Synthesis of low-molecular-weight telechelic polyisobutylene

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Telechelic polyisobutylene with chlorine end-groups was made by cationic polymerization of isobutylene in methylene chloride with difunctional initiators (2,5-dimethoxy-2,5-dimethylhexane and 2,5-diacetyl-2,5-dimethylhexane). The structure of these initiators resembles that of polyisobutylene. BC13 and TiCl4 were used as catalysts. The molecular weight of these telechelics was varied between 600 and 3500 g mol\(^{-1}\), with a molecular-weight distribution ranging from 1.3 to 2.1. The materials were characterized by high-performance liquid chromatography, nuclear magnetic resonance and infra-red spectroscopy.

(Keywords: telechelic polymers; polyisobutylene; 2,5-dimethoxy-2,5-dimethylhexane; BC13; synthesis; characterization)

INTRODUCTION

The first reported synthesis of polyisobutylene dates from 1873\(^1\). It was made by heating isobutylene monomer (IB) in the presence of strong acids. With this method only low-molecular-weight oligomers with one functional end-group could be made. Later it was discovered that the molecular weight could be increased considerably by lowering the temperature of the synthesis down to \(-75^\circ C\).\(^2\)

TELECHELIC POLYISOBUTYLENE

An alternative way to use polyisobutylene (PIB) is not as a homopolymer, but as a segment in a block copolymer. If PIB is to be used as a soft block in a copolymer, it is necessary that it can be polymerized in a controlled way such that it gets defined end-groups that are reactive or can be converted to reactive end-groups. There are two different processes known to make block copolymers. One is living polymerization, where a two- or three-block copolymer is made of isobutylene and a second monomer in one pot by successively adding the components; for example, the polymerization of isobutylene followed by that of \(\alpha\)-methylstyrene.\(^3\) A second way is to make a difunctional oligomer that is polymerized in a second step to a block copolymer. This second step is normally a condensation polymerization. The difunctional oligomers that are used usually have molecular weights in the range of 500–5000 g mol\(^{-1}\).

Initiating system

The literature reports different methods for the synthesis of difunctional PIB. Some of them start from the monomers\(^4\), while others make the oligomers from high-molecular-weight polymer\(^5\). Kennedy\(^6,7\) has shown that it is possible to make dichloro-terminated oligomers by using special difunctional initiators. The main structure of these initiators that will form a carbocation with a Lewis acid are:

![Chemical structure](image)

R may be an alkyl or aryl group. In the case when a difunctional initiator is needed, R must have two of these functional groups. The initiators form stable tertiary carbocations, while the functional group and the Lewis acid together form the anion.

The best initiators are those which initiate easily so that less reactive catalysts can be used. These catalysts will initiate slowly or not at all with impurities. The best-described initiators are 1,4-dicumyl chloride and the ether or ester derived from it. A disadvantage of this initiator is that rather low temperatures are necessary for the synthesis. At temperatures above \(-60^\circ C\) the growing chain can react with the aromatic ring of the initiator with the formation of indanyl structures.\(^10\) This arylation will not happen with aliphatic initiators, so that temperatures up to 0°C can be used.\(^11\) Aliphatic initiators with a chlorine functionality are poor initiators.\(^12\)

This reaction is catalysed by Lewis acids.\(^13\) The reactivity for the polymerization decreases in the order: AlCl3 > BF3 > AlBr3 > TiCl4 > TiBr4 > BC13 > BBr3 > SnCl4. A catalyst that is too reactive is less...
suitable as it will easily form an initiating complex with
traces of impurities and will give chain transfer at higher
temperatures. It was also found that a very reactive
catalyst like AlCl₃ can use the solvent CH₂Cl₂ as an
initiator. This is, however, undesired, as it will lead to
monofunctional polymers. This initiating effect was,
however, not observed with Lewis acids less reactive than
AlCl₃. Lewis acids stronger than BC₃ will initiate
polymerization in the presence of traces of water, giving
monofunctional polymers.

With BC₃ in methylene chloride the initiation with
water will go slowly, so that the initiation of water relative
to an aliphatic ester or ether catalyst is slow. With
BC₃ the chain transfer in methylene chloride is very slow,
even at temperatures above -60°C. For these two
reasons boron trichloride is very suitable to carry out
the reaction at temperatures above -60°C.

The telechelic made with the above-described difunctional
initiator and BC₃ are chlorine end-capped with the chlorine at a tertiary carbon atom. Another way to
proceed with this end-group is to remove the chlorine as hydrogen chloride. In this case an olefinic end-group is formed to which other different reactive components can be added.

The reaction can be initiated with water but the
resulting polymer then has only one functional
end-group. The other end bears an inert head-group. If
all traces of water are removed and a difunctional initiator is added, both ends will start growing, and at
the end of the reaction two chlorine end-groups are obtained. The initiation with the difunctional initiator is similar to that with water. The initiator molecule is only
much bigger than H₂O.

This activated initiator will now start a polymerization in the same way as the water-initiated system. Only the
anion-cation complex of the polymerizing chain is different with the different initiators. With water a
BC₃OH⁻ anion is formed, while the above ether
initiator gives BC₃OCH₃ as an anion. Initiators with
ester functionalities can also be used. They will give
BC₃OCOCH₃ as anions. In the literature it is claimed
that the initiator with an ether or an ester functionality gives living polymers in the presence of
BC₃.

**Termination**

In the termination reaction with PIB it is difficult to
get an end-group other than the Cl from catalyst. This
is in contrast to most other ionic polymerizations, where
the end-group is determined by the quenching agent.

It was the aim of our study to synthesize reactive,
difunctional PIB oligomers in the molecular-weight range
of 500-5000 with a method described by Kennedy.

**EXPERIMENTAL**

**Materials**

Methylene chloride (Merck, stabilized with 2-methyl-1-butene) was refluxed for 24 h over CaH₂ and then
distilled and stored over sodium wire.

Isobutylene (Ucar) was passed through columns filled
with BaO and molecular sieves to dry it and was then condensed into the reaction vessel.

Boron trichloride (Ucar), titanium tetrachloride (Merck), methanol (Merck) and 2,5-dihydroxy-2,5-
dimethylhexane (Fluka) were all used as received.

**Analytical techniques**

**High-performance liquid chromatography.** The polymers
were characterized with a Waters HPLC. The HPLC
was provided with 100, 500 and 1000 Å Styrage-columns and an RI detector. The system was calibrated with
polystyrene. The determined molecular masses were
therefore not accurate and the h.p.l.c. results were mainly
used to establish the polydispersity of the PIB samples.
Measurements were all carried out with 10 g l⁻¹ solutions of the telechelic in tetrahydrofuran (THF) at an
operating speed of 2 ml min⁻¹.

**Infra-red analysis.** A Perkin-Elmer 1310 IR and a
Biorad 60 FTIR were used for infra-red analyses of the
PIB telechelics. The viscous telechelics were put as a paste
on one side of an NaCl plate.

**Nuclear magnetic resonance.** End-groups of the
telechelics were studied with a Bruker 80 MHz
apparatus. Solutions of 5–10% telechelic in d-chloroform were used. Tetramethylsilane (TMS) was used as an
internal reference. The samples were scanned 100 times
in order to reduce the noise.

**Initiator synthesis**

2,5-Dimethyl-2,5-hexane diacetate. The synthesis was
carried out according to a similar synthesis of t-butyl
acetate in Vogel. To 100 g 2,5-dihydroxy-2,5-dimethyl-
hexane (Fluka) was added 300 ml of acetic acid
anhydride (Fluka). After all the anhydride had been
added, the mixture was heated to the reflux temperature.
When all the 2,5-dihydroxy-2,5-dimethylhexane had
dissolved, 1 g SnCl₃ was added as a catalyst. This mixture
was held at reflux temperature for 2 h and then cooled
to room temperature. The acetic acid formed was distilled
off at 125–130°C. After the distillation of the acetic acid,
the pressure was reduced and the excess of anhydride was removed at 40–50°C (20 mmHg). The residue was
dissolved in 100 ml ether and washed with 100 ml water.
After separation of the water layer, the ether was
evaporated and the product was distilled (121°C,
22 mmHg). The yield was 44.7 g 2,5-dimethyl-2,5-hexane
diacetate (27.6%).

The analysis with n.m.r. showed three peaks:
- CH₃ 1.73 ppm
- CH₂ 1.34 ppm
- OOCOCH₃ 1.87 ppm. The synthesis of this initiator was described earlier, but with
a different method, and no yield or physical data were
given.

2,5-Dimethyl-2,5-dimethoxyhexane. To 100 g diol
(2,5-dihydroxy-2,5-dimethylhexane) 15 g sodium was
added. This mixture was intensively stirred for 7 h at
140°C and cooled to room temperature. The unreacted
sodium was removed and 55 ml MeI was added. The temperature rose several degrees after the MeI addition. After 3.5 h 100 ml hexane was added to dissolve the product. The mixture was now filtered and a liquid product remained when all the hexane was vaporized. By comparing the peak at 1.22 ppm from the –CH₃ near the hydroxyl group and the peak at 3.18 ppm from the –CH₂ near the ether group in the n.m.r. spectrum, it was established that this product contained about two-thirds monoether and one-third diol.

The above-mentioned synthesis was repeated twice with the product of the previous synthesis. First it was repeated with 9 g Na and 28 ml MeI, and then a second time with 7 g Na and 14 ml MeI. N.m.r. analysis showed that after three repeated reaction cycles most of the diol was converted to the diether. The results of this synthesis are given in Table 1. The total yield was 37%. The liquid product obtained after the third cycle was distilled at 85°C (18 mmHg). In a second initiator synthesis the product of the first cycle was used in the second reaction cycle and the total yield became higher (59% after distillation).

N.m.r. analysis showed three peaks: –CH₃ 1.49 ppm; –CH₂ 1.15 ppm; –OCH₃ 3.18 ppm. A synthesis of this initiator was not found in the literature.

**Telechelic polyisobutylene synthesis**

The synthesis was carried out in glass equipment that was dried overnight in an oven at 120°C and then mounted together while it was still hot. It was flushed with nitrogen gas and when it had reached room temperature the glass vessel was immersed in a cooling bath of −30 to −40°C. The isobutylene was condensed into this 250 ml flask for 0.5 to 2 h. During the condensation, the gas flow was measured with a calibrated flowmeter. When the required amount of IB was condensed, 150 ml precooled methylene chloride (−20°C) was added under stirring. The mixture was given 10 min to cool further to the temperature of the cooling bath. After these 10 min the initiator or initiator solution and 50 ml precooled 1 mol l⁻¹ BCl₃ solution in methylene chloride (−20°C) were added successively.

The BCl₃ solution was added fast under intensive mixing with a magnetic stirrer. The reaction was allowed to proceed for 1 h. Then the polymer was quenched by adding 10 ml methanol to the reaction flask. After quenching, more methanol was added to precipitate all the PIB. The polymer precipitated as a viscous liquid. The exact amount of condensed isobutylene was now calculated by weighing the reaction vessel with the precipitated polymer. This calculation afterwards was carried out because not all the IB that went through the flowmeter may have been condensed.

Most of the methylene chloride was removed with a Rotavapor. After this the polymer was washed a few times with fresh methanol by adding methanol and then decanting it. The remaining methanol was removed with a Rotavapor at 30°C. Care was taken to make sure that the temperature stayed low, as at higher temperatures the chlorine could evolve as HCl.

**RESULTS AND DISCUSSION**

**Influence of catalyst**

BCl₃ as well as TiCl₄ have been used as a catalyst under identical conditions. Titanium chloride is a stronger Lewis acid than BCl₃. Because of this it will more easily form an initiating complex with impurities like water, and the stronger cation will also give chain transfer more easily. Both effects are undesirable because this will give monofunctional chains. The experiments with TiCl₄ were carried out at −40 and −80°C. The syntheses at −40°C gave products with peaks at 895 cm⁻¹ in the i.r. spectrum, which means that it contained unsaturated groups (Table 2). The unsaturation was not found if BCl₃ was used. An experiment carried out at −80°C with TiCl₄ as catalyst did not show any unsaturation, but the chlorine content of this telechelic was much lower than would be expected on the basis of the initiator/monomer ratio as shown in Table 2. The ratio of end-groups to bulk groups should be 0.056 under the chosen conditions. This means that only a few of the end-groups contained a chlorine atom. The nature of the other groups was not examined. Based on these results and the good results obtained with BCl₃ (values close to the theoretical 0.056) it was decided to use only BCl₃ as a catalyst.

**Influence of initiator type**

The dimethyl ether and the diacetic acid ester of 2,5-dimethyl-2,5-hexanediol were examined as initiators for isobutylene polymerization. The above aliphatic initiators were chosen as initiators because their structure

![Diagram](https://via.placeholder.com/150)

Table 1: Synthesis of 2,5-dimethyl-2,5-dimethoxyhexane in three cycles

<table>
<thead>
<tr>
<th>Cycle</th>
<th>Diol (mol)</th>
<th>Monoether (mol)</th>
<th>Diether (mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start</td>
<td>0.68</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>1st cycle</td>
<td>0.12</td>
<td>0.30</td>
<td></td>
</tr>
<tr>
<td>2nd cycle</td>
<td></td>
<td>0.19</td>
<td>0.14</td>
</tr>
<tr>
<td>3rd cycle</td>
<td></td>
<td>0.02</td>
<td>0.23</td>
</tr>
</tbody>
</table>

Table 2: Polymerization with different catalysts

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Temperature (°C)</th>
<th>Unsaturation (i.r.)</th>
<th>Chlorine (n.m.r.)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Peak ratio*</td>
<td>CH₂ end/CH₃ bulk</td>
</tr>
<tr>
<td>TiCl₄</td>
<td>−40</td>
<td>0.44</td>
<td>0.005</td>
</tr>
<tr>
<td>TiCl₄*</td>
<td>−40</td>
<td>0.36</td>
<td>0.000</td>
</tr>
<tr>
<td>TiCl₄</td>
<td>−80</td>
<td>0</td>
<td>0.016</td>
</tr>
<tr>
<td>BCl₃</td>
<td>−40</td>
<td>0</td>
<td>0.05</td>
</tr>
</tbody>
</table>

*Ratio between infra-red peaks at 895 and 930 cm⁻¹

*With ester-type initiator. The initiator/monomer ratios were chosen to give a molecular weight of 2000
resembles that of polyisobutylene. The use of the dimethyl ether has not been reported before. The diacetate was previously used by Kennedy.

Another reason for the use of an aliphatic initiator is the absence of side-reactions like indanyl ring formation that can occur when aromatic rings are present in the initiator part. The molecular-weight distribution of polymers initiated with the different initiators is given in Figure 1 and Table 3 together with the results of an experiment without initiator.

Figure 1 and Table 3 show that the ester initiator gives a molecular-weight distribution that is much broader than the corresponding ether initiator. A broad distribution for this ester-type initiator was also found by Kennedy. The reason for a broad distribution is a low ratio between the initiation and polymerization rates.

The experiment with adding only the catalyst and no initiators shows that, without initiator, polymerization still occurs and is probably initiated by impurities. This polymer had a high molecular weight.

Because the diether gave the narrower molecular-weight distribution, it was decided to use 2,5-dimethyl-2,5-dimethoxyhexane for the polymerizations. Unless otherwise stated the reactions were carried out in the same way as PIB 2.

**Influence of initiator/catalyst ratio**

The initiator/catalyst ratio was changed to evaluate the influence of this ratio on the polymerization (Table 4). The ratio was changed between 0.2 and 1.22. Low ratios seemed to have no effect on the yield, which was always above 90% up to an initiator/BCl3 ratio of 0.5. At higher ratios no polymer at all was obtained within 2 h of reaction.

A ratio above 0.5 means that there were more ether groups from the initiator than there were BCl3 groups to form a complex with them. This ratio was specific for this ether initiator; with the ester type of initiator, polymerization took place at all ratios. The uncomplexed ether groups that are present in the case of a ratio above 0.5 seem to inhibit the polymerization in some way. It is known from the work of Polanyi et al. that diethyl ether acts as an inhibitor in polyisobutylene polymerization. This inhibition might be one reason for the fact that the molecular-weight distribution with the ether is smaller than with the ester type of initiator. The polymerization does not start immediately but only after the activation of most of the initiator.

**Molecular-weight distribution**

The molecular-weight distributions that were found with the ether initiator are equal to or lower than 2.0. Only the telechelics made with the ester type of initiator gave broader distributions, as mentioned before (Figure 1). In Figure 2 the molecular-weight distribution ($M_w/M_n$) plotted against the molecular weight ($M_n$). One sees a narrowing of the molecular-weight distribution with increasing molecular weight, as would be expected for living polymer systems. This narrowing is caused by the fact that the initiation time becomes less significant at longer chain lengths.

**Continuous monomer addition**

Isobutylene was polymerized with continuous monomer addition to examine the living character of the polymerization. This polymerization was carried out by
condensing a continuous stream of monomer into a reaction vessel with methylene chloride in it as a solvent. At 15 min after the start of the condensation the initiator and the catalyst were added. The monomer addition was continued over the next 3 h. At certain time intervals samples were taken from the reaction vessel with a syringe and then put in flasks filled with methanol. The bulk volume and concentration were corrected for the withdrawn sample volume. After standing overnight the methanol was decanted from the viscous polymer. The polymer was rinsed with fresh methanol and then dried in a vacuum oven. From the weight of the samples the amount of polymer that had been formed in the reaction flask could be calculated. The results are presented in Figure 3. The molecular-weight distribution of these samples was examined with h.p.l.c. These results are presented in Figure 4.

Figure 3 shows a linear increase of the formed polymer mass with time. This linear increase means that the polymerization is fast enough to polymerize immediately all the freshly added monomer. After about 1 h of monomer addition the polymer yield becomes less than 100% as shown by a discrepancy from the straight line in Figure 3. This decreasing yield becomes larger at the end of the experiment where the reaction had lasted for 3 h. This means that at the end of the reaction time not all added monomer is polymerized any more. The polymer concentration might be too high. It is also possible that the living head-groups have a short lifetime and that the reaction time is too long at this point. It was shown in the literature that PIB chains that were end-capped with chlorine can only slowly be reinitiated in the presence of BCl₃.

Figure 4 shows the h.p.l.c. curves from each sample point in Figure 3. In Figure 5 we have plotted the molecular weight as a function of time. One sees an increase of molecular weight during the polymerization. The increase of molecular weight becomes less after 1 h. This is also the time at which the yield becomes less than 100%. In Figure 6 we see an increase of molecular-weight distribution with time. If the polymerization was completely living during the whole experiment, an initially broad distribution would be expected, followed by a narrowing of the molecular-weight distribution with time. That the molecular-weight distribution increases with time means that some chains are still growing, but other chains have stopped growing or grow at a lower rate.
speed. This means that the polymerization under these conditions is not living. Chain transfer to monomer has not occurred, as was shown by the absence of double bonds in the i.r. and n.m.r. spectra.

**I.r. analyses**

I.r. measurements of the telechelics gave peaks that correspond with peaks for the polymer found in the literature. Only a few extra peaks originating from the end-groups were found in the telechelics, a weak chlorine peak at 645 cm⁻¹ and in some cases an olefinic peak at 890 cm⁻¹.

**N.m.r. analyses**

The telechelics were all examined with n.m.r. to determine the ratio of chlorine groups relative to the amount of isobutylene. The chlorine itself could not be seen, but the peaks of the monomer adjacent to the chlorine were shifted enough to determine this chlorine content. The bulk CH₂ and CH₃ groups give peaks at respectively 1.11 and 1.42 ppm, while the end-groups show peaks at respectively 1.68 and 1.96 ppm.

**CONCLUSIONS**

It was shown that telechelic polyisobutenes can be synthesized with chlorine end-groups. Both 2,5-dimethoxy-2,5-dimethylhexane and 2,5-dimethoxy-2,5-hexane diacetate can be used as initiators for this reaction. Even at low molecular weights a small molecular-weight distribution is obtained by using 2,5-dimethoxy-2,5-dimethylhexane as initiator, BCl₃ as catalyst and dichloromethane as solvent at -40°C. If the BCl₃/initiator ratio becomes too small, no polymerization occurs at all.

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