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Subtle Microwave-Induced Overheating Effects in an Industrial Demethylation Reaction and Their Direct Use in the Development of an Innovative Microwave Reactor

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Abstract: A systematic study of the conventional and microwave (MW) kinetics of an industrially relevant demethylation reaction is presented. In using industrially relevant reaction conditions the dominant influence of the solvent on the MW energy dissipation is avoided. Below the boiling point, the effect of MWs on the activation energy $E_a$ and $k_0$ is found nonexistent. Interestingly, under reflux conditions, the microwave-heated (MWH) reaction displays very pronounced zero-order kinetics, displaying a much higher reaction rate than observed for the conventionally thermal-heated (CTH) reaction. This is related to a different gas product (methyl bromide, MeBr) removal mechanism, changing from classic nucleation into gaseous bubbles to a facilitated removal through escaping gas/vapors. Additionally, the use of MWs compensates better for the strong heat losses in this reaction, associated with the boiling of HBr/water and the loss of MeBr, than under CTH. Through modeling, MWH was shown to occur inhomogeneously around gas/liquid interfaces, resulting in localized overheating in the very near vicinity of the bubbles, overall increasing the average heating rate in the bubble vicinity vis-à-vis the bulk of the liquid. Based on these observations and findings, a novel continuous reactor concept is proposed in which the escaping MeBr and the generated HBr/water vapors are the main driving forces for circulation. This reactor concept is generic in that it offers a viable and low cost option for the use of very strong acids and the managed removal/quenching of gaseous byproducts.

INTRODUCTION

The 20th century has been mainly dominated by the use of conventional thermal energy to drive chemical reactions. The use of alternative energy sources, such as microwave (MW) technology, appeared in the late 1970s, and its use could overcome existing bottlenecks in chemical manufacturing and improve the carbon footprint of many reactions. While it is clear that poor instrumentation has led to erroneous reports and set unrealistic expectations,* a systematic and controlled investigation of the influence of MWs on chemical reactions and their kinetics has been lacking. Such background information is however indispensable for possible future implementation of MW technology on an industrial scale. To date, many types of thermal and nonthermal MW effects have been reported; as a guide to past research in these reports, the interested reader is referred to the critical review by De La Hoz and co-workers. Presently, many of the “nonthermal MW effects” have been disproven as being the result of an incorrect temperature measurement. Also, the energy contained in MWs is far too low to break even hydrogen bonds. In the presence of non-MW absorbing solvents the groups of Dudley and Stiegman argue that both substrate and product can temporarily store energy, leading to localized temperature increases at the reactive site, accounting for the observed MW rate enhancements. More recently they also showed that poorly MW-absorbing molecules can be selectively heated by MWs provided association to a nonreactive polar molecule as a good MW absorber, albeit the effect is then less pronounced.

In a further advancement the Kappe group used Si–C vessels which were proclaimed to impede the penetration of MWs into the reaction vessel, heating the Si–C material instead, and thus administering in essence conventional heating. That way no difference in reactivity could be observed between conventional
and MW heated reactions.\(^1\) However, very recently the Strouse group showed convincingly that 3 mm Si—C tube walls only retain \(~48\%\) of the MWs, pointing at significant MW transmission into, and dissipation inside, the reaction vessel.\(^2\) This MW leakage through Si—C tube walls is particularly acute when a strongly absorbing solvent is used.\(^3\) Fan et al. have pioneered the use of MW technology to the hydrothermal depolymerization of cellulose to glucose, demonstrating a distinct influence of the applied MW density and a MW heat input via molecular radiators;\(^4\) notably, they used proton/deuterium exchange techniques to also obtain structural information. Many research groups continue to propose differing activation energies for reactions run in the presence of CTH vis-à-vis MWH. However, the determination of the kinetic parameters \((E_a, k_0)\) of a reaction strongly depends on the applied model and thus requires a thorough analytical method. In this study we have investigated the MW activation of an industrially (pharmaceutically) relevant demethylation reaction, converting (3-methoxyphenyl)methylammonium bromide (3MPMA) into (3-hydroxyphenyl)methylammonium bromide (3HPMA) (see Scheme 1)\(^5,6\) with a detailed kinetics study. For this purpose, we have used both a SAIREM MiniFlow200SS\(^7\) with TM monomode cavity and an Anton Paar Monowave 300, both of which are equipped with fiber-optic temperature measurement. The MiniFlow200 uses a solid-state MW generator, which enables precise frequency and microwave power control; in addition, it features forward and reflected power measurement so that an energy balance can be obtained. The absence of this latter feature in most commonly used MW heating equipment has been demonstrated to result in significant misreadings of the actual MW power transferred to the sample under investigation.\(^8\) The experimental study is complemented by a simulation study to obtain additional insight into the interacting physical phenomena: electromagnetics, fluid dynamics, and heat transfer. The demethylation reaction in this study is typically run on multitonne scale and generally employs limited amounts of solvent and reagents, therewith increasing the efficiency of the process and its productivity. This reaction however has received little mechanistic attention, especially under relevant reaction conditions. From a MW point of view, the investigation of a polar reaction with high substrate/product dipoles and continuously changing dielectric properties, as the reaction progresses, presents a great opportunity to gain more knowledge and understanding of how the use of MWs could potentially benefit such chemical reactions.\(^9,10\)

**Scheme 1. Overview of the Reaction**

To assess the potential influence of different heating methods on the 3MPMA to 3HPMA demethylation reaction, kinetic reaction profiles (as conversion—time plots) were first established for the case of an open vessel CTH covering the 90—118 °C temperature range (Figure 1A). The reaction mixture consisted of 5.038 g of 3MPMA (0.0367 mol) and 12.38 g of 48% HBr (0.0734 mol HBr; twice excess vis-à-vis substrate), and no additional solvent was added. Temperature measurement was performed in a dual way, recording both the oil bath temperature and, by fiber-optic temperature probe (FOTP), the internal reaction temperature. Interestingly, Figure 1A shows that the rate of the reaction becomes equal for oil bath temperatures ≥115 °C, equating by FOTP to an effective internal maximal reaction temperature of 112—112.5 °C. A linear correlation between the target (oil bath) and the recorded fiber-optic temperatures is observed only up to 113 °C (oil bath) (Figure 1B).

Kinetic analysis of the reaction profiles in Figure 1A shows a complex kinetic behavior which can generally not be explained as a single first- or second-order process for the entire conversion range. More specifically, the beginning and the end of the reaction appear to behave as different consecutive second-order processes, which interchange at the 50—60% conversion level (Figure 1S). In the 90 °C case, the conversion level is below 50% showing thus only the first process. To explain this complex behavior, it was hypothesized that throughout the reaction the number of available protons is reduced by (reversible) protonation of 3HPMA (Scheme 1S). This becomes particularly important in the later stages of the reaction, as close to stoichiometric amounts of reagents are employed in this reaction. As shown in Figure 2, this model fits the experimental kinetic data very well for all temperatures. In a second approach, conversion—time plots...
plots were also established for the MWH reaction. Kinetic analysis of this data shows that the two-step model presented in Scheme 1S (eq 15a) can only be applied up to 111.2 °C (FTOP) (Figure 3A). Furthermore, in the 90−113 °C range the E_a/k_0 values of the CTH and MWH reactions vary only slightly (Figures 3B/2B). As shown in Figure 4A, the reaction rate observed at ~98 °C is independent of the type of heating while at ~113 °C, MW operation leads to a markedly higher rate of reaction than observed with CTH (Figure 4B). Moreover, Figure 4B shows pronounced zero-order behavior under MW operation at 113.85 °C while under CTH, classic second-order is observed. As the recorded temperature for both heating types (MWH, CTH) is 113 °C, within experimental error, this does demonstrate a pronounced change in the reaction mechanism between the MW and the conventional thermally run reaction.

A multiphysics simulation, primarily using Comsol Multiphysics 5.2 (see also the section on numerical simulation in Supporting Information), was conducted to create additional insight into the electromagnetics, the fluid dynamics, and the heat transfer. To the best of our knowledge, only two studies have previously been published covering the combined simulation of electromagnetic heating and conductive/convective heat transfer in bench-scale MW chemistry systems. These studies both concerned heating in a CEM Discover MW device. More specifically, Robinson et al. studied the MW heating of a variety of solvents in a stirred vial, but their conducted simulation did not include a complete fluid dynamics model. In contrast, Sturm et al. focused on the heating of water in a MW field, including a laminar fluid dynamics model to simulate the free convection under nonstirring conditions. In addition, for the continuous flow case, Patil et al. presented an experimentally verified numerical study into a MW heated millireactor; in particular, they demonstrated temperature measurement deviations due to large thermal gradients around the sensor. To model the MWH demethylation reaction case adequately, a much more advanced methodology was required, covering all the relevant physical phenomena and stirring. Furthermore, to correctly predict the electromagnetic field inside the cavity and the reactor, knowledge was required of the dielectric properties of the reaction mixture at the relevant temperatures. The medium...
properties were measured and determined to be $\varepsilon' = 17$ and $\sigma = 3.3 \, \text{S/m}$. The dissipative ($\sigma$) term is best described as an electrical conductivity due to the high concentration of ions in solution. A visual setup is provided in Figure 5a, and additional details are available from Supporting Information and Table 1S. The fluid agitation in the reactant mixture was simulated by applying a rotating geometrical domain, to account for stirrer bar rotation, in combination with the $k$-$\varepsilon$ Reynolds-averaged Navier–Stokes model accounting for turbulence and turbulent heat transfer. Figure 5b–e shows the main modeling results for the multiphysics simulation. Additionally, an animated video is available from Supporting Information (video 1S). More specifically Figure 5b shows the MW field (by means of the electric field intensity on logarithmic scale) in and around the cavity applicator and reactor. The simulation shows that the electromagnetic emission and dissipation in the reactor and cavity construction materials are negligible. In addition, it also reconfirms the general nonuniformity of MW fields.\(^{13}\) The latter feature also expresses itself in the heat generation simulation presented in Figure 5c, where it can be seen that the main zone of heat generation occurs close to the MW antenna of the cavity. However, in Figure 5d it is shown that for the applied stirring speed, the temperature variations in the reactor are too small to have any significant effect on the reaction rate, i.e., less than 1.5% rate variation for the highest temperatures and less than 0.4% below 111 °C. Figure 5e shows the overall electromagnetic dissipation and heat loss to the surroundings versus temperature for both the simulated and the experimental case. It can be seen that the simulation correctly approximates the characteristics of the experimental energy balance. The curves lie close to each other, and both have an accelerating heat loss as the temperature approaches the boiling point. Figure 5f shows a quasi-electrostatic analysis of the MW field around a bubble. The MW field is deformed by the presence of the bubble. Red zones, representing localized overheating, and blue zones, indicative of relatively cooler zones, can be observed. However, these do not cancel out: there is a \(\sim40\%\) average increase in heat generation in a layer of \(\sim0.1\) times the bubble radius. Generally the evaporation of hydrobromic acid into a bubble extracts heat from the reactant liquid adjacent to this bubble due to the expansion of the bubble, which can result in a reduction of vapor pressure and consequently a potential bubble collapse. Though the flow regime in the reactor is turbulent due to stirring, the Kolmogorov length scale is calculated to be 30 to 240 µm.\(^{17}\) Below this scale, bubbles do not benefit from turbulent convective heat transfer, and their growth potential is limited unless a directly adjacent heat source is present. For the CTH case, only the bubbles contacting the heated reactor wall can grow, so that fewer and larger bubbles are formed. In comparison, for the MWH case, the presence of a locally enhanced volumetric heat source enables many more bubbles to form. This mechanism agrees well with the observed differences in boiling regime observed during experimentation; illustrative videos of the MWH (video 2S) and CTH (videos 3S and 4S) demethylation reaction at 112 °C are included in Supporting Information.

As the removal of MeBr gas through water/HBr vapor will be governed by the available surface between the reaction mixture and the water vapor bubbles, the occurrence of zero-order kinetics may relate to the available interface becoming insufficient to remove all produced MeBr formed at one given point in time. Scheme 2 represents the proposed mechanism for higher temperature MW operation. The derived rate equation fits the high temperature MW kinetic data very well (see Scheme 2S, eq 28b, and Figure 2S). An additional observation in the MW transmission transient during experiments can be made; Figure 3S shows that fluctuations in the power regulation occur less rapidly with increasing reaction temperature, which indicates a change in medium properties as...
the reactant mixture progresses from mechanism 1 to 2. Further
258 to the use of open vessel reactors, we also evaluated the use of a
259 closed vessel, i.e., using a pressure NMR tube for CTH and an
Anton Paar MW closed vessel for the MWH reaction. As
260 shown in Table 1, no distinct MW influence is observed when
261 the reaction is performed in a closed vessel reactor. Indeed, in
262 closed vessel operation no mass-transfer limitation problem
263 arises as the produced MeBr builds up a pressure of ~110 psi
264 (54.7% conversion) (Figure 4S), in that way shifting the
265 equilibrium from MeBr gas to MeBr liquid.
266
267 The observation of intense steam/gas bubble production in
268 the high temperature MWH reaction provided an interesting
269 opportunity for the development of a novel continuous MW
270 reactor concept in which MW energy is actually converted to
271 kinetic energy, i.e., the escaping MeBr, and the generated HBr/
272 water vapor can drive the reaction mixture around a loop. This
273 concept is similar to gas/air-lift reactors, which find common
274 application in industrial biotechnology and multiphase
275 processes, but contrary to the concept proposed here, these
276 rely on the introduction of a separate gas/air stream. The
277 development of a continuous MW reactor for the demethylation
278 reaction presented here holds distinct industrial advantages, notably (1) a controlled release and thus manageable
279 scrubbing of toxic MeBr, (2) a continuous all-glass reactor
280 concept tailored to the use of strongly corrosive acids, avoiding
281 the need for expensive specialty alloys (e.g., Hastelloy), (3) the
282 avoidance of an expensive pumping system capable of
283 withstanding MeBr/HBr, (4) the absence of moving parts,
284 and (5) enhanced mass transfer properties. Figures 5S and 6S
285 show respectively the schematics of the circular and the
continuous MW reactor. A video of the circular MW reactor in
286 operation, employing a PI of 140 W, is included in Supporting
287 Information (video SS), and the conversion–time plots are
288 shown in Figure 7S.

Table 1. Conversion Levels for the Demethylation Reaction
Performing at 118 °C in an Anton Paar MW Closed Vessel and,
for the Conventional Heated Counterpart, an NMR Pressure Tube

<table>
<thead>
<tr>
<th>entry</th>
<th>reactor type</th>
<th>2 h reaction time</th>
<th>4 h reaction time</th>
<th>6 h reaction time</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Anton Paar MW closed vessel reactor*</td>
<td>49.5</td>
<td>64.4</td>
<td>70.9</td>
</tr>
<tr>
<td>2</td>
<td>conventional by NMR pressure tube</td>
<td>47.6</td>
<td>64.5</td>
<td>70.7</td>
</tr>
</tbody>
</table>

*Ratio gas to liquid phase in Anton Paar reaction vessel and NMR pressure tube are the same.

CONCLUSION
291 In summary, we have shown that the main influence of MWH,
292 vis-à-vis CTH, on the kinetic parameters of an industrially
293 relevant demethylation reaction occurs only under reflux
294 conditions. Thus, the use of MWs opens a different mechanism
295 for the elimination of gaseous byproducts (e.g., MeBr), by the
296 creation of vast amounts of bubbles, therewith changing the
297 observed reaction order of the demethylation reaction from 2
298 to 0. Through modeling, the origin of this change in reaction
299 order was shown to relate to a deformation of the microwave
300 field in the presence of bubbles, leading to localized overheating
301 in the close vicinity of the bubbles. Based on these insights, a
302 novel continuous MW reactor concept could be proposed in
303 which MW energy is converted into kinetic energy, making the
304 production and removal of MeBr the driving force for the reactor. This offers a generic reactor concept for reaction types
305 in which significant amounts of gaseous byproducts (e.g., de(m)ethylation, metathesis, dehydration) are created.

ASSOCIATED CONTENT
309
Supporting Information
310, and illustrative videos are available in the Supporting
311 Information. The Supporting Information is available free of
312 charge on the ACS Publications website at DOI: 10.1021/
313 jacs.7b00689.
314
Experimental details, additional figures, and reaction
315 schemes/models (PDF)
316 Simulation results of SAIREM TM cavity for demethyla-
317 tion of (3-methoxyphenyl)methylammonium bromide
318 (MPG)
319 Illustrative video of the MWH demethylation reaction at
112 °C (MPG)
320 Illustrative video of the CTH demethylation reaction at
112 °C (MPG)
321 Illustrative video of the CTH demethylation reaction at
112 °C (MPG)
322 Video of the circular MW reactor in operation employing
323 a PI of 140 W (MPG)
324

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