Title: CONTROLLED FREE-RADICAL POLYMERIZATION PRODUCTS USING NEW CONTROL AGENTS

Abstract: The present invention relates to a new process for the production of polymers by means of pseudo-living free-radical polymerization using nitrogen monoxide, as well as to intermediate products and polymers obtainable by the process according to the invention and the use thereof.
**Controlled free-radical polymerization products using new control agents**

The present invention relates to a new process for the preparation of polymers via controlled "pseudo-living" free-radical polymerization of vinyl monomers using nitrogen monoxide.

Nowadays, the demand for homopolymers, random copolymers and block copolymers of a specific molecular weight, a narrow molecular weight distribution and/or well-defined end groups has continuously increased in a number of industries. The controlled structure of these macromolecules provides them with novel properties and allows a tailor-made property profile to be obtained. Thus many new technologies require controlled polymer structures, such as for example in the fields of electronics, computer science, communications, genetic engineering, biotechnology and materials science.

Well-defined polymers can for example be produced by ionic living polymerization processes. However, ionic processes require drastic reaction conditions, such as for example temperatures of about -78°C, extreme dryness and the absence of protic species and only a few monomers can be used.

In contrast to ionic processes, free-radical polymerization can be carried out under mild conditions and a wide range of monomers can be used. Significant progress in the control of free-radical polymerization has been made.

Otsu et al., for example, were the first to report on the possible control of the free-radical polymerization of some vinyl monomers based on the iniferter concept (see: *J. Polym. Sci.: Part A: Polym. Chem.* 2000, 38, 2121). Meanwhile three main methods of controlled free-radical polymerization, sometimes described as "living" free-radical polymerization, have been developed:
Nitroxide-mediated polymerization (NMP), which is based on the reversible capture of the propagating radicals by nitroxide radicals to form dormant chains. This strategy is disclosed for example in US-A 4,581,429. This process is handicapped by slow polymerization kinetics, a limited range of suitable monomers and the high cost of the required nitroxide radicals.

Atom transfer radical polymerization (ATRP), which involves the reversible trapping of the propagating chains by halogen atoms and is mediated by metallic catalysts, is described extensively in Acc. Chem. Res. 1999, 32, 895. A serious drawback of this method is the use of transition metal catalysts (Fe, Cu) which are potentially toxic, impart colour to the resulting polymers and lead to corrosion problems. Removal of the catalyst from the resulting polymers is possible but relatively costly.

Reversible addition-fragmentation chain transfer (RAFT) using dithio esters as transfer agents, which is disclosed for example in WO 98/01478. This method is limited to specific monomers and uses costly dithio esters which impart an unpleasant smell to the resulting polymers.

Although controlled free-radical polymerization represents an attractive method of obtaining new polymer structures there is still a need for an inexpensive, versatile control method that can be applied to a broad range of monomers.

One method of obtaining alkoxyamine initiators as nitroxide precursors has been described in WO 98/13392 and in EP-A 0 891 986. Conventional free-radical initiators are decomposed thermally in the presence of nitrogen monoxide NO. The resulting NO compounds can be used as initiators for controlling the free-radical polymerization of vinyl monomers. A disadvantage of these initiators is that polymerization appears to be controlled only for short polymer chains with number average molecular weights smaller than 15,000. Most industrially relevant polymers have to possess a number average molecular weight in excess of 50,000 in order to display the desired me-
chanical properties such as rigidity, elasticity or toughness. In addition, the methods described in the cited prior art still require the synthesis and isolation of the initiators in a step separate from the polymerization.

When added directly during free-radical polymerization nitrogen monoxide is reported (such as for example in DE-A 2 660 230, US-A 4,040,912, US-A 3,964,979) to be a polymerization inhibitor.

Under appropriate conditions sodium nitrite can be used as a precursor of nitroxide compounds. Since sodium nitrite is inexpensive, this could provide a cheap method of access to nitroxide radicals.

The prior art has already mentioned sodium nitrite as a control agent for free-radical polymerization. For instance, EP-A 1 061 059 mentions sodium nitrite as an inhibitor of the free radical polymerization of butadiene when used in combination with specific phosphorus-containing compounds. The possible control of polymerization under such conditions is not disclosed.

In attempts to reduce gel formation during the polymerization of concentrated solutions of sodium acrylate in water, Bortel et al. (*J.M.S.-Pure Appl. Chem. 1998, A35(2), 401*) observed that adding sodium nitrite inhibits polymerization up to a polymerization temperature of 60°C. It was found that although the molecular weight of the polymer increases slightly with time, the polydispersity \( D = \bar{M}_w / \bar{M}_n \) is very high (typically \( D \) higher than 6), this being a clear indication of "classical" free-radical polymerization. Other monomers than sodium acrylate are not disclosed.

The in-situ formation of NO compounds from free radical initiators and nitroso compounds is reported in *Mendeleev Comm. 1999, 6, 250*. Similar compounds can also be formed by the reaction of free radical initiators with nitrones as reported in WO 99/03894. These methods still however require the use of preformed reagents which may be toxic.
The present invention is based on the problem of providing homo- and copolymers of controlled molecular weight and controlled molecular structure by a simple and inexpensive method of controlling the free-radical polymerization of vinyl monomers that overcomes the drawbacks encountered in the prior art.

It has now been found that the free radical polymerization of vinyl monomers and of vinyl monomer mixtures can be easily controlled with respect to the molecular weight or molecular weight distribution of the resulting homopolymers, random and block copolymers if it is conducted in the presence of a reaction product of at least one vinyl monomer with nitrogen monoxide and at least one free-radical initiator.

The present invention relates to a process for producing polymers by means of controlled pseudo-living free-radical polymerization of olefinically unsaturated monomers using nitrogen monoxide.

In the present process, in a first step a free-radical intermediate product of the general formula (I),

\[ \text{Init}_1^1 \left( M \right)_p \text{N-O} \left( M \right)_q \text{Init}_2^2 \]  

in which

\begin{align*}
M & \quad \text{represents any desired vinyl monomer,} \\
p & \quad \text{is an integer between 1 and 15,} \\
q & \quad \text{is an integer between 1 and 15 and} \\
\text{Init}_1^1 \text{ and } \text{Init}_2^2 & \quad \text{are radicals of a free-radical initiator, wherein } I_1 \text{ and } I_2 \text{ can be identical or different within the compound,}
\end{align*}
is produced by reacting at least one monomer with nitrogen monoxide and at least one initiator and then in a second step the polymerization is carried out optionally with the addition of an additional monomer or monomer mixture and/or free-radical initiator.

The free-radical intermediate product of the general formula (I) can optionally be isolated. Preferably the process according to the invention is carried out in the form of a one-pot synthesis.

The polymerization process of the present invention can be used to prepare various polymers of low polydispersity, which include for example end-functionalized polymers, block, multiblock or gradient polymers, star-shaped polymers or graft and branched polymers.

The first and second steps of the polymerization can generally be carried out in different temperature ranges. The polymerization step of the process according to the invention can also optionally be carried out in several stages, it being possible to add the required monomer or monomer mixture at any stage. Using this method block copolymers can for example be produced. The excess monomer or monomer mixture can optionally be removed at the end of each stage before continuing with the addition of further monomer.

Monomers or combinations of monomers known for free-radical polymerization can be polymerized according to the invention.

Monomers M that can be used in the first step of the present invention are any polymerizable vinyl monomers such as styrene, α-methylstyrene, p-methylstyrene and other alkylstyrenes, acrylic and methacrylic acid and alkyl esters thereof, (meth)acrylamide, (meth)acrylonitrile, acrolein, vinyl acetate, vinyl chloride, conjugated dienes and vinylypyridine. Preferred monomers are styrene, α-methylstyrene and
alkylstyrenes, acrylic and methacrylic acid and alkyl esters thereof, acrylonitrile, butadiene or isoprene. Particularly preferred monomers are acrylic and methacrylic alkyl esters and acrylonitrile. In particular M can also represent different vinyl monomers, such as for example where the compound (I) is formed by reacting nitrogen monoxide with at least one initiator and a mixture of vinyl monomers.

Typical monomers M suitable for the second step of the present invention are for example styrene, α-methylstyrene, p-methylstyrene and other alkylstyrenes, acrylic and methacrylic acid and alkyl esters thereof, (meth)acrylamide, (meth)acrylonitrile, acrolein, vinyl acetate, vinyl chloride, conjugated dienes, vinylpyridines, maleic acid and the anhydride thereof and fumaric acid. Preferred monomers are styrene, α-methylstyrene and alkylstyrenes, acrylic and methacrylic acid and alkyl esters thereof, acrylonitrile, butadiene, isoprene and maleic anhydride. Particularly preferred monomers are acrylic and methacrylic acid and alkyl esters thereof, styrene, α-methylstyrene, acrylonitrile and butadiene.

In the process according to the invention nitrogen monoxide is for example either introduced into the reaction medium directly in the form of a nitrogen oxide gas or produced by an in situ reaction between a nitrite compound and an activator component. Suitable activator components are for example reducing or oxidizing agents or acids.

The gaseous nitrogen monoxide used in the process according to the invention can be used in pure form or as a mixture with gaseous nitrogen oxides of the general formula (II)

\[ \text{N}_x\text{O}_y \]  

(II)

in which

\[
\begin{align*}
\text{a) } & x \text{ is 1 and } y \text{ is 2 or} \\
\text{b) } & x \text{ is 2 and } y \text{ can have the values 1, 3, 4 or 5,}
\end{align*}
\]
in any desired ratios.

In a preferred embodiment of the invention nitrogen monoxide is produced in situ by a reaction between a nitrite compound and an activator component.

Suitable nitrite components are both inorganic and organic nitrite compounds. Inorganic nitrite compounds are for example compounds of the general formula (III),

$$Z(NO_2)_n$$  

(III)

in which

$Z$ is selected from the group comprising alkaline, alkaline earth, earth or transition metals or lanthanide or actinide or from one of the cations $Sn^{2+}$, $Sn^{4+}$, $In^{3+}$, $Ti^+$, $Ti^{3+}$, $Pb^{2+}$, $Bi^{3+}$, $Sb^{3+}$, $Sb^{5+}$ or $NH_4^+$ and

$n$ is a number from 1 to 5.

Suitable organic nitrite compounds are for example compounds of the general formula (IV),

$$R-O-N=O$$  

(IV)

in which

$R$ is an organic radical containing at least 2 carbon atoms and other functional groups, selected from the group comprising esters, thioesters, ethers, thioethers, alcohols, thiols, amines, amides, imines, imides, urethanes, halogen, nitro, nitroso, nitrates, nitrites, sulfates, sulphones, nitriles, carboxylic acids, carboxylic anhydrides, aldehydes and ketones.
Preferred compounds of formula (III) are for example water-soluble salts, where \( Z \) is a mono- or divalent cation. Particularly preferred are for example those where \( Z \) is an alkaline cation or NH\(_4^+\).

Preferred compounds of formula (IV) are for example those where \( R \) is an organic radical containing 2 to 6 carbon atoms. Particularly preferred compounds of formula IV are for example isoamyl nitrite, propyl nitrite, n-butyl nitrite or tert-butyl nitrite.

Suitable reducing agents for use as activators for the present invention are all known reducing agents, such as for example Fe(II), Cu(I) and Ti(III) salts, ascorbic acid, formic acid, tartaric acid, oxalic acid and salts thereof, alkaline hydroxymethanesulfinate salts, dextrose, formaldehyde, hydrogen peroxide, sodium sulfite, sodium and potassium thiosulfate, 2-mercaptoethanol and other thiols. In a preferred embodiment the reducing agents are Fe(II) salts, ascorbic acid, formic acid, oxalic acid, formaldehyde or dextrose. Particularly preferred reducing agents are Fe(II) sulfate, ascorbic acid and dextrose.

Suitable oxidants for use as activators for the present invention are for example Fe(III), Cr(III), Cu(II), Ti(IV), Sn(IV), Hg(II) salts, alkaline permanganates, alkaline chromates, hydrogen peroxide, alkaline hypochlorites. Preferred oxidants are Fe(III), Cu(II) and Cr(III) salts. Particularly preferred oxidants are Fe(III) and Cr(III) sulfates.

Suitable acids for use as activators are for example strong and weak inorganic and organic acids, such as for example hydrochloric acid, sulfuric acid, phosphoric acid, formic acid, ascorbic acid, acetic acid, tartaric acid, oxalic acid and methanesulfonic acid. In a preferred embodiment the acid is for example sulfuric acid, phosphoric acid, acetic acid or methanesulfonic acid. Particularly preferred acids are sulfuric and acetic acids.
The activator or mixture of activators is used in amounts of 0.01 to 500 mol %, preferably 0.1 to 300 mol %, most preferably 1 to 200 mol %, based on the nitrite component.

Suitable free-radical initiators for both steps of the present invention are any suitable agents producing free radicals, including precursors such as azo compounds, peroxides or peroxy esters, which generate radicals for example by thermolysis, or precursors such as styrene, which generate radicals by autopolymerization. Also suitable are redox initiating systems such as $K_2S_2O_8 + FeSO_4$, photochemical initiating systems or high energy radiation such as electron beam or X- or $\gamma$- radiation. In the latter case the molar amount of radicals generated in the polymerization medium represents the molar amount of initiator and can be calculated based on the total amount of radiation energy and the radical generation efficiency of the type of radiation employed.

Initiators can also be molecules containing free radicals such as oxygen, nitrogen dioxide and ozone.

Examples of free radical initiators generating radicals by thermolysis are $2,2'$-azo-bis(isobutyronitrile), $2,2'$-azo-bis(isovaleronitrile), $2,2'$-azo-bis(methyl isobutyratate), $4,4'$-azo-bis(4-cyanopentanoic acid), $1,1'$-azo-bis(1-cyclohexancarbonitrile), 2-tert-butylazo-2-cyanopropane, $2,2'$-azo-bis[2-methyl-N-(1,1-bis(hydroxymethyl)-2-hydroxyethyl) propionamide], $2,2'$-azo-bis[2-methyl-N-(2-hydroxyethyl) propionamide], $2,2'$-azo-bis(isobutyramidine hydrochloride), $2,2'$-azo-bis(N,N'-dimethylene isobutyramine), $2,2'$-azo-bis[2-methyl-N-(1,1-bis(hydroxymethyl)-2-ethyl) -propionamide], $2,2'$-azo-bis[2-methyl-N-(2-hydroxyethyl) propionamide], $2,2'$-azo-bis(isobutyramide) dihydrate, $2,2'$-azo-bis(2,2,4-trimethylpentane), $2,2'$-azo-bis(2-methylpropane), tert-butyl peroxyacetate, tert-butyl peroxybenzoate, tert-butyl peroxyoctoate, tert-butyl peroxydecanoate, tert-butyl peroxyisobutyrate, tert-amyl peroxypropionate, tert-butyl peroxypropionate, diisopropyl peroxydicarbonate, dicyclohexyl peroxydicarbonate, dicumyl peroxide, dibenzoyl peroxide, dilauroyl peroxide, potassium
peroxydisulfate, ammonium peroxysulfate, di-tert-butyl hyponitrite and dicumyl hyponitrite.

Initiators generating radicals by photolysis include for example benzoin derivatives, benzophenone, acyl phosphine oxides and photoredox systems.

Initiators generating radicals as a result of a redox reaction consist in general of a combination of an oxidant and a reducing agent. Suitable oxidants are for example potassium peroxysulfate, hydrogen peroxide, tert-butyl hydroperoxide, cumyl hydroperoxide, benzoyl peroxide and p-menthane hydroperoxide. Suitable reducing agents are for example Fe(II) salts, Ti(III) salts, potassium thiosulfate, potassium bisulfite, ascorbic acid and its salts, oxalic acid and salts thereof, hydrogen peroxide and dextrose.

The first step of the process according to the invention is carried out at a temperature between 0 and 120°C, preferably between 20 and 100°C, and most preferably between 40 and 80°C.

In the second step the polymerization is carried out in the presence of the intermediate product formed in the first step at temperatures between 20 and 180°C, preferably between 50 and 150°C, and most preferably between 60 and 130°C.

The monomer is added in an amount of at least 100 mol %, preferably at least 300 mol %, and most preferably at least 500 mol %, based on nitrogen monoxide.

Where the intermediate product is formed in a completely separate step from the polymerization step, such as for example when the intermediate product is isolated before further use, or where the monomer or monomer mixture to be polymerized is different from the monomer or monomer mixture used in the formation of the intermediate product, the monomer used in the formation of the intermediate product is
added in an amount of at most 15,000 mol %, preferably at most 10,000 mol %, and most preferably at most 5,000 mol %, based on nitrogen monoxide.

Where the intermediate product is formed in a "one-pot" process before the polymerization step is begun, the maximum amount of monomer or monomer mixture used in the first step may already be the amount of monomer required for the polymerization step and it depends on the molecular weight of the desired polymers. Typically this amount of monomer will be no more than 1,000,000 mol %, preferably no more than 500,000 mol %, and most preferably no more than 300,000 mol %, based on nitrogen monoxide.

If the monomer or monomer mixture to be polymerized is not present in the first step, the monomer or monomer mixture to be polymerized according to the invention is added in a later stage in amounts depending on the desired molecular weight of the polymer, and typically in amounts of between 500 and 1,000,000 mol %, preferably between 1,000 and 500,000 mol %, and most preferably between 2,000 and 300,000 mol %, based on the initiator used in the polymerization step.

The initiator compound is added in an amount of between 0.01 and 200 mol %, preferably between 0.05 and 150 mol %, and most preferably between 0.1 and 100 mol %, based on nitrogen monoxide.

The polymerization of the second step of the present invention is initiated with a free radical initiator or a mixture of free radical initiators, in amounts between 0.1 and 500 mol %, preferably between 0.5 and 200 mol %, and most preferably between 1 and 100 mol %, based on nitrogen monoxide used in the first step for the formation of the intermediate product.

The process may be carried out in the presence of an organic solvent or in the presence of water or in mixtures thereof. When a nitrite component is used as a source of nitrogen monoxide, water promotes the reaction between the nitrite component and
the activator component. Additional cosolvents or surfactants, such as glycols or salts of fatty acids or salts of quaternary ammonium compounds or ethylene-oxide based amphiphilic compounds, may be used.

If organic solvents are used, suitable solvents or mixtures of solvents are typically pure alkanes, aromatic hydrocarbons, halogenated hydrocarbons, alkanols, esters, ethers and ketones or mixtures thereof. Preferred solvents are for instance toluene, tetrahydrofurane, ethyl benzene, diethylether, hexane and methyl ethyl ketone.

Water can be used in the process of the present invention and both water-soluble monomers and water-insoluble monomers can be used. The type of polymerization used can be bulk, solution, emulsion, dispersion or suspension polymerization and it can be carried out both batchwise and continuously.

The present invention also relates to intermediate products of the general formula (I) which are obtainable by the process according to the invention.

The intermediate products according to the invention of the general formula (I) can be present as mixtures with oligomeric nitroso compounds.

The invention also relates to polymers of the general formula (V),

\[ \text{Init}^1 \left( \frac{M}{\text{Pol}} \right)^p \]
\[ \text{Init}^2 \left( \frac{M}{\text{Pol}} \right)^q \]

in which

\[ \text{Pol} \] is a polymeric radical,
\[ M \] represents any desired vinyl monomer,
p is an integer between 1 and 15,
q is an integer between 1 and 15 and

Init\textsuperscript{1} and Init\textsuperscript{2} are radicals of a free-radical initiator, wherein I\textsubscript{1} and I\textsubscript{2} can be identical or different within the compound.

The polymeric radical Pol contains monomer units selected from the monomers used for step 2. Pol can be a linear-, block-, random-, graft- or star-copolymer.

The polymers according to the invention display a polydispersity $D = \overline{M}_w / \overline{M}_n < 2$, preferably $D < 1.8$, and most preferably $D < 1.5$.

The present invention also relates to the use of the intermediate product of the general formula (I) in free-radical polymerization.

The present invention also relates to the use of polymers of the general formula (V) for the production of coatings and as compatibilizers in thermoplastic polymer blends.

The polymerization process according to the present invention has several benefits over the prior art:

Polymers with low polydispersity $D$ can be prepared compared to the high polydispersity observed in case of conventional free radical polymerization ($D > 2$). The polydispersity $D$ of the polymers synthesized according to the invention is usually lower than 2 and can be significantly lower at low monomer conversion rates ($< 30\%$).

Another aspect of the present invention is that the number average molecular weight of the polymer chains increases linearly with the monomer conversion, which allows a tailor-made polymer molecular weight to be obtained. Furthermore, the molecular weight of the polymers can be controlled by varying the amount of initiator and/or
the amount of nitrogen monoxide and/or the amount of monomers used in the reaction.

A further benefit of the present invention is that after removal of the non-polymerized monomers or after reaching a conversion of 100 %, a second polymerization step can be initiated simply by adding to the polymer synthesized in the first polymerization step a portion of fresh vinyl monomer or monomer mixture that can be different from the vinyl monomer or monomer mixture used in the first polymerization step. The polymerization of the vinyl monomer or monomer mixture added in the second step is then initiated by the polymer chains synthesized in the first polymerization step and di-block copolymers can for example be generated if the polymer chains synthesized in the first polymerization step were linear chains with one single growing site. The molecular weight and molecular weight distribution of each block can be controlled independently during the respective polymerization step. The process of polymerization of a vinyl monomer or monomer mixture initiated by polymer chains synthesized in a previous polymerization step carried out according to the invention can be repeated, whereby for example multi-block-copolymers of controlled molecular weight and molecular weight distribution for each block can be obtained.
Examples

In the examples the following abbreviations are used:

AIBN: azobisisobutyronitrile
BeMA: benzyl methacrylate
\textit{t-BMA}: tert-butyl methacrylate
MMA: methyl methacrylate

Controlled free-radical polymerization of tert-butyl methacrylate in the presence of sodium nitrite and iron(II) sulfate.

Example 1

\textit{In the presence of sodium nitrite and iron(II) sulfate:}

0.0414 g NaNO\textsubscript{2} (6 \textit{\times} 10\textsuperscript{-4} mol), 0.0834 g FeSO\textsubscript{4}.7H\textsubscript{2}O (3 \textit{\times} 10\textsuperscript{-4} mol) and 0.0492 g AIBN (3\textit{\times}10\textsuperscript{-4} mol) were introduced into a 50 ml reaction flask that had been evacuated by three nitrogen-vacuum cycles. 20 ml of degassed water were then added and the flask was thermostated at 80°C. Finally, 8 ml of t-BMA (5 \textit{\times} 10\textsuperscript{-2} mol) were added to the reaction flask. Samples were regularly removed from the polymerization medium and the % monomer conversion was calculated by \textit{\textsuperscript{1}H-NMR} analysis of the organic phase.

Comparative example 1

\textit{With sodium nitrite and without iron(II) sulfate:}

0.0492 g AIBN (3 \textit{\times} 10\textsuperscript{-4} mol) and 0.0414 g NaNO\textsubscript{2} (6 \textit{\times} 10\textsuperscript{-4} mol) were introduced into a 50 ml reaction flask that had been evacuated by three nitrogen-vacuum cycles. 20
ml of degassed water were then added and the flask was thermostated at 80°C. Finally, 8 ml of t-BMA (5 *10^{-2} mol) were added to the reaction flask. Samples were regularly removed from the polymerization medium and the % monomer conversion was calculated by ¹H-NMR analysis of the organic phase.

Comparative example 2

*Without sodium nitrite and without iron(II) sulfate:*

0.0492 g AIBN (3 *10^{-4} mol) were introduced into a 50 ml reaction flask that had been evacuated by three nitrogen-vacuum cycles. 20 ml of degassed water were then added and the flask was thermostated at 80°C. Finally, 8 ml of t-BMA (5 *10^{-2} mol) were added to the reaction flask. Samples were regularly removed from the polymerization medium and the % monomer conversion was calculated by ¹H-NMR analysis of the organic phase.

The molecular weight and polydispersity of the resulting polymers are displayed in Figures 1 and 2.

Without sodium nitrite (comparative example 1), the polymerization takes place very rapidly (55% monomer conversion after 30 min), a high molecular weight polymer is formed in the early stages of the polymerization, and the polydispersity is high, as is consistent with a non-controlled process.

Whenever sodium nitrite alone is added (comparative example 2), the polydispersity is slightly lower compared to the previous non-controlled polymerization, and the \( \bar{M}_n \) increases as the % monomer conversion increases. The polymerization rate is not basically different from that observed in the absence of sodium nitrite (41% monomer conversion after 30 min).
Only when both sodium nitrite and FeSO$_4$ are added to the polymerization medium (example 1) is nitrogen monoxide generated and the number average molecular weight increases linearly from the outset as the % monomer conversion increases, as is consistent with a controlled process. Also the molecular weight of the polymers synthesized is lower than in comparative examples 1 and 2. Furthermore, the polymerization carried out in the presence of sodium nitrite and FeSO$_4$ takes place more slowly than in the two previous cases (a monomer conversion of only 5% and 49% after 2 and 6h, respectively) and an induction period of 1 to 2 hours, corresponding to the formation of the intermediate product, is observed.

**Example 2**

*Controlled free-radical polymerization of tert-butyl methacrylate with a sodium nitrite/FeSO$_4$/K$_2$S$_2$O$_8$ system (effect of the amount of sodium nitrite):*

Experiment with a molar ratio of sodium nitrite to K$_2$S$_2$O$_8$ of 2:1.

0.0414 g of NaNO$_2$ (6 *10$^{-4}$ mol) and 0.0834 g of FeSO$_4$.7H$_2$O (3 *10$^{-4}$ mol) were introduced into a 50 ml reaction flask that had been evacuated by three nitrogen-vacuum cycles. 15 ml of degassed water and 8 ml of t-BMA (5 *10$^{-2}$ mol) were then added and the flask was thermostated at 80°C. Finally, 0.0811 g of K$_2$S$_2$O$_8$ (3 *10$^{-4}$ mol) in 5 ml of water were added. Samples were regularly removed from the polymerization medium and the monomer conversion was calculated by $^1$H-NMR analysis of the organic phase.

The same method is used for the other sodium nitrite/K$_2$S$_2$O$_8$ molar ratios by modifying the sodium nitrite amounts accordingly and keeping constant the amounts of the other constituents.
In Figure 3 the molecular weight of the resulting polymers is plotted as a function of the monomer conversion.

As can be seen from Figure 3, in all cases the molecular weight of the polymers increases linearly as the monomer conversion increases. In this case the molecular weight obtained at a given conversion can be controlled by adjusting the amount of sodium nitrite.

Example 3

Resumption experiment in the case of the polymerization of tert-butyl methacrylate

Synthesis of the Poly(t-BMA) macroinitiator:

0.1656 g NaNO₂ (2.4 *10⁻³ mol) were introduced into a 250 ml reaction flask that had been evacuated by three nitrogen-vacuum cycles. 40 ml of degassed water and 32 ml of t-BMA (2 *10⁻¹ mol) were then added, and the flask was thermostated at 80°C. When the temperature was reached a degassed solution of 0.4226 g ascorbic acid (2.4 *10⁻³ mol) in 20 ml water was added to the flask and finally a degassed solution of 0.3244 g K₂S₂O₈ (1.2 *10⁻³ mol) in 20 ml of water was added. After a reaction time of 1 h, the water and residual monomer were removed in vacuo and the intermediate product was dried overnight at room temperature. The conversion rate was between 5 and 10%.

Resumption experiment:

10 ml of t-BMA (6.3 *10⁻² mol) were added to the non-purified macroinitiator described above and the flask was placed in an oil bath at 50°C. Samples were regularly removed from the polymerization medium and the % monomer conversion was cal-
culated gravimetrically after drying at 80°C in vacuo, taking into account the amounts of macroinitiator initially introduced.

As can be seen from Figure 4, the molecular weight of the polymers increased linearly as the % monomer conversion increased and the polydispersity remained low. For conversions up to 40% the polydispersity D is very low, for example D < 1.5.

Example 4

*Controlled free-radical polymerization of tert-butyl methacrylate in the presence of sodium nitrite and iron(III) sulfate:*

0.0414 g NaNO₂ (6 *10⁻⁴ mol) and 0.0735 g Fe₂(SO₄)₃·5H₂O (1.5 *10⁻⁴ mol) were introduced into a 50 ml reaction flask that had been evacuated by three nitrogen-vacuum cycles. 15 ml of degassed water and 8 ml of t-BMA (5 *10⁻² mol) were then added and the flask was thermostated at 80°C. Finally, 0.0811 g of K₂S₂O₈ (3 *10⁻⁴ mol) in 5 ml of water were added. Samples were regularly removed from the polymerization medium and the % monomer conversion was calculated by ¹H-NMR analysis of the organic phase.

It can be seen from Figure 5 that the use of Fe(III) is equivalent to the use of Fe(II) for controlling the free-radical polymerization of vinyl monomers in the presence of nitrites.
Example 5

*Controlled free-radical polymerization of t-BMA in the presence of tert-butyl nitrite and 2-mercaptoethanol in an aqueous medium:*

In a 50 ml one-necked round bottom flask fitted with a three-way stopcock connected to either a nitrogen line or a vacuum pump were added 16 ml t-BMA (10^4 mol) and 11 ml water. The flask was then immersed in an oil bath thermostated at 80°C. Then 2 ml of a 0.6 mol/l degassed solution of 2-mercaptoethanol in toluene (1.2 *10^{-3} mol) and 2 ml of a 0.66 mol/l degassed solution of tert-butyl nitrite in toluene (1.3 *10^{-3} mol) were added to the flask. After a reaction time of 5 minutes the degassed initiator solution (0.1974 g K_2S_2O_8 in 5 ml of water; 7.3 *10^{-4} mol) was added to the reaction flask. Samples were regularly removed from the polymerization medium and the % monomer conversion was calculated by 1H-NMR. After the evaporation of the residual monomer and solvent, the polymer was dried in vacuo at 80°C.

Figure 6 shows that the combination of an organic nitrite and a reducing agent is also suitable for controlling the free-radical polymerization of vinyl monomers.

Example 6

*Controlled free-radical polymerization of t-BMA in the presence of sodium nitrite and acetic acid in an aqueous medium:*

0.0207 g NaN_2O_2 (3 *10^{-4} mol) were introduced into a 50 ml reaction flask that had been evacuated by three nitrogen-vacuum cycles. 10 ml of degassed water and 8 ml of t-BMA (5 *10^{-2} mol) were then added, and the flask was thermostated at 80°C. Finally 2.5 ml of a solution of 0.2 ml of acetic acid in 50 ml of water (1.75 *10^{-4} mol) were added, followed by a solution of 0.0811 g of K_2S_2O_8 in 5 ml of water (3 *10^{-4} mol)
mol). Samples were regularly removed from the polymerization medium and the monomer conversion was calculated by gravimetry after drying at 80°C in vacuo. Figure 7 shows the linear increase of the average molecular weight of the polymer as the % monomer conversion increases. This indicates the control of the free-radical polymerization in the presence of sodium nitrite and acetic acid.

Example 7

Reaction of methyl methacrylate (MMA) with sodium nitrite and sulfuric acid at 0°C. Controlled free-radical polymerization of MMA in the presence of intermediate compound I:

I) Synthesis of the intermediate compound I:
A solution of non-degassed sodium nitrite (12.903 g of sodium nitrite in 30 ml of water; 0.187 mol) was added to 20 ml of non-degassed MMA (0.187 mol) in 80 ml of diethyl ether at 0°C. The reaction flask was connected to a funnel containing a solution of 10 ml of concentrated sulfuric acid in 30 ml of water (0.177 mol) and sealed with a bubbling apparatus. Then, the sulfuric acid solution was added drop-wise to the solution of MMA. After a reaction time of 3h at 0°C, the organic phase was extracted by CH₂Cl₂ and washed three times with water. The organic phase was finally dried with MgSO₄, filtered and the solvent and residual monomer were eliminated in vacuo at room temperature. 5.14 g of intermediate compound I (white solid) was collected. In this case the initiator for the formation of the intermediate compound I consists of traces of nitrogen dioxide (1 % or less) produced by the decomposition of the sodium nitrite in the presence of sulfuric acid.

II) Polymerization:
0.067 g of AIBN (4.1 *10⁻⁴ mol) and 0.096 g of the intermediate compound I were introduced into the reaction flask that had been previously evacuated by three nitrogen-vacuum cycles. 6 ml of degassed toluene and 6 ml of MMA (5.64 g; 5.6 *10⁻²
mol) were then added to the flask that was thermostated at 60°C. Samples were regularly removed from the polymerization medium and the % monomer conversion was calculated gravimetrically after drying at 80°C in vacuo, taking into account the amounts of initiator and intermediate product I initially used.

Figure 8 shows the linear increase in the average molecular weight of the polymer as the % monomer conversion increases. This indicates the control of the free-radical polymerization.

Example 8

*Synthesis of intermediate compound II in the presence of NO-gas and controlled free-radical polymerization of t-BMA initiated by AIBN in the presence of intermediate compound II.*

*Synthesis of intermediate compound II:*

Nitrogen monoxide is synthesized according to the literature (*Blanchard, A.A.; Inorg. Syn. 1946, 2, 126*) by the dropwise addition over 2h of 64.4 ml of a 1M degassed solution of sodium nitrite in water \((6.4 \times 10^{-2} \text{ mol NaNO}_2)\) to 72 ml \((5.5 \times 10^{-3} \text{ mol FeSO}_4)\) of a degassed solution prepared by dissolving 278 g FeSO_4\(_4\)\(7\)H_2O and 55 ml 95 % H_2SO_4 in 1 litre of water. A slow stream \((100 \text{ ml/min})\) of nitrogen is passed through the reaction vessel in order to transport the non-purified nitrogen monoxide into 30 ml \((0.19 \text{ mol})\) of degassed t-BMA in a second vessel at room temperature. The monomer rapidly turns blue. After a reaction time of 3h, the residual monomer is eliminated in vacuo to leave 1.5 g of the intermediate compound II as a viscous light-green solid. Similar results are obtained if the monomer and the various solutions are not degassed.
Polymerization using the intermediate compound II:

In a typical procedure, 0.1 g of the intermediate compound II is introduced into a 50 ml reaction flask that had previously been evacuated by three nitrogen-vacuum cycles. 4 ml of degassed toluene and 5 ml of t-BMA (3.2 *10^{-2} mol) are then added to the flask that is thermostated at 60°C. Finally, 1 ml (3.4 *10^{-4} mol) of a solution of 0.555 g AIBN in 10 ml toluene is added to the reaction flask. Samples are regularly removed from the polymerization medium and the monomer conversion is calculated gravimetrically after drying at 80°C in vacuo, taking into account the amounts of initiator and intermediate product II initially used. The plots of experimentally determined $M_n$ versus conversion (Figure 9) and ln[$M_n$/[M]] versus time (Figure 10) are linear, which indicate that the polymerization is controlled.

Example 9

Synthesis of intermediate compound III in the presence of NO-gas and controlled free-radical polymerization of t-BMA initiated by AIBN in the presence of intermediate compound III:

Using the same method as in example 1 but using MMA instead of t-BMA in the first reaction step (synthesis of intermediate compound III (white solid) instead of intermediate compound II), the results reported in Figure 11 and 12 are obtained for the polymerization of t-BMA.

Example 10

Block copolymer poly(phenylmethacrylate-b-t-butyl methacrylate):

1) Polybenzylmethacrylate macroinitiator
0.5 g of the intermediate compound III from example 9 is introduced into a 50 ml reaction flask that had previously been evacuated by three nitrogen-vacuum cycles. 10 ml of degassed toluene and 10 ml of BeMA (3.2 \*10^{-2} \text{ mol}) are then added to the flask that is thermostated at 60°C. Finally, 0.0984 g (3.4 \*10^{-4} \text{ mol}) AIBN are added to the reaction flask. After a reaction time of 17 hours the resulting polymer is recovered and purified by precipitating twice in heptane and drying 24h in vacuo at room temperature. 1.8 g (20 % conversion) of PBeMA (\( \bar{M}_n = 16000 \), D= 1.8) are obtained.

2) Resumption with t-BMA

0.2 g of the PBeMA obtained in step 1 were dissolved in 2 ml toluene and 2 ml t-BMA. The system was heated at 80 °C and the polymerization was carried out for 18 h. After precipitation in heptane 1.7 g of a white polymer displaying \( \bar{M}_n = 70000 \) and D= 2.7 are obtained (85 % conversion for the second monomer). SEC analysis with UV and refractive index detectors shows that the molecular weight distribution is bimodal and that part of the PBeMA from step 1 initiated the polymerization of t-BMA with formation of a poly(benzylmethacrylate-b-t-butylmethacrylate) block copolymer. The second peak in the SEC-chromatogram is due to PBeMA deactivated during the purification step from step 1 and accounts for the increased polydispersity of the crude polymerization product from step 2.
Claims


2. A process according to Claim 1, characterized in that in a first step a free-radical intermediate product of the general formula (I),

\[
\text{Init}^1 \left\{ \begin{array}{c}
M \\
p
\end{array} \right\} \text{p}
\]

\[
\text{Init}^2 \left\{ \begin{array}{c}
M \\
q
\end{array} \right\} N-O.
\]

in which

\(M\) represents any desired vinyl monomer,
\(p\) is an integer between 1 and 15,
\(q\) is an integer between 1 and 15 and
\(\text{Init}^1\) and \(\text{Init}^2\) are organic radicals of a free-radical initiator, wherein \(I_1\) and \(I_2\) can be identical or different within the compound,

is produced by reacting at least one monomer with nitrogen monoxide and at least one initiator and then in a second step the polymerization is carried out optionally with the addition of additional monomer or monomer mixture and/or free-radical initiator.

3. A process according to Claim 1, characterized in that the process is carried out in the form of a one-pot synthesis.
4. A process according to Claim 1, characterized in that nitrogen monoxide is introduced either directly into the reaction medium in the form of a nitrogen oxide gas or it is produced by an in situ reaction between a nitrite compound and an activator component.

5. A process according to Claim 4, characterized in that nitrogen monoxide is produced in situ by a reaction between a nitrite compound and an activator component.

6. A process according to Claim 4, characterized in that the activator components are reducing or oxidizing agents or acids.

7. A process according to Claim 4, characterized in that gaseous nitrogen monoxide is used in pure form or as a mixture with gaseous nitrogen oxides of the general formula (II)

\[ \text{N}_x\text{O}_y \]  \hspace{1cm} (II)

in which

a) \( x \) is 1 and \( y \) is 2 or

b) \( x \) is 2 and \( y \) can have the values 1, 3, 4 and 5,

in any desired ratios.

8. A process according to Claim 5, characterized in that the nitrite components are either inorganic or organic nitrite compounds.

9. A process according to Claim 8, characterized in that the inorganic nitrite compounds are compounds of the general formula (III)

\[ \text{Z(NO}_2)_n \]  \hspace{1cm} (III)
in which

\[ Z \text{ is selected from the group comprising alkaline, alkaline earth, earth or transition metals or lanthanide or actinide or from one of the cations} \]

\[ \text{Sn}^{2+}, \text{Sn}^{4+}, \text{In}^{3+}, \text{Tl}^+, \text{Tl}^{3+}, \text{Pb}^{2+}, \text{Bi}^{3+}, \text{Sb}^{5+}, \text{Sb}^{5+} \text{ or NH}_4^+ \text{ and} \]

\[ n \text{ is a number from 1 to 5.} \]

10. A process according to Claim 8, characterized in that the organic nitrite compounds are compounds of the general formula (IV),

\[ R-O-N=O \quad (IV) \]

in which

\[ R \text{ is an organic radical containing at least } 2 \text{ carbon atoms and other functional groups, selected from the group comprising esters, thioesters, ethers, thioethers, alcohols, thiols, amines, amides, imines, imides, urethanes, halogen, nitro, nitroso, nitrates, nitrites, sulfates, sulphones, nitrile, carboxylic acids, carboxylic anhydrides, aldehydes and ketones.} \]

11. Intermediate products of the general formula (I),

\[ \text{Init}^1 \text{—} \left( \text{M} \right)_p \]
\[ \text{N—O—} \quad (I) \]
\[ \text{Init}^2 \text{—} \left( \text{M} \right)_q \]

in which
M represents any desired vinyl monomer,

\( p \) is an integer between 1 and 15,

\( q \) is an integer between 1 and 15 and

\[ \text{Init}^1 \text{ and } \text{Init}^2 \text{ are radicals of a free radical initiator, wherein } I_1 \text{ and } I_2 \text{ can be identical or different within the compound,} \]

which are obtainable according to Claims 1 to 10.

10 12. Polymers of the general formula (V),

in which

\[ \text{Init}^1 - (M)^p \]

\[ \text{N} \longrightarrow \text{O} \longrightarrow \text{Pol} \quad (V) \]

\[ \text{Init}^2 - (M)^q \]

15 \( \text{Pol} \) is a polymeric radical,

\( M \) represents any desired vinyl monomer,

\( p \) is an integer between 1 and 15,

\( q \) is an integer between 1 and 15 and

\( I_1 \) and \( I_2 \) are radicals of a free radical initiator, wherein \( I_1 \) and \( I_2 \) can be identical or different within the compound,

which are obtainable according to Claims 1 to 10.

20 13. Polymers according to Claim 12 having a polydispersity \( D = \bar{M}_w / \bar{M}_n \) of lower than two.
14. Use of the free-radical intermediate product according to Claim 11 in free-radical polymerization.

15. Use of the polymers according to Claim 12 for the production of coatings and as compatibilizers in thermoplastic polymer blends.
**Figure 1**

![Graph showing Mn,SEC vs conversion (%)]

- ▼ AIBN
- ■ AIBN/NaNO₂
- ● AIBN/NaNO₂/FeSO₄

**Figure 2**

![Graph showing Mw/Mn vs conversion (%)]

- ▼ AIBN
- ■ AIBN/NaNO₂
- ● AIBN/NaNO₂/FeSO₄