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Photolatent Ring-Opening Metathesis Polymerization in Miniemulsion: a Powerful Approach to Produce Polynorbornene Latexes

Loïc Pichavant, Patrick Lacroix-Desmazes, Abraham Chemtob, Julien Pinaud, and Valérie Héroguez

The use of miniemulsion Ring-Opening Metathesis Polymerization (ROMP) to form polyunsaturated latexes is still a largely unexplored field. The main obstacle remains the preparation of a chemically-stable monomer/catalyst miniemulsion, which is generally jeopardized by the high reactivity of ROMP catalysts at ambient temperature. To overcome this drawback, a photo-latent ROMP catalytic system has been employed for the first time. Our approach starts with a N-heterocyclic carbene (NHC) photogenerating system (1,3-bis(mesityl)imidazolium tetraphenylborate / 2-isopropylthioxanthone) which enables the in situ formation of an active NHC-derived Ru catalyst. The ability to initiate the ROMP in miniemulsion by a photoactive way depends on the absorption conditions of the initial formulation due to irradiation attenuation by scattering. In this contribution, the optimization of an aqueous norbornene miniemulsion in order to improve the irradiation penetration depth is presented in a first part. In a second part, the miniemulsion ROMP is investigated by using the NHC photogenerating system and two ruthenium complex pre-catalysts [RuX(ν-cymene)] dimer (X = Cl or I). Stable PNb latexes with particle size in the range of 100 nm were obtained.

Introduction

Aqueous polymer latexes have high industrial interests, firstly, because water is a cheap, safe and environmentally friendly solvent. Secondly, because the aqueous polymer dispersions feature specific properties such as low viscosity and film-forming ability (provided proper adjustment of the Tg of the polymer), which make them relevant for many applications including coatings, adhesives, paper coating, inks, additives for construction materials, and non-woven textiles. Although there is no commercial application known today, miniemulsion free radical polymerization is a powerful way to obtain polymer latexes thanks to the tolerance of free radicals towards water. In addition, miniemulsion polymerization relies on droplet nucleation allowing the polymerization of very hydrophobic monomers as well as the preparation of organic/inorganic nanocomposites, which are difficult to achieve with conventional ab initio emulsion polymerization. In a typical miniemulsion route, a mixture containing the monomer(s), a water-insoluble initiator, and a costabilizer (to avoid Ostwald ripening) forms an organic phase. This latter is then mixed with the aqueous phase containing a surfactant before being emulsified by high energy input usually through an ultrasonifier. Miniemulsion droplets typically display average diameters ranging from 40 to 400 nm, and are kinetically stable. Polymerization is generally triggered upon heating when using thermal radical initiators. Contrary to radical polymerization, only few contributions have been reported regarding the formation of latexes by Ring-Opening Metathesis Polymerization (ROMP) in aqueous dispersed media. The challenges of such route are firstly the water sensitivity of most ROMP catalysts, and secondly, their non-latent character. Therefore, the active catalyst jeopardizes the preparation of a chemically-stable monomer miniemulsion, causing premature polymerization.

ROMP in miniemulsion was introduced by Claverie et al. in 2001. To circumvent the issue of premature polymerization, they first prepared a miniemulsion containing a hydrophobic catalyst (1st generation Grubbs’ catalyst dissolved in a small amount of toluene) and added dropwise the cycloolefin monomer. Nevertheless, only the least reactive monomers (cyclooctene and cyclooctadiene) could be used without miniemulsion destabilization. In previous contributions, we reported the miniemulsion ROMP of norbornene (Nb) by the addition of well-defined poly(ethylene oxide)-based ruthenium carbene as water-soluble macroinitiators to monomer miniemulsion. Stable polynorbornene (PNb) latex particles with average diameter of 200-500 nm were obtained with almost quantitative monomer conversion. Following a similar strategy, a tandem ROMP and Atom Transfer Radical Polymerization (ATRP) in miniemulsion was also investigated. However, the issue was the cumbersome and tricky synthesis of the macroinitiator. At the same time, miniemulsion ROMP of an amphiphilic PEO-based cyclobutene macromonomer was reported by Le et al. using Grubbs’ third generation catalyst. The initial miniemulsion was stable and no coagulation appeared, but the system led to partial macromonomer conversion after 48 h of reaction at room temperature.

In this context, we envision that a major improvement for miniemulsion ROMP could be made by incorporating a hydrophobic latent catalytic system into the organic droplet...
phase. Several pathways are described in the literature to generate latent ROMP catalytic systems using chemical, thermal or mecanochemical means (ultrasound), but the convenient method would be the use of an irradiation stimulus. In most examples in solution, photolatent systems consist in ruthenium complexes made inactive by chelating ligands or electron-rich carbene moieties. Conversion into an active form proceeds by photoinduced ligand dissociation/rearrangement or isomisation reactions. In contrast, we recently proposed a completely different approach based on two consecutive steps: the photogeneration of N-heterocyclic carbene (NHC) ligand, and its subsequent reaction with an inactive Ru precatalyst to form in situ an active NHC-derived Ru ROMP catalyst. The combination of 1,3-dimesitylimidazolium tetraphenylborate salt (IMesH^+BPh^-) (NHC protected form) with 2-isopropylthioxanthone (ITX) (sensitizer) proved to be able to generate the 1,3-dimesitylimidazol-2-ylidene (IMes) NHC upon UV irradiation. In regard to the second step, the active Ru precatalyst, formed in situ the active RuCl_2(p-cymene)IMes) Noels' catalyst in presence of IMes to initiate the ROMP of Nb.

We show herein that such tricomponent catalytic system [RuCl_2(p-cymene)] dimer/IMesH^+BPh^-/ITX can also serve for photoROMP in miniemulsion. To the best of our knowledge, no photolatent system for ROMP of cycloolefin in aqueous dispersed media has been reported yet. In a first part of this study, particular attention has been paid to miniemulsion formulation in order to optimize incident irradiation, and increase the uniformity of irradiation distribution within the reactor. Indeed, a photochemical reaction is only possible in irradiated volumes. In a heterogeneous or dispersed medium, the incident photon flux is attenuated because of light absorption and scattering. To mitigate the detrimental effect of scattering on photopolymerization in dispersed system, Nb miniemulsions with an average droplet smaller than 100 nm have been prepared and their optical properties (extinction) have been evaluated by UV/Vis Spectroscopy, based on measurements in regular transmission. Finally, photo-induced ROMP of Nb has been triggered through the water-insoluble tricomponent catalytic system dissolved in the monomer droplets, and the results have been compared with photo-ROP in solution.

**Experimental section**

**Materials**

Ethanol (EtOH; 96%), chloroform (CHCl_3; 99.4%), dichloromethane (DCM; 99.8%) from VWR Chemicals and 1,2-dichloroethane (DCE; 99.8%) from Aldrich were used without further purifications. 1,3-bis(mesityl)imidazolium chloride (IMesH^+Cl^-; 97%) and sodium tetraphenylborate (NaBPh_4; 99.5%) were purchased from ABCR. Dichloro(p-cymene) ruthenium(II) dimer ([RuCl_2(pCy)_2]); 98%) and diiodo(p-cymene) ruthenium(II) dimer ([Ru_2(pCy)_2]) were purchased from Alfa Aesar. 2-isopropylthioxanthone (ITX; 98%) was obtained from TCI Europe. CREMOPHOR CO40 and CREMOPHOR RH60 (steric surfactants) were obtained from BASF. Merp®HCS, Tween®20, Brij®S100 (steric surfactants), norbornene (Nb) and hexadecane were purchased from Aldrich. The structure of the steric surfactants is described in the results and discussion part. Deionized water was obtained with a Millipore Milli-Q Biocel A10 purification unit.

**Irradiation source**

UV irradiation was performed with a LIGHTNINGCURE® Series apparatus from HAMAMATSU equipped with a light guide, a medium pressure mercury-xenon lamp and a filter at 365 nm. Samples were placed at 10 cm of the light guide. Irradiance was measured with a Power Puck II radiometer from EIT® (20 mW/cm^2 at 365 nm).

**Characterization methods**

^1^H NMR studies were acquired using a Bruker spectrometer 400 MHz employing CDCl_3 or DMSO-d_6 as solvent at 25°C. Conversions of Nb were determined by gas chromatography (GC) with hexadecane as internal standard, using a VARIAN GC3900 (apolar capillary column BP1-30m; injector temperature: 250°C; detector temperature: 300°C; ramp temperature: 2 min at 50°C + 10°C/min until 250°C; GC retention times: t_R^+Nb = 1.77 min; t_R^+dodecane = 13.25 min). UV/Vis measurements were performed with a UV/Vis spectrometer Agilent Carry 4000 (optical path length: 1 mm). Dynamic light scattering (DLS) measurements were performed using a MALVERN Zetasizer Nano ZS equipped with He-Ne laser (4 mW; 633 nm). All spectra are the average of three measurements. Before measurements, latexes were diluted about 250 times to minimize multiple scatterings caused by high concentration. The scattering angle used was 90°. Size-exclusion chromatography (SEC) was performed in tetrahydrofuran (THF) (1 mL min^-1) with trichlorobenzene as the flow marker, using both refractometric and UV detectors. TEM pictures were performed with a HITACHI H7650 microscope operating at an accelerating voltage of 80 kV. Latexes were diluted 70 times and 10 mL of the final latexes were poured on carbon-carbon grids and dried in normal atmosphere. The pictures were recorded with Digital Micrograph (GATAN) and analyzed with the software ImageJ to determine the particle average diameter and the polydispersity (PDI). PDI is calculated with the following formula (Equation 1):

\[
PDI = \left( \frac{\sigma}{D} \right)^2
\]

With:

- D: the average diameter
- \(\sigma\): the standard deviation

**Synthesis of 1,3-dimesitylimidazolium tetraphenylborate (IMesH^+BPh^-)**

In a typical experiment, 2.28 g of IMesH^+Cl^- (341 g/mol; 6.7 \(10^{-3}\) mol) are dissolved in 67.5 mL of degassed ethanol (Freeze-Pump-Thaw method). A solution of NaBPh_4 (3.59 g; 342 g/mol; 10.5 \(10^{-3}\) mol) in degassed ethanol (67.5 mL) is added dropwise under a nitrogen flux. The formed precipitate (IMesH^+BPh^-) is filtrated, washed with water and ethanol and dried overnight.
under vacuum. Mass Yield: 85% - $^1$H NMR data in DMSO-$d_6$: $\delta$ (ppm) (400 MHz): 2.12 (s; 12H); 2.36 (s; 6H); 6.79 (t; 4H); 6.94 (m; 8H); 7.21 (m; 12H); 8.26 (s; 2H); 9.64 (s; 1H). (Supplementary information, Figure S1)

**Miniemulsion formation**

Aqueous phase was prepared by adding a variable amount (0.01-1 g) of surfactant (CREMOPHOR CO40, CREMOPHOR RH60, Merpol® HCS, Tween®20 or Brij®S100) to 10 mL of milliQ water. The organic phase was prepared by mixing Nb (25 w-%) with hexadecane (10 w-%) in the solvent (DCM or DCE) (65 w-%). Then, the organic phase (0.5, 1 or 2 g for miniemulsions at 5, 10 or 20 w-% of dispersed phase versus the continuous phase respectively) was added to the aqueous phase and the pre-miniemulsion was stirred during 1 hour. Then a miniemulsion was formed by ultrasonication with a BioBlock Vibra-Cell 75043 (750 W) equipped with an ultrasonic tip (from 2 min to 20 min; pulse mode, 5.0 s on 5.0 s off). During ultrasonication, samples were placed in an ice bath to avoid any temperature increase.

**Polymerization in solution**

A solution of Nb (25 w-%; 823 mg; 8.75 $10^{-3}$ mol; 510 eq.) and hexadecane (10 w-%; 329 mg; 427 µL) in dichloromethane (2.47 mL; 2 mL) was prepared and degassed (Freeze-Pump-Thaw method). Then, [RuCl$_2$(p-Cy)$_2$] (1 eq.; 1.71 $10^{-5}$ mol; 10 mg) or [Ru$_2$(p-Cy)$_2$] (1 eq.; 1.71 $10^{-5}$ mol; 17 mg) and IMes$^+$$\text{BPh}_4^-$ (5 eq.; 8.6 $10^{-5}$ mol; 54 mg)/ITX (2.5 eq.; 4.3 $10^{-5}$ mol; 11 mg) were added under inert atmosphere (glovebox). This solution was then transferred inside a quartz cuvette (optical path length: 1 mm) and irradiated during different times. After irradiation, the samples were dissolved in a minimum of dichloromethane and analyzed by gas chromatography (Nb conversion). Finally, the samples were precipitated in acetone, filtrated, dried under vacuum and the obtained polymers were analyzed by $^1$H NMR and SEC (data are given in the Results and discussion part, Figure S3 and Figure S5).

**Polymerization in miniemulsion**

10 mL of a solution of Brij®S100 in milliQ water (100 g/L) were degassed by bubbling nitrogen during 1 hour. Then, 0.5, 1 or 2 g (for miniemulsions at 5, 10 or 20 w-% respectively) of degassed organic phase containing the catalytic system were added under nitrogen (preparation of the organic solution are described in the previous section “polymerization in solution »). The pre-miniemulsion was stirred during 1 hour. Then, the miniemulsion was formed by ultrasonication during 2 min (pulse mode, 5.0 s on 5.0 s off). The miniemulsion was then transferred inside a quartz cuvette (optical path length: 1 mm) and irradiated during different times. After irradiation, the samples were dissolved in a minimum of THF and analyzed by gas chromatography (Nb conversion). Finally, the samples were precipitated in acetone, filtrated, dried under vacuum and the obtained polymers were analyzed by $^1$H NMR and SEC (data are given in the Results and discussion part Figure S4 and Figure S6).

**Results and discussion**

In a first stage of this study, attention has been paid to optimize the formulation of the miniemulsion to obtain droplets sizes lower than 100 nm (to minimize radiation scattering). For the sake of simplicity, this part of the study was performed without ROMP initiator.

**Formulation of the Nb miniemulsion**

The aqueous miniemulsion is composed of a continuous phase – water and surfactant – and a dispersed phase – organic solvent, monomer (Nb) and hexadecane used as costabilizer to prevent Ostwald ripening. The miniemulsion is formed by first stirring the two-phases (pre-emulsion) and second by ultrasonication. Several parameters have been screened in order to optimize the formulation.

**Influence of the type of surfactant:** In this part, the dispersed phase concentration was 10 w/w-% and was composed of Nb (80 w-%) and hexadecane (20 w-%). The type of surfactant must be chosen with care. Indeed, we have shown that convensional ionic surfactants such as SDS can interact with IMes$^+$$\text{BPh}_4^-$ by counterion exchange having a negative effect on the initiator formation (Figure S2, supplementary information). This kind of reaction is well known in ionic liquid synthesis. Consequently, nonionic surfactants were employed. A series of five polymeric surfactants has been selected as a function of their hydrophilic/lipophilic balance (HLB); a high HLB being required to stabilize an oil-in-water emulsion. The miniemulsion was formed by ultrasonication during 10 minutes at 50 % of the maximal power. For each surfactant, the droplet size determined by DLS has been plotted as a function of the surfactant concentration. From these curves can be extracted two fundamental parameters: the critical micelle concentration (CMC), from this concentration the droplet stabilization is optimal and the addition of more stabilizer would induce the undesirable formation of micelles in the continue phase in the presence of the organic dispersed...
The results for the five tested surfactants are summarized in the Table 1. The smallest droplets were obtained with the Brij®S100 and the CREMOPHOR CO40 with a concentration of 100 g/L (D_{min} around 120 nm). As the Brij®S100 is a more common and cheaper surfactant, it has been chosen for the rest of the study.

**Influence of the type and amount of solvent in the dispersed phase:** For these experiments, Brij®S100 was used as surfactant and the organic phase was composed of the solvent (65 w-%; minimal concentration for which the catalytic systems is still soluble), Nb (25 w-%) and hexadecane (10 w-%). For the preparation of the organic phase, the two main parameters were studied: the nature of the solvent (DCM or DCE), and its weight content (5, 10 or 20 w/w-%). The results are presented in Table 2.

Regarding the nature of the solvent (entries (1) and (5) and entries (3) and (6)) the use of DCE leads to smaller droplets. The organic phase content has also an influence on droplet size (entries (5), (7) and (8)). As expected, the less the organic phase ratio is, the smaller the droplets are. With a dispersed phase at 5 w/w-%, very small droplets are obtained (36 nm; entry (7)).

**Ultrasonication parameters optimization:** The last studied parameters for the optimization of the miniemulsion were the ultrasonication conditions such as the ultrasonication time and the power of the ultrasonication probe (Table 2). Regarding the first parameter (entries (1), (3) and (4) and entries (5) and (6)), an increase of the droplet size was observed with the ultrasonication time. The same behavior was observed with the increase of the sonication power (entries (1) and (2)). This phenomenon is very surprising because droplet size usually decreases with these parameters essentially due to the greater efficiency of the emulsification step but sometime partially due to an increase of the temperature which involves a partial evaporation of the organic phase homogenization. In our case, this increase in size of the droplets could be explained by the partial precipitation of the surfactant with the increase of the medium temperature particularly marked beyond 60% ultrasonication power. Indeed, even if an ice bath is used during the sonication step to avoid the temperature increase it is not possible to prevent it completely and the reactor is a little bit hot after the homogenization step. Final temperature can locally reach 70°C. Supplementary experiments, consisting in placing a miniemulsion of 83 nm droplets (entry (1), Table 2) diameter in a hot water-bath during one hour confirmed the increase of the droplet size with the temperature (final droplet size =104 nm). By contrast, no destabilization occurred within 24 hours at room temperature for the same miniemulsion, which confirmed the miniemulsion stability during the polymerizations. Because miniemulsions prepared using DCE as solvent and 2 min ultrasonication at a power of 50% (entries (5), (7) and (8), Table 2) displayed the smallest droplets sizes, this system was selected for the rest of the study.

**Evaluation of the miniemulsion optical properties**

In IMesH\(^{+}\)BPh\(_2\)/ITX 2-component NHC generating system, the primary photochemical step is electron transfer sensitized by excited triplet ITX. To match with ITX absorption in the UV-A range and avoid IMesH\(^{+}\)BPh\(_2\) absorption, a medium pressure Hg lamp equipped with a filter at 365 nm has been used. In order to assess the level of extinction (due to the combined effect of scattering and absorption of ITX), the miniemulsions were analyzed by spectrophotometric methods based on measurements in regular transmission at 365 nm. Measurements were performed on three miniemulsions with variable organic phase content: 5, 10 and 20 w-% contained in a 1 mm thick cell (Table 2 entry (5), (7) and (8)) (Figure 2). UV-visible analysis was also carried out on the organic phase before its emulsification as a reference (blue curve, Figure 2). By following the definition of the absorbance (Equation 2), the percentage of transmitted irradiation can be determined to evaluate the extent of scattering.

\[
A = \log(\frac{I_0}{I}) \quad \text{Equation 2}
\]

With:

- A: the measured absorbance
- I: the transmitted intensity
- I₀: the initial intensity

As observed on the Figure 2, in the non-absorbing region (300–800 nm), the decrease in organic phase content is manifested by a dramatic increase of the miniemulsion transparency arising from a diminished scattering coefficient. Below 300 nm, monomer absorption domain is very large, involving an important fall of the transparency regardless of the droplet size and consequently of the scattering. At 20 w/w-% in droplet phase (D=103 nm), the transmittance is very low (I/I₀=0.3%). At 10 w/w-%, the miniemulsion is slightly transparent but only 8% of the light is transmitted. The best conditions for a highly penetrative miniemulsion is 5 w/w-% (D=36 nm; I/I₀=57%).

<table>
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<tr>
<th>Surfactant</th>
<th>Description /formula</th>
<th>HLB</th>
<th>CMC (g/L) (DLS)</th>
<th>D_{min} (nm) (DLS)</th>
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<td>16</td>
<td>5</td>
<td>170</td>
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<tr>
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<td>C_{12}H_{25}O(C_{12}H_{24})_n</td>
<td>18</td>
<td>10</td>
<td>119</td>
</tr>
</tbody>
</table>
Polymerization tests

The formulation study for the formation of the miniemulsion was performed with droplets containing only Nb, solvent and hexadecane. The tricomponent catalytic system \([\text{RuCl}_2(\text{p-cymene})]\) dimer/IMesH\(^+\)BPh\(_4^-\)/ITX needs to be added in the organic phase for the photoROMP. As previously reported, this catalytic system is very effective for the photoROMP of Nb in solution using DCM as solvent, allowing to reach full conversion in less than 5 min.\(^{33}\) Nevertheless, because DCE was selected as solvent for the organic phase of the miniemulsion, we decided to study the behavior of the catalytic system in this solvent. In addition, the ruthenium dimer pre-catalyst \([\text{RuCl}_2(\text{pCy})_2]_2\) instead of \([\text{RuCl}_2(\text{pCy})_2]_2\), was also evaluated for the photoROMP of Nb (Scheme 1).

System IMesH\(^+\)BPh\(_4^-\)/ITX/[RuCl\(_2(\text{pCy})_2\)]\(_2\): First, polymerization tests were carried out with IMesH\(^+\)BPh\(_4^-\)/ITX (5 eq./2.5 eq.) in presence of \([\text{RuCl}_2(\text{pCy})_2]_2\) (1 eq.) as pre-catalyst in solution and in miniemulsion with different dispersed phase contents (5, 10 and 20 w/w-%). For this study, the formulations were irradiated at 365 nm (20 mW/cm\(^2\)) during 30 min. Nb conversions as a function of irradiation times were plotted in Figure 3 while values of molecular weights were reported in Table 3.

In solution, using DCE as solvent, Nb conversion reached 90% after 20 min irradiation. Experimental average molar masses were 10 times higher than the theoretical ones, indicating a poor initiation efficiency of the polymerization (Table 3). Moreover, the polymer obtained exhibit broad molecular weight dispersity decreasing with the conversion, which is characteristic of an initiation rate much lower than that of propagation. The \(^1\)H NMR spectrum of the obtained polymer showed the characteristic signals of PNB, with in particular, the presence of the signal corresponding to the unsaturations of the PNB chains with the suitable integrations. The cis/trans ratios of the obtained polymers were about 50/50. (Figure S3, supplementary information).

A control experiment was also carried out with \([\text{RuCl}_2(\text{pCy})_2]_2\) alone (without IMesH\(^+\)BPh\(_4^-\)/ITX), without the NHC precursor (Figure S7, blue curve). Indeed, this precatalyst has proved to be active for the ROMP of Norbornene in DCM under UV irradiation at 365 nm in the presence of ITX.\(^{27}\) In DCE, it appears that \([\text{RuCl}_2(\text{pCy})_2]_2\), alone, is able to polymerize Nb (Nb conversion is higher than 40 % after 30 minutes of irradiation). Thus, during the Nb ROMP initiated with IMesH\(^+\)BPh\(_4^-\)/ITX/[RuCl\(_2(\text{pCy})_2\)]\(_2\), a secondary ROMP initiated by \([\text{RuCl}_2(\text{pCy})_2]_2\) alone occurs. However, it is much less active than with the IMesH\(^+\)BPh\(_4^-\)/ITX system: 70% monomer conversion is reached after 5 min irradiation with IMesH\(^+\)BPh\(_4^-\)/ITX/[RuCl\(_2(\text{pCy})_2\)]\(_2\) versus 10% at the same time with \([\text{RuCl}_2(\text{pCy})_2]_2\) alone.

![Figure 2](image-url)  
**Figure 2:** UV-visible analysis of the organic phase (DCE 65 w-%, Nb 25 w-%, hexadecane 10 w-%) and of the miniemulsions at 5, 10 and 20 w-%. Ultrasonication: 2 min at 50 %; optical path length: 1mm.

![Figure 3](image-url)  
**Figure 3:** Nb conversion as a function of the irradiation time with the catalytic system IMes\(^+\)BPh\(_4^-\)/ITX/[RuCl\(_2(\text{pCy})_2\)]\(_2\) in solution (blue curve) and in miniemulsion at 5 w-% (green curve) at 10 w-% (red curve) and at 20 w-% (purple curve).
Miniemulsion polymerization was first carried out with a 5 w/w-% organic phase content (Figure 3, green curve). Unfortunately, no conversion was observed although this miniemulsion exhibited the best penetration depth. The organic dispersed phase was probably too diluted and the droplets size too small to be activated by irradiation. By contrast, at 10 w/w-% (Figure 3, red curve), conversion reached 40 % after 30 minutes of irradiation (versus 90 % in solution). The characteristic signals of PNB were observed by \(^1\)H NMR spectroscopy (cis/trans ratio also around 50/50) (Figure S4, supplementary information). As in solution ROMP, both ROMP initiated by the formed Noels' catalyst after irradiation and by \([\text{RuCl}_2(p\text{Cy})]^2\) alone take place (Figure S7, red curve). Experimental average molar masses are also much higher than the theoretical ones and after 20 minutes of irradiation, they became so high that the PNB could not be solubilized in THF for SEC analysis (Table 3). The absence of correlation between the molar masses and the irradiation time probably comes from the difficulty of solubilizing PNB chains of high molar masses and the uncontrolled polymerization initiation. The DLS measurements showed that the particle size after polymerization was very close to the initial droplet size, attesting the good colloidal stability of the miniemulsion upon polymerization (Table 3; Figure 4). In addition, TEM pictures (Figure 4) analyses showed spherical particles with sizes consistent with the sizes measured by DLS (D=103 nm (0.14) for TEM and D=105 nm (0.23) for DLS).

Finally, the polymerization of the miniemulsion at 20 w/w-% of dispersed phase was performed in order to check if an increase of the droplet concentration could further improve the initiation process and thus Nb conversion (Figure 3, purple curve). Nevertheless, no significant improvement was observed. Indeed, the miniemulsion ROMP at 20 w/w-% exhibits almost the same conversion profile as the miniemulsion at 10 w/w-%. This phenomenon can be explained by the fact that the increase of nucleation sites (droplets nucleation) is counterbalanced by a decrease of the transparency (initial droplet size: D=143 nm (0.240) involving a very low miniemulsion transparency). The final particle size (D=147 nm (0.202)) close to the initial droplet size, proved the good stability of the emulsion.

<table>
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<th>Radiation time (min)</th>
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<th>15</th>
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<td><strong>Polymerization in solution</strong></td>
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<tr>
<td><strong>Polymerization in miniemulsion 10 w-%</strong></td>
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</tr>
<tr>
<td>(M_{\text{nth}}) (kg/mol)*</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>2.4</td>
<td>4.1</td>
<td>5.5</td>
<td>8.6</td>
<td>10.1</td>
</tr>
<tr>
<td>(M_{\text{exp}}) (kg/mol)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>486</td>
<td>203</td>
<td>Not soluble in THF</td>
<td></td>
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</tr>
<tr>
<td>(M_{\text{exp}}) (SEC)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>3.43</td>
<td>8.9</td>
<td></td>
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<tr>
<td>(D)</td>
<td>82</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>79</td>
<td>88</td>
<td>88</td>
<td>104</td>
<td>105</td>
</tr>
<tr>
<td>Average diameter (nm) (DLS)</td>
<td></td>
<td></td>
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<td></td>
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</table>

* \(M_{\text{nth}}=\pi_{\text{Nb}}M_{\text{NB}}x_{\text{H}}x_{\text{H}}/\left(n_{\text{[RuCl}_2(p\text{Cy})]}^2/2\right)\) where \(x_{\text{H}}\) is the Nb conversion, \(M_{\text{NB}}\) is the molecular weight of Nb, \(n_{\text{Nb}}\) is the initial amount of Nb, \(n_{\text{[RuCl}_2(p\text{Cy})]}\) is the initial amount of \([\text{RuCl}_2(p\text{Cy})]^2\).
Polymerization with \([\text{RuI}_2(p\text{Cy})]_2\) as ruthenium pre-catalyst: As shown in previous section, \([\text{RuCl}_2(p\text{Cy})]_2\) alone can catalyze without NHC the ROMP of Nb in solution and in miniemulsion, leading to a competitive initiation mechanism. To overcome this issue, \([\text{RuI}_2(p\text{Cy})]_2\) was proposed as a good candidate to gain complete photolatency in absence of NHC. Indeed, changing the chloride ligands by iodide ligands could induce latency of this system because of an increase of the steric hindrance and thus a limited coordination of Nb.\(^{34}\) Figure 5 (blue and green curves) displays the Nb conversion versus the irradiation time for the NHC precursor system IMesH\(^+\)BPh\(_4^–\)/ITX together with the precatalyst \([\text{RuI}_2(p\text{Cy})]_2\). In solution (Figure 5, blue curve), this tricomponent system allowed monomer conversion to reach only 45% after 30 minutes of irradiation, revealing a less active catalytic system for ROMP of Nb than the tricomponent system employing \([\text{RuCl}_2(p\text{Cy})]_2\). Nevertheless, in sharp contrast with \([\text{RuCl}_2(p\text{Cy})]_2\), no Nb conversion was observed when \([\text{RuI}_2(p\text{Cy})]_2\) ruthenium complex was used alone (red curve, Figure 5). Thus, the combination of IMesH\(^+\)BPh\(_4^–\)/ITX with \([\text{RuI}_2(p\text{Cy})]_2\) provides a fully photolatent system for the ROMP of Nb, with no competitive initiation mechanism.

Regarding the ROMP in miniemulsion at 10 w/w-% with this new catalytic system IMesH\(^+\)BPh\(_4^–\)/ITX/[\text{RuI}_2(p\text{Cy})]_2, polymers were obtained with slightly lower conversion (30% after 30 minutes of irradiation) than in solution. Good colloidal stability of the final dispersion could also be noted with particle sizes \(D = 74\ \text{nm} \ (0.49)\) close to the initial droplet size \(D = 83\ \text{nm} \ (0.32)\)).

Conclusions

In this contribution, Nb ROMP in miniemulsion was performed through a new photo-latent catalytic system. First, the miniemulsion formulation was optimized in order to obtain aqueous Nb miniemulsions with initial droplets smaller than 100 nm. Droplets with diameter of 36 nm (PDI=0.271) and 64 nm (PDI=0.228) for miniemulsion at 5 w/w-% and 10 w/w-% respectively were obtained. The UV attenuation of these emulsions was checked by UV-visible spectroscopy. Then, polymerizations were carried out by adding in the organic phase IMesH\(^+\)BPh\(_4^–\), ITX and \([\text{RuCl}_2(p\text{Cy})]_2\), in order to generate in situ a Noels’ catalyst type complex in the droplets. Thanks to this new initiator, P NB were obtained by ROMP of Nb in solution and in miniemulsion with conversions up to 98% and 55% respectively. Despite these encouraging results, \([\text{RuCl}_2(p\text{Cy})]_2\) alone appears to initiate the ROMP of Nb under UV irradiation, giving rise to a competition between two initiation mechanisms. In order to overcome this drawback, its iodine homologue \([\text{RuI}_2(p\text{Cy})]_2\) was used and proved to be fully inactive alone for the ROMP of Nb under UV irradiation (fully latent system), while in the presence of IMesH\(^+\)BPh\(_4^–\) and ITX it allowed reaching Nb conversions of about 50% in solution and about 35% in miniemulsion at 10 w/w-% of dispersed phase.

In this first study, we thus demonstrated the potential of photo-latent catalytic systems to obtain PNB by ROMP in aqueous miniemulsion. In a future work, use of another irradiation system, composed of a LED lamp and an annular reactor will be explored with this photoROMP system, with the aim to scale-up the production of PNB particles. Particular attention will be paid to the influence of the reactor geometry and the lamp power on the reactions efficiencies.
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