Investigation of isophorone diisocyanate oligoisocyanurate effect on water dispersible polyurethane properties

Oligoisocyanurates (OIC) are the products of various diisocyanates cyclotrimerization reaction, with their functionality being more than three (potential crosslinking agents). They are applicable in chemical polymer industry as an additive to diisocyanates for polyurethane coatings obtaining with various properties, as well as an isocyanate-containing component on its own for polyurethane synthesis. The OIC-based polymers possess a number of unique properties, e.g., improved thermal stability, fire resistance, UV resistance, also they are not expected to undergo hydrolysis. We studied the effect of isophorone diisocyanate oligoisocyanurate content on hydrodynamic, thermo-mechanical and thermal properties of resulting water dispersible polyurethanes. Samples of water dispersible polyurethanes containing isophorone diisocyanate (IPDI), poly (butylene adipate) (PBA) and 2,2-dimethylolpropionic acid (DMPA) were prepared following acetone method for this purpose, with IPDI OIC amount being 10–30 wt.% per pure IPDI. IPDI OIC was partially blocked with morpholine to exclude crosslinking in WDPU synthesis. Dynamic mechanical analysis (DMA), differential scanning calorimetry (DSC), and dynamic light scattering (DLS) were used to characterize the polymers obtained. It was established that oligoisocyanurates improve the characteristics of both water dispersible polyurethanes and their polymer films. Moreover, oligoisocyanurates prevent polyurethane coatings crystallization due to its branched structure, which without a doubt will affect their performance positively.

Keywords: polyurethane, water dispersible polyurethane, isophorone diisocyanate, 2,2-bis(hydroxymethyl)propionic acid, poly(butylene adipate), isophorone diisocyanate oligoisocyanurate, dynamic light scattering, dynamic mechanical analysis

Introduction

Polyurethane is one of the most demanded polymers in many industries nowadays. It stands out by a sufficiently large operating temperature range and high strength characteristics. The application of polyurethanes varies from sheets for making surfaces of spindles, rollers, wheels to base for paints, varnishes and binders for important composites.

Water-based polyurethane dispersions, namely one-(1K) and two-component (2K) waterborne polyurethane compositions are reported to be a relatively new eco-friendly type of polyurethane coating compounds. Their application tends to increase in the painting of wood, plastic, metals and other materials [1–3]. Despite the high quality, unique durability and environmental benefits of 2K waterborne polyurethanes, they have significant disadvantages related to the limited viability and relative toxicity of the isocyanate component. 1K waterborne polyurethane based on prepolymer of physical drying or self-crosslinking film forming agents appeared to be much more technologically advanced and safe.

The preparation of a prepolymer is the main stage in the synthesis of 1K water dispersible polyurethanes (WDPU). Bifunctional isocyanates, e.g. isophorone diisocyanate (IPDI) [4], hexamethylene diisocyanate (HDI) [5], toluene diisocyanate (TDI) [6] etc., polyols of various molecular weight [7], as well as hydroxyl- or amino-terminated low molecular weight bifunctional chain extenders [8] are used for that purpose. It is also necessary to reach the water dispersibility for the resulting prepolymer — to that end, a hydrophilic agent is introduced into the composition. Salt of 2,2-bis(hydroxymethyl)propionic acid (DMPA) is one of the most commonly used hydrophilic agents [9, 10].

Oligoisocyanurates (OIC), which are the products of the cyclotrimerization reaction of various diisocyanates [11, 12], are possible to use in the synthesis of WDPU. OIC found application in chemical industry as an additive to diisocyanates in order to obtain polyurethane coatings with refined properties [13], as well as an isocyanate-containing component on its own in the synthesis of polyurethane and hybrid composites [14, 15], and aerogels. It was reported [16, 17] that polyurethanes obtained using OIC had high thermal resistance and fireproof, also resistance to UV and hydrolysis. Furthermore, OIC can be modified with various low molecular...
weight compounds containing groups reactive to isocyanate ones. The above-mentioned fact also allows vary WDPU properties over a wide range.

After having reviewed the available sources, a limited number of works dedicated to WDPU synthesis using HDI [13, 18–20] and IPDI-based [3, 15] OIC were found. Moreover, the effect of the HDI OIC content on WDPU properties was studied thoroughly in the first articles, while only basic opportunity to obtain WDPU was shown in the latter ones.

In our opinion, the use of IPDI-based OIC in the WDPU synthesis seems preferable due to IPDI containing aliphatic and cycloaliphatic isocyanate groups of different reactivity. The reactivity of isocyanate groups can vary widely [21, 22] depending on the synthesis conditions. This provides the ability to control the prepolymer structure and hence the properties of the final product. Furthermore, IPDI-based OIC are completely soluble even at NCO conversion of ~73 % as noted in the [3]. In contrast, the critical gelation conversion of HDI cyclotrimerization reaction products is only 50 %. This experimental fact allows changing WDPU properties to a larger extent due to the various degree of OIC modification.

There is no doubt that 1K WDPU properties with IPDI-based OIC depend on the isocyanurate content in the system, the degree of modification and the IPDI NCO groups conversion in OIC.

The objective of this study thereby is to investigate the effect of IPDI-based OIC content in WDPU on hydrodynamic, thermal and thermo-mechanical properties of resulting products.

**Experimental**

**Materials and instruments**

IPDI (98 %) was purchased from Sigma-Aldrich and then purified by vacuum distillation at T = 90 ºC (10 mmHg); Tₘ = 158–159 ºC; d₁₂ = 1,049 g/cm³; nₚ₀ = 1,4840.

2,2-bis(hydroxymethyl)propionic acid (98 %) (DMPA) was purchased from Sigma-Aldrich, 1,2-diaminopropane (99 %) (DAP) and N,N,N-triethylamine (99 %) (TEA) were purchased from Acros Organics. These chemicals were used without further purification.

Poly(butylene adipate) (PBA) was purchased from Aldrich; the content of hydroxyl groups is 1.7 wt.% (Mₙ ~ 2000), determined by chemical method.

Acetone, morpholine (99 %) and 1,4-butanediol (99 %) (BD) were purchased from Sigma-Aldrich. These chemicals were purified by standard method [23].

Catalysts: 1,4-diazobicyclo[2.2.2]octane (98 %) (DABCO), propylene oxide (98 %) (PO) and dibutyltin dilaurate (95 %) (DBTDL) were purchased from Aldrich and were used without further purification.

**Synthesis of isophorone diisocyanate oligoisocyanurates**

Polycyclotrimerization of IPDI was carried out in the mass in presence of DABCO (0.4 wt.%) + PO (2 wt.%) catalytic system, T = 60 ºC. The reaction was controlled by FTIR spectroscopy using ALPHA spectrometer (Bruker, Germany) using NaCl cuvettes by a decrease in absorption line intensity (v = 2270 cm⁻¹) which corresponds to NCO asymmetric stretching vibrations.

IPDI-based OIC with NCO conversion of 20 % for the WDPU synthesis was partially blocked with morpholine. 0.31 g of morpholine per 1 g of IPDI OIC. The content of aliphatic and cycloaliphatic NCO groups in the IPDI OIC determined by the method described in [24] was 33 % and 67 %, respectively.

**Synthesis of waterborne polyurethanes**

Synthesis of WDPU-10 containing 10 wt.% of IPDI OIC (per pure IPDI) went as it follows. DMPA (1.20 g) and TEA (0.90 g) were put into a 250 mL flask with ground glass neck. A sample of PBA (14.89 g) and BD (0.58 g) were added to the flask and were dissolved with constant stirring and T = 55 ºC after the obtained DMPA salt was dissolved in acetone (32 g), pre-drawn on molecular sieves. Then, IPDI (10.00 g) and DBTDL (0.06 g) were added to the flask. The reaction of urethane formation was carried out for 3 hours with constant stirring and T = 55 ºC and was controlled by carrying out the titrimetric analysis [25]. After the reaction being done, partially blocked IPDI OIC (1.31 g) dissolved in acetone (5 g) was added into the flask. The reaction was carried out for 30 minutes.

After the first stage, the reaction mass was allowed to cool to the room temperature. Then, a chain extension was performed, with PDA sample (1.78 g) dissolved in 93 g of distilled water added to the flask with intense stirring ([NH₃]/[NCO] = 1). The excess acetone was removed in vacuum. The solid content in the final product was 25 %.
WDPU-20, WDPU-30 and OIC-free WDPU-0 were prepared by the described technique. The precise formulation of WDPUs obtained is reflected in Table 1. DMPA content in every sample was 4 wt.%, first stage of synthesis was characterized with the reactive groups ratio [NCO]/[OH] = 2.0.

### Table 1

<table>
<thead>
<tr>
<th>Sample</th>
<th>DMPA, g</th>
<th>TEA, g</th>
<th>PBA, g</th>
<th>BD, g</th>
<th>IPDI, g</th>
<th>IPDI OIC, g</th>
<th>Blocking agent, g</th>
<th>PDA, g</th>
<th>PBA, wt.%</th>
</tr>
</thead>
<tbody>
<tr>
<td>WDPU-0</td>
<td>1.16</td>
<td>0.87</td>
<td>14.89</td>
<td>0.58</td>
<td>10.00</td>
<td>0</td>
<td>0</td>
<td>1.80</td>
<td>50.8</td>
</tr>
<tr>
<td>WDPU-10</td>
<td>1.20</td>
<td>0.90</td>
<td>14.89</td>
<td>0.58</td>
<td>10.00</td>
<td>1.0</td>
<td>0.31</td>
<td>1.78</td>
<td>48.6</td>
</tr>
<tr>
<td>WDPU-20</td>
<td>1.25</td>
<td>0.94</td>
<td>14.89</td>
<td>0.58</td>
<td>10.00</td>
<td>2.0</td>
<td>0.62</td>
<td>1.75</td>
<td>46.5</td>
</tr>
<tr>
<td>WDPU-30</td>
<td>1.30</td>
<td>0.98</td>
<td>14.89</td>
<td>0.58</td>
<td>10.00</td>
<td>3.0</td>
<td>0.93</td>
<td>1.72</td>
<td>44.6</td>
</tr>
</tbody>
</table>

### Researching methods

The viscoelastic properties of WDPU samples were investigated by dynamic mechanical analysis (DMA) using a DMA 242 C device (Netzsch-Gerätebau GmbH, Germany): uniaxial tension on film samples, working length of 10 mm, width of 3 mm, thickness of ~0.2 – 0.3 mm, continuous scanning, T = –170 + 170 °C, a speed of 2º/min, helium atmosphere. A sinusoidal oscillating force was applied to the samples, allowing one to develop strain amplitude of up to 30 μm at fixed frequency of 1 Hz. The temperatures of relaxation transitions were determined by the inflection points on the curves of the safe modulus (E').

Thermal analysis of WDPU samples was carried out using a METTLER TOLEDO DSC822e differential scanning calorimeter with a heating rate of 20º/min.

The particle size distribution in WDPU dispersions was determined by dynamic light scattering at 25 and 60 °C using a Photocor Compact.

### Results and Discussion

As shown in [3], polycyclotrimerization can be presented as follows:

\[
\begin{align*}
&\text{Trifunctional cycles are formed at the first stage during cyclotrimerization of diisocyanates. They form n-mers and then a crosslinked polymer at subsequent reaction stages. As previously mentioned, the critical gelation conversion (CGC) in IPDI polycycloaddition is able to reach 73%. However, using IPDI OIC in WDPU synthesis with NCO conversion not having yet reached CGC will result in obtaining crosslinked polymers highly likely. This deals with the obvious fact that different compounds that contain more than three functional groups, act as possible crosslinking agents. The part of polyfunctional compounds will obviously increase with the conversion of isocyanate groups. It is necessary to block partially the NCO groups of OIC to eliminate possible crosslinked structures formation during the WDPU synthesis. We determined [3] the critical gelation conversion in reaction between IPDI OIC and bifunctional compounds at different NCO group conversion values to define the minimum of NCO groups (\(NCO_{\text{blocked}}\)) that are to be blocked. So, \(NCO_{\text{blocked}}\) vs. NCO group conversion of IPDI OIC graph was obtained based on the data mentioned above (Fig. 1). The minimum required value of \(NCO_{\text{blocked}}\) increases with increasing of isocyanate groups conversion of IPDI OIC as expected. However, polymer crosslinking and, accordingly, coating hardening after application means an open road to damp- and heat-proof coatings. Attention therefore should be paid to hydroxyl- and amine-terminated components with other functional groups as IPDI OIC blocking agents, e.g. double bonds (allyl alcohol, allylamine etc.). Self-crosslinking 1K WDPU can be obtained using that kind of IPDI OIC blocking agent. The degree of } \\
\end{align*}
\]
crosslinking for WDPU-based coatings will obviously be determined by functional groups content in the composition. Thus, there are two ways to control crosslinking. The first one is selecting highly converted IPDI OIC, and the second way is varying the IPDI OIC content in WDPU synthesis. We have investigated the second way in the context of this work.

To determine how IPDI OIC content influences waterborne polyurethanes, we prepared 3 samples containing IPDI OIC by weight (per free IPDI): 10% (WDPU-10), 20% (WDPU-20) and 30% (WDPU-30) (Table 1). OIC-free WDPU-0 obtained under similar conditions was used as a reference sample.

The size of dispersed particles and their distribution in dispersion medium is one of the main characteristics of any colloidal solution. Dynamic light scattering of all obtained dispersions was carried out to establish the IPDI OIC content impact in WDPU on their hydrodynamic characteristics. Table 2 and Figure 2 show results of the analysis.

The trend that is observed for WDPU-0 and WDPU-10 samples (Table 2) is worth mentioning, where a decrease in average hydrodynamic radius ($R_h$) and the diminishing of particle size distribution (Fig. 2) with the increase in temperature can be seen. This observed pattern is apparently the result of aggregates destruction. It should be noted that distribution narrowing occurs due to the destruction of the aggregates of dispersion
particles larger than 100 nm. In contrast, the distribution broadens with increasing temperature for WDPU-20 and WDPU-30 samples. This may indicate their lower resistance to the temperature increase.

### Hydrodynamic characteristics of WDPU

<table>
<thead>
<tr>
<th>Sample</th>
<th>$R_N$, nm</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$T = 25^\circ C$</td>
</tr>
<tr>
<td>WDPU-0</td>
<td>78.7</td>
</tr>
<tr>
<td></td>
<td>(16.0; 201.2)*</td>
</tr>
<tr>
<td>WDPU-10</td>
<td>40.8</td>
</tr>
<tr>
<td>WDPU-20</td>
<td>20.4</td>
</tr>
<tr>
<td>WDPU-30</td>
<td>36.0</td>
</tr>
</tbody>
</table>

Note. * $R_N$ was determined for every peak.

It can be seen from the Table 2 that IPDI OIC-containing WDPU are characterized by a lower value of $R_N$ and by a narrower particle size distribution in comparison to WDPU-0 at room temperature. Thus we can conclude that IPDI OIC introduction (from 10 wt.% to 30 wt.% per pure IPDI) in to WDPU positively affects on their hydrodynamic characteristics.

The analysis of IPDI OIC content influence on WDPU thermo-mechanical properties using the DMA method was the next step.

Table 3 shows the transition temperatures measured by inflection points on dynamic elastic modulus ($E''$) vs. temperature graph and by maxima on dynamic loss modulus ($E''$) vs. temperature graph. There are also elastic modulus values for sample temperature 30 °C higher than glass transition temperature.

### Thermo-mechanical and thermal properties of WDPU

<table>
<thead>
<tr>
<th>Sample</th>
<th>$E_{Tg+30^\circ C}$, MPa</th>
<th>$T_1$, °C (by $E''$)</th>
<th>$T_2$, °C (by $E''$)</th>
<th>$T_3$, °C (by $E'$)</th>
<th>$T_4$, °C (by $E'$)</th>
<th>$T_g$, °C</th>
<th>$T_m$, °C</th>
<th>$\Delta H_m$, kJ/g</th>
</tr>
</thead>
<tbody>
<tr>
<td>WDPU-0</td>
<td>1194</td>
<td>$-148$</td>
<td>$-56$</td>
<td>37</td>
<td>115</td>
<td>$-52$</td>
<td>119</td>
<td>$-22.6$</td>
</tr>
<tr>
<td>WDPU-10</td>
<td>1250</td>
<td>$-148$</td>
<td>$-55$</td>
<td>37</td>
<td>112</td>
<td>$-53$</td>
<td>116</td>
<td>$-11.1$</td>
</tr>
<tr>
<td>WDPU-20</td>
<td>1297</td>
<td>$-148$</td>
<td>$-53$</td>
<td>37</td>
<td>102</td>
<td>$-54$</td>
<td>107</td>
<td>$-13.1$</td>
</tr>
</tbody>
</table>

It is well known from the kinetic theory of elasticity [26] that the molar mass of interstitial polymer chains is inversely proportional to the dynamic elastic modulus for sample temperature 30 °C higher than glass transition temperature ($E''_{Tg+30^\circ C}$). Therefore, we can suggest relying on the modulus variation data (Table 3), that the molecular weight of polyurethane interstitial chains decreases when the IPDI OIC percentage in WDPU composition increases, with the assumption of constant density of WDPU-0 — WDPU-30 polymer samples. This is most likely due to the fact that, following its high branching and polarity, IPDI OIC enhances the polyester mesh formation possibility mainly by reason of hydrogen bonds.

All four samples revealed qualitatively equal results for temperature transitions. The first transition ($T_1$) is most likely a β-transition. The second peak on the loss modulus ($E''$) vs. temperature graph resulted by the ‘thawing’ of polyester block segmental mobility. Thereby $T_2$ is the glass transition temperature ($T_g$). It can be seen from the Table 3 that $T_g$ for studied polyurethanes there hardly depends on the IPDI OIC content. Temperature transitions $T_3$ and $T_4$ are most likely associated with initial melting of WDPU samples crystalline phase (first), as well as with their transition to a viscous flow state (second).

The nature of observed temperature transitions was confirmed by DSC analysis (Fig. 3, Table 3).

Degree of crystallinity is an important characteristic of polymer films. It can be estimated indirectly by melting enthalpy ($\Delta H_m$) magnitude due to close chemical nature of prepared WDPU. So, Table 3 indicates more than double decrease in $\Delta H_m$ value with an increasing in IPDI OIC content in WDPU. The obtained experimental fact can be apparently explained by the branched structure of OIC, limiting the polyester block ‘packing’. It should be expected that a further increase in IPDI OIC content in WDPU can lead to a completely amorphous polymer yielding.
We have investigated the effect of isophorone diisocyanate oligoisocyanurate content in water-dispersible polyurethane compositions on their hydrodynamic, thermo-mechanical, and thermal properties in this work. According to the obtained results it was found that the OIC introduction improves properties of both aqueous polyurethane dispersions and their polymer films. This is indicated, in particular, by decreasing of dispersed particle average size. Furthermore, isophorone diisocyanate oligoisocyanurates block the crystallization of coatings they involved in due to its branching. It will undoubtedly positively affect the performance of these coatings when used at reduced temperatures. Along with this, oligoisocyanurate introduction to water dispersible polyurethane compositions has no significant effect on decreasing of glass transition temperatures. It should provide high performance characteristics at low operating temperatures.

Acknowledgments
This work was performed in accordance with the state task, state registration No. AAAA-A19-119041090087-4 using the equipment of the Multi-User Analytical Center of IPCP RAS.

We also wish to express our appreciation to Senior Research, Bubnova M.L., for performing DMA analysis and helping with results explanation.

References
Исследование влияния изофorbitилдиизоцианата на свойства полиуретана и его композитов

Изображение документа и его содержание:

Доклад о влиянии изофorbitилдиизоцианата на свойства полиуретана и его композитов.

Введение:
Изофorbitилдиизоцианат (ОДИ) — турецкий диизоцианат, используемый в качестве катализатора или ускорителя реакции.

Материалы и методы:
Исследования проводили на образцах полиуретана с различным содержанием ОДИ.

Результаты и обсуждение:
Получены данные о влиянии ОДИ на структуру и свойства полиуретана.

Заключение:
Изображение документа и его содержание:

Изображение документа и его содержание:
термомеханический и жгут физико-кинетические суперисприверских полимеризаций курантыды изофорондиизоцианат (ИФДИ ОИЦ) олигоизоцианурат курантын жер зерттелген. Осы мазакта актентыйд эсептенен изофорондиизоцианаты (ИФДИ), полибутиленадипинат (ПБА) және еркін ИФДИ салыстырылғаны түрде 10–30 % салмақ ОИЦ ИФДИ, полибутиленадипинат (ПБА) және 2,2-бис(гидроксиметил)пропион кышқылы (ДМПА) және жарықтың динамикалық қышқылы (ДМПА) негізінде суперисприверских полимеризацийың ултегері синтезделді. NCO-иілді ОИЦ тобынан ВДПУ синтезі кезінде тігілген құрылымдардың пайда болуы болды. Мысалы, ушін морфологиямен қоршаған. Алынған дисперсиялар мен құрылдың орта хабының динамикалық қышқылы (ДМПА) негізінде суперисприверских полимеризацийың ултегері синтезделді. Изофорондиизоцианаттар ортаңдың негізінде жабық құрылдың динамикалық қышқылы (ДМПА) негізінде суперисприверских полимеризацийың ултегері синтезделді. Изофорондиизоцианат, 2,2-бис(гидроксиметил)пропион кышқылы, полибутиленадипинат, олигоизоцианурат изофорондиизоцианат, динамикалық қышқылы (ДМПА) негізінде суперисприверских полимеризацийың ултегері синтезделді.

Исследование влияния олигоизоциануратов изофорондиизоцианата на свойства вододисперсных полимерных покрытий

Олигоизоцианураты (ОИЦ) — продукты реакции циклодимеризации различных дисцианатов с функциональностью больше трех (потенциально сшивающие агенты). Они нашли применение в химической полимерной промышленности в качестве добавки к дисцианатам для получения полимерных покрытий с различными свойствами, а также непосредственно в качестве изоцианатсодержащего компонента для синтеза полимеров. Полимеры, синтезированные на основе ОИЦ, обладают рядом уникальных свойств, а именно: повышенной термо-, огнестойкостью, а также устойчивостью к УФ-излучению и гидролизу. В статье исследовано влияние содержания олигоизоциануратов изофорондиизоцианата (ОИЦ ИФДИ) в составе вододисперсных полимерных покрытий на гидродинамические, термомеханические и теплофизические свойства получаемых продуктов. С этой целью актентовым методом были синтезированы образцы вододисперсных полимерных покрытий на основе изофорондиизоцианата (ИФДИ), полибутиленадипината (ПБА) и 2,2-бис(гидроксиметил)пропионовой кислоты (ДМПА), содержащие 10–30 % весовых ОИЦ ИФДИ относительно свободного ИФДИ. Для исключения образования сбитых структур при синтезе ВДПУ NCO-группы ОИЦ ИФДИ частично блокировали морфолином. Полученные дисперсии и покрытия на их основе были охарактеризованы с использованием методов динамического механического анализа (ДМА), дифференциальной сканирующей калориметрии (ДСК) и дифференциального рассеяния света (ДРС). Установлено, что олигоизоцианураты улучшают характеристики как водных дисперсий полимерных покрытий, так и полимерных пленок на их основе. Кроме того, ввиду своей разветвлённости, олигоизоцианураты препятствуют кристаллизации покрытий на их основе, что, несомненно, положительно скажется на их эксплуатационных характеристиках.

Ключевые слова: полимер, водная дисперсия полимерных покрытий, изофорондиизоцианат, 2,2-бис(гидроксиметил)пропионовая кислота, полибутиленадипинат, олигоизоцианурат изофорондиизоцианата, динамическое светорассеяние, динамический механический анализ.

References


