Aryltriazene Photopolymers for UV-Laser Applications: Improved Synthesis and Photodecomposition Study

Matthias Nagel,* Roland Hany, Thomas Lippert, Martin Molberg, Frank A. Nüesch, Daniel Rentsch

An improved synthesis of photosensitive homopolymers containing aryltriazene chromophores covalently incorporated into the polymer backbone is reported. Such photopolymers proved to have promising properties for novel UV-laser applications. A homologous series of new aryltriazene polymers with increasingly branched side chains ($R = \text{Me, Et, iPr, tBu}$) was synthesized and characterized. Homogeneous thin films with thicknesses from $\approx 15$ to $>150$ nm were prepared by spin-coating. Photodecomposition was studied in solution and on thin films. Polymers with increasingly branched and bulky substituents showed decreasing photodissociation rates. NMR studies suggested an enhanced hindrance of the $N(2)\text{–}N(3)$ bond rotation in the aryltriazene moiety with increasing steric demand of the substituents.

Introduction

Since the first reports about laser ablation of polymers in 1982, a wide variety of photopolymers tailored for laser-induced polymer processing, ranging from surface modification, ablative photodecomposition, to thin film deposition have been investigated. [1] Designed polymers, used as sacrificial absorbing layers for laser-induced forward transfer (LIFT) techniques, are increasingly finding applications in direct-writing operations for the controlled transfer and microdeposition of materials. [2] Various polymeric composite materials (usually a binder matrix doped with dispersed absorber dyes) have been applied as energy-absorbing dynamic release layer (DRL) systems, e.g., for high-resolution full-color printing, [3] transfer of single molecules [4] as well as biologic cellular systems, [5a–e] release of prefabricated parts in the assembly of microelectromechanical systems [5f] or structured transfer of semiconducting polymers used in organic light-emitting diodes. [5g,5h] One of the common issues of these light-to-heat conversion layers based on IR-lasers is the intrinsically high thermal load on sensitive materials to be transferred. [6] In order to avoid overheating defects and heat-induced side-reactions, [6] we developed a modified concept of the LIFT process using a thin intermediate sacrificial photopolymer layer which is distinctly UV-sensitive.
and decomposes photochemically more gentle into small and volatile fragments.\(^7\)

Aryl-dialkyltriazene polymer films can be efficiently decomposed and ablated when irradiated with UV lasers.\(^2\) Two Ar–N=N–NAr moieties as photolabile chromophores per repeating unit are covalently integrated into the polymer backbone (Scheme 2). Photolysis leads to a controlled photofragmentation with evolution of elemental nitrogen. Synthesis of the first arytriazene homopolymer TP-6a was published in 1993,\(^9\) and, due to its favorable ablation characteristics,\(^3\) it became an important reference compound within the last decade.\(^8,10\) Related photodegradable polymers based on the arylazo parent system were synthesized, such as poly(alkylarylazo sulfonate)s\(^11\) and poly(arylazophosphonate)s,\(^12\) diazosulfone polymer films,\(^13\) triazene containing copolymers,\(^14\) and triazene-unit containing polyurethanes\(^15\) as well as triazene polyacrylates.\(^15\) However, with regard to practical aspects such as shelf-life, chemical compatibility, and their specific absorption range, triazene polymers are still the most attractive candidates for applications as the key part in photoinduced release layer systems.

For LIFT applications which require excellent film-forming properties,\(^2\) only thin polymer films with thickness of a few hundred nanometers at most are used. Until now, use of TP-6a as thin-film DRL was impeded due to its poor solubility and the formation of inhomogeneous solutions with crosslinked gel-type swollen lumps that are caused by undesirable side reactions. Here, we report the optimization of the synthetic protocol for TP-6a and a homologous series of new triazene homopolymers derived from it which all allow the fabrication of smooth thin films. We compare the photolytic decomposition of triazenes in solid state as well as in solution and discuss the influence of side-chain groups on the relative polymer stability subsequent to absorption.

**Experimental Part**

**Synthesis of Triazene Homopolymers TP-6a–TP-6d**

To 3.0 g of 4,4’-oxydianiline (DAO, 15 mmol) were added 30 mL of water and 7.5 mL of 37 wt.-% hydrochloric acid with stirring at room temperature. After complete dissolution of DAO, the solution was cooled to 0 to −5 °C and then 2.1 g sodium nitrite (30 mmol) dissolved in 20 mL of water was added dropwise within 10–15 min, keeping the temperature below 0 °C and after complete addition, the yellow bisdiazonium salt solution was stirred for a further 15–20 min. The corresponding diamine component DA-6 (15 mmol) was dissolved in 50 mL of ice-cold water containing 3.5 mL of conc. HCl. The two cold solutions were then added to a mixture of 150 g crushed ice and 100 mL of hexane. The biphasic mixture was mechanically stirred vigorously and 60 mL of a 2 M ice-cooled aqueous K$_2$HPO$_4$ solution (pH ≈ 8.5) was added from a dropping funnel. The color of the mixture changed slightly to dark yellow (pH 7–8). Then 60 mL of a cooled 2 M K$_2$PO$_4$ buffer solution (pH ≈ 12) was added dropwise. With rising pH values, the mixture became turbid and the precipitation of the polymer started from pH ≈ 8.5–9. The mixture became more viscous and after complete addition of the buffer solution, the pH of the reaction mixture was adjusted to 10–11 by the dropwise addition of 2 M KOH. The suspension was then stirred for 2–3 h while warming to room temperature. The light beige polymeric product was then filtered off and washed subsequently with hexane, tert-butyl methyl ether, isopropanol, methanol, and finally several times with water to remove inorganic salt residues. After drying overnight in a vacuum chamber, the raw polymer was dissolved in 120–150 mL of THF and the slightly viscous orange solution was then separated by centrifugation from a small fraction of insoluble viscous and dark residues. The supernatant clear polymer solution was then precipitated from 800 mL of methanol as sponge-like lumps, which were separated from the turbid solution by sieving. The product was washed with methanol and water and dried to a constant weight (yields > 85% related to the DAO starting material).

**Synthesis of Disecondary Diamines DA-6a–DA-6d**

Reactions were performed in a 1 L three-necked reaction vessel equipped with a 250 mL dropping funnel, thermometer, magnetic stirrer, and a chiller-cooled reflux condenser (−5–10 °C) with a gas reservoir on top to keep volatile amines in the reactor during the exothermic reaction. The monoamine (3 mol) dissolved in 100 mL THF was heated to 45–50 °C and a solution of 122 g 1,6-dibromo-hexane (0.5 mol) in 75 mL THF was added dropwise, such that a slight reflux is maintained. After addition of about half of the dibromoalkane, the mixture became more and more turbid and crystalline mass started to precipitate and after complete addition of 1,6-dibromohexane, the mixture was stirred for 3 h at a slight reflux. A concentrated aqueous solution of 68 g of KOH (1.2 mol) dissolved in ≈40 g crushed ice was added and the resulting mixture stirred over night at room temperature. The excess of monoamine and THF were distilled off at atmospheric pressure, the remaining waxy mass was suspended in 300 mL of toluene and water was then removed by azotropic distillation. The crystalline mass of KBr could easily be separated from the organic phase by filtration. The solid phase was washed twice with toluene and the solvent distilled off from the collected organic filtrate fractions under reduced pressure. The remaining oil was purified by fractional distillation which gave the diamines as highly hygroscopic colorless liquids.

**Characterization**

NMR experiments were performed on a Bruker Avance-400 spectrometer using a 5 mm broadband inverse probe with z-gradient. Measurements were carried out at 297 K with ≈30 mg of sample dissolved in 0.7 mL of solvent. Chemical shifts are given in ppm relative to the remaining signals of chloroform at 7.26 ppm.
(1H) and 77.7 ppm (13C). For 1H, 13C 2D correlation experiments, Bruker standard pulse programs and parameters were used.

**TP 6a.** 1H NMR: δ = 1.12 (–NH2), 1.27 [m, –N(CH2)6CH2–], 1.41 (m, –N(CH2)2CH2–), 2.35 [s, –N(CH3)2], 2.49 (t, J = 7.1 Hz, –N–CH2–), 1.35 [m, –N(CH2)2CH2–], 2.46 (t, J = 7.2 Hz, –N–CH2–), 2.50 (q, J = 7.2 Hz, –N–CH2–), 3.05 [–N–CH2–CH2–], 3.71 [2–CH2–CH2–], 52.7 [–N–CH2–CH2–], 6.96 (t, J = 7.2 Hz, –N–CH2–), 2.10 [m, –N(CH2)2CH2–], 1.00 [m, –N(CH2)2CH2–], 4.17 (m, –N(CH2)6CH2–), 2.46 (t, J = 7.2 Hz, –N–CH2–), 2.50 (q, J = 7.2 Hz, –N–CH2–), 3.05 [–N–CH2–CH2–], 3.71 [2–CH2–CH2–], 29.8 [–N–CH2–CH2–], 44.4 [–N–CH2–CH2–], 119.7, 122.2, 148.2, 155.6 (Ar–C), [–N(CH3)2], not detected.

DA 6a: 1H NMR: δ = 1.12 (–NH2), 1.27 [m, –N(CH2)6CH2–], 1.41 (m, –N(CH2)2CH2–), 2.35 [s, –N(CH3)2], 2.49 (t, J = 7.1 Hz, –N–CH2–), 1.35 [m, –N(CH2)2CH2–], 2.46 (t, J = 7.2 Hz, –N–CH2–), 2.50 (q, J = 7.2 Hz, –N–CH2–), 3.05 [–N–CH2–CH2–], 3.71 [2–CH2–CH2–], 52.7 [–N–CH2–CH2–], 6.96 (t, J = 7.2 Hz, –N–CH2–), 2.10 [m, –N(CH2)2CH2–], 1.00 [m, –N(CH2)2CH2–], 4.17 (m, –N(CH2)6CH2–), 2.46 (t, J = 7.2 Hz, –N–CH2–), 2.50 (q, J = 7.2 Hz, –N–CH2–), 3.05 [–N–CH2–CH2–], 3.71 [2–CH2–CH2–], 29.8 [–N–CH2–CH2–], 44.4 [–N–CH2–CH2–], 119.7, 122.2, 148.2, 155.6 (Ar–C), [–N(CH3)2], not detected.

CHN contents were determined on a varioEL (Foss-Heraeus). **TP 6a** (C20H24N8O8), 366.46 g mol−1. Calcul. C 55.55, H 7.15, N 22.93; Found C 65.48, H 7.18, N 23.05. **TP 6b** (C22H26N10O8), 394.51 g mol−1. Calcul. C 66.98, H 7.66, N 23.10; Found C 66.78, H 7.63, N 23.13. **TP 6c** (C24H26N10O8), 422.57 g mol−1. Calcul. C 68.22, H 8.11, N 19.89; Found C 68.04, H 8.09, N 20.06. **TP 6d** (C30H24N10O8), 450.63 g mol−1. Calcul. C 69.30, H 8.50, N 18.65; Found C 69.63, H 8.46, N 18.75.

**Results and Discussion**

**Reported Polymer Synthesis**

According to the literature protocol[9a,9b] summarized in Scheme 1, the synthesis of **TP 6a** starts with the transformation of bisaminol precursor, DAO, into its bisdiazo- nitrite derivative, BDS, by a diazotation reaction. The acidification of the freshly prepared BDS is then treated with an excess of sodium carbonate dissolved in water, raising the pH to strongly alkaline values. Subsequently, a solution of diamino compound DA 6a dissolved in hexane is added, and the interfacial polycondensation reaction starts immediately. The polymer is insoluble in both solvents and precipitates as beige flakes which are filtered.
For purification, a solution of the polymeric reaction products in THF is precipitated in methanol, and the reported yields are 63% of TP-6a related to DAO.\[9b\] Following the protocol repeatedly, the colors of our polymeric products ranged from light-beige to brown. The darker products had significantly lower solubility in THF and the solutions were more viscous or even gel-like (Figure 1). Therefore, the purification step was impeded by the low solubility of some polymer batches and yields were low. Attempts to spin-coat such viscous polymer solutions into homogeneous thin films failed.

We attribute these findings to side reactions that induce crosslinking reactions in parallel to polymerization (Scheme 1): One significant drawback of the reported synthesis is the inhomogeneous reaction conditions during the addition of DA-6a. It can only be added slowly, since the elimination of HCl formed during the condensation step leads to the formation of CO$_2$ bubbles. The precipitating polymer suspension, therefore, forms a foaming mixture that is difficult to homogenize, and insufficient neutralization conditions can induce a protolytic back-cleavage of the formed triazene polymer.\[8,9c,18\]

In addition, alkaline solutions of BDS are unstable and we noticed a progressive darkening after addition of sodium carbonate. When DA-6a was added to “aged” solutions of BDS, mainly dark-colored and only partly soluble products were obtained. In alkaline solutions, aryldiazonium compounds decompose with the evolution of molecular nitrogen, forming aryl radicals that recombine to biaryl species (Gomberg–Bachmann reaction): In the presence of hydroxide ions, BDS is transformed in part into diazohydroxide intermediates (Scheme 1, species A) that, with increasing pH values, are deprotonated and transformed into diazotates (B). B reacts with a further diazonium moiety into diazoanhydride compounds C that decompose quickly with evolution of N$_2$ into arene radicals D. The interfering side reactions (A → D) start immediately after reaching alkaline pH values, and thus before DA-6 itself is added. Therefore, depending on the speed of addition of the coupling compound, the competing reactions lead to arene radical species D at the same time. These attack either aryl moieties of the starting
materials or of already formed triazene-polymer chains, causing crosslinking and chain termination.

**Improvement of the Synthetic Protocol**

In order to circumvent these problems, the reaction conditions for the polycondensation were adapted (Scheme 2). First, the carbonate solution was replaced by a non-effervescent neutralizing agent that allows a reproducible pH control of the reaction mixture. In order to avoid local high alkali concentrations, we chose the phosphate buffer system that allows to adjust a controlled pH gradient by adding solutions of K2HPO4 (pH ≈ 8.5) and K3PO4 (pH ≈ 12, each 2M in water). Furthermore, DA-6a was converted into its water-soluble bishydrochloride derivative. This component can then be added in aqueous solution as a protected reactant directly to the acidic solution of the bisdiazonium salt without any interaction. With both reaction components in the reaction mixture, the polycondensation reaction can be triggered by raising the pH of the reaction mixture up to ≈ 8, where the diamino compound is liberated from its corresponding “protected” ammonium precursor form. The in situ liberated bisamine DA gets dissolved in the organic phase (hexane) of the biphasic reaction mixture and the interfacial polycondensation starts. Subsequently, the pH is increased by addition of K3PO4 and the precipitation of the polymer typically started at pH ≈ 9, where the unwanted formation of diazotates in the equilibrium is negligible (Scheme 1, B). The polymeric products were dissolved in THF and precipitated from methanol (yields relative to DAO > 85–90%).

**Physical Properties**

All GPC curves showed a similar profile with average polydispersity values D in the range of 4.5–5.0 (Table 1); these are typical values for an interfacial polycondensation. Number- and mass-average molar masses ranged from 25 000 to 45 000 g mol⁻¹ for $M_n$, and 140 000 to 180 000 g mol⁻¹ for $M_w$. For TP-6a, these values are different from those of former batches synthesized according to the published protocol,[9a] where we obtained $M_n$ and $M_w$ of ≈ 15 000 and ≈ 70 000 g mol⁻¹, in accordance with the published values.[9a,9b] The higher molar masses of the new TP-6 series suggest that the modified reaction conditions avoid uncontrolled chain terminations. All polymers started to decompose at a temperature of ≈ 230 °C.

**NMR Studies**

The $^1$H NMR spectra of TP-6a–TP-6d in Figure 2 demonstrate the purity of the triazene polymers. The protons of

<table>
<thead>
<tr>
<th>TP</th>
<th>Yield</th>
<th>$M_n$ a,b) g mol⁻¹</th>
<th>$M_w$ a,b) g mol⁻¹</th>
<th>$M_{max}$ a,b) g mol⁻¹</th>
<th>$D$ a)</th>
<th>$T_g$ c) °C</th>
<th>$T_D$ d) °C</th>
</tr>
</thead>
<tbody>
<tr>
<td>TP-6a-R e)</td>
<td>63%</td>
<td>15 000</td>
<td>71 000</td>
<td>30 000</td>
<td>4.7</td>
<td>63</td>
<td>227</td>
</tr>
<tr>
<td>TP-6a</td>
<td>85–95% a)</td>
<td>28 000</td>
<td>140 000</td>
<td>105 000</td>
<td>5.0</td>
<td>47</td>
<td>225–230</td>
</tr>
<tr>
<td>TP-6b</td>
<td>90–95% a)</td>
<td>35 300</td>
<td>162 000</td>
<td>115 000</td>
<td>4.5</td>
<td>29</td>
<td>225–230</td>
</tr>
<tr>
<td>TP-6c</td>
<td>80–90% a)</td>
<td>43 000</td>
<td>182 000</td>
<td>138 500</td>
<td>4.4</td>
<td>38</td>
<td>225–230</td>
</tr>
<tr>
<td>TP-6d</td>
<td>85–90% a)</td>
<td>30 000</td>
<td>143 000</td>
<td>120 300</td>
<td>4.8</td>
<td>87</td>
<td>225–235</td>
</tr>
</tbody>
</table>

a) Average values of at least three batches; b) $M_n$, $M_w$: number- and mass-average molar masses; $M_{max}$: molar mass at the maximum of the GPC curve; c) Glass transition temperature; d) Beginning of weight loss; e) Reference polymer corresponding to TP-6a; values taken from ref.[9a,9b].

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*Figure 1. Visualization of the improvement process: Solutions of different batches of TP-6a (each 5 wt-% in cyclohexanone/chlorobenzene, 1:1) after turning the samples once upside down. From left to right: three flasks with polymer synthesized according to the reported synthesis show dark color, high viscosity, and inhomogeneous gel-type swollen lumps sticking at the glass walls. The flask on the right finally contains TP-6a prepared according to the optimized synthetic procedure which exhibits excellent film-forming properties.*
the methylene groups attached to N(3) of the triazene moieties show broadened signals. As for most 1-aryl-3,3-dialkyltriazenes, the N(2)–N(3) bond rotation is restricted.[20d] This effect is ascribed to partial \( \pi \)-bond formation between N(2) and N(3).[14d,20a,20b] Therefore, NMR resonances for –N(1)–N(2)–N(3) < alkyl substituents are broadened, and the rotation barrier has been quantitatively evaluated with variable-temperature NMR experiments for various 1-aryl-3,3-dialkyltriazenes derivatives (coalescence temperatures \( T_C \approx 230 \rightarrow 310 \) K).[20c,20d] The hindered rotation of the dialkylamino group fixes the N(3)-alkyl substituents in a position either \( \text{cis} \) or \( \text{trans} \) to the azo group which itself tends to adopt the sterically less hindered \( \text{trans} \) configuration.[14d,20b,20c] The rotation barrier is strongly dependent on electronic effects of the substituent groups. Strong electron-withdrawing groups on the aromatic ring favour the 1,3-dipolar resonance structure and the \( \text{cis} \) and \( \text{trans} \) rotamers are distinct even at room temperature.[20b–d] Increasing line widths of the –N(3)–CH\(_2\)–CH\(_2\)– resonances from \( \Delta \nu_{1/2} \approx 1.7 \) Hz for TP-6a and TP-6b, to 2.2 Hz for TP-6c and 2.6 Hz for TP-6d were observed. This suggests that increasing rotational hindrance also occurs with increasing steric demand and mass of the N(3)-alkyl substituents.[20d,21] The hindered rotation affected the \( ^{13} \text{C} \) NMR spectra much stronger. For example, line widths of \( \Delta \nu_{1/2} \approx 100 \) Hz for –N(3)–CH\(_2\)– and \( \approx 70 \) Hz for –N(3)–CH\(_3\) of TP-6a were determined from the 1D \( ^{13} \text{C} \) NMR spectrum. For other polymers, the alkyl carbon resonances were significantly broader, and \( ^{13} \text{C} \) chemical shifts were obtained from 2D correlated \(^1\text{H}, ^{13} \text{C}\)-HSQC spectra.

Parts of \(^1\text{H} \) NMR spectra of TP-6c at different temperatures are shown in Figure 3. At 297 K, the two nonequivalent i-C\(_3\)H\(_7\) methine protons are close to coalescence with a broad signal at 3.80 ppm (\( \Delta \nu_{1/2} \approx 90 \) Hz). At higher temperatures the signal narrows, whereas distinct signals for the two isomers appear at 3.72 ppm and 5.17 ppm at 280 K. Due to steric requirements, we suggest that the iso-propyl group of the favoured isomer (\( \approx 90\% \)) is in \( \text{trans} \) position relative to the azo group (Figure 3). The large chemical shift difference (\( \Delta \nu \approx 590 \) Hz) for the two methine protons results in a coalescence temperature around 290 K.
The other alkyl resonances broaden with decreasing temperature, but coalescence was not reached. The line widths of the aryl proton signals were not temperature dependent.

Photodegradation Studies

UV-Vis spectra of all polymers are similar and show three strong main absorption bands at \( \approx 200, 290, \) and 330 nm,
and a weaker band at 220 nm appears as a pronounced shoulder (Figure 4). The absorption maximum at 330 nm is attributed to the characteristic aryltriazene band of a $\pi-\pi^*$ transition in the azoamine chromophore.[1] Values for the maximum molar decadic absorption coefficient $\varepsilon$ vary between 31 400 and 33 400 L mol$^{-1}$ cm$^{-1}$ related to the molar mass of the repeating units (Table 2). Excitation of the triazene band leads to cleavage of the photolabile chromophore with release of elementary nitrogen.[1,9c,22]

The absorption measurements show the progressive decomposition of the photolabile triazene chromophore when the polymers were exposed to irradiation: both bands at 290 and 330 nm, as well as the shoulder at 220 nm get weaker, whereas the aromatic absorption bands at 200 nm increase upon irradiation (Figure 4).

### Photodecomposition in Thin Films

Polymer films with thickness between $\approx 15$ and $\approx 100$ nm were investigated. Average roughness of the film surfaces were typically below 10 nm, which corresponds to the roughness of the substrate surfaces. A typical example for the change in the UV absorption curve during photodecomposition is shown in Figure 4(a) for a 70 nm thick film of TP-6d. For the kinetic analysis, the decrease of absorbance $A$ at the maximum wavelength $\lambda_{\text{max}}$ was followed as a function of irradiation time $t$. From the corresponding kinetic equation $A(t) = A_0 e^{-kt}$, rate constants $k$ for each polymer were derived according to $kt = \ln [A_0 - A_{\infty}] - \ln [A(t) - A_{\infty}]$, where $A_0$ is the initial absorbance before photolysis, and $A_{\infty}$, the remaining absorption after complete photodecomposition. Linear fits in such plots indicating first-order kinetics were found for absorbance values below $\approx 0.3$ in the later part of the irradiation experiments, or for films thinner than $\approx 30$ nm [cf. Figure 4(e)]. For thicker films, the initial stages of the photolysis proceeded according to transient pseudo zero-order kinetics due to the high optical density of the films. The specific half-life time $t_{1/2} = \ln 2/k_{\text{rel}}$ of reference polymer TP-6a, derived from the linear part in the plot, was 2.96 min, and the corresponding $t_{1/2}$ values increased steadily to 10.56 min for TP-6d (Table 2). Absolute rate constants are dependent on the experimental setup and only partly meaningful. Therefore, $k_{\text{rel}}$ ratios relative to $k_{\text{rel}}$ of reference polymer TP-6a = 1.0 as the standard reference are summarized in Table 2 as well. The photochemical reactivity within the polymer series is decreasing with the increasing steric demand of side-chain substituents $R$.

### Photodecomposition in Solution

For comparison, photolysis was performed in THF solutions. The experimental setup (lamp, distance of target from light source) of the experiments described above were maintained. Therefore, the kinetic results of the two sets of experiments should remain comparable. Figure 4(b) depicts the photodecomposition of polymer TP-6d with time intervals of 2 min each. With a similar initial absorbance $A_0$, the photolysis is significantly faster in solution compared to films (Table 2). The absorption band of the triazene chromophore flattens first and then disappears completely, indicating a high degree of polymer photodecomposition. The aromatic band at $\approx 200$ nm increases significantly but cannot be measured reliably, since the strong absorption of the solvent THF interferes in this spectral domain.

As for the films, kinetic analysis revealed linear fits for absorbances less than $\approx 0.3$. Literature results[9b] showed that in the related aryltriazene polymers, an increasing degree of substitution of linear side chains $R$ (with $R = \text{Me, Et, Pr}$) resulted in enhanced photodecomposition ($k_{\text{rel}} > 1$).

#### Table 2. UV Spectral Characteristics and Photolysis Kinetic Data.

<table>
<thead>
<tr>
<th>TP</th>
<th>$\lambda_{\text{max}}^{a}$</th>
<th>$\varepsilon(\lambda_{\text{max}}^{b})$</th>
<th>$k_{\text{rel}}^{c,d}$</th>
<th>$t_{1/2}^{e}$ (solution)</th>
<th>$t_{1/2}^{f}$ (thin film)</th>
<th>$t_{1/2}^{g}$ (film)/$t_{1/2}^{h}$ (soln)</th>
<th>$k_{\text{rel}}^{i}$ (thin film)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TP-6a-R$^f$</td>
<td>332</td>
<td>30 600</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>TP-6a</td>
<td>332</td>
<td>32 500</td>
<td>$\Xi$1.0; (1.45)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>TP-6b</td>
<td>335</td>
<td>33 200</td>
<td>1.12; (1.30)</td>
<td>0.79; (3.73)</td>
<td>2.9</td>
<td>0.68</td>
<td></td>
</tr>
<tr>
<td>TP-6c</td>
<td>336</td>
<td>33 400</td>
<td>0.64; (2.29)</td>
<td>0.44; (6.75)</td>
<td>2.9</td>
<td>0.52</td>
<td></td>
</tr>
<tr>
<td>TP-6d</td>
<td>333</td>
<td>31 400</td>
<td>0.40; (3.5)</td>
<td>0.28; (10.56)</td>
<td>3.0</td>
<td>0.42</td>
<td></td>
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</tbody>
</table>

$^a$Wavelength at the absorption maximum in THF; $^b$Related to the molar mass of repeating units; $^c$Relative ratio of absolute values $k_{\text{rel}}$ for identical irradiation conditions; $^d$Irradiation with a mercury lamp; $^e$Monochromatic irradiation with a XeCl$^*$ excimer lamp at 308 nm; $^f$Reference polymer corresponding to TP-6a; values taken from ref.[9a,b].

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This behavior was also observed for the polymers TP-6a and TP-6b (Table 2). According to the given explanation that the increasing steric demand of side chain substituents at N(3) of the triazene chromophore decreases the ability of recombination of the formed radicals, we expected that photodecomposition of TP-6c and TP-6d also exhibited values $k_{rel} > 1$. However, for the branched iso-propyl and tert-butyl substituted polymers, the decomposition rates significantly slowed down. The recombination is influenced by the stability and lifetime of radical intermediates. Increasing inductive effects of substituents R will increase the stability of these radicals, allowing for more recombination before N$_2$ is released. This then explains that rates for photodecomposition of polymers with branched side chains slowed down as observed.

A further hypothesis to explain the decreasing photodecomposition rates from TP-6a to TP-6d is based on photolysis and thermolysis studies of structurally related poly(alkylaryldiazosulfide) and diarylpentadiene compounds. The authors revealed an enhanced photolability of the $\sim$N–N–Z-configuration and suggested a photoinduced $E \leftrightarrow Z$ rearrangement as an intermediate reaction step. If a Z-configuration is also favoring photodissociation for triazenes, we assume that increasing steric demand and mass of the N(3)-alkyl substituents hinder the E/Z isomerization step. This would also be in qualitative agreement with our NMR studies where we observed an increasing rotational hindrance about the N(2)–N(3) bond from TP-6a to TP-6d.

A significant difference between decomposition in solution and in films consists in the remaining basic absorbance of the characteristic triazene twin bands [Figures 4(a) and 4(b)]. The residual absorbance of the triazene bands is minimal in solution [Figure 4(b)], since most of the formed radicals are immediately quenched by disproportionation reactions with the surplus of the surrounding solvent THF. In contrast, photodegraded polymer films show a significant remaining absorption even after long irradiation times [Figure 4(a)]. We attribute this to the formation of conjugated polyarene species. After cleavage and release of N$_2$, the remaining closely packed ary radical species can recombine within the film, forming crosslinked conjugated arene networks that absorb at longer wavelength. We found a reduced solubility of irradiated film areas when the substrates were cleaned by rinsing with THF. Additionally, microscopic analysis confirmed a slightly shady and turbid clouding of irradiated domains, indicating a change in homogeneity and refractive index of the former clear films.

**Excimer-Lamp Irradiation Experiments**

With regard to practical applications, polymer films were also investigated with quasi-monochromatic irradiation of a XeCl$^+$ excimer lamp at 308 nm. A plot of UV curves is shown in Figure 4(c) for a 100 nm thick film of TP-6a. With the narrow-band excitation, all curves intersect properly at four isosbestic points, indicating a uniform mechanism of the photodecomposition process. TP-6b–TP-6d showed similar curves with sharp isosbestic points. Only the main absorption bands of the triazene chromophores are excited at 308 nm, whereas the polychromatic UV light of the mercury arc lamp excites several electronic transitions simultaneously below 400 nm. Therefore, more complex photodecomposition processes are induced, which have an influence on the pathway of photofragmentation reactions. In agreement with the mercury lamp photolysis experiments, decreasing rate constants were found for TP-6a to TP-6d with increasing steric demand of the alkyl substituents attached to the triazene moiety (Table 2).

**Conclusion**

The modified synthesis provides access to triazenes as pure and soluble high-molecular mass compounds in high yields. Photodecomposition of films decreases with the increasing steric demand of the side-chain substituents, and a clean first order decomposition kinetic was found for monochromatic irradiation at 308 nm for all polymer films with an absorbance below 0.3. The promising potential of these materials in LIFT applications was demonstrated with highly sensitive materials recently: assisted by a 100 nm thick sacrificial DRL of polymer TP-6a, living mammalian neuroblast cells were transferred and gently deposited on a receiver substrate. In the same way, multi-spectral nanocrystal quantum dots (NCQD) were successfully transferred by the TP-6a-film-assisted LIFT setup into laterally patterned arrays.

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