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by

S. Floyd
K. Y. Choi
T. W. Taylor
W. H. Ray
CHEMICAL PROCESS SYSTEMS
ENGINEERING LABORATORY

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W.H. Ray

A CONSTITUENT LABORATORY OF
THE SYSTEMS RESEARCH CENTER

THE UNIVERSITY OF MARYLAND
COLLEGE PARK, MARYLAND 20742
Polymerization of Olefins through Heterogeneous Catalysis. III. Polymer Particle Modelling with an Analysis of Intraparticle Heat and Mass Transfer Effects

S. FLOYD, K. Y. CHOI, T. W. TAYLOR and W. H. RAY*, Department of Chemical Engineering, University of Wisconsin, Madison, Wisconsin 53706

Synopsis

Propylene and ethylene polymerization in liquid and gas media are described by a multigrain particle model. Intraparticle heat and mass transfer effects are investigated for a range of catalyst activities. For slurry polymerization, intraparticle mass transfer effects may be significant at both the macroparticle and microparticle level; however, for normal gas phase polymerization, microparticle mass transfer effects appear more likely to be important. Intraparticle temperature gradients would appear to be negligible under most normal operating conditions.

INTRODUCTION

The kinetics of polymerization of olefins over heterogeneous catalysts is not yet understood in detail. There is still great controversy concerning the surface phenomena and the nature of the active centers for reaction. In addition, from a reaction engineering standpoint, it is of significance that the effects of heat and mass transfer limitations during polymerization have not been clarified. Recent theoretical and experimental studies by our research group1-7 have dealt with questions of kinetics and physical transport limitations in these polymerizations. These and other studies provide evidence that, in some circumstances, significant diffusion resistance to monomer transport will exist, and this can mask the intrinsic rate constants of the catalyst. In addition to mass transfer effects, there exists the possibility of inadequate removal of the heat of polymerization from the growing polymer particle. This may result in temperature gradients within the particle.

In modern polyolefin processes which utilize highly active Ziegler-Natta catalyst systems, it is often reported that particle sintering or agglomeration occurs in the polymerization reactor due to poor heat removal from the reacting catalyst/polymer particles. This suggests that there may also exist a significant temperature difference between the solid phase and bulk fluid phase during the reaction. This series of papers will be concerned with the quantitative analysis of both external film and intraparticle heat and mass transfer limitations through detailed mathematical modelling.

Several recent papers have reported modelling studies with nonisother-
Polymer particles. McGeary and Rawlings simulated both particle fluid and intraparticle temperature nonuniformities and concluded from their model that nonisothermal effects could be important during the first 30–60 min of reaction. Nagel et al. analyzed the temperature history of low activity catalysts (such as Stauffer AA 1.1) in slurry polymerization and concluded that temperature gradients in the particle were negligible. On the other hand, Wisseroth and Brockmeier studied particle/fluid temperature differences and concluded that significant temperature differences could exist between the particle and the surrounding fluid under some circumstances. Choi et al. carried out a detailed analysis of nonisothermal effects and summarized their results at the 1982 IUPAC meeting. Most recently, Laurence and Chiovetta analyzed heat and mass transfer effects and concluded that temperature transients are an important factor during catalyst breakup. Both Choi et al. and Laurence and Chiovetta showed that the temperature inside the particle could reach the melting point of the polymer under some circumstances for gas phase polymerization. It is the purpose of the present series of papers to extend the analysis reported in Refs. 1 and 4 so as to more clearly define the conditions under which intraparticle gradients are expected to arise and when particle–fluid mass transfer limitations and particle overheating are expected to occur. This paper will deal with intraparticle gradients, and the companion paper will treat boundary layer heat and mass transfer resistances.

**POLYMER PARTICLE MODELS**

Polymer particles in Ziegler–Natta polymerization have several resistances to heat and mass transfer. Assuming the macroscopic reactor is completely mixed and at uniform temperature, the first heat and mass transfer resistances are present in the particle boundary layer. Next, monomer must diffuse through the particle to the catalyst active sites to react. Heat generated from the polymerization is transferred by conduction to the catalyst particle surface and then by convection through the boundary layer to the fluid. The analysis of this system is complicated by the fact that as the reaction proceeds polymer is formed and the particle grows.

The catalyst particle model used for detailed simulations of these effects is the multigrain model shown schematically in Figure 1. This model was originally suggested by Yermakov et al. to estimate concentration profiles in the polymer particle and has been used in our work to predict the yield and the molecular weight distribution of the polymer product. This model was also adopted by Laurence and Chiovetta for detailed simulations of behavior at short times. The model structure is based on numerous experimental observations that the original catalyst particle quickly breaks up into many small catalyst fragments ("primary crystallites") which are dispersed throughout the growing polymer. Thus, the large macroparticle is comprised of many small polymer particles (microparticles), which encapsulate these catalyst fragments. In this idealized picture, all microparticles at a given large particle radius are assumed to be the same size. As illustrated in Figures 1–3, for monomer to reach the active sites, there is
THE MULTIGRAIN MODEL

Fig. 1. The multigrain model.

Fig. 2. Concentration and temperature gradients in the macroparticle.
both macrodiffusion in the interstices between microparticles and microdiffusion within the microparticles. In general, the effective diffusion coefficients for the two regimes are not equal. We also include the possibility of an equilibrium sorption of monomer at the surface of the microparticle.

**Material Balances**

To model the particle, relations must be developed between the monomer concentrations in the large and small particles and the radial shell growth, particle yield, and temperature. The governing equation for the diffusion of monomer in the macroparticle is

\[
\epsilon_i \frac{\partial M_i}{\partial t} = \frac{1}{r^2} \frac{\partial}{\partial r_i} \left( D_i r_i \frac{\partial M_i}{\partial r_i} \right) - R_c
\]

where \( \epsilon_i \) is the large particle porosity, \( M_i(r_i, t) \) is the monomer concentration in the pores of the macroparticle, and \( D_i \) is the pseudobinary macrodiffusion coefficient. The reaction rate term \( R_c \) represents the total rate of consumption of monomer in an infinitesimal spherical shell at a given radius of the macroparticle. The boundary and initial conditions are

\[
r_i = 0, \quad \frac{\partial M_i}{\partial r_i} = 0
\]

\[
r_i = R_i, \quad D_i \frac{\partial M_i}{\partial r_i} = k_p(M_b - M_i)
\]

where \( k_p \) is the propa catalyst sites, and \( M_c \)

As mentioned above assumed to be imper. shell. The polymerize the catalyst core. An takes the form

\[
\rho_p C_v
\]

with boundary condit

\[
\text{Pol' where } M_s \text{ is the bulk transfer coefficient in tration at the external}
\]

For the microparticle

\[
\text{where } M(r, t) \text{ is the m pseudobinary microparticle, all of the catalyst core at } r = r_c \text{ by}
\]

\[
\text{where boundary cond}
\]

\[
\text{librium at the surface particle radius, } R_i \text{ is i merization at the catalyst cores, and } M_c \]

\[
The polymerization rate at the catalyst core. An takes the form
\]

\[
\rho_p C_v
\]

with boundary condit

\[
r = \]
where $M_\ast$ is the bulk monomer concentration in the reactor, $k_0$ is the mass transfer coefficient in the external film, and $M_s$ is the monomer concentration at the external solid surface.

For the microparticles, the monomer diffusion equation is given as

$$
\varepsilon_0 \frac{\partial M}{\partial t} = \frac{1}{r^2} \frac{\partial}{\partial r} \left( D_0 r^2 \frac{\partial M}{\partial r} \right)
$$

\hspace{1cm} (r_c \leq r \leq R_s)

where $M(r, t)$ is the monomer concentration in the microparticle, $D_0$ is the pseudobinary microdiffusion coefficient, and $\varepsilon_0$ is the porosity. In the microparticles, all of the active sites are assumed to be at the surface of the catalyst core at $r = r_c$. Thus, the boundary and initial conditions are given by

$$
r = r_c, \quad 4\pi r^2 D_0 \frac{\partial M}{\partial r} = \frac{4}{3} \pi r^3 R_m
$$

\hspace{1cm} (6)

$$
r = R_s, \quad M = M_{\infty}(M_i) \leq M_i
$$

\hspace{1cm} (7)

$$
t = 0, \quad M = M_{\infty}
$$

\hspace{1cm} (8)

where boundary condition (7) allows for the possibility of a sorption equilibrium at the surface of the microparticles. Here $r_c$ is the catalyst primary particle radius, $R_s$ is the microparticle radius, and $R_m$ is the rate of polymerization at the catalyst particle surface given by

$$
R_m = k_p C_\ast M_c
$$

\hspace{1cm} (9)

where $k_p$ is the propagation rate constant, $C_\ast$ is the concentration of active catalyst sites, and $M_c$ is the monomer concentration at the catalyst surface.

### Energy Balances

As mentioned above, the microparticle consists of a solid catalyst core assumed to be impermeable and encapsulated by a catalyst-free polymer shell. The polymerization reaction occurs only at the external surface of the catalyst core. Analogous to eq. (5), the microparticle energy balance takes the form

$$
\rho_\ast c_\ast \frac{\partial T}{\partial t} (r, t) = k_\ast \left[ \frac{1}{r^2} \frac{\partial}{\partial r} \left( r^2 \frac{\partial T}{\partial r} \right) \right]
$$

\hspace{1cm} (r_c < r \leq R_s)

\hspace{1cm} (10)

with boundary conditions

$$
r = r_c, \quad 4\pi r^2 k_\ast \frac{dT}{dr} = (-\Delta H_r) \frac{4}{3} \pi r^3 R_m
$$

\hspace{1cm} (11)

$$
r = R_s, \quad T = T_i
$$

\hspace{1cm} (12)
Similarly, an energy balance over the macroparticle takes the form

\[ \rho_r C_r \frac{\partial T_l}{\partial t} = \left[ \frac{1}{r_l^2} k_r \frac{\partial}{\partial r_l} \left( r_l \frac{\partial T_l}{\partial r_l} \right) \right] + (-\Delta H_p) R_v \]

\[ 0 < r_l < R_l \]

where the volumetric reaction rate \( R_v \) may include the effects of macroparticle heat and mass transfer resistance. For the macroparticle, the boundary and initial conditions are

\[ r_l = 0, \quad \frac{\partial T_l}{\partial r_l} = 0 \]

\[ r_l = R_l, \quad k_r \frac{\partial T_l}{\partial r_l} = h \left( T_b - T_l \right) \] (15a)

or

\[ r_l = R_l, \quad T_l = T_s \]

\[ t = 0, \quad T_l = T_{ib} \] (15b)

These equations must be solved together with eqs. (1-4) to predict the concentration and temperature distribution in the macroparticle. This is illustrated pictorially in Figures 2 and 3.

**Particle Parameter Values**

In the analysis to follow, the practical conclusions will depend on the range of parameter values one might encounter for a polymer particle. For ethylene polymerization and propylene polymerization in slurry or gas phase reactors, these are tabulated in Table I. For some parameters (e.g., for \( k_p, C_0, D_a, D_l, R_v, R_i \)) there are a range of values which arise, and this range is indicated in the table. Both the microscale pseudobinary diffusivity \( D_a \) and macroscale pseudobinary diffusivity \( D_l \) play crucial roles in estimating intraparticle temperature and concentration gradients. However, it is difficult to determine the value of these diffusivities precisely. Thus let us discuss how one may arrive at the most likely values.

For macroscale diffusion through the interstices between the microparticles, we may estimate the effective diffusivity \( D_l \) as one would for more conventional heterogeneous catalysts.\(^4\) The effective diffusion coefficient in a solid may be represented by the bulk diffusivity in the fluid, \( D_a \), multiplied by \( \epsilon/\tau \), where \( \epsilon \) is the porosity of the solid and \( \tau \) is a "tortuosity factor," i.e., \( D_l = D_a (\epsilon/\tau) \). In liquid slurry polymerization, the slurry diluent permeates the pores of the macroparticle and the appropriate bulk diffusivity, \( D_a \), is the diffusivity of monomer in the diluent, which is of the order of \( 8 \times 10^{-6} \text{ cm}^2/\text{s} \) under industrial conditions (see Table I). The porosity of the catalyst or polymer particle is 0.4 or less, and values of \( \tau \) in the range 2–7 are common.\(^4\) Thus, the value of macroparticle diffusivity for slurry polymerization would be in the range of \( 10^{-6} \leq D_l \leq 10^{-5} \text{ cm}^2/\text{s} \). For gas
<table>
<thead>
<tr>
<th>Property</th>
<th>Slurry (n-heptane)</th>
<th>Gas</th>
<th>Slurry (n-hexane)</th>
<th>Gas</th>
<th>References</th>
</tr>
</thead>
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<tr>
<td>$M$, mol·L⁻¹</td>
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<td>2.0</td>
<td>1.0</td>
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<td>6</td>
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<td>28</td>
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<tr>
<td>Mol fraction monomer</td>
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<td>0.26</td>
<td>1</td>
<td>19.22</td>
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<td>0.3</td>
<td>0.3</td>
<td>0.3</td>
<td>17.18, 23</td>
</tr>
<tr>
<td>$E$, kcal·mol⁻¹</td>
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<td>0.3</td>
<td>0.3</td>
<td>0.3</td>
<td>17.18, 23</td>
</tr>
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<td>2640</td>
<td>2000-4000</td>
<td>4000</td>
<td>6</td>
</tr>
<tr>
<td>$k_s$ (mol sites·L·cat⁻¹)</td>
<td>$10^{-10}·10^{-1}$</td>
<td>$5·10^{-5}$</td>
<td>$10^{-3}·10^{-1}$</td>
<td>$4·10^{-4}$</td>
<td>17.18, 20</td>
</tr>
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<td>$2.6·10^{-4}$</td>
<td>$2.3·10^{-4}$</td>
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<td>0.61</td>
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<td>0.07</td>
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<tr>
<td>$\mu_{icp}$</td>
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<td>0.135</td>
<td>0.135</td>
<td>0.135</td>
<td>14.15</td>
</tr>
</tbody>
</table>

---

1. Values correspond to conditions of industrial operation\(^22,24\) (pressures may be higher or lower for individual processes).
2. Calculated using Russell's equation\(^6\) with a porosity of 0.37.
3. Calculated using the Kendal-Monroe equation.\(^5\)
4. Polymerizing particles grow in diameter by a factor of 10 or more compared to original catalyst size. Some modern catalysts are capable of producing polymer particles 2 mm or more in diameter.\(^7\)
cm\(^2\)/s for vapors such as n-hexane, cyclohexane, and benzene. The value of \(D_0\) in amorphous polypropylene was somewhat lower in the case of benzene. The diffusion coefficient for such vapors depends significantly on the concentration; Michaels and Bixler\(^{27}\) found that the diffusion coefficient increased by 2 orders of magnitude between zero penetrant concentration and saturation for vapors such as benzene and hexane in polyethylene at 25°C. For linear polyethylene, the measured \(D_s\) values at saturation were of the order of \(10^{-7}\) cm\(^2\)/s. \(D_s\) for heptane in polypropylene was found to be \(\sim 10^{-8}\) cm\(^2\)/s at reaction conditions.\(^{30}\) In slurry polymerization, the microparticles are in contact with the diluent liquid and are probably saturated by it. Thus, swelling may be considered to occur, and the microdiffusion coefficient might be higher than for gas phase polymerization.

In summary, it is reasonable to surmise that the value of \(D_s\) lies in the range \(10^{-8}-10^{-4}\) cm\(^2\)/s for diffusion of monomer ethylene or propylene in polyethylene or polypropylene and copolymers under reaction conditions. For highly isotactic polypropylene or high density polyethylene, the lower end of the range would apply, while for copolymers (such as ethylene-propylene copolymers) the upper end would apply. Polymerization at low temperatures would mean lower values of \(D_s\), while polymerization at high temperatures would indicate higher \(D_s\) values. Because increased monomer sorption increases \(D_s\), polymerizing under high pressures in gas phase or slurry would give higher \(D_s\) values while low pressure lab reactors should have a lower \(D_s\) value. Also since diluents such as heptane or hexane also swell the polymer and aid diffusion, slurry phase values of \(D_s\) would be expected to be larger than for gas phase polymerization. Obviously larger comonomers such as hexane and octene would have diffusivities towards the lower end of the range of \(D_s\). Higher values of the diffusivity would be expected for hydrogen, \((D_s, \sim 10^{-8})\) while smaller values are anticipated for diffusion of organoaluminum compounds. For HDPE or highly isotactic polypropylene at typical industrial temperatures and pressures in gas phase, one would expect \(D_s\) to lie in the range \(1-5 \times 10^{-8}\) cm\(^2\)/s.

An additional issue is the process of sorption of monomer (or other fluids) by the polymer of the microparticles. According to Michaels and Bixler,\(^{31}\) there is negligible sorption in the crystalline portion of the polymer although macroscopically the polymer is completely homogeneous and isotropic with respect to dissolution and diffusion. Thus, the solubility in partially crystalline polymer will be proportional to the amorphous content. From the work by Michaels and Bixler,\(^{31}\) one may approximately model the equilibrium sorption of monomers from the gas phase as

\[
M_m = kP
\]

where \(k\) is a Henry's law constant and \(P\) is the gas partial pressure in the pores. For partially crystalline polymer, \(k = \alpha k^*\), where \(\alpha\) is the amorphous content of the polymer and \(k^*\) is the solubility constant for purely amorphous polymer. From literature data,\(^{30,31}\) values of \(k^*\) are \(~0.04\) mol/L atm for solubility of ethylene in polyethylene and \(~0.16\) mol/L atm for propylene in polyethylene. Values of \(k^*\) for propylene in polypropylene are probably comparable. From the experiments in Ref. 31, the solubility of
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Fig. 9. Regimes for macroparticle diffusion resistance and temperature gradients. $D_i \Phi_e$ vs. observed catalyst activity. Approximate values for typical catalyst if $M_g = M_e$: (a) ethylene slurry polymerization, low and high activity catalyst; (b) propylene gas phase polymerization, low and high activity catalyst; (c) ethylene slurry polymerization, low and high activity catalyst; (d) ethylene gas phase polymerization, low and high activity catalyst (low activity, $R_{ob} = 400$ g/g cal h, high activity, $R_{ob} = 4000$ g/g cal h under representative industrial conditions).

0.02, in terms of the polymerization parameters. Here we may define an overall observed reaction rate $R_{ob}$, which accounts for both microparticle and macroparticle diffusion limitations as well as for equilibrium sorption

$$R_{ob} = \eta_i \eta_i \eta_{in} R_{kin}$$

(40)

where $R_{kin}$ is defined by eq. (24) with $T_{ref} = T_s, M_{ref} = M_g$ in this case. Using the definition of $a_i$, we see that

$$a_i = \left( \frac{p_i R_{ob} (R_i)^2}{3.6 M_0 M W D_i \Phi_e \eta_i} \right)^{1/2}$$

(41)

so that lines corresponding to $a_i = 1, \eta_i = 1$ can be represented in simple terms. Also plotted is the line $D_i = 10^{-4}$ corresponding to the criterion $\beta < 0.02$ for which there are negligible temperature gradients. Note that it is only for extremely high catalyst activities and large catalyst particles that significant macroparticle temperature gradients could exist (indicated by curves in the shaded region of Figure 9).

To illustrate how to use Figure 9, suppose that one has a 20 $\mu$m diameter catalyst particle with $R_{ob} = 4000$ g/g cat h for ethylene gas phase polymerization when the ethylene pressure is 27 atm ($M_e = 1$ mol/L). If we assume $M_g = M_e$, then $R_{ob}/M_g MW = 143$. Thus this catalyst corresponds to point 1 in Figure 9 where $D_i \Phi_e = 10^4, D_i = 10^4$. Thus, for $D_i \Phi_e$ values $> 10^4$, there is negligible heat and mass transfer resistance in the macroparticle. If we assume $D_i \sim 10^4$ as a conservative estimate, then we conclude that there will be no significant internal concentration and temperature gradients in the macroparticle such that $D_i \Phi_e > 10^{-2}$. If significant internal temperature had grown by a factor of 10, the macroparticle diameter catalyst will have some effect.

If we consider gas phase polymerization at the Stauffer rate represented by point 3, $D_i \Phi_e = 10^{-6}$ for the onset of gas phase polymerization, we will have no significant mass transfer resistance in the macroparticle; however, the particle size will be significant mass transfer resistance. Having shown that in the macroparticle unanalyzed for macroparticle an isothermal particle, (37) has the solution

$$\frac{R_{ob}/M_g MW (g/g - cat \cdot hr)/(mol/fl) (g/mol)}{10^2}$$

where $z$ and $a_i$ are defined factor, $\eta_i$, defined by $a_i$.

From the definition of (41) and (42) graphically illustrate catalyst particles, one smaller catalyst particle size, growth factor $\Phi_e$.

It is interesting to note that in propylene slurry polymerization (e.g. size of 60 $\mu$m), reasonable to cause macroparticle growth rate catalyst in g not expect to see this behavior where a catalyst polymerization with high activity is true for high activi...
temperature gradients in the macroparticle. However, note that a 60 \mu m diameter catalyst particle with the same observed productivity (point 2) will be expected to have negligible internal heat and mass transfer resistance for \( D_I \Phi_y > 10^{-3} \). Thus if we assume \( D_I = 10^{-4} \), then one would expect significant internal temperature and concentration gradients until the polymer had grown by a factor of 10 (\( \Phi_y = 10 \)).

If we consider gas phase propylene polymerizations with a low activity catalyst of the Stauffer AA type with a catalyst diameter of 60 \mu m, this is represented by point 3 in Figure 9 corresponding to \( D_I \Phi_y, D_I \) values of \( \sim 5 \times 10^{-6} \) for the onset of diffusion resistance. Thus since \( D_I > 10^{-4} \) for gas phase, we will have negligible temperature and concentration gradients in the macroparticle for this catalyst in gas phase. If we use a 60 \mu m diameter catalyst for the slurry polymerization of propylene with the same observed productivity, \( R_{\text{mac}} \), this corresponds to point 4 in Figure 9 where \( D_I = 10^{-5}, D_I \Phi_y = 10^{-6} \) for the onset of diffusion resistance. For slurry \( 10^{-4} \leq D_I \leq 10^{-5} \text{ cm}^2/\text{s} \). Thus there will not be any temperature gradients in the macroparticle; however, with an actual value of \( D_I \sim 10^{-6} \), there will be significant mass transfer resistance due to diffusion—at least until the particle size exceeds \( \Phi_y > 10 \).

Having shown that intraparticle temperature gradients will be negligible in the macroparticle under most normal operating conditions, we may now analyze for macroparticle mass transfer limitations in more detail assuming an isothermal particle. For this case, the macroparticle material balance (37) has the solution

\[ c = \sinh(a_1 z) / z \sinh(a_1) \]

where \( z \) and \( a_1 \) are defined by Eq. (38). Thus, the macroparticle effectiveness factor, \( \eta_1 \), defined by eq. (39), has the solution \(^{41}\)

\[ \eta_1 = \frac{3}{a_1} \left[ \frac{1}{\tanh(a_1)} - \frac{1}{a_1} \right] \]  

(42)

From the definition of \( a_1, R_{\text{mac}} \) in eqs. (40) and (41) one may represent eqs. (41) and (42) graphically as shown in Figures 10–12. Note that, for larger catalyst particles, one expects more serious diffusion limitations than for smaller catalyst particles at the same growth factor. However, for a fixed catalyst particle size, diffusion limitations are reduced with increasing growth factor \( \Phi_y \).

It is interesting to note from Figure 11 that for a low activity catalyst in propylene slurry (e.g., Stauffer AA type catalyst with average catalyst size of 60 \mu m), reasonable values of \( D_I \) (e.g., \( 10^{-6} - 10^{-5} \text{ cm}^2/\text{s} \)) are sufficient to cause macroparticle diffusion limitations. However, for the same productivity catalyst in gas phase (where \( D_I \approx 10^{-4} - 10^{-3} \text{ cm/s} \)), one would not expect to see macroparticle diffusion limitations. As indicated in Figure 10, where a catalyst particle size of \( d_p = 20 \mu m \) is used for ethylene polymerization with a high activity catalyst, one sees that the same conclusion is true for high activity catalysts; i.e., the possibility of significant intrap-
particle concentration gradients for slurry polymerization, but not for gas phase polymerization. However, as indicated in Figure 12, for gas phase polymerization with large high activity catalysts, internal concentration gradients can be significant. Furthermore, if catalyst activity continues to improve and catalyst particle sizes increase dramatically, then internal gradients will become even more significant in gas phase processes.

In this paper, graphical form to all tration and tempera merization. Because activity, these crite rions when there is macroparticle diffus and 6. Similarly, ma if there is a significa estimates of \( M_0 \) are

From the analysis ditions normally en temperature gradien the macroparticles would be for large l the polymer particle concentration gradit
Fig. 12. Regimes for isothermal macroparticle diffusion resistance, $D_i/\Phi_i$ vs. observed catalyst activity for catalyst with initial particle size $d_i = 100\,\mu$m. Approximate values for typical catalyst if $M_S = M_S$: (a,A) propylene slurry polymerization, low and high activity catalyst; (b,B) propylene gas phase polymerization, low and high activity catalyst; (c,C) ethylene slurry polymerization, low and high activity catalyst; (d,D) ethylene gas phase polymerization, low and high activity catalyst (low activity, $R_{an} = 400\,\text{g/g cat h}$, high activity, $R_{an} = 4000\,\text{g/g cat h}$ under representative industrial conditions).

It is clear from Figures 10–12 that as the polymer particle size increases, the diffusion resistance becomes smaller. Thus, diffusion control in slurry could manifest itself in terms of an acceleration or hybrid-type rate behavior, as has been observed under many circumstances, even for catalysts of low activity. Note that these results predict that a significant effect of catalyst particle size on the polymerization rate may be anticipated for slurry polymerization. This effect would primarily be seen in the initial rate for low activity catalyst, but there could be a significant effect on the overall yield for high activity catalysts.

CONCLUSIONS

In this paper quantitative criteria have been developed and presented in graphical form to allow one to evaluate the extent of intraparticle concentration and temperature gradients in polymer particles during olefin polymerization. Because we use observed reaction rates as a measure of catalyst activity, these criteria may underestimate microparticle diffusion limitations when there is significant resistance due to sorption equilibrium or macroparticle diffusion unless one uses good estimates of $M_{01}$ in Figures 5 and 6. Similarly, macroparticle diffusion resistance may be underestimated if there is a significant boundary layer mass transfer resistance unless good estimates of $M_S$ are used in Figures 9–12.

From the analysis presented, it may be concluded that under most conditions normally encountered in industry or the laboratory, intraparticle temperature gradients should be negligible for both the microparticles and the macroparticles in gas or slurry polymerization reactors. Exceptions would be for large highly active catalyst particles early in the lifetime of the polymer particles in gas phase reactors. On the other hand, intraparticle concentration gradients in the microparticles can be significant for high
activity catalyst systems having large primary crystallites of catalyst, especially in gas phase polymerization. By contrast, concentration gradients in the macroparticles will normally be negligible for gas phase polymerization, but are expected to be significant in many slurry systems, even for catalysts of relatively low activity. For gas phase reactors, concentration gradients in the macroparticle could be important for high activity catalysts of large size especially early in the life time of the polymer particle. As the polymer particle grows, this macroparticle diffusion resistance will be reduced, contrary to some suggestions in the literature which ascribe the catalyst rate decay to an increase in the resistance to monomer transfer. Diffusion control in these systems can be detected by an acceleration type rate behavior, or by an effect of catalyst particle size on the rate or yield.

The sequel will discuss the question of heat and mass transport resistances in the particle boundary layer. More general conclusions regarding the importance of heat and mass transfer resistances will also be presented in a companion paper.

The analysis of heat and mass transfer resistance in the microparticle has assumed a spherical microparticle (globular microstructure). For cases of other microparticle morphologies, this will serve as an approximation. Further detailed modelling of other structures would be worthwhile if the present analysis suggests that microscale diffusion limitations may be important.

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APPENDIX: NOMENCLATURE

\[
\begin{align*}
A_s & \quad \text{surface area of polymer particle} \\
c & \quad \text{dimensionless monomer concentration} = \frac{M_1}{M_2} \\
C_p & \quad \text{heat capacity of polymer} \\
C_w & \quad \text{heat capacity of fluid} \\
C_{ac} & \quad \text{concentration of active sites (mol sites/L cat)} \\
d & \quad \text{diameter of catalyst particle} \\
d_p & \quad \text{diameter of polymer particle} \\
D_a & \quad \text{diffusivity at zero penetrant concentration} \\
D_b & \quad \text{bulk diffusivity of monomer} \\
D_{de} & \quad \text{effective diffusivity in macroparticle} \\
D_{sm} & \quad \text{effective diffusivity in microparticle} \\
E & \quad \text{activation energy for propagation (cal/mol)} \\
\Delta H_p & \quad \text{heat of polymerization (cal/mol)} \\
h & \quad \text{external film heat transfer coefficient (cal/cm² s K)} \\
k & \quad \text{solubility constant (mol/L atm)} \\
k^* & \quad \text{solubility constant for purely amorphous polymer (mol/L atm)} \\
k_1 & \quad \text{thermal conductivity of polymer particle (cal/cm s K)} \\
k_2 & \quad \text{thermal conductivity of fluid (cal/cm s K)} \\
k_p & \quad \text{propagation rate constant (L/mol sites s)} \\
k_s & \quad \text{external film mass transfer coefficient (cm/s)} \\
M & \quad \text{monomer concentration in macroparticle} \\
M_s & \quad \text{bulk monomer concentration} \\
M_{ca} & \quad \text{monomer concentration at catalyst surface} \\
M_{sm} & \quad \text{monomer concentration at surface of macroparticle} \\
M_{mp} & \quad \text{monomer concentration in pores of macroparticle} \\
M_{mp} & \quad \text{monomer concentration at macroparticle surface} \\
\Delta M & \quad \text{concentration drop across external film (mol/L)} \\
\end{align*}
\]
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MW molecular weight of monomer
Nu Nusselt number = hdv/k,
ρ pressure (atm)
r microparticle radius
r1 macroparticle radius
rC catalyst primary crystallite radius
R gas constant = 1.987 cal/mol K
R1 radius of catalyst particle
Rm rate of polymerization of catalyst surface
Rk kinetic reaction rate
R1 radius of macroparticle
Robs observed polymerization rate (g/g cat h)
Rc radius of microparticle
Rv volumetric reaction rate in macroparticle
Re Reynolds number = ρdv/Ds
Sc Schmidt number = μ/ρDs
Sh Sherwood number = k,dv/Ds
T temperature in microparticle
Tb temperature in bulk fluid
Ts temperature at catalyst surface
T1 temperature in macroparticle
Ts temperature at macroparticle surface
ΔT temperature rise across external film
u particle-fluid relative velocity
un terminal velocity of particle
Vc volume of catalyst particle
z dimensionless radius = r1/R1

GREEK SYMBOLS

α1 dimensionless modulus = R1νk, k
α2 dimensionless modulus = r, √(k, C)/D, k
ρ porosity
γ dimensionless activity energy = E/RTs
φs microparticle growth factor = R, r
Φ macroparticle growth factor = R1/R
ρs density of slurry liquid
ρd density of slurry diluent
ρm density of monomer
ρp density of polymer
μs viscosity of slurry diluent
μm viscosity of monomer
μl viscosity of slurry liquid
ηs sorption equilibrium effectiveness factor
ηφ microparticle effectiveness factor
ηΦ macroparticle effectiveness factor
θ dimensionless temperature rise = γ (T, − Ts)/Ts
τ tortuosity factor
τm time constant for concentration equilibrium
τT time constant for temperature equilibrium

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Synthesis of
Aron

ALMERIA NATA
Weizman.

Novel aromatic polyphosphonates (methyl, phenyl) phosphonic acid and phase transfer catalysis. Mixtures of the polyphosphonate and the chloromethyl graft confer flame resistance to plastics.

Polycondensation of aromatic polyphosphonates for achieving melt, interfacial, and solid-state reactions. Of the phosphenyl (PPD) is the most wasterable stability by bringing aromatic Chloromethylphosphonate to melt polycondensation.

These polyphosphonates that are generally useful for performance plastics.

In the present study bisphenols: 4,4'-sulf phenoxyphenylpropane (tetra 2,7-dihydroxynaphthyl with PPD and methyl characterized.

The influence of the molecular weight is of the polymers obt the glass transition polyphosphonates a patibility and flammability.

* Present address: Mexico, PQ H3A 2K6, Car.
up to ~100 μm; however, it may not be valid for slurry polymerization concentration transients in ultralarge particles. In this case, neglecting these transients provides a conservative bound on intraparticle temperature gradients; thus we shall use the QSSA in the analysis here.

By invoking the QSSA, one may combine eqs. (1)-(4) and (13)-(16) in their steady state form to yield the relationship for the intraparticle temperature rise in the macroparticle,

\[ T_i(0) - T_s = \frac{(-\Delta H_p) D_i (M_s - M_i(0))}{k_s} \]

(30)

where \( T_s \) and \( M_s \) are the temperature and monomer concentrations at the surface of the macroparticle and \( T_i(0) \) and \( M_i(0) \) are the values at the center.

For the slurry polymerization of propylene and ethylene (using the range of parameters in Table I), one may obtain conservative estimates by assuming \( M_i(0) = 0 \) and \( M_s = M_x \), to estimate the maximum temperature rise in the macroparticle. This analysis predicts at most a 2-3 K temperature rise for propylene polymerization, and less than 1 K for ethylene polymerization. The actual values will normally be considerably less than this, since the center particle monomer concentration value will not be zero and there may be mass transfer resistance in the external boundary layer which causes \( M_x < M_s \). Detailed calculations indicate that for typical conditions in slurry even with high activity catalyst, the internal temperature rise is a fraction of a degree centigrade. Thus our analysis supports the assumption that macroparticle internal temperature gradients should be negligible for slurry polymerization.

The situation for gas-phase polymerization is much more complicated. First, for homopolymerization without the presence of an inert or chain transfer agent, mass transfer in the macropores would not be properly described by a diffusion process but would have a substantial convective transport contribution driven by a pressure gradient in the particle. However, this is an unusual case because most gas-phase reactors have inerts, chain transfer agents, and sometimes comonomer which allow macroscale counterdiffusion mass transfer to apply. To analyze this situation, we shall consider both cases.

First, we assume that convective mass transport in a one-component gas leads to a negligible intraparticle mass transfer resistance because the macropores are large (~1 μm) and the absolute pressure is large (15-30 atm) so that the intraparticle pressure drop required to overcome convective flow resistance is a small fraction of the total pressure. Thus as a conservative bound we assume that \( M_i = M_s \) everywhere in the macropores. In this case the macroscale reaction rate \( R_x \) is only a function of \( T_i \) and

\[ R_x = \eta_x \eta_{eq} k_p(T_i) C_x M_s (R_x) \eta / (R_i)^3 \]

(31)

where \( \eta_x \) accounts for microparticle diffusion limitations. \( \eta_{eq} \) represents the sorption equilibrium. This means that eq.

where

\[ z = \frac{1}{j} \]

\[ \delta = \frac{1}{j} \]

and the well-known exp

\[ \exp \left[ -\frac{E}{RT} \right] \]

has been made. Equatio

whose solution has been center temperature, \( \theta(0 \)

then for the parameter

macroparticle will be \( \Phi \) macro particle mass tran

\[ d_x = \]

Here \( \Phi_x = R_x / R_i \) is the reaction rate derived fr

7. Note that for low to internal temperature
the sorption equilibrium, and \( R_e \) is the radius of the original catalyst particle. This means that eq. (13) may be rewritten

\[
\frac{1}{z^2} \frac{\partial}{\partial z} \left( z^2 \frac{\partial \delta}{\partial z} \right) + \delta e^* = 0
\]

(32)

\[
\theta'(0) = 0
\]

\[
\theta(1) = 0
\]

where

\[
\cdot z = \frac{r_e}{R_i}, \quad \theta = \frac{E}{RT_s} \left( \frac{T_i - T_\delta}{T_\delta} \right),
\]

(33)

\[
\delta = \frac{(-\Delta H_p) n \eta \kappa_p R(T_3) C_m M_s (R_i)^2 E}{1000 k_c R(T_3)^2 \left( \frac{R_1}{R_2} \right)}
\]

and the well-known exponential approximation

\[
\exp \left[ -\frac{E}{RT_s} \left( \frac{T_3 - T_i}{T_i} \right) \right] \approx \exp \left[ -\frac{E}{RT_s} \left( \frac{t_3 - T_t}{T_t} \right) \right]
\]

(34)

has been made. Equation (32) is the classical Frank–Kamenetski equation whose solution has been tabulated in Ref. 41. In particular, the particle center temperature, \( \theta(0) \), is less than 0.1 for \( \delta < 1 \). Since

\[
T_i(0) - T_s = \frac{\theta(0) T_s}{(E/RT_s)}
\]

(35)

then for the parameters in Table I, the internal temperature rise in the macroparticle will be less than 2 K if \( \delta < 1 \) in the case where there is no macroparticle mass transfer limitation. Thus we may use the definition for \( \delta \) and set \( \delta = 1 \), to obtain the relationship between catalyst diameter, and \( R_{ab} \) as

\[
d_c = 2R_e = \left[ \frac{14400k_c(T_e)^2 M W \Phi_s}{(-\Delta H_p k_c E/R) \Phi_s, R_{ab}} \right]^{1/2}
\]

(36)

Here \( \Phi_s = R_c/R_i \) is the macroparticle growth factor and \( R_{ab} \) is an observed reaction rate derived from eq. (31). This is illustrated graphically in Figure 7. Note that for low to medium catalyst activities, there is no significant internal temperature rise if the catalyst particle is below 100 \( \mu \text{m} \) in di-
Fig. 7. Regimes for significant macroparticle temperature gradients for ethylene and propylene polymerization. Catalyst size \( d_c \) vs. observed rate for various macroparticle growth factors \( \Phi_p = R_p / R_1 \).

ameter. However, for high catalyst activities (> 1000 g/g cat h), the catalyst particle diameter must be smaller to insure no internal temperature rise. This is a happy situation because, as catalyst activities increase, less catalyst is required for the same size polymer particle, and one naturally reduces the catalyst particle size. For the most active catalysts in use today, catalyst particle diameters below 20 \( \mu \) m would insure no intraparticle temperature gradients. Note that, as the polymer particle grows in size (i.e., \( \Phi_p \) increases), intraparticle temperature gradients become less significant. In any case, the assumption of no mass transfer resistance is a very conservative bound. Thus one may conclude that, for one-component gas phase polymerization, there would be negligible internal temperature gradients in the macroparticle except for very large, high-activity catalyst particles.

Now let us consider the case where the presence of inert, transfer agent, or slowly reacting comonomer requires a diffusion description of mass transfer in the macroparticle. Carrying out the bounding analysis using eq. (30) for gas phase polymerization is not as conclusive as it was for slurry. For the parameters in Table I, gas phase diffusivities can be higher than for slurry, so that, using eq. (30), large macroparticle temperature gradients cannot be ruled out. Thus more detailed studies of the gas phase situation are required.

If we put the macroparticle equations (1)–(4) and (13)–(16) in dimensionless form we obtain for the material balance

\[
\frac{1}{2} \frac{\partial}{\partial z} \left( z^2 \frac{\partial c}{\partial z} \right) - \frac{\alpha c}{c \exp \left( \frac{\theta}{1 + \theta/\gamma} \right)} = 0, \quad 0 < z < 1
\]

\[\frac{dc(0)}{dz} = 0\]

\[c(1) = 1\]
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where eqs. (1) and (25) can be combined to yield

\[ \theta = \beta \gamma (1 - c) \]

Here we have defined the parameters

\[ z = \frac{r_1}{R_1}, \quad \alpha_i = R_1 \left[ \frac{\eta_i}{\eta} \frac{k_b(T_1)C_0}{D_i \Phi_i} \right]^{1/2}, \]

\[ \gamma = \frac{E}{RT_S}, \quad \beta = \left( \frac{(-\Delta H_p) D_i M_S}{k_c T_S} \right) \]

\[ c = \frac{M_i}{M_S}, \quad \theta = \left( \frac{T_1 - T_S}{T_S} \right) \gamma, \quad \Phi_\gamma = \frac{R_1}{R_c} \]

This is the classical nonisothermal reaction–diffusion problem whose solution has been tabulated many places (e.g., Refs. 41 and 42). The solution to this equation may be represented by an effectiveness factor \( \eta_i \) which applies to the macroparticle and is defined by

\[ \eta_i = \frac{\text{reaction rate with macroscale diffusion}}{\text{reaction rate in absence of macroscale diffusion}} = \frac{3(dz/dz)_{z=1}}{(\alpha_i)^2} \]  

(39)

For the parameters in Table I, \( \gamma < 20 \), and at most \( \beta = 0.2 \). The effectiveness factor plots for this range of parameters are shown in Figure 8. Note that, for \( \alpha_i < 1 \), there is no significant diffusion limitation or temperature rise in the pellet for the range of \( \beta, 0 < \beta < 0.2 \). Because the curves for \( \beta = 0, \beta = 0.02 \) are virtually superimposed, for \( \beta < 0.02 \), there is no significant temperature rise for any value of \( \alpha_i \). For the parameter ranges in Table I for gas phase polymerization, one sees that \( \beta < 0.02 \) for \( D_i < 10^{-4} \) cm²/sec. Figure 9 provides a quick check on the criteria \( \alpha_i < 1, \beta < \)

![Graph](image_url)

**Fig. 8** Nonisothermal macroparticle effectiveness factors for range of \( \beta, 0 < \beta < 0.2 \), corresponding to gas phase olefin polymerization.
temperature gradients in diameter catalyst particle will be expected to have ample for $D_i \Phi_x > 10^{-3}$. The significant internal temperature had grown by a factor If we consider gas phase catalyst of the Stauffer $\Phi_x < 10^{-4}$ for the onset of gas phase, we will have $r$ in the macroparticle for the diameter catalyst for the observed productivity, $I$ $D_i = 10^{-3}$, $D_i \Phi_x = 10$ $10^{-6} \leq D_i \leq 10^{-8}$ cm$^2$/s in the macroparticle; $h$ will be significant mass the particle size exceed.

Having shown that in the macroparticle unanalyze for macroparticle an isothermal particle, (37) has the solution

$$R_{ob} = \eta_i \eta_{m} R_{kin}$$

(40)

where $R_{kin}$ is defined by eq. (24) with $T_{ref} = T_s$, $M_{ref} = M_s$ in this case. Using the definition of $a_i$, we see that

$$a_i = \left( \frac{p_{ob} (R_i)^2}{3.6 M_s MW D_i \Phi_x \eta} \right)^{1/2}$$

(41)

so that lines corresponding to $a_i = 1$, $\eta_i = 1$ can be represented in simple terms. Also plotted is the line $D_i = 10^{-4}$ corresponding to the criterion $\beta < 0.02$ for which there are negligible temperature gradients. Note that it is only for extremely high catalyst activities and large catalyst particles that significant macroparticle temperature gradients could exist (indicated by curves in the shaded region of Figure 9).

To illustrate how to use Figure 9, suppose that one has a 20 $\mu$m diameter catalyst particle with $R_{ob} = 4000$ g/g cat h for ethylene gas phase polymerization when the ethylene pressure is 27 atm ($M_s = 1$ mol/L). If we assume $M_s = M_h$, then $R_{ob} / M_s MW = 143$. Thus this catalyst corresponds to point 1 in Figure 9 where $D_i \Phi_x = 10^{-4}$, $D_i = 10^{-4}$. Thus, for $D_i \Phi_x$ values $> 10^{-4}$, there is negligible heat and mass transfer resistance in the macroparticle. If we assume $D_i \sim 10^{-4}$ as a conservative estimate, then we conclude that there will be no significant internal concentration and
temperature gradients in the macroparticle. However, note that a 60 \( \mu \text{m} \) diameter catalyst particle with the same observed productivity (point 2) will be expected to have negligible internal heat and mass transfer resistance for \( D_i \Phi_s > 10^{-3} \). Thus if we assume \( D_i = 10^{-4} \), then one would expect significant internal temperature and concentration gradients until the polymer had grown by a factor of 10 (\( \Phi_s = 10 \)).

If we consider gas phase propylene polymerizations with a low activity catalyst of the Stauffer AA type with a catalyst diameter of 60 \( \mu \text{m} \), this is represented by point 3 in Figure 9 corresponding to \( D_i \Phi_s \), \( D_i \) values of \( \sim 5 \times 10^{-6} \) for the onset of diffusion resistance. Thus since \( D_i > 10^{-4} \) for gas phase, we will have negligible temperature and concentration gradients in the macroparticle for this catalyst in gas phase. If we use a 60 \( \mu \text{m} \) diameter catalyst for the slurry polymerization of propylene with the same observed productivity, \( R_{\text{obs}} \), this corresponds to point 4 in Figure 9 where \( D_i = 10^{-5}, D_i \Phi_s = 10^{-5} \) for the onset of diffusion resistance. For slurry \( 10^{-4} \leq D_i \leq 10^{-2} \text{ cm}^2/\text{s} \). Thus there will not be any temperature gradients in the macroparticle; however, with an actual value of \( D_i \sim 10^{-6} \), there will be significant mass transfer resistance due to diffusion—at least until the particle size exceeds \( \Phi_s > 10 \).

Having shown that intraparticle temperature gradients will be negligible in the macroparticle under most normal operating conditions, we may now analyze for macroparticle mass transfer limitations in more detail assuming an isothermal particle. For this case, the macroparticle material balance (37) has the solution

\[
c = \frac{\sinh(\alpha_i z)}{z \sinh(\alpha_i)}
\]

where \( z \) and \( \alpha_i \) are defined by Eq. (38). Thus, the macroparticle effectiveness factor, \( \eta_i \), defined by eq. (39), has the solution

\[
\eta_i = \frac{3}{\alpha_i} \left[ \frac{1}{\tanh(\alpha_i)} - \frac{1}{\alpha_i} \right]
\]  

(42)

From the definition of \( \alpha_i, R_{\text{obs}} \) in eqs. (40) and (41) one may represent eqs. (41) and (42) graphically as shown in Figures 10-12. Note that, for larger catalyst particles, one expects more serious diffusion limitations than for smaller catalyst particles at the same growth factor. However, for a fixed catalyst particle size, diffusion limitations are reduced with increasing growth factor \( \Phi_s \).

It is interesting to note from Figure 11 that for a low activity catalyst in propylene slurry (e.g., Stauffer AA type catalyst with average catalyst size of 60 \( \mu \text{m} \)), reasonable values of \( D_i \) (e.g., \( 10^{-6} - 10^{-5} \text{ cm}^2/\text{s} \)) are sufficient to cause macroparticle diffusion limitations. However, for the same productivity catalyst in gas phase (where \( D_i \approx 10^{-4} - 10^{-3} \text{ cm}^2/\text{s} \)), one would not expect to see macroparticle diffusion limitations. As indicated in Figure 10, where a catalyst particle size of \( d_p = 20 \mu \text{m} \) is used for ethylene polymerization with a high activity catalyst, one sees that the same conclusion is true for high activity catalysts; i.e., the possibility of significant intra-
particle concentration gradients for slurry polymerization, but not for gas phase polymerization. However, as indicated in Figure 12, for gas phase polymerization with large high activity catalysts, internal concentration gradients can be significant. Furthermore, if catalyst activity continues to improve and catalyst particle sizes increase dramatically, then internal gradients will become even more significant in gas phase processes.

In this paper, qualitative graphical form to all temperature gradient, the macroparticles would be for large h the polymer particle concentration gradients...