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Novel Investigations on Kinetics and Polymerization

Mechanism of Oxazolines Initiated by Iodine

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TITLE RUNNING HEAD. Polymerization of oxazoline initiated by iodine

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ABSTRACT. The cationic ring-opening polymerization of 2-methyl-2-oxazoline (MOx) using iodine initiating system was reported. Uncolored polyoxazolines were produced in spite of the use of iodine initiator, well-known dye molecule. Low molecular weight asymmetric telechelic polyoxazolines carrying an acetylimine head group and an oxazolinium end group were synthesized, with a good control over the molecular weight. NMR and MALDI-Tof experiments allowed the full determination of the chemical structures of the produced poly(2-methyl-2-oxazoline)s, notably with the acetylimine head group, explained by spontaneous α-HI elimination. Finally, to our knowledge, we reported the first detailed mechanism of cationic ring-opening polymerization of 2-methyl-2-oxazoline in acetonitrile according to UV experiments. An ionic-type mechanism was involved with the dissociation of molecular iodine to active initiating species.

KEYWORDS. cationic ring-opening polymerization; iodine; mechanistic study; polyoxazoline
INTRODUCTION

Poly(oxazoline)s\textsuperscript{1,2} are nowadays the subject of great attention due to their properties such as non-toxicity, hydrophilicity, and biocompatibility\textsuperscript{3,4} which made them good candidates for biological and biomedical applications\textsuperscript{5,6} and the use of poly(2-methyloxazoline)s (POx)s was even discussed to substitute poly(ethylene oxide).\textsuperscript{7} To date, the cationic ring-opening polymerization of 2-oxazolines has been accomplished using many initiators including Lewis acids, such as boron trifluoride, and alkyl esters such as tosylates, triflates and halides as summarized in Scheme 1.\textsuperscript{1,8,9} The alkyl halide initiators ranged from chloride,\textsuperscript{10} bromide\textsuperscript{11} to iodide\textsuperscript{12-14} as well as acetyl halide.\textsuperscript{15} The alkyl iodide initiators were mostly converted \textit{in situ} from chloride\textsuperscript{16-18} or bromide\textsuperscript{19} analogues using NaI or KI reactants. The structure of iodide (macro)initiators could also be more sophisticated\textsuperscript{20} with, for example, metalloinitiators using Fe(II) and Rh(II) tris-bipyridines,\textsuperscript{21} polyhedral oligomeric silsesquioxanes (POSS),\textsuperscript{22} gold nanoparticles\textsuperscript{23} or carbon nanotubes.\textsuperscript{24}

\textbf{Insert Scheme 1}

Widespread used tosylate or iodine initiators showed drawbacks that must be taken in consideration. For instance, the partial initiation from \textit{p}-toluene-sulfonic acid (H\textsuperscript{+}), formed by inadvertent hydrolysis of tosylate initiator\textsuperscript{25} led to side products during the polymerization reaction. On the other hand, alkyl iodide initiators proved to be not stable under storage due to the weak C-I bond. They are known to be light-sensitive and temperature-sensitive, and are prone to decomposition to give free iodine (I\textsubscript{2}) or hydroiodic acid (HI) by elimination. Riffle and Saegusa studied the polymerization of 2-oxazoline (Ox), 2-methyl-2-oxazoline (MOx) and 2-ethyl-2-oxazoline (EtOx) in presence of MeI and others iodide initiators like benzyl iodide and 1-iodobutane. Saegusa showed the influence of the nucleophilic reactivity of the monomer on the nature of the mechanism, and proved that MOx > I > Ox.\textsuperscript{12,26,27} High monomer nucleophilic reactivity promoted covalent mechanism (Ox monomer) where the attack of the stronger nucleophile of iodide at the oxazolinium ring led to the covalent-bonded species.\textsuperscript{28,29} In the case of MOx, the mechanism differed and nucleophilic attack of the monomer at the oxazolinium ring
reproduced an oxazolinium ring. Riffle studied the polymerization of EtOx in details using various iodide initiators. Both covalent and ionic active species were present in these systems. The less reactive alkyl halide, 1-iodobutane, yielded much smaller initiation rate constants than benzyl iodide with incubation period of several hours.

In this contribution, we describe the first polymerization of MOx using molecular iodine as initiator, in acetonitrile at 60 °C. The main benefit of the use of I2 was that no preliminary synthesis was required in comparison with triflate, tosylate and halide initiators. We managed to prepare well defined poly(2-methyl-2-oxazoline)s with control over the molecular weight and low polydispersities. Additionally, the chemical structures of polymers were fully established as the chain end-groups were clearly identified using NMR and MALDI-TOF analyses. Finally, UV experiments permitted us to report the mechanism of the polymerization of the 2-methyl-2-oxazoline in acetonitrile for the first time in the literature.
EXPERIMENTAL SECTION

Materials

Molecular iodine, diethylether, methanol, chloroform, potassium hydroxide and calcium hydride (CaH₂) were purchased from ACROS and were used as received. Acetonitrile was dried and distilled according to standard procedures. 2-Methyl-2-oxazoline (MOx) was dried, distilled from CaH₂ and stored under a dry nitrogen atmosphere. Deuterated solvents (CDCl₃ and CD₃CN) were purchased from SDS and were used without further purification.

Analytical techniques

¹H and ¹³C NMR spectra were recorded using a Bruker AC 200 with CDCl₃ as solvent. Chemical shifts (¹H NMR) were referenced to the peak of residual CHCl₃ at 7.26 ppm. Chemical shifts (¹³C NMR) were referenced to CDCl₃ at 77 ppm. Size exclusion chromatography (SEC) was performed on a PL-GPC 50 Plus equipped with an RI refractive index detector. Three PL aquagel-OH columns (25, 7.5 and 4.6 mm ID) were used at 40 °C with a 0.8 mL·min⁻¹ flow rate of H₂O/CH₃OH: 7/3 (+ 0.1M LiNO₃), calibrated using PEO standards. Mass spectrometry analyses were conducted with a Bruker Ultra-Flex MALDI-Tof mass spectrometer, equipped with a nitrogen laser (LSI, 337 nm, 10 ns pulse length) and one detector. Mixture of peptides was used for external calibration. The ions were accelerated by a potential of 25 kV and reflected with a 26.3 kV potential. All measurements were recorded in the reflection mode using α-cyano-4-hydroxycinamic acid (HCCA) as matrix with NaI. For each spectrum 300 transients were accumulated. The resolution at m/z = 1650 was 331. The polymer was dissolved in acetonitrile at a concentration of 10 mg.mL⁻¹. UV-visible analyses were conducted with a Perkin Elmer - lambda 35 UV/Vis spectrometer equipped with PTP-1+1 Peltier System. Experimental conditions: [M]₀=2.35 × 10⁻⁴ M, [I₂]₀=7.88 × 10⁻⁶ M, 60 °C, blank: acetonitrile. To avoid any saturation of the absorbance, the reaction mixture was diluted until 2.10⁻⁴ M instead of 4 M in monomer for a typical polymerization.
Typical procedure for the polymerization of the 2-methyl-2-oxazoline (MOx) using iodine: $PO_{x_{n-1}} + I_2 \overset{H_2O}{\rightarrow} PO_x$. Targeted degree of polymerization: 10.

Reactions were carried out under a dry nitrogen atmosphere. Iodine (0.60 g, 2.35 mmol) and MOx (2 g, 23.5 mmol) were dissolved in dry acetonitrile (5 mL). The solution was vigorously stirred at 60 °C. The product was quenched by addition of an adequate amount of methanolic potassium hydroxide (0.2 mL, 8.91M). The flask was maintained for 4 hours at 30 °C. After cooling, the polymer was isolated by slow precipitation from cold diethylether.

$^1$H NMR (CDCl$_3$) χ (ppm): 7.0-6.7 (m, N=CH), 4.93 (t, N-CH$_2$ oxazolinium), 4.25 (t, O-CH$_2$ oxazolinium), 3.5-3.1 (m, CH$_2$ POx), 2.6 (s, CH$_3$ oxazolinium), 2.3-2.0 (m, CH$_3$ POx).

$^{13}$C NMR (CDCl$_3$) χ (ppm): 176.2 (C=O acetylimine), 171.5-169.8 (C=O amide), 119.7 (N=CH-CH$_2$-N), 117.7 (N=C), 49.3-40.6 (CH$_2$), 21.8-19.7 (CH$_3$), 12.3 (CH$_3$ acetylimine).

Determination of the conversion by $^1$H NMR spectroscopy

The conversion from MOx to corresponding POx was determined by $^1$H NMR spectroscopy (in deuterated acetonitrile) comparing the signals from the released CH$_2$ of polymer at 3.5-3.1 ppm with the signals from the remaining CH$_2$ of monomer at 4.4 and 3.7 ppm.
RESULTS AND DISCUSSION

The present study aims at examining (i) the synthesis of well-defined poly(2-methyl-2-oxazoline)s, and (ii) the mechanism of cationic ring-opening polymerization of MOx in acetonitrile, in the presence of molecular iodine initiator. The latter was already well-known to act as initiator for cationic polymerization.\(^{34}\) For example, it was proved that the living/controlled character of the cationic polymerization of numerous monomers such as isobutyl vinyl ether was inefficient in the presence of I\(_2\) and that it was improved by adding others co-initiating species (HI, SO\(_2\), ZnI\(_2\)).\(^{35-38}\) On the other hand, to date, molecular iodine was never applied for the polymerization of oxazolines.

**Polymerization kinetics.** A kinetic study of the cationic ring-opening polymerization of 2-methyl-2-oxazoline was first achieved. To avoid any confusion between the terms of initiator and iodine, the abbreviations I* and I were attributed to the initiator and the iodine atom, respectively. MOx conversion *versus* time was first performed by \(^1\)H NMR experiment in CD\(_3\)CN. The CH\(_2\) monomer protons resonated at \(\delta\) 4.4 and 3.7 ppm shifts and broadened in the corresponding polymer at 3.5-3.1 ppm. These intensities were used to calculate the monomer concentration as a function of reaction time. The influence of the temperature on kinetic polymerization was studied at 40, 60 and 80 °C. Whatever the temperature, conversions were high and the final reaction time logically decreased with the temperature (Table 1). As expected, the kinetics showed a decrease in polymerization rate with [M]\(0\)/[I*]\(0\) ratio.

**Insert Table 1**

The slope of the ln([M]\(0\)/[M]) *versus* reaction time led to the determination of the rate constant of polymerization (k\(_{\text{app}}\)) (Figure 1). Theoretically, this k\(_{\text{app}}\) can be calculated according to Equation 1:

\[
-k_{\text{app}}[P^*][M] = \frac{d[M]}{dt}
\]

**Equation 1**

where [M] and [P*] are the concentrations of the monomer and propagating species, respectively. Assuming that the concentration of propagating species was equal to the initial initiator concentration
Equation 1 was integrated in Equation 2. In the last equation [I]₀ replaced [I*]₀ because it was not known whether each iodine molecule initiated one polymer chain.

\[
\ln \frac{[M]_0}{[M]} = k_{\text{app}} [I]_0 \cdot t \quad \text{Equation 2}
\]

\[
\ln([M]_0/[M]) \text{ versus time was found to be linear as evidenced in Figure 1. This linear behavior showed that the concentration of active chains remained constant throughout the polymerization indicating that termination reactions were not significant. We noted that the } k_{\text{app}} \text{ values depended on } [I]_0 \text{ and decreased with } [M]_0/[I]_0. \text{ The trend revealed that } [I^*] \text{ did not linearly scale with } [I]_0 \text{ (Table 2). The difference between } [I^*] \text{ and } [I] \text{ increased with } [M]_0/[I]_0. \text{ Additionally, for the same } [M]/[I] \text{ ratio, the rate constant propagation for the cationic polymerization of } \text{MOx using iodine initiator (0.82 \times 10^{-3} L.mol}^{-1}.s^{-1}) \text{ was consistent with the values obtained for MeOTs and MeI initiators, in dimethylacetamide (2.40 \times 10^{-3} \text{ and } 2.22 \times 10^{-3} \text{ L.mol}^{-1}.s^{-1}, \text{ respectively}).}^{39} \text{ From the graphical resolution of the Arrhenius equation (equation 3) the frequency factor and the activation energy were deduced for the } \text{MOx/I}_2/\text{CH}_3\text{CN system. The calculation of } K_p \text{ from } \ln([M]_0/[M])=f(t) \text{ at } 40, 60 \text{ and } 80 \degree C \text{ was achieved. The values fitted with the literature data [system; } E_a (\text{KJ.mol}^{-1}); A \text{ (L.mol}^{-1}.s^{-1})]. \text{ Our initiator system seemed more reactive than MeI initiator, with the following values: } [\text{MOx/I}_2/\text{CH}_3\text{CN}; 60.0; 3.5x10^6] \text{ and } [\text{MOx/MeI/CH}_3\text{CN}; 72.9; 1.7x10^8].^{12,40} \]

Insert Figure 1

Insert Table 2

\[
k_p = Ae^{-E_a/RT} \quad \text{Equation 3}
\]

All obtained polymers were characterized by size exclusion chromatography to determine the molecular weights and the polydispersity indexes (PDI). The molecular weight versus global conversion was reported in Figure 2 for two different theoretical molecular weights (2550 and 4250 g.mol^{-1}). The } M_n \text{ values increased linearly until 80 } \% \text{ conversion showing the control of the polymerization reaction. For higher conversion, the experimental values deviated from the targeted ones probably due to the presence of transfer reactions. The study of the mechanism revealed the HI was released at the}
beginning of the polymerization (according MALDI-Tof and NMR part). HI could reinitiate new chains, induced transfer reactions and might explain the limited DPs even if the HI initiated chains were not detected by MALDI-TOF spectroscopy.

**Insert Figure 2**

The influence of the temperature on the molecular weight was studied at 40, 60 and 80 °C (Table 3 runs 1-3). The lowest value of polydispersity index was obtained at 60 °C. As a consequence, a series of POx polymers were synthesized at this temperature, with a \([M]_0/[I^*]_0\) ratio ranging from 10 to 100 (Table 3 runs 3-7). In all cases, the molecular weight distribution of the obtained poly(2-methyl-2-oxazoline)s was lower than 1.5. Up to an \([M]/[I^*]\) ratio of 40, the molecular weights versus DP\(_n\) increased linearly as depicted in Figure 3. As already mentioned in the literature, the deviation could be attributed to polymer adsorption onto some component of the SEC apparatus.\(^{41}\) For higher M\(_n\) values, the decrease of molecular weight and the enhancement of molecular weight distribution meant the presence of transfer reactions (Table 3 run 7). We concluded that the synthesis of high molecular weight polyoxazolines using iodine was limited, as already described for others initiators.\(^{42}\)

**Insert Table 3**

**Insert Figure 3**

**Chain-end functionalities. Determination by MALDI-Tof and NMR measurements.** It is important to notice that polymerization reactions were stopped at about 80 % conversion for the determination of chain-end functionalities in order to minimize transfer. Moreover, for the sake of clarification of this work, the following abbreviation was attributed to each possible chain end-group, taking into account the head and end groups of the poly(2-methyl-2-oxazoline) considered chain:

\[ \text{POx}^{\text{head group}}_{\text{end group}} \]

In many cases \(^1\text{H}\) NMR analysis was suitable for the determination of end-groups and also permitted to identify the type of mechanism. The cationic ring-opening polymerization of oxazoline proved to
proceed via two different types of species, *i.e.* ionic and covalent types, depending on the nature of the initiator and on the monomer. Ionic and covalent polymerization mechanisms were explained by the balance of nucleophilicities between monomers and $X^-$ counteranions. In both mechanisms the first step dealt with the reaction between the RX initiator and the $\text{MOx}$ monomer (Step A of Scheme 2). In the case of the covalent mechanism, a further step occurred with the reaction between the initiation product and the $X^-$ counteranion (Step B of Scheme 2).

**Insert Scheme 2**

The cationic propagating specie of an oxazolinium salt was not fragile and could be isolated. Thus the ionic-type mechanism was detected by NMR technique showing the oxazolinium group. The presence of typical peaks of oxazolinium species were detected at 4.9 and 4.4 ppm corresponding to CH$_2$-O and CH$_2$-N, respectively. Similar results were obtained in our case, clearly demonstrating that an ionic-type mechanism was involved for the CROP of $\text{MOx}$ using iodine (Figure 4). An additional signal was displayed around 7.0 ppm and was attributed to the CH=N proton of an acetylimine group obtained by the elimination of hydroiodic acid (HI) at the end-group of the polymer chain. The HI elimination was explained by the weakness of the N-I bond, and led to the formation of more stable conjugated specie as shown in Scheme 3. The peak corresponding to the CH= appeared from the beginning of the polymerization and its integration kept constant with regard to oxazolinium group during the polymerization process. We concluded that the spontaneous $\alpha$-HI elimination occurred at the very beginning of the polymerization and revealed the existence of the $\text{PO}_{x-n-2}^{\text{POx}} \overset{\text{HI}}{\rightarrow}_{x}$, corresponding to a polyoxazoline main chain with an acetylimine head group and an oxazolinium end group. By the integration of respective $^1\text{H}$ NMR signals the CH=N/oxazolinium ratio was 89 %, This ratio was underestimated since a complete broken of N-I was supposed. Thus the covalent route represented lower than 11 % and the ionic species were the most important ones. $^13\text{C}$ NMR spectroscopy confirmed this $\text{PO}_{x-n-2}^{\text{POx}} \overset{\text{HI}}{\rightarrow}_{x}$ chemical structure with signals at 171.5-169.8 ppm for the carbonyl group of the POx main chain whereas the downfield peak at 176.2 ppm was assigned to the C=O acetylimine group. Peaks at 119.7 and 117.7 ppm were attributed to the units adjacent to imine group and CH=N, respectively.
Finally, peaks at 21.8-19.7 and 12.3 ppm were assigned to POx main chain and the methyl group of acetylimine, respectively.

**Insert Figure 4**

**Insert Scheme 3**

MALDI Tof mass spectroscopy was used to confirm the nature of the POx end-groups and to study the polymerization mechanism. The theoretical values were compared to the experimental ones (M, RM and ΔM) as summarized in Table 4. The theoretical mass was expressed following the Equation 4:

$$M_{th} = \Delta M_{th} \times n + RM_{th}$$  \hspace{1cm} \text{Equation 4}

where $$M_{th}$$ is the calculated mass of the polymer of degree of polymerization nearest to the measured value, $$\Delta M_{th}$$ is the mass of the monomer unit, and $$RM_{th}$$ representing the calculated residual mass. The determined Mp values (mass at peak maximum of the most intensive signal) and the number-average molecular weight (Mn) are also given in Table 4.

**Insert Table 4**

The polyoxazoline chemical structures were analyzed before and after the termination stage by potassium hydroxide (Scheme 4). Spectra were recorded in reflectron mode using Na+ cations in an α-cyano-4-hydroxycinamic acid matrix. The detected ions were MNa+ type. Two distributions of peaks from m/z in the range of 356 (n = 3; POx_n−1^{−\text{HI}}) to 1419 Da (n = 13, POx_{n−1}^{1} \text{OH}) were observed with an interval between two principal peaks of the same family being 85.10, which corresponded to the molecular weight of the 2-methyl-2-oxazoline monomeric unit. Before the termination stage POx_{n−2}^{−\text{HI}} was detected with HI α-elimination and oxazolinium head group according to $^1$H NMR experiment (Figure 4). The MALDI Tof mass study confirmed the HI elimination during the polymerization stage with a good correlation between theoretical and experimental Mn values. We noted the agreement between RM_{exp} and RM_{th} values as well as the close value of M_{p,exp} and M_{p,th}. In the MALDI Tof mass spectrum of POx terminated by the potassium hydroxide, a monomodal distribution was attributed to POx_{n−1}^{−\text{HI}} where a scission of HI labile group occurred. No other family was mentioned thus no side
reaction was detected. In order to avoid a complete HI scission and to show the N-I bond an experiment was achieved in the darkness. A multimodal distribution displayed two different populations of the same intensity as compiled in Figure 5. POx$_n$$_{\text{OH}}^1$ corresponded to the substitution of oxazolinium head chain by the hydroxyl group.

**Insert Scheme 4**

**Insert Figure 5**

Finally, the originality in this work was also due to the obtaining of uncolored poly(2-methyl-2-oxazoline)s despite the use of iodine as initiator. This was achieved by both the addition of the potassium hydroxide and the elimination of hydroiodic acid during the polymerization reaction. Iodine was easily eliminated to afford well-defined products with an acetylimine head group and a hydroxyl end group.

**Investigations on the polymerization mechanism by UV experiments.** During the polymerization process the evolution of iodide species was monitored by UV spectroscopy as function of time (Figure 6). The electronic spectrum of iodine was firstly recorded in cyclohexane and acetonitrile to evaluate the influence of the dielectric constant D on iodine. Both solvents have a dielectric constant of 2.0 and 37.5 and a dipole moment of 0 and 3.84 Debye, respectively. In non-polar solvent, the absorption band of free iodine was observed at 480-600 nm. In the very polar and polarizable acetonitrile, the absorption of iodine disappeared and additional transition appeared before the addition of MOx. This indicated that acetonitrile interacted with iodine according to partial bimolecular initiation and that other iodide species were formed such as triiodide anion. The presence of the latter was clearly shown by UV experiments with the absorption bands at 290 and 366 nm whereas the pentaiodide anion did not appear at 442 nm. The addition of MOx induced a total shift of the equilibrium towards I$_3^-$ and I$^+$ and gave oxazolinium species with triiodide counteranion: MOx$^{\text{Ium}}$...I$_3^-$. This trend was suggested by the increase of the typical bands of I$_3^-$ and the disappearance of iodine band as soon as the addition of MOx. The I$^+$ species acted as active initiator species and reacted with the nitrogen atom of monomeric unit.
After 5 minutes of reaction the decrease of I$_3^-$ species bands could be explained by the conversion of triiodide to I$^-$. The conversion released molecular iodine and produced I$_3^-$ and I$^+$ ions which reacted with MOx again. This step should explain the relatively large polydispersity index around 1.2-1.4 caused by a long time of initiation stage. The UV experiments concluded on the partial ionization of I$_2$ to I$^+$ and I$_3^-$ in a polar solvent and the complete conversion with the addition of MOx (Scheme 5). Then the active species I$^+$ reacted with MOx and I$_3^-$ was converted in I$^-$ counteranion. The released iodine was ionized in I$^+$ and I$_3^-$ until complete depletion of iodine. The next stage was the propagation with the formation of POx$_{n-2}$Ox$^{-}$H$I$. 

Insert Figure 6

Insert Scheme 5
CONCLUSION

Low molecular weight poly(2-methyl-2-oxazoline)s were synthesized with good control over the chain length and low polydispersity using molecular iodine as initiating system. The main kinetic features were described. The polyoxazoline end-groups were also identified using NMR and MALDI Tof MS measurements which notably showed the formation of N-I bond as head group of the polymer chain. The spontaneous α-HI elimination due to the unstable N-I bond was also demonstrated and uncolored products were isolated. The mechanism of the cationic ring-opening polymerization of MOx was finally discussed. All experimental results allowed us to propose an ionic-type mechanism for the ring opening polymerization of 2-methyl-2-oxazoline in acetonitrile.

ACKNOWLEDGMENT. The authors would like to thank the French Ministry of Education and Research for grant (BG)
FIGURE CAPTIONS

Figure 1. Ln([M]₀/[M]) versus time determined by $^1$H NMR experiment ($k_{app}$ values were expressed in L.mol$^{-1}$.s$^{-1}$).

Figure 2. Molecular weight versus conversion determined by size exclusion chromatography (left: $M_{n,th}$ = 2550 g.mol$^{-1}$; right: $M_{n,th}$ = 4250 g.mol$^{-1}$).

Figure 3. Mn and polydispersity indexes obtained by means of SEC for MOx initiated by iodine.

Figure 4. $^1$H NMR spectrum of the poly(2-methyl-2-oxazoline)s showing both the covalent mechanism of the polymerization and the elimination of HI.

Figure 5. MALDI Tof mass spectrum of POx after the termination reaction by the potassium hydroxide.

Figure 6. UV-Vis spectra as a function of time for the polymerization of MOx in acetonitrile using iodine.

SCHEME TITLES

Scheme 1. Cationic ring-opening polymerization of MOx using various initiators.

Scheme 2. Covalent and ionic mechanism types for the cationic ring-opening polymerization of MOx.

Scheme 3. Elimination of hydroiodic acid during the polymerization reaction.

Scheme 4. Poly(2-methyl-2-oxazoline) after the termination reaction by the potassium hydroxide.

Scheme 5. Proposed mechanism of MOx polymerization using iodine initiator.
Tables

Table 1. Experimental conditions and conversion data for the polymerization of 2-methyl-2-oxazoline.

Table 2. Experimental values of $k_{\text{app}}$ for various $[M]_0/[I]_0$ ratios.

Table 3. Experimental and theoretical values of $\text{DP}_n$ and $M_n$.

Table 4. Analytical data derived from MALDI Tof MS spectroscopy.
Scheme 1. Cationic ring-opening polymerization of MOx using various initiators.
**Table 1.** Experimental conditions and conversion data for the polymerization of 2-methyl-2-oxazoline.

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Figure 1. Ln([M]₀/[M]) versus time determined by $^1$H NMR experiment ($k_{app}$ values were expressed in L.mol$^{-1}$.s$^{-1}$).
Table 2. Experimental values of $k_{app}$ for various $[M]_0/[I]_0$ ratios.

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<th>$[M]_0/[I]_0$</th>
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Figure 2. Molecular weight versus conversion determined by size exclusion chromatography (left: $M_{n,\text{th}} = 2550 \text{ g.mol}^{-1}$, right: $M_{n,\text{th}} = 4250 \text{ g.mol}^{-1}$).
### Table 3. Experimental and theoretical values of DP$_n$ and M$_n$.

<table>
<thead>
<tr>
<th>Run</th>
<th>Temp. (°C)</th>
<th>Time (min)</th>
<th>DP$_{n,th}^a$</th>
<th>conv (%)</th>
<th>M$_{n,th}$ (g/mol)</th>
<th>DP$_{n,SEC}^b$</th>
<th>M$_{n,SEC}^b$ (g/mol)</th>
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<td>1400</td>
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<td>20</td>
<td>1700</td>
<td>1.6</td>
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</table>

$^a$: $DP_{n,th} = \frac{[M]}{\langle M \rangle_{conv}}$

$^b$: from SEC data in MeOH/H$_2$O: 3/7 relative to PEO standards.
Figure 3. $M_n$ and polydispersity indexes obtained by means of SEC for MOx initiated by iodine.
Scheme 2. Covalent and ionic mechanism types for the cationic ring-opening polymerization of MOx.
Figure 4. $^1$H NMR spectrum of the poly(2-methyl-2-oxazoline)s showing both the ionic mechanism of the polymerization and the elimination of HI.
Scheme 3. Elimination of hydroiodic acid during the polymerization reaction.
Table 4. Analytical data derived from MALDI ToF MS spectroscopy.

<table>
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<th>assignment</th>
<th>RM&lt;sub&gt;formula&lt;/sub&gt;</th>
<th>ΔM&lt;sub&gt;exp&lt;/sub&gt;&lt;sup&gt;a&lt;/sup&gt;</th>
<th>RM&lt;sub&gt;th&lt;/sub&gt;&lt;sup&gt;a&lt;/sup&gt;</th>
<th>RM&lt;sub&gt;exp&lt;/sub&gt;&lt;sup&gt;a&lt;/sup&gt;</th>
<th>M&lt;sub&gt;p,th&lt;/sub&gt;&lt;sup&gt;a&lt;/sup&gt;</th>
<th>M&lt;sub&gt;p,exp&lt;/sub&gt;&lt;sup&gt;a&lt;/sup&gt;</th>
<th>calculation</th>
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<td>PO&lt;sub&gt;x&lt;/sub&gt;_n−2&lt;sup&gt;−H&lt;sub&gt;i&lt;/sub&gt;&lt;/sup&gt;</td>
<td>C&lt;sub&gt;8&lt;/sub&gt;H&lt;sub&gt;13&lt;/sub&gt;N&lt;sub&gt;2&lt;/sub&gt;O&lt;sub&gt;2&lt;/sub&gt;I</td>
<td>85.05</td>
<td>296.10</td>
<td>296.10</td>
<td>976.90</td>
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<td>85.10(n-2) + RM</td>
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<td>144.02</td>
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<td>824.50</td>
<td>85.10n + RM</td>
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<tr>
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<td>C&lt;sub&gt;4&lt;/sub&gt;H&lt;sub&gt;7&lt;/sub&gt;NO&lt;sub&gt;2&lt;/sub&gt;</td>
<td>85.05</td>
<td>101.10</td>
<td>101.00</td>
<td>781.90</td>
<td>781.40</td>
<td>85.10(n-1) + RM</td>
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</table>

ΔM = mass of the monomer unit. ΔM<sub>th</sub> = 85.10 g.mol<sup>−1</sup>. RM = residual mass.

<sup>a</sup>: value expressed in g.mol<sup>−1</sup>. 
Scheme 4. Poly(2-methyl-2-oxazoline) after the termination reaction by the potassium hydroxide.
Figure 5. MALDI Tof mass spectrum of POx after the termination reaction by the potassium hydroxide.
Figure 6. UV-Vis spectra as a function of time for the polymerization of MOx in acetonitrile using iodine.
Scheme 5. Proposed mechanism of MOx polymerization using iodine initiator.
REFERENCES


(34) Matyjaszewski, K. Cationic polymerizations: mechanisms, synthesis, and applications; Marcel Dekker, Inc.: New York, 1996.


Cationic ring-opening polymerization of 2-methyl-2-oxazoline was performed in the presence of iodine. The synthesis of uncolored products was explained by MALDI-Tof experiment which revealed the spontaneous α-H-I elimination and the absence of iodated by-products. An ionic-type mechanism of polymerization was proposed according to UV experiments.