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Influence of redox initiator component ratios on the emulsion copolymerisation of vinyl acetate and neodecanoic acid vinyl ester†

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Redox initiated emulsion polymerisation of vinyl acetate and neodecanoic acid vinyl ester was investigated at temperatures ranging from $-1\text{ }^{\circ}\text{C}$ to $87\text{ }^{\circ}\text{C}$ (initiation temperature between $-1\text{ }^{\circ}\text{C}$ and $60\text{ }^{\circ}\text{C}$), using varying molar ratios of the following redox components: L-ascorbic acid, *tert*-butyl hydroperoxide and ammonium iron(III) sulfate dodecahydrate as a catalyst. The high flexibility of redox initiators enables product properties, as well as space-time-yield, to be adjusted as required. Polymers being products by process, it was presumed that modifying the conversion rate would lead to a different product. However, it was shown that the reaction rate is adjustable by varying the catalyst amount without changing the product properties, such as molecular weight, particle size, glass transition temperature and polymer structure, while reducing the overall process time by 40–86% (at equimolar ratios of reducing and oxidising agent). In contrast, variation of the *tert*-butyl hydroperoxide content resulted in changes of the molecular weight. The influence of the initiation temperature and of the redox system on the reaction rate was determined, enabling control over the reaction rate in the whole temperature range. Meanwhile, overall process times of approximately 2–240 min and high conversions of 90–99% could be achieved. Statistical modelling confirmed the results and facilitated predictions, enabling the conversion rate to be adjusted to the desired properties. The possibility of being able to adjust the conversion rate and product properties independently of each other creates additional degrees of freedom in process design.

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1. Introduction

Poly vinyl acetate is widely produced *via* bulk, solvent and emulsion polymerisation for use in many applications, for example, in paint or adhesives.^{1–5} However, though polymerisation processes in bulk or solvent may still be necessary to obtain the best adhesive properties, they are less suited for paint and varnish, as monomer and organic solvent residues can be harmful to users, as well as to the environment. The emulsion polymerisation process uses water as a solvent and is suitable for any application where the polymer is applied as a latex dispersion.⁶ Therefore, emulsion polymerisation is the most commonly used process for the production of water-borne latex polymers.^{7–11} The importance of emulsion polymerisation in industry and in a continuously growing field, leads to a high demand for different process optimisation strategies. Most of

the radical based emulsion polymerisation processes are initiated by using thermal initiators.^{12–16} Due to the high temperatures required to achieve sufficient radical generation, while at the same time staying below the boiling point of water, the remaining temperature range required for the process is limited ($60\text{--}95\text{ }^{\circ}\text{C}$), unless pressure is applied to increase the temperature even further.^{12,17–19} The initiation temperature (ϑ_0) mainly affects the conversion rate of the polymerisation and therefore impacts the number of particles, as well as the resulting product properties, such as the molecular weight and its distribution. According to Lovell and Schork, reducing initiation temperature (ϑ_0) also lowers the rate of chain transfer into polymer, as the activation energy is higher than the propagation energy, reducing the possibility of side chain reactions.^{19,20} In order to extend the accessible temperature range to lower temperatures, and to broaden the variety of final product properties, water soluble redox initiators have been introduced to emulsion polymerisation. Redox initiation enables the generation of free radicals by a one-electron transfer reaction, using an oxidising and reducing agent, sometimes complemented by a catalyst, depending on the system. The redox initiation results in a sufficient radical generation rate at milder conditions, since the activation energy is significantly lower compared to thermal initiators (only $40\text{--}80\text{ kJ mol}^{-1}$ instead of $125\text{--}60\text{ kJ mol}^{-1}$).²⁰

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One substantial optimisation aim in emulsion polymerisation is the increase in space-time-yield, while, at the same time, keeping the product properties constant. However, conversion rate and product properties cannot be adjusted independently of one another when using conventional thermal initiators, since variation of temperature commonly results in changes in the molecular weight (higher molecular weights at lower temperatures).^{19,21} The use of redox initiators is promising in this regard, as they offer the possibility of adjusting the radical generation rate (and thus the conversion rate of the polymerisation) by varying the component ratios and environmental settings (such as pH value), while keeping the initiation temperature constant. It has indeed been shown that the initiation temperature of emulsion polymerisations (*e.g.* of methyl methacrylate, styrene and butyl acrylate) can be lowered to 40 °C, without significantly affecting product properties, such as particle size or conversion.^{22–24} A further decrease of the process initiation temperature down to 30 °C and room temperature was achieved in the production of polyaniline.^{25,26} Reducing the initiation temperature without altering the product contributes positively to process intensification and process safety.^{27–29} For example, reducing the temperature can thereby result in a reduction of the production costs, since less heating of the reactor is required (which saves energy). Slowing down the conversion rate through process optimisation also reduces the risk of leakage when approaching the boiling point of the chemicals used, therefore increasing the safety of employees or workforce.³⁰ Yet, until now, very few emulsion polymerisations have been successfully performed at lower temperatures while achieving high overall conversions. For instance, although the reaction temperature for poly vinyl acetate was reduced to the range of 8–22 °C by using a redox system composed of potassium persulfate and dimethylethanolamine, the overall conversion was limited to 48–88% after 4 h of reaction, therefore resulting in a comparatively low space-time-yield.^{31,32} Considering that the emulsion polymerisation of poly vinyl acetate at 60 °C can achieve a conversion of 99% after 15 min, 4 h of reaction for less conversion cannot be considered an efficient alternative.^{33,34} Even the use of Macro RAFT agents in combination with ascorbic acid and *tert*-butyl hydroperoxide at 25 °C sufficed only for a conversion of 55–89% after 24–48 h reaction time.³⁵ Since redox initiated emulsion polymerisation at low temperatures has only scarcely been investigated, it is of great interest to expand research in this direction.

In addition to the benefits that result from the extension of an applicable temperature range, the use of redox initiator systems considerably widens the possibilities of affecting the radical generation rate and therefore control of the whole polymerisation kinetics. For instance, adjustments of the pH, the type of peroxide, the ratio of oxidant to reducing agent, the amount of oxygen, the amount of catalyst, as well as the type and amount of other metal ions present in the dispersion can all affect the radical generation.³⁶ While the above factors result in a high flexibility of the systems, they demand precise reaction control and fundamental understanding of the redox initiator kinetics under different reaction conditions.³⁷

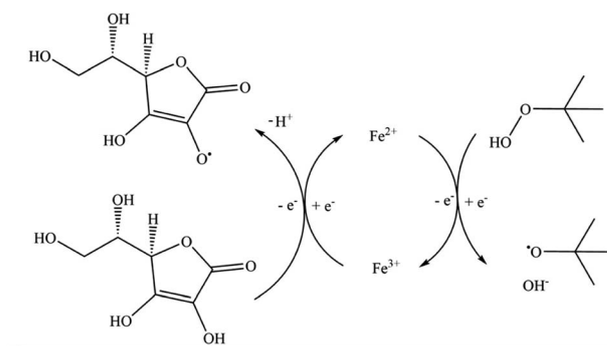


Fig. 1 Schematic representation of the reaction mechanism of the redox system (AsAc/tBHP/Fe_{cat.}).³⁷

In this work, we have chosen to apply a redox initiator consisting of L-ascorbic acid (AsAc) as reducing agent, ammonium iron(III) sulfate dodecahydrate (Fe-cat.) as catalyst and *tert*-butyl hydroperoxide (*t*BHP) as oxidising and radical generating agent, since AsAc based systems have proven to be successful for emulsion polymerisation in the past, not only in batch or semi-batch but also in continuous processes (see Fig. 1).^{21,33–35} Although the reaction mechanism has not yet been fully proven, Fig. 1 shows a formal reaction scheme based on the iron catalyzed reaction of H₂O₂ and Fe(II) as well as AsAc with Fe(III).^{37–39}

The monomer system chosen for this study consists of vinyl acetate and neodecanoic acid vinyl ester (Versa@10), since vinyl acetate based latices are industrially relevant and have already been investigated in conjunction with the chosen redox components.^{34,37,40,41} Previous studies showed, that controlling the decomposition kinetic of ascorbic acid permits control of the reaction rate of emulsion polymerisation of vinyl acetate and Versa@10 in the temperature range of 10–70 °C.³⁷ While this evidence indicates a high flexibility and achievable process control, the effect of reaction rate and temperature on the properties of the produced products were not documented.

There are different ways of influencing the process, either through modifying the set-up, the process parameters, or the kinetics of the reaction. In this work, the focus was placed on the influence of the reaction kinetics on the obtained product properties, so that the reactor set-up, as well as the different parameters, such as chemical compounds and stirrer speed, were kept constant. Ensuring consistent product qualities requires not only a thorough analysis of the final product, but the process must also be robust and reproducible. Therefore, most factors influencing the radical generation were also kept constant, and the focus was placed on the ratio of ascorbic acid to peroxide, on the amount of catalyst and on the initiation temperature.

The aim of this work is to significantly lower the initiation temperature, while achieving a full conversion and maintaining control over the desired product properties, such as the molecular weight (MW) and distribution (MWD), the particle size, or the glass transition temperature (*T*_g). Additionally, sufficient reaction rates and their control should be achieved in the whole temperature range.



Table 1 Overview of the used chemicals, their supplier and purity

Component	Abbreviation	Supplier	Purity
Vinyl acetate	VA	Wacker Chemie AG	99%
Versa@10	—	Wacker Chemie AG	99%
Mowiol 4-88	—	Sigma-Aldrich	MQ200
L-Ascorbic acid	AsAc	Sigma-Aldrich	ACS
tert-Butyl hydroperoxide	tBHP	Sigma-Aldrich	MQ200
Ammonium iron(III) sulfate dodecahydrate	Fe-cat.	Merck KGaA	ACS

The influence of the initiation temperature and of the amount of tBHP and Fe-catalyst is quantified in terms of product properties and polymerisation rate in order to achieve better process control in the whole temperature range from -1 to 87 °C.

2. Materials and methods

All chemicals were used directly and without further purification (Table 1).

The initial charge (Table 2) of the reaction was first degassed with nitrogen for at least 30 minutes. The gas flow was fed into the initial charge through a metal pipe with a diameter of 2 mm and with a flow of approximately 120 bubbles per minute. The initial charge was stirred at 300 rpm during degassing. The finished emulsion was continually flushed with nitrogen throughout the entire reaction to avoid interferences due to oxygen influx. The amount of water evacuated by the continuous nitrogen flow was not taken into account, as it affects only 0.5–1.5% of the total amount, depending on the initiation temperature and reaction time.

2.1. Emulsion polymerisation – 0.5 L RC1mx calorimeter

The emulsion polymerisations were carried out in a 0.5 L RC1mx glass calorimeter of Mettler Toledo as a batch process at 1 bar. The process was controlled *via* the IControl 6.1 software.

The reactor is 16.5 cm deep, has a diameter of 6.88 cm and a dish-like bottom with the sample outlet in the middle. The reaction solution was stirred with a 45° pitched six blade stainless steel turbine with a diameter of 4.5 cm and at a stirring rate of 300 rpm. The stirrer was located in the center and 3.5 cm above the bottom of the reactor (*cf.* Fig. 2). The reactor was filled with 312 mL demineralised water with a conductivity of 0.8 uS cm^{-1} , 80 g of a monomer mixture consisting of vinyl acetate and Versa@10 in a molar ratio of 9 : 1 and 15 g L^{-1} Mowiol 4-88 as an emulsifier, representing 7.5 wt% of the monomer amount (Table 2). Prior to initiation, the emulsion was heated/cooled to the initiation temperatures of -1 , 5, 15, 25 and 60 °C. The temperature was controlled by the RC1mx calorimeter and supported by a Julabo FP50 Cryostat.

A redox initiator system was used to initiate the polymerisation reaction. It consisted of L-ascorbic acid (AsAc) acting as reducing agent, tert-butyl hydroperoxide (tBHP, 70%) acting as an oxidising agent and ammonium iron(III) sulfate dodecahydrate (Fe-cat.) as a catalyst. AsAc was previously dissolved in 3 g of demineralised water. tBHP was used without further modification and the Fe-cat. was dissolved in 1 g of demineralised water. The oxygen brought into the reaction through the initiation was insignificant due to the small amount compared to the initial charge and due to the continuous degassing throughout the reaction.

All three redox initiator components were each added as a one shot by using a polyethylene syringe from Henke-Sass, Wolf GmbH.

The amounts of oxidising agent and catalyst were varied, whereas the amount of AsAc was kept constant (Table 3). In the following Table 3, the molar ratio between the components AsAc:tBHP:Fe-cat. is defined as r_{redox} . The ratio of the Fe-cat. component is defined as $r_{\text{Fe-cat.}}$ and the ratio of tBHP as r_{tBHP} . All ratios are given with respect to the component AsAc.

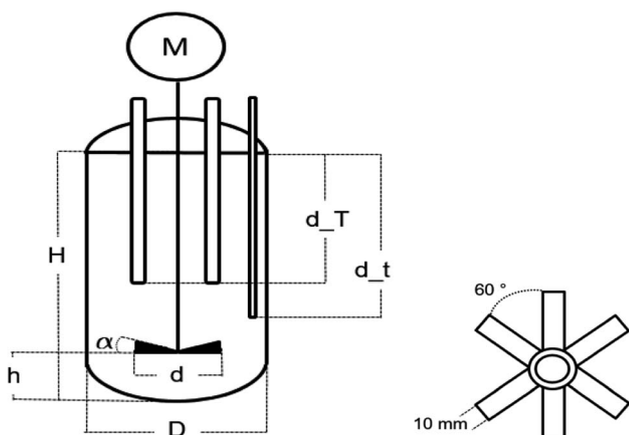


Fig. 2 Schematic representation of the 0.5 L RC1mx glass calorimeter with two temperature sensors and a gas supply, and one of the stirrer (on the right). The specified dimensions are: $H = 165.0$ mm, $h = 35.0$ mm, $D = 68.8$ mm, $d = 45.0$ mm, $\alpha = 45^\circ$, $d_T = 50$ mm, $d_t = 70$ mm, $\varnothing_T = 12.0$ mm and $\varnothing_t = 3.0$ mm.

Table 2 Overview of the initial charge of the redox initiated emulsion polymerisation in batch process. Total amount: 400 g

Component	Water	Vinyl acetate/ Versa@10	Mowiol 4-88	AsAc
Molar ratio		9 : 1		
Mass [g]	312	80	6	0.83
Concentration [g L^{-1}]	—	—	15	2.08
wt% (based on total mass)	78	20	1.5	0.2
wt% (based on monomer)	—	—	7.5	1.04



Table 3 Molar ratio and amounts of redox components used for the emulsion polymerisation

r_{redox}^a	t_{BHP} (g L ⁻¹)	Fe (g L ⁻¹)
1 : 0.7 : 0.01	1.08	0.03
1 : 1.3 : 0.01	1.98	0.03
1 : 1 : 0.003	1.52	0.01
1 : 1 : 0.01	1.52	0.03
1 : 1 : 0.03	1.52	0.09

^a Concentration of AsAc in each batch = 2.08 g L⁻¹. Each polymerisation was performed at least twice to ensure the reproducibility of the results.

2.2. Determination of conversion

To determine the conversion of each reaction, a sample was taken at the end of each reaction through the outlet at the bottom of the reactor, inhibited with a 7 wt% hydroquinone solution, and then measured by gas chromatography (GC). In addition to the offline determination methods, an inline monitoring of the amount of monomer was performed by Raman spectroscopy.

The end of the reaction was defined by the time at which the temperature of the reactor and the temperature of the jacket remained constant at the initial temperature and confirmed by Raman spectroscopy.

2.2.1. Gas chromatography. The samples were measured with an Agilent 7820A using hydrogen as a carrier gas (column: CP-Sil 5CB fused silica, 30 m, 1.0 μm, detector: FID, injector temperature: 200 °C, detector temperature: 250 °C, sample volume: 0.4 μL). After the sample was taken through the outlet of the reactor, 500 mg were withdrawn with a 100–1000 μL Eppendorf research micropipette, weighed, and then dissolved in 5 mL of *N,N*-dimethylacetamide. Approx. 70 mg toluene were added and used as an internal standard. Once fully prepared and dissolved, 1.5 mL of the solution were transferred into an amber glass vial, which was then sealed with a PTFE/silicone septum. The device has an accuracy of ±5%.

2.2.2. Raman spectroscopy. Inline monitoring of the conversion was performed *via* Raman spectroscopy (RamanRxn1-785 system, Kaiser Optical Systems, IO-1/2S-NIR probe, laser power 10.1 mW at the probe, Software: ICRaman). Spectra were recorded every 32 seconds with an integration time of 30 seconds throughout the entire process.

The signal at 1650 cm⁻¹ shows an overlap of the C=C bond of both monomers, vinyl acetate and Versa®10, and can therefore be tracked during the reaction to determine the total monomer conversion.⁴²

2.2.3. Reaction calorimetry. The thermal conversion was determined by the IControl 6.1 software (Mettler Toledo) and based on the RTCal calibration of the device (RC1mx, Mettler Toledo). The temperature of the reactor and of the jacket were tracked during the experiment and the reaction enthalpy was calculated based on the previously performed calibration. The thermal conversion was calculated according to eqn (1), where $\Delta H_{\text{R}}(t)$ is the standard enthalpy of the reaction at time t , $Q_{\text{R}}Q^{\text{R}}$ the heat transfer coefficient of the reaction and $Q_{\text{baseline}}Q^{\text{baseline}}$ that of the baseline.

$$\alpha(t) = \frac{\Delta H_{\text{R}}(t)}{\Delta H_{\text{R}}(t_{\text{ende}})} = \frac{\int_{t_0}^t (\dot{Q}_{\text{R}} - \dot{Q}_{\text{baseline}}) dt}{\int_{t_0}^{t_{\text{ende}}} (\dot{Q}_{\text{R}} - \dot{Q}_{\text{baseline}}) dt} \times 100 \quad (1)$$

2.3. Determination of mean particle size

The mean particle size was determined by dynamic light scattering (Zetasizer Nano ZS, Malvern Instruments). One drop of the sample previously used to determine the conversion was diluted in 5 mL demineralised water and then measured in a polyethylene UV-cuvette. Each particle size presented in the following work was obtained by a triple determination each consisting of 18 measurements at 25 °C.

The refractive index (1.48) and density (1.167 g cm⁻³) of the copolymer were determined in a previous work and implemented in the dynamic light scattering software for all measurements.³⁴

2.4. Determination of glass transition temperature

The T_{g} was determined by differential scanning calorimetry (DSC 1, Mettler Toledo). Approximately 3 g of the sample were dried in an aluminium sample pan in a drying cabinet at 60 °C for five days. 7–14 mg of the dried polymer were weighed into the sample crucible and measured with a temperature gradient of -30 °C to +250 °C. The heating rate was 10 °C min⁻¹ and the cooling rate was 20 °C min⁻¹. The accuracy of the measurements is approximately ±2 °C.⁴³

2.5. Determination of molecular weight distribution

The molecular weight distribution was determined by gel permeation chromatography at room temperature using a Knauer Autosampler Smartline 3800, a Knauer K-4002 2-channel Degasser, an AI-12-13 intelligent pump from Flom, an RI 2000 Detector from Schambeck SFD GmbH and three columns (one PLgel 10 μm guard column and two PLgel 10 μm mixed-B) from Agilent. Tetrahydrofuran was used as eluent (flow rate: 1 mL min⁻¹) and the device was calibrated with different molecular weight distributions of linear polystyrene. Adjusting the obtained molecular weight distribution by using the universal calibration and the Mark–Houwink equation, with the parameters summarised in Table 4, resulted in an increase of the molecular weight by approximately 5%.^{44–47}

For the measurements, 200 mg of the sample, previously homogenised by shaking, were withdrawn with a 100–1000 μL Eppendorf research micropipette and then dissolved in 5 mL

Table 4 Mark–Houwink parameters for the universal calibration of molecular weights⁴⁷

	a	K [10 ⁻⁵ cm ³ g ⁻¹]
Poly styrene	0.703	16
Polyvinyl acetate	0.791	5.1



Table 5 Set temperature sections of the Minimal Film Forming Temperature (MFFT) measurements

Temperature section	1	2	3	4	5	6	7	8	9	10
Temperature [°C]	12.6	13.8	14.9	16	17.1	18.2	18.9	20.1	21.1	22

N,N-dimethylacetamide. After complete dissolution of the polymer, 1.5 mL of the solution were transferred into an amber glass vial and sealed with a PTFE/silicone septum.

2.6. Determination of minimal film forming temperature (MFFT)

The minimal film forming temperature (MFFT) was determined using an MFFT 10 device from Coesfeld Materialtest.

The MFFT was determined by spreading a 0.3 mm thick film onto the device, which was previously set to a temperature gradient ranging from 12.6 °C to 22.0 °C. The section where the previously applied and subsequently dried latex formed an even and transparent film was identified as the MFFT. The determination of the MFFT through this method has an accuracy of ± 1 section, which, in this case, is equal to approximately ± 2 °C.

The different temperature sections of the gradient are summarised in Table 5.

2.7. Nuclear magnetic resonance (NMR) spectroscopy analysis

^1H and 2D COSY NMR spectra were measured with an AV4001 spectrometer (Bruker Biospin GmbH, 400 MHz, 295 K). 10–13 mg of the dried polymer (previously dried for T_g measurements) were dissolved in 0.8 mL of DMSO- d_6 and measured in a disposable NMR tube (Deutero BORO eco 5 mm 7"). The spectra were analysed using MestReNova 14.0.1.

2.8. Determination of the number of particles

The number of particles (N_p) was calculated based on the total conversion and on the average particle size (d_p) at each temperature. The following equation was used (eqn (2))

$$N_p \left(\frac{\text{particle}}{\text{L}} \right) = \frac{3\alpha(t) g_{\text{monomer}}}{4\pi\rho \left(\frac{d_p}{2} \right)^3} 10^{21} \frac{1}{V_R} \quad (2)$$

where $\alpha(t)$ is the monomer conversion determined as in eqn (1) (–), g_{monomer} the initial total monomer weight (g), ρ the density of the copolymer (g cm^{-3}),⁴⁸ d_p the particle size (nm) and V_R the total batch volume (L).

3. Results and discussion

3.1. Conversion

All experiments were tracked inline *via* Raman spectroscopy (*cf.* Appendix Fig. S1† for Representative Raman spectrum) and *via* reaction calorimetry to determine whether the monomer was fully consumed during the reaction. After completion of the reaction, a sample of the dispersion was taken and analysed by gas chromatography to confirm the final conversion of the emulsion polymerisation. In order to validate these methods, both methods were compared to the thermal conversion and assessed as to whether they were all suitable for an adequate representation of the conversion.

Fig. 3 shows an overview of the normalised thermal and Raman conversion at different initiation temperatures, as well

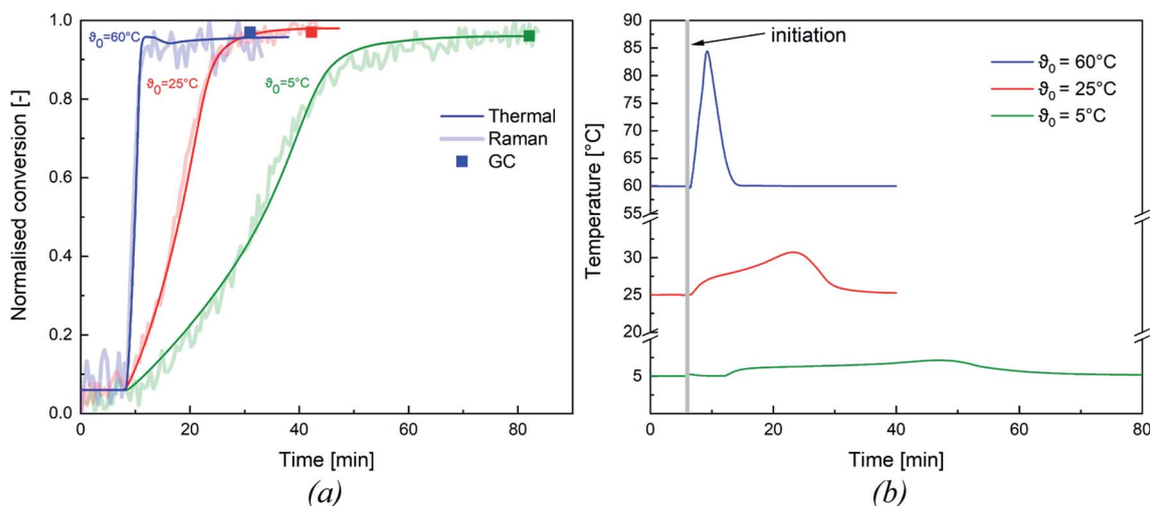


Fig. 3 Overview of the normalised thermal, Raman, and GC conversions (a) and temperature profile of the emulsion polymerisation (b) at a constant molar ratio of 1 : 0.7 : 0.01 and at different initiation temperatures of $\vartheta_0 = 5$ °C (green), $\vartheta_0 = 25$ °C (red) and $\vartheta_0 = 60$ °C (blue).



Table 6 Overview of the average conversion of all reactions determined by GC and given in %

r_{redox}	ϑ_0 [°C]				
	60	25	15	5	-1
1 : 0.7 : 0.01	95.5	98.1	94.3	95.5	90.0
1 : 1.3 : 0.01	99.0	99.4	99.6	99.7	—
1 : 1 : 0.01	98.4	99.4	99.4	99.1	—
1 : 1 : 0.03	97.1	98.0	98.5	99.1	99.1
1 : 1 : 0.003	99.1	99.8	99.6	99.4	—

as the corresponding conversion determined by GC. Both trends concur and the conversion determined by GC also matches the inline conversions. To simplify the following analysis of our results, we have focused on the thermal conversion, as this method could be validated by the other two methods. An overview of the thermal conversion of all emulsion polymerisations considered in this work can be found in the ESI (cf. Appendix Fig. S2†).

While the presented example clearly demonstrates the influence of the initiation temperature on the reaction rate at constant redox initiator component ratios, the following sections further analyses the factors influencing polymerisation kinetics and product properties.

The conversion of all experiments are summarised in Table 6: all experiments show a near full conversion in the whole temperature range from $\vartheta_0 = 60$ °C to $\vartheta_0 = -1$ °C. It is therefore possible to achieve full conversion in reasonable time, even at a considerably reduced initiation temperature (cf. Fig. 3, conversion >95% after 80 min at $\vartheta_0 = 5$ °C). While no significant influence of ϑ_0 and $r_{\text{Fe-cat}}$ was observed, slightly lower conversions (average = $94.7 \pm 3.0\%$) were obtained at $r_{\text{tBHP}} = 0.7$. Since emulsion polymerisation is a complex process and overall conversion can be influenced by many factors (e.g. mixing,

initiation, emulsifier type & content, temperature, choice of monomer, etc.), a direct comparison of our results with other works is possible only to a limited extend. For instance, slightly lower conversions were achieved in a continuous process (residence time 5–15 min) where the same material system was polymerised in a tubular reactor: increasing the monomer content from 20 wt% to 40 wt% resulted in a decrease of the average conversion from 92% to 81%.³³ Guan *et al.* obtained conversions in the range of 48–88%, but required longer reaction times (240 min) for the redox initiated emulsion polymerisation of vinyl acetate, using a potassium persulfate/*N,N*-dimethylethanolamine based initiator system.³¹ However, they did identify a significant influence of the reaction temperature on the overall conversion. This would indicate a significant advantage in using AsAc based redox systems compared to the example above. Guan *et al.* also noted an influence of the emulsifier content, which, however, was kept constant in this work and therefore cannot be verified.

In the context of this work, we can conclude that the initiator provides a sufficient radical generation rate at all initiation temperatures. Furthermore, at least an equimolar ratio of tBHP and AsAc should be used to maximise the overall conversion. However, a high conversion is not the only criterion to assess a successful polymerisation at lower temperatures. It is also essential that the obtained product properties meet the desired requirements (see next section).

3.2. Product properties

In this section, the correlation between the redox composition and initiation temperature on the product properties, such as particle size, molecular weight (MW), minimal film forming temperature (MFFT), as well as glass transition temperature (T_g), is discussed.

Fig. 4 shows a representative comparison of the MW distributions obtained at a constant r_{redox} and at different

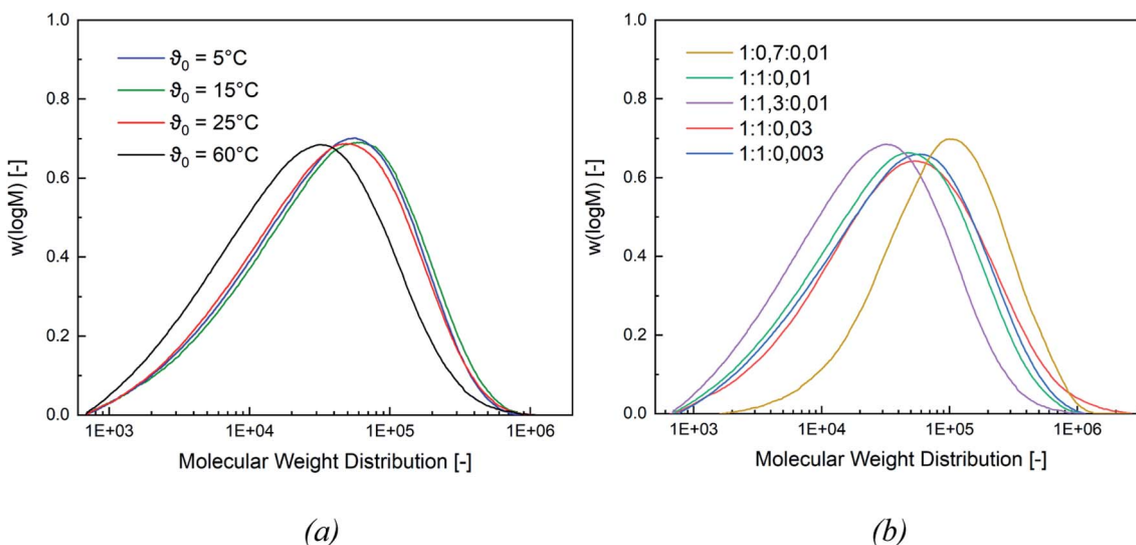


Fig. 4 Representative influence of different operating conditions such as ϑ_0 (a), molar ratio 1 : 1.3 : 0.01 and the molar ratio of the redox components (b), temperature $\vartheta_0 = 60$ °C on the MW distribution.



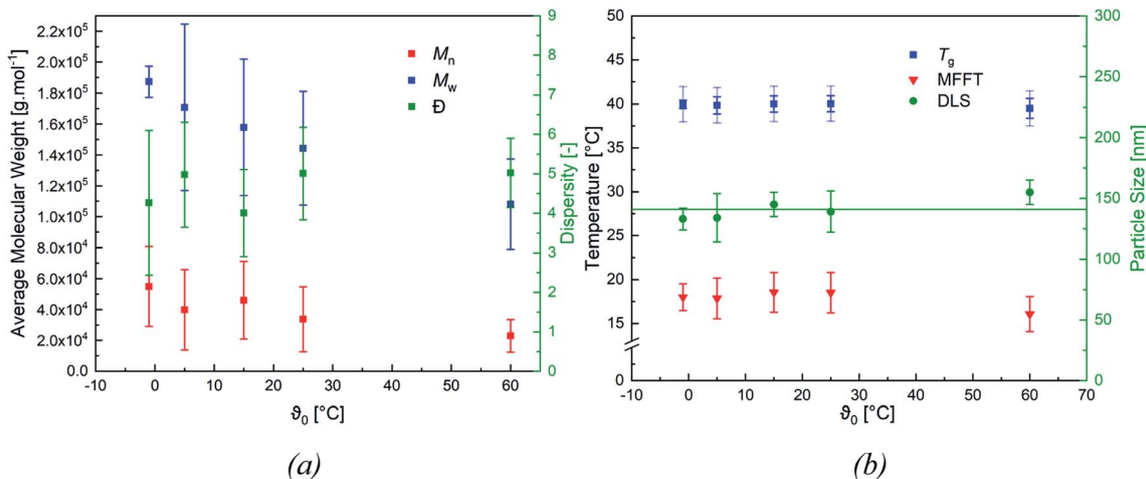


Fig. 5 Overview of the mean molecular weight (M_n and M_w) and the dispersity index (a) and the average particle size (green line corresponds to the average particle size = 141 nm), MFFT and T_g (b), standard deviation includes the deviation of the measurements as well as the accuracy of the device) as a function of ϑ_0 . Each point includes all experiments at the given temperature, at different r_{redox} , the error bars correspond to the standard deviation ($n = 3$).

Table 7 Overview of the calculated number of particles present in the dispersion based on the measured particle size (DLS)

ϑ_0 (°C)	Number of particles in 0.4 L (10^{16})	Number of particles in 1 L (10^{17} L ⁻¹)
60	3.5	0.9
25	4.8	1.2
15	4.3	1.1
5	5.4	1.4
-1	5.5	1.4
Average	4.7 ± 0.8	1.2 ± 0.2

temperatures (for MW distributions of the remaining molar ratios *cf.* Appendix Fig. S5 and S6†). While a lower MW is obtained at $\vartheta_0 = 60$ °C, the average molecular weight stays constant at lower initiation temperatures. At the same molar ratio of the redox components, lowering the temperature will lead to higher MW, while a higher temperature favours termination reactions and therefore results in a lower MW. However, the change in ϑ_0 has no effect on the MW below an initiation temperature of ≤ 25 °C.

When varying the ratio of the redox initiator components at constant ϑ_0 , two effects are observed. Firstly, adjusting $r_{\text{Fe-cat}}$ showed no influence on MW nor on its distribution. However secondly, varying r_{BHP} results in changes of the MW: reducing the amount of peroxide has shifted the MWD to higher MW, while an increase has shifted the MWD to lower MW (see Fig. 4, (b)). When considering the MW solely as a function of ϑ_0 , an overall decreasing trend of the average M_n and M_w values of all experiments can be observed with higher temperatures (*cf.* Fig. 5, (a)), whereas the polydispersity index ($\text{DI} = M_w/M_n$) is not affected.

Fig. 5 (b) features the measured particle size as a function of ϑ_0 , as well as the average particle size (141 ± 8 nm), which was used to calculate the number of particles ($(4.7 \pm 0.8) \times 10^{16} \hat{=} (1.2 \pm 0.2) \times 10^{17}$ L⁻¹, *cf.* Table 7) present in the dispersion.

The average number of particles shows no effect when varying $r_{\text{Fe-cat}}$, nor ϑ_0 .

Fig. 5 (b) also includes the DSC (accuracy: ± 2 °C) and MFFT measurements which revealed an average T_g of 40 ± 1 °C and an equally constant film forming ability. This indicates an independency of the polymer structure towards a varying ϑ_0 and r_{redox} . Although this was expected, due to the choice of monomer having the most significant influence on polymer structure and flexibility, an additional ¹H NMR spectroscopy analysis was conducted to verify this assumption (*cf.* Fig. 7, (a)).⁴⁹ The ¹H-NMR analysis showed a 98.9% superposition of the polymer signals at 4.75 ppm, 1.95 ppm and 1.75 ppm. The smaller but visible signals at around 1 ppm can be attributed to the neo-C₉H₁₉ chain originating from Versa@10, which has been added in a small amount, even though ¹H-COSY NMR showed no coupling between these signals (*cf.* Fig. 7(b)).⁵⁰ Furthermore, ¹³C- and HSQC-NMR spectra showed no coupling of these signals (Appendix, Fig. A7†), most probably due to the small ratio of vinyl acetate to Versa@10 and low signal intensities in ¹³C spectra.

According to literature, the small signals at 3.75–3.93 ppm, 4.21 ppm, 4.45 ppm and 4.66 ppm are related to the OH-group of the protective colloid polyvinyl alcohol.^{51,52} The CH₂-group of polyvinyl alcohol overlaps with the signals related to the neo-

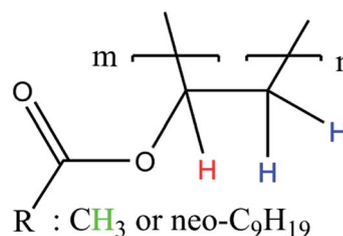


Fig. 6 Simplified polymer structure for the assignment of NMR signals in Fig. 7 to their corresponding H-atoms.



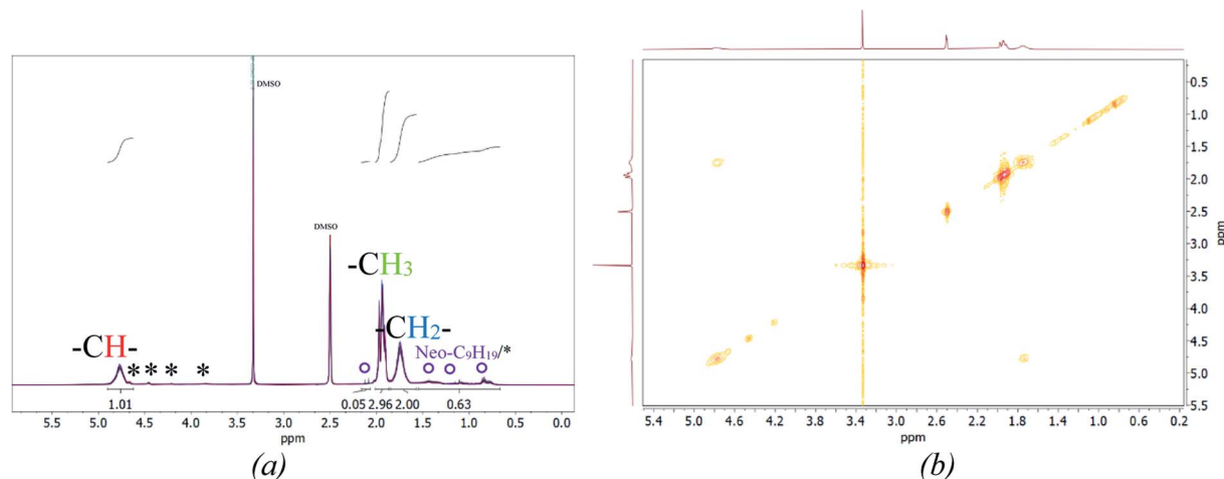


Fig. 7 Stacked ^1H NMR spectra of all products (a, *: signals associated to Mowiol 4-88, °: signal associated to R = Neo- C_9H_{19}) and a representative 2D ^1H -COSY spectrum (b).

C_9H_{19} chain originating from the Versa@10 monomer and is located between 1.26 and 1.55 ppm.^{51,52} Since the signals belonging to the protective colloid partially overlap with those of the copolymer, the signal related to the CH-group (cf. Fig. 6) at 4.75 ppm has an average integral higher than 1, as will be shown hereafter.

To ensure a comparability of the spectra, the signal at 1.75 ppm was normalised to 2 (CH_2 -group, cf. Fig. 6) to be able to compare the resulting integral of the signals at 4.75 ppm and 1.75 ppm. This resulted into an average integral of 1.04 ± 0.01 for the CH-signal at 4.75 ppm and of 2.96 ± 0.03 for the CH_3 -signal at 1.95 ppm. The NMR analysis based on 30 samples showed an average deviation in the polymer structure of approximately 1.14%.

Examination of the signals related to the neo- C_9H_{19} chain originating from the Versa@10 monomer indicates that both monomers may not be incorporated to the final copolymer in the original ratio of 9 : 1. An estimation is possible by integrating the signals related to the neo- C_9H_{19} chain and subtracting the signal intensity related to the $-\text{CH}_2$ signal from the protective colloid, which overlaps with the neo- C_9H_{19} signals. Accordingly, an average of $4 \pm 1\%$ of Versa@10 has been incorporated in to the copolymer. Interestingly, vinyl acetate to Versa@10 ratio is estimated to be highest at an initiation temperature of 60 °C, while a low initiation temperature of -1 or 5 °C led to a lower incorporation (cf. Appendix Table A4† for detailed intensities of the signals). This could be due to the very low water solubility of Versa@10 (2.58 wt% vinyl acetate against 7.5×10^{-4} wt% Versa@10 at room temperature⁵⁰), which would

increase at higher temperature and therefore facilitate the implementation of the monomer into the final polymer.

A more accurate, quantitative determination of the exact ratio *via* NMR-methods is, however, not possible due to overlapping of signals in ^1H spectra and low signal intensities in $^{13}\text{C}/\text{HSQC}$ spectra. Summarised, no significant differences in polymer structure could be identified *via* NMR-methods, whereas slight influence of polymerisation temperature on Versa@10 incorporation is indicated.

3.3. Statistical evaluation

The statistical evaluation of the correlations described in the previous sections enables access to a broad range of interpolated reaction times and the related molecular weights of the products and validates the statistical significance of the empirically detected dependencies. The overall polymerisation rate is described by the time at which 99% of the overall conversion of the reaction is achieved ($t_{99\%}$, min). All models are statistically relevant, as indicated by the analysis of variances (cf. Appendix Tables A1–A3†), and provide an adequate description in the interpolated range (Table 8), as shown by the parameter plots (R^2 0.988–0.999, Fig. 8, (a)).

The relations follow quadratic process orders, whereas the reaction time is exponentially related to ϑ_0 and linearly to $r_{\text{Fe-cat.}}$, as well as to r_{tBHP} (eqn (3)).

$$\frac{1}{\sqrt{t_{99\%}}} = -4.48 \times 10^{-2} + 5.22r_{\text{Fe-cat.}} + 1.03 \times 10^{-1}r_{\text{tBHP}} + 1.98 \times 10^{-3}\vartheta_0 + 8.49 \times 10^{-5}\vartheta_0 T^2 \quad (3)$$

Table 8 Significance of statistical models, influence factors and predicted range of model responses

Model output	R_{model}^2	$R_{\text{adj.}}^2$	$R_{\text{pred.}}^2$	Significant influence factors	Range
M_n	0.962	0.956	0.942	ϑ_0 , r_{tBHP}	17 400–100 600 g mol^{-1}
M_w	0.851	0.825	0.775	ϑ_0 , r_{tBHP}	97 000–238 000 g mol^{-1}
$t_{99\%}$	0.968	0.961	0.942	ϑ_0 , $r_{\text{Fe-cat.}}$, r_{tBHP}	4–236 min



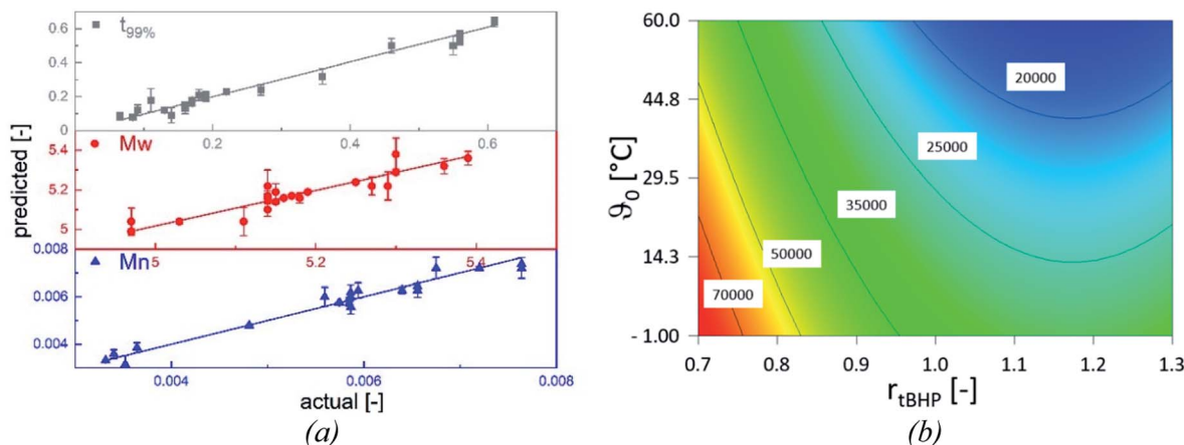


Fig. 8 Parameter plots of statistical models (a). Values of x- and y-axis are given including the individual model transformations. Straight lines represent linear plots, error bars the standard deviation of the model prediction. Contour plot of M_n in dependence of ϑ_0 and r_{tBHP} (b). Values in white boxes represent M_n [g mol⁻¹] of the respective contour, the plot is independent of r_{Fe-cat} .

In general, with equimolar ratios of reducing and oxidising agents, the overall process time can be reduced by 40–86% by varying r_{Fe-cat} . For example, at equimolar ratios of AsAc and *t*BHP, low initiation temperatures ($\vartheta_0 \leq 25$ °C) and high catalyst concentration ($r_{Fe-cat} = 0.03$) a complete conversion can be achieved in shorter reaction times ($t_{99\%} = 10$ –22 min), whereas the reaction time increases to $t_{99\%} = 32$ –194 min when r_{Fe-cat} is lowered by a factor ten. Increasing *t*BHP would also result in higher reaction rates but would also lead to changes in the molecular weight at the same time, since M_n and M_w show linear relations to initiation temperature but exponential relations to r_{tBHP} (eqn (4) and (5)).

$$\frac{1}{\sqrt{M_n}} = -1.11 \times 10^{-2} + 2.9 \times 10^{-2} r_{tBHP} + 2.68 \times 10^{-5} \vartheta_0 - 1.24 \times 10^{-2} r_{tBHP}^2 \quad (4)$$

$$\log_{10}(M_w) = +6.00 - 1.21 r_{tBHP} - 3.31 \times 10^{-3} \vartheta_0 + 4.49 \times 10^{-1} r_{tBHP}^2 \quad (5)$$

In contrast to thermal initiators, additional degrees of freedom are obtainable in a certain range, which allows the product properties as well as the reaction time to be adjusted, simply by varying r_{redox} : in particular, different molecular weights can be targeted by varying r_{tBHP} at a constant ϑ_0 , while the reaction rate can be controlled independently by adjusting r_{Fe-cat} .

It is interesting to note that the DI solely depends on r_{tBHP} and not on ϑ_0 , with the highest DI (6.5) obtained at max. r_{tBHP} and the lowest (2.4) at min. r_{tBHP} . Our results are within the same range as previous works, where DIs of 5.2–6.0 were obtained for the same redox/monomer system at $r_{tBHP} = 1.0$ and in

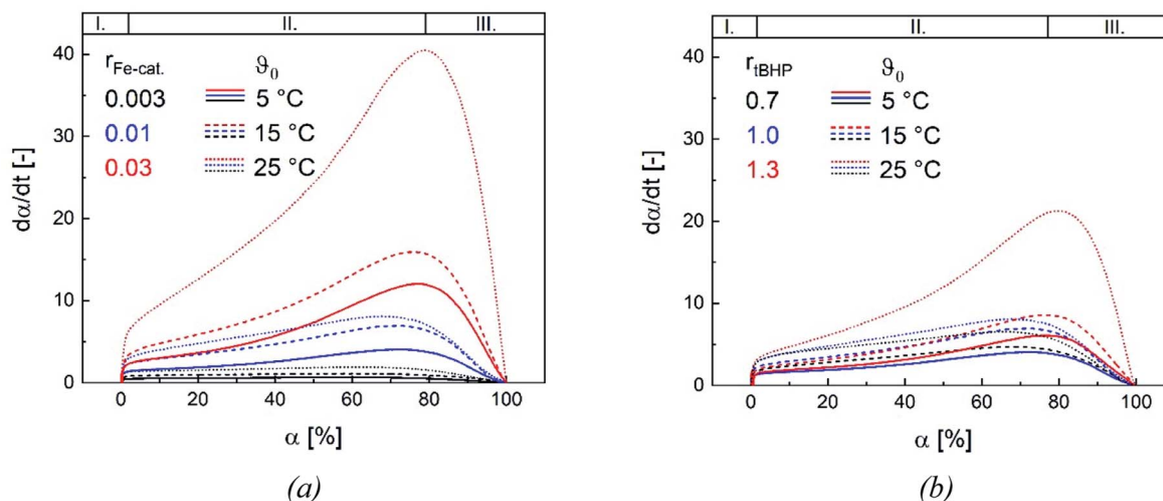


Fig. 9 Dependency of the reaction rate on ϑ_0 and on r_{Fe-cat} at a fixed $r_{tBHP} = 1.0$ (a) and dependency of the reaction rate on ϑ_0 and on r_{tBHP} at a fixed $r_{Fe-cat} = 0.01$ (b), shown for three initiation temperatures. I., II., and III. represent the three intervals of the emulsion polymerisation.



a continuous process with slightly higher monomer contents (20–40 wt%).³³

While the data given in this section enables a prediction of the final product properties and reaction times, the influence of r_{redox} and ϑ_0 on the kinetics of the polymerisation are discussed in the next section.

3.4. Reaction kinetics

Changes in product properties and overall reaction times are connected to the individual reaction profiles. The reaction rate increases for all settings with higher ϑ_0 , as well as with changes regarding $r_{\text{Fe-cat.}}$ and r_{tBHP} (cf. Fig. 9), which demonstrates that all of these factors contribute to the radical generation rate of the initiator system. The kinetic profile of all reactions clearly shows three intervals (I, II and III, cf. Fig. 9): After adding the initiator, a rapid increase in polymerisation rate occurs without any induction period, which is related to the process of micellar nucleation (Interval I). As soon as the conversion reaches approx. 3%, a less pronounced but still significant further increase in reaction rate is observed (Interval II). After 66–76% conversion, the system enters interval III (disappearance of monomer droplets) without showing a gel effect. The uniformity of the profiles indicates that the polymerisation rate in the binary vinyl acetate and Versa@10 system can be described *via* a pseudo-homopolymerisation approach in the temperature range of $\vartheta_0 \leq 25^\circ\text{C}$ (Fig. 9).⁹

The additional increase of the reaction rate in interval II shows that significant nucleation occurs until interval III is reached: This result is consistent with former works, in which significant secondary nucleation was reported for redox initiated emulsion polymerisations, and in contrast to processes using thermal initiators.¹⁴ Since both a higher ϑ_0 and a higher $r_{\text{Fe-cat.}}$ lead to a significant increase in the reaction rates during interval II, we can conclude that the secondary nucleation is directly related to the radical generation rate. Interestingly, no changes in MW, particle number nor particle size were observed, when varying $r_{\text{Fe-cat.}}$: this shows that in this case, the

overall process and product properties are dominated by one influence factor. Due to the relatively high hydrophobicity of VAc, Versa@10, tBHP and the (most probably) generated *tert*-butoxy radicals, as well as the low monomer and high emulsifier contents in this work, it can be assumed that particle generation occurs predominantly *via* micellar nucleation. Therefore, the main factor determining the reaction rate when varying $r_{\text{Fe-cat.}}$ should be the entry of radicals into monomer-swollen micelles, whereas events during particle growth, such as radical entry/exit, which may affect chain propagation, chain transfer or termination, are not influenced. The rate of chain termination has been reported to be relatively slow compared to the rates of transfer to monomer and polymer in emulsion polymerisation of vinyl acetate. The influence of the termination rate on the molecular weight can therefore be considered negligible even at high initiation rates.³³ In general, Fig. 10 shows that the reaction rate increases with higher r_{tBHP} as well as $r_{\text{Fe-cat.}}$. This trend is consistent with previous results on redox initiated emulsion polymerisations and can be explained by higher reaction rates at higher oxidising component ratios.³⁷ Furthermore, a higher initiator concentration has been shown to lead to lower MW (*e.g.* for AsAc/H₂O₂ initiated polymerisation of *n*-butyl methacrylate).¹⁴ This trend could also be explained by a higher radical entry rate into the micelles, although the question arises as to why a variation of r_{tBHP} results in changes of the MW, while variation of $r_{\text{Fe-cat.}}$ does not (Fig. 4(b)).

Our results suggest that the maximum polymerisation rate is linearly related to $r_{\text{Fe-cat.}}$ and exponentially related to r_{tBHP} : this trend is in accordance with former results regarding the decomposition rate constant of AsAc in the redox initiator system, which depends linearly on $r_{\text{Fe-cat.}}$ and exponentially on the ratio of oxidising to reducing agent.³⁷ While the present data provides kinetic information in dependency of r_{redox} , it is still not precisely clarified exactly which radicals are formed and whether, in addition to changes in the decomposition rate, the formation of different radical species occurs due to changes in r_{tBHP} , as is suggested by the observed change of MW in this

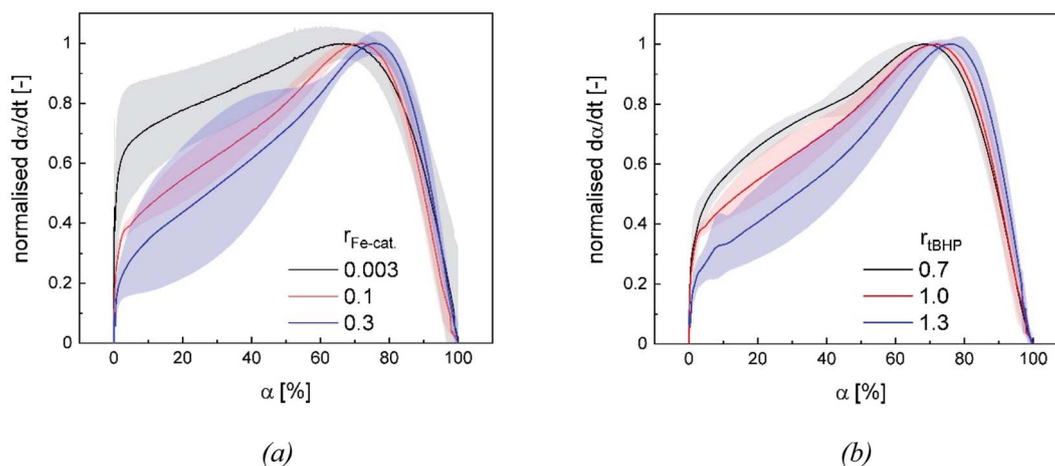


Fig. 10 Normalised dependency of the reaction rate on $r_{\text{Fe-cat.}}$ at fixed $r_{\text{tBHP}} = 1.0$ (a) and of the reaction rate on r_{tBHP} at fixed $r_{\text{Fe-cat.}} = 0.01$ (b). Lines represent the average trend of the experiments at all ϑ_0 and the error bands represent the standard deviation.



work. In this sense, our results emphasize the importance of further investigations on redox initiator decomposition mechanisms in dependency of process conditions and r_{redox} .

4. Conclusion

It was shown that AsAc/tBHP/Fe-cat. redox initiated emulsion copolymerisation of vinyl acetate and Versa®10 in the temperature range of -1 to 87 °C results in high conversion (95–99%) as well as variable and fast reaction times in the whole temperature range. The process was investigated with respect to the influence of different molar ratios of the redox system and variation of initiation temperatures on the conversion rate and product properties. While the conversion rate is adjustable by both the r_{redox} as well as ϑ_0 , the product properties were found to be independent of the amount of catalyst used to initiate the polymerisation, and therefore independent of the radical generation rate. Modifying the amount of oxidising agent resulted in changes in MW, but not in its distribution nor particle size. Overall, this work has proved that the process can be modified to a certain extent to increase process safety or reduce production costs, without compromising on the desired product properties.

Conflicts of interest

The authors declare no competing financial interest.

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