Supramolecular polymer additives to improve the crystallization behavior and optical properties of polybutylene terephthalate and polyamides

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Florian Richter

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Chairman: Prof. Dr.
First Examiner: Prof. Dr. Hans-Werner Schmidt
Second Examiner: Prof. Dr.
Third Examiner: Prof. Dr.
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1. Introduction

1.1. Nucleating agents

As a result of their wide range of applications combined with their low costs, polymeric materials have nowadays become indispensable in our daily life. The use of additives opens up new application areas, is essential for polymer processing and ensures the specific property profile for the end consumer. The stability of polymers, their mechanical, electrical and optical properties can be effectively enhanced by the addition of specific additives. Examples for major classes of additives include antioxidants, UV-stabilizers, processing aids, flame retardants, optical brighteners, colorants, acid scavengers and nucleating agents. The latter play an important role in semi crystalline polymers.\textsuperscript{[1]}

Nucleating agents can increase the crystallization temperature and thus reduce cycle times during melt processing\textsuperscript{[2, 3]}, affect the physical properties and can in some cases improve the optical properties (clarity and haze).\textsuperscript{[3, 4, 5–9]} In heterogeneous nucleation the amount of crystal nuclei increases drastically having a strong influence on the macroscopic properties of the polymer solid state.\textsuperscript{[1, 10]} As the growth rate of the spherulites remains constant an increase in nuclei density decreases the crystal size, which enhances the elongation at break and the impact resistance. \textsuperscript{[1, 11]} The influence of nucleation on the morphology of neat isotactic polypropylene (iPP) and iPP comprising 0.15 wt\% of the sorbitol based additive bis(3,4-dimethylbenzylidene)sorbitol (DMDBS) is shown in Figure 1. The spherulite size is strongly reduced by the addition of DMDBS compared to neat iPP shown in the right micrograph of Figure 1.

\textbf{Figure 1.} Polarized optical micrographs of isotactic polypropylene without additive (left) and comprising 0.15 wt\% of DMDBS (right) after cooling from the melt to room temperature at 10 K/min.\textsuperscript{[12]}
Introduction

Typically in low molar mass compounds crystallization upon cooling occurs rather fast, whereas the crystallization of polymeric materials requires a much larger undercooling and time. The required undercooling to initiate crystallization is shortened by the addition of a nucleating agent. Usually the undercooling is determined by non-isothermal differential scanning calorimetry (DSC).\textsuperscript{[13]} Here the \textit{peak crystallization temperatures} of the exothermic polymer crystallization peak at a defined cooling rate is shifted to higher temperatures by the incorporation of a nucleating agent compared to the neat polymer. In Figure 2 the effect of different concentrations of a commercial nucleating agent on the peak crystallization temperatures ($T_{c,p}$) of iPP is shown. Neat iPP crystallizes at 110°C whereas upon the addition of a nucleating agent the crystallization temperature of iPP is distinctly increased. For lower additive concentrations the $T_{c,p}$ values decrease approaching the values for neat iPP. Here it is important to note, that with a few exceptions\textsuperscript{[14, 4]} the degree of crystallinity remains constant upon additivation.

\textbf{Figure 2.} Differential scanning thermographs of neat iPP and iPP comprising different concentrations of the commercial nucleating agent NA 11 at a cooling rate of 10 K/min.\textsuperscript{[15]}
The **nucleation efficiency** of an additive can be calculated by comparing the increase in crystallization temperature induced by a nucleating agent to a sample nucleated by its own crystal fragments.\[^{[16, 17]}\] This involves the partial melting of the polymer. Upon cooling the remaining crystal fragments act as perfect nuclei for the crystallization of the polymer. The nucleation efficiency (NE) is calculated by:

\[
\text{NE (\%)} = 100\left(\frac{\Delta T_{c,p}}{\Delta T_{c,p \text{ max}}}\right) = 100\left(\frac{T_{c,p \text{ nucl}} - T_{c,p \text{ neat}}}{T_{c,p \text{ theo}} - T_{c,p \text{ neat}}}\right)
\]

with \(T_{c,p \text{ nucl}}\) being the crystallization temperature of the polymer induced by a nucleating agent, \(T_{c,p \text{ neat}}\) being the crystallization temperature of the neat polymer and \(T_{c,p \text{ theo}}\) being the maximum crystallization temperature obtained by self nucleation experiments. According to Lotz et. al. four thermal steps are necessary to determine the highest theoretical crystallization temperature of a polymer:\[^{[16]}\]

- The first step involves the complete *erasure of the thermal history* of the polymer. Therefore the polymer is heated well above the equilibrium melting temperature for 5 min to exclude the nucleation by unmolten self nuclei upon cooling.
- An *initial standard state* is created by cooling the “erased” melt at a defined cooling rate. This step is important to easily obtain reproducible starting material for the partial melting of the polymer.
- The next step involves the partial *melting of the polymer* at a temperature \(T_s\) between the maximum and the offset of the endothermic melting peak to create stable crystal fragments in the polymer melt.
- Subsequent cooling of the melt induces *self-nucleation* and increases the crystallization temperature of the polymer.

By repeating the four steps for different \(T_s\) values, the value for the maximum crystallization temperature (\(T_{c,p \text{ theo}}\)) is obtained at the peak minimum from the highest temperature exotherm.
1.1.1. Nucleating agents for isotactic polypropylene

Isotactic polypropylene combines slow crystal growth rates with a high degree of undercooling making it a perfect system to study controlled heterogeneous nucleation.\cite{18, 19} Due to its high commercial interest a vast number of substances have been investigated as potential nucleating agents for iPP. First research was conducted on inorganic salts such as alkaline earth metal oxides, titan dioxide, potassium and sodium phosphates or minerals such as talc, kaolin and silica.\cite{20–22} However these compounds exhibit the drawback of being insoluble in the polymer matrix and thus have to be finely distributed during melt processing.\cite{7, 23} In addition organic pigments such as γ-quinacridone, anthrachinone or perylene\cite{21, 22, 24} and polymers like for example polyethylene terephthalate, polycyclopentene and polyethylene have been found to promote the crystallization of iPP.\cite{25} Apart from talcum, commercially applied nucleating agents comprise salts of organic compounds. Examples are organic salts with aluminium and alkali metals such as sodium benzoate, sodium 2,2'-methylene-bis-(4,6-di-tert-butylphenyl)phosphate (NA 11, Asahi-Denka Chemical Company)\cite{26} and disodium bicyclo[2.2.1]heptane-2,3-dicarboxylate, (Hyperform HPN 68, Milliken Chemical).\cite{27} The chemical structures of the latter two nucleating agents based on organic salts are shown in Figure 3.

![Chemical structures of important commercial nucleating agents for iPP; α-nucleating agent, NA 11, Asahi-Denka Chemical Company (left) and γ-nucleating agent, Hyperform HPN 68, Milliken Chemical (right).]
1.1.2. Supramolecular nucleating agents

An efficient heterogeneous nucleating agent has to provide a preferably large epitaxial surface to the polymer and thus has to be finely distributed within the polymer matrix. Recently, organic nucleating agents that are soluble in the polymer melt eliminating the dispersion issues gained growing interest. Upon cooling from the melt these additives crystallize prior to the polymer into fine pronounced objects, usually in the nanometer scale. These additives are called supramolecular polymer additives. The formed supramolecular aggregates provide an extremely high density of nucleation sites due to their high surface-to-volume ratio thus maximizing the epitaxial surface.\cite{28} Concerning the tailored design of new supramolecular additives some general requirements have to be met:\cite{1, 6, 29}

- Additive should bear structural moieties to allow supramolecular aggregation by intermolecular forces.
- Additive should be soluble at the processing conditions achieving an optimal distribution in the polymer melt.
- Additive becomes insoluble above the crystallization temperature of the neat polymer upon cooling from the melt.
- Additive has an appropriate surface for the epitaxial crystallization of a particular polymer.

The nucleation ability of supramolecular polymer additives is strongly dependent on their self-assembly behavior, which is among others determined by the processing conditions, the individual chemical structure and concentration of the additive.\cite{6, 30}

A general scheme for the self-assembly concept of supramolecular nucleating agents is presented in Figure 4. The characteristic of supramolecular polymer additives is their solubility in the corresponding media polymer melt not necessary on a molecular level (Figure 4 A). Upon cooling the additive molecules self organize into so-called pre-aggregates due to intermolecular forces (B). Upon further cooling these aggregates assemble to form larger nano-objects (C), which are capable of providing an appropriate surface for the nucleation of the polymer (D).
The formation of such supramolecular structures is very sensitive towards the processing conditions, the additive concentration, and the type of polymer. Accordingly for the molecular design of new supramolecular additives several factors have to be taken into account.\cite{6,31} For the one thing the strength of the intermolecular interactions governed by the symmetry of the molecule and the amount and type of hydrogen bonding units. On the other hand the choice of the peripheral substituents that influence the solubility in the particular polymer and fine tunes the additive morphology and crystallographic order. In the following developments in the field of supramolecular nucleating agents will be presented.

1.1.2.1. Sorbitol based supramolecular additives

Dibenzylidene sorbitol derivatives represent a commercially interesting and versatile class of supramolecular additives that was particularly developed with regard to the nucleation and clarification of isotactic polypropylene.\cite{4,5,32} The most important sorbitol-based nucleating agents for the α-phase of iPP are 1,3:2,4-dibenzylidenesorbitol (DBS, Millad 3905, Milliken Chemical and Irgaclear D, Ciba Speciality Chemicals), 1,3:2,4-bis(p-methylbenzylidene) sorbitol (MDBS, Millad 3940, Milliken Chemical and Irgaclear DM, Ciba Speciality Chemicals), bis(3,4-dimethylbenzylidene)sorbitol (DMDBS, Millad 3988, Milliken Chemical) and the lastest generation clarifier 1,2,3-trideoxy-4,6:5,7-bis-O-[(4-propylphenyl)methylene]-nonitol.
(TBPMN, Millad NX8000, Milliken Chemical). Owing to their excellent solubility in the polypropylene melt at different temperatures they bypass the dispersion issues associated with inorganic particles. Upon cooling a three-dimensional fibrillar nano network is formed, providing a high surface area for nucleation of the polymer.\textsuperscript{[4, 33]} Despite its commercial interest, sugar based (i.e. sorbitol or nonitol) nucleating agents have severe drawbacks, such as limited chemical and thermal resistance, the latter excluding their use as additives for other high melting semi crystalline polymers.\textsuperscript{[34]} Figure 5 presents the chemical structures of important commercial nucleating agents based on sorbitol or nonitol derivatives.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{structures.png}
\caption{Chemical structures of important sorbitol based nucleating agents for iPP; \textit{\alpha}-nucleating agents 1,3:2,4-dibenzylidenedesorbitol, DBS, Millad 3905 (a), 1,3:2,4-bis(p-methylbenzylidene) sorbitol, MDBS, Millad 3940 (b), bis(3,4-dimethylbenzylidene)sorbitol, DMDBS, Millad 3988 (c) and 1,2,3-trideoxy-4,6:5,7-bis-O-[(4-propylphenyl)methylene]-nonitol, TBPMN, NX8000, all from Miliken Chemical (d).}
\end{figure}
1.1.2.2. Trisamide based supramolecular additives

1,3,5-benzenetrisamide derivatives represent a recently developed class of nucleating agents that solve the aforementioned issues of sorbitol derivatives, showing remarkable nucleation properties in iPP\(^6, 9, 35-37\) and polyvinylidene fluoride (PVDF)\(^{38}\). Besides their use as nucleating agents, these compounds can, depending on their structural details, be applied in electret materials based on iPP to improve their charge storage behavior\(^{39}\) or as processing aids to improve the flow properties of the polymer melt.\(^{40}\) Additionally selected 1,3,5-benzenetrisamide derivatives exhibit liquid crystalline behavior\(^{41}\) and can act as gelators for various organic solvents\(^{42}\) or water \(^{43}\). The schematic structure of 1,3,5-benzenetrisamide derivatives is shown in Figure 6. These compounds consist of a central core with 1,3,5 substitution, moieties that are able to form intermolecular hydrogen bonds and nonpolar peripheral substituents, being responsible for the crystallographic order and the dissolution behavior in the polymer melt. Owing to their amide moieties these compounds can establish intermolecular forces via hydrogen bonds and ensure one-dimensional columnar self-assembly. By contrast to sorbitol acetal derivatives, 1,3,5-benzenetrisamides combine excellent chemical resistance with high thermal stability and thus can also be applied in high melting technical thermoplastics such as polyamides or polyesters.

![Figure 6. Schematic structure of 1,3,5 benzenetrisamide derivatives.](image)

By this structural concept a new family of supramolecular nucleating agents for iPP and PVDF was developed over the past decade (Figure 7). So far the following cores were investigated: 1,3,5-benzene tricarboxylic acid (a)\(^6, 12, 35, 37, 44\), 5-amino isophthalic acid (b)\(^6, 12, 44\), 3,5-
diamino benzoic acid \((\text{c})\)^{[6, 12, 44]}, 1,3,5-triamino benzene \((\text{d})\)^{[6, 36, 44]}, cis,cis-1,3,5-cyclohexane tricarboxylic acid \((\text{e})\)^{[45]} and trisamides based on the melamine core \((\text{f})\)^{[46]}

![Figure 7. Basic structures of nucleating agents based on the motive of C3-symmetrical trisamides; 1,3,5-benzenetrisamide \((\text{a})\), 5-amino isophthalic acid \((\text{b})\), 3,5-diamino benzoic acid \((\text{c})\), 1,3,5-triamino benzene \((\text{d})\), cis,cis-1,3,5-cyclohexane tricarboxylic acid \((\text{e})\) and trisamides based on the melamine core \((\text{f})\).](image)

Out of the numerous compounds synthesized and investigated only selected derivatives are capable to induce the \(\alpha\) or \(\beta\) crystal structure within iPP, or enhance the optical properties of iPP. Blomenhofer et al. reported on the use of substituted 1,3,5-benzenetrisamides as highly efficient nucleating and clarifying agents in iPP.\(^{[6]}\) The additives were, depending on their chemical structure, found to selectively induce the \(\alpha\)-phase of iPP and enhance the optical properties even at very low concentrations. Based on these structural motifs the latest generation clarifying agent “IRGACLEAR® XT 386” was introduced by Ciba Holding SA in the year 2004.\(^{[47]}\) In 2009 Abraham et. al were the first to study the effect of 1,3,5-benzenetrisamides on the nucleation of PVDF as another semi-crystalline polymer.\(^{[38]}\) The polymer crystallization temperature and the dissolution and crystallization behavior of the additive in the PVDF melt was investigated as function of the concentration (Figure 8). As shown in Figure 8 the temperature composition diagrams are divided into three regions. In region I the additive is soluble in the polymer melt and does not crystallize upon cooling. In region II the additive is soluble at the processing temperature and crystallizes upon cooling
in fibrillar nano-objects that provide a surface for epitaxial crystallization of the polymer and induce nucleation. In region III the additive is not completely soluble under the applied temperature and nucleation both takes place on the surface of the crystallized supramolecular structures as well as on the surface of the not dissolved additive. It was found that certain 1,3,5-benzenetrisamides were soluble in the PVDF melt and self assembled prior to the polymer upon cooling promoting nucleation of PVDF.

Figure 8. Crystallization temperature of PVDF (top graphs) and the dissolution temperature of the additive in the PVDF melt (triangles up) and crystallization temperature of the additive from the PVDF melt (triangles down) (bottom graphs) as function of the additive concentration. The dashed line in the top graphs represents the crystallization temperature of extruded neat PVDF $T_{c,p} = 139.3$ °C. The dotted line in the bottom graphs represents the maximum temperature (250 °C) during DSC measurements.
Lightfoot et al. were first to study the crystal structure of 1,3,5-benzenetrisamides. As example 1,3,5-tris(2,2-dimethylpropionylamino)benzene was investigated.\textsuperscript{[48]} The crystal structure reported was composed of infinite π-stacked rods supported by a triple helical network of hydrogen bonds (Figure 9 A). Due to the conjugation of the amide linkages with the aromatic core, the amide bonds are not arranged perpendicular to the benzene core, but tilted resulting in a staggered arrangement of the benzenetrisamide molecules.

The directional orientation of substituted 1,3,5-benzenetrisamides was discussed by Kristiansen et al. who examined the stacking based on the tert-butyl substituted derivative N,N',N''-tris(tert-butyl)benzene 1,3,5-tricarboxamide (Figure 9 B).\textsuperscript{[44, 49]} It was found that the linear aggregates can possess two different helical hands, depending on the c-direction of the amide moieties (Figure 9 C). In addition, the repeat distance in the arrangement was given, which is in close proximity to the spacing of the two methyl groups in iPP, thus favoring epitaxial growth.

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**Figure 9.** (A) Crystal structure of N,N',N''-tris(2-methoxyethyl)benzene 1,3,5-tricarboxamide composed of infinite π-stacked rods supported by a triple helical network of hydrogen bonds \textsuperscript{[48]}; (B) staggered arrangement of N,N',N''-tris(tert-butyl)benzene 1,3,5-tricarboxamide; (C) Possible configurations of N,N',N''-tris(tert-butyl)benzene 1,3,5-tricarboxamide with the direction of the helical bonding patterns showing up (left) and down (right).\textsuperscript{[49]}
1.1.2.3. Bisamide based supramolecular polymer additives

Examples of supramolecular structures based on bisamides, bisureas and amide-ureas have been described in the literature. These molecules usually comprise a linear structure containing amid- and/or urea-groups. Here self assembly occurs generally into a sheet-like supramolecular structure. In most cases these compounds were investigated as organogelators. Their application in the field of supramolecular nucleating agents for iPP has also been reported. Furthermore these compounds were investigated to improve the charge storage properties of iPP.

The schematic structure of linear molecules with two units capable for hydrogen bonding is shown in Figure 10. The molecules consist of a central unit being substituted symmetrically with hydrogen bonding units to promote sheet-like self assembly. By varying the peripheral substituents, the packing of the molecules and thus their dissolution properties in the polymer melt can be adjusted specifically to the polymer. Furthermore these substituents allow a fine tuning of crystallographic order and surface morphology of the additives. In addition the substituents control the distance between the individual layers.

Based on this schematic structure, Mohmeyer et al. studied the effect of low molecular-weight 1,4-phenylene-bisamides as nucleating agents for iPP. The solubility behavior of the additives, the nucleation efficiency and the ratio of the α- to the β-phase was...
investigated with symmetrical and asymmetrical substituted bisamides, also as function of the concentration (Figure 11).

![Chemical structures of selected additives: isomers of dicyclohexyl-substituted 1,4-phenylene bisamides 1-3 and cyclohexyl/n-alkyl-substituted 1,4-phenylene bisamides with a variation of the length of the alkyl chain 4-6.](image)

**Figure 11.** Chemical structures of selected additives: isomers of dicyclohexyl-substituted 1,4-phenylene bisamides 1-3 and cyclohexyl/n-alkyl-substituted 1,4-phenylene bisamides with a variation of the length of the alkyl chain 4-6.\(^{[55]}\)

Above a critical concentration, the additives were found to induce the β-polymorph of iPP that exhibits enhanced mechanical properties.\(^{[56]}\) Nucleated iPP with the β-phase strongly scatters light. The haze is about 100 % for a 1.1 mm sample. The highest amount of β-phase was observed for the symmetrically substituted bisamides. However subtle changes in the chemical structures had significant influence on the nucleation efficiency. As seen from the wide angle X-ray patterns in Figure 12, the inversion of one amide group from compound b to c resulted in a distinct increase in β-content.
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Figure 12. Wide-angle X-ray diffraction patterns of injection molded iPP plaques (1.1 mm thickness) comprising the dicyclohexyl-substituted 1,4-phenylene bisamides b (left) and c (right) at a concentration of 0.13 wt%.\[55\]

Similar studies were conducted by Lu et al. who investigated the effect of eight N,N´-diphenyl bisamides on the formation of the β-crystalline form in iPP.\[57\] Here N,N´-diphenyl succinamide and N,N´-diphenyl glutaramide were found to be the most efficient β-nucleators. New Japan Chemical investigated dicarboxylic acid derivatives for the use as β-nucleating agents in iPP. The N,N-dicyclohexyl-2,6-naphtalene dicarboxamide was commercialized under the trade name NJ STAR NU-100 (New Japan Chemical/Rika International, Figure 13)\[7\]. The additive was found to be one of the most efficient β-nucleating agents.

![Chemical structure of the β-nucleating agent NJ STAR NU-100, New Japan Chemical/Rika International.](image)

Figure 13. Chemical structure of the β-nucleating agent NJ STAR NU-100, New Japan Chemical/Rika International.
1.2. Visual appearance of transparent polymers

The appearance of a transparent object in the visible spectrum mainly derives from the transmitted light that is passing through a sample as schematically shown in Figure 14.\textsuperscript{[58]} As already mentioned above, the optical properties such as gloss, transmission, haze and clarity can be improved by the addition of the so-called “clarifying agents”. According to ASTM D 1003 the latter two parameters haze and clarity are defined through the diffuse component of the total transmitted light in the visible spectral range. The amount of light that is scattered in angles larger than 2.5° is referred to as “haze” and is a measure of the cloudiness of a sample, where high haze values cause a decrease in contrast.\textsuperscript{[59]} Plastics with a haze of 100 % strongly scatter light and appear optically opaque whereas plastics with a haze of 0 % appear “water clear”.

![Figure 14. Parameters of the visual appearance of transparent objects (top) and influence of haze (bottom left) and clarity (bottom right) on an image.\textsuperscript{[58]}](image-url)
Clarity is defined by the amount of light that is scattered in angles smaller than 2.5°, where low clarity values cause a loss in sharpness of an image. Clarity of 0 % would mean a total loss in sharpness, whereas sheet glass exhibits clarity values of 100 %. The influence of haze and clarity on the visual appearance of image is illustrated in Figure 14 (bottom). In semi-crystalline polymers differences in refractive indices of the crystalline and amorphous phase leads to scattering which reduces the optical transparency of a sample. Furthermore, scattering on anisotropic structures (i.e. spherulites) plays an important role in the optical appearance of a material. The total amount of scattered light can be decreased by reducing the spherulites size or by preventing spherulite structures to be formed.\cite{3, 7, 60, 61} However, a reduction in crystallite size alone by the addition of a nucleating agent cannot be regarded as only one reason for the improvements in optical properties.\cite{6, 9} In 2009 Bernland et al. studied the solid-state structures of nucleated and clarified iPP by means of small-angle light scattering (SALS). It was found that an efficient clarifying agent for iPP has to be capable of preventing the formation of spherulite structures but induce rod- or shish-kebab like crystalline entities.\cite{61}

In Figure 15 the optical appearance of injection molded iPP platelets are presented as function of the additive concentration for 1,3,5-tris(2,2-dimethylpropionylamino)benzene. The optical properties haze and clarity critically depend on the composition of the mixture. Usually in these kind of diagrams an “optimum” concentration for minimum haze can be observed, that coincides with the solubility limit of the additive at the processing conditions.\cite{6, 9} At the optimum concentration at around 200 – 300 ppm the best optical performance in haze and clarity is found (middle). At lower concentrations a modest reduction in haze can be observed presumably due to less available nucleation sites. Here the optical properties deteriorate approaching the values for neat iPP (left). However if the amount of additive exceeds the “optimum” concentration, scattering on undissolved additive particles increases the haze and reduces clarity (right).
Figure 15. Optical properties (haze ⭐, and clarity ⬤) of iPP (thickness 1.1 mm) comprising different amounts of 1,3,5-tris(2,2-dimethylpropionylamino)benzene; left: neat iPP; middle: iPP with optimum additive concentration; right: iPP with 0.25 wt% of additive exceeding the optimum concentration.
1.3. Laser transparency of polymers

The steady progress in the development of polymeric products increases the demand for more complex and sophisticated components. The demands on the complex geometries, especially in small parts, can only be hardly satisfied by classical joining technologies, such as adhesive bonding or mechanical joining. Were these conventional procedures come up against their limits, welding processes can fulfill the requested requirements.\(^{[62]}\) Commonly the conjunction of two components by the influence of heat is referred to as welding. The required energy can either be supplied by convection, conduction, friction or radiation. In case of laser beam welding the energy is applied by radiation in the near infrared range (NIR) absorbed in the bulk material.\(^{[63]}\)

In this connection, two main techniques are of commercially applied. Firstly the butt welding process, where the junction occurs temporally separated to the plasticization of the polymer. Second, the more commonly applied transmission laser beam welding. In this process the two joining partners are brought in contact and held together with a moderate clamping force. The upper part has to be preferably transparent to the wave length of the irradiating laser, the lower part being absorbent. The absorption of the transmitted beam results in heating at the interfaces and melting of both parts due to thermal conduction (Figure 16).\(^{[64]}\)

![Figure 16. Schematic representation of the transmission laser beam welding process of a laser-transparent (top) and a laser-absorbing (bottom) polymer. \(^{[63, 65]}\)](image)

In spite its procedural advantages and the ability to join different materials, transmission welding of polymers exhibits limitations given by the laser transparency of the base material that varies strongly for different polymers. Figure 17 shows the transmission spectra for
injection molded samples with a thickness of 2 mm of polyamide 66 (PA66), polyamide 6 (PA6) and polybutylene terephthalate (PBT). The transmittance for light in the near infrared range is considerably lower for PBT compared to the polyamides. The use of PBT as laser transparent material especially in thick parts is therefore limited.

Figure 17. Transmission spectrum of PBT (Ultradur B4520), PA6 (Ultramid B3K) and PA66 (Ultramid A3k) measured on injection molded samples with a thickness of 2 mm.

The interactions of electromagnetic radiation with polymeric materials can be described by the three parameters “Transmission” (T), “Absorption” (A) and “Reflection” (R) (Figure 18). As light strikes the surface of a polymer a small amount of the incoming light is reflected. The transmitted portion of light gets partly absorbed by the polymer molecules or possible inhomogenities. The remaining fractions are either transmitted directly (specular transmission) or after being scattered on the polymer spherulites (diffuse transmission). These spherulites usually cause back-scattering associated with a loss in transmitted energy or a broadening of the laser beam. As demonstrated in Figure 17 these phenomena play a strong role in the laser transmittance of polybutylene terephthalate (PBT) limiting its usability for transmission welding techniques.
**Figure 18.** Interactions of electromagnetic radiation with polymers.
2. Objectives and scope of this thesis

The objectives and scope of this thesis is to explore for the first time supramolecular nucleating agents for the technical important semi-crystalline polymers polybutylene terephthalate and polyamides. The principle aim is to increase the polymer crystallization behavior, to control the polymer morphology and to improve the optical properties.

Supramolecular polymer additives based on 1,3,5-benzenetrisamides

The first part of this thesis is dedicated to the nucleation of PBT and semi-crystalline polyamides with supramolecular nucleating agents based on 1,3,5-benzenetrisamides. The schematic structure of the investigated trisamides is shown in the following.

- Peripheral nonpolar substituent
- Moiety forming hydrogen bonds
- Central core with 1,3,5-substitution

Given the current state of the art, a prediction of the nucleation properties based on the molecular structure of the additive is not possible. In order to achieve the aim to identify suitable nucleating agents and improve the crystallization behavior and optical properties of PBT and polyamides the following issues have to be addressed:

- As the individual chemical structure of the additives is crucial for the self-assembly/nucleation behavior, the core and peripheral substituents have to be systematically varied to establish structure property relations.
- A large number of 1,3,5-benzenetrisamide derivatives have to be tested with respect to their nucleation potential in PBT and polyamides.
- In order to evaluate the nucleation ability of the large number of additives, a rapid and reliable screening method is of advantage. In this work a screening process developed at the Chair Macromolecular Chemistry I utilizing temperature dependent polarized light microscopy has to be adapted to PBT and polyamides.
Objectives and scope of this thesis

• To explore the nucleation properties in more detail for selected compounds, the *additive dissolution and crystallization behavior* in the melt and the *crystallization temperature of the polymers* as function of the additive concentration have to be investigated. The additive dissolution and crystallization temperatures in the melt are determined by polarized optical microscopy and the polymer crystallization temperature is determined by DSC.

• Another aim of this thesis is to obtain more insight in the *morphology and size* of the *supramolecular nano-structures* which are formed upon cooling within the polymer melt. Experiments have to be developed to selectively remove the polymer matrix, in order to reveal the supramolecular structures. To visualize the supramolecular nano-structures scanning electron microscopy is applied.

• Finally the capability of 1,3,5-benzene-trisamides to increase the laser transparency of PBT is evaluated in collaboration with the BASF SE.

Supramolecular polymer additives based on bisureas

The aim of the second part is the synthesis and characterization of a new class of supramolecular nucleating agents, tailored to semi-crystalline polyamides. The objective is to improve the crystallization behavior and the optical properties. The linear molecules consist of a central unit being substituted symmetrically with two hydrogen bonding units to favor a sheet-like self-assembly.
To vary the self-assembly properties in the polyamide melt additives with amide and urea moieties are investigated. An additional structural variation of fundamental interest is the introduction of trans-1,4-cyclohexane and cis-1,4-cyclohexane as central unit to vary the linearity of the additives.

- Based on the schematic structure the \textit{synthesis} of novel bisamide and bisurea derivatives has to be carried out.
- Standard characterization methods such as NMR-spectroscopy, mass spectroscopy, thermogravimetry and differential thermal analysis are applied for \textit{characterization of the compounds}.
- To establish \textit{structure property relations} concerning the \textit{nucleation and clarification of semi-crystalline polyamides}, the peripheral substituents are systematically varied in length and degree of branching. The polymer crystallization temperature is determined by DSC. The optical properties are measured on injection molded samples with a standard ASTM procedure.
- To obtain a more detailed insight in the \textit{solid state morphology} the \textit{spherulite size} and the \textit{crystal morphology} of the polyamides have to be investigated on injection molded specimens. As main methods, polarized optical microscopy and wide angle x-ray scattering are applied.
- The \textit{effect of cooling rate and sample thickness on the optical properties} is studied in detail by varying mold temperature and mold thickness during the injection molding process.
- Finally the capability to increase the laser transmittance of PA6 is evaluated and compared to the commercially available nucleating agent talc.
3. Methods and procedures

The main focus of this work is on the investigation of the nucleation and clarification ability of supramolecular polymer additives in semi-crystalline polymers. The basic molecular characterization was carried out by standard techniques such as NMR- and mass spectroscopy, thermogravimetric analysis and differential thermal analysis. In order to evaluate the nucleating and clarifying potential of the supramolecular additives in various semi-crystalline polymers, efficient and reliable methods and procedures were adapted and optimized. Most of the procedures presented in the following have already been established by coworkers from the Chair Macromolecular Chemistry I for iPP\textsuperscript{[12, 46, 31]}, and PVDF\textsuperscript{[30]} and were specifically adjusted to suit the specific conditions of PBT and polyamides. The advantage of these procedures resides in the possibility to obtain significant information with only small amounts of material. The polymer crystallization and melting temperatures were characterized by differential scanning calorimetry after microscale polymer processing. From injection molded platelets the optical characteristics (transmission, haze, clarity, laser transparency) were obtained and the polymer morphology was investigated. Optical and electron microscopy permit a deeper insight into the self-assembly of the supramolecular additives.

In the following the applied techniques and conditions will be briefly summarized particularly in view of the specifics of PBT and polyamides:

- Compounding
- Injection molding
- Thermal analysis
- Polarized optical microscopy
- Additive screening method
- Optical characterization
- Laser transparency measurements
- Wide angle X-ray diffraction
- Scanning electron microscopy
3.1. Compounding

To ensure a preferably homogenous distribution of the additives, the polymer granulate was first pulverized in a freezer mill. The obtained powder was blended with the additive powder. A glass bottle with the polymer powder comprising the initial additive concentration was clamped in a tumble mixer and blended for 24 h at 40 - 45 rpm (Figure 19).

![Figure 19. Tumble mixer for the preparation of polymer powder/additive powder blends.](image)

The so obtained powder mixtures were compounded in a co-rotating twin-screw compounder (DSM Xplore 15 mL) under nitrogen atmosphere (Figure 20).

![Figure 20: DSM Xplore 15 mL micro-compounder with the marked heating zones H1, H2, and H3.](image)
The processing parameters had to be adjusted to the respective polymer and are presented in Table 1. In order to prevent the material from partial melting and adhesion to the feeding hopper, the temperature in the first heating zone (H1) was lower compared to the melting temperature of the polymer. After compounding the melt was discharged and collected either as polymer string or directly filled into the injection molding unit. Different additive concentrations were prepared by successively diluting the initial concentration with defined amounts of neat material. The neat polymer was treated in the same way to obtain blank control which was extrudated. Within the thesis the values reported for the crystallization temperatures of PBT and polyamides are an average of the extruded blank control.

Table 1: Processing parameters for the investigated polymers and their measured melt temperatures.

<table>
<thead>
<tr>
<th></th>
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</thead>
<tbody>
<tr>
<td>PBT</td>
<td>230-270-270</td>
<td>255</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PA6</td>
<td>230-260-260</td>
<td>250</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PA66</td>
<td>260-300-300</td>
<td>280</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PA6/66</td>
<td>230-250-250</td>
<td>245</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PA6/12</td>
<td>230-250-250</td>
<td>245</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PA12</td>
<td>180-240-240</td>
<td>210</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The preparation of a concentration series is exemplarily shown for PA6 in Table 2. For two cleaning runs and the first sample run a polymer/additive powder blend with a concentration of 1.5 wt% was used. For the following runs the initial additive concentration was diluted with a mixture of the polymer/additive powder blend and neat PA6 as shown in Table 2. After determining the dead volume of the compounder, the exact additive concentration in the extrudate could be calculated as follows:

With a dead volume of 5.4 g (for PA6) and an initial additive concentration of 1.5 wt%, the amount of additive within the dead volume is 0.081 g. In the first dilution run 6.8 g of the initial powder mixture with 1.5 wt% of additive and 1.8 g of neat PA6 are added. The amount of additive in 6.8 g of the initial powder mixture is 0.102 g. Thus the total amount of additive within the compounder in the first dilution runs is 0.183 g. The exact additive concentration can be calculated from the total amount of additive divided by the amount of polymer:

\[ \frac{(0.081 \text{ g} + 0.102 \text{ g})}{(5.4 \text{ g} + 6.8 \text{ g} + 1.8 \text{ g})} \times 100 \% = \frac{0.183 \text{ g}}{14 \text{ g}} \times 100 \% = 1.307 \% \]
Table 2. Initial weights and additive concentration for a concentration series in PA6.

<table>
<thead>
<tr>
<th>Run</th>
<th>Comment</th>
<th>$m_{\text{powder mixture}}$ [g]</th>
<th>$m_{\text{neat polymer}}$ [g]</th>
<th>$c_{\text{Additive}}$ [wt%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>cleaning</td>
<td>14.0</td>
<td>-</td>
<td>1.500</td>
</tr>
<tr>
<td>2</td>
<td>cleaning</td>
<td>8.6</td>
<td>-</td>
<td>1.500</td>
</tr>
<tr>
<td>3</td>
<td>1. sample</td>
<td>8.6</td>
<td>-</td>
<td>1.500</td>
</tr>
<tr>
<td>4</td>
<td>2. dilution</td>
<td>6.8</td>
<td>1.8</td>
<td>1.307</td>
</tr>
<tr>
<td>5</td>
<td>3. dilution</td>
<td>4.6</td>
<td>4.0</td>
<td>0.997</td>
</tr>
<tr>
<td>6</td>
<td>4. dilution</td>
<td>3.9</td>
<td>4.7</td>
<td>0.802</td>
</tr>
<tr>
<td>7</td>
<td>5. dilution</td>
<td>2.7</td>
<td>5.9</td>
<td>0.599</td>
</tr>
<tr>
<td>8</td>
<td>6. dilution</td>
<td>1.6</td>
<td>7.0</td>
<td>0.402</td>
</tr>
<tr>
<td>9</td>
<td>7. dilution</td>
<td>0.4</td>
<td>8.2</td>
<td>0.198</td>
</tr>
<tr>
<td>10</td>
<td>8. dilution</td>
<td>0.3</td>
<td>8.3</td>
<td>0.109</td>
</tr>
<tr>
<td>11</td>
<td>9. dilution</td>
<td>0.1</td>
<td>8.5</td>
<td>0.053</td>
</tr>
<tr>
<td>12</td>
<td>10. dilution</td>
<td>0.0</td>
<td>8.6</td>
<td>0.020</td>
</tr>
</tbody>
</table>
3.2. Injection molding

Platelets for optical characterization were prepared by injection molding the compounded blends using a micro-injection molding machine (DSM Xplore 12 mL) under nitrogen, yielding specimens with a diameter of 25 mm and different thicknesses (0.5 mm, 1 mm, 2 mm, 3 mm, 5 mm). The molds for the different thicknesses and the corresponding injection molded specimens are presented in Figure 21.

![Figure 21](image)

Figure 21. Injection molds with different thicknesses (0.5 mm, 1 mm, 2 mm, 3 mm and 5 mm) (left) and injection molded specimen with a diameter of 25 mm and a thickness of 1 mm (right).

The injection unit was directly filled with the polymer melt, discharged from the twin-screw compounding. The injection molding conditions for the different polymers are listed in Table 3.

Table 3: Injection molding conditions of the investigated polymers.

<table>
<thead>
<tr>
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<tbody>
<tr>
<td>PBT</td>
<td>250</td>
<td>80</td>
<td>6</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>PA6</td>
<td>250</td>
<td>100</td>
<td>6</td>
<td>10</td>
<td>10</td>
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<tr>
<td>PA66</td>
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<td>10</td>
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<td>10</td>
<td>10</td>
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<td>8</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>PA12</td>
<td>230</td>
<td>60</td>
<td>6</td>
<td>10</td>
<td>10</td>
</tr>
</tbody>
</table>
3.3. Thermal analysis

DSC (differential scanning calorimetry) measurements for the determination of the polymer melting and crystallization temperatures were conducted on a Perkin Elmer Diamond DSC and a Mettler Toledo DSC/SDTA 821e under nitrogen at standard heating and cooling rates of 10 K/min. To erase the thermal history, samples were heated well above the equilibrium melting temperature for 5 min before each cooling run. For each sample two heating and cooling scans were performed. The starting and end temperatures of the polymers used in the frame of this work are presented in Table 4. Values for the polymer crystallization temperature ($T_{c,p}$) were determined at the exothermic peak minimum in the second cooling scan. Melting temperatures ($T_{m}$) were determined at the maximum of the endothermic peak of the second heating scan. Thermogravimetric analyses (TGA) of the additives were performed on a Mettler Toledo TGA/SDTA851e under nitrogen blanket at a heating rate of 10 K/min. Melting temperatures of the additives were determined by simultaneous differential thermal analysis (DTA).

Table 4. Starting and end temperatures for the DSC measurements of the investigated polymers. For each sample two heating and cooling scans were performed under nitrogen at 10 K/min. Samples were held at the end temperature for 5 min before each cooling run.

<table>
<thead>
<tr>
<th>Polymer</th>
<th>Starting temperature [°C]</th>
<th>End temperature [°C]</th>
</tr>
</thead>
<tbody>
<tr>
<td>PBT</td>
<td>50</td>
<td>260</td>
</tr>
<tr>
<td>PA6</td>
<td>30</td>
<td>250</td>
</tr>
<tr>
<td>PA66</td>
<td>30</td>
<td>280</td>
</tr>
<tr>
<td>PA6/66</td>
<td>30</td>
<td>240</td>
</tr>
<tr>
<td>PA66/6</td>
<td>30</td>
<td>250</td>
</tr>
<tr>
<td>PA6/12</td>
<td>30</td>
<td>240</td>
</tr>
<tr>
<td>PA12</td>
<td>30</td>
<td>230</td>
</tr>
</tbody>
</table>

To determine the nucleation efficiency of the additives, self-seeding experiments according to Lotz et al.\textsuperscript{[9, 6]} were adapted for PBT and polyamides at standard heating and cooling rates of 10 K/min. The nucleation efficiency scale is calculated by:

$$NE(\%) = 100(\Delta T_{c,p}/\Delta T_{c,p \ max}) = 100(T_{c,p \ nucl} - T_{c,p \ neat})/(T_{c,p \ theo} - T_{c,p \ neat})$$  \hspace{1cm} (1)

The crystallization temperature of neat melt-processed polymer ($T_{c,p \ neat}$) was determined at the exothermic peak minimum upon cooling from the melt after the thermal history was erased above the equilibrium melting temperature. After partial melting for 5 min at a
temperature $T_s$ between the maximum and the offset of the endothermic melting peak, the remaining crystal fragments act as perfect nuclei for the crystallization of the polymer and increase the crystallization temperature upon cooling. The value for the highest theoretical polymer crystallization temperature ($T_{c,p\,\text{theo}}$) is obtained at the exothermic peak minimum upon cooling from $T_s$. The increased polymer crystallization temperature by the addition of a nucleating agent is described as $T_{c,p\,\text{nuc}}$. 

3.4. Polarized optical microscopy

Polarized light microscopy was performed using an optical microscope (Nikon, DIAPHOT 300) equipped with a hot stage (Mettler, FP82HT). Optical micrographs were recorded by a Nikon ACT-1 software using a digital camera (Nikon, DMX1200). The crystallization ($T_{c,a}$) and dissolution temperatures ($T_{d,a}$) of the additives were determined at the disappearance and reappearance of birefringent structures in compounded polymer samples at a heating and cooling rate of 5 K/min. The morphology of the injection molded platelets was determined on thin sections with a thickness of 10 μm. Sections were cut on a Leica RM 2255 rotary microtome parallel to the flow direction as indicated in Figure 22.

![Figure 22. Thin section (10 μm thick) from injection molded specimen cut parallel to the flow direction.](image)

3.5. Additive screening method

In order to evaluate the nucleation ability of a large number of compounds a rapid and reliable screening method is of great interest. In this work we utilized an additive screening process established by Abraham et. al.\[4\] to identify suitable compounds for the use as nucleating agents. For this purpose a compression molded polymer film was placed on a microscopic slide. A small amount of additive was positioned in the middle of the film and the whole setup was covered and positioned in a hot stage under a polarized optical microscope. The samples were heated well above the melting temperature of the polymer for 5 min whereas the additives partly dissolve and diffuse into the surrounding polymer melt. The setup was slowly cooled at 10 K/min to monitor the crystallization processes of both the additives and the polymer. Optical micrographs of the screening process were recorded by a Nikon ACT-1 software using a digital camera (Nikon, DMX1200).
3.6. Optical properties

The optical properties Transmission, Haze, and Clarity were determined according to ASTM D-1003 on injection molded platelets using a Haze-Gard Plus instrument (BYK Gardner GmbH, Germany). All reported values are an arithmetic average of at least three measured specimens. The optical properties of the samples were measured 24 h after their preparation.

3.7. Laser transparency measurements

Laser transparency (LT) measurements were conducted at the BASF SE using a FOBA DP50 Nd:YAG laser marking system. The irradiating laser wave length was 1064 nm with a power output of 2 W. For determination of the laser transmittance the beam was divided into two discrete rays that were detected separately by a reference sensor (signal 1) and a measurement sensor (signal 2) after passing the polymer sample. All measurements were conducted in the middle of the injection molded specimens on at least three samples per additive and concentration. The LT values were calculated according to the following formula, as the ratio from signal 2 and signal 1:[66]

\[
Transmission [\%] = \frac{\text{signal 2}}{\text{signal 1}} \times 100 \quad (2)
\]

Figure 23. Schematic representation of the measurement set-up for determining the laser transparency of polymers. The laser passes through a beam splitter and the separate rays are detected by a reference sensor and a LT-sensor after penetrating the sample.
3.8. Wide angle X-ray diffraction

The crystal morphology of injection molded polyamide samples (thickness of 1.1 mm) was determined by wide angle X-ray diffraction. Measurements were performed with a Bruker D8 Advance X-ray diffractometer using CuKα radiation (λ = 1.54 Å). Data was recorded in the range of 5-45° (2θ) with a step size of 0.025° and a step time of 10 sec. The crystalline peak and the amorphous halo were separated according to a method described by Murthy et al.\[^{67}\] using amorphous templates with Origin8G software. After profile analysis of the diffraction scans the content of α- and γ-crystal modification was calculated by

\[
CI_{\alpha} (\%) = \frac{\sum A_{\alpha-\text{form}}}{\sum (A_{\alpha-\text{form}} + A_{\gamma-\text{form}}) + A_{\delta-\text{form}}} \tag{3}
\]

\[
CI_{\gamma} (\%) = \frac{\sum A_{\gamma-\text{form}}}{\sum (A_{\alpha-\text{form}} + A_{\gamma-\text{form}}) + A_{\delta-\text{form}}} \tag{4}
\]

\[
CI(%) = CI_{\alpha} + CI_{\gamma} \tag{5}
\]

with \(A_{\alpha-\text{form}}\) and \(A_{\gamma-\text{form}}\) being the area under the crystalline peaks of the α- and γ-modification, respectively and \(A_{\delta-\text{form}}\) being the area under the amorphous halo.

3.9. Scanning electron microscopy

Samples for the morphological studies were prepared by melting PBT comprising different concentrations of trisamide in DSC pans under nitrogen at 280°C. To ensure complete dissolution and distribution of the trisamides the samples were held for 5 min at this temperature. The samples were cooled to room temperature with defined linear cooling rates of 10, 40, 80, and 110 K/min. After removal of the solid samples from the DSC pans, the surface was hydrolyzed in a carefully stirred solution of 20 wt% sodium hydroxide in water for 2.5 h at 110 °C. Residual NaOH was removed by carefully stirring the specimens in water. The dried samples were sputtered with platinum (0.8 nm) by a Cressington Sputter Coater 208HR. Scanning electron micrographs were recorded using a Zeiss 1530 FESEM (SEM imaging). Histograms were obtained by measuring the lateral dimensions of at least 250 objects with Zeiss AxioVision LE software. The size distribution is a mean value from the lateral dimensions of all measured objects.
4. Nucleation of polybutylene terephthalate

4.1. Introduction

Polybutylene terephthalate (PBT) ranks among the semi-crystalline engineering thermoplastics and is distinguished by its good mechanical properties including high strength, rigidity, and toughness.\cite{68}

![PBT structure](image)

PBT

Its wide continuous service temperature and high chemical resistance combined with excellent electrical properties qualifies PBT for applications in electronics, automotive engineering and mechanical engineering.\cite{69} However PET dominates the market of fiber, film, and bottle molding applications by far, whereas PBT is mostly used for injection molding applications due to its high crystallization rate.\cite{68} For PBT, two crystal modifications can be formed, the triclinic α-phase and the also triclinic β-phase.\cite{70, 71} While the molecular packing of both modifications are very similar, differences in the fiber identity period were observed that can be attributed to conformational changes in the four-methylene group sequence (G−G−TTGG, α-phase and TSTS−T, β-phase) as shown in Figure 24.\cite{71}

![Figure 24. Triclinic unit cell and the corresponding cell dimensions of the α-form (left) and β-form (right) of PBT in the projection along the 010 plane as reported by Yokouchi.\cite{71}](image)
The \( \alpha \)-polymorph is commonly obtained by cooling from the melt whereas the \( \beta \)-phase occurs under strain and stress. By mechanical deformation and relaxation, the transition between the two polymorphs takes place reversibly.\[^{[71]}\]

Depending on the crystallization conditions different types of spherulites can be found (Figure 25).\[^{[72, 73]}\] A 45° maltese cross pattern is obtained upon fast cooling from the polymer melt. In contrast to this unusual type of spherulites, the 90° pattern (usual type) develops most likely from solution for example with a mixture of tetrafluoro acetic acid (TFA) in CCl_{4}. Isothermal crystallization at temperatures above 180°C or slowly cooled material yields a mixture of both types of spherulites.

![Figure 25. Types of spherulites formed at different crystallization conditions: (a) Unusual type from crystallization at low temperatures; (b) Usual type formed from solution; (c) Mixed type from isothermal crystallization above 180 °C.\[^{[73]}\]

In the literature only few examples for the application of nucleating agents in polybutylene terephthalate could be found. Heuseveldt et al.\[^{[74]}\] reported on a mixture consisting of PBT and a nucleating agent, e.g. talc, carbon black and certain inorganic phosphates. Zhang et al.\[^{[75]}\] studied the effect of elastomer-modified nano-SiO₂ [SiO₂-(E-MA-GMA)], a commercial nucleating agent (Bruggolen® P 250), and talc on the non-isothermal crystallization at different cooling rates. It was observed that all three compounds acted as heterogenous nuclei for the crystallization of PBT. The influence of pigments and mineral fillers on the non-isothermal crystallization behavior was investigated by Pillin et al.\[^{[76]}\] Generally a nucleating effect with 0.1% (w/w) pigment content was observed. Phthalocyanine Green showed the strongest influence on the crystallization of PBT. Soluble supramolecular nucleating agents for PBT have not been reported yet.
4.2. Results and discussion

In this chapter we report on 1,3,5 benzenetrisamides as supramolecular nucleating agents for PBT. To evaluate structurally different 1,3,5 benzenetrisamides as potential nucleating agents, a screening method described by Abraham et al.\textsuperscript{[4]} for PVDF was adapted for PBT. Promising compounds were investigated in a concentration range from 0.006 wt% (60 ppm) to 0.8 wt% (8000 ppm) and compared with respect to their dissolution and crystallization behavior of the additive in the PBT melt and the crystallization temperature of PBT. To obtain more insight in the morphology and size of the formed nano-objects, the PBT matrix was carefully hydrolyzed in alkaline solution. The supramolecular objects were examined and the average size was analyzed by scanning electron microscopy (SEM).

4.2.1. Chemical structures and thermal properties of the investigated 1,3,5-benzenetrisamides

43 derivatives of 1,3,5 benzenetrisamides were screened with respect to the nucleation of PBT. In order to find nucleating agents for PBT the central core and peripheral substituents were systematically varied. The additives are based on four different central cores, 1,3,5 benzenetricarboxylic acid, 1,3,5-triaminobenzene, 2,4,6-trimethyl-1,3,5-trisaminobenzene and 1,3,5-trisaminotoluene. The peripheral substituents were selected from branched aliphatic, cycloaliphatic and aromatic moieties. The chemical structures of the substituted 1,3,5-benzenetrisamides, their melting temperatures and temperatures at 10 % weight loss (T\textsubscript{10 wt%}) are summarized in Table 5.

The investigated compounds comprise for all four cores short aliphatic (1-6, 19-22, 28-31, 37, 38), cycloaliphatic (7, 8, 23, 24, 32, 39) or aromatic moieties (9-11, 25, 26, 33-35, 40-42). Additionally 1,3,5-benzenetrisamide derivatives with adamantyl (12, 13, 27, 36, 43) or polar carboxylic acid- or amino-groups as well as their salts (14-18) have been investigated. Due to their stable amid linkage, all investigated compounds exhibit excellent thermal stability reflected by high melting and weight loss temperatures. The thermal behavior was obtained from TGA measurements, reported as the temperature at a weight loss of 10 % (T\textsubscript{10 wt%}). The additive melting temperatures were simultaneously determined by DTA. Owing to the formation of strong intermolecular hydrogen bonds, a large number of the investigated 1,3,5 benzenetrisamides sublime completely without melting, indicated also by a weight loss of 100 %. As example the compounds 1-5 with short branched aliphatic substituents exhibit no
melting endotherm and show direct sublimation with a 100 % weight loss. The $T_{-10 \ wt\%}$ ranged between 350°C and 380°C. Increasing the number of C atoms resulted in derivatives which showed melting behavior and simultaneous evaporation from the liquid phase. Compounds 6-8 have melting temperatures between 315 and 419°C. The derivatives 9-11 with aromatic substituents showed similar melting behavior with melting temperatures between 377 and 403°C. It is interesting to note that all compounds with adamantyl substituents (12, 13, 27, 36, 43) sublimed directly and showed an even higher thermal stability ($T_{-10 \ wt\%}$) above 440°C. A classification of the 1,3,5 benzenetrisamides 14-18 with polar substituents according to their thermal properties is difficult due to their high water uptake at ambient conditions. Therefore only the values for the melting temperatures are reported in Table 5.

Inversion of the amide linkages in the central core unit has an influence on the melting temperature of the 1,3,5 benzenetrisamides. For example the cyclohexyl derivative 23 showed a melting transition about 85°C lower compared to the corresponding compound 7 based on a 1,3,5 benzenetricarboxylic acid core. Comparing the iso-propyl substituted derivatives 1 and 19, the compound based on 1,3,5-triaminobenzene (19) melted at 290°C whereas 1 sublimed. The aromatic derivatives 25 and 26 showed a melting endotherm at 278 and 304°C.

The 1,3,5-benzenetrisamides 29 and 30 with methyl substitution at the core in 2,4,6-position showed comparably low melting temperatures of 288°C and 280°C, whereas their corresponding derivatives 20 and 21 sublimed directly. Out of the nine derivatives based on 2,4,6-trimethyl-1,3,5-trisaminobenzene, seven sublimed without melting. It is interesting to note that the aromatic compounds 33-35 sublimed directly whereas the aromatic derivatives based on a 1,3,5-benzenetricarboxylic acid and a 1,3,5-triaminobenzene core exhibit a melting transition.

From the seven compounds based on the 1,3,5-trisaminotoluene core the derivatives 40-42 with aromatic substituent melted at temperatures between 332 and 353°C, whereas the remaining compounds sublimed and exhibit no melting endotherm.
Table 5. Chemical structures of the screened 1,3,5-benzenetrisamides, their melting temperatures $T_m$ (DTA) and temperatures at 10% weight loss $T_{-10\text{ wt} \%}$ (TGA, $N_2$ atmosphere). The results on the nucleation ability of PBT were obtained by a screening method with polarized optical microscopy.

<table>
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<tr>
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| Substituent (R) | \[
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R \ R \\
O \ O \\
H \ H \\
\end{array}
\] | \[
\begin{array}{c}
N \ \ \ N \\
R \ \ \ R \\
\end{array}
\] | \[
\begin{array}{c}
N \ \ \ N \\
R \ \ \ R \\
\end{array}
\] | \[
\begin{array}{c}
N \ \ \ N \\
R \ \ \ R \\
\end{array}
\] | \[
\begin{array}{c}
N \ \ \ N \\
R \ \ \ R \\
\end{array}
\] |
| $T_m$ [°C] | subl. | subl. | subl. | subl. | subl. |
| $T_{-10\text{ wt} \%}$ [°C] | 353 | 380 | 358 | 350 | 362 |
| PBT nucleation | + | + | + | + | + |

<table>
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<tr>
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<th>7</th>
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\end{array}
\] | \[
\begin{array}{c}
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R \ \ \ R \\
\end{array}
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\begin{array}{c}
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R \ \ \ R \\
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\] | \[
\begin{array}{c}
N \ \ \ N \\
R \ \ \ R \\
\end{array}
\] | \[
\begin{array}{c}
N \ \ \ N \\
R \ \ \ R \\
\end{array}
\] |
| $T_m$ [°C] | 315 | 371 | 419 | 377 | 403 |
| $T_{-10\text{ wt} \%}$ [°C] | 342 | 380 | 420 | 402 | 418 |
| PBT nucleation | - | + | + | + | + |

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\end{array}
\] | \[
\begin{array}{c}
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R \ \ \ R \\
\end{array}
\] | \[
\begin{array}{c}
N \ \ \ N \\
R \ \ \ R \\
\end{array}
\] |
| $T_m$ [°C] | 334 | 371 | 419 |
| $T_{-10\text{ wt} \%}$ [°C] | 393 | 448 | 455 |
| PBT nucleation | + | + | + |

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OH \\
N \ \ \ H \\
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\] | \[
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OH \\
N \ \ \ H \\
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\] | \[
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Na^+ \\
OH \\
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\] | \[
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Na^+ \\
OH \\
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\] | \[
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O \ \ \ OH \\
Na^+ \\
OH \\
\end{array}
\] |
| $T_m$ [°C] | 329 | 394 | 395 | 374 | 390 |
| $T_{-10\text{ wt} \%}$ [°C] | - | - | - | - | - |

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<th>22</th>
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\end{array}
\] | \[
\begin{array}{c}
N \ \ \ N \\
R \ \ \ R \\
\end{array}
\] |
| $T_m$ [°C] | 290 | 278 | 304 | subl. | |
| $T_{-10\text{ wt} \%}$ [°C] | 320 | 374 | 372 | 344 | 402 |
| PBT nucleation | - | + | + | + | + |

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R \ \ \ R \\
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\] | \[
\begin{array}{c}
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R \ \ \ R \\
\end{array}
\] | \[
\begin{array}{c}
N \ \ \ N \\
R \ \ \ R \\
\end{array}
\] | \[
\begin{array}{c}
N \ \ \ N \\
R \ \ \ R \\
\end{array}
\] |
| $T_m$ [°C] | 285 | 278 | 304 | subl. |
| $T_{-10\text{ wt} \%}$ [°C] | 378 | 424 | 424 | 474 |
| PBT nucleation | - | - | - | + |
Table 5 (continued). Chemical structures of the screened 1,3,5-benzenetrisamides, their melting temperatures $T_m$ (DTA) and temperatures at 10% weight loss $T_{-10\,\text{wt\%}}$ (TGA, N$_2$ atmosphere). The results on the nucleation ability of PBT were obtained by a screening method with polarized optical microscopy.

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<td>438</td>
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<tr>
<td>PBT nucleation</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
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<td>![Structure 35]</td>
<td>![Structure 36]</td>
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<td>subl.</td>
<td>subl.</td>
<td>subl</td>
</tr>
<tr>
<td>$T_{-10,\text{wt%}}$ [°C]</td>
<td>418</td>
<td>439</td>
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<td>PBT nucleation</td>
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<td>-</td>
<td>+</td>
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<table>
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<td>405</td>
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<td>423</td>
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<td>PBT nucleation</td>
<td>-</td>
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<td>-</td>
<td>-</td>
<td>-</td>
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<td>![Structure 43]</td>
</tr>
<tr>
<td>$T_m$ [°C]</td>
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<td>subl.</td>
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<tr>
<td>$T_{-10,\text{wt%}}$ [°C]</td>
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<td>450</td>
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<tr>
<td>PBT nucleation</td>
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<td>+</td>
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</table>

a) subl.: sublimation; b) nucleation of PBT (+); c) no nucleation of PBT (-); d) n.d.: not determined due to the hygroscopicity of the compound;
4.2.2. Nucleation properties

In order to evaluate the nucleation ability of the large number of 1,3,5-benzenetrisamides investigated, a rapid and reliable screening method is of an advantage. In this work we adapted a screening process described by Abraham et al.\textsuperscript{[38]} utilizing temperature dependent polarized light microscopy. For this purpose a compression molded film of PBT was placed on a microscopic slide (Figure 26). A small amount of additive was positioned in the middle of the film and the whole setup was covered and placed in a hot stage under a polarized optical microscope. The samples were held at 280°C for 5 min whereas the 1,3,5-benzenetrisamide derivatives partly dissolve and diffuse into the surrounding polymer melt. The setup was then slowly cooled at 10 K/min to monitor the self-assembly process of the dissolved additive molecules into supramolecular nano-objects. The crystallization processes of both the additives and the polymer were recorded by a digital camera attached to the polarized optical microscope.

![Microscopic slide](image)

**Figure 26.** Setup for the screening process as established by Abraham et. al.\textsuperscript{[38]}

Figure 27 compares as an example for compound 6 and 21 micrographs recorded between crossed polarizers at different temperatures. Both micrographs at 280°C show on the right the birefringent undissolved additive in the PBT melt. The black region corresponds to the optical isotropic melt of PBT. Upon cooling to 230°C, both trisamides crystallize into fine needles. This starts from the region of the non dissolved additive. In case of 21 at 200°C the nucleation of PBT is induced on the surface of the additive needles and adjacent in the PBT melt. This indicates that very fine supramolecular structures of 21 are present, which are not visible by optical light microscopy, but capable to nucleate PBT. At lower temperature of 195°C, the PBT crystallization in the region of the additive is completed and first crystallites of PBT are formed in the remaining additive free melt. Derivative 6 is a trisamide which does
not nucleate PBT. At 230°C the trisamide crystallizes in form of fine needles, which continue to grow upon lowering the temperature (200°C). The needles do not act as nucleation sites for the crystallization of PBT. At 195°C the PBT nucleation occurs simultaneously in the additive free PBT melt and in between the needles.
Figure 27. Optical micrographs from polarized light microscopy of the 1,3,5-benzenetrisamides 21 and 6 (scale bar 200 μm). Samples were heated, kept at 280°C for 5 minutes, cooled with a rate of 10 K/min and observed at different temperatures. The additive is visible in the isotropic PBT melt (black) on the right. The non-dissolved additive diffuses into the PBT melt and crystallizes upon cooling into fine needles. In the case of 21 nucleation of PBT is induced on the surface of the additive, whereas 6 is not capable to nucleate.
The screening results for the 1,3,5-benzenetrisamides 1-43 are also presented in Table 5. Nucleation of PBT is denoted with (+) and no visible nucleation with (-). With exception of compound 6 all compounds based on the 1,3,5-benzenetricarboxylic acid core were found to be capable of nucleating PBT. Regarding the derivatives 19 to 27 based on the 1,3,5-triaminobenzene core, five out of nine compounds showed nucleation of PBT. Usually derivatives based on 1,3,5-triaminobenzene are better soluble than their corresponding analogues from the first additive series. Therefore 19 and the compounds 24-26 were too good soluble and showed no nucleation effect in PBT. From the investigated derivatives based on the 2,4,6-trimethyl-1,3,5-trisaminobenzene core six out of nine nucleate PBT. It is interesting to note that the derivatives 29 and 30 based on 2,4,6-trimethyl-1,3,5-trisaminobenzene do not nucleate, whereas trisamides 2 and 3 based on the 1,3,5-benzenetricarboxylic acid core and 20 and 21 based on the 1,3,5-triaminobenzene core bearing the same substituent do nucleate PBT. Due to their asymmetric structure the mono-substituted 1,3,5-benzenetrisamides 37-43 can only establish weaker hydrogen bonds and should be more soluble in the polymer melt. This reflects in lower nucleation ability as only three out of seven screened derivatives were capable to nucleate PBT (38, 42, 43).

Beside the compounds with nonpolar aliphatic or aromatic moieties, further 1,3,5-benzenetrisamides derivates bearing polar carboxylic acid- or amino-groups as well as their salts have been investigated. As the 1,3,5-benzenetrisamides with nonpolar substituents have been tailored with respect to the nucleation of the also nonpolar iPP, the aim was to offer an appropriate surface to the more polar PBT for the epitaxial crystallization on the surface of the additive. The selected compounds 14-18 are shown in Table 5. Compounds 14, 16, and 17 bearing polar carboxylic acid- or amino substituents exhibited excellent solubility in the polymer melt and hence were not capable to nucleate PBT. Sodium or chloride salts of these derivatives were nearly insoluble but also showed no nucleation effect in PBT.

The nucleation efficiency depends also on the additive concentration. Therefore selected 1,3,5-benzenetrisamides were investigated in a concentration range from 0.006 wt% (60 ppm) to 0.8 wt% (8000 ppm). The different concentrations were prepared by subsequently diluting the initial concentration of 0.8 wt% in the melt with neat PBT. By this dilution process the following concentrations of 0.4 wt%, 0.2 wt%, 0.1 wt%, 0.05 wt%, 0.025 wt%, 0.0125 wt% and 0.006 wt% were obtained. From each concentration the additive
dissolution and crystallization temperatures in the melt were determined by polarized optical microscopy and the crystallization temperature of PBT was determined by DSC. The crystallization temperature of melt processed neat PBT was determined at 188.5°C.

In Figure 28 the DSC cooling curves of neat PBT and PBT comprising 0.2 wt% and 0.4 wt% of 3 are shown. Due to the presence of nuclei at higher temperatures the crystallization temperature of PBT is distinctly increased upon addition of 3. As less nucleation sites are available at lower concentrations, the \( T_{c,p} \) values decrease with the amount of additive added. The crystallization temperatures reported in the following section were determined from the exothermic peak minimum of the second cooling scan.

![Figure 28](image)

**Figure 28.** DSC cooling scans of neat PBT and PBT comprising 0.2 wt% of 3.

In the following, three sets of selected compounds will be compared to discuss structure-property relations as function of the concentration. The first set compares the influence of different substituents of four trisamides based on the 1,3,5-benzenetricarboxylic acid core. The second set compares trisamides with the same substituents based on the 1,3,5-triaminobenzene core. The third set is based on the 2,4,6-trimethyl-1,3,5-trisaminobenzene as core. The top row of figures 2-4 shows the crystallization temperature (\( T_{c,p} \); dots) of PBT as function of the additive concentration. The dashed horizontal line indicates the \( T_{c,p} \) of neat PBT as reference. The temperature composition diagrams of the binary mixtures can be divided into three regions. Region I represents the concentration range where the additive is soluble in the polymer melt and does not crystallize upon
Nucleation of polybutylene terephthalate

cooling. In these cases no birefringence is observed in the melt. It should be pointed out that the lack of birefringence in the polarized optical microscope does not necessarily mean a molecular solution. Small aggregates of trisamide molecules may still be present in the melt, similar to amphiphilic molecules in optically isotropic micellar solutions. In region II the additive is soluble at the processing temperature and crystallizes upon cooling in fibrillar nano-objects, visible as birefringent structures in the melt. These supramolecular structures provide a surface for epitaxial crystallization of the polymer and induce nucleation. In region III the additive is not completely soluble under the applied temperature. Therefore nucleation both takes place on the surface of the crystallized supramolecular structures as well as on the surface of the not dissolved additive. The bottom graphs show the additive dissolution \( (T_{d,a}, \text{triangles up}) \) and crystallization temperature \( (T_{c,a}, \text{triangles down}) \) as determined by polarized light microscopy. In the bottom graphs the dashed horizontal line indicates the processing temperature of the mixing process at 260°C.

In the following, sets of selected compounds will be compared to reveal structure property relations and to investigate the concentration dependence. The first set compares the influence of different substituents of four trisamides based on the 1,3,5-benzenetricarboxylic acid core. The second and third set compares four trisamides with the same substituents based on the 1,3,5 triaminobenzene core and the 2,4,6-trimethyl-1,3,5-trisaminobenzene core respectively. For each compound the top row of Figure 29, Figure 31 and Figure 33 show the crystallization temperature \( (T_{c,p}, \text{dots}) \) of PBT as function of the additive concentration. The dashed horizontal line indicates the \( T_{c,p} \) of neat PBT as reference. The bottom graphs show the dissolution \( (T_{d,a}, \text{triangles up}) \) and crystallization temperature \( (T_{c,a}, \text{triangles down}) \) of the additive. Here, the dashed horizontal line indicates the processing temperature of 260°C.

The temperature composition diagram of a binary mixture of a polymer and an additive that is soluble in the polymer melt can be divided into three concentration regions. Region I represents the concentration range where the additive is soluble in the polymer melt and does not crystallize upon cooling prior to the polymer as determined by polarized optical microscopy. In these cases no birefringence is observed in the melt. It should be pointed out that the lack of birefringence does not mean a molecular solution because very small aggregates of trisamide molecules can be present, similar to optically isotropic micellar solutions. In region II the additive is again soluble at the processing temperature but
crystallizes in fibrillar nano-objects upon cooling from the melt. These supramolecular structures provide a surface for the epitaxial crystallization of the polymer and thus induce nucleation. In region III the additive is only partly soluble in the processing range and nucleation can both take place on the surface of the crystallized additive whiskers as well as the undissolved additive moieties.

Figure 29 compares the data of the first set of additives based on the 1,3,5-benzenetricarboxylic acid core with the substituents iso-propyl 1, tert-butyl 2, 1,1-dimethylpropyl 3 and cyclohexyl 7. In agreement with the observations from the screening test, additive 1 with iso-propyl substituent is readily soluble in the PBT melt. In the concentration range from 0.006 - 0.4 wt% (I) the additive is soluble and does not crystallize prior to the polymer. Consequently no nucleation of PBT occurs. At the concentration of 0.8 wt% and the processing temperature of 260°C additive 1 is to the largest extent soluble in the PBT melt. Only a small fraction is insoluble as indicated by a weak birefringence. Upon cooling the dissolved fraction of the additive crystallizes prior to the polymer and nucleation takes place mainly at the formed supramolecular structures, but also at the not dissolved additive. The crystallization temperature of PBT is increased by 9.2°C from 188.5°C to 198.2°C. The small change in the chemical structure of the peripheral substituent from iso-propyl to tert-butyl results in a less soluble compound. Only up to 0.1 wt% additive 2 is soluble and does not crystallize upon cooling and no increase in T_{c,p} is observed. In the concentration range from 0.2 - 0.8 wt% the additive is partially soluble and nucleates PBT with a T_{c,p} of 192.9°C at only 0.2 wt%. Additive 3 with 1,1-dimethylpropyl substituent is slightly better soluble than 2 indicated by comparably lower T_{d,a} and T_{c,a} values in the range from 0.2 - 0.8 wt%. At 0.2 wt% (II) the additive is soluble at the processing temperature and crystallizes prior to the polymer. Consequently nucleation takes place only at the formed supramolecular structures with a T_{c,p} of 195.7°C. At higher concentrations (III) the maximum T_{c,p} of additive 3 was found to be 198.0°C at a concentration of 0.8 wt%. The cyclohexyl substituted derivative 7 is completely soluble over the entire investigated concentration range, does not crystallize prior to the polymer and consequently is not capable to nucleate PBT.
Figure 29. Crystallization temperatures of PBT \( (T_{c,p}) \) containing the 1,3,5-benzenetrisamide derivatives 1, 2, 3 and 7 (top graphs) and additive dissolution \( (T_{d,a}) \) and crystallization temperatures \( (T_{c,a}) \) (bottom graphs) as function of the additive concentration. The dashed lines at 188.5°C in the top graphs represent the crystallization temperature of neat PBT. The dashed lines in the bottom graphs at 260°C indicate the processing temperature.
The structure and appearance of the supramolecular entities within the polymer matrix is strongly dependent on the individual chemical structure of the additive. At high concentrations compounds 1-3 based on 1,3,5-benzenetricarboxylic acid all showed fibrillar structures in the polymer melt upon cooling from the dissolved state as shown exemplarily for 3 in Figure 30. Here at 280°C the additive is fully soluble in the polymer melt and visible crystallization in from of fine filaments can be observed at 235°C. Upon further cooling the supramolecular structures grow larger and later act as nuclei for the polymer crystallization at temperatures below 200°C. However no conclusions can be drawn from the appearance of the fibrillar structures on the nucleation ability in the polymer.

![Figure 30. Polarized optical micrographs of PBT comprising 0.4 wt% of 3 cooled at 10 K/min at different stages of the cooling process.](image-url)
Figure 31 presents the results of the 1,3,5-benzentrisamide derivatives 19-21 and 23 based on the 1,3,5-triaminobenzene core with the same substituents as the compounds in the first set. Inversion of the amide linkages has a pronounced influence on the solubility in the polymer melt. Trisamides based on the 1,3,5-triaminobenzene core are generally more soluble. Hence the $T_{d,a}$ and $T_{c,a}$ of the additives are shifted towards lower temperature values. This observation was also made for trisamide based nucleating agents of similar structure in isotactic polypropylene [78, 79] and polyvinylidene fluoride [4]. Compared to compound 1, which is not soluble at the concentration of 0.8 wt%, compound 19 based on the 1,3,5-triaminobenzene core is soluble over the entire investigated concentration range. Although no birefringent structures are visible in polarized optical microscopy, a slight increase in $T_{c,p}$ of 2°C up to 190.5°C was observed at a concentration of 0.8 wt%. This indicates that very fine supramolecular nano-structures are present capable to induce nucleation. Also compound 20 based on the 1,3,5-triaminobenzene core with tert-butyl substituents is much more soluble compared to compound 2. Only at a concentration of 0.4 wt% additive 20 self assembles upon cooling prior to PBT. But only a marginal increases of $T_{c,p}$ was observed. At a concentration of 0.8 wt% the additive is not completely soluble and an increase in $T_{c,p}$ to 193.2°C was found. Also compound 21 with 1,1-dimethylpropyl substituent is much more soluble than compound 3. Up to a concentration of 0.4 wt% the additive remains soluble. At a concentration of 0.8 wt% 21 is soluble at the processing conditions, self assembles upon cooling and nucleates PBT very efficiently with a $T_{c,p}$ of 199.1°C, which corresponds to an increase of 10.6°C. This is the highest $T_{c,p}$ value found in this study. Compound 23 with the cyclohexane substituent is soluble over the entire investigated concentration range similar to compound 7.
Figure 31. Crystallization temperatures of PBT ($T_{cp}$) containing the 1,3,5-benzenetrisamide derivatives 19, 20, 21 and 23 (top graphs) and additive dissolution ($T_{da}$) and crystallization temperatures ($T_{ca}$) (bottom graphs) as function of the additive concentration. The dashed lines at 188.5°C in the top graphs represent the crystallization temperature of neat PBT. The dashed lines in the bottom graphs at 260°C indicate the processing temperature.
The additive crystallization process for the compounds based on 1,3,5-triaminobenzene is exemplarily shown for 21 in Figure 32. At 210°C the additive forms finely distributed structures that are only faintly visible in polarized optical microscopy. Although 21 is present at a higher concentration compared to 3 in Figure 30 the onset for the additive crystallization is considerably lower than for the corresponding derivative based on a 1,3,5-benzenetricarboxylic acid core.

![Polarized optical micrographs](image)

**Figure 32.** Polarized optical micrographs of PBT comprising 0.8 wt% of 21 cooled at 10 K/min at different stages of the cooling process.
Figure 33 compares the nucleation properties of the trisamides 28 – 30 and 32 based on the 2,4,6-trimethyl-1,3,5-trisaminobenzene core. This core was initially selected with the intention to reduce the solubility in PBT, by the addition of three additional methyl groups. This is the case if compound 28 is compared to compound 1 and 19. Compound 28 with additional core substitution in 2,4,6-position is indeed only soluble in concentrations up to 0.05 wt% and nucleation is observed down to a concentration of 0.0125 wt%. In the concentration range from 0.1 – 0.8 wt% the additive is partially soluble and nucleates PBT with a $T_{c,p}$ of 194.8°C at 0.8 wt%. In contrast 1 and 19 are soluble almost over the entire investigated concentration range. Surprisingly, compounds 29 and 30 are soluble, where the before discussed structural analogues are much less soluble and showed the best nucleation properties. Surprisingly compounds 7 and 23 with cyclohexyl substituents are soluble over the entire concentration range compound 32 based on the 2,4,6-trimethyl-1,3,5-trisaminobenzene core is only soluble in the range from 0.006 – 0.1 wt%. The nucleation properties are very similar to compound 28.

The above discussion on structure property relations with respect to the nucleation clearly shows that a prediction based on just the chemical structure is not possible. It is not the individual molecule, rather the structure of the supramolecular assembly, which determines the nucleation properties.
Figure 33. Crystallization temperatures of PBT ($T_{c,p}$) containing the 1,3,5-benzentrisamide derivatives 28, 29, 30 and 32 (top graphs) and additive dissolution ($T_{d,a}$ ▲) and crystallization temperatures ($T_{c,a}$ ▼)(bottom graphs) as function of the additive concentration. The dashed lines at 188.5°C in the top graphs represent the crystallization temperature of neat PBT. The dashed lines in the bottom graphs at 260°C indicate the processing temperature.
The formation of supramolecular structures is exemplarily shown for 28 in Figure 34. Although 28 is only soluble at high temperatures above 300°C, first additive structures become visible at considerably low temperatures of 235°C.

Figure 34. Polarized optical micrographs of PBT comprising 0.2 wt% of 28 cooled at 10 K/min at different stages of the cooling process.
4.2.3. Nucleation efficiency

To determine the nucleation efficiency of the 1,3,5-benzenetrisamides in PBT, self-seeding experiments as established by Lotz et al.[6] for polypropylene were performed. This involves the partial melting of the polymer at a temperature $T_s$ between the maximum of the endothermic melting peak and its offset, where upon cooling the remaining crystal fragments act as perfect nuclei for the crystallization of PBT and increase the crystallization temperature. The value for the highest theoretical polymer crystallization temperature ($T_{c,p\,theo}$) is obtained at the exothermic peak minimum upon cooling from $T_s$. The increased polymer crystallization temperature by the addition of a nucleating agent is described as $T_{c,p\,nucl}$.

The nucleation efficiency scale is calculated by:

$$NE\ (%) = 100(\Delta T_{c,p}/\Delta T_{c,p\,max}) = 100(T_{c,p\,nucl} - T_{c,p\,neat})/(T_{c,p\,theo} - T_{c,p\,neat})$$

The crystallization exotherms of neat melt processed PBT for various $T_s$ values are shown in Figure 35. Maximum self nucleation of PBT was induced after melting for 5 min at 229.0°C yielding a maximum polymer crystallization temperature $T_{c,p\,theo}$ of 211.0°C.

![Figure 35. Crystallization exotherms of neat melt processed PBT for different $T_s$ values.](image-url)
In the following the efficiencies of the best additives from the three additive series will be compared with the reference compound talc. As presented in Figure 36 talc shows remarkable nucleation behavior over the entire concentration range with a maximum crystallization temperature of 199.7°C at 0.8 wt% (NE = 49.8 %). In comparison, compound 1 (NE = 42.2 %, $T_{c,p\text{ nucl}} = 198.0^\circ\text{C at 0.8 wt%}$), compound 3 (NE = 45.3 %, $T_{c,p\text{ nucl}} = 198.7^\circ\text{C at 0.4 wt%}$), and compound 21 (NE = 47.1 %, $T_{c,p\text{ nucl}} = 199.1^\circ\text{C at 0.8 wt%}$) were found to be the most efficient nucleating agents from the investigated 1,3,5-benzenetrisamide derivatives. Even though 3 and 21 corresponds with the nucleation efficiency of the reference compound at the same concentration, benzenetrisamide derivatives only considerably affect the crystallization of the polymer when present at high additive contents.
Figure 36. Crystallization temperatures of PBT ($T_{cp}$, ●) comprising talcum (top) and additive dissolution ($T_{da}$ ▲) and crystallization temperatures ($T_{ca}$ ▼)(bottom) as function of the additive concentration in comparison to the three most efficient 1,3,5-benzenetrisamide nucleating agents 1, 3, and 21. The dashed line at 188.5°C in the top graphs represents the crystallization temperature of neat PBT and the line in the bottom graphs at 260°C indicates the processing temperature.
4.2.4. Morphology of 1,3,5-benzenetrisamides

To obtain more insight in the morphology and size of the formed supramolecular nanostructures which are formed upon cooling within PBT, experiments were conducted to hydrolyze PBT. It was possible to selectively hydrolyze the PBT matrix in alkaline solution without affecting the formed supramolecular structures. Due to the nonpolar character of the surface and in particular to the fibrillar structure, hydrolysis can only occur at both ends. Therefore the self assembled trisamides are very resistant to hydrolysis. Samples for the morphological studies were prepared by melting compositions containing 0.1 wt%, 0.2 wt% and 0.4 wt% of the 1,1-dimethyl propyl substituted trisamide 3 in DSC pans under nitrogen at 280°C. To ensure complete dissolution and distribution of the trisamides the samples were held for 5 min at this temperature. The samples were cooled to room temperature with four defined linear cooling rates of 10 K/min, 40 K/min, 80 K/min and 110 K/min. After removal of the solid samples from the DSC pans, the surface was carefully hydrolyzed in alkaline solution. Scanning electron microscopy was used to investigate the fibrillar supramolecular nano-structures. Figure 37 compares SEM micrographs of the formed supramolecular nano-structures and shows the corresponding histograms of the lateral size distribution. The first series of experiments was carried out at a concentration of 0.4 wt%. At a cooling rate of 10 K/min and after hydrolysis very long fibrillar structures, with average lateral dimensions of 142 ± 49 nm were observed. Upon increasing the cooling rate to 40 K/min at the same concentration of 0.4 wt%, a few long objects with lateral dimensions around 200 nm and numerous shorter assemblies between 25 and 65 nm are present. The average lateral dimension is with 72 ± 53 nm by a factor of two smaller. At faster cooling rates of 80 K/min and 110 K/min the fraction of long objects decreases further in favor of shorter and smaller assemblies. The average lateral dimensions are 56 ± 31 nm at 80 K/min and 50 ± 24 nm at 110 K/min.
Figure 37. SEM images of supramolecular structures of 3 at a concentration of 0.4 wt% after hydrolysis of PBT and corresponding histograms of the lateral dimensions. Samples were prepared at different cooling rates of 10 K/min, 40 K/min, 80 K/min and 110 K/min.
In the second series of experiments the concentration of 3 was lowered to 0.2 wt% (Figure 38). At a cooling rate of 10 K/min, compared to the higher concentration of 0.4 wt%, lowering the concentration leads to by a factor of two smaller assemblies with narrower distribution (average lateral dimensions of 70 ± 25 nm). Increasing the cooling rate to 40 K/min at 0.2 wt% results in shorter assemblies with a narrower size distribution, similar to the observations made for 0.4 wt%. Here the average lateral dimensions are 48 ± 12 nm. Further increase in cooling rate to 80 K/min and 110 K/min results in very fine homogeneous nano-objects (sticks) with a very narrow size distribution (37 ± 6 nm). Compared to the cooling rate of 10 K/min the assemblies are by a factor of two smaller. It is interesting to note that the dimensions of the formed supramolecular structures for the highest two cooling rates are similar, featuring almost the same values for the average lateral dimensions.

In conclusion the formation of supramolecular nano-objects in a polymer melt is influenced by cooling rate and additive concentration but can be controlled. Remarkably well-defined supramolecular nano-objects can be formed.
Figure 38. SEM images of supramolecular structures of 3 at a concentration of 0.2 wt% after hydrolysis of PBT and corresponding histograms of the lateral dimensions. Samples were prepared at different cooling rates of 10 K/min, 40 K/min, 80 K/min and 110 K/min.
The third series of experiments compares SEM micrographs of the formed supramolecular nano-structures at a concentration of 0.1 wt% (Figure 39). For all cooling rates very short and pronounced supramolecular nano-objects are observed. The average lateral dimensions are in the range of 40 nm and surprisingly are independent from the applied cooling conditions. Interestingly at a concentration of 0.1 wt% no nucleation of PBT is observed as indicated by the DSC measurements presented in 4.2.2. This indicates that the observed supramolecular structures are either too small to induce the nucleation of PBT or are not present in the PBT melt prior to the crystallization of the polymer.
Figure 39. SEM images of supramolecular structures of 3 at a concentration of 0.1 wt% after hydrolysis of PBT and corresponding histograms of the lateral dimensions. Samples were prepared at different cooling rates of 10 K/min, 40 K/min, 80 K/min and 110 K/min.
4.2.5. Laser transparency

The transparency of a visible object is mainly governed by the amount of transmitted light. Scattering on spherulites in semi-crystalline polymers decreases the portion of specular transmitted light accompanied by an increase in diffuse transmission. This phenomenon is strongly pronounced in PBT that, even in thin parts, appears fully opaque. The use of PBT in applications such as laser transmission welding is therefore only possible to a limited extent. As the transparency of polymers for light in the near infrared range should be dependent and capable of being influenced by its morphology, we investigated the influence of nucleation on the laser transparency of injection molded samples. In the course of this work two different PBT batches from the same polymer grade (Ultradur® B4500) (hereinafter referred to as “batch1” and “batch2”) were used. The reference value for the laser transparency of batch1 was found to be 48.2 % ± 0.4 % and thus considerably higher in comparison to batch2 (41.8 % ± 0.3 %). The origin of the reproducible higher values for batch1 is not due to differences in molecular weight or spherulite size and could not be explained up to now.

The LT values at a wave length of 1064 nm of PBT comprising selected 1,3,5-benzenetrisamides that showed promising nucleation behavior in polarized optical microscopy are presented in Figure 40 and Figure 41. Laser transparency was determined at a concentration of 0.1 wt% and 0.8 wt% on injection molded specimens with a thickness of 1.1 mm. Figure 40 shows the results of the laser transparency measurements of PBT (batch2) comprising 0.1 wt% of additive. While all compounds showed positive nucleation behavior in the screening test none of additives were capable to improve the laser transmittance. A very similar trend was observed regarding batch1. All transmission values are mainly within the range of the reference material (Figure 41). Lower LT values occurring for several additives in both batches can be attributed to scattering on undissolved additive particles.
Figure 40. Laser transparency of PBT batch1 comprising selected 1,3,5-benzenetrisamides at a concentration of 0.1 wt%.

Figure 41. Laser transparency of PBT batch2 comprising selected 1,3,5-benzenetrisamides at a concentration of 0.1 wt%.
Nucleation of polybutylene terephthalate

In view of the excellent solubility of the 1,3,5-benzenetrisamides in the PBT melt, few derivatives were investigated at concentrations as high as 0.8 wt%. In spite of the high amount of additive no improvements in transmittance could be achieved (Figure 42). Taking these results into account, 1,3,5-benzenetrisamides do not, despite their ability to promote the nucleation of PBT, improve the laser transparency of PBT.

Figure 42. Laser transparency of PBT batch2 comprising selected 1,3,5-benzenetrisamides at a concentration of 0.8 wt%.
5. Nucleation and clarification of polyamides

5.1. Introduction

Long before the first synthesis of polyamides, Carothers and his coworkers paved the way for modern polymer science with their studies on polycondensation reactions in 1929.\textsuperscript{[80]} In search of new high melting polymer analogues for aliphatic polyesters, polyamides gained growing interest over the years, leading to the invention of PA66 in 1935 and PA6 in 1938.\textsuperscript{[81]}

\[
\begin{align*}
\text{PA66} & \quad \ quad
The unit cell of the α-modification of PA6 is shown in Figure 43. The molecules are in the fully extended zig-zag conformation and form planar H-bonded sheets. The chains in these planes are orientated antiparallel allowing all hydrogen bonds to be developed perfectly.

Figure 43. Monoclinic unit cell of the α-phase of PA6 in the projection along the c direction (top) and cross section of the chain positions (bottom). Images reprinted from Kohen et al. [82]
In contrast, PA6 crystallized in the monoclinic γ-phase is composed of sheets of parallel chains that deviate from the fully planar conformation. H-bonds in this polymorph can be formed without any strain due to a 30° tilt of the amide with respect to the chain axes.

**Figure 44.** Monoclinic or pseudohexagonal unit cell of the γ-phase of PA6 in the projection along the a direction (top left) and projection along the c direction (top right) and cross section of the chain positions (bottom). Images reprinted from Kohen et al.[82]
Both polymorphs can be converted into each other by treatment with aqueous solutions of \( \text{I}_2 \) and KI (\( \alpha \)-phase into \( \gamma \)-phase) or by annealing at high temperatures (\( \gamma \)-phase into \( \alpha \)-phase). Due to the arrangement of hydrogen bonds the \( \alpha \)-phase is thermally more stable (higher melting temperature and enthalpy) compared to the \( \gamma \)-form. The \( \gamma \)-modification in turn exhibits higher fracture toughness with lower elastic modulus.\[^{85}\] The different phases and their relative amounts can be characterized using wide-angle X-ray diffractometry. Diffractograms of the two main crystal forms of PA6 \( \alpha \) and \( \gamma \) are shown in Figure 45.

![Wide-angle X-ray diffraction pattern of the \( \alpha \) - and \( \gamma \)-phase of PA6, and a mixture of both forms. Image reprinted from Kyotani et al.\[^{86}\].](image)

**Figure 45.** Wide-angle X-ray diffraction pattern of the \( \alpha \) - and \( \gamma \)-phase of PA6, and a mixture of both forms. Image reprinted from Kyotani et al.\[^{86}\].

Nucleating agents for polyamides usually include inorganic particles, organic salts and polymeric nucleating agents, all being insoluble in the polymer melt and hence have to be finely dispersed during processing.\[^{1, 82}\] For example, Hitch et al. reported on a mixture of PA66 including fumed alumina, talc, graphite or calcium fluoride. Calcium fluoride showed remarkable nucleation behavior, but decreased impact strength.\[^{87}\] In contrast, organic salts like zinc phenylphosphinate and zinc phenylphosphonate showed nucleation ability in PA66 bypassing the issue of brittleness.\[^{88}\] Several publications describe polymeric materials as nucleating agents for polyamides. PA4.6 was found to efficiently nucleate both PA6 and PA66.\[^{89}\] Du et al. described the use of maleated poly(ethylene-co-vinyl acetate) as nucleating agent for PA6.\[^{90}\]
5.2. 1,3,5-Benzetrisamides as nucleating agents for polyamides

It is worth mentioning that a large proportion of the research on nucleating agents, besides following some general trends, still mainly relies on trial and error. If lead structures evolve, still a fine-tuning of the molecular structure in detail is necessary to find the best suited nucleating agent. Therefore a screening of a large number of potential candidates with respect to their nucleation ability in the particular polymer which has to be nucleated is indispensible.

In this thesis 1,3,5-benzetrisamide derivatives were investigated for the first time as potential nucleating agents for PA6 and PA66. As polyamides exhibit both hydrophobic aliphatic chain segments as well as hydrophilic amid groups, both can either interact with a polar or nonpolar surface of the nucleating agent. On this account 1,3,5-benzetrisamide derivatives bearing nonpolar and polar peripheral substituents were investigated.

The chemical structures of the 1,3,5-benzetrisamides screened in PA6 and PA66 by polarized optical microscopy are summarized in Table 6. The additives are based on four different cores:

- 1,3,5-Benzetetricarboxylic acid (1-15)
- 1,3,5-Triaminobenzene (16-19)
- 2,4,6-Trimethyl-1,3,5-triaminobenzene (20-24)
- 1,3,5-Triaminotoluene (25-27)

The peripheral substituents include short aliphatic, cycloaliphatic and aromatic moieties as well as polar substituents. The thermal properties of most of the screened 1,3,5-benzetrisamide derivatives have already been discussed in detail in chapter 4.2.1.
Table 6. Chemical structures of the 1,3,5-benzenetrisamides screened in PA6 and PA66, their melting temperatures $T_m$ (DTA) and temperatures at 10 % weight loss $T_{-10\text{ wt\%}}$ (TGA, N$_2$ atmosphere).

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a) subl.: sublimation. b) n.d.: not determined due to hygroscopicity of the compound;
In contrast to the nucleation experiments in polybutylene terephthalate all screened 1,3,5-benzenetrisamides showed in the screening experiments no indication of nucleation of PA6 and PA66.

The typical results of the screening experiments are exemplarily shown for compound 9 in Figure 46. With compound 9 an additive was selected that showed a distinct nucleation ability in PBT. The screening results are illustrated by the micrographs recorded between crossed polarizers at different temperatures for PA6 (left) and PA66 (right). The first micrograph of PA6 at 230°C presents on the right the birefringent undissolved additive in the polymer melt. The black area corresponds to the optical isotropic polymer melt. Upon heating to 270°C, the additive dissolves and diffuses into the surrounding polymer melt. The needles partially disappear. Upon cooling with a rate of 10 K/min to again 230°C the additive crystallizes in the polymer melt into fine needles, starting from the region of the non-dissolved additive. Upon further cooling at 190°C the polymer starts to crystallize independently and not influenced by the supramolecular needles or the additive residuals. This clearly shows that no nucleation of PA6 occurs at the surface of the additive. These screening results demonstrate that even though supramolecular aggregations of the trisamides are present in the polyamide melt, the surface of the 1,3,5-benzenetrisamide derivative exhibit no epitaxial matching with the polymer and consequently no nucleation takes place.

In case of PA66 upon heating from 260°C to 290°C the additive dissolves in the melt and partially disappears. Upon cooling to again 260°C the additive remains in the dissolved state and does not crystallize in the polymer melt. At 240°C first polymer crystallites are formed independent from the dissolved or non-dissolved additive. Consequently no nucleation is induced by the additive.

Compounds 11, 13, and 14 bearing polar carboxylic acid- or amino-groups are too good soluble in the melt and hence did not crystallize from the polymer melt upon cooling. The sodium salt 12 or hydrochloride salt 15 of these derivatives however are nearly insoluble, but also not capable to nucleate PA6 or PA66.

Due to the absence of epitaxial matching with the polymer combined with their excellent solubility in the melt, 1,3,5-benzenetrisamides are an inappropriate class of additives for the nucleation of polyamides.
Figure 46. Optical micrographs from polarized light microscopy of the 1,3,5-benzenetrisamide 9 in PA6 (left) and PA66 (right) (scale bar 100 μm). The samples were heated, kept at 270°C (PA6) and 290°C (PA66) for 5 min, cooled with a rate of 10 K/min and observed at different temperatures. The additive is visible as birefringent structures in the isotropic polyamide melt (black). The non-dissolved additive diffuses into the melt and crystallizes in the case of PA6 upon cooling into fine needles, whereas in the case of PA66 the additive remains dissolved. Compound 9 is not capable to nucleate PA6 and PA66.
5.3. Bisamides as nucleating agents for polyamides

Polyamides tend to crystallize in a planar zig-zag structure as shown in chapter 5.1.\textsuperscript{[82]} Accordingly as potential nucleating agents, molecules that favor a sheet-like self assembly, similar to the schematic structure shown in Figure 47, were tested. These linear compounds consist of a central unit with hydrogen bonding units and are substituted symmetrically with aliphatic moieties.

Based on the work of Mohmeyer et al.\textsuperscript{[55]} on the nucleation of iPP with 1,4-phenylene-bisamides, two symmetrically substituted bisamides $0a$ and $0b$ were evaluated with respect to their nucleation potential in PA6. Both compounds were investigated in a concentration range reaching from 1.0 wt% (10000 ppm) to 0.025 wt% (250 ppm). The chemical structures of the additives and crystallization temperatures of the polymer as function of the additive concentration are shown in Figure 48. The average crystallization temperature of melt processed neat PA6 (10 samples) was at 186.5°C and is indicated by the dashed horizontal lines in Figure 48.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure47.png}
\caption{Schematic structure of linear bisamides consisting of a central unit, two amide linkages, and peripheral substituents (top). The bisamides $0a$ and $0b$ were tested as potential nucleating agents in PA6 (bottom).}
\end{figure}
As revealed in Figure 48 the bisamide derivatives 0a and 0b were not capable to nucleate PA6. Similar to the class of 1,3,5-benzenetrisamides, linear bisamides exhibit excellent solubility in the melt preventing the formation of supramolecular structures upon cooling. Consequently bisamides of this type are inappropriate for nucleating polyamides.

\[ \begin{align*}
\text{0a} & \quad \text{0b}
\end{align*} \]

Figure 48. Crystallization temperatures of PA6 comprising the bisamide derivatives 0a and 0b as function of the additive concentration. The dashed lines indicate the crystallization temperature of neat PA6.

To favor supramolecular aggregation in polar ambience such as a polyamide melt the strength of the intermolecular hydrogen bonds has to be increased. Therefore in the following chapter bisurea derivatives will be discussed.
5.4. Bisureas as nucleating agents and clarifiers for polyamides

It is well known that, in contrast to the weaker amide or urethane moieties urea groups are capable to establish two hydrogen bonds between the hydrogens of the nitrogen atoms and the carbonyl oxygen. This association via so-called bifurcated hydrogen bonds is schematically shown in Figure 49. Due to the strong intermolecular interactions, urea compounds combine excellent aggregation properties with high thermal stability and therefore have the potential to be applied in high melting thermoplastics.

![Figure 49. Schematic illustration of bifurcated hydrogen bonds formed between adjacent urea groups.](https://example.com/fig49)

Low molecular weight compounds based on the bisurea motif have extensively been studied in the literature. Most the work conducted on supramolecular structures through hydrogen bonding interactions of urea groups focused on the formation of organo- and hydrogels. Also the formation of polymeric assemblies has been reported. However, the use of bisurea molecules as supramolecular polymer additives and nucleating agents has not been investigated yet.

Therefore we synthesized a new class of C2 symmetric bisurea derivatives schematically illustrated in Figure 50 for the use as supramolecular nucleating agents in semi-crystalline polyamides.

![Figure 50. Schematic structure of linear bisureas consisting of a central unit, two urea linkages, and peripheral substituents.](https://example.com/fig50)

In the following the influence of the bisurea derivatives on the crystallization behavior of PA6 and additional semi-crystalline polyamide homopolymers and copolymers will be
discussed. To establish structure property relations the structural units of the additives were systematically varied. The compounds were investigated in a concentration range from 1.5 wt% (15000 ppm) to 0.025 wt% (250 ppm) and compared with respect to their nucleation efficiency and their influence on the crystal morphology. Furthermore the influence of nucleation on the optical properties was evaluated.
5.4.1. Synthesis and Characterization

In the frame of this work, C2 symmetric bisurea derivatives were synthesized based on 1,4-substituted trans- and cis-cyclohexane central units. The synthesis can be carried out by two different synthetic routes, either starting from 1,4-diaminocyclohexane or from 1,4-cyclohexane diisocyanate. The central units were reacted to the bisurea derivatives by the addition of the corresponding isocyanates or amines under dry and inert conditions. The synthesized compounds were characterized using $^{1}$H-NMR-spectroscopy, mass spectroscopy, differential scanning calorimetry, and thermo gravimetrical analysis. A summary of the characterization data for each compound is given in the Experimental Section in chapter 8.3 of this thesis.

**Synthesis of trans-1,4-cyclohexyl-bisurea derivatives**

The trans-1,4-cyclohexyl-bisurea derivatives were synthesized by two synthetic routes (Figure 51) employing commercially available starting materials. Route A comprised the reaction of an isocyanate with trans-1,4-diaminocyclohexane in dry THF. The reaction of an amine with trans-1,4-cyclohexane diisocyanate is denoted route B. In this case the cyclohexane core was first dissolved in THF and an equimolar amount of the corresponding substituent was slowly added under heavy stirring.

![Figure 51. Synthetic routes A and B to trans-1,4-cyclohexyl-bisurea derivatives starting from trans-1,4-diaminocyclohexane and trans-1,4-cyclohexane diisocyanate.](image-url)
The purification was carried out by recrystallization from organic solvents. The presence of water should be avoided as the isocyanate group reacts with traces of water to form CO$_2$ and the corresponding amine decreasing the total reaction yield.

The following symmetrically substituted trans-1,4-cyclohexyl-bisurea derivatives were synthesized. As substituents linear, branched aliphatic and cycloalipahtic substituents were employed. For comparison also the phenyl substituted bisurea derivative was synthesized.

![Synthetic route A](attachment:image)

**Synthetic route A:**

<table>
<thead>
<tr>
<th>R =</th>
<th>1a</th>
<th>1b</th>
<th>1c</th>
<th>1m</th>
</tr>
</thead>
<tbody>
<tr>
<td>R =</td>
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<td>1q</td>
<td>1r</td>
<td>1s</td>
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</tbody>
</table>

**Synthetic route B:**

<table>
<thead>
<tr>
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<th>1e</th>
<th>1f</th>
<th>1g</th>
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</thead>
<tbody>
<tr>
<td>R =</td>
<td>1h</td>
<td>1i</td>
<td>1j</td>
<td>1k</td>
</tr>
<tr>
<td>R =</td>
<td>1l</td>
<td>1n</td>
<td>1o</td>
<td></td>
</tr>
</tbody>
</table>
Synthesis of cis-1,4-cyclohexyl-bisurea derivatives

The synthesis of the cis-1,4-cyclohexyl-bisurea derivatives involved the reaction of cis-1,4-diaminocyclohexane with the corresponding substituted isocyanate, (Figure 52) similar to the synthetic route A, described for the trans-1,4-cyclohexyl-bisurea compounds. As cis-1,4-cyclohexane diisocyanate is not commercially available, a similar reaction via route B could not be realized. The purification was carried out similar to the trans-bisureas by recrystallization.

Figure 52. Synthetic route to cis-1,4-cyclohexyl-bisurea derivatives based on cis-1,4-diaminocyclohexane.

The following cis-1,4-cyclohexyl-bisurea derivatives were synthesized:

<table>
<thead>
<tr>
<th>R =</th>
<th>2a</th>
<th>2b</th>
<th>2c</th>
<th>2d</th>
</tr>
</thead>
<tbody>
<tr>
<td>R =</td>
<td>2e</td>
<td>2f</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Synthesis of asymmetrically substituted trans-1,4-cyclohexyl-bisurea derivatives

The synthesis of the asymmetric substituted bisurea compounds was carried out in two steps (Figure 53). The first step involved the mono substitution of trans-1,4-diaminocyclohexane with cyclohexyl isocyanate yielding trans-1-(4-aminocyclohexyl)-3-cyclohexylurea, followed by an addition reaction with the second isocyanate. The synthesis of the mono-substituted compound was the more complicated reaction step. Therefore trans-1,4-diaminocyclohexane was dissolved in dried THF. The mixture was cooled in a cooling bath (isopropyl alcohol/ dry ice) and an equimolar amount of cyclohexyl isocyanate, strongly diluted in THF, was slowly added under heavy stirring. The precipitated white solid was suspended in water and acidified with HCl whereupon trans-1-(4-aminocyclohexyl)-3-
cyclohexylurea hydrochloride dissolves immediately. After filtration, the filtrate was brought to pH>8 by slowly adding NaOH and the trans-1-(4-aminocyclohexyl)-3-cyclohexylurea precipitated and could be used without further purification. The second step involved the reaction of trans-1-(4-aminocyclohexyl)-3-cyclohexylurea with the respective isocyanates in dry NMP similar to the synthesis of the symmetric substituted 4-cyclohexyl-bisurea derivatives. The purification was carried out by precipitating the solution in 1M HCl and recrystallization of the obtained solid from organic solvents.

**Figure 53.** Synthetic route to asymmetric substituted trans-1,4-cyclohexyl-bisurea derivatives based on trans-1,4-diaminocyclohexane.

The following asymmetrically substituted trans-1,4-cyclohexyl-bisurea derivatives were synthesized. As fixed substituent for all asymmetric derivatives, cyclohexane was employed. As variable substituent branched aliphatic and phenyl substituents were employed.

<table>
<thead>
<tr>
<th>R =</th>
<th>3a</th>
<th>3b</th>
<th>3c</th>
<th>3d</th>
</tr>
</thead>
</table>
NMR characterization

The spectroscopic data of all synthesized bisurea compounds is summarized in the experimental section. In the following as an example, the results of the spectroscopic characterization of the trans-bisurea derivative 1b, the cis-bisurea 2d, as well as the asymmetric substituted compound 3d will be discussed in detail.

1,1’-(trans-1,4-cyclohexylene)bis(3-n-propylurea) (1b)

The $^1$H-NMR spectrum of 1,1’-(trans-1,4-cyclohexylene)bis(3-n-propylurea) 1b measured in CDCl$_3$/CF$_3$COOD is shown in Figure 54. The N-H protons of the urea groups are not visible, due to exchange processes with the solvent. Also the signal for trifluoroacetic acid at 10.6 ppm is not presented in the spectrum. With exception of the equatorial and axial Hs of the cyclohexane core all signals could be clearly assigned to the corresponding protons. The signals of the CH-group 2 and methylene groups 3 are shifted towards higher ppm values due to their spatial proximity to the urea groups and can be found at 3.46 ppm and 3.24 ppm, respectively. The methylene group 4 can be assigned to a sextet between the signals of the equatorial and axial protons of cyclohexane core. The methyl group 5 could be assigned to the triplett at 0.96 ppm. Full substitution of trans-1,4-diaminocyclohexane was confirmed by the integration ratio of 2 and 5.

![Figure 54. $^1$H-NMR spectrum of 1,1’-(trans-1,4-cyclohexylene)bis(3-n-propylurea) 1b (CDCl$_3$/CF$_3$COOD).]
As the cis-1,4-cyclohexylene-bisureas are more better than their corresponding trans-derivatives, $^1$H-NMR spectra of these compounds were recorded in DMSO. Figure 55 shows the spectra of compound $2d$. The N-H protons of the urea groups do not exchange with the solvent, thus being visible in the range of 5.50-5.72 ppm. Similar to $1b$ the signals of the equatorial and axial protons of the cyclohexane core are not distinguishable. A duplet for the two methyl groups 6 of the isoproyl substituents can be found at 1.00 ppm. The two CH-groups 2 and 5 are again shifted to higher ppm values due to their proximity to the urea protons but can be clearly distinguished owing to their splitting.

![Figure 55. $^1$H-NMR spectrum of 1,1'-(cis-1,4-cyclohexylene)bis(3-isopropylurea) 2d (DMSO).](image)
1-phenyl-3-[4-(cyclohexylcarbamoylamino)cyclohexyl]urea (3d)

The $^1$H-NMR spectrum of 1-phenyl-3-[4-(cyclohexylcarbamoylamino)cyclohexyl]urea 3d recorded in DMSO is shown in Figure 56. The protons of the aromatic substituent can be assigned to the signals at 6.86, 7.2, and 7.35 ppm. Due to the proximity of the phenyl-substituent to the N-H proton of the urea group 4, the signal is shifted towards higher ppm values and can be found as a singlet at 8.29 ppm. The remaining three urea protons can be clearly distinguished owing to their integrals. The signal for the N-H protons of 5 is splitted into two duplets at 5.59 ppm because of the asymmetry of the bisurea molecule. The remaining protons of the cyclohexane core and substituent can be found as a multiplet in the range of 0.9-1.9 ppm.

Figure 56. $^1$H-NMR spectrum of 1-phenyl-3-[4-(cyclohexylcarbamoylamino)cyclohexyl]urea 3d (DMSO).
5.4.2. Thermal properties

The thermal properties of the bisurea derivatives were simultaneously determined using combined thermogravimetric (TGA) and differential thermal analysis (DTA). With respect to the thermal properties the synthesized bisureas can be categorized into three categories that will be described in detail in the following.

Compounds in category A show a melting transition followed by the evaporation from the liquid phase. In the TGA/DTA diagram a typical melting transition can be clearly detected directly followed by a sharp and continuous weight loss. Complete evaporation is indicated by a weight loss of 100 % upon further heating. Figure 57 shows the TGA/DTA diagram of 2f, which had a melting transition at 239°C. The 10 percent weight loss (T_{10 wt%}) is at 256°C. The endotherm at 150°C corresponds to a solid state phase transition which could be observed by polarized optical microscopy. Similar thermal behavior was only observed for compound 2e (T_m = 252°C; T_{10 wt%} = 266°C).

![Thermogravimetric and simultaneous differential thermal analysis](attachment:figure_57.png)

**Figure 57.** Thermogravimetric (solid line) and simultaneous differential thermal analysis (dashed line) analysis of 2f, measured under nitrogen atmosphere at a heating rate of 10 K/min.
In the second category B, the bisurea derivatives exhibit higher melting temperatures, but melting occurs after initial sublimation. Figure 58 shows the TGA/DTA of 2d. The first endothermic peak of the DTA curve represents the sublimation of the compound, followed by a melting transition at 319°C and subsequent evaporation from the liquid phase. The melting transition was confirmed by the disappearance of birefringent structures of the additive in polarized optical microscopy and is also indicated by a bend in the TGA curve. Similar sublimation and melting behavior was found for 1e, 1f, 1h, 2a, 2b, 2c, and 3c. Their melting temperatures were in the range from 309 (2a) to 379°C (3c).

![Figure 58. Thermogravimetric (solid line) and simultaneous differential thermal analysis (dashed line) analysis of 2d, measured under nitrogen atmosphere at a heating rate of 10 K/min.](image-url)
In category C no melting endotherm, but sublimation of the bisureas can be detected followed by thermal decomposition at elevated temperatures. Thermal decomposition was confirmed by polarized optical microscopy and is indicated by the bend in the TGA curve. Figure 59 shows the TGA/DTA diagram of 1l which starts to sublime at around 250°C and thermal decomposition occurs above 365°C. The endotherm at 210°C corresponds to a solid state phase transition. As sublimation and thermal degradation coincide, a total weight loss of nearly 100% is observed. The same trend was found for the compounds 1a, 1b, 1c, 1d, 1g, 1i, 1j, 1k, 1m, 1n, 1o, 1p, 1q, 1r, 1s, 3a, 3b and 3d.

![Figure 59. Thermogravimetric (solid line) and simultaneous differential thermal analysis (dashed line) analysis of 1l, measured under nitrogen atmosphere at a heating rate of 10 K/min.](image-url)
In order to establish structure property relations a short comparison of the thermal properties of the three bisurea types (symmetric trans-, cis-, and asymmetric trans-bisureas) will be given in the following.

The chemical structures of the symmetric substituted trans-bisureas, their melting temperatures and temperatures at 10 % weight loss are listed in Table 7. The investigated substituents ranged from linear and branched aliphatic to cycloaliphatic moieties. Due to the thermally stable urea linkage, all investigated compounds exhibit excellent stability reflected by high melting and weight loss temperatures. Owing to the formation of strong intermolecular hydrogen bonds, a large number of the investigated trans-bisureas sublime completely without melting, indicated by a weight loss of 100 %. Out of 16 symmetric trans-bisureas, 13 sublimed without melting. In contrast the compounds with more flexible long linear (1e, 1f) or long branched aliphatic substituents (1h), melted at high temperatures. The highest melting transition was observed for 1e with a T<sub>m</sub> of 370°C.
Table 7. Chemical structures of the symmetric substituted trans-bisureas, their melting temperatures $T_m$ (DTA) and temperatures at 10% weight loss $T_{-10\,\text{wt\%}}$ (TGA, N$_2$ atmosphere).

<table>
<thead>
<tr>
<th>Abbr.</th>
<th>1a</th>
<th>1b</th>
<th>1c</th>
<th>1d</th>
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<tbody>
<tr>
<td>Substituent (R)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$T_m$ [°C]</td>
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<td>subl.</td>
<td>subl.</td>
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</tr>
<tr>
<td>$T_{-10,\text{wt%}}$ [°C]</td>
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<td>328</td>
<td>329</td>
<td>306</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Abbr.</th>
<th>1e</th>
<th>1f</th>
<th>1g</th>
<th>1h</th>
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<tbody>
<tr>
<td>Substituent (R)</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$T_m$ [°C]</td>
<td>370</td>
<td>345</td>
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<td>355</td>
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<tr>
<td>$T_{-10,\text{wt%}}$ [°C]</td>
<td>315</td>
<td>278</td>
<td>286</td>
<td>299</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Abbr.</th>
<th>1i</th>
<th>1j</th>
<th>1k</th>
<th>1l</th>
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<tbody>
<tr>
<td>Substituent (R)</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>$T_m$ [°C]</td>
<td>subl.</td>
<td>subl.</td>
<td>subl.</td>
<td>subl.</td>
</tr>
<tr>
<td>$T_{-10,\text{wt%}}$ [°C]</td>
<td>290</td>
<td>312</td>
<td>319</td>
<td>323</td>
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</table>

<table>
<thead>
<tr>
<th>Abbr.</th>
<th>1m</th>
<th>1n</th>
<th>1o</th>
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<tbody>
<tr>
<td>Substituent (R)</td>
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<td></td>
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<tr>
<td>$T_m$ [°C]</td>
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<td>subl.</td>
<td>subl.</td>
<td>subl.</td>
</tr>
<tr>
<td>$T_{-10,\text{wt%}}$ [°C]</td>
<td>317</td>
<td>310</td>
<td>319</td>
<td>303</td>
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</table>

<table>
<thead>
<tr>
<th>Abbr.</th>
<th>1q</th>
<th>1r</th>
<th>1s</th>
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</thead>
<tbody>
<tr>
<td>Substituent (R)</td>
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<td></td>
</tr>
<tr>
<td>$T_m$ [°C]</td>
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<td>subl.</td>
<td>subl.</td>
</tr>
<tr>
<td>$T_{-10,\text{wt%}}$ [°C]</td>
<td>332</td>
<td>309</td>
<td>339</td>
</tr>
</tbody>
</table>

a) subl.: sublimation.
The thermal properties of the symmetric substituted cis-bisureas are shown in Table 8. Due to the cis-configuration of the cyclohexane core, the urea groups are either oriented in axial or equatorial position (chair conformation) or both in axial position (boat conformation). In both conformations the molecules are less symmetric than their trans-analogues, leading to a weakening of the intermolecular hydrogen bonds. Therefore all cis-bisureas undergo a melting transition independent from their substituent. The lowest melting temperatures were found for the cyclohexyl- and phenyl-substituted compounds 2e and 2f with a $T_m$ of 252 and 236°C. All cis-bisureas showed comparably low thermal stability ($T_{-10 \text{ wt\%}}$) between 243 and 266°C, whereas their corresponding trans-derivatives exhibit $T_{-10 \text{ wt\%}}$ between 290 and 339°C.

**Table 8.** Chemical structures of the symmetric substituted cis-bisureas, their melting temperatures $T_m$ (DTA) and temperatures at 10% weight loss $T_{-10 \text{ wt\%}}$ (TGA, N$_2$ atmosphere).

<table>
<thead>
<tr>
<th>Abbr.</th>
<th>2a</th>
<th>2b</th>
<th>2c</th>
<th>2d</th>
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<tbody>
<tr>
<td>Substituent (R)</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$T_m$ [°C]</td>
<td>308</td>
<td>325</td>
<td>309</td>
<td>319</td>
</tr>
<tr>
<td>$T_{-10 \text{ wt%}}$ [°C]</td>
<td>254</td>
<td>243</td>
<td>265</td>
<td>264</td>
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</tbody>
</table>

<table>
<thead>
<tr>
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<th>2f</th>
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</thead>
<tbody>
<tr>
<td>Substituent (R)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$T_m$ [°C]</td>
<td>252</td>
<td>236</td>
</tr>
<tr>
<td>$T_{-10 \text{ wt%}}$ [°C]</td>
<td>266</td>
<td>258</td>
</tr>
</tbody>
</table>

The asymmetric substituted trans-bisurea derivatives 3a-3d showed similar thermal stability as the symmetric substituted derivatives 1a-1s (Table 9). With exception of 3c all bisureas sublimed directly without melting. 3c exhibits the highest melting temperature of all investigated bisureas with 379°C. In all cases $T_{-10 \text{ wt\%}}$ was above 300°C.
**Table 9.** Chemical structures of asymmetric substituted trans-bisureas, their melting temperatures $T_m$ (DTA) and temperatures at 10 % weight loss $T_{-10\text{wt\%}}$ (TGA, $N_2$ atmosphere).

![Chemical structures of asymmetric substituted trans-bisureas](image)

<table>
<thead>
<tr>
<th>Abbr.</th>
<th>3a</th>
<th>3b</th>
<th>3c</th>
<th>3d</th>
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</thead>
<tbody>
<tr>
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<td><img src="image" alt="Structure" /></td>
<td><img src="image" alt="Structure" /></td>
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</tr>
<tr>
<td>$T_m$ [$^\circ$C]</td>
<td>subl. a)</td>
<td>subl.</td>
<td>379</td>
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</tr>
<tr>
<td>$T_{-10\text{wt%}}$ [$^\circ$C]</td>
<td>302</td>
<td>346</td>
<td>308</td>
<td>323</td>
</tr>
</tbody>
</table>

a) subl.: sublimation.
5.4.3. Nucleation and optical properties of PA6

As described before the nucleation ability of a supramolecular polymer additive depends on the individual chemical structure. In the self-assembled structure the surface induces nucleation via epitaxial matching. In addition the additive concentration and the solubility of the additive in the polymer melt has an influence. The optical properties are usually governed by scattering due to differences in refractive index of the crystalline and amorphous phase or anisotropic structures. A reduction in spherulite size by the addition of a nucleating agent usually leads to a certain decreased scattering enhancing the transmission and clarity of a polymeric material to some degree. In some cases special compounds are capable to improve transmission, clarity and in particular haze to a very large extent. These additives are called “clarifiers”.

In the following the influence of the chemical structure on the polymer crystallization temperature and the optical properties of PA6 will be investigated in detail. Structure property relations will be systematically discussed. The additives were investigated in a concentration range from 1.5 wt% (15000 ppm) to 0.025 wt% (250 ppm). The different concentrations were prepared by diluting the initial additive concentration of 1.5 wt% in the melt with a mixture of the initial PA6/additive powder blend and neat PA6, yielding the following dilution series: 1.3 wt%, 1.0 wt%, 0.8 wt%, 0.6 wt%, 0.4 wt%, 0.2 wt%, 0.1 wt%, 0.05 wt%, and 0.025 wt%. Optical properties were measured on injection molded platelets with a thickness of 1.1 mm.

In order to explore possible effects of transamidification or decomposition reactions of the polyamide under processing conditions on the crystallization behavior, a blank “concentration” series with neat PA6 was compounded. As shown in Figure 60 the crystallization temperatures for all ten processing runs are mainly within 1°C. The increased crystallization temperature of the first run can originate from impurities in the miniextruder. Therefore cleaning runs were conducted. The calculated mean value for the $T_{c,p}$ of neat PA6 is 186.5°C ± 0.5°C. The degree of crystallinity of PA6 was determined from the enthalpy of the first melting endotherm and is 25.2 % ± 0.5°C.\[13\]
It is well-known that the processing conditions, such as melt temperature, undercooling and shear stress have a strong influence on the crystallization kinetics of semi-crystalline polymers and particular polyamides.\cite{82, 101} The effect of melt processing, as described in the experimental section, on the crystallization temperature of PA6 is shown in Figure 61. The peak polymer crystallization temperature of the PA6 granulate at 174.1°C is considerably lower compared to the compounded material. The compounded PA6 has a $T_{c,p}$ at 186.5°C. In addition the crystallization exotherm was found to be broader. This result can be attributed to the formation of orientation induced chain alignment in the polyamide melt during processing.\cite{102} This so-called “memory effect” is particularly marked in polyamides due to the formation of H-bonds between the functional amide groups that even in the melt remain fixed to some extent.\cite{82, 102, 103} Consequently, as the crystallization kinetics are very sensitive towards the processing history, all samples have to be prepared under identical processing conditions, also the reference

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure60.png}
\caption{Crystallization temperature of neat PA6 versus the number of processing runs. For each run the samples were compounded for 4 min at 250°C.}
\end{figure}
Figure 61. Differential scanning exotherms of PA6 granulate (black) and melt processed PA6 (red) at a cooling rate of 10 K/min.
In the following, the synthesized bisurea compounds will be compared systematically in sets to reveal structure property relations with respect to the nucleation and optical properties of PA6. These properties will be discussed as function of the additive concentration. The first set compares *symmetrical trans-bisureas with linear aliphatic substituents* that vary in chain length from C2 to C18. The second to the fourth set compares *symmetrical trans-bisureas with branched substituents*. The fifth set discusses *symmetrical trans-bisureas with cycloaliphatic substituents*. The sixth set compares *asymmetrically substituted trans-bisureas* and the seventh set *symmetrically substituted cis-bisurea* derivatives.

5.4.3.1. Symmetrical trans-1,4-cyclohexane bisurea derivatives with linear substituents

In this chapter the nucleation and optical properties of the bisurea derivatives 1a – 1f with linear substituents in PA6 will be discussed. The polymer crystallization temperature ($T_{c,p}$) was obtained from DSC measurements, reported as the temperature at the exothermic minimum upon cooling from the melt. Exemplary DSC cooling traces of neat PA6 and PA6 comprising different concentrations of the n-octyl substituted bisurea 1e are presented in Figure 62. Upon addition of 1e the crystallization temperature of PA6 is distinctly increased. At a high concentration of 1.0 wt% the polymer crystallization temperature is at 192.1°C. At lower additive concentrations the $T_{c,p}$ decreases. At a concentration of 0.1 wt% $T_{c,p}$ is at 191.5°C.

![Differential scanning exotherms of neat PA6 and PA6 comprising different concentrations of 1e as indicated.](image)

*Figure 62.* Differential scanning exotherms of neat PA6 and PA6 comprising different concentrations of 1e as indicated.
The optical properties clarity and haze were determined according to ASTM D-1003 on injection molded platelets with a thickness of 1.1 mm. Figure 63 illustrates the effect of the addition of the propyl-substituted bisurea derivative 1b on the visual appearance of injection molded PA6 platelets. On the left is neat PA6 for comparison with a clarity of 68 % and a haze of 98 %, in the middle PA6 containing 0.2 wt% of 1b with a clarity of 96% and a haze of 79 % and on the right PA6 containing 1.3 wt% of 1b with a clarity of 96 % and a haze of 44 %.

Figure 63. Effect of the addition of 1b on the visual appearance of injection molded PA6 platelets. Left: neat PA6; middle: PA6 comprising 0.2 wt% of 1b; right: PA6 comprising 1.3 wt% of 1b.

The chemical structures of the bisurea derivatives 1a – 1f with linear substituents, their nucleation and optical properties are compared in detail in Figure 64 A and B. The additives are presented with increasing length of the aliphatic moieties. The substituents were varied from ethyl 1a, n-propyl 1b, n-butyl 1c, n-hexyl 1d, n-octyl 1e, to n-octadecyl 1f. The top row of Figure 64 shows the polymer crystallization temperature (T_{c,p}) of PA6 as function of the additive concentration. Remarkably with all trans-bisureas the crystallization temperature of PA6 could be distinctly increased.

For example the T_{c,p} values for PA6 steeply increases by the addition of the ethyl substituted bisurea derivative 1a. At a concentration of 0.02 wt% the crystallization temperature of PA6 is increased by 6.3°C from 186.5°C to 192.8°C. At higher additive concentrations the T_{c,p} further increases to a maximum of 194.6°C at 0.4 wt%. Above 0.4 wt% the polymer crystallization temperature only varies marginally with the amount of additive. A plateau is reached with a mean T_{c,p} value of 194.2°C indicated by the dashed horizontal line in Figure 64. Increasing the chain length from C2 to C18 has a noticeable influence on the nucleating ability of the bisurea derivatives. The small change in the chemical structure of the peripheral substituents from ethyl to n-propyl results in a decrease of the mean T_{c,p} plateau value to 193.5°C. However 1b is capable to efficiently nucleate PA6 at a concentration as low
as 0.02 wt% with a $T_{c,p}$ of 191.9°C. Additive 1c with n-butyl substituents is a less efficient nucleating agent for PA6 indicated by a lower mean $T_{c,p}$ plateau value of 192.5°C. With further increasing the chain length and flexibility of the linear aliphatic substituents the mean plateau values for the polymer crystallization temperature decrease in the series from n-hexyl 1d (192.5°C) over n-octyl 1e (192.3°C) to n-octadecyl 1f (191.5°C).

The optical properties clarity and haze of injection molded samples with a thickness of 1.1 mm are presented in the bottom graphs of Figure 64. In the case of the ethyl substituted derivative 1a the clarity increases dramatically upon addition of already slight amounts of additive. The clarity reaches a plateau at around 97% for concentrations exceeding 0.5 wt%. The change in length and flexibility of the peripheral substituent to n-propyl, n-butyl or n-hexyl does not result in a noticeable change in clarity. The values remain remarkably high. Upon further increasing the length of the substituent to n-octyl or n-octadecyl the clarity slightly decreases. For the n-octyl substituted derivative 1e the values deteriorate from 96.1% to 90.1% with increasing additive concentration which can be attributed to scattering on nondissolved additive particles due to an insufficient mixing during the powder blending and processing steps. For 1f the highest clarity is 92.4 % at a concentration of 1.5 wt%.

The haze values are presented in the bottom row of Figure 64. In the case of iPP in these kind of diagrams usually an “optimum” concentration for minimum haze is observed. At very low concentrations the optical properties deteriorate due to less available nucleation sites. However if the amount of additive exceeds the “optimum” concentration, scattering occurs on undissolved additive particles and the haze increases. Regarding the bisureas investigated in this thesis, a deviant behavior was found. For 1a the values for haze steadily decrease with increasing additive concentration in agreement with the excellent solubility of the bisureas in the polyamide melt. The lowest haze was 47 % at a concentration of 1.5 wt%. Compound 1b with n-propyl substituent displayed an even lower haze with 41 % at 1.5 wt%. If the haze values at 1.0 wt% are compared with increasing length of the peripheral substituents an increase of the haze values is observed. The haze values increase from n-propyl < ethyl < n-butyl < n-hexyl < n-octyl < n-octadecyl. It is interesting to note that although 1e and 1f are still efficient nucleating agents for PA6 the haze remains at a very high value.
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Figure 64A. Polymer crystallization temperatures (\(T_{c,p}\) (top graphs) and the optical properties haze (\(\star\)) and clarity (\(\ast\)) (bottom graphs) of PA6 comprising the trans-bisureas 1a – 1c as function of the additive concentration. The dashed lines indicate the plateau in the polymer crystallization temperature PA6.
Figure 64B. Polymer crystallization temperatures ($T_{c,p}$, ⋆) (top graphs) and the optical properties haze (★) and clarity (●) (bottom graphs) of PA6 comprising the trans-bisureas 1d – 1f as function of the additive concentration. The dashed lines indicate the plateau in the polymer crystallization temperature PA6.
5.4.3.2. Symmetrical trans-1,4-cyclohexane bisurea derivatives with branched substituents

The second set of additives compares longer branched substituents that are structural isomers of the n-octyl derivative 1e described before. The chemical structures and their nucleation and optical properties are presented in Figure 65. For 1g with 6-methylheptane substituents the mean T_{c,p} plateau value was 192.9°C and thus in the range of the n-butyl or the n-hexyl substituted derivatives from the above discussed series of additives. However, the values for the polymer crystallization temperature deteriorate for concentrations exceeding 1.0 wt%, which can be attributed to an insufficient distribution of the additive in the PA6 melt during processing. The change in chemical structure of the peripheral substituent from 6-methylheptane to 1,5-dimethylhexyl substituents 1h does not result in improved nucleation properties. The mean T_{c,p} plateau value for 1h is 192.9°C. For the 2-ethylhexyl substituted derivative 1i the mean polymer crystallization temperature at the plateau is approximately in the same range with 193.3°C. In contrast to the nucleation properties that are only little affected by the subtle changes in chemical structures, the introduction of branches has strong influence on the optical properties of PA6. The less beneficial nucleation ability of 1g at higher additive concentrations is also reflected in lower clarity values. Similar to the n-octyl substituted derivative 1e from the first series of additives, the clarity decreases slightly with the amount of additive due to an insufficient distribution in the polymer matrix. The clarity decreases from 95.1 % at 0.2 wt% to 89.1 % at a concentration of 1.5 wt%. For 1h and 1i high clarity values are obtained that are persistent throughout the investigated concentration range.

For 1g with 6-methylheptane substituent the values for haze steadily decrease with increasing amount of additive. The lowest haze is 73 % at a concentration of 1.5 wt% and thus similar to the values obtained for 1e (74 %) from the first series of additives. It is important to note that branching in proximity to the α-C atoms of the substituents as in 1h and 1i had severe effect on the optical properties of PA6. For 1h the introduction of branches decreases the haze from 98 % of neat PA6 to 22 % at an additive concentration of 1.5 wt%, which corresponds to an overall decrease of 76 %. The introduction of an ethyl group in β-position as in 1i results in a decrease of haze to 44 %, also at 1.5 wt%. Following these results it is likely to assume that branching of the peripheral substituents in close proximity to the urea group, especially in α-position to the nitrogen atom, strongly favors the clarification ability of the bisurea derivatives.
Figure 65. Polymer crystallization temperatures (T\text{c,p}) (top graphs) and the optical properties haze (*) and clarity (•) (bottom graphs) of PA6 comprising the trans-bisureas 1g-1i as function of the additive concentration. The dashed lines indicate the plateau in the polymer crystallization temperature of PA6.
The third set of additives compares four bisurea derivatives with branched peripheral substituents, that were varied from 3-pentyl 1l, 1,2-dimethylpropyl 1m, neopentyl 1n to iso-propyl 1o. The derivatives 1l, 1m and 1n are structural isomers with 5 C-atoms each. All compounds within this series were capable to efficiently promote the nucleation of PA6. For 1l with 3-pentyl substituent the mean $T_{c,p}$ plateau values was about 192.1°C. The 1,2-dimethylpropyl substituted derivative 1m at higher concentrations was less distributed in the polymer melt, thus the $T_{c,p}$ values decrease with the amount of additive. The neopentyl and iso-propyl substituted derivatives displayed the highest mean $T_{c,p}$ plateau values within the series. For 1n the value is 193.1°C and for 1o 192.8°C.

Within this set of additives all compounds displayed clarity values above 90 %. However the lack of distribution in the polymer matrix of 1m is also reflected in the values for clarity. The values vary slightly between 95 and 98 % in the investigated concentration range, whereas the values remained almost constant for 1l, 1n and 1o. One of the most important observations is that the introduction of branches in proximity to the urea groups strongly improves the haze values. For 1l the values for haze steadily decrease with the additive concentration, with 30 % at a concentration of 1.5 wt%. Surprisingly, despite the missing branch in α-position to the nitrogen atom, 1l has the most pronounced effect on the optical properties of PA6 within this series of additives. This result is even more interesting as 1l shows lower nucleation ability compared to 1l, 1n and 1o. This once more demonstrates that a direct correlation between the nucleation efficiency and clarifying of an additive cannot be drawn. Compound 1m with 1,2-dimethylpropyl substituent displayed similar good optical properties with a haze as low as 42 %. For 1n with neopentyl substituent the haze is higher with 57 % at 1.5 wt%, whereas 1o featured a haze value at 1.5 wt% of around 36%.

To further corroborate the observation that the introduction of branches in close proximity to the urea groups are responsible for the good clarification ability of the bisureas, derivatives with highly branched substituents, preferably in α-position to the urea groups are discussed in the following.
Figure 66. Polymer crystallization temperatures \( T_{c,p} \) (top graphs) and the optical properties haze (★) and clarity (●) (bottom graphs) of PA6 comprising the trans-bisureas 1l-1o as function of the additive concentration. The dashed lines indicate the plateau in the polymer crystallization temperature of PA6.
The chemical structures, the nucleation and optical properties of the bisurea derivatives 1j, 1k and 1p with quaternary α-C atoms, are presented in Figure 67. The additives are arranged with decreasing length of the peripheral substituents varied from tert-octyl 1j, over 1,1-dimethylpropyl 1k, to tert-butyl 1p. All three compounds significantly increase the polymer crystallization temperatures of PA6. For 1j the mean polymer crystallization temperature plateau value was found to be the lowest within this series of additives with 192.3°C. The one $T_{c,p}$ value at 1.3 wt% was distinctly higher than for the remaining concentrations, which seems to be an artifact with the DSC measurement. Compound 1k with 1,1-dimethylpropyl substituent features the highest plateau in the polymer crystallization temperature at 194.3°C. This is the highest mean $T_{c,p}$ value found in this thesis. It is also interesting to note that the highest $T_{c,p}$ plateau value with 194.5°C was found for 1k at a comparatively low concentration of 0.2 wt%. The tert-butyl substituted derivative 1p shows similar nucleation ability with a mean $T_{c,p}$ plateau value of 193.7°C. While 1k and 1p displayed excellent nucleation behavior, the polymer crystallization temperatures of PA6 comprising 1j were considerably lower, most likely due to the increased length of the peripheral substituent which slightly changes the epitaxy.

The clarity values for 1j were found to be relatively low. They decrease also with the additive concentration. Remarkably 1k and 1p exhibit clarity values around 97 % even at the lowest investigated concentration of 0.02 wt%. It is important to note that despite its comparably low nucleation ability, 1j caused a distinct reduction in haze. The lowest value is 26 % at 0.8 wt%. This again demonstrates that a direct correlation between the nucleation and clarifying ability of an additive cannot be drawn. It is interesting to note that for 1j and 1p the haze reaches a plateau above 0.8 wt% where the values remain constant upon further addition of additive. For 1j the plateau is around 28 % and for 1p 19 %. In contrast the haze for 1k steadily decreases with the additive concentration as seen for most of the additives. As already expected from the previous series of additives these findings support strongly the assumption that an efficient clarifying agent for PA6 has to bear highly branched peripheral substituents, particular in the α-position.
Figure 67. Polymer crystallization temperatures ($T_{c,p}$ *) (top graphs) and the optical properties haze (*) and clarity (**) (bottom graphs) of PA6 comprising the trans-bisures 1j, 1k, and 1p as function of the additive concentration. The dashed lines indicate the plateau in the polymer crystallization temperature of PA6.
Figure 68 illustrates the optical effect of the addition of 1p on injection molded platelets of PA6. Already at a very low additive concentration of 0.05 wt% high clarity values of 97% could be achieved. For concentrations exceeding 0.8 wt% the haze reaches a plateau whereupon further addition of 1p the optical properties remain unchanged. Here haze values of around 20% and clarity values above 95% were obtained.

Figure 68. Emblem of the University of Bayreuth viewed through injection molded platelets of PA6 (thickness 1.1 mm) containing 0.0, 0.05 and 1.0 wt% of 1p (top).
5.4.3.3. Symmetrical trans-bisurea derivatives with cyclic substituents

In an additional series of additives, bisurea derivatives with cycloaliphatic substituents were investigated. The peripheral substituents comprise 1-adamantyl \(1s\), cyclohexyl \(1q\) and phenyl \(1r\) moieties. The chemical structures of the investigated compounds \(1s-1r\), their nucleation and optical properties are presented in Figure 69. Within this series of additives all bisurea derivatives are efficient nucleating agents for PA6. The mean \(T_{c,p}\) plateau value of \(1s\) is 192.7°C. However the crystallization temperature scatters strongly, why the assignment of a proper plateau is complicated. This can be attributed to a worse solubility in the PA6 melt. The compounds \(1q\) and \(1r\) exhibit similar plateau values at around 192.8°C.

As for all trans-bisureas the additives within this series exhibit high clarity values. For \(1s\) the values are above 95 % even at the lowest investigated concentration of 0.02 wt%. For \(1q\) the increase in clarity is less marked compared to \(1s\). Here the plateau in clarity is reached for concentrations above 0.1 wt%. This effect is even more pronounced with \(1r\). Regarding the haze of PA6 \(1s\) was found to be the most efficient clarifier with a remarkable low haze of 16 % at 1.5 wt%. Regarding the shape of the haze curve it can be assumed that further addition of \(1s\) leads to an even higher transparency. However, similar to the 1,1-dimethylpropyl substituted derivative \(1k\) from the above series of additives, the haze strongly decreases with the additive concentration due to the excellent solubility in the polymer melt. The introduction of the less branched cyclohexyl \(1q\) substituent yields somewhat poorer haze with 64 % at 1.5 wt%. For \(1r\) the lowest haze values is at 51 %, also at 1.5 wt%.
Figure 69. Polymer crystallization temperatures ($T_{c,p}$) (top graphs) and the optical properties haze (•) and clarity (•) (bottom graphs) of PA6 comprising the trans-bisureas 1s, 1q, and 1r as function of the additive concentration. The dashed lines indicate the plateau in the polymer crystallization temperature of PA6.
5.4.3.4. Asymmetrically substituted trans-bisurea derivatives

In addition to the above discussed compounds, the asymmetrically substituted trans-bisureas 3a-3d were synthesized and the properties in PA6 will be discussed in the following. All derivatives exhibit one cyclohexyl substituent and differ concerning the second peripheral substituent. The chemical structure of the asymmetric bisureas with tert-octyl 3a, iso-propyl 3b, tert-octyl 3c and phenyl substituent, their nucleation and optical properties are presented in Figure 70.

In accordance with the symmetrically substituted bisureas, all derivatives in this series were capable to significantly increase the crystallization temperature of PA6. The mean $T_{c,p}$ plateau values of the asymmetric derivative 3a is 192.6°C and thus almost similar compared to the corresponding symmetric derivative 1j (192.3°C) with two tert-butyl substituents. For 3b and 3d also similar mean $T_{c,p}$ plateau values are found. In contrast the nucleation ability of 3c is considerably lower compared to the corresponding symmetrical derivative 1p with tert-butyl substituents. Here the $T_{c,p}$ plateau values scatter strongly why a plateau value can only be estimated.

The clarity was for all asymmetric compounds above 95 % and no distinctions between the different substituents can be made. All derivatives significantly decreased the values for haze, however this effect was less pronounced in comparison to the corresponding symmetric bisureas. For example 3a displayed minimum haze values of 42 % at 1.5 wt% whereas the corresponding symmetric analogues 1j featured values as low as 28 % at this concentration. This effect is likely to originate from defects in the self assembly of the bisureas due to the asymmetric structure of the additives. Despite the only minor nucleation ability 3c is the most efficient clarifier investigated in this series. The lowest haze value is 31 % at 1.5 wt%.

It is again interesting to note that a substitution of the α-hydrogen atoms with two methyl groups resulted in a stronger decrease in haze, as already discussed before.
Figure 70. Polymer crystallization temperatures ($T_{c,p}$) (top graphs) and the optical properties haze (★) and clarity (♦) (bottom graphs) of PA6 comprising the trans-bisureas 3a-3d as function of the additive concentration. The dashed lines indicate the plateau in the polymer crystallization temperature of PA6.
5.4.3.5. Symmetrically substituted cis-bisurea derivatives

The chemical structures of the cis-bisurea derivatives $2a - 2f$ and their nucleation and optical properties are compared in detail in Figure 71A and B. The substituents are varied from n-octyl $2a$, tert-octyl $2b$, isopropyl $2c$, tert-butyl $2d$, cyclohexyl $2e$, to phenyl $2f$. The conformational change in the central unit from the trans- to the cis-bisureas has dramatic influence on the self-assembly and nucleation ability.

In comparison to the trans-analogues all derivatives are less efficient as nucleating agents. For $2a$ with n-octyl substituents the mean $T_{c,p}$ plateau value is $190.1^\circ C$ and thus considerably lower than for the corresponding trans-bisurea $1e$ ($192.3^\circ C$). For compound $2b$ the crystallization temperature steadily increases with the additive concentration and no real plateau can be assigned. With $2b$ the highest $T_{c,p}$ is $190.1^\circ C$ at 1.5 wt% and thus $2.8^\circ C$ lower compared to the trans-bisurea $1j$ with the same substituent ($192.9^\circ C$). The isopropyl substituted derivative $2c$ is not capable to nucleate PA6 and no increase in $T_{c,p}$ is observed, whereas the trans analogue $1o$ has a $T_{c,p}$ plateau value of $192.8^\circ C$. The compounds $2d$ with tert-butyl substituents and $2e$ with cyclohexyl substituents are again capable to promote nucleation of PA6. The mean $T_{c,p}$ plateau values are again distinctly lower compared to the trans-bisurea analogues $1p$ and $1q$. The phenyl substituted derivative $2f$ in contrast does not nucleate PA6.

It is interesting to note that for $2a$ the values for clarity seem to deteriorate at lower additive concentrations compared to neat PA6. However clarity values obtained from specimens with a haze of close to 100 % are prone to errors and thus may be inaccurate. Similar behavior is observed for $2c$, $2e$, and $2f$, all compounds with a haze of close to 100 %. The small nucleation ability of the cis-bisureas is also reflected in the values for haze that are distinctly higher compared to the investigated trans-bisureas. Despite the nucleation potential of $2a$ no improvements in optical properties can be observed and the haze remains around the value of neat PA6 (98%) for all concentrations. Similar behavior is found for $2c$, $2e$, and $2f$.

The tert-octyl substituted derivative $2b$ was only capable to improve the haze by 5 % from 98 % to 93 % at 1.5 wt%. For $2d$ the haze is decreased by 14 % to 84 % at 1.5 wt%. In this connection it is again apparent that derivatives with highly branched substituents are the most beneficial substituents to improve the properties of PA6.
Figure 71A. Polymer crystallization temperatures ($T_{c,p}$) (top graphs) and the optical properties haze (⋆) and clarity (●) (bottom graphs) of PA6 comprising the cis-bisureas 2a-2c as function of the additive concentration. The dashed lines indicate the plateau in the polymer crystallization temperature of PA6.
Figure 71B. Polymer crystallization temperatures ($T_{c,p}$) (top graphs) and the optical properties haze ($\times$) and clarity ($\circ$) (bottom graphs) of PA6 comprising the cis-bisureas 2d-2e as function of the additive concentration. The dashed lines indicate the plateau in the polymer crystallization temperature of PA6.
5.4.4. Nucleation efficiency and optical properties in comparison to talc

To determine the nucleation efficiency of the bisurea derivatives, self-seeding experiments as described by Lotz et al.\textsuperscript{[6]} for polypropylene were conducted. The self seeding procedure has already been described in chapter 4.2.3 on polybutylene terephthalate. Figure 72 shows the crystallization exotherms of melt processed neat PA6 for various self nucleation temperatures ($T_s$). Maximum self nucleation temperature of PA6 was induced after melting for 5 min at 220.5\degree C yielding a maximum achievable polymer crystallization temperature $T_{c,p} \text{theo}$ of 195.4\degree C.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure72.png}
\caption{Crystallization exotherms of melt processed neat PA6 for different self nucleation temperatures.}
\end{figure}

In the following the nucleation behavior and optical properties of the most efficient bisureas will be compared to the literature known nucleating agent talc. In this study Micro-Talc IT Extra from Mondo Minerals was used.\textsuperscript{[104]} Figure 73 compares the results obtained for the crystallization temperature and the optical properties haze and clarity for talc in comparison to the most efficient clarifiers 1p and 1k discussed before. As already observed in the case of PBT, talc exhibits an excellent nucleation behavior in PA6. The best polymer crystallization temperature was 194.8\degree C at a concentration of 1.5 wt\%. This corresponds to a nucleation
efficiency of 93.3 %. The values observed for talc are only slightly higher than those for the two most efficient trans-bisurea nucleating agents 1a with ethyl ($T_{c,p} = 194.6°C$, NE = 90.0 %, at 0.4 wt%) and 1k with 1,1-dimethylpropyl substituent ($T_{c,p} = 194.5°C$, NE = 88.9 %, at 0.2 wt%). It is important to note that the required concentration to reach the $T_{c,p}$ plateau is for the bisurea derivatives lower compared to talc. As the theoretical polymer crystallization temperature ($T_{c,p \text{ theo}}$) of PA6 is only located 9°C above the value for the equilibrium crystallization temperature, slight changes in $T_{c,p}$ strongly affect the calculated nucleating efficiencies. Besides the discussed compounds 1a and 1k, the n-propyl substituted derivative 1b with a nucleation efficiency of 84.4 % at a concentration of 1.5 wt% and the tert-butyl substituted derivative 1p with a nucleation efficiency of 83.3 % at a concentration of 1.0 wt% are the most efficient nucleating agents found in this thesis.

Regarding the optical properties the comparison with talc is different and very much in favor for the bisureas. In contrast to the 1,1-dimethylpropyl substituted trans-bisurea 1k and the tert-butyl substituted trans-bisurea 1p, talc improves the optical properties of PA6 only with respect to the clarity. The values for clarity show a maximum at 0.2 wt% with 93 %. With increasing additive concentration the clarity is reduced again. The haze values are only reduced to about 85 % at the highest concentration of 1.5 wt%. The comparably poor haze can be attributed to the insufficient distribution of the insoluble talcum particles in the polymer melt during processing and consequently increased scattering of light. This demonstrates once more the advantage of supramolecular polymer additives compared to insoluble polymer additives particular with respect to the optical properties.
Figure 73. Polymer crystallization temperatures ($T_{\text{cr}}$) (top graphs) and the optical properties haze (●) and clarity (●) (bottom graphs) of PA6 comprising the bisurea derivative 1p, 1k and talc as function of the additive concentration. The dashed lines indicate the plateau in the polymer crystallization temperature of PA6.
5.4.5. Stability of nucleation effect

In the literature the nucleation properties of an additive are not only determined by the increase in polymer crystallization temperature but also its stability.\textsuperscript{105, 26} Therefore several heating and cooling cycles are performed and the polymer crystallization temperatures are studied. In this thesis the nucleation stability was investigated by multiscanning DSC experiments according to Mai et. al.\textsuperscript{26} PA6 samples comprising two selected bisureas were repeatedly heated and cooled in the DSC. Samples were heated up to 250°C and held for 5 min before each cooling run to prevent self nucleation on remained non-molten crystal fragments.

![Diagram of PA6 comprising 0.4 wt% of 1r for 13 heating and cooling cycles in a multiscanning DSC experiment.](image_url)

**Figure 74.** Cooling curves of PA6 comprising 0.4 wt% of 1r for 13 heating and cooling cycles in a multiscanning DSC experiment. Samples were held at 250°C for 5 min before each cooling run.
Figure 75. Cooling curves of PA6 comprising 0.4 wt% of 2a for 13 heating and cooling cycles in a multiscanning DSC experiment. Samples were held at 250°C for 5 min before each cooling run.

The nucleation efficiencies were calculated from the polymer crystallization temperatures of the polymer for each cooling run and are shown in Figure 76. The crystallization temperature of PA6 comprising 1r is nearly unaffected by the number of measuring cycles confirming the reversibility of the supramolecular self assembly process and demonstrating that no degradation occurred. The $T_{c,p}$ of neat PA6 is increased by the additive from 186.5°C to 191.9°C (first run). After 13 runs the $T_{c,p}$ decreases from 191.9°C to 191.1°C. This is a decrease of only 0.8°C. Despite its lower thermal stability the nucleation ability of the cis-derivative 2a exhibits similar behavior (Figure 75). From the first DSC run to the 13th run the $T_{c,p}$ decreases from 190.1°C to 189.4°C. This corresponds to a decrease of only 0.7°C.

This becomes more visible if the nucleation efficiency depending on the number of measuring cycles is plotted (see Figure 76). Both bisureas are nearly unaffected by the number of heating and cooling scans if taken into account the thermal stress after 1 h in the melt. As already mentioned, slight changes in $T_{c,p}$ strongly affect the calculated nucleating efficiencies due to the small temperature range between $T_{c,p}$ and the theoretical crystallization temperature of PA6. After 13 cycles the values for nucleation efficiency
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decreased by only 9% for 1r and 8% for 2a. In contrast sorbitol derivatives such as bis(3,4-dimethylbenzylidene)sorbitol (Millard 3988) only exhibit less thermal stability even in iPP of the heterogeneous nucleation, the latter losing most of its nucleation ability already after the 5th scan in a similar experiment.\[^{26}\]

![Figure 76. Nucleation efficiencies of 1p (○) and 2a (●) in PA6 as function of the number of measuring cycles in a multiscanning DSC experiment.](image)
5.4.6. Influence of the mold temperature on optical properties

The macroscopic properties of semi-crystalline polymers are strongly dependent on the spherulite size and can also be influenced by the processing conditions. As it is evident from Figure 77 that upon fast cooling from the melt a higher nuclei density occurs and smaller spherulites are formed. That in turn leads to less scattering increasing the total transmission of visible light. When a polymer is cooled rather slowly, nucleation only occurs sporadically and fast crystal growth of individual particles leads to larger spherulites.\cite{1}

![Figure 77. Schematic representation of the rate of nucleation and the rate of crystal growth as function of the temperature.\cite{1}](image)

The influence of the cooling rate on the optical properties of PA6 was investigated at different mold temperatures during injection molding. The mold temperatures were varied from 40°C to 120°C. The haze values of the injection-molded specimens (1.1 mm thickness) are presented in the top graph of Figure 78. The open stars represent the values for neat PA6 and the solid squares the haze values for PA6 comprising 0.8 wt% of the tert-butyl substituted trans-bisurea 1p. As expected the haze value decreases as faster cooling is applied, both for neat and clarified PA6. At a mold temperature of 40°C the haze values are about 12% lower compared to the specimens injected in a mold at 120°C. Over the whole investigated temperature range 1p acts as efficient clarifying agent and distinctly improves the haze compared to the neat PA6. Upon faster cooling, the bisureas self assemble into supramolecular nano-objects in the polyamide melt capable to nucleate and clarify. The
values for clarity are presented in the bottom graph of Figure 78. Increasing the mold temperature from 40°C to 120°C resulted for neat PA6 in a strong decrease in clarity from 90% to only 29.2%. Regarding the clarified samples no improvements were observed upon faster cooling.

Figure 78. Values for haze (top graph) and clarity (bottom graph) for neat PA6 (solid squares) and PA6 comprising 0.8 wt% of the tert-butyl substituted trans-bisurea 1p (open squares) as function of the mold temperature.
5.4.7. Influence of sample thickness on optical properties

Variations in sample thickness have an impact on the morphology of injection molded specimens. Fewer but larger spherulites are formed where the melt cools slowly, whereas faster cooling leads to more but small crystalline entities.\textsuperscript{[106]} As a result of the temperature gradient from the outside to the inside of the specimens the spherulite size increases from the skin to the core and is on average larger for higher mold thicknesses.\textsuperscript{[107]} As the transparency of objects is mainly determined by the amount of transmitted light, values for haze increase with thickness. The influence of the mold thickness on haze of PA6 comprising the 1,1-dimethylpropyl substituted bisurea 1k, the tert-butyl substituted bisurea 1p and talc is compared in Figure 79. As usual the haze values increase with the sample thickness.

As discussed in chapter 5.4.4 talc caused only a very modest reduction in haze. At a thickness of 0.5 mm a plateau in haze was reached at around 55 % for concentrations above 0.1 wt%. For 1.1 mm the haze values steadily decrease with higher additive concentrations to a value of only 85 % at the highest concentration of 1.5 wt%. For specimens thicker than 1.1 mm the haze values remained at 100 % independent of the additive concentration.

In contrast both bisurea derivatives 1k and 1p were capable to enhance the optical properties of PA6 for all thicknesses. For a mold thickness of 0.5 mm both additives showed minimum haze values of around 8 % at a concentration of 1.3 wt%. In thicker samples, further addition of 1k resulted in a steady decrease in the haze values, whereas for 1p a plateau was reached for 1.1 mm and 2.0 mm samples. Thus it can be assumed an increase of the 1k concentration would even lead to lower haze values. In platelets with a thickness of 1.1 and 2.0 mm, 1p with tert-butyl substituent was found to be the more efficient clarifier for PA6 up to a concentration of 0.8 wt%. This trend is reversed for concentrations exceeding 1.0 wt%. Here 1k featured lowest haze values of 11 % at 1.1 mm and 21 % at 2.0 mm, whereas 1p displayed values at 19 % and 30 % respectively. At a mold thickness of 3.0 mm the lowest haze achieved is 44 % for 1p (1.5 wt%) and 53 % for 1k (1.3 wt%). Remarkably even for very thick samples (5.0 mm), the lowest haze value obtained with 1k was 77 % (1.5 wt%) whereas 1p caused only a modest reduction in haze (93 %, 1.5 wt%).
Figure 79. Haze values of PA6 comprising A: talc, B: the tert-butyl substituted bisurea 1p and C: the 1,1-dimethylpropyl substituted bisurea 1k for different mold thicknesses (0.5 mm, ●; 1.1 mm, ■; 2.0 mm, ▲; 3.0 mm, ★ and 5.0 mm, ○) as function of the additive concentration.
5.4.8. Laser transparency

Laser transmittance of PA6 comprising the tert-butyl substituted trans-bisurea 1p and talc was determined at a wavelength of 1064 nm on injection molded plaques. The investigated thickness is varied from 1.1 mm, 2.0 mm, to 3.0 mm. In Figure 80 the values for laser transparency are plotted as function of the additive concentration. The LT values for neat PA6 were found to be 82 % for a thickness of 1.1 mm, 72 % for a thickness of 2.0 mm and 59 % for a thickness of 3.0 mm.

Similar to the values for haze presented in chapter 5.4.3.2 one would expect steady improvements in laser transparency with increasing amounts of additive. Surprisingly for all thicknesses at low concentrations in the range from 0.02 – 0.1 wt% the LT-values are lower compared to the neat material. For a thickness of 1.1 mm at a concentration of 0.02 wt% the LT-value is only 78 % and thus 4 % lower compared to the neat PA6. At concentrations exceeding 0.2 wt% 1p is capable to enhance the laser transparency of PA6. Here the transmittance is increased by 3 % to 86 % for a thickness of 1.1 mm at a concentration of 1.5 wt%. A very similar trend is observed for the 2.0 mm and 3.0 mm thick specimens. The lowest LT-values were 63% for a thickness of 2.0 mm at 0.02 wt% and thus 9 % lower compared to the neat PA6. For a thickness of 3.0 mm at an additive concentration of 0.02 wt% the LT-value is 53 %, 6 % lower as the neat polymer. Similar to 1.1 mm above 0.2 wt% the LT increases. The increase for 2.0 mm is 10 % to a LT-value of 82 % at a concentration of 1 wt%. For 3.0 mm the LT increases by 13 % to 72 % at a concentration of 0.8 wt%. These results demonstrate that 1p is capable to distinctly increase the LT in PA6 especially in thick parts.

The different shapes of the curves for haze and laser transparency at low additive concentrations can be explained by the different measurement procedures for haze and laser transmittance. Haze is determined using an integrating sphere that collects the total hemispheric transmittance when the sample is placed directly on the entrance port of the sphere. Values for haze are obtained by dividing the diffuse by the total transmittance.[59] In contrast, laser transparency is determined by a sensor located 1.1 mm below the polymer specimen and calculated as the ratio of the reference and the measurement beam.[66] While haze is calculated with regard to the total transmittance the former can improve while the total transmittance decreases compared to the neat material.
While **talc** caused only modest improvements in haze as already shown in chapter 5.4.4 laser transmittance is reduced throughout the whole investigated concentration range and for all mold thicknesses. It is interesting to note that the transmittance values reach a plateau above an additive concentration of 0.1 wt%. For a thickness of 1.1 mm the LT-value decreases by 2 % to values of 80 %. For a thickness of 2.0 mm the LT-plateau value is at 65 % and 12 % lower than the neat PA6. For 3.0 mm the plateau value is at 50 % and 9 % lower than neat PA6. The origin of the poor laser transparency of **talc** is irregular and due to insufficient distribution in the polyamide melt causing additional scattering of the irradiating laser.

![Figure 80. Laser transparency values of PA6 comprising 1p (left) and talc (right) for different mold thicknesses (1 mm, •; 2 mm, ▲ and 3 mm, ◆) as function of the additive concentration.](image-url)
5.4.9. Morphology of injection molded samples

In the following the influence of nucleation on the spherulitic morphology is investigated on injection molded samples of a 1.1 mm thickness. A comparison of neat PA6 and PA6 comprising the tert-butyl substituted trans-bisurea 1p and the reference nucleating agent talc will be made. Figure 81 and Figure 82 present thin sections cut parallel to the flow direction and investigated in an optical microscope between crossed polarizers. The images were merged from two separate micrographs in order to present the complete cross section of the injection molded parts. The haze values for the respective concentrations are reported in the top left of each image.

Polyamides tend to crystallize rather fast exhibiting a usually relatively fine and homogeneous spherulitic morphology. This effect is even amplified in injection molding due to the extremely fast cooling rates. A determination of concrete values for the crystallite size is at this level not possible. Figure 81 presents in the top left micrograph the spherulite structure of neat PA6. Resulting from the temperature gradient from the outside to the inside of the specimens during solidification the specimens are composed of highly oriented skin layers and a less oriented spherulitic core region. The black stripe in the middle of the specimens is found for all samples and is a consequence of the injection molding process. The spherulitic morphology of PA6 was significantly influenced by the addition of 1.0 wt% of 1p. Here single crystallites are so small to be resolved separately in polarized optical microscopy. The haze in this sample is reduced from 98 % of neat PA6 to 20 %. The thin stripes oriented parallel to the cutting direction originate from the cutting process. Upon lowering the concentration of 1p from 1.0 wt% to 0.2 wt% the crystallite size remains relatively unaffected. The haze value increases to 47 %. Still at a concentration as low as 0.02 wt% the morphology of the injection molded specimens is significantly affected. However, the spherulites are slightly larger compared to a concentration of 0.2 wt%. As expected, the modifications in morphology are also accompanied by drastic changes regarding the optical properties. Scattering is reduced with decreasing spherulite size thus lowering the values for haze even at very low concentrations.
Figure 81. Thin sections of injection molded samples (thickness 1.1 mm) parallel to the flow direction and the corresponding haze values of neat PA6 (top left) and PA6 comprising 1p at different concentrations. The images presented consist of two separate pictures taken between crossed polarizers.
**Talc** displays a remarkable nucleation ability in PA6, exceeding even slightly the polymer crystallization temperature values of the most efficient bisurea compounds. The effect of **talc** on the spherulitic morphology in injection molded specimens is considerably different. In contrast to the tert-butyl substituted bisurea 1p nucleation with **talc** has less influence on the crystallite size. At a concentration 1.0 wt% and 0.6 wt% the spherulitic morphology is less homogeneous compared to 1p. Here agglomerates of **talcum** particles are visible in optical microscopy that originate from the not ideal distribution of **talc** in the PA6 matrix. At a concentration of 0.2 wt% less agglomerated particles are visible in optical microscopy. However larger spherulites were formed compared to 1p at the same concentration. The less homogeneous morphology and the larger spherulites are also reflected in the haze values of PA6. As already described in chapter 5.4.4 the values for haze are considerably lower compared to 1p, which is a clarifier for PA6.
Figure 82. Thin sections of injection molded samples (thickness 1.1 mm) parallel to the flow direction and the corresponding haze values of PA6 comprising talc at different concentrations. The images presented consist of two separate pictures taken between crossed polarizers.
5.4.10. Crystal modification of PA6

PA6 crystallizes, depending on the processing conditions, into three main polymorphs, the α-, γ- and the metastable β-phase, with the α- and γ-form being the most prominent modifications. Both polymorphs differ concerning their mechanical properties as well as their thermal properties and can be detected by DSC measurements. Figure 83 shows the DSC diagram of a PA6 sample comprising 0.1 wt% talc revealing a mixture of both the α and γ form. In the heating scan two melting transitions are observed for the γ-phase at 218°C and for the α-phase at 223°C.

![DSC Diagram of PA6](image)

*Figure 83. DSC diagram of PA6 comprising 0.1 wt% of talc.*

Wide-angle X-ray diffractometry is a more accurate way to characterize the different phases and their relative amounts. The most difficult step in determining the phase content from a WAXD scan is the separation of the amorphous halo from the crystalline peaks. Other than for PE or iPP, PA6 does not show well pronounced crystalline peaks, making it more difficult to resolve the diffraction pattern especially for poorly crystallized samples. In order to bypass these difficulties a method described by Murthy et. al. using amorphous templates was used. The use of an amorphous film as template material must be discarded due to post-crystallization of the polyamide at ambient conditions. Alternatively the amorphous template was obtained by subtracting the crystalline peaks from a diffraction scan of a highly crystalline sample, enriched in α-phase. Furthermore the starting values for the positions of the crystalline diffraction peaks as well as their full width at half maximum
Nucleation and clarification of polyamides

(FWHM) were estimated from enriched patterns. The so obtained values were used for profile fitting of the injection molded specimens using Origin8G software. To simplify the fitting routine all profiles were considered to be Gaussian.\textsuperscript{[109]}

In order to separate the contribution of the crystalline peaks from the diffraction pattern of PA6, a highly crystalline sample enriched in the α-phase was obtained by heating an injection molded specimen in water at 180°C for 3h. The so obtained diffraction pattern and the corresponding deconvolution curves are shown in Figure 84. The relevant peak positions are labeled with the designation of the lattice planes in brackets. The profile parameters are summarized in Table 10. From the integrals of the deconvolution curves the percentage of the α- and γ-form and the degree of crystallinity can be calculated using the following formulae:

\begin{align}
CI_{\alpha}(\%) &= \frac{\sum A_{\alpha\text{-form}}}{\sum (A_{\alpha\text{-form}} + A_{\gamma\text{-form}} + A_{\delta\text{-form}})} \\
CI_{\gamma}(\%) &= \frac{\sum A_{\gamma\text{-form}}}{\sum (A_{\alpha\text{-form}} + A_{\gamma\text{-form}} + A_{\delta\text{-form}})} \\
CI(\%) &= CI_{\alpha} + CI_{\gamma}
\end{align}

with $A_{\alpha\text{-form}}$ and $A_{\gamma\text{-form}}$ being the area under the crystalline peaks of the α- and γ-modification, respectively and $A_{\delta\text{-form}}$ being the area under the amorphous halo.
Figure 84. Diffractometer scan and corresponding profile analysis of a highly crystalline PA6 sample enriched in the α-phase.

Table 10. Profile parameters for PA6.

<table>
<thead>
<tr>
<th></th>
<th>α₁ (200)</th>
<th>α₂ (002+202)</th>
<th>γ₁ (001)</th>
<th>δ (amorphous)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak position</td>
<td>19.99</td>
<td>24.19</td>
<td>21.30</td>
<td>22.11</td>
</tr>
<tr>
<td>FWHM [°]</td>
<td>0.61</td>
<td>0.91</td>
<td>0.77</td>
<td>5.63</td>
</tr>
<tr>
<td>Area</td>
<td>5054</td>
<td>7948</td>
<td>290</td>
<td>8241</td>
</tr>
</tbody>
</table>

The value for the FWHM of the amorphous halo was used in the following to analyze the crystal morphology of the injection molded specimens. As proposed by Murthy et. al.\(^{[67]}\) the full width of δ was allowed to vary in a small range of 0.5°. The diffraction scan of neat PA6 is shown in Figure 85. Due to the only less pronounced crystalline peaks an exact profile analysis for the neat polymer could not be drawn. However it appears clearly that PA6 crystallizes predominantly in the γ-phase under the given processing conditions.
The effect of nucleation with bisurea derivatives on the crystal structure of PA6 is exemplarily shown for different concentrations of 1r in Figure 86. The so obtained crystal phase composition is summarized in Table 11. The results of the XRD patterns significantly differ from the neat polymer. Remarkably the addition of 1r had significant influence on the phase content of PA6. In contrast to the neat polymer, nucleation predominantly promotes the formation of α-phase crystals under equal processing conditions. An increase in the fractions of the γ-polymorph was detected at lower additive concentrations, however the values associated with the γ-form are still distinctly lower compared to the neat polymer Figure 85.
Figure 86. Diffractometer scans and corresponding profile analysis of PA6 comprising different concentrations of 1r.

Table 11. Content of the different crystal phases of PA6 comprising different concentrations of 1r obtained from profile analysis shown in Figure 86.

<table>
<thead>
<tr>
<th>Concentration [wt%]</th>
<th>Clα [%]</th>
<th>Clγ [%]</th>
<th>Cl_total [%]</th>
<th>Clδ [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.0 wt%</td>
<td>24.1</td>
<td>0.8</td>
<td>24.9</td>
<td>75.1</td>
</tr>
<tr>
<td>0.6 wt%</td>
<td>25.6</td>
<td>1.4</td>
<td>27.0</td>
<td>73.0</td>
</tr>
<tr>
<td>0.2 wt%</td>
<td>25.4</td>
<td>1.8</td>
<td>27.2</td>
<td>72.8</td>
</tr>
<tr>
<td>0.02 wt%</td>
<td>21.6</td>
<td>5.6</td>
<td>27.2</td>
<td>72.8</td>
</tr>
</tbody>
</table>

Additionally, all bisureas from the three additive series (symmetric trans-, symmetric cis- and asymmetric trans-bisureas derivatives) were investigated regarding their influence on the crystal modification of PA6 at a concentration of 0.2 wt%. The diffraction patterns for eight selected symmetrically substituted trans-bisureas and two asymmetrically substituted trans-bisureas are summarized in Figure 87 arranged with increasing haze values from bottom to top. Consistent with the observations for 1r all trans-bisureas promoted the
formation of α-phase crystals, even at low concentrations and can thus be regarded as efficient α-phase nucleators for PA6. A correlation between the chemical structure of the trans-additives and the ability to induce the α-polymorph in PA6 could not be drawn as the diffraction patterns only differ slightly. The same applies for the optical characteristics. As example for 1f and 1r haze values above 90 % are observed at a concentration of 0.2 wt%. For 1p the haze is 47 % at the same concentration. However the α-phase content of the three additives is relatively the same. In conclusion fractions of the γ-form cannot solely account for the higher haze values observed for several additives.

Figure 87. Diffractometer scans of neat PA6 and PA6 comprising different symmetric trans-bisurea additives at 0.2 wt% arranged from top to bottom with decreasing haze values. The haze values for plaques of 1.1 mm are presented close to the left margin.
The lower nucleation ability of the cis-bisurea derivatives is also reflected in the crystal phase composition of PA6. Only the compounds 2d, 2b and 2e were capable to promote the formation of a larger amount of the α-polymorph. However, in spite of the modest increase in α-phase content, the values for haze have not changed. With 2a, 2c and 2f the diffraction patterns resemble those of neat PA6, with mainly γ-crystals formed.

Figure 88. Diffractometer scans of neat PA6 and PA6 comprising different symmetric cis-bisureas and the corresponding haze values, arranged with increasing γ-phase content from bottom to top. The haze values for plaques of 1.1 mm are presented close to the left margin.
5.4.11. Nucleation and optical properties of PA66 and PA12 homopolymers and polyamide copolymers

In the following the influence of the most efficient bisurea derivative 1p on the crystallization behavior and the optical properties of PA66, PA12 and polyamide copolymers is discussed. The additive concentration was varied from 1.5 wt% (15000 ppm) to 0.025 wt% (250 ppm) in the following semi-crystalline polyamides:

- Polyamide 66 (PA66), Ultramid® A27 E 01
- Copolyamide 6/66 (PA6/66), Ultramid® C33
- Copolyamide 66/6 (PA66/6), Ultramid® 9A E 01
- Copolyamide 6/12 (PA6/12), Zytel® 1512
- Polyamide 12 (PA12), Vestamid® L2101F

Self-seeding experiments to determine the nucleation efficiency as described for PA6 in 5.4.4 were performed for each polyamide. Figure 89 to Figure 93 show the crystallization exotherms of the above polyamides for various $T_s$ values.

\[ \text{Figure 89. Crystallization exotherms of neat PA66 for different self nucleation temperatures ($T_s$).} \]

$T_{c,p}$ theo is at 243.5°C indicated by the dashed red vertical line. The $T_{c,p}$ of the neat resin heated above the equilibrium melting temperature is at 234.2°C indicated by the dashed blue vertical line.
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**Figure 90.** Crystallization exotherms of neat PA6/66 for different self nucleation temperatures \(T_s\). \(T_{c,p\text{,theo}}\) is at 175.3°C indicated by the dashed red vertical line. The \(T_{c,p}\) of the neat resin heated above the equilibrium melting temperature is at 157.5°C indicated by the dashed blue vertical line.

**Figure 91.** Crystallization exotherms of neat PA66/6 for different self nucleation temperatures \(T_s\). \(T_{c,p\text{,theo}}\) is at 223.3°C indicated by the dashed red vertical line. The \(T_{c,p}\) of the neat resin heated above the equilibrium melting temperature is at 210.7°C indicated by the dashed blue vertical line.
**Figure 92.** Crystallization exotherms of neat PA6/12 for different self nucleation temperatures ($T_s$). $T_{c,p}$ is at 207.0°C indicated by the dashed red vertical line. The $T_{c,p}$ of the neat resin heated above the equilibrium melting temperature is at 188.8°C indicated by the dashed blue vertical line.

**Figure 93.** Crystallization exotherms of neat PA12 for different self nucleation temperatures ($T_s$). $T_{c,p}$ is at 147.5°C indicated by the dashed red vertical line. The $T_{c,p}$ of the neat resin heated above the equilibrium melting temperature is at 161.3°C indicated by the dashed blue vertical line.
The self nucleation temperature ($T_s$), the polymer crystallization temperature ($T_{c,p}$), the maximum polymer crystallization temperature ($T_{c,p \, \text{theo}}$) and the maximum crystallization temperature gain ($\Delta T_{c,p}$) for each polymer are summarized in Table 12.

**Table 12.** Self-nucleation temperature ($T_s$), polymer crystallization temperature ($T_{c,p}$), maximum polymer crystallization temperature ($T_{c,p \, \text{theo}}$) and maximum polymer crystallization temperature gain ($\Delta T_{c,p}$) for each polymer.

<table>
<thead>
<tr>
<th>Polymer</th>
<th>$T_s$ [°C]</th>
<th>$T_{c,p}$ [°C]</th>
<th>$T_{c,p , \text{theo}}$ [°C]</th>
<th>$\Delta T_{c,p}$ [°C]</th>
</tr>
</thead>
<tbody>
<tr>
<td>PA66</td>
<td>261.0</td>
<td>234.2</td>
<td>243.5</td>
<td>9.3</td>
</tr>
<tr>
<td>PA6/66</td>
<td>196.5</td>
<td>157.5</td>
<td>175.3</td>
<td>17.8</td>
</tr>
<tr>
<td>PA66/6</td>
<td>241.0</td>
<td>210.7</td>
<td>223.3</td>
<td>12.6</td>
</tr>
<tr>
<td>PA6/12</td>
<td>217.5</td>
<td>188.8</td>
<td>207.0</td>
<td>18.2</td>
</tr>
<tr>
<td>PA12</td>
<td>175.0</td>
<td>147.5</td>
<td>161.3</td>
<td>13.8</td>
</tr>
</tbody>
</table>

PA12 and the copolyamides PA6/12 and PA6/66 exhibit multiple crystallization exotherms in the self-seeding experiments as both polymers crystallize into separate polymorphs showing two melting transitions upon heating. An exact determination of the self-nucleation temperature $T_s$ is therefore complicated. Heating above the second melting endotherm yields to an only minor raise in polymer crystallization temperature as the first phase is in the fully molten state. However, if $T_s$ is selected too low the crystals of the second phase do not melt. Thus for the determination of the maximum theoretical polymer crystallization temperature, the polymer has to be carefully heated to a temperature $T_s$ located between the first and second melting endotherm. The values for $T_{c,p \, \text{theo}}$ of these two polymers were determined at the exothermic minimum of the first peak occurring upon cooling from $T_s$. Considering this, the highest values for the temperature gain are obtained for PA6/66 and PA6/12 with 17.8°C and 18.2°C respectively. In contrast, due to its very fast crystal growth rates, the lowest $\Delta T_{c,p}$ values were found for PA66, making this polymer much more difficult to nucleate.\(^{[19]}\)
The concentration dependence of the nucleation properties (top graphs) and the optical properties (bottom graphs) of 1p in the polyamide copolymers PA6/66, PA66/6 and PA6/12 are presented in Figure 94. Consistent with the observation for PA6 in chapter 5.4.3.2, 1p shows remarkable nucleation behavior in the investigated copolyamides. For PA6/66 (Ultramid® C33) the T_{c,p} values steeply increase by the addition of 1p. At a concentration of as low as 0.02 wt% the polymer crystallization temperature increases by 7.1°C to 164.6°C. At higher amounts of additive the T_{c,p} values further increase to a maximum of 173.9°C at 1.3 wt%. This corresponds to a nucleation efficiency of 92 %. The leap in T_{c,p} from 1.0 to 1.3 wt% can be explained by the formation of two exothermic crystallization peaks in the DSC measurements. For the two highest concentrations an additional peak at higher temperatures evolves and was used to determine the T_{c,p} values. For PA66/6 (Ultramid® 9A) the T_{c,p} also increases with increasing additive concentration. Above 0.4 wt% the polymer crystallization temperature only varies marginally with the additive concentration. A plateau with a mean T_{c,p} value of 219.2 °C is reached, what corresponds to a nucleation efficiency of 67 %. For PA6/12 (Zytel® 1512) the mean T_{c,p} plateau value is at 192.1°C and also a pronounced nucleation effect is observed. It is interesting to note that the plateau is already reached at concentrations above 0.1 wt%. Here the nucleation efficiency is considerably low with only 20 %.

In addition to the crystallization kinetics, 1p has strong influence on the optical properties of injection molded specimens. For PA6/66 the clarity values remain at the very high initial values of around 96 %. For PA66/6 clarity increases from 60 % to over 95 % with very low additive amounts. For PA6/12 the clarity values increase from 78% of the neat resin to also over 95 %. The values remain at these high values for all investigated concentrations. The values for haze steadily decrease with further addition of 1p demonstrating the excellent solubility of the additive at the processing conditions. For PA6/66 the haze decreases to values as low as 6 % for a concentration of 1.3 wt%. For the neat resin no haze value could be obtained, due to the limited processability of neat PA6/66. For PA66/6 the haze values decrease by 67 % from 90 % of the neat polymer to 23 % at a concentration of 1.0 wt%. For the copolymer PA6/12 the haze values decease from 75 % to 37 % at 0.6 wt%. For higher amounts of additive the haze increases again slightly, most likely to insufficient solubility or
liquid phase separation in the polymer melt at higher concentrations. This phenomenon was also observed for iPP with trisamide additives.\textsuperscript{[6, 110]}

As expected at lower additive concentrations the crystallization temperatures were found to decrease for all polymers, approaching the values for the corresponding neat melt processed polyamides. For PA66/6 and PA6/12 at high amounts of additive a plateau was reached where further addition of 1\textsuperscript{p} showed no influence on the $T_c$. Interestingly, for PA6/66 a sharp increase in crystallization temperature of around 5°C was observed for concentrations exceeding 1.0 wt\%. The highest nucleation efficiencies were found for PA6/66 with 92 % at a concentration of 1.3 wt\%. For this polymer, 1\textsuperscript{p} represents an almost ideal nucleating agent. For PA66/6 and PA 6/12 the highest efficiencies were found to be 74 % at 0.8 wt\% and 20 % at 1.3 wt\% respectively.

In addition to the crystallization kinetics, 1\textsuperscript{p} had significant influence on the optical properties of injection molded specimens (Figure 94, bottom graphs). The values for haze steadily decreased with further addition of 1\textsuperscript{p} demonstrating the excellent solubility of the additive at the processing conditions. Values for clarity maintained on a very high level, even at the lowest additive concentrations. However, the results obtained for PA6/66 are rather different. Here the optical properties deteriorate compared to the neat reference material for concentrations below 0.2 wt\%. 


The nucleation behavior of the tert-butyl substituted bisurea 1p in PA66 (Ultramid® A27) and PA12 (Vestamid® L2101F) is shown in Figure 95. Despite the higher processing temperatures, 1p is capable to considerably promote the nucleation of PA66 without degradation. Derivative 1p increases the $T_{c,p}$ value from 234.2°C of neat PA66 to 240.8°C at 1.3 wt%. This corresponds to a nucleation efficiency of 71%. However, the NE values rapidly decrease with less amount of additive most likely due to the enhanced solubility at the higher temperatures. At a concentration of 0.02 wt% the $T_{c,p}$ is only slightly increased to 235.7°C.

For PA12 again a plateau was reached at concentrations above 0.4 wt%. The mean $T_{c,p}$ plateau value is at 154.0°C. This is a nucleation efficiency of 47%. It is interesting to note that for PA6/12 and PA12 the lowest nucleation efficiencies of 1p in the investigated
polymers were obtained. This indicates that the length of the repeating unit is essential in determining whether the additive is capable to significantly promote the nucleation of the polymer. The bisureas in this study are mainly of advantage if PA6 repeating units are present.

Nucleation with 1p also has an influence on the optical properties of PA66 and PA12. In case of PA66 the clarity increases dramatically upon addition of already small amounts of additive. The clarity reaches a plateau for concentrations above 0.2 wt% at 95%. For PA12 the clarity values remain at the initially high values of 95%. Despite the excellent nucleation behavior in PA66, 1p has only minor effects on haze at first glance. However, considering the complete opaque appearance of neat PA66 with a haze of 100% the values obtained for haze (52%) at 1.3 wt% are not as bad as one might initially believe. Even though 1p is only capable to nucleate PA12 with comparably low efficiencies, the optical properties of the specimens are still improved. The lowest haze value is 27% at 1.5 wt% and thus 43% lower than for neat PA12. It is interesting to note that the values for haze reach a plateau for concentrations above 0.4 wt%, which most likely seems to be a result of the minor solubility in the polymer matrix due to the lower processing temperature of PA12. In conclusion, the tert-butyl substituted trans-bisurea 1p is capable to effectively nucleate and clarify PA6. In addition a transfer was possible to further semi-crystalline polyamides. Here, the additive is most beneficial in polyamides with PA6 repeating units.
Figure 95. Crystallization temperatures ($T_{cp}$ ⋆) (top) and the optical properties haze (★) and clarity (∗) (bottom) of PA66 and PA12 comprising 1p as function of the additive concentration.
6. Summary

The use of additives to enhance the processability and end properties is one of the main measures to enlarge the property profile of commercial thermoplastic polymers. The physicochemical properties of semi crystalline polymers depend to a large extend on the crystal structure and spherulite size, which can be controlled by nucleating agents. These additives can increase the crystallization temperature and thus reduce cycle times during melt processing, affect the physical properties and can in some cases improve the optical properties (clarity and haze). This thesis describes the use of supramolecular nucleating agents to increase the polymer crystallization behavior and to improve the optical properties of polybutylene terephthalate and polyamides.

In the first chapter the influence of the molecular structure of 1,3,5-benzenetrisamides on the nucleation behavior of polybutylene terephthalate was investigated. In order to explore structure property relations, 43 derivatives based on the 1,3,5-benzenetricarboxylic acid core, 1,3,5-triamino benzene core, 2,4,6-trimethyl-1,3,5-benzenetrisamide core and 1,3,5-trisaminotoluene core were screened with regard to their nucleation ability of PBT. For promising compounds the additive dissolution and crystallization behavior in the polymer melt and crystallization temperature of the polymer were investigated in the concentration range from 0.8 wt% down to 0.006 wt%. It was revealed that certain 1,3,5-benzenetrisamides are capable to nucleate PBT. The 1,3,5-benzenetrisamides feature outstanding high temperature stability and chemical resistance to be processed under the commonly applied conditions for PBT. They can act as supramolecular polymer additives, are soluble in the PBT melt under the processing conditions and upon cooling self-assemble into fine fibrillar supramolecular nano-objects inducing the nucleation of PBT. The investigated 1,3,5-benzenetrisamide additives are much more soluble in PBT compared to the same additives in isotactic polypropylene and polyvinylidene fluoride. It was found that particularly the trisamide derivatives based on the 1,3,5-triaminobenzene core are much better soluble than the derivatives based on the 1,3,5-benzenetricarboxylic acid core and 2,4,6-trimethyl-1,3,5-trisaminobenzene core. As a result additives featuring too good solubility do not self-assemble prior to the polymer crystallization and hence do not induce nucleation.
Summary

This demonstrates that the nucleation efficiency strongly depends on the individual chemical structure of the additives and slight structural changes have large impact. However a prediction of the nucleation ability based on the molecular structure is not possible. We found that the most efficient additive, namely 1,3,5-tris(2,2-dimethylbutyrylamino)benzene, was capable to increase the crystallization temperature of PBT by 10.6°C from 188.5°C to 199.1°C. In addition for the first time it was possible to visualize the supramolecular structures within the PBT matrix by selectively hydrolyzing PBT. The morphology of the formed supramolecular nano-objects is determined by cooling rate and additive concentration. Faster cooling and lower additive concentrations leads to the formation of needle-like structures with smaller lateral dimensions and a narrower size distribution. In case of the concentration of 0.2 wt% and a cooling rate of 110 K/min supramolecular nano-objects with a diameter of 37 ± 6 nm were observed. An attempt to increase the laser transparency of PBT by nucleation with 1,3,5-benzenetrisamides was not successful.

In the second chapter a new class of supramolecular nucleating agents on the structural motif of 1,4-cyclohexane bisureas was synthesized specifically for semi-crystalline polyamides. These compounds combine excellent aggregation behavior with high thermal stability, due to the strong intermolecular interactions between the urea groups. The bisureas are temperature stable enough to be applied in high melting polyamides. In total, 29 derivatives were synthesized and investigated with respect to their nucleation and clarification behavior in PA6. The chemical structure of the additives was systematically varied by changing the conformation of the central unit and the length and degree of branching within the peripheral substituents. Bisureas based on the trans-conformation of 1,4-cyclohexane are capable to efficiently nucleate the α-form of PA6 even at concentrations below 200 ppm, with extremely high nucleation efficiencies of around 90 %. In addition, selected derivatives were found to distinctly improve the optical properties haze and clarity, as well as the laser transparency, in injection molded samples. This is the first time semi-crystalline polyamides were clarified. The most efficient clarifiers for PA6, namely 1,1''-(trans-1,4-cyclohexylene)bis(3-tert-butylurea) and 1,1''-(trans-1,4-cyclohexylene)bis(1,1-dimethylpropylurea) were found to decrease the values for haze from nearly 100 % (1.1 mm thick samples) for neat PA6 to 19 % and 11 % at a concentration of 1.5 wt%. Cis-bisureas exhibit limited nucleation and no clarification ability. By varying the peripheral substituents of the trans-bisureas interesting structure-property relations could be revealed. The
investigated derivatives with longer flexible substituents were efficient nucleating agents for PA6, but displayed only modest improvements in optical properties. Short, highly branched substituents, preferably in α-position to the urea groups, were found to be a key segment for the clarification of PA6. Additionally we demonstrated that the process conditions, namely cooling rate and mold thickness, strongly determine the final optical properties. The use of trans-1,4-cyclohexane bisureas could also be expanded to other semi-crystalline polyamide homo- and copolymers. Here, \(1,1'-(\text{trans-1,4-cyclohexylene})\text{bis}(3\text{-tert-butylurea})\) was capable to efficiently nucleate PA66, PA12, PA6/66, PA66/6 and PA6/12. For PA66/6 and PA12 haze values of about 30 % were observed. The effect on the optical properties was more pronounced if PA6 repeating units are present. For PA6/66 haze values below 10 % could be achieved. For PA66 and PA6/12 the haze was between 40 and 50 %.

For the first time the concept of supramolecular nucleating and clarifying agents was successfully transferred to semi-crystalline polyamides. With the new class of compounds it could be demonstrated that trans-1,4-cyclohexane bisurea derivatives are capable to increase the crystallization behavior and improve the optical properties, particularly the haze values, remarkably.
7. Zusammenfassung


Im ersten Kapitel wurde der Einfluss der molekularen Struktur von 1,3,5-Benzoltrisamiden auf die Nukleierung von Polybutylenterephthalat untersucht. Um Struktur-Eigenschaftsbeziehungen aufzuzeigen wurden 43 Derivate basierend auf 1,3,5-Benzoltricarbonsäure, 1,3,5-Triaminobenzol, 2,4,6-Trimethyl-1,3,5-triaminobenzol und 1,3,5-Triaminotoluol hinsichtlich ihres Vermögens PBT zu nukleieren gescreent. Die Löslichkeits- sowie die Kristallisationstemperatur ausgewählter Additive in der Polymerschmelze und die Kristallisationstemperatur des Polymers wurden in einem Konzentrationsbereich von 0.8 bis zu 0.006 Gew.-% untersucht. Es wurde gefunden, dass bestimmte 1,3,5-Benzoltrisamide in der Lage sind PBT zu nukleieren. Diese Verbindungen zeigen außergewöhnliche thermische Stabilität und Chemikalienbeständigkeit und können somit bei den für PBT üblichen Bedingungen verarbeitet werden. Ausgewählte Verbindungen sind in der PBT Schmelze löslich und kristallisieren beim Abkühlen bereits über der Kristallisationstemperatur des Polymers zu homogen verteilten Nanoobjekten aus, welche die Nukleierung des Polymers induzieren. Die untersuchten 1,3,5-Benzoltrisamide sind in der PBT Schmelze löslicher verglichen mit denselben Verbindungen in isotaktischem Polypropylen oder Polyvinylidenfluorid. Insbesondere die Derivate basierend auf dem 1,3,5-Triaminobenzolkern zeigten eine erhöhte Löslichkeit. Demzufolge findet der Selbstorganisationsprozess bei zu gut löslichen Verbindungen nicht vor der Kristallisation des Polymers statt, wodurch keine Nukleierung induziert wird.
Zusammenfassung


Im zweiten Kapitel wurde eine neue Klasse von supramolekularen Nukleierungsmitteln basierend auf 1,4-Cyclohexan-Bisharnstoffen, zur Verwendung in teilkristallinen Polyamiden, synthetisiert. Aufgrund der starken intermolekularen Wechselwirkungen zwischen den Harnstoffgruppen zeigt diese Verbindungsklasse ein exzellentes Aggregationsverhalten und hohe thermische Stabilität, weshalb sie auch in hochschmelzenden Polyamiden eingesetzt werden können. Insgesamt wurden 29 Derivate synthetisiert und hinsichtlich der Nukleierung und Klarmodifizierung von PA6 untersucht. Die chemische Struktur der Additive wurde systematisch durch Veränderung der Konformation der zentralen Einheit und der Länge und dem Verzweigungsgrad der peripheren Substituenten, variiert. Bisharnstoffe basierend auf trans-1,4-Cyclohexan waren in der Lage die α-Form von PA6 bei sehr niedrigen Konzentrationen von 200 ppm mit extrem hohen Nukleierungseffizienzen von 90 % zu nukleieren. Ausgewählte Verbindungen verbesserten die optischen Eigenschaften Haze und Clarity, sowie die Lasertransparenz von spritzgegossenen Probekörpern deutlich. Somit war es zum ersten Mal gelungen die Transparenz von teilkristallinen Polyamiden zu verbessern. Mit 1,1'-(trans-1,4-Cyclohexan)bis(3-tert-butylurea) und 1,1'-(trans-1,4-Cyclohexan)bis(1,1-di-methylpropylurea)
wurden zwei Verbindungen gefunden welche den Haze von PA6 von fast 100 % (1.1 mm Probendicke) auf 19 % und 11 % bei einer Konzentration von 1.5 Gew.-% senken. Cis-Bisharnstoffe sind hingegen nur geringfügig in der Lage die Kristallisation von PA6 zu induzieren und verbessern nicht die optischen Eigenschaften des Polymers. Durch systematische Variation der peripheren Substituenten konnten interessante Struktur Eigenschaftsbeziehungen aufgedeckt werden. Die untersuchten Verbindungen mit längeren beweglichen Resten stellten effiziente Nukleierungsmittel für PA6 dar, waren aber nur im geringen Maße in der Lage die optischen Eigenschaften zu verbessern. Derivate mit kurzen, hochverzweigten Substituenten, bevorzugt in α-position zu den Harnstoffgruppen, verbesserten den Haze hingegen deutlich. Es konnte zudem gezeigt werden, dass die Verarbeitungsbedingungen, insbesondere die Abkühlrate und die Probendicke, starken Einfluss auf die Endeigenschaften des Polymerfestkörpers ausüben. Trans-1,4-Cyclohexan-Bisharnstoffe wurden ebenfalls in anderen teilkristallinen Polyamid Homo- und Copolymeren untersucht. 1,1′-(trans-1,4-Cyclohexan)bis(3-tert-butylurea) war in der Lage effizient PA66, PA12, PA6/66, PA66/6 und PA6/12 zu nukleieren. Für PA66/6 und PA12 wurden Hazewerte von etwa 30 % erreicht. Eine größere Klarmodifizierung war in Gegenwart von PA6 Wiederholeinheiten möglich. Für PA6/66 konnte der Haze unter 10 % gesenkt werden. Bei PA66 und PA6/12 lagen die besten Hazewerte zwischen 40 und 50 %.

Zum ersten Mal konnte somit das Konzept der supramolekularen Nukleierungsmittel auf teilkristalline Polyamide übertragen werden. Mit der neu entwickelten Verbindungsklasse konnte gezeigt werden, dass trans-1,4-Cyclohexan-Bisharnstoffe in der Lage sind das Kristallisationsverhalten zu beschleunigen und die optischen Eigenschaften, insbesondere den Haze, deutlich zu verbessern.
8. Experimental part

8.1. Materials and equipment

*Chemicals and solvents*

cis-1,4-diaminocyclohexane  
TCI Europe 98%, used as received

Dimethylformamide  
Grüssing 99.5%, used as received

Methanol  
Technical grade, purification by distillation

N-methyl-2-pyrrolidinone  
BASF, dried over CaH₂, distilled under vacuum

Talc Micro-Talc IT Extra  
Mondo Minerals, used as received

Tetrahydrofurane  
Acros 99.9%, used as received

trans-1,4-cyclohexane diisocyanate  
ABCR 97%, used as received

trans-1,4-diaminocyclohexane  
Aldrich 98%, used as received

All solvents used for recrystallization (except DMF) were purified by distillation. All other chemicals were commercially available and used without further purification.

*Equipment*

Compounding  
Co-rotating twin-screw compounder  
DSM Xplore 15 mL

DSC  
Perkin Elmer Diamond DSC and Mettler Toledo DSC/SDTA 821e; Standard heating and cooling rates of 10 K/min

Haze-Meter  
BYK Gardener Haze Gard plus  
Calibration by clarity and haze standard

Injection molding  
Micro-injection molding machine  
DSM Xplore 12 mL

Laser transparency  
FOBA DP50 Nd:YAG laser marking system  
Wave length: 1064 nm

Mass spectroscopy  
Finnigan Mat 112S (70 eV)

Microtome  
Leica RM 2255 rotary microtome

¹H-NMR-spectroscopy  
Bruker AC 300 (300 MHz)

Polarized light microscopy  
Nikon, DIAPHOT 300 (microscope)  
Mettler, FP82HT (hot stage)
Experimental part

- **Nikon, DMX1200** (digital camera)
- **Powder mill**
  - Retsch ZM100, freezer mill, Schieritz & Hauenstein AG
- **SEM**
  - Zeiss 1530 FESEM
  - Cressington Sputter Coater 208HR
- **TGA**
  - Mettler Toledo TGA/SDTA851e
  - Standard heating and cooling rates of 10 K/min
- **WAXD**
  - Bruker D8 Advance
  - CuKα radiation (λ = 1.54Å)
8.2. Polymers

In the frame of this work the influence of supramolecular additives on the crystallization behavior of semi-crystalline polymers was studied based on a thermoplastic polyester (polybutylene terephthalate) and semi-crystalline polyamides.

The following polymers were investigated:

- Polybutylene terephthalate (PBT), Ultradur® B4500, an intermediate-viscosity, extrusion grade for the production of flat films and thin-walled profiles.
- Polyamide 6 (PA6), Ultradim® B27 E, a low-viscosity grade for compounding and the production of monofilaments.
- Polyamide 66 (PA66), Ultradim® A27 E 01, a low-viscosity grade for compounding and the production of monofilaments and bristles.
- Co-polyamide 6/66 (PA6/66), Ultradim® C33, an intermediate-viscosity grade for the production of multilayer films and monofilaments.
- Co-polyamide 66/6 (PA66/6), Ultradim® 9A E 01, a low-viscosity grade suitable for film, monofilament, extrusion, and molding applications.
- Co-polyamide 6/12 (PA6/12), Zytel® 1512, suitable for molding applications.
- Polyamide 12 (PA12), Vestamid® L2101F, a high-viscosity film grade used for packaging applications.

With exception of PA12, supplied by Evonik Industries AG, all polymer resins were provided by BASF SE. Except for PA6/12 and PA12 where no information about the specific additivation was available, all polymer grades contained no further additives. The polymer granulate was pulverized in a freezer mill (Retsch ZM100, Schieritz & Hauenstein AG) prior to use through a 1 mm sieve.
8.3. Synthesis and characterization

Synthesis and characterization of bisurea derivatives

All solvents were purified and dried prior to use according to standard procedures. The starting materials were purchased from ABCR, Acros, Aldrich, and TCI Europe and used as received. \(^1\)H-NMR spectra were recorded on a Bruker Avance 300 spectrometer. The chemical shifts are reported in ppm (δ) with the coupling constant values J given in Hz. Mass spectra were recorded on a Finnigan Mat 112S instrument (70 eV) with direct probe inlet at the central analytic laboratory of the University of Bayreuth. All spectra were recorded at 298 K.

8.3.1. General synthetic route to trans-1,4-cyclohexyl-bisurea derivatives

*Synthesis of trans-1,4-cyclohexyl-bisurea derivatives from trans-1,4-diaminocyclohexane*

Trans-1,4-diaminocyclohexane was added in a flame-dried Schlenk flask and dissolved in THF under argon atmosphere. The solution was cooled to 0°C in an ice bath and the corresponding isocyanate, diluted in THF, was added slowly under heavy stirring. The reaction mixture was heated to reflux and, unless indicated otherwise, maintained at this temperature for 12 h. After cooling the precipitated white solid was filtered off, dried under vacuum for 2 h (70°C, 100 mbar) and recrystallized.
**1,1′-(trans-1,4-cyclohexylene)bis(3-ethylurea) 1a**
Molecular formula: C_{12}H_{24}N_{4}O_{2}; M = 256.19 g/mol
Internal notebook number: JF262

**Reaction batch:**
1.78 g (16 mmol) trans-1,4-diaminocyclohexane
2.22 g (32 mmol) ethylisocyanate
150 mL THF
Purification: Recrystallization from MeOH

**Characterization:**
\[ ^1H \text{-NMR (300 MHz, DMSO d6): } \delta \text{ [ppm] = 5.6 (m, 4H), 3.26 (m, 2H), 2.97 (q, J = 6.5 Hz, 4H), 1.76 (m, 4H), 0.96 (t, J = 7.1 Hz, 6H) } \]
MS (70 eV), m/z (%): 256 (35); 212 (21); 184 (81); 168 (93); 139 (27); 96 (59); 89 (100); 68 (35); 56 (41); 44 (45)
Schmelzpunkt: T_{m} = subl.
Zersetzungspunkt: T_{-10 \text{ wt%}} = 316°C

**1,1′-(trans-1,4-cyclohexylene)bis(3-n-propylurea) 1b**
Molecular formula: C_{14}H_{28}N_{4}O_{2}; M = 284.22 g/mol
Internal notebook number: JF274

**Reaction batch:**
1.50 g (13 mmol) trans-1,4-diaminocyclohexane
2.40 g (28 mmol) propylisocyanate
200 mL THF
Purification: Recrystallization from DMF

**Characterization:**
\[ ^1H \text{-NMR (300 MHz, CDCl}_3/CF}_3\text{COOD): } \delta \text{ [ppm] = 3.45 (s, 2H), 3.24 (t, J = 7.3 Hz, 4H), 2.06 (d, J = 6.3 Hz, 4H), 1.38-1.72 (m, 8H), 0.96 (t, J = 7.4 Hz, 6H) } \]
MS (70 eV), m/z (%): 284 (27); 226 (39); 198 (68); 183 (93); 182 (73); 139 (25); 113 (10); 103 (100); 97 (52); 96 (54); 59 (44); 44 (52)
Schmelzpunkt: T_{m} = subl.
Zersetzungspunkt: T_{-10 \text{ wt%}} = 328°C

**1,1′-(trans-1,4-cyclohexylene)bis(3-n-butylurea) 1c**
Molecular formula: C_{16}H_{32}N_{4}O_{2}; M = 312.25 g/mol
Internal notebook number: JF269

**Reaction batch:**
1.50 g (13 mmol) trans-1,4-diaminocyclohexane
2.90 g (29 mmol) buylisocyanate
500 mL THF
Purification: Recrystallization from DMF

**Characterization:**
\[ ^1H \text{-NMR (300 MHz, CDCl}_3/CF}_3\text{COOD): } \delta \text{ [ppm] = 3.45 (s, 2H), 3.28 (t, J = 7.3 Hz, 4H), 2.06 (d, J = 6.3 Hz, 4H), 1.28-1.65 (m, 12H), 0.95 (t, J = 7.3 Hz, 6H) } \]
MS (70 eV), m/z (%): 312 (28); 240 (38); 212 (72); 197 (100); 139 (26); 117 (97); 96 (46); 74 (31); 57 (30)
Schmelzpunkt: T_{m} = subl.
Zersetzungspunkt: T_{-10 \text{ wt%}} = 329°C
**Experimental part**

**1,1′-(trans-1,4-cyclohexylene)bis(3-(1,2-dimethylpropyl)urea) 1m**

- **Molecular formula:** $C_{18}H_{36}N_4O_2$; $M = 440.28$ g/mol
- **Internal notebook number:** TH01
- **Reaction batch:**
  - 1.46 g (9 mmol) trans-1,4-diaminocyclohexane
  - 1.54 g (18 mmol) 1,2-dimethylpropylisocyanate
  - 200 mL THF
- **Purification:** Recrystallization from DMF
- **Characterization:**
  - $^1H$ NMR (300 MHz, CDCl$_3$/CF$_3$COOD): $\delta$ [ppm] = 3.61 (m, 2H), 3.42 (m, 2H), 2.06 (m, 4H), 1.76 (q, $J = 6.7$ Hz, 2H), 1.50 (m, 4H), 1.20 (d, $J = 6.6$ Hz, 6H), 0.94 (d, $J = 6.6$ Hz, 12H)
  - MS (70 eV), m/z (%): 340 (4); 297 (26); 254 (100); 211 (6); 141 (4); 93 (12); 45 (54)
  - Schmelzpunkt: $T_m$ = subl.
  - Zersetzungspunkt: $T_{-10\ \text{wt%}} = 317^\circ$C

**1,1′-(trans-1,4-cyclohexylene)bis(3-tert-butylurea) 1p**

- **Molecular formula:** $C_{16}H_{32}N_4O_2$; $M = 312.25$ g/mol
- **Internal notebook number:** HH7, JF272, JF277
- **Reaction batch:**
  - 5.00 g (44 mmol) trans-1,4-diaminocyclohexane
  - 9.90 g (100 mmol) tert-butylisocyanate
  - 500 mL THF
- **Purification:** Recrystallization from MeOH
- **Characterization:**
  - $^1H$-NMR (300 MHz, DMSO d6): $\delta$ [ppm] = 5.44-5.54 (m, 4H), 3.23 (m, 2H), 1.75 (m, 4H), 1.19 (s, 18H), 1.04 (m, 4H)
  - MS (70 eV), m/z (%): 312 (18); 240 (7); 212 (44); 197 (32); 156 (5); 139 (10); 97 (25); 61 (15); 58 (100)
  - Schmelzpunkt: $T_m$ = subl.
  - Zersetzungspunkt: $T_{-10\ \text{wt%}} = 303^\circ$C

**1,1′-(trans-1,4-cyclohexylene)bis(3-cyclohexylurea) 1q**

- **Molecular formula:** $C_{20}H_{36}N_4O_2$; $M = 364.28$ g/mol
- **Internal notebook number:** JF246
- **Reaction batch:**
  - 2.51 g (22 mmol) trans-1,4-diaminocyclohexane
  - 4.49 g (44 mmol) cyclohexyl isocyanate
  - 300 mL THF
- **Purification:** Recrystallization from DMF
- **Characterization:**
  - $^1H$ NMR (300 MHz, CDCl$_3$/CF$_3$COOD): $\delta$ [ppm] 3.56 (s, 2H), 3.4 (s, 2H), 2.33-0.9 (m, 28H)
  - MS (70 eV), m/z (%): 364 (11); 266 (8); 238 (46); 223 (55); 143 (38); 98 (45); 57 (100); 44 (29)
  - Schmelzpunkt: $T_m$ = subl.
  - Zersetzungspunkt: $T_{-10\ \text{wt%}} = 332^\circ$C
1,1'-(trans-1,4-cyclohexylene)bis(3-phenylurea) \textit{1r}

Molecular formula: C$_{20}$H$_{24}$N$_4$O$_2$; M = 352.19 g/mol

Internal notebook number: JF247

\textit{Reaction batch:}

1.50 g (13 mmol) trans-1,4-diaminocyclohexane
2.90 g (26 mmol) phenylisocyanate
300 mL THF

Purification: Recrystallization from MeOH

\textit{Characterization:}

$^1$H-NMR (300 MHz, DMSO d$_6$): $\delta$ [ppm] = 8.3 (s, 2H), 7.4 (m, 4H), 7.18 (t, J = 7.8 Hz, 4H), 6.9 (t, J = 7.3 Hz, 2H), 6.06 (d, J = 7.9 Hz, 2H), 3.4 (m, 2H), 1.88 (m, 4H) 1.25 (m, 4H)

MS (70 eV), m/z (%): 352 (3); 259 (8); 232 (4); 141 (5); 99 (10); 93 (100); 66 (6); 44 (6)

Schmelzpunkt: T$_{m}$ = subl.

Zersetzungspunkt: T$_{10 \text{ wt\%}}$ = 309°C

1,1'-(trans-1,4-cyclohexylene)bis(3-(1-adamantyl)urea) \textit{1s}

Molecular formula: C$_{28}$H$_{44}$N$_4$O$_2$; M = 468.70 g/mol

Internal notebook number: JF268

\textit{Reaction batch:}

1.50 g (13 mmol) trans-1,4-diaminocyclohexane
3.40 g (29 mmol) 1-adamantyl isocyanate
200 mL THF

Purification: Extraction with DMF

\textit{Characterization:}

$^1$H-NMR (300 MHz, DMSO d$_6$): $\delta$ [ppm] = 5.52 (d, J = 7.9 Hz, 2H), 5.37 (s, 2H), 3.26 (m, 2H), 1.97 (s, 6H), 1.45-2.08 (m, 34H), 1.03 (m, 4H)

MS (70 eV), m/z (%): 468 (18); 290 (65); 275 (29); 177 (30); 135 (74); 113 (50); 94 (100); 58 (25)

Schmelzpunkt: T$_{m}$ = subl.

Zersetzungspunkt: T$_{10 \text{ wt\%}}$ = 339°C
Experimental part

Synthesis of trans-1,4-cyclohexyl-bisurea derivatives from trans-1,4-cyclohexane diisocyanate

Trans-1,4-cyclohexane diisocyanate was added in a flame-dried Schlenk flask and dissolved in THF under argon atmosphere. The solution was cooled to 0°C in an ice bath and the corresponding amine, diluted in THF, was added slowly under heavy stirring. The reaction mixture was heated to reflux and, unless indicated otherwise, maintained at this temperature for 12 h. After cooling the precipitated white solid was filtered off, dried under vacuum for 2 h (70°C, 100 mbar) and recrystallized.

1,1'-(trans-1,4-cyclohexylene)bis(3-n-hexylurea) 1d
Molecular formula: C_{20}H_{40}N_{4}O_{2}; M = 368.32 g/mol
Internal notebook number: FR84
Reaction batch:
1.35 g (8 mmol) trans-1,4-cyclohexane diisocyanate
1.46 g (16 mmol) hexylamine
200 mL THF
Purification: Recrystallization from DMF
Characterization:
\(^1\)H NMR (300 MHz, CDCl\(_3\)/CF\(_3\)COOD): \(\delta [ppm] = 3.44 \, (s, 2H), \, 3.24 \, (m, 4H), \, 2.05 \, (m, 4H), \, 1.18-1.65 \, (m, 20H), \, 0.90 \, (t, J = 7.0 \, Hz, 6H)\)
MS (70 eV), m/z (%): 368 (23); 268 (30); 240 (78); 225 (100); 145 (76); 102 (27); 97 (41); 57 (28); 44 (37)
Schmelzpunkt: T\(_m\) = 370°C
Zersetzungspunkt: T\(_{-10 \text{wt}}\%\) = 306°C

1,1'-(trans-1,4-cyclohexylene)bis(3-n-octylurea) 1e
Molecular formula: C_{24}H_{48}N_{4}O_{2}; M = 424.38 g/mol
Internal notebook number: SG460
Reaction batch:
3.30 g (20 mmol) trans-1,4-cyclohexane diisocyanate
5.69 g (44 mmol) octylamine
200 mL THF
Purification: Recrystallization from DMF
Characterization:
\(^1\)H NMR (300 MHz, CDCl\(_3\)/CF\(_3\)COOD): \(\delta [ppm] = 3.56 \, (m, 2H), \, 3.30 \, (t, J = 7.3 \, Hz, 4H), \, 2.12 \, (m, 4H), \, 1.64 \, (m, 4H), \, 1.52 \, (m, 4H), \, 1.33 \, (m, 20H), \, 0.91 \, (m, 6H)\)
MS (70 eV), m/z (%): 424 (21); 296 (30); 268 (66); 263 (100); 173 (58); 130 (27); 97 (36); 44 (27)
Schmelzpunkt: T\(_m\) = 370°C
Zersetzungspunkt: T\(_{-10 \text{wt}}\%\) = 315°C
1,1’-(trans-1,4-cyclohexylene)bis(3-n-octadecylurea) 1f
Molecular formula: C_{44}H_{88}N_{4}O_{2}; M = 705.20 g/mol
Internal notebook number: FR79
Reaction batch:
0.71 g (4 mmol) trans-1,4-cyclohexane diisocyanate
2.29 g (9 mmol) octadecylamine
200 mL THF
Purification: Recrystallization from DMF
Characterization:
^{1}H NMR (300 MHz, CDCl_{3}/CF_{3}COOD): \delta [ppm] = 3.45 (s, 2H), 3.25 (m, 4H), 2.06 (m, 4H), 1.17-1.68 (m, 68H), 0.90 (t, J = 7.0 Hz, 6H) 
MS (70 eV), m/z (%): 704 (4); 546 (34); 501 (6); 437 (4); 409 (10); 367 (12); 313 (10); 270 (16); 238 (4); 170 (4); 141 (15); 99 (40); 44 (100)
Schmelzpunkt: T_{m} = 345°C
Zersetzungspunkt: T_{-10 wt\%} = 278°C

1,1’-(trans-1,4-cyclohexylene)bis(3-(6-methylheptane)urea) 1g
Molecular formula: C_{24}H_{48}N_{4}O_{2}; M = 424.38 g/mol
Internal notebook number: JF285
Reaction batch:
3.09 g (18 mmol) trans-1,4-cyclohexane diisocyanate
5.29 g (40 mmol) 1-amino-6-methylheptane
150 mL THF
Purification: Recrystallization from DMF
Characterization:
^{1}H NMR (300 MHz, CDCl_{3}/CF_{3}COOD): \delta [ppm] = 3.52 (s, 2H), 3.19 (d, J = 6.5 Hz, 4H), 2.09 (m, 4H), 1.17-1.62 (m, 22H), 0.91 (m, 12H)
MS (70 eV), m/z (%): 424 (29); 313 (8); 296 (91); 268 (95); 253 (100); 173 (52); 130 (39); 96 (35); 58 (39); 44 (31)
Schmelzpunkt: T_{m} = subl.
Zersetzungspunkt: T_{-10 wt\%} = 286°C

1,1’-(trans-1,4-cyclohexylene)bis(3-(1,5-dimethylhexyl)urea) 1h
Molecular formula: C_{24}H_{48}N_{4}O_{2}; M = 424.38 g/mol
Internal notebook number: JF113
Reaction batch:
3.30 g (20 mmol) trans-1,4-cyclohexane diisocyanate
5.65 g (44 mmol) 1,5-dimethylhexylamine
200 mL THF
Purification: Recrystallization from DMF
Characterization:
^{1}H NMR (300 MHz, CDCl_{3}/CF_{3}COOD): \delta [ppm] = 3.80 (sx, J = 6.6 Hz, 2H), 3.46 (s, 2H), 2.08 (m, 4H), 1.11-1.69 (m, 24H), 0.88 (d, J = 6.6 Hz, 12H) 
MS (70 eV), m/z (%): 425 (18); 340 (15); 297 (90); 269 (44); 254 (52); 211 (18); 174 (19); 128 (26); 97 (22); 44 (100)
Schmelzpunkt: T_{m} = 355°C
Zersetzungspunkt: T_{-10 wt\%} = 299°C
1,1’-(trans-1,4-cyclohexylene)bis(3-(2-ethylhexyl)urea) 1i
Molecular formula: C_{24}H_{48}N_4O_2; M = 424.38 g/mol
Internal notebook number: JF112

**Reaction batch:**
3.30 g (20 mmol) trans-1,4-cyclohexane diisocyanate
5.65 g (44 mmol) 2-ethylhexylamine
200 mL THF
Purification: Recrystallization from MeOH/i-PrOH (2:1)

**Characterization:**
^1^H NMR (300 MHz, CDCl_3/CF_3COOD): δ [ppm] = 3.53 (s, 2H), 3.21 (d, J = 6.5 Hz, 4H), 2.10 (m, 4H), 1.17-1.62 (m, 22H), 0.91 (t, J = 7.2 Hz, 12H)
MS (70 eV), m/z (%): 425 (39); 314 (18); 297 (96); 269 (92); 254 (100); 174 (34); 131 (18); 97 (14); 57 (13)
Schmelzpunkt: T_m = subl.
Zersetzungspunkt: T_{-10 wt%} = 290°C

1,1’-(trans-1,4-cyclohexylene)bis(3-(tert-octyl)urea) 1j
Molecular formula: C_{24}H_{48}N_4O_2; M = 424.38 g/mol
Internal notebook number: JF124

**Reaction batch:**
3.23 g (19 mmol) trans-1,4-cyclohexane diisocyanate
5.53 g (43 mmol) tert-octylamine
300 mL THF
Purification: Recrystallization from DMF

**Characterization:**
^1^H-NMR (300 MHz, DMSO d6): δ [ppm] = 5.48 (d, J = 7.9 Hz, 2H), 5.39 (s, 2H), 1.75 (m, 4H), 1.64 (s, 4H), 1.22 (s, 12H), 1.04 (m, 4H), 0.93 (s, 18H)
MS (70 eV), m/z (%): 425 (2); 354 (34); 314 (3); 397 (15); 241 (5); 225 (4); 184 (10); 114 (14); 97 (6); 58 (100)
Schmelzpunkt: T_m = subl.
Zersetzungspunkt: T_{-10 wt%} = 312°C

1,1’-(trans-1,4-cyclohexylene)bis(3-(1,1-dimethylpropy)urea) 1k
Molecular formula: C_{18}H_{36}N_4O_2; M = 340.5 g/mol
Internal notebook number: JF283

**Reaction batch:**
2.80 g (17 mmol) trans-1,4-cyclohexane diisocyanate
3.40 g (38 mmol) 1,1-dimethylpropylamine
400 mL THF
Purification: Recrystallization from DMF

**Characterization:**
^1^H-NMR (300 MHz, DMSO d6): δ [ppm] = 5.53 (d, J = 7.9 Hz, 2H), 5.36 (s, 2H), 3.23 (m, 2H), 1.75 (m, 4H), 1.54 (m, 4H), 1.13 (s, 12H), 1.04 (m, 4H), 0.75 (t, J = 7.3 Hz, 6H)
MS (70 eV), m/z (%): 340 (6); 311 (9); 254 (9); 226 (11); 211 (7); 184 (6); 96 (8); 72 (24); 59 (100); 44 (11)
Schmelzpunkt: T_m = subl.
Zersetzungspunkt: T_{-10 wt%} = 319°C

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1,1'-(trans-1,4-cyclohexylene)bis(3-(3-pentyl)urea) 1l
Molecular formula: C_{18}H_{36}N_{4}O_{2}; M = 340.5 g/mol
Internal notebook number: JF273
Reaction batch:
2.00 g (12 mmol) trans-1,4-cyclohexane diisocyanate
2.20 g (25 mmol) 3-pentylamine
200 mL THF
Purification: Recrystallization from DMF
Characterization:
\^1H NMR (300 MHz, CDCl_{3}/CF_{3}COOD): δ [ppm] = 3.57 (m, 2H), 3.44 (m, 2H), 2.07 (m, 4H), 1.66 (m, 4H), 1.50 (m 8H), 0.94 (t, J = 7.4 Hz, 12H)
MS (70 eV), m/z (%): 340 (19); 311 (26); 254 (100); 226 (38); 211 (46); 131 (22); 96 (19); 86 (21); 59 (99)
Schmelzpunkt: T_{m} = subl.
Zersetzungspunkt: T_{-10 wt%} = 310°C

1,1'-(trans-1,4-cyclohexylene)bis(3-neopentylurea) 1n
Molecular formula: C_{18}H_{36}N_{4}O_{2}; M = 340.5 g/mol
Internal notebook number: JF267
Reaction batch:
2.00 g (12 mmol) trans-1,4-cyclohexane diisocyanate
2.30 g (26 mmol) neopentylamine
200 mL THF
Purification: Recrystallization from DMF
Characterization:
\^1H-NMR (300 MHz, DMSO d6): δ [ppm] = 5.68 (m, 4H), 2.78 (m, 4H), 1.78 (m, 4H), 1.09 (m, 4H), 0.8 (s, 18H)
MS (70 eV), m/z (%): 340 (16); 283 (8); 254 (100); 226 (33); 211 (65); 139 (15); 131 (41); 96 (26); 88 (23); 58 (17)
Schmelzpunkt: T_{m} = subl.
Zersetzungspunkt: T_{-10 wt%} = 323°C

1,1'-(trans-1,4-cyclohexylene)bis(3-isopropylurea) 1o
Molecular formula: C_{14}H_{28}N_{4}O_{2}; M = 284.4 g/mol
Internal notebook number: HH6
Reaction batch:
1.75 g (11 mmol) trans-1,4-cyclohexane diisocyanate
1.25 g (21 mmol) isopropylamine
400 mL THF
Purification: Recrystallization from DMF
Characterization:
\^1H-NMR:
MS (70 eV), m/z (%): 284 (13); 226 (35); 198 (48); 183 (63); 139 (18); 113 (13); 103 (70); 97 (48); 96 (50); 58 (70); 45 (100)
Schmelzpunkt: T_{m} = subl.
Zersetzungspunkt: T_{-10 wt%} = 319°C
8.3.2. General synthetic route to cis-1,4-cyclohexyl-bisurea derivatives

Cis-1,4-diaminocyclohexane was added in a flame-dried Schlenk flask and dissolved in THF under argon atmosphere. The solution was cooled to 0°C in an ice bath and the corresponding isocyanate, diluted in THF, was added slowly under heavy stirring. The reaction mixture was heated to reflux and, unless indicated otherwise, maintained at this temperature for 12 h. After cooling the precipitated white solid was filtered off, dried under vacuum for 2 h (70°C, 100 mbar) and recrystallized.

1,1’-(cis-1,4-cyclohexylene)bis(3-n-octylurea) 2a
Molecular formula: C_{24}H_{48}N_{4}O_{2}; M = 424.38 g/mol
Internal notebook number: SG462

Reaction batch:
3.30 g (28 mmol) cis-1,4-diaminocyclohexane
9.89 g (64 mmol) octylisocyanate

200 mL THF

Purification: Recrystallization from MeOH

Characterization:
\(^1\)H-NMR (300 MHz, DMSO d6): \(\delta \text{ [ppm]} = 5.73 \text{ (m, 4H), 2.95 (q, J = 6.4 Hz ,4H), 1.08-1.65 (m, 34H), 0.85 (m, 6H)}\)

MS (70 eV), m/z (%): 424 (21); 296 (39); 252 (82); 213 (28); 173 (81); 141 (45); 97 (100); 59 (58); 57 (61); 44 (57)

Schmelzpunkt: \(T_m = 308°C\)
Zersetzungspunkt: \(T_{-10 \text{ wt%}} = 254°C\)

1,1’-(cis-1,4-cyclohexylene)bis(3-tert-octylurea) 2b
Molecular formula: C_{24}H_{48}N_{4}O_{2}; M = 424.38 g/mol
Internal notebook number: JF286

Reaction batch:
0.66 g (5 mmol) cis-1,4-diaminocyclohexane
2.0 g (12 mmol) tert-octylisocyanate

100 mL THF

Purification: Recrystallization from Cyclohexane

Characterization:
\(^1\)H-NMR (300 MHz, DMSO d6): \(\delta \text{ [ppm]} = 5.60 \text{ (d, J = 7.5 Hz, 2H), 5.52 (s, 2H), 1.64 (s, 6H), 1.27-1.57 (m, 8H), 1.23 (s, 12H), 0.93 (s, 18H)}\)

MS (70 eV), m/z (%): 426 (11); 354 (34); 297 (44); 184 (20); 141 (15); 114 (5); 58 (100)

Schmelzpunkt: \(T_m = 325°C\)
Zersetzungspunkt: \(T_{-10 \text{ wt%}} = 243°C\)
1,1’-(cis-1,4-cyclohexylene)bis(3-isopropylurea) 2c
Molecular formula: C\textsubscript{20}H\textsubscript{24}N\textsubscript{4}O\textsubscript{2}; M = 352.40 g/mol
Internal notebook number: JF254

Reaction batch:
1.50 g (13 mmol) cis-1,4-diaminocyclohexane
2.21 g (26 mmol) isopropyl isocyanate
150 mL THF
Purification: Recrystallization from MeOH

Characterization:
\(^1\)H-NMR (300 MHz, DMSO d6): \(\delta [ppm] = 5.65 \text{ (d, J = 7.5 Hz, 2H)}\), 5.57 \(\text{ (d, J = 7.6 Hz, 2H)}\), 3.63 \(\text{ (m, 2H)}\), 3.46 \(\text{ (m, 2H)}\), 1.00 \(\text{ (d, J = 6.5 Hz, 12H)}\)
MS (70 eV), m/z (%): 284 (8); 226 (23); 198 (12); 182 (100); 141 (39); 113 (8); 103 (75); 97 (67); 68 (36); 59 (88); 45 (81); 44 (59);
Schmelzpunkt: \(T_m = 309^\circ C\)
Zersetzungspunkt: \(T_{-10 \text{ wt\%}} = 265^\circ C\)

1,1’-(cis-1,4-cyclohexylene)bis(3-tert-butylurea) 2d
Molecular formula: C\textsubscript{16}H\textsubscript{32}N\textsubscript{4}O\textsubscript{2}; M = 312.50 g/mol
Internal notebook number: JF248

Reaction batch:
1.50 g (13 mmol) cis-1,4-diaminocyclohexane
2.57 g (26 mmol) tert-butylisocyanate
150 mL THF
Purification: Recrystallization from MeOH

Characterization:
\(^1\)H-NMR (300 MHz, DMSO d6): \(\delta [ppm] = 5.53 \text{ (m, 2H)}\), 5.68 \(\text{ (s, 2H)}\), 3.43 \(\text{ (bm, 2H)}\), 1.49 \(\text{ (m, 4H)}\), 1.35 \(\text{ (m, 4H)}\), 1.20 \(\text{ (s, 18H)}\)
MS (70 eV), m/z (%): 297 (3); 240 (9); 196 (24); 184 (2); 141 (28); 117 (14); 97 (27); 59 (100); 42 (11);
Schmelzpunkt: \(T_m = 319^\circ C\)
Zersetzungspunkt: \(T_{-10 \text{ wt\%}} = 264^\circ C\)

1,1’-(cis-1,4-cyclohexylene)bis(3-cyclohexylurea) 2e
Molecular formula: C\textsubscript{20}H\textsubscript{36}N\textsubscript{4}O\textsubscript{2}; M = 364.53 g/mol
Internal notebook number: JF249

Reaction batch:
1.50 g (13 mmol) cis-1,4-diaminocyclohexane
3.25 g (26 mmol) cyclohexylisocyanate
150 mL THF
Purification: Recrystallization from MeOH

Characterization:
\(^1\)H NMR (300 MHz, DMSO d6): \(\delta [ppm] = 5.66 \text{ (m, 4H)}\), 3.45 \(\text{ (s, 2H)}\), 1.82-0.89 \(\text{ (m, 28H)}\)
MS (70 eV), m/z (%): 364 (5); 266 (13); 238 (8); 222 (30); 192 (8); 143 (29); 98 (39); 57 (100); 44 (62); 42 (42);
Schmelzpunkt: \(T_m = 252^\circ C\)
Zersetzungspunkt: \(T_{-10 \text{ wt\%}} = 266^\circ C\)
1,1’-(cis-1,4-cyclohexylene)bis(3-phenylurea) 2f
Molecular formula: C_{20}H_{24}N_{4}O_{2}; M = 352.43 g/mol
Internal notebook number: JF250

Reaction batch:
1.50 g (13 mmol) cis-1,4-diaminocyclohexane
3.1 g (26 mmol) phenylisocyanate
150 mL THF
Purification: Recrystallization from MeOH

Characterization:
\(^1\text{H-NMR (300 MHz, DMSO d6)}\): δ [ppm] = 8.33 (s, 2H), 7.37 (m, 4H), 7.22 (t, J = 7.8 Hz, 4H), 6.88 (t, J = 7.3 Hz, 2H), 6.20 (d, J = 7.9 Hz, 2H), 3.61 (m, 2H), 1.66 (m, 4H) 1.51 (m, 4H)

MS (70 eV), m/z (%): 352 (9); 260 (15); 141 (10); 94 (13); 93 (100); 81 (6)

Schmelzpunkt: T\(_m\) = 236°C
Zersetzungspunkt: T\(_{-10 \text{wt%}}\) = 258°C
8.3.3. General synthetic route to asymmetric substituted trans-1,4-cyclohexyl-bisurea derivatives

*Preparation of trans-1-(4-aminocyclohexyl)-3-cyclohexylurea*

Trans-1,4-diaminocyclohexane was added in a flame-dried Schlenk flask and dissolved in THF under argon atmosphere. The solution was cooled to -40°C in a cooling bath (isopropyl alcohol/dry ice) and isocyanatocyclohexane, diluted in THF, was added slowly under heavy stirring. The reaction mixture was stirred for 12 h at room temperature. The precipitated white solid was filtered off, suspended in water and acidified to pH 2 (HCl). The emerging clear solution was again filtrated and the filtrate was brought to pH 8 (NaOH) whereupon trans-1-(4-aminocyclohexyl)-3-cyclohexylurea precipitated as a white solid.

**trans-1-(4-aminocyclohexyl)-3-cyclohexylurea**

Molecular formula: C\(_{13}\)H\(_{25}\)N\(_3\)O; M = 239.63 g/mol
Internal notebook number: FR86, JF258

*Reaction batch:*
2.39 g (21 mmol) trans-1,4-diaminocyclohexane
2.61 g (21 mmol) cyclohexyl isocyanate
300 mL THF

*Characterization:*
MS (70 eV), m/z (%): 239 (26); 222 (5); 183 (6); 143 (22); 113 (18); 97 (100); 59 (44); 57 (48); 44 (27)
Schmelzpunkt: \(T_m = \text{subl.}\)
Zersetzungspunkt: \(T_{10\text{ wt\%}} = 229°C\)

*Preparation of asymmetric substituted trans-1,4-cyclohexyl-bisurea derivatives*

Trans-1-(4-aminocyclohexyl)-3-cyclohexylurea was added in a flame-dried Schlenk flask and dissolved in NMP under argon atmosphere. The corresponding isocyanate, diluted in NMP, was added slowly under heavy stirring. The resulting mixture was heated to 80°C and stirred for 12 h. The solution was precipitated in 1M HCl and filtered off. The resulting white solid was washed with THF, dried under vacuum for 2 h (70°C, 100 mbar) and recrystallized.
Experimental part

1-tert-octyl-3-[4-(cyclohexylcarbamoylamino)cyclohexyl]urea 3a
Molecular formula: C_{22}H_{42}N_{4}O_{2}; M = 394.59 g/mol
Internal notebook number: JF257
Reaction batch:
1.10 g (5 mmol) trans-1-(4-aminocyclohexyl)-3-cyclohexylurea
0.70 g (5 mmol) tert-octylisocyanate
80 mL NMP
Characterization:
\(^1\)H-NMR (300 MHz, DMSO d6): \(\delta\) [ppm] = 5.57 (d, \(J = 7.7 \text{ Hz}, 2H\)), 5.47 (d, \(J = 7.9 \text{ Hz}, 1H\)), 5.42 (s, \(1H\)), 3.27 (m, \(2H\)), 1.22 (s, \(6H\)), 1.04 (m, \(4H\)), 0.93 (s, \(9H\)), 0.84-1.86 (m, \(33H\))
MS (70 eV), m/z (%): 394 (5); 379 (3); 323 (80); 283 (10); 266 (52); 238 (14); 223 (9); 184 (5); 141 (6); 114 (28); 98 (15); 59 (100)
Schmelzpunkt: \(T_m = \text{subl.}\)
Zersetzungspunkt: \(T_{-10 \text{ wt\%}} = 302^\circ C\)

1-isopropyl-3-[4-(cyclohexylcarbamoylamino)cyclohexyl]urea 3b
Molecular formula: C_{17}H_{32}N_{4}O_{2}; M = 324.50 g/mol
Internal notebook number: JF255
Reaction batch:
2.00 g (8 mmol) trans-1-(4-aminocyclohexyl)-3-cyclohexylurea
0.70 g (8 mmol) isopropyl isocyanate
120 mL NMP
Characterization:
\(^1\)H NMR (300 MHz, CDCl\(_3\)/CF\(_3\)COOD): \(\delta\) [ppm] = 3.94 (m, \(1H\)), 3.56 (m, \(1H\)), 3.39 (m, \(2H\)), 1.08-2.14 (m, \(24H\))
MS (70 eV), m/z (%): 324 (27); 266 (21); 248 (35); 223 (47); 198 (60); 183 (55); 143 (45); 103 (46); 97 (67); 68 (26); 59 (49); 57 (100); 45 (49); 44 (50)
Schmelzpunkt: \(T_m = \text{subl.}\)
Zersetzungspunkt: \(T_{-10 \text{ wt\%}} = 346^\circ C\)

1-tert-butyl-3-[4-(cyclohexylcarbamoylamino)cyclohexyl]urea 3c
Molecular formula: C_{18}H_{34}N_{4}O_{2}; M = 338.49 g/mol
Internal notebook number: FR87
Reaction batch:
2.83 g (12 mmol) trans-1-(4-aminocyclohexyl)-3-cyclohexylurea
1.17 g (12 mmol) tert-butylisocyanate
80 mL NMP
Characterization:
\(^1\)H NMR (300 MHz, CDCl\(_3\)/CF\(_3\)COOD): \(\delta\) [ppm] = 3.59 (m, \(1H\)), 3.30 (m, \(2H\)), 1.43 (s, \(9H\)), 1.08-2.12 (m, \(18H\))
MS (70 eV), m/z (%): 338 (14); 266 (11); 238 (51); 222 (31); 197 (29); 184 (7); 143 (22); 97 (39); 61 (21); 59 (100); 57 (62);
Schmelzpunkt: \(T_m = 379^\circ C\)
Zersetzungspunkt: \(T_{-10 \text{ wt\%}} = 308^\circ C\)
1-phenyl-3-[4-(cyclohexylcarbamoylamino)cyclohexyl]urea 3d
Molecular formula: $C_{20}H_{30}N_4O_2$; $M = 358.48$ g/mol
Internal notebook number: JF251

Reaction batch:
3.00 g (13 mmol) trans-1-(4-aminocyclohexyl)-3-cyclohexylurea
1.50 g (13 mmol) phenylisocyanate
130 mL NMP

Characterization:
$^1$H-NMR (300 MHz, DMSO d6): $\delta$ [ppm] = 8.29 (s, 1H), 7.35 (m, 2H), 7.20 (t, $J = 7.9$ Hz, 2H),
6.87 (t, $J = 7.2$ Hz, 1H), 6.02 (d, $J = 7.8$ Hz, 1H), 5.59 (m, 2H), 0.93-1.94 (m, 18H)
MS (70 eV), m/z (%): 358 (21); 265 (7); 248 (15); 232 (19); 141 (10); 93 (100); 81 (9); 57 (34);
44 (13)
Schmelzpunkt: $T_m = \text{subl.}$
Zersetzungspunkt: $T_{10 \text{ wt%}} = 323^\circ$C
9. Literature


b) R. L. Mahaffey, US 4371645.


c) M. J. Mannion, US 5198484.


[58] BYK, Qualitätskontrolle für Lacke und Kunststoffe. 2007.
c) W. Korte, Einfluss der Spritzgießherstellbedingungen und Möglichkeiten der prozessnahen Qualitätssicherung mittels statistischer Methoden beim Ultraschallschweißen.: Dissertation, RWTH Aachen. 1996.


Erklärung:

Hiermit erkläre ich, dass ich die Arbeit selbstständig verfasst und keine als die von mir angegebenen Quellen und Hilfsmittel benutzt habe.


Bayreuth, den

(Florian Richter)