proposed by Korcek et al [<sup>26</sup>]:

374 
$$\log_{10}k_3^{sec-POO^{\circ}}(30 \circ C) = 16.4 - 0.2 * BDE$$

375 
$$E_{3}^{sec-POO^{\circ}} = 0.55 \times (BDE - 62.5)$$

376 
$$\log_{10} k_3^{tert-POO^{\circ}}(30 \circ C) = 15.4 - 0.2 * BDE$$

377 
$$E_3^{tert-POO^\circ} = 0.55 \times (BDE - 65)$$

- The rate constants for termination  $(k_{4ij}, k_{5ij}, k_{6ij})$  should in principle be determined from a study under varying oxygen pressures [<sup>27</sup>]. In the absence of this latter, we used the following assumptions: (i)  $10^{10} >$  $k_{4ij} > k_{5ij} >> k_{6ij}$ , (ii)  $k_{5ij} = 5 \times 10^8$  l mol<sup>-1</sup> s<sup>-1</sup> and  $k_{6ij} = 10^5$  l mol<sup>-1</sup> s<sup>-1</sup> keeping in mind that the rate constant for the termination between two POO° ( $k_{6ij}$ ) mainly influences the kinetics under high oxygen pressures, (iii)  $k_{412} = (k_{411} \times k_{422})^{1/2}$  [23,24]. The effect of  $k_{4ij}$  is illustrated by the parametric study given in Figure 8f. Let us precise that the termination product of two radicals can be a double bond, consistently with experimental observations (Figure 5).

- Finally,  $k_{1ij}$ ,  $k_{2ij}$ ,  $k_{0i}$  and  $k_{4ij}$ ,  $\Box_{ki}$  were the remaining adjustable parameters. We assumed that  $k_{21} = k_{22}$ . The results of a parametric study for those constants are depicted in Figures 8b, 8c and 8d. For summarizing:
- k<sub>1i</sub> was adjusted basing on the initial concavity of the curve in Figure 2. We also checked its consistency
  with comparable results obtained by Colin et al [17] (Figure 10b).
- k<sub>2i</sub> was adjusted on the mass loss rate at early stages. It must however be recognized that several k<sub>2</sub>,
  k<sub>4</sub> sets would lead to comparable simulations.
- $k_{3ij}$  were fixed but their influence is depicted for example in Figure 8e.

393

Another attempt to validate the model was to fit the data by Colin et al [17], despite the fact that the
 formulation used in their work may not exactly match with the BMI and BMI DABPA 2-1 studied in this

work. Results are given in Figure 9 where two cases were considered: the fictive case of a pure BMI
resin and the case of BMI DABPA 2-1 (on which rate constants were adjusted in a consistent manner
with rate constants determined from Figure 9).

399

Finally, it can be seen that kinetic model describes the mass loss kinetics (Figures 2 and 9) in a wide range of temperatures (180 to  $375^{\circ}$ C) with elementary rate constants obeying Arrhenius law with reasonable activation energies (Figure 10):  $E_{1i} \sim 140 \text{ kJ mol}^{-1}$ ,  $E_{2i} \sim E_{4i} \sim 15 \text{ kJ mol}^{-1}$ .

403 Further improvements can be envisaged in order to optimize the fits under oxidative conditions:

404 - here, we assumed quasi ideal structures which may not the case for real networks. For example, BMI-

405 DABPA networks can possibly BMI or DABPA homopolymerized sequence. Their reactivity must be

406 understood and some reactions should be added to the model. However, it is first needed to evidence

- 407 the presence of such sequences in networks.
- 408 the participation of sites having a lower reactivity (such as the -CH-CH-CH-CH- sequence in pure BMI).

409 - the actual effect of curing degree could also be taken into account [<sup>28</sup>].

410 - we assumed that some parameters (namely  $\upsilon_1 M_{v1}$ ,  $\upsilon_2 M_{v2}$ ,  $\upsilon_{01} M_{v01}$ ,  $\upsilon_{02} M_{v02}$ ) remain constant during

411 the whole timescale under consideration which is clearly an approximation.

412 - in particular, the possibility of retro Diels Alder occurring under nitrogen for BMI-DABPA is to413 investigate.



	180°C	210°C	240°C	350°C	375°C	E <sub>a</sub> (kJ mol⁻¹)	
k <sub>11</sub> (s <sup>-1</sup> )	4×10 <sup>-7</sup>	2.75×10 <sup>-</sup>	2.5×10 <sup>-5</sup>	1.1×10 <sup>-2</sup>	2.75×10 <sup>-2</sup>	142.4	
k <sub>12</sub> (s <sup>-1</sup> )	2.7×10 <sup>-6</sup>	1.65×10 <sup>-</sup> 5	1.3×10 <sup>-5</sup>	5.0×10 <sup>-2</sup>	1.4×10 <sup>-1</sup>	139.3	
k <sub>21</sub> (I mol <sup>-1</sup> s <sup>-1</sup> )	2.5×10 <sup>4</sup>	3.1×10 <sup>4</sup>	3.75×10 4	6.5×10 <sup>4</sup>	7.5×10 <sup>4</sup>	13.6	
k <sub>22</sub> (I mol <sup>-1</sup> s <sup>-1</sup> )	2.5×10 <sup>4</sup>	3.1×10 <sup>4</sup>	3.75×10 4	6.5×10 <sup>4</sup>	7.5×10 <sup>4</sup>	13.6	
k <sub>311</sub> (I mol <sup>-1</sup> s <sup>-1</sup> )	366	768	1912	9450	13200	44.9	
k <sub>321</sub> (I mol <sup>-1</sup> s <sup>-1</sup> )	17	46	58	292	391	39.1	
k <sub>312</sub> (I mol <sup>-1</sup> s <sup>-1</sup> )	588	1100	1475	9201	12210	38.0	
k <sub>322</sub> (I mol <sup>-1</sup> s <sup>-1</sup> )	28	33	75	285	362	32.2	
k <sub>411</sub> (I mol <sup>-1</sup> s <sup>-1</sup> )	5×10 <sup>8</sup>	7×10 <sup>8</sup>	9×10 <sup>8</sup>	2.2×10°	2.4×10 <sup>9</sup>	20.0	
k <sub>422</sub> (I mol <sup>-1</sup> s <sup>-1</sup> )	8×10 <sup>8</sup>	1.15×10°	1.5×10°	3.2×10°	3.2×10 <sup>9</sup>	18.4	
k <sub>412</sub> (I mol <sup>-1</sup> s <sup>-1</sup> )		0					
k <sub>511</sub> (I mol <sup>-1</sup> s <sup>-1</sup> )	5×10 <sup>8</sup>	5×10 <sup>8</sup>	5×10 <sup>8</sup>	5×10 <sup>8</sup>	5×10 <sup>8</sup>	0	
k <sub>512</sub> (I mol <sup>-1</sup> s <sup>-1</sup> )	5×10 <sup>8</sup>	5×10 <sup>8</sup>	5×10 <sup>8</sup>	5×10 <sup>8</sup>	5×10 <sup>8</sup>	0	
$k_{512} = k_{521} (I \text{ mol}^{-1} \text{ s}^{-1})$		(k <sub>511</sub> ×k <sub>522</sub> ) <sup>0.5</sup>					
k <sub>611</sub> , k <sub>612</sub> , k <sub>622</sub>	10 <sup>8</sup>	10 <sup>8</sup>	10 <sup>8</sup>	10 <sup>8</sup>	10 <sup>8</sup>	0	
Ук1	1	1	1	1	1		
$\gamma_1$	0	0	0	0	0		
Ук2	1	1	1	0.8	0.8		
$\gamma_2$	0	0	0	0	0		
$\upsilon_1 M_{v1}$ (g mol <sup>-1</sup> )	25	29	32.5	85	117.5		

$\upsilon_2 M_{v_2}$ (g mol <sup>-1</sup> )	45	50	51.5	130	95	
0	1200	1200	1200	1200	1200	
P P	1300	1300	1300	1300	1300	
k <sub>01</sub> (s <sup>-1</sup> )	6×10 <sup>-9</sup>	4×10 <sup>-8</sup>	2×10 <sup>-7</sup>	5.5×10 <sup>-5</sup>	1.5×10 <sup>-4</sup>	126.9
υ <sub>01</sub> Μ <sub>ν01</sub> (g mol⁻¹)	0	0	0	1	4	
k <sub>02</sub> (s⁻¹)	7×10 <sup>-8</sup>	5×10 <sup>-7</sup>	3.0×10 <sup>-6</sup>	8.5×10 <sup>-4</sup>	1.6×10 <sup>-3</sup>	129.8
υ <sub>02</sub> Μ <sub>ν02</sub> (g mol⁻¹)	0	0	0	1.75	2	

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# Table 3. Kinetic parameters for simulations given in Figures 2 and 9.



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Figure 9. Simulation of results by Colin et al [17] with kinetic model working with  $[P_1H]_0 = 2.5$  and  $[P_2H]_0 = 5 \text{ mol } I^{-1} (BMI-DABMA 2-1) \text{ or } [P_1H]_0 = 3.6 \text{ mol } I^{-1} \text{ and } [P_2H]_0 = 0 \text{ (pure BMI)}.$ 





432

Figure 10. Arrhenius diagram for rate constants.

# 433 CONCLUSIONS

In this paper, networks made from bismaleimide and diallylbisphenol A were synthesized and 434 435 submitted to isothermal ageing at various temperatures under inert atmosphere and or under air. 436 Results clearly show that diallyl comonomer lowers thermal stability of BMI resins. This was confirmed 437 by an estimation of the Bond Dissociation Energies of structural units of BMI and BMI-DABPA since 438 DABPA displays some weak bonds that can more easily undergo homolysis and radical attack. Basing on 439 those observations, a kinetic model for co-oxidation was proposed. The complexity of such modelling 440 tools is the high number of parameters that can influence the simulations. In order to decrease the 441 number of adjustable parameters, some rate constants values were and some other were identified from a parametric study. Finally, the kinetic model displays an encouraging capability to simulate mass 442 443 loss induced by thermal ageing of BMI based networks as confirmed by comparing its prediction with 444 literature data. Its validation process can be pursued by performing long term ageing runs at lower 445 temperatures.

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