Semicontinuous emulsion copolymerization to obtain styrene-methyl acrylate copolymers with predetermined chemical composition distributions

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Semicontinuous Emulsion Copolymerization to Obtain Styrene—Methyl Acrylate Copolymers with Predetermined Chemical Composition Distributions

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SYNOPSIS

Four types of emulsion copolymerization processes were applied to produce various styrene—methyl acrylate copolymer latices. (1) Batch emulsion copolymerization. (2) Monomer starved semicontinuous emulsion copolymerization, i.e., a monomer mixture of constant composition was fed to the reactor at a constant rate. Sufficiently low addition rates led to homogeneous copolymers. (3) Optimal addition rate profiles were used to produce homogeneous emulsion copolymers in relatively short times. The profiles were determined in a semiempirical way, and applied to three different copolymer compositions. (4) We made an attempt to determine an addition rate profile to produce a heterogeneous emulsion copolymer with predetermined heterogeneity, i.e., a copolymer of which the chemical composition distribution (CCD) did not consist of one narrow peak, as with homogeneous copolymers, but had a predetermined broadness profile. Strategies (2) and (3) were used to produce homogeneous emulsion copolymers with varying fractions of styrene. Strategy (3) was also used to demonstrate the influence of the monomer ratio on the kinetics. The advantage of this method is that the monomer ratio is constant over the whole conversion range in the latex particles. High performance liquid chromatography (HPLC) was used to determine the homogeneity/heterogeneity of the copolymers produced, and proved to be invaluable in determining the optimal addition rate profile. The final goal will be to study the influence of the chemical composition distribution (CCD) on mechanical properties of these copolymers, as both homopolymers used in this study have quite different glass transition temperatures; the results will be published later. © 1993 John Wiley & Sons, Inc.

INTRODUCTION

Since there is an increasing demand for special polymers, more and more copolymers and even terpolymers are being used in industry. The problem one can run into is that when these copolymers are made batchwise, composition drift generally influences the chemical composition distribution. Due to the monomer partitioning, this problem can be even worse when the copolymers are made in emulsion, in those cases where the more reactive monomer is also the less water-soluble monomer.¹

In order to produce a copolymer latex with desired properties, one needs to know how to control the chemical composition distribution of the copolymers. This has previously been realized by many researchers, by producing emulsion copolymers semicontinuously, which involves adding the monomers to the reaction mixture during the course of the reaction.²⁻⁵ The easiest way to achieve is to add the monomer mixture (pure or pre-emulsified) at a constant, low rate (starved conditions). Many investigators have applied this strategy.²⁻¹⁰ However, working under starved conditions means that the monomer concentrations are low, and this leads to...
low polymerization rates and long reaction times. Therefore, Arzamendi and Asua\textsuperscript{11,12} proposed a method to make homogeneous copolymers in emulsion at high monomer concentrations, by adding the more reactive monomer (via an optimal addition rate profile) to the reaction mixture containing all of the less reactive monomer [vinyl acetate (VAc); they copolymerized VAc with methyl acrylate (MA)]. By using this method they could keep the ratio of the monomers inside the latex particles constant. Since the method of the optimal addition rate profile leads to short reaction times, heat removal can be a limiting factor. Therefore, they also looked at the situation where the reactor had limited capacity for heat removal, adding the less reactive monomer to the reactor at a constant rate, and adding the second monomer via an optimal addition rate profile.\textsuperscript{13} In our laboratory,\textsuperscript{14} this technique was applied to the emulsion copolymerization of styrene (S) and methyl acrylate (MA), but a somewhat different procedure was utilized to calculate the optimal addition rate profile. Arzamendi et al.\textsuperscript{15} applied this technique to the emulsion copolymerization of methyl methacrylate and ethyl acrylate (two different compositions) and Urquiola et al.\textsuperscript{16} also to VAc–MA copolymerization. Very recently, Arzamendi et al.\textsuperscript{17} published results of simulations of reactions with optimal addition rate profiles for homogeneous terpolymers of styrene, methyl methacrylate, and ethyl acrylate. A more elaborate approach to modeling semicontinuous (emulsion) polymerizations was published by Hamielec et al.\textsuperscript{18,19} They modeled polymerizations, including molecular weight distributions, with empirically measured kinetic parameters. They were able to model batch, semicontinuous, and continuous polymerizations, even taking into account heat removal. However, if few of the kinetic parameters are known or can be estimated, the approach presented by Asua et al.\textsuperscript{11,12} is more appropriate, depending on what one wants to look at.

In the present investigation we have applied a procedure very similar to that developed by us previously\textsuperscript{14} to produce homogeneous S–MA copolymer latices with varying amounts of styrene. A similar procedure was followed to produce a copolymer with an average composition of 25\% S, but with a predetermined broad CCD. In all cases we added $\pm 1\%$ chain transfer agent (CTA) to minimize the influence of addition strategy on the molecular weight distribution, since this has a significant influence on mechanical properties, which might obscure the influence of the CCD, which is what we want to study eventually. The use of CTA might also prevent gel formation during the starved conditions process. Note that the goal is to produce several different CCDs, and therefore we did not consider other aspects like the effect of high polymerization rates on heat removal from the reactor, etc.

We compared batch and semicontinuous reactions with respect to composition drift, i.e., how the factors that govern composition drift in batch reactions also control the homogeneity of the semicontinuously formed copolymers. We also studied the kinetics of the reactions with optimal addition rate profiles.

**EXPERIMENTAL**

Applied chemicals are: styrene (S), methyl acrylate (MA), sodium persulfate (SPS, initiator), n-dodecyl mercaptan (NDM, chain transfer agent), sodium bicarbonate (SB, buffer) (all p.a., Merck), sodium dodecyl sulfate (SDS, emulsifier, Fluka, 99\%). All experiments were carried out in a 1.3-L stainless-steel reactor. Temperature was always kept at 323 K. The reactor configuration has been described elsewhere.\textsuperscript{1} During semicontinuous reactions the monomer feed (never emulsified) was added with a Dosimat 665 (Metrohm), and the feed was computer-controlled (Atari 1040 ST).

Conversion was followed by solid content analysis. Copolymer composition was sometimes checked with $^1$H-NMR. The spectra were recorded with a Bruker AM 400 spectrometer (400 MHz) at 298 K using CDCl$_3$ as a solvent. High performance liquid chromatography (HPLC) was used to determine the chemical composition distribution (CCD) of the styrene–methyl acrylate copolymers. Gradient elution was applied to obtain the CCDs. The procedure has been described elsewhere.\textsuperscript{20} The column was a silica column [Zorbax SIl (Rockland) Silica column with Zorbax SIl precolumn 4.6 mm ID $\times$ 15 cm]. The column was used at room temperature. We applied an eluent consisting of $n$-heptane and THF: $n$-Heptane is a nonsolvent for all copolymers. The gradient was linear, and the composition shifted from 80\% $n$-heptane/20\% THF to 60\% $n$-heptane/40\% THF after 1 min and 100\% THF after 16 min. Flow rate was 0.6 mL/min. We used a flame ionization detector with a moving wire (FID).\textsuperscript{21} UV detection of polycrylates is not feasible due to the low absorption of polycrylates. However, FID responses of poly (methyl acrylate) and polystyrene are comparable. Molecular weights were determined with gel permeation chromatography (GPC). The chromatographic system (Waters Associates) was equipped with a differential refractometer and a 2 Shodex linear column, kept at 40°C. The eluent was
THF, and the elution rate was 1 mL/min. The samples for $^1$H-NMR and HPLC and GPC were taken from samples used for solid content analysis. The particle sizes, needed to calculate the number of particles, were determined with dynamic light scattering (DLS). We could not check the diameters with transmission electron microscopy.

RESULTS AND DISCUSSION

During the semicontinuous reactions, the total amounts of monomers added was such that all latices had a final solids content of about 17%, corresponding to an initial monomer to water ratio (M/W) of 0.2 g/g for batch reactions. All reactions were carried out at 50°C, and the stirring rate was 250 rpm. NDM was, as stated before, added in all cases in order to (i) limit the molecular weights and (ii) diminish the effect of addition strategy on the molecular weight distribution (MMD); it is known from the literature that there can be an effect. Addition of NDM could decrease the extent of this by exhibiting a much larger effect on MMD than the addition strategy. To check this, we measured the MMDs of the final products. The batch reactions and reactions under starved conditions were performed \textit{ab initio} (without seed); the reactions with (optimal) addition rate profiles were started from a seed latex. This was done to avoid irreproducibility due to the nucleation stage. By doing so, the number of particles could be controlled and also the reaction rate, which is very important in reactions where both monomers are fed to the reactor at different rates. The seed always had the same average composition as the desired copolymer. The size of the seed particles was small enough to ensure that the polymer in the seed particles had a negligible contribution to the final CCD compared with the contribution of the copolymer that was polymerized onto them. We found that neither DLS nor transmission electron microscopy could give reliable diameters of the seed particles. However, the exact size of the seed particles is not important. The seed latices were prepared by semicontinuous addition of a monomer mixture with the same composition as the desired composition of the copolymer in the seed latex. These reactions were carried out at 80°C. The amount of seed latex in the recipes was such that the particle number was about $5 \times 10^{17}$ L$^{-1}$ (the calculation was based on the diameter that was obtained from DLS). The particle diameters of the final latices could be measured more reliably.

Batch reactions and reactions under starved conditions were carried out according to the recipes of Tables I and II. In the experiments with (optimal) addition rate profiles, the reactor was always precharged with all ingredients [including seed and all of the methyl acrylate (MA) and part of the styrene (S)], and the remaining part of the S was added during the reaction. The recipes for these reactions are shown in Table III. The total volume decrease of the reaction volume due to sampling for analyses of solid content and particle diameter was always less than 5% in semicontinuous reactions. Corrections were made to account for this decrease.

Batch Reactions

In order to put the results of the semicontinuous reactions into perspective, we did three batch reactions with M/W = 0.2, but with varying overall molar compositions, namely $F_S$ (overall fraction of S in copolymer) = 0.25, 0.50, and 0.80. Figure 1 shows the CCDs of these reactions as determined with HPLC/FID. The CCD of the copolymer with 25 mol % is significantly different from that measured previously with UV detection. In that case,
Table III Recipes for the Reactions with the Addition Rate Profiles

<table>
<thead>
<tr>
<th></th>
<th>$F_s = 0.25$</th>
<th>$F_s = 0.50$</th>
<th>$F_s = 0.80$</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>H$_2$O</strong></td>
<td>900*</td>
<td>900*</td>
<td>900*</td>
</tr>
<tr>
<td><strong>SDS</strong></td>
<td>1.00*</td>
<td>1.00*</td>
<td>1.00*</td>
</tr>
<tr>
<td><strong>SPS</strong></td>
<td>0.264</td>
<td>0.264</td>
<td>0.264</td>
</tr>
<tr>
<td><strong>SB</strong></td>
<td>0.100</td>
<td>0.100</td>
<td>0.100</td>
</tr>
<tr>
<td><strong>Seed latex</strong></td>
<td>27.939</td>
<td>20.492</td>
<td>12.658</td>
</tr>
<tr>
<td><strong>MA</strong></td>
<td>126.323</td>
<td>80.626</td>
<td>30.635</td>
</tr>
<tr>
<td><strong>S</strong></td>
<td>8.715</td>
<td>30.976</td>
<td>115.969</td>
</tr>
<tr>
<td><strong>SD</strong></td>
<td>42.22</td>
<td>66.53</td>
<td>32.273</td>
</tr>
<tr>
<td><strong>NDM</strong></td>
<td>1.35</td>
<td>1.116</td>
<td>1.466</td>
</tr>
<tr>
<td><strong>NDM</strong></td>
<td>0.422</td>
<td>0.665</td>
<td>0.323</td>
</tr>
</tbody>
</table>

* The amount of SDS in the seed was subtracted from 1.00 g so that the total amount of SDS always was 1.00 g.

it was impossible to detect the poly(methyl acrylate) peak with HPLC. We were now able to show unambiguously that there is indeed a homopolymer peak in the CCD as a result of composition drift. Concerning the CCD of the copolymer with 50% of S, we have to point out that the average fraction of S in the copolymer as calculated from this CCD is higher than 50%. The reason for this is probably that polymer is formed with fractions of S much lower than 50% due to composition drift, but this polymer is not detected because of its low concentration. In fact, calculating the average composition from a broad CCD is only possible if the measurement is very accurate. The exact position of the baseline is important in this case, because a small shift of the baseline can result in a relatively large change in apparent concentration of polymer, which has a low concentration. $^1$H-NMR showed that the average composition was 50%.

It can be clearly seen that the breadth of the CCD depends strongly on the overall composition. The lower $F_s$, the broader the CCD, which means that composition drift is strongest in the case of the lowest $F_s$, even leading to a bimodal distribution. This can be explained qualitatively using the monomer partitioning data and reactivity ratios. The monomer partitioning behavior of this system is such that the composition drift due to the difference between the monomer ratio in the particles and the overall monomer ratio is not strongly dependent on the overall fraction of S. Mass balance calculations with monomer partitioning data as determined by van Doremaele et al. show this. The strong composition drift with low fractions of S must therefore primarily result from the copolymerization kinetics as described by the reactivity ratios ($r_S = 0.73$ and $r_{MA} = 0.19$ at 323 K). As can be seen in Figure 2, where we plotted the instantaneous fraction of S in the copolymer ($F_s$) versus the fraction of S in the feed ($f_s$), the lower $f_s$, the higher the relative difference between $F_s$ and $f_s$ and the stronger the composition drift. Judging from the above results, we expect the copolymers with 80 mol % S to be quite homogeneous in all process modes.

**Reactions under Starved Conditions**

We did reactions under starved conditions with two different fractions of S (50 and 80 mol %), and each reaction was carried out using three different addition times (4, 8, and 32 h, in all cases adding 120 g of monomer mixture to 600 g of water). These reactions are very simple because of the constant addition rates. The reactions were performed for comparison with the results of reactions with optimal addition rate profiles, which are less simply carried out. We have done similar reactions previously with 25 mol % S. $^4$ Figures 3a and 3b show the overall conversions ($X_0$) of these reactions versus fractional time $t/t_0$, $t_0$ being the total addition time. Our results are comparable to those found earlier, i.e., the longer the addition times, the higher the instantaneous conversions of monomer to polymer.

Adding the monomer mixture at low rates leads to a situation where the addition rate determines the reaction rate. In this way, one can effectively influence the concentrations of the monomers inside the latex particles, where polymerization takes place throughout the greater part of the reaction. Early on in the reaction, the system reaches a steady state in which the polymerization rate equals the addition rate for both monomers. This must mean that the ratio of the monomers in the particles corresponds to the value that can be calculated from the reactivity ratios and copolymer composition. It is obvious that it takes some time before this situation is
reached, and it is during this time that composition drift occurs. There is a relatively large difference between reaction rate and addition rate in the beginning of the reaction as can be seen in Figures 3a and 3b. Later during the reaction, the curve of conversion versus time is parallel to the line, indicating 100% instantaneous conversion showing that the reaction rate equals the addition rate. One can also assume that particle nucleation takes place over a period of time and that the number of particles is not fixed in the beginning. If the number of particles increases with time, then it is evident that in the beginning polymerization rate is lower than the addition rate which leads to composition drift. The composition drift is clearly shown in Figures 4a and 4b. Here we plotted the cumulative fractions of S in the copolymer as determined with \(^1\)H-NMR versus fractional time. In all cases we see that the conversion of S in the beginning is larger than the conversion of MA, leading to S-rich copolymers, except for the very beginning, where the absence of a significant number of particles could cause monomer consumption processes other than those normally occurring in polymer particles to prevail. In the literature,\(^{1,14}\) aqueous phase polymerization has often been assumed to explain the relatively high consumption rate of MA. However, poly(methyl acrylate) (PMA) is not water-soluble, so that it is hard to see that aqueous phase polymerization could occur to a large enough extent. Note that in these figures \(t/t_0\) ranges from 0 to 0.5 and \(F_s\) from 0.4 to 1.

Figures 5a and 5b show the CCDs of the reactions under starved conditions. In Figure 5a we see that the average composition of the copolymer formed in the 4-h addition, as calculated from the CCD, is about 60% S, which we also found for the corresponding batch copolymer. The explanation might be the same in this case, there could be a tail stretching towards lower fractions of S. This then indicates that there is still considerable composition drift occurring, and this is not surprising if we com-

**Figure 1** Experimentally determined CCDs of the batch reactions with \(F_s = 0.25, 0.50, \) and 0.80. \(R\) designates the relative abundance and is in arbitrary units.

**Figure 2** Depiction of the instantaneous copolymer composition \((F_s)\) as a function of composition of the feed \((f_s)\) for an S-MA free radical copolymerization at 50°C with \(r_S = 0.73\) and \(r_{MA} = 0.19\) (ultimate model).
Figure 3 Overall conversion versus fractional time ($t_0 = \text{addition time}$) of the experiments under starved conditions with different addition times: (+) 4 h; (○) 8 h; (□) 32 h; (a) $F_S = 0.50$; (b) $F_S = 0.80$.

Figure 4 Copolymer composition (measured with $^1$H-NMR) versus fractional time ($t_0 = \text{addition time}$) of the reactions under starved conditions with different addition times: (+) 4 h; (○) 8 h; (□) 32 h; (a) $F_S = 0.50$; (b) $F_S = 0.80$. (The solid line represents the overall fraction of S.)

Figure 5 Overall conversion versus fractional time ($t_0 = \text{addition time}$) of the experiments under starved conditions with different addition times: (+) 4 h; (○) 8 h; (□) 32 h; (a) $F_S = 0.50$; (b) $F_S = 0.80$. (The solid line represents the overall fraction of S.)
STYRENE-METHYL ACRYLATE COPOLYMERS

Optimal Addition Rate Profiles

As stated in the Introduction, Arzamendi and Asua\textsuperscript{11,12} were the first to report on the possibility to produce homogeneous emulsion copolymers via a method that is referred to as addition with an optimal addition rate profile. This way the ratio of the monomer concentrations inside the polymer particles can be fixed at a value that ensures the formation of a homogeneous copolymer of the desired composition. Furthermore, the concentrations of the monomers in the particles are much higher than they are in monomer starved reactions. This leads to higher polymerization rates and therefore to shorter reaction times.

In order to comply with the above conditions (constant ratio and high concentrations), the following strategy was applied by Arzamendi and Asua, and in our group by van Doremaele et al.\textsuperscript{11,14}; we shall discuss both methods and adopt the strong points of both. The reactor is precharged with a seed to ensure relatively constant numbers of particles. All ingredients are added as well as all of the less reactive monomer [\(\text{VAc}\) in the case of \(\text{MA-VAc}\) copolymerizations (Arzamendi and Asua\textsuperscript{11,12}), \(\text{MA}\) in the case of \(\text{S-MA}\) copolymerizations (van Doremaele et al.\textsuperscript{11,14})] and part of the more reactive monomer. The amount of precharged more-reactive monomer is crucial. It is calculated so that the amount ensures formation of copolymer of the desired composition as soon as the reaction is started. This calculation is based on mass balances, monomer partitioning data, and reactivity ratios.

The remaining part of the more reactive monomer is then added during the reaction at a rate that is variable in time, depending on the reaction rate and the change in volume of the various phases (monomer partitioning). The difficulty of this strategy lies in the determination of the optimal addition rate profile. Arzamendi and Asua and van Doremaele et al. looked at the problem from different angles, but arrived at methods that are intrinsically the same. Arzamendi and Asua\textsuperscript{11} first developed a mathematical model for calculating the optimal addition rate profile for a seeded reaction where they precharged all of the \(\text{VAc}\) and part of the \(\text{MA}\). The calculation of \(\eta\) was based on the equation derived by Ugelstad and Hansen.\textsuperscript{24} However, in a second paper,\textsuperscript{12} the same authors stated that several parameters needed for the calculation of \(\eta\) are not known and that it is therefore not possible to calculate the optimal addition rate profile without further experiments. To overcome this problem, they developed a semiempirical method to calculate the time evolution of \(\eta\). Using thermodynamic equilibrium equations according to Ugelstad et al.\textsuperscript{25} and the reactivity ratios, they calculated the amount of \(\text{MA}\) needed in the initial charge. Then they carried out an experiment where they added the remaining part of \(\text{MA}\) at a constant rate and calculated \(\eta\) with help of a model and the experimental results of this first reaction. Then \(\eta\) was correlated with the volume fraction of polymer in the particles. This correlation was used to calculate the optimal addition rate profile. If the results of the reaction that was carried out with the new addition rate profile indicated that the copolymer composition deviated from the desired value, then the procedure was repeated. The authors pointed out that the dependence of \(\eta\) on the volume fraction of polymer was the critical one. This was refuted by us,\textsuperscript{14} and we stated that \(\eta\) is also dependent on a number of other variables, especially when \(\eta\) is lower than 0.5. Arzamendi et al.\textsuperscript{15} also studied a system where \(\eta\) was lower than 0.5, and they correlated \(\eta\) to the volume fraction of polymer and particle diameter. We used another approach, i.e., we
compared the experimentally found addition profile with the calculated optimal addition profile in each iteration step. (Note that an addition rate profile is not the same as an addition profile. This will be explained below.) This means that it is not necessary to calculate \( \bar{n} \) in each iteration step. The procedure is recalled briefly hereafter, but a more detailed description was given earlier.\(^{14}\)

(i) We calculated the amount of S that is needed to produce the right copolymer at the beginning of the reaction (as did Arzamendi and Asua\(^{12}\)), and we determined a correlation between amount of S to be added \((A^*_S)\) and instantaneous molar conversion \((x_{\text{inst}})\): \(A^*_S = g(x_{\text{inst}})\), the calculated optimal addition profile (where * indicates profile is optimal). This correlation is unique and dependent only on reactivity ratios and monomer partitioning. Only if the reaction is carried out in such a way that the experimental dependency of added amount of S on instantaneous molar conversion corresponds exactly to the calculated dependency \(A^*_S = g(x_{\text{inst}})\), the fraction of S in the particles will be constant and have the desired value, and the copolymer will be homogeneous. To meet this condition, one needs to find an optimal addition rate profile \([A^*_S = h(t)]\).

(ii) The next step is to carry out a reaction and determine the conversion-time curve, \(x_{\text{inst}} = f(t)\).

(iii) The experimental conversion-time curve is combined with \(A^*_S = g(x_{\text{inst}})\) to give a better addition rate profile \(A = h(t)\).

This new addition rate profile is then used in step (ii) until the profile is not significantly different from the previous one.

In the present study, we applied this procedure with the exception that in the present case \(x_{\text{inst}}\) was not the instantaneous molar conversion but the instantaneous weight conversion, which means that it is not necessary to follow the partial molar conversions of the monomers with gas chromatography. Furthermore, the initial runs that were carried out in each iteration, were not based on the assumption that \(\bar{n} = 0.5\), as did van Doremaele et al.,\(^{14}\) but were done with a constant addition rate of S, as did Asua et al.\(^{12}\)

We determined the optimal addition rate profile for homogeneous copolymers with 25, 50, and 80 mol % S. As stated in the Introduction, the addition rate profile for 25 mol % S was determined previously,\(^{14}\) but in this case we used a different seed latex (different particle diameter). Therefore, the polymerization rate is different and also the addition rate profile. We used sodium persulfate as initiator in stead of potassium persulfate, but we applied the same molar concentration.

In Figure 6, we plotted the results of the calculations with help of a model simulation (SIEMCO\(^1\)) to obtain the correlation \(A^*_S = g(x_{\text{inst}})\) for each composition. Then we performed the iteration procedure as described above. In the initial runs for 50 and 80 mol % S we added the S at a constant rate based on the reaction rate of the corresponding batch reactions. The addition rate profile of the initial run for 25 mol % S was based on the addition rate profile that was obtained before.\(^{14}\) In all cases we obtained an optimal addition rate profile after three or four iteration steps, comparable to what was found earlier for 25 mol % S. Figure 7 shows the experimental results of the iteration procedures, i.e., \(A_S\) versus instantaneous conversion for the three compositions. For comparison we also depicted the calculated optimal addition profiles. As can be seen from both Figures 6 and 7, the difficulty with the profile for 80 mol % S was to find the time at which the reaction reached the conversion at which the addition had to be started.

The best impression of such an iteration procedure is obtained by looking at the CCDs of all these reactions as depicted in Figures 8a, 8b, and 8c corresponding to the reactions with 25, 50, and 80 mol % S. One can clearly see that the homogeneity of the copolymers produced improves with every iter-
number of runs could have been smaller. So it seems that judging the optimality is best done by looking at the CCDs rather than at the addition profiles. The reactions performed to obtain copolymers with 80 mol % S, as mentioned before, being carried out under almost azeotropic conditions, all resulted in homogeneous copolymers, notwithstanding the fact that the addition profiles differed considerably.

Figure 7  Representation of the successive experimental reactions in the iteration procedure to find the optimal addition profiles. Amount of S to be added ($A_S$) as function of instantaneous conversion ($x_{\text{inst}}$). (△) 1st run; (○) 2nd run; (□) 3rd run; (+) 4th run; calculated optimal addition profile (●) (see also Fig. 6). (a) $F_S = 0.25$; (b) $F_S = 0.50$; (c) $F_S = 0.80$.

Figure 8  Experimentally determined CCDs of successive reactions in the iteration procedure to find the optimal addition rate profiles. Run numbers are indicated in the figures, except in Figure 8(c). (a) $F_S = 0.25$; (b) $F_S = 0.50$; (c) $F_S = 0.80$. $R$ designates relative abundance (arbitrary units).
In Figure 9 we depicted the dependency of \( \bar{n} \) on conversion for the reactions with optimal addition rate profiles. The calculations were based on the penultimate model.\(^{23} \) Clearly this dependency is influenced by the ratio \( S/MA \). However, if the ratio \( S/MA \) is different in the particles, it is also different in the water phase. So by changing the overall ratio, the ratios in both particle and aqueous phase change, and therefore it is not possible to draw any mechanistic conclusions. Furthermore, the particle number is not the same in each reaction. Only a more detailed kinetic study could perhaps reveal some kinetic parameters.

**Addition Rate Profile for Heterogeneous Copolymers**

Heretofore all efforts were aimed at producing homogeneous copolymers. However, it might equally well be that one wishes to make copolymers that have a broad composition distribution rather than a narrow distribution. Whatever the shape of the CCD one wants to obtain, the reaction will have to be controlled in a similar way as shown, necessary to produce homogeneous copolymers. In order to illustrate this hypothesis, we attempted to make a copolymer with a CCD of predefined broadness profile.

In order to be able to compare the properties of such an emulsion copolymer with a copolymer with the same average composition but with a narrow distribution, we tried to make a copolymer consisting of 25 mol % \( S \) on the average, but with a composition distribution ranging from 0 to 50 mol % \( S \). In Figure 10 we depicted schematically the CCD of such a copolymer.

**Figure 10** Schematic CCD of a copolymer of hypothetical heterogeneity.

To accomplish this, we applied the following strategy, comparable to that applied for the preparation of the homogeneous copolymers. In the beginning of the reaction, poly(methyl acrylate) (PMA) is formed, and slowly the instantaneous composition of the copolymer is shifted towards higher fractions of \( S \), in fact, a composition drift that is reversed compared to the drift occurring in batch emulsion copolymerizations. The highest instantaneous fraction of \( S \) incorporated has to be 50 mol %. If one would try to make the same copolymer in reversed order, i.e., starting at 50 mol % \( S \) and ending at 0 mol % \( S \), one would have to add too much MA in the beginning, leaving no MA to add to control the reaction. This is a direct consequence of the copolymerization behaviour of \( S \) and MA.

With help of mass balances, thermodynamic cor-

**Figure 9** Average number of radicals per particle (\( \bar{n} \)) as a function of overall conversion for the reactions with the optimal addition rate profiles: (+) \( F_S = 0.25 \); (○) \( F_S = 0.50 \); (□) \( F_S = 0.80 \).

**Figure 11** Experimentally determined CCDs of successive copolymer products in the iteration procedure to find the addition rate profile for the heterogeneous copolymer defined in Figure 10. Run numbers are indicated in the figure. \( R \) designates relative abundance (arbitrary units).
relations, and reactivity ratios, we calculated the dependency $A_S^x = g(x_{\text{inst}})$. The further procedure is the same as with the optimal addition rate profiles.

The first run was based on a reaction where the total amount of MA was added to the reactor, and S was added at a constant rate. Figure 11 shows the CCD of this reaction (run no. 1). One can see that the distribution already resembles vaguely the one we are looking for. However, the difficulty in the present case seems to be the presence of all the MA at the beginning of the reaction. Since MA is a very fast reacting monomer, it is very hard to find the optimal addition rate profile for the first part of the reaction. The CCDs of the first reactions manifest this by showing a relatively large PMA peak. To improve the CCD, we carried the iteration procedure (runs 2 and 3 in Fig. 11). But these runs did not seem to get much closer to the desired CCD. In Figure 12 we depicted the amount of S to be added versus instantaneous conversion for the three successive reactions. The figure illustrates that the first profile is already comparable to the calculated addition rate profile and that the first part of the profiles improve with increasing run number. This may indicate that the iteration procedure is only very slow. Considering this, we decided not to continue it. We think that if one were to make a heterogeneous copolymer with an average composition of 50% S, with $F_S$ ranging from 0.25 to 0.75, the procedure would be easier to carry out.

**Molecular Weights**

As stated earlier in this work we added NDM to minimize the effect of addition strategy on the

![Figure 12](image)

**Figure 12** Representation of the successive experimental reactions in the iteration procedure to find the addition rate profile for the heterogeneous copolymer defined in Figure 10. Amount of S to be added ($A_S$) as function of instantaneous conversion ($x_{\text{inst}}$). (Δ) 1st run; (O) 2nd run; (□) 3rd run; (●) calculated addition rate profile.

**Table IV Molecular Weights of Some of the S-MA Copolymers Measured with GPC**

<table>
<thead>
<tr>
<th></th>
<th>$F_S$</th>
<th>$M_n$ (kg/mol)</th>
<th>$M_w/M_n$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Batch</td>
<td>0.25</td>
<td>48.6</td>
<td>2.3</td>
</tr>
<tr>
<td>Batch, no NDMb</td>
<td>0.25</td>
<td>476</td>
<td>7.5</td>
</tr>
<tr>
<td>Starved conditions (8 h)</td>
<td>0.25</td>
<td>56.6</td>
<td>2.5</td>
</tr>
<tr>
<td>Optimal addition</td>
<td>0.25</td>
<td>29.0</td>
<td>2.1</td>
</tr>
<tr>
<td>Batch</td>
<td>0.50</td>
<td>67.7</td>
<td>2.5</td>
</tr>
<tr>
<td>Starved conditions (8 h)</td>
<td>0.50</td>
<td>54.6</td>
<td>2.4</td>
</tr>
<tr>
<td>Optimal addition</td>
<td>0.50</td>
<td>33.1</td>
<td>2.1</td>
</tr>
<tr>
<td>Batch</td>
<td>0.80</td>
<td>34.6</td>
<td>2.8</td>
</tr>
<tr>
<td>Starved conditions (8 h)</td>
<td>0.80</td>
<td>44.9</td>
<td>3.3</td>
</tr>
<tr>
<td>Optimal addition</td>
<td>0.80</td>
<td>60.7</td>
<td>3.3</td>
</tr>
</tbody>
</table>

* In all cases NDM was added unless indicated otherwise. $M_n$ is the number average molecular weight, $M_w$ the weight average molecular weight, and the polydispersity is the ratio of these.

b The recipe for this reaction is the same as for the batch reaction with NDM.

MMD. We measured the MMDs of all products mentioned in this work with gel permeation chromatography (GPC). In Table IV, the molecular weights are given of several copolymers. It can be clearly seen that the molecular weights are relatively small, compared to a batch reaction carried out without NDM, but, more importantly, there is no obvious effect of addition strategy on MMD. The number average molecular weights ($M_n$) vary somewhat, but show no clear trend. One could perhaps suggest a trend in the polydispersity ($M_w/M_n$), which seems to increase going from $F_S = 0.25$ to $F_S = 0.80$. The polydispersity is in all cases much lower than what is obtained without NDM, which indicates that adding NDM is a good way to avoid large differences in MMD.

The copolymers mentioned in this work will be used in a study of the effect of the shape of the CCD on the mechanical properties.

**CONCLUSIONS**

Batch emulsion copolymerization of styrene and methyl acrylate leads to copolymers with broad chemical composition distributions or even bimodal distributions. Homogeneous emulsion copolymers of styrene and methyl acrylate with various fractions of styrene can be produced via two different strategies. Using constant addition rates under starved conditions leads to homogeneous copolymers if long addition times are used. Applying optimal addition rate profiles (adding all the methyl acrylate to the reactor at the beginning and adding the styrene at a rate that varies in time) results in homogeneous
copolymers but with reaction times comparable to
the times needed for the batch reactions. The in-
formation about composition drift in batch reactions
is seen to be useful in explaining how quickly an
optimal addition rate profile can be found. It is
shown that HPLC is of great value in determining
whether an optimal addition rate profile is really
optimal. The reactions with optimal addition rate
profiles did not give enough information to draw
conclusions about the influence of the ratio styrene/
methyl acrylate on $\bar{n}$. The strategy used to produce
homogeneous copolymers with optimal addition rate
profiles can be applied to the production of copol-
ymers with a predefined broad CCD. However, in
our case the iteration procedure seemed to be very
slow. If NDM is used, the molecular weights are quite
low in all instances, and there is no obvious effect
of addition strategy, though the polydispersity seems
to increase with increasing mole fraction S.

Eventually the homogeneous and heterogeneous
copolymers with varying fractions of styrene will be
used to demonstrate the effect of the shape of the
CCDs and average chemical composition on the me-
chanical properties of these kinds of emulsion co-
polymers.

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