8th PhD-Workshop on Polymer Reaction Engineering

14th June to 16th June 2019 in Hamburg
Dear PhD-students, professors and company representatives,

the PhD-Student Workshop on Polymer Reaction Engineering represents an international platform where young researchers in the field of polymer reaction engineering have the opportunity to get into contact with fellow PhD-students but also to meet various industry representatives and potential employers personally. Moreover, PhD-students can benefit from useful feedback on their work and improve their soft skills by practicing the presentation of complex scientific topics.

The workshop is being held for the eighth time now. It has been launched in Lyon in 2012 with a total number of 35 participants. The second workshop has been organized in Hamburg. The third one took place in San Sebastian and in 2015 we met in Fürstenfeldbruck near Munich. The WPPRE student workshop in 2016 was hosted again by the University of Hamburg. Last year we recorded a number of 48 registrations in Prague. In this year we are expecting 47 participants from universities as well as industry.

Having participants from ten universities and various research fields we can look forward to an interesting and diverse workshop whose contributions include the following topics:

- process advancements as well as new methods of inline-spectroscopy
- synthesis of homopolymers, copolymers, blends and hybridpolymers
- bulk-, emulsion-, catalytic- and high-pressure polymerization
- modeling of chemical reaction networks: deterministic and stochastic
- focusing on structure-property relationships and thermodynamics

With all these topics ahead we are looking forward to an informative workshop in Hamburg with you.

The Organizing Committee

Organizing Committee
Markus Busch  Chair of WPPRE, TU Darmstadt
Werner Pauer  Secretary of WPPRE, University of Hamburg
Paul Peikert  Technical University Darmstadt
Elisabeth Schulz  Technical University Darmstadt
Sarah-Franziska Stahl  University of Hamburg
Sebastian Eller  University of Hamburg

Sponsors

Novolen Technology
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**General Information**

**Language**

The official language of the workshop is English.

**Insurance**

The organizers do not accept responsibility for individual medical or personal insurance. Participants are strongly advised to take out their own insurance policies.

**Telephone**

The international code for Germany is +49.

**Emergency phone numbers**

- General Emergency Number: 112
- Firebrigade: 112
- Police: 110
- Ambulance: 112

**Taxi phone numbers**

- (+49) 40 211 211
- (+49) 40 66 66 66
- (+49) 40 211 522

**Main Station**

Glockengießerwall / Kirchenallee, 20095 Hamburg
Arrival in Hamburg: Public Transport & Your Way to the Workshop

Transfer from the Airport to the Venue

There are several possibilities to get to the University of Hamburg. Taking public transport to the University takes about 50 minutes (see below for two possible options). Next to public transport, Taxis are available at the airport.

Contact

If you have any further questions, please feel free to contact us via email or visit our homepage.

wppre@pre.tu-darmstadt.de
http://efce.info/PhDStudent_Workshop_WPPRE.html

Workshop Venue Address

Institute of Technical and Macromolecular Chemistry
University of Hamburg
Bundesstraße 45
20146 Hamburg

Map with directions from Hamburg Airport to the University of Hamburg.
Places to Know

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**Conference Venue**
Adress: University of Hamburg, Bundesstraße 45, 20146 Hamburg

**Dinner on Friday at Hamborger Veermaster**
Adress: Reeperbahn 162, 20359 Hamburg

**Dinner Cruise on Elbe River on Saturday**
Adress: St. Pauli-Landungsbrücken 8-10, 20359 Hamburg

**Sunday Lunch at Blockbräu**
Adress: Bei den St. Pauli-Landungsbrücken 3, 20359 Hamburg
### Program by Days
**Friday, 14th June 2019**

<table>
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<th>Time</th>
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| 14:00 - 15:00 | Registration at the conference site  
Poster wall preparation |
| 15:00 - 15:15 | Welcome  
Prof. Dr. M. Busch |
| 15:15 - 15:35 | **Session I: Thermodynamics**  
Chair: Amin Nasresfahani (Queen’s University)  
Lenka Krajáková (University of Chemistry and Technology Prague)  
Sorption and swelling equilibria relevant for slurry ethylene polymerization |
| 15:35 - 15:55 | Maryam T. Agboluaje (Queen’s University)  
The effect of polar organic and aqueous solvent on the copolymerization kinetics of an acrylate and functional methacrylate system |
| 15:55 - 16:15 | Sebastian M. Dron (University of San Sebastian)  
Improvement of film formation abilities of waterborne latex particles for coating applications |
| 16:15 - 16:35 | Jana Sartorius (TU Darmstadt)  
Kinetic measurements by calorimetry and high-pressure: Decomposition of peroxides depending on solvent and pressure |
| 16:35 - 16:55 | Václav Pelc (University of Chemistry and Technology Prague)  
Mathematical Modelling of SAN Copolymerization including Thermal Initiation and Oligomer Formation |
| 16:55 - 18:00 | Coffee Break and Poster Session |
| 18:00 - 18:30 | Dr. Timo Melchin (Wacker) |
| 19:30 – 21:00 | St. Pauli Night Guard Tour |
| 21:00 – 23:00 | Dinner at Hamborger Veermaster |

Event kindly sponsored by: [Covestro](#)  
Dinner kindly sponsored by: [سابك](#)
**Session II: Modelling**

*Chair: Fabian Wenzel (University of the Basque Country)*

09:00 – 09:20  **Paul Peikert** (TU Darmstadt)
*Predicting the Branching Structure as well as the Radius of Gyration of LDPE-Homo- and Co-Polymers by Stochastic Modeling*

09:20 – 09:40  **Christian Zander** (University of Stuttgart)
*Propagation of terminal double bonds in the polymerization of N-Vinylpyrrolidone in aqueous solution: modelling strategies and comparison to experimental results*

09:40 – 10:00  **Felix Kandelhard** (University of Hamburg)
*Combined CFD- and Heat-Transfer-Modelling of an Adiabatic Storage Calorimeter*

10:00 – 10:20  **Francisco Arraez** (University of Ghent)
*Model-based study of 2-oxazolines CROP with special attention to side reactions*

10:20 - 11:00  **Coffee Break and Poster Session**

11:00 - 11:30  **Dr. Stefan Spiegel (Wiley)**

**Session III: Polyolefines**

*Chair: Sarah-Elisabeth Dechent (University of Hamburg)*

11:30 – 11:50  **Sabrine Kardous** (University Lyon 1 - LAGEP)
*Optimization of grade transition in fluidized-bed reactors accounting for condensed agent effects*
11:50 – 12:10  **Yashmin Blazzio** (University Lyon 1 - LAGEP)
*A novel stopped-flow reactor for gas-phase olefin polymerization*

12:10 - 12:30  **Jorik Hill** (Martin-Luther-University Halle-Wittenberg)
*Kinetic study of the gas phase polymerization of propylene with two Ziegler-Natta catalysts*

12:30 - 13:00  **Dr. Rolf Bachmann** (Covestro)

13:15  **Lunch: Catering served at conference venue**

14:30 – 14:50  **Stephanie Schlappa** (University of Potsdam)
*In-line monitoring of growth of nano-scaled particles in turbid emulsion polymerization using Photon Density Wave spectroscopy*

14:50 – 15:10  **Fabian Wenzel** (University of San Sebastian)
*Waterborne Nanoparticles by thiol-ene (Mini)Emulsion Polymerization*

15:10 – 15:30  **Laurence Jacob** (University of Hamburg)
*In-Line Monitoring of Latex-Particle size distribution during emulsion polymerizations up to 63 wt%*

15:30 – 16:30  **Coffee Break and Poster Session**
Program by Days
Saturday, 15th June 2019

16:30 – 17:00  Dr. David Eckes (Evonik)

Session V: Process Development

Chair: Lenka Krajáková (University of Chemistry and Technology Prague)

17:00 – 17:20  Isabel Kronshorst (TU Darmstadt)
Implementation of Hydrogen-Injection into a LLDPE-Mini-Plant

17:20 – 17:40  Amin Nasresfahani (Queens’s University)
Starved Intervals: Optimization of Semi-Batch Solution Radical Copolymerization via Comprehensive Stochastic Modeling and Derivative-Free Algorithms

17:40 – 18:00  Ikenna H. Ezenwajiaku (Queens’s University)
Influence of Ionic Strength from monomer and sodium chloride on aqueous-phase radical copolymerization kinetics of cationic monomers

19:45  Dinner Cruise at Elbe River

Dinner Cruise kindly sponsored by:

WACKER
Program by Days
Sunday, 16th June 2019

09:30  Miniaturwunderland

Event kindly sponsored by:

12:15  Lunch at Blockbräu

Lunch kindly sponsored by:

12:30  Sebastian Fries (Novolen)

13:00  Poster Prize and Closing Ceremony

The Prize for the best contribution is kindly sponsored by:

WILEY
List of Posters

1. Mathematical Modelling of SAN Copolymerization including Thermal Initiation and Oligomer Formation
   Václav Pelc, Juraj Kosek

2. Predicting the Branching Structure as well as the Radius of Gyration of LDPE-Homo- and Co-Polymers by Stochastic Modeling
   Paul Peikert, Markus Busch

3. Combined modelling and experimental study of triboelectric charging of polyethylene powders: wall-sheeting and agglomeration
   Simon Jantač, Jarmila Kučerová, Ladislav Konopka, Juraj Kosek

4. Experimental and modeling study of ethylene polymerization in gas phase reactors
   Amel Ben Mrad, Nida Sheibat-Othman, Timothy F. L. McKenna

5. Sorption and swelling equilibria relevant for slurry ethylene polymerization
   Lenka Krajáková; Barbora Tučková; Juraj Kosek

6. Estimation of Diffusion Coefficients for Multiple Penetrant/Polymer Systems Based on Sorption Data
   Rita Ferreira Alves, Timothy F.L. McKenna

7. Combined CFD- and Heat-Transfer-Modelling of an Adiabatic Storage Calorimeter
   F. Kandelhard, W. Pauer, H.-U. Moritz

8. Kinetic measurements by calorimetry and high-pressure: Decomposition of peroxides depending on solvent and pressure
   J. Sartorius, M. Busch

9. Power compensation calorimetry for studying the kinetics of bulk-phase polymerization of propylene
   Sina Valaei, M. Bartke

10. Entanglement Formation mechanism of Ethylene Polymerization by the Heterogeneous Ziegler-Natta Catalysts
    Yuming Chen, Wei Lib, Lei Huib, Binbo Jianga, Jingdai Wanga, Yongrong Yang

11. Optimization of grade transition in fluidized-bed reactors accounting for condensed agent effects
    Sabrine Kardous, Timothy Mckenna, Nida Sheibat-Othman

12. Kinetic study of the gas phase polymerization of propylene with two Ziegler-Natta catalysts
    Jorik Hill, Michael Bartke

13. The real-time detection and control of the atmosphere composition: Application in preparation of the PP/EPR in-reactor alloy
    Zheng Zheng, Jijiang Hu, Bo-geng Li

    Timothy F.L. McKennaa, Yashmin Blazzioa, Nida Othmanb, Sebastian Norsica
15. Ring-Opening Polymerization of Epoxides by DMC in Micro-reactor
   Jing Zhao, Hong Fan, Bo-geng Li

16. Implementation of Hydrogen-Injection into a LLDPE-Mini-Plant
   Isabel M. Kronshorst, Julia Eigenseer, Markus Busch

17. Waterborne Nanoparticles by thiol-ene (Mini)Emulsion Polymerization
   F. Wenzel, M. Aguirre, J. R. Leiza

18. In-line monitoring of growth of nano-scaled particles in turbid emulsion polymerization using Photon Density Wave spectroscopy
   Stephanie Schlappa, Roland Hass, Oliver Reich, Lena Bressel

19. In-Line Monitoring of Latex-Particle size distribution during emulsion polymerizations up to 63 wt%
   Laurence Jacob; Werner Pauer

20. Controlled Synthesis of Polyolefin Copolymers with Rapid Curability for Photovoltaic Cell Encapsulation
   Ligiong Luo, Kailun Zhang, Pingwei Liu, Bo-Geng Li, Wen-Jun Wang, Guangda Zhou

21. Influence of Particle Morphology on Film Drying Stresses
   Hesham Abdeldaim, José M. Asua

22. Fully bio-derivable CO2 fixating polymers for HNIPU applications
   S.-E. Dechent, A. W. Kleij, G. A Luinstra

23. High Barrier Waterborne Polysulfide Dispersions
   J. Elgoyhen, R. Tomovska.

24. Determination of micromixing efficiency in viscous media
   E. Arian, W. Pauer

25. Refractive Index Increments of EVA-Copolymers
   Elisabeth Schulz, Markus Busch
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Sorption and swelling equilibria relevant for slurry ethylene polymerization

Lenka Krajáková; Barbora Tučková; Juraj Kosek
University of Chemistry and Technology Prague, Department of Chemical Engineering, Technická 5, 166 28 Prague 6, Czech Republic

Depending on the polymerization process different PE grades are produced. Low density PE is produced by free radical polymerization, whereas high density PE and linear low density PE are produced by catalytic polymerization. Our research focuses on polyethylene produced by catalytic polymerization conducted in gas-dispersion, solution or liquid-dispersion reactors. Liquid-dispersion (suspension) polymerization is often called „slurry“ because of the mushy consistency of the reaction mixture, where a solid polymer grows on polymer/catalyst particles dispersed in the liquid diluent. Gaseous monomer thus needs to bubble, sorb and diffuse through the liquid diluent and the growing polymer to reach the active catalyst sites and react there. The knowledge of phase equilibria for the species present in the reactor is crucial for overall process design, for its optimization and innovation. These equilibria have significant effect on polymerization rate, the polymer structure formation, heat transfer inside/out of the reactor and also on the further polymer product processing.

Thermodynamic data relevant for “slurry” polymerization are of our interest in this contribution. We present sorption and swelling equilibria in polymer/diluent(1) systems, which were measured using new methodology developed in our laboratory [1]. The method is based on recording the mass decrease during desorption, where the equilibrium sorption is obtained by extrapolation by the initial slope method. We present results for various PE grades (from LLDPE to HDPE) and liquid hydrocarbon (C6 or C8). Solubilities of hydrocarbons in PE are measured at different temperatures.

Next, we present swelling equilibria of PE particles in various liquid diluents measured by video-microscopy in pressure cell and complement thus the solubility measurements. The highly demanded experimental data for slurry polymerization are sorption data for ternary system: PE/liquid diluent/gaseous monomer. Gravimetric measurement of co-solubilities is complicated, thus we use video-microscopic method to observe the co-swelling equilibria for ternary system. Preliminary results of co-swelling experiments for the system PE-hexane-ethylene are presented and discussed.

In the next part of our research the effect of swelling by liquid diluent on polyethylene microstructure was investigated by Atomic Force Microscopy and provided interesting findings.

The effect of polar organic and aqueous solvent on the copolymerization kinetics of an acrylate and functional methacrylate system

Maryam T. Agboluaje, Michelle Gelber and Robin A. Hutchinson*,
Department of Chemical Engineering, Queen’s University, Kingston, Canada;
maryam.agboluaje@queensu.ca

Monomers with hydroxyl functional groups are important components in both solvent-borne and waterborne copolymer systems. As well as introducing hydrophilicity to the copolymer, the reactive functionality can be used to form a high molecular weight crosslinked network upon application as for example, vehicular coatings. Previous studies have shown that polar solvents that are capable of forming or disrupting the hydrogen bonding of the functional monomer can cause a significant deviation in the copolymerization kinetics compared to what is observed in the bulk system. What remains poorly known, however, is how the copolymerization kinetics of functional/non-functional monomer pairings changes as the solvent changes from organic to aqueous.

In this work, NMR is used to analyze copolymer composition and pulsed laser polymerization coupled with size exclusion chromatography (PLP-SEC) is used to determine the rate coefficients that describe homo- and copolymerization chain growth, $k_p$ and $k_p^{cop}$, respectively. The influence of methanol, butanol, and methanol/water mixtures on the homopolymerization kinetics of 2-methoxymethyl acrylate (MEA) and 2-hydroxyethyl methacrylate (HEMA), and MEA/HEMA copolymerization kinetics is investigated by comparison to their bulk kinetic behavior.

It is found that hydrogen bonding systematically increases the value of $k_p$ for MEA as the monomer concentration in alcohol is decreased, with a greater increase in butanol than in methanol. The largest increase is seen in methanol/water mixtures, where $k_p$ increased by more than a factor of three compared to the bulk value. For MEA/HEMA copolymerization, the presence of methanol and butanol causes a similar increase in $k_p^{cop}$ compared to the bulk system for low HEMA content in the monomer mixture, but this effect diminishes as the HEMA content increases. A similar observation is made for MEA/HEMA systems in methanol/water mixtures, although at higher HEMA contents $k_p^{cop}$ remains higher than for the bulk case. Thus, the unique ability of water to enhance homopropagation rate coefficients is also seen in copolymerization.

In terms of copolymer composition, the presence of methanol, butanol and methanol/water mixtures at higher quantities leads to an increased incorporation of MEA into the copolymer compared to the bulk case. However, unlike what was observed for $k_p^{cop}$, the addition of water to the system does not affect copolymer composition: at the same quantity of solvent relative to monomer, the increase in MEA incorporation into the copolymer was similar, regardless of the solvent used. The terminal model represented the copolymer composition well, however, it does not provide a good representation of $k_p^{cop}$ as a function of comonomer fraction.

Session I
Improvement of film formation abilities of waterborne latex particles for coating applications

Sebastian M. Dron, M. Paulis
Polymat, University of the Basque Country San Sebastian, Spain

Waterborne latex dispersions accessible through emulsion polymerisation play an important role in the application field of coatings, especially for paints. However, even with benefits like the reduced volatile organic compounds (VOC) level, compared with solvent borne coating systems, only 16% of industrial paints are waterborne latex coatings. This could be owed to some technical challenges, which occur to reach properties such as hardness, block and print resistance. However, considering new environmental regulations the dramatic decrease of VOCs in coating products will be one of the most important goals during the development of new applications in this area. Even in waterborne emulsion polymers, the use of a certain amount of VOCs is necessary to obtain films with good properties (coming from a relative high glass transition temperature (T_g) polymer) at ambient temperature (with low minimum film formation temperature (MFFT)). Instead of VOCs temporary plasticizers can be used. In this project, Temporary Hydro Plasticizers (THPs) and Temporary Oligomer Plasticizers (TOP) are investigated to lower the MFFT without influencing the T_g. THPs use water to obtain the plasticizing effect, whereas TOPs use short chains, consisting out of the same monomers as the latex. As basic polymer a BA/MMA (40/60) latex with 50% solids content (T_g = 30°C, MFFT = 25°C) was used. As THP amongst others, polyethylene glycol methacrylate (PEGMA) and itaconic acid (ITA) were used. With these THPs the MFFT could be decreased down to 14 ± 2 °C, obtaining films with suitable mechanical and water resistance properties. A DSC measurement method was developed to study how much water causes the plasticization effect and its behaviour in the polymer matrix. For the TOPs, the oligomers were synthesized in situ during the emulsion polymerization using a chain transfer agent (CTA). The TOPs decreased the MFFT down to 17 ± 1 °C. The combination of both approaches allowed synthesizing a latex with a MFFT of 5 ± 1 °C and a T_g of 31 °C which reach his optimal mechanical performance independent from the temperature. Currently the produced latices are used, in cooperation with the industry, as binders for paint formulation to investigate their application performance.

Your notes:
Radical polymerization is widely applied in industrial production using peroxides as initiators. To guarantee a smooth start-up of polymerization a mixture of peroxides is used. Therefore, peroxides with complementary temperatures of decomposition are combined to stabilize the number of radicals during temperature changes. To calculate suitable peroxide concentration, kinetic parameters such as reaction rate and activation energy has to be known at reaction conditions in temperature and pressure dependence. Data sets are often not available or do not include the influence of solubility or pressure.

Polymers like LDPE are industrially synthesized under high pressure up to 3000 bar and temperature around 300°C. At these conditions, some classical techniques of kinetic measurement are difficult to perform, but high-pressure calorimetry can be carried out using massive and specialized devices. Calorimetry is sensitive to reaction progress and slight changes between different conditions because decomposition of peroxides is highly exothermic. This can be carried out in high-pressure crucibles in a DSC device withstanding pressure up to 800 bar or in a Transitometer withstanding pressure up to 2000 bar. Great advantage of calorimetric measurements in a Transitometer is the feasibility of extreme conditions and the simultaneous control of pressure, temperature and volume. Thus, the pressure is kept constant to prevent critical pressure build up and enables to determine pressure dependence.[1]

Kinetics of radical polymerization to LDPE are a highly complex reaction network, so that data needs to be measured partwise of single reactions getting reliable results. First step of the reaction network is the decomposition of peroxides creating radicals. Therefore, the decomposition of peroxides is measured without monomer surrounding. This way the overlap of decomposition reaction with propagation, termination or transfer is prevented.

The peroxide solution is heated up in tightly closed and pressurized crucibles with a certain heating rate while the calorimetical signal is measured. The scan starts below the onset-temperature and cover the complete decomposition creating an exothermal peak. Analysing this, the temperature of decomposition, the enthalpy, activation energy and pre-exponential factor is obtained. The enthalpy is the integral of the time-dependent calorimetric signal. Through Kissinger-method or Borchardt-Daniels-method as a n th order-method, the activation energy and the pre-exponential factor is calculated.

For the Kissinger-method, measurements with varying heating rates are necessary and an order of reaction of n=1 is assumed. With the Borchardt-Daniels-method, the reaction order is an additional variable and activation energy and pre-exponential factor are calculated through a multilinear fit out of a single measurement. Using wide ranges of heating rates for the Kissinger-method the reliability is increased and the assumed order of reaction n=1 can be verified with the n th order-method. Combining kinetic and enthalpy data, the heat input by a reaction is known.

Present results of DSC scans show a correlation of decomposition temperature with the polarity and heat capacity of the solvent. Besides solvent effects, kinetic parameters show a dependency on pressure. Solvent properties also influence the enthalpy of decomposition. The enthalpy increases with increasing polarity of the solvent and decreases with increasing pressure. Combining results of DSC and Transitometer, wide ranges of heating rates and pressure can be monitored and results of both devices can be compared.

Under high-pressure conditions of radical polymerization, peroxides are surrounded by a high viscose matrix of a polymer-monomer mixture. Replacing solvent by polymer melt or waxes for measurements could be a step forward to more practical reaction conditions.


Your notes:

Session I
Styrene-acrylonitrile (SAN) copolymer can be found in many applications: from plastic bottles, through kitchenware to reflectors in cars, lamps etc. It is of great importance to ensure the highest possible quality of the produced copolymer. Hence various mathematical models were developed to describe the SAN copolymerization process and some of these can be found in the literature, e.g. [1]. One of the drawbacks of available models is usually omission of implementation of oligomer formation. The presence of oligomers in the final product can negatively affect properties of the product. For the polymer producer, it is thus advantageous to have a mathematical model enabling prediction of the final product quality (including content of oligomers) based on the defined reaction and down-stream processing conditions.

A new process model incorporating the detailed description of thermal initiation based on experimental studies by Buchholz and Kirchner [2, 3] was developed. This model includes a new detailed initiation scheme, which can lead to oligomer formation, and which considers both thermal and initiator-based formation of free radicals. The model is implemented in MATLAB and, using this implementation, the unknown values of kinetic parameters are estimated from available experimental data including: (i) conversion of comonomers, (ii) number and weight average molecular weights of copolymer, (iii) copolymer composition, (iv) reactor conditions, and (v) oligomer content. Experimental data were taken either from open literature, e.g. [4], or provided by the industrial partner. The model with estimated parameters was subsequently implemented in the process simulator Aspen Plus and validated against additional sets of experimental data.


Predicting the Branching Structure as well as the Radius of Gyration of LDPE-Homo- and Co-Polymers by Stochastic Modeling

Paul Peikert, Markus Busch*, TU Darmstadt, Darmstadt, Germany;

*markus.busch@pre.tu-darmstadt.de

With a production of more than 90 million tons per year Low-Density Polyethylene (LDPE) and its co-polymers represent one of the most important plastic materials. The relevance of these polymers produced at high-pressure up to 3000 bar and high temperatures up to 300 °C can be traced back to their unique product properties resulting from their branching structure. While occurring short-chain branches (SCB) mainly affect crystallinity and density, long-chain branches (LCB) are crucial for the viscoelastic behavior and processability of the product. Therefore, depending on the microstructure, LDPE can be employed as packaging material for food or pharmaceuticals. The area of application can even be extended by producing LDPE-co-polymers. Consequently, EVAs (polyethylene-vinyl acetate) can be applied as encapsulating material in the photovoltaic industry for example.

Due to its extraordinary process conditions, simulation-aided product design is frequently used in order to obtain a product tailor-made for its application. Therefore, in this contribution a coupled deterministic and stochastic hybrid approach has been developed in order to model the branching structure of LDPE-homo and co-polymers. The stored exact position of SCBs and LCBs of an ensemble of macromolecules can afterwards for example be employed to calculate the radius of gyration in dependence on the chain length. Thereby, a newly developed and validated self-avoiding random walker enables the prediction of the macromolecular dimension in dependence on the operation at the plant. In order to evaluate the influence of polar co-monomers in LDPE-co-polymers on the radius of gyration, several EVA- and EMA-co-polymers (polyethylene-methyl acrylate) with different branching structures and co-monomer-content in the polymer have been produced in a mini-plant autoclave. On basis of these co-polymer samples, the impact of polar co-monomers on the radius of gyration can be determined in comparison to LDPE-homo-polymers.

Thus, with this contribution a more detailed view on the structure-property relationship for the high-pressure ethene (co)-polymerization is demonstrated.

Session II
Propagation of terminal double bonds in the polymerization of N-Vinylpyrrolidone in aqueous solution: modelling strategies and comparison to experimental results

C. Zander¹, K.-D. Hungenberg⁵, T. Schall¹, C. Schwede¹, U. Nieken²
¹Universität Stuttgart, Institut für Chemische Verfahrenstechnik, Stuttgart, Germany; ²Hungenberg Consultant, Birkenau, Germany; ³BASF SE, Ludwigshafen, Germany

Specialty polymers, e.g. Polyvinylpyrrolidone, are mostly produced in batch or semi-batch processes due to the need for flexibility in the production of this product class. In the course of process intensification, new processes in smart-scale reactor systems which can be operated continuously are under investigation to improve process control and energy efficiency. One of the major problems in the development of processes in such reactor systems is the occurrence of fouling deposits which prohibit the continuous operation. The latter may be formed by side reactions leading to branched or crosslinked polymer chains which form an insoluble polymer gel. To predict process windows in which reactors can be operated without the formation of fouling deposits, kinetic models which predict polymer properties quantitatively are essential. Since these models are to be used in extensive parameter variations or CFD simulations, the computational effort should be as small as possible while the relevant structural property information must be captured as accurately as necessary.

The propagation and termination kinetics of the system of interest are well known and published in literature [1,2]. While it is clear that side reactions are relevant in the reaction mechanism [3], their nature and especially their rates are still in the focus of research. Recent results suggest that production of and propagation of terminal double bonds are important parts of the branching mechanism [4]. The work presented in this contribution is based on these findings. Starting from a multi-dimensional property distribution, different reduced models have been derived using so-called pseudo-distributions [5,6] and validated. Further reduction of model complexity has been achieved by transferring information from more detailed models into simplified models which are suitable for fast calculations.

Different models with different levels of complexity and detail of information will be introduced and compared. The simplified models have been used for parameter estimations and will finally be compared to CSTR experiments. The model predictions and experimentally observed values are in very good agreement.

The work has partially been funded by the German Federal Ministry for Economic Affairs and Energy as part of the ENPRO Initiative.

Combined CFD- and Heat-Transfer-Modelling of an Adiabatic Storage Calorimeter

F. Kandelhard, W. Pauer, H.-U. Moritz
University of Hamburg, Institute of Technical and Macromolecular Chemistry, Bundestraße 45
20146 Hamburg

The adiabatic storage test (AST) is an adiabatic calorimetric method that can be used to gather safety relevant data of chemical processes. This data include activation energies of the main and side reaction, pressure behavior or decomposition temperatures. In this study a new high-pressure-AST-calorimeter was modelled with an combined CFD (computational fluid dynamic) and heat transfer approach in COMSOL Multiphysics® in order to understand and subsequently improve the heat transfer properties and heat transport paths in the actual calorimeter. The model is capable to simulate the heat up process as well as simple calorimetric experiments with small deviation compared to the experimental data. It also enables an analysis of the temperature- and flow-fields inside the calorimeter during such processes, which can open new ways to improve the device.

Your notes:
Model-based study of 2-oxazolines CROP with special attention to side reactions

Francisco J. Arraez, Paul H. M. Van Steenberge, Xiaowen Xu, Valentin-Victor Jerca, Richard Hoogenboom, Dagmar R. D’hooge, Ghent University, Belgium

Poly(2-oxazolines) (PAOx) are an interesting bioinspired class of polymers whose biocompatibility allows for drug, protein and gene delivery applications. PAOx are available through cationic ring-opening polymerization (CROP) of 2-oxazolines representing an easy and key strategy for the synthesis of well-defined polymers with controlled average polymer composition, narrow size exclusion chromatography (SEC) trace and suitable end-group functionalities. Due to the living nature of CROP and by the incorporation of the correct 2-oxazoline comonomer, a wide variety of linear as well as branched/network (co)polymers can be synthesized with well-tailored structures and less abrupt transitions from one comonomer type to the other. Even so, a key challenge to be dealt with consists of evaluating the PAOx synthesis success at the molecular level, hence, beyond experimentally accessible average CROP characteristics.

In this contribution [1-3], a combination of an advanced kinetic Monte Carlo modeling technique with meticulous experimental analysis is covered, allowing the kinetic analysis of CROP of 2-oxazolines, with specific focus on functionality design per chain length and the effect of side reactions such as chain transfer to monomer (β-elimination) and macropropagation. A novel parameter tuning is introduced with for the first time reliable macropropagation rate coefficients based on complete SEC data. Model-based design is shown to be an effective strategy to identify optimal synthesis conditions that maximize the functionality efficiency for both low and high targeted chain lengths.


Session II
Polyethylene (PE) is the most widely produced polymer in the world. The variety of PE applications requires different grade specifications to suit the market demand. Most processes used to make linear low density polyethylene (LLDPE) are gas phase processes to avoid problems linked to the solubility of amorphous polymers in liquid hydrocarbons. These processes are clean, solvent-free and less energy consuming than free radical processes (lower temperature and pressure), but their productivity is limited by the heat generated by the reaction. Fluidized-bed reactors (FBRs) are the only reactors that can be used, from an economic point of view, to make many grades of LLDPE or HDPE by changing the operating conditions (e.g. Pressure, Temperature...). [1]

In order to overcome the issue of heat transfer in gas processes, “condensed mode cooling” is frequently employed, where induced condensing agents (ICAs), such as isobutane, are injected in either liquid or vapor form in order to absorb part of the reaction heat. [2] However, it was observed that the presence of ICA influences the absorption of monomer into the polymer and consequently the reaction rate and the polymer properties (e.g. the polymer molecular weight and density).

In this work, a combined kinetic and thermodynamic model (based on Sanchez-Lacombe equation of state) is developed for the catalytic copolymerization of ethylene with 1-butene in an FBR. This model is able to predict the co-solubility effect of the ICA (isobutene) and thus its effect on the reaction rate and the polymer properties. However, the Sanchez-Lacombe model is not available for a quaternary system (here PE, ethylene, ICA and co-monomer) and no thermodynamic data is available in the literature for this system. Therefore, the co-solubility effect of ICA (isobutene) on ethylene was evaluated in a ternary system (PE, ethylene and ICA) and was assumed to remain unchanged when adding the co-monomer (i.e. assuming no interaction between ICA and the co-monomer). Moreover, the solubility data of the co-monomer (1-butene) in the polymer is available only in a binary system, so it was approximated to obey the same trend as the ICA in a ternary system (as they have similar free volumes and thermodynamic properties). These two assumptions allow us to estimate the concentrations of ethylene and co-monomer in the amorphous polymer, and therefore to calculate the copolymerization rate and the copolymer properties.

This model was then used to optimize the transition between different grades of LLDPE using a dynamic optimization procedure, which provides the optimal flow rate profiles of hydrogen and co-monomer. The objective function allows for minimization of the quantity of off-spec during grade transitions, where the primary controlled variables are the melt index (MI), which is related to the polymer molecular weight (that is determined by transfer to hydrogen and the co-monomer fraction in the copolymer), and the polymer density (ρ) which is controlled by the co-monomer fraction in the copolymer. This strategy demonstrates the importance of accounting for the effect of the induced condensing agent within grade transition optimization, as it has a direct effect on the polymer properties and the reaction rate.

The proposed model is expected to be a powerful tool for fundamental understanding of gas-phase polyolefin polymerization, and can be used for improving PE product quality in the polymer industry, with minimized waste and energy consumption during the transition while maintaining the process safety.


Your notes:

Session III
Given the industrial relevance of polyolefin production, it is of great interest to understand the phenomena taking place during the nascent phase of the polymerization. It is well known that the first few moments of the polymerization can be crucial in ensuring satisfactory properties of the final polymer (morphology formation), as well obtaining stable reaction conditions (risk of particle overheating)\(^1\). Given the promising results obtained with simple fixed bed stopped flow reactors\(^2,3\), a novel stopped-flow reactor has been developed for studying nascent phase in gas-phase olefin polymerizations over a broader range of conditions. The reactor was conceived with a fritted annular configuration to improve (in relation to previous versions) heat removal with a more homogeneous distribution of the gas flow.

Reactor prototypes were conceived and tested in our facilities, followed by the production of a professionally engineered version of the set-up, developed with ILS (Germany). Studies performed using the prototypes with supported metalloocene catalysts indicate the set-up allows one to follow the development of polymer thermal properties, molecular weights and particle morphology at short reaction times with satisfactory heat transfer control in the reactor. Furthermore, preliminary results from the mathematical modelling of the set-up indicate minimal heat accumulation in the reactor.


Kinetic study of the gas phase polymerization of propylene with two Ziegler-Natta catalysts

Jorik Hill, Michael Bartke
Martin-Luther-University Halle-Wittenberg, Institute of Chemistry, Polymer Reaction Engineering Group, Heinrich-Damerow-Str. 4, 06120 Halle (Saale), Germany

Polypropylene (PP) is a versatile polymeric material, which is used for many high-volume applications such as consumer goods, packaging, automotive parts and medical products. With a global market share of about 20%, PP is one of the most commercially important polymers. Industrially, heterogeneous Ziegler-Natta (ZN) catalysts are predominantly used for the polymerization of propylene. The catalyst greatly defines the physical properties of the final product as well as the productivity of the process. Since the development of ZN catalysts in the early 1950s, continuous improvements were performed with regard to higher catalyst activity and better product performance.

During the coordinative polymerization of propylene, the polymer chains grow at the active sites of the catalyst. The active center determines the rate of monomer incorporation and thus the polymerization kinetics are catalyst dependent. The ongoing industrial catalyst development requires kinetic evaluation of newly designed catalysts. One important aspect is the lab-scale kinetic investigation of these catalysts under industrial relevant conditions. The gathered results are essential for the next scale-up step. Combining the experimental output with modeling and simulation enhances process development. For instance, lab-scale kinetics can be combined with pilot-scale process dynamics to precisely plan pilot-scale tests.

The current contribution deals with the kinetic investigation of two commercial Ziegler-Natta catalysts. Polymerization experiments were conducted under industrial relevant conditions in a 5 L stirred tank gas phase reactor. The reactor was operated in semi-batch mode enabling to monitor the reaction rate under constant pressure and temperature. The effects of hydrogen and a prepolymerization step were studied. Based on the experimental results, a kinetic model was developed. The model could adequately describe the experimental findings within the measurement errors.

Your notes:

Session III
In-line monitoring of growth of nano-scaled particles in turbid emulsion polymerization using Photon Density Wave spectroscopy

Stephanie Schlappa, Roland Hass, Oliver Reich, Lena Bressel
University of Potsdam – innoFSPEC, Institute of Chemistry, Physical Chemistry, Am Muehlenberg 3, 14476 Potsdam, Germany

Highly concentrated liquid dispersions with particles and droplets in the nanometer and micrometer scale are of wide technical importance. Examples are found in polymer and coatings industry, in food industry and in biotechnology. Such emulsions or suspensions often exhibit strong light scattering, thus limiting conventional optical process analytical technologies in their application range e.g. for particle sizing or concentration determination.

A contamination- and calibration-free process analytical technology suitable for highest concentrations (i.e. > 40 vol%), stirred or flowing systems and for potentially explosive environments is found in in-line or on-line Photon Density Wave (PDW) spectroscopy [1-2]. It allows for the independent quantification of the absorption and scattering properties of disperse materials, i.e. the absorption and reduced scattering coefficients, respectively. The absorption coefficient exhibits information about the chemical composition and/or the temperature of the material. The reduced scattering coefficient however is related to the size of the dispersed particles. Sizes are determined applying Mie theory for spherical particles and theory of dependent light scattering [3].

As a chemical model process the radical emulsion polymerization of vinyl esters stabilized by vinylalcohols (protective colloid) was investigated by PDW spectroscopy. Vinylesters were polymerized to have specifically set particle sizes and particle size distributions by changing the type of redox initiator, amount of initial protective colloid and addition rates of educts.

With a time resolution in the minute scale the growth of formed nanoparticles could be monitored during emulsion polymerization by PDW spectroscopy. Highly turbid polyvinylester dispersions with mean particle sizes in the nanometer or lower micrometer range and with solid contents of approx. 50% were achieved. Particle size distributions obtained without dilution in the final product by PDW spectroscopy were in good agreement with conventional off-line and dilution based analysis techniques as Scanning Electron Microscopy, Static Light Scattering and Dynamic Light Scattering.


Session IV
Waterborne Nanoparticles by thiol-ene (Mini)Emulsion Polymerization

F. Wenzel, M. Aguirre, J. R. Leiza
POLYMAT and Kimika Aplikatua Saila, Kimika Fakultatea, University of the Basque Country
UPV/EHU, Joxe Mari Korta Zentroa, Tolosa Hiribidea 72, 20018 Donostia-San Sebastián, Spain

Thiol-ene polymerization takes place between a thiol functional group and a diene functional group. Despite being a radical polymerization, it follows step-growth mechanism. Therefore, generally stoichiometric amounts of the monomers are used. Thiol-ene polymerization exhibits many advantages, such as the extremely rapid reactions due to the insensitiveness of the reactants to oxygen and moisture, which are near to quantum yields. Thiol-ene polymerizations show properties of so called "click" reaction. Furthermore, a broad range of thiols and dienes can be used. Due to this versatility and robustness thiol-ene polymerization is feasible for a wide range of applications such as protective coatings, optical and biomedical devices, coupling and bioorganic modifications.\(^1\)

On the other hand, waterborne polymerizations are gaining more and more attention over solvent borne systems due to increasing restrictions regarding volatile organic compounds (VOC). However, because of the rather hydrophobic nature of the dienes it is challenging to introduce thiol-ene polymerization into a conventional emulsion polymerization process, whereas the possibility to do so by using a less hydrophobic diene was reported by D. Shipp et al.\(^2\), in most cases a miniemulsion polymerization process is necessary.

The aim within this project is to investigate the suitability of thiol-ene polymerization for the production of waterborne pressure sensitive adhesives (PSAs). PSAs are viscoelastic materials that can adhere strongly to solid surfaces upon application of a light contact pressure and in short contact times.\(^3\) Mobility of the polymers chains is needed for the PSA to be able to adhere to the substrate and to provide tack. But also a certain degree of stability to provide cohesion and resistance against creep is needed. Polymers coming from thiol-ene polymerization exhibit very low \(T_g\)'s because of their linear nature. Therefore, different kinds of thiol and diene monomers were investigated in this work, either in emulsion or miniemulsion polymerization, to tune the \(T_g\) of the polymer. Furthermore, monomers with a functionality higher than two were used to introduce crosslinking in order to provide increased cohesive strength to the waterborne PSA.


In-Line Monitoring of Latex-Particle size distribution during emulsion polymerizations up to 63 wt%

Laurence Jacob, Werner Pauer
University of Hamburg, Institute for Technical and Macromolecular Chemistry, Bundesstraße 45
20146 Hamburg

The most common methods for the determination of mean particle size and particle size distribution mainly perform offline analysis in diluted solutions, which poses numerous disadvantages. Therefore, an inline measurement method would be preferable. Thus, the relatively new method of photon density wave (PDW) spectroscopy [1] was investigated for its suitability for inline particle size measurement in high solid content emulsion polymerization.

For this reason, emulsion copolymerization of vinyl acetate and Versa® 10 in a molar ratio of 9:1, and with varying emulsifier content, were carried out with different target particle sizes, both in batch and semi-batch, achieving a size range of 50 to 325 nm. For the initiation of the copolymerization, a redox initiator system, consisting of L-ascorbic acid, tert-butyl hydroperoxide and ammonium iron (III) sulfate, was used. The results of PDW spectroscopy were compared with those of conventional offline measurement methods, such as dynamic light scattering (DLS) and sedimentation analysis, by means of a disc centrifuge.

In order to attest the suitability of the PDW method, the measured particle diameters were plotted against each other. A linear fit of the results comparing PDW spectroscopy with a disc centrifuge gave a slope of (0.94 ± 0.06) and a y-intercept value of (7 ± 12) nm. The fit has a correlation quality of 98% and a corrected R² of 95%. This corresponds to a very good agreement of the data. The linear fit of the comparison between PDW spectroscopy and DLS gave a slope of (0.99 ± 0.14) and a y-intercept value of (-47 ± 24) nm. The correlation quality of the fit is 92% and the corrected R² is 84%. The suitability of PDW-spectroscopy was proven experimentally with a polymer content of up to 63 wt%.

Implementation of Hydrogen-Injection into a LLDPE-Mini-Plant

Isabel M. Kronshorst, Julia Eigenseer, Markus Busch
TU Darmstadt, Ernst-Berl-Institut für Technische und Makromolekulare Chemie, Alarich-Weiss-Str. 8, D-64287 Darmstadt, Germany,
e-mail: markus.busch@pre.tu-darmstadt.de

As plastics play a fundamental role in modern society, controlling its properties is of major importance. Linear low-density polyethylene (LLDPE) is a co-polymer of ethylene and an α-olefin. It is prepared in a catalytic solution polymerization and offers a variety of properties depending on molecular weight, used co-monomer and amount of implemented co-monomer. During the polymerization process, changing one of these parameters influences the others. As the aspects interfere with each other mutually, a reduction of the system’s complexity is necessary in order to regulate a parameter separately. The important aspects that influence the properties of LLDPE are the amount of co-monomer, the type of co-monomer, the conversion and the degree of polymerization. In order to compare different side chain lengths different co-monomers must be accessible. Therefore, an existing LLDPE solution mini-plant which has a pre-existing liquid monomer dosing is expanded by a liquefied gas dosing.

For the analysis of the impact of the short-chain branching length on the polymer properties, the molecular weights of the polymers have to be similar. Therefore, a chain transfer dosing is introduced using hydrogen. By this the degree of polymerization can be adjusted to a certain value. By oppressing the influence of degree of polymerization on polymer properties, the influence of the side chain lengths can be estimated separately.

The implementation of both adjustments is separately validated on the correct operation. Figure 1 shows the validation of the transport of the liquefied gas. It depicts the ratio between the actual feed rate and the pre-set feed rate as a measure of the pump head’s temperature. The deviation which is caused by cavitation can be reduced and a proper transport of the liquefied gas is validated.

Additional to the verification of a correct operation, for the hydrogen dosage a proper method of feeding has to be chosen. The impact of hydrogen concentration transients is shown in figure 2. As part of this work a method of feeding hydrogen to the system without transients was developed.

Figure 1: Deviation of feed rate from the pre-set feed rate as a measure of temperature.

With both of these adjustments, the mini-plant can now be used to prepare a variety of co-polymers differing in amount and length of side branching at controlled molecular weights.

Acknowledgement: Borealis AG

Your notes:
Starved Intervals: Optimization of Semi-Batch Solution Radical Copolymerization via Comprehensive Stochastic Modeling and Derivative-Free Algorithms

Amin Nasresfahani, Robin A. Hutchinson
Department of Chemical Engineering, Queen’s University, Kingston, ON K7L 3N6, Canada

Theoretical models are vital in bridging kinetic and process procedures to the resulting structures and to tackle the design complexities in synthesizing engineered materials. Discretizing the polymerization time into multiple constant starved-feed intervals allows kinetic Monte Carlo simulations to be utilized in semi-batch recipe optimizations where manipulating the independent feed streams are regarded as a gateway of improving the process. This concept is coupled with an accelerated stochastic model [1] developed for semi-batch solution free radical copolymerization of acrylate/methacrylates synthesized at high-temperatures; Considering secondary reactions such as backbiting, macromonmer production/consumption, and midchain radicals provides a realistic off-line optimization. It uses the reaction time intervals to regulate flowrates of the feed streams. Derivative-free optimization algorithms (e.g., Pattern Search[2] and Particle Swarm[3]) are utilized to effectively derive the polymer product properties to the target value. The developed concept of starved-intervals is found to be a promising strategy compared to the traditional starved-semi-batch recipes often used for obtaining low-molecular-mass high-value polymer resins, reducing the total polymerization time by 75% while simultaneously improving the quality of the product and the productivity of the process.

Figure 1: A Graphical abstract depicting the concept of the technique compared to the traditional starved-feed operation.


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INFLUENCE OF IONIC STRENGTH FROM MONOMER AND SODIUM CHLORIDE ON AQUEOUS-PHASE RADICAL COPOLYMERIZATION KINETICS OF CATIONIC MONOMERS

Ikenna H. Ezenwajiaku, Robin A. Hutchinson*
Department of Chemical Engineering, Queen’s University, Kingston, ON, K7L 3N6, Canada; ikenna.ezenwajiaku@queensu.ca

Aqueous-phase radical polymerization kinetics of industrially relevant water-soluble polymers have been critically investigated using the in-situ NMR technique. This specialized experimental technique has greatly improved the understanding of these systems and allows for reliable data collection at monomer concentrations up to 40 wt%, following both overall monomer conversions as well as composition drift in a copolymerization system.

Our work focuses on the effects of varying ionic strength contributions from monomer contents and sodium chloride (NaCl) on kinetics of [2-(methacryloyloxy)ethyl]trimethylammonium chloride (TMAEMC) and 3-(methacryloylamino)propyl]trimethylammonium chloride (MAPTAC), cationic monomers containing an ester and amide functional groups, respectively, copolymerized with both non-ionized (AA) and fully-ionized (NaA) acrylic acid to form copolymers with applications as cationic flocculating agents in water treatment. The drift in monomer composition was found to be enhanced with increasing ionic strength of monomer and NaCl. However, as molar fractions of TMAEMC and MAPTAC decrease in solution, this effect diminished with more pronounced effects in copolymerization with NaA (as shown in Fig. 1 for AA-TMAEMC). This could be attributed to NaA cross propagation being more favoured than AA due to electrostatic interactions while homopropagation of TMAEMC remains relatively unchanged.

A model to capture the composition drifts of AA/NaA-TMAEMC copolymerization has been successfully completed which accurately predicts the effects of ionic strength from monomer contents and NaCl as well.

Your notes:

Figure 1: Comonomer composition drifts at varied monomer contents in batch copolymerizations of TMAEMC with non-ionized AA (left) and fully ionized AA (right) at 50°C and 0.4 wt. % V-50, and different initial monomer compositions.
Combined modelling and experimental study of triboelectric charging of polyethylene powders: wall-sheeting and agglomeration

Simon Jantač, Jarmila Kučerová, Ladislav Konopka, Juraj Kosek*
University of Chemistry and Technology Prague, Department of Chemical Engineering, Technická 5, 166 28 Prague 6, Czech Republic, e-mail: Juraj.Kosek@vscht.cz

In the processing and transportation of insulator powders like polyethylene, collisions occur among powder particles and between powder particles and device walls. During such collisions, charge carriers are redistributed asymmetrically among the particles, which results in charging. Charged particles can then agglomerate. Apart from agglomeration, fouling can play a role in industrial systems, as PE particles of any charge polarity are attracted towards the metal walls of reactors and conveyors due to the effect of electrostatic induction. Therefore, sheeting does not strictly depend on the polarity of charge but rather on the charge magnitude. The mechanism of electrostatic charging is still not fully understood and therefore we systematically investigate what parameters affect the charging and how it can be explained. Besides experimental studies of charging dynamics, we focus on the modelling of particle charging and its comparison with measured charge produced by particle-particle and particle-wall collisions.

The central part of our research is the free-fall electrostatic separator that allows us to acquire valuable information about the charge distribution in powders (i.e., weight fraction of particles vs. particle charge). We managed to measure separately particle-particle and particle-wall charging, as well as the combined charging of both types as it typically happens in industrial processes. We observed that solid volume fraction, particle size distribution and the type of contact significantly affect the bipolar charge distribution in powders. Such bipolarity in charging due to particle-particle collisions, as well as the formation of agglomeration centres was also predicted and quantified using our particulate mathematical model based on discrete element method. The charge is not the only parameter affecting the dynamics of agglomeration or wall-sheeting; the amount of dissipated energy (restitution coefficient) during contact is equally important to the particle charge. To further increase the accuracy of our model we measured the restitution coefficient as a function of impact velocity, particle size, shape and temperature, and subsequently we developed the mathematical model enabling us to predict restitution coefficients of colliding particles of various shapes and sizes. Our work demonstrates the influence of many parameters on the generated charge during particle-particle and particle-wall collisions, as well as the effect of dissipated energy on the agglomeration and wall-sheeting.

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EXPERIMENTAL AND MODELING STUDY OF ETHYLENE POLYMERIZATION IN GAS PHASE REACTORS

Amel Ben Mrad, Nida Sheibat-Othman, Timothy F. L. McKenna
C2P2 (LCPP), UCL/CNRS-CPE, UMR 5265, Villeurbanne, France

A widely-used approach to control overheating in the gas phase polyethylene systems is the so-called “condensed operating mode” where liquid species are injected together with the monomer feed [1]. These liquid species, usually alkanes, are called “induced condensing agents” (ICA). Upon entering the reactor, the liquefied components vaporize and the latent heat of evaporation helps to cool the system. It has recently been demonstrated that the inert species most typically used for this purpose can strongly influence the solubility of all species in the growing polymer particles [2]. Different thermodynamic models are available that can capture this type of behavior, but all of them rely on a set of adjustable parameters that cannot be predicted a priori. To add to the complications, very limited solubility data is available for multicomponent systems; it is therefore very difficult to obtain realistic model parameters for olefin polymerization systems.

We have chosen to work with the Sanchez-Lacombe equation of state, as it is one of the most widely applied thermodynamic models in polymer industry [3]. The interaction parameters used in the Sanchez-Lacombe EoS will be identified by fitting equilibrium solubility data which are measured experimentally using a pressure-decay technique in standard laboratory equipment. Experiments are carried out in a 2 liter stirred reactor, operated under a range of pressures from atmospheric to 30 bars, and over a range of temperatures. A simple operating protocol allows us to generate solubility data for a limited cost. After introducing a known quantity of polymer into the reactor, an insoluble gas (e.g. Argon) is used to carefully measure the volume of the void space among the polymer particles and in the feed lines. Then, mixtures of soluble gases are injected into the reactor, the pressure is measured and since we know the volume of the gas phase, we know the number of moles of gas in the void. The difference between the moles fed to the reactor and the moles in the gas phase gives us the solubility. Gas phase composition is measured with an upgraded Micro GC, allowing us to estimate individual solubilities in mixes of different process gases.


Estimation of Diffusion Coefficients for Multiple Penetrant/Polymer Systems Based on Sorption Data

Rita Ferreira Alves, Timothy F.L. McKenna
Université de Lyon, Univ. Lyon 1, CPE Lyon, CNRS, C2P2-UMR 5265, Villeurbanne, France.

The ability to accurately estimate diffusion coefficients in polymers is extremely important for the design of reactors, unit operations, and in the modelling of polymerization kinetics, particularly in multicomponent systems. However, a very limited amount of data is found in the literature, and the non-ideal nature of the polymer-penetrant(s) interactions makes it complicated to estimate diffusivities à priori, more so if the co-solubility [1] effect is significant as is the case in olefin polymerization for example.

In this work the Sanchez-Lacombe Equation of State (SL-EoS) [1][2] is used to calculate the free volume of binary and ternary systems (solute(s)-polymer). A Free Volume Theory [3][4] (FVT) based model is then employed to estimate the diffusion coefficients of the solute(s) in the amorphous phase of the polymer. The advantages of the proposed methodology are that it relies on available experimental sorption data of gases in polymer and an equation of state, providing an attractive alternative to attain FVT parameters which otherwise would require arduous experimental data to estimate. The model is validated with published experimental diffusion results and shows good agreement.


Power compensation calorimetry for studying the kinetics of bulk-phase polymerization of propylene

Sina Valaei, Michael Bartke
Martin Luther Universität Halle-Wittenberg, Naturwissenschaftliche Fakultät II, Heinrich Damerow Straße 4, 06108 Halle(Saale), Germany

Polypropylene is one of the most important plastics in the market. Most common methods to produce polypropylene are gas phase- and bulk polymerization with supported Ziegler-Natta or metallocene catalysts.

For lab scale studies of gas-phase polymerization, reaction kinetics can be obtained by semi-batch operation of the reactor feeding monomer in a pressure control-loop at isothermal conditions. For bulk phase polymerization, this approach is not feasible and hence often the only kinetic information obtained from an experiment is the yield obtained by weighing of the polymer formed.

Here we report application of power compensation calorimetry for detailed online monitoring of reaction kinetics in bulk phase polymerization of propylene.

A power compensation calorimetry setup consists of an internal electrical heater submerged in the contents of a partially filled reactor (in this case liquid propylene) paired with an external cooler. The external cooler keeps the jacket temperature a few degrees below the desired reactor temperature, therefore the cooler is constantly removing energy from the reactor by conduction through the walls of the reactor. This heat flow to the cooling system is compensated by the electrical heater in order to keep the reactor in thermal equilibrium. Once an exothermic reaction such like polymerization takes place, less electrical power is needed in order to keep the reactor in isothermal conditions. In addition to an excellent temperature control, from the difference between the power needed in absence and presence of the exothermic reaction, one is able to calculate the heat released by the reaction.

One challenges for application of power compensation for bulk-phase polymerization of propylene is the significant change of filling level throughout the reaction due to the much higher density of the polymer compared to the monomer. Due to this change of filling level, the heat transfer area thus also the energy flow to the cooling circuit changes with conversion, which needs to be accounted for.

In this work, addition of power compensation calorimetry to an existing 5 Liter Büchi reactor for bulk phase polymerization of propylene is reported. For different filling levels, calibration experiments have been performed in order to estimate the heat transfer coefficient.

An online-sensor has been developed in order to estimate on the basis of the power compensation measurements, material balances and partitioning calculations, both the filling level, the heat flow to the cooling system as well the heat released by the reaction.

This online-sensor is used to study the kinetics of homo polymerizations of propylene with a supported Ziegler-Natta catalyst.

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Entanglement Formation mechanism of Ethylene Polymerization by the Heterogeneous Ziegler-Natta Catalysts

Yuming Chen\textsuperscript{a,b}, Wei Li\textsuperscript{b}, Lei Hu\textsuperscript{b}, Binbo Jiang\textsuperscript{a}, Jingdai Wang\textsuperscript{a}, Yongrong Yang\textsuperscript{a*}

\textsuperscript{a} State Key Laboratory of Chemical Engineering, Department of Chemical and Biological Engineering, Zhejiang University, Hangzhou, 310027, Zhejiang, P. R. China
\textsuperscript{b} Department of Polymer Science and Engineering, School of Material Science and Chemical Engineering, Ningbo University, Ningbo, 315211, Zhejiang, P. R. China

Chain entanglement was very important for adjusting the processability and mechanical property of nascent ultrahigh molecular weight polyethylene (UHMWPE). So far, it is still a mystery to unravel the formation mechanism of entanglements when the ethylene polymerization is conducted by the supported heterogeneous catalysts. In this study, a series of weakly entangled UHMWPE was synthesized by the polyhedral oligomeric silsesquioxane/MgCl\textsubscript{2} nanoaggregates modified Ziegler-Natta catalysts. The structure of nanoaggregates was evaluated by X-ray photoelectron spectroscopy (XPS), Density functional theory (DFT) simulations, and Scanning probe microscope (SPM) experiments, where the coordination strategy of MgCl\textsubscript{2} and hydroxyl of POSS was investigated. These nanoaggregates presented extremely low activity on ethylene polymerization and were proved to serve as isolators for separating the active sites and growing chains, where the entanglement density of nascent UHMWPE was non-linearly decayed with the increased numbers of nanoaggregates. Importantly, this non-linearly decayed effect contributed by increasing numbers of POSS/MgCl\textsubscript{2} isolators offset the exponential increment of entanglements upon rising the temperature, which was the essential reason for the successful synthesis of weakly entangled UHMWPE even at 85 °C. Finally, we have proposed the dependence and sensitivity of G\textsubscript{n} (t=0) (i.e., indicating the initial entanglement density of nascent polymers) on the polymerization activity, which was able to quantitatively trace the formation of entanglements during polymerization through the heterogeneous catalyst.

![Scheme 1. The entanglement formation mechanism diagram of nascent polyethylene with different polymerization conditions.](image)


Your notes:
The real-time detection and control of the atmosphere composition: Application in preparation of the PP/EPR in-reactor alloy

Zheng Zheng, Jijiang Hu, Bo-qing Li
State Key Laboratory of Chemical Engineering, Zhejiang University, Hangzhou 310027, China

In our recent work, a method based on cascade flow meters (i.e. a Coriolis flow meter, a thermal flow meter and a differential pressure flow meter in series on the drain line output of the reactor) has been developed for real-time detecting and controlling the atmosphere composition of the reaction system in multi-component gas phase copolymerization of olefins[1]. Here, we would like to report the application of this method in preparation of the in-reactor alloy of polypropylene/ethylene-propylene rubber (PP/EPR). The preparation process consists of slurry homopolymerization of propylene and gas phase copolymerization of ethylene and propylene with hydrogen modulation. As shown in Fig. (a), the atmosphere composition in the gas phase during the copolymerization stage has been well controlled in constant level. Compared with the feed mode of constant-proportion intake, the feed mode with constant-composition in reactor gives products a more uniform composition distributions and better mechanical properties. Using the real-time detection and control technology, the monomer consumption rates (Fig. (b)) and the copolymer composition of the rubber phase were obtained also. And thus, the monomer reactivity ratios during the gas phase copolymerization were investigated.

Ring-Opening Polymerization of Epoxides by DMC in Micro-reactor

Jing Zhao, Hong Fan, Bo-geng Li*

State Key Laboratory of Chemical Engineering, Zhejiang University, Hangzhou 310027, China

Polyols obtained by ring-opening polymerization of epoxides catalyzed by Co-Zn double metal cyanides (Co-Zn DMC) have many advantages such as narrow molecular weight distribution, lower unsaturated degree and less in impurities, which benefits the subsequent polyurethane foaming process. However, the reaction system is quite dangerous in tank reactor because it is highly exothermic, so it has to be conducted in way of semi-continuous feed-starving of monomer in industry. This severely influences the production efficiency and still needs huge investments in equipments. Micro-reactor improves the operation safety for the highly exothermic reaction due to its small volume, large surface to volume ratio. Herein, we report the ring-opening polymerization of propylene oxide catalyzed by Co-Zn DMC in micro-reactor system, propylene oxide and prepolymer of polypropylene glycol (PPG) containing Co-Zn DMC catalyst were pumped into the preheat lines, mixed in the micromixer and flowed through the delayed lines to the outlet where a back pressure valve was connected, and as a result, polyols of different molecular weight were obtained within several minutes and their molecular weight distribution (characterized by polydispersity index, abbreviated as PDI) was usually quite narrow (PDI<1.1). In addition, we studied the effects of residence time, temperature, catalyst concentration, reactants ratio and pressure on conversion, molecular weight and molecular weight distribution. These results show that the micro-reactor has good application prospects in industry of polyols.

Your notes:

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Controlled Synthesis of Polyolefin Copolymers with Rapid Curability for Photovoltaic Cell Encapsulation

Liqiong Luo, Kailun Zhang, Pingwei Liu, Bo-Geng Li, Wen-Jun Wang, Guangda Zhou
Zhejiang University, College of Chemical and Biological Engineering, Hangzhou, P.R. China
Hangzhou First Applied Material Co. Ltd, Hangzhou, P.R. China

Polymers such as ethylene–vinyl acetate (EVA) copolymers have been widely used as encapsulants in photovoltaic cell modules to prolong their outdoor lifetime. However, EVAs are prone to yellowing after long-term usage, which significantly reduces the photoelectric conversion efficiency of the cells. Alternatively, polyolefins (POs) with outstanding outdoor stability have been considered as a highly promising encapsulant material for the photovoltaic cell encapsulation. The inertness of the POs, however, results in a slow curing during the cell encapsulation process.

In this study, we address the problem through a polymer product engineering approach by controllably introducing pendant vinyl groups into ethylene-octene copolymers via terpolymerization of ethylene, octane-1, and an asymmetric diene. The vinyl functionalities are ready to react with crosslinking agents, which distinctly increases the PO curing rate measured by storage modulus (G’). We are able to precisely tune the vinyl content in the POs from 0 to 4.01 mol% through engineering the polymerization processes. Moreover, under the same crosslinking reaction condition, G’ was significantly increased from 11294 pa to 57090 pa. Interestingly, we also observed that the crosslinking reaction was beneficial to the product transparency and volume resistivity, and did not affect the damp-heat and UV aging performances of the resulting PO films.


Your notes:
Influence of Particle Morphology on Film Drying Stresses
Hesham Abdeldaim, José M. Asua
POLYMAT and Departamento de Química Aplicada, Facultad de Ciencias Químicas, University of the Basque Country UPV/EHU, JoxeMarí Korta Zentroa, Tolosa Hiribidea 72, Donostia-San Sebastián 20018, Spain

Adverse impacts on health and the environment are forcing paint and coating manufacturers to replace the volatile organic compounds in traditional formulations with water-based formulations. One of the approaches to achieve high film forming ability with excellent mechanical properties from aqueous dispersions is the use of heterogeneous particles with two phases. One phase with low Tg for coalescence forming the layer and the other phase with high Tg for providing the mechanical strength to the final film. During film drying, the material experiences a sequence of transformations, involving water loss, particle deformation, and interparticle coalescence. Depending on the drying conditions, the formulation, and the thickness of the coating, the films may suffer number of shrinkage cracks, which compromises the mechanical integrity of the film. Therefore, Understanding the evolution of the film formation stresses will provide manipulation and control over the film-formation process. In the present research, the effect of particles size and layer thickness on the film mechanical properties and cracking phenomena were studied. The film induced stresses during drying were estimated using WAXS and Beam bending techniques. Moreover, a physical mechanical finite element mathematical model was developed to predict the induced stresses through deformed two particles during drying process.

Correspondence to: hesham.abdeldaim@polymat.eu

Your notes:
Fully bio-derivable CO$_2$ fixing polymers for HNIPU applications

S.-E. Dechent, A. W. Kleij, G. A Luinstra

University of Hamburg, Institute for Technical and Macromolecular Chemistry, Bundesstr. 45, 20464 Hamburg, Germany, Institute of Chemical Research of Catalonia (ICIQ), The Barcelona Institute of Science and Technology (BIST), Av. Països Catalans 16, 43007 – Tarragona, Spain, Catalan Institute of Research and Advanced Studies (ICREA), Pg. Lluis Companys 23, 08010 – Barcelona, Spain

Non-isocyanate polyurethanes (NIPUs) are phosgene and isocyanate-free alternatives to standard industry polyurethanes. They are easily accessible from cyclic carbonates and amines. NIPUs and corresponding hybrid materials (HNIPUs) which are derived from epoxy resins partly modified with cyclic carbonates, hold an interesting promise of innovation as they in part show superior mechanical and chemical properties. However, finding an easy process remains a hurdle in NIPU synthesis as long reaction times and rather harsh reaction conditions are required.[1] A route involving polybutadiene (PBD) as a starting material for the sequence of post-functionalization of epoxidation, cyclic carboxylation, and diamine cross-linking could have an interesting potential as a future-proof/green route of routinely synthesizing HNIPUs.[2] Here it is reported how several low molecular polybutadiene were partial epoxidized and how a consecutive carboxylation with CO$_2$ can be achieved. This particular way of evolving polybutadienes allows for tailored HNIPU preparation; various functionalities are introduced onto the polymer backbone. We have tested a set of metal-free catalysts including conventional, widely used tetra-n-butylammonium bromide (TBAB) for CO$_2$ insertion reactions into partly epoxidized PBDs optimizing conditions to target quantitative conversions and high selectivities. Eco-friendly NIPU cross-linking reactions with (bio-derivable) diamines further exemplified the capability of polybutadienes in forming a backbone of a tunable, potentially fully sustainable hybrid NIPU thermoset.


High Barrier Waterborne Polysulfide Dispersions

J. Elgoyhen, R. Tomovska.
POLYMAT and Kimika Aplikatua Saila, Kimika Fakultatea, University of the Basque Country UPV/EHU, Joxe Mari Kortu Zentroa, Tolosa Hiribidea 72, 20018 Donostia-San Sebastián, Spain

Today, the polymer industry is going through significant changes in priorities driven by the growing awareness of ecological issues. In this context, the development of eco-efficient and sustainable processes and products has become an absolute necessity.

As an example, many industrial coating processes involve the use of waterborne products obtained by polymerization in dispersed media. These VOC-free polymer products are indeed a greener alternative to the traditional solvent borne coatings, as water is used as the disperse phase. Although the market of waterborne coatings is constantly increasing, the variety of polymer dispersions commercially available is restricted and relies mainly on chain growth radical polymerization using thermal or redox radical initiator. Hence, the development of polymerization technologies to expand the range of products and to make the process even more efficient and environmental friendly is needed. To answer to this need, we propose an approach that unites a photopolymerization in dispersed media technique with thiol-ene step-growth radical processes to produce polysulfide aqueous dispersed products.

Thiol-ene chemistry has been widely studied, and its application in coating has recently emerged [1] because of its numerous advantages such as rapid kinetics, control of the polymer architecture, and insensitivity to oxygen. [2] Thiol-ene photopolymerization has been used in UV-cured coating formulations [1], although its application is restricted to surface and thin-layer. Since the last decade, the feasibility of thiol-ene photopolymerization in miniemulsion has been reported [3-6]. While minimizing cost and environmental impact because of the lower consumption of energy and the use of a VOC-free process, thiol-ene photopolymerization in miniemulsion turns out to be a privileged mean to obtain high sulfur content products with controlled architecture.

Herein, we present how the polymer architecture in thiol-ene chemistry may be fine-tuned by adjusting the monomer functionalities. For that aim, a wide range of thiol and diene monomers was photopolymerized in miniemulsion. The bifunctional monomers resulted in linear structures prone to form crystalline domains within the polymer films, whereas, more than two functional groups per monomer molecule promoted controlled branching and cross-linking and network formation. This could pave the way to a new portfolio of polysulfide coatings, with versatile properties implied by the structure of polymers.

Determination of micromixing efficiency in viscous media

E. Arian, Hamburg, Germany; W. Pauer, Hamburg, Germany

University of Hamburg/ Institute of Technical and Macromolecular Chemistry,
Hamburg, Germany, pauer@chemie.uni-hamburg.de

Mixing, especially micromixing processes, have an influence on the yield, selectivity and quality of the final product. After determining the micromixing efficiency, the design of chemical reactors, impeller geometries and the operation conditions can be improved in order to obtain a high yield and a high selectivity of a desired product. One of the most common chemical test reaction is the Villermaux-Dushman reaction containing a neutralization reaction in a borate buffer and a comproportionation reaction yielding iodine due to the injection of an acid. While carrying out this reaction in water is simple, straightforward, and well established, the determination of the micromixing efficiency in higher viscosities is challenging. On the one hand, a suitable viscous media has to be selected and on the other hand, a compatible test reaction has to be found to operate in the selected viscous media.

A high concentrated sucrose solution has been applied, as it is a Newtonian fluid, which can achieve viscosities up to 1200 mPas. The borate buffer was replaced with a phosphate buffer giving two major advantages. Firstly, the handling of the chemical compounds was safe by abstaining from boric acid and secondly the buffer could be completely dissolved in concentrated sucrose solution. Initially, experiments were carried out in a stirred tank reactor yielding the segregation index as a function of the stirring speed at different viscosities using a sucrose solution. Furthermore, the results were transferred into a continuous tubular reactor, where the micromixing efficiency was also determined. Finally, an invert sugar solution was successfully tested as a viscous media for the test reaction with the advantage of reaching viscosities up to 15000 mPas. The result of these experiments highly suggests that sugar solutions can be used as a viscous media to determine the micromixing efficiency at different viscosities.


Your notes:
Refractive Index Increments of EVA-Copolymers

Elisabeth Schulz, Markus Busch
Technische Universität Darmstadt, Darmstadt, Germany

Due to their high versatility, lightweight, ease of processing as well as moldability to complex architectures, polymers have outgrown most human-made materials with a production of 322 million tons in 2015.[1,2] An example for a widely applicable polymer is polyethylene-vinyl acetate (EVA). EVA-copolymers hold a diverse spectrum of materials with a broad range in applications. Next to others, EVA-copolymers are employed in the wire and cable industry for heat shrinkable and flame retardant insulation.[3] Due to their complex structure-property-relationship in regard to suitable applications, it is mandatory to examine (co-)polymers as precisely as possible.

During the last decades, high temperature GPC-MALLS analysis of polymers has been well established. Hereby, the multi-angled laser light scattering analysis (MALLS) is employed to determine absolute molecular weights and other characteristics such as the radius of gyration ($R_g$) or long chain branching of polymers.[4] However, in order to apply this technique, the refractive index increment ($dn/dc$) of the respective polymer is essential. This polymer specific parameter is a function of the employed solvent, temperature and wavelength, which has to be carefully selected for appropriate evaluation of light scattering experiments.[5] While $dn/dc$ data of most homopolymers for various temperatures, wavelengths and solvents are available in literature, the accessibility of respective literature data for copolymers is scarce.

The presented work is an effort in order to supplement the published information of $dn/dc$ values of EVA-copolymers to enable a more accurate characterization in GPC-MALLS. For this, EVA-copolymers with varying vinyl acetate content are investigated. Measurements are conducted in a differential refractometer at a temperature of 150 °C with (1,2,4)-trichlorobenzene as solvent.

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