Flow chemistry experiments in the undergraduate teaching laboratory: synthesis of diazo dyes and disulfides

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Received: 15 July 2020 / Accepted: 21 September 2020 / Published online: 7 October 2020 © The Author(s) 2020

Abstract
By embedding flow technology in the early phases of academic education, students are exposed to both the theoretical and practical aspects of this modern and widely-used technology. Herein, two laboratory flow experiments are described which have been carried out by first year undergraduate students at Eindhoven University of Technology. The experiments are designed to be relatively risk-free and they exploit widely available equipment and cheap capillary flow reactors. The experiments allow students to develop a hands-on understanding of continuous processing and give them insights in both organic chemistry and chemical engineering. Furthermore, they learn about the benefits of microreactors, continuous processing, multistep reaction sequences and multiphase chemistry. Undoubtedly, such skills are highly valued in both academia and the chemical industry.

Keywords Flow chemistry · First-year undergraduate · Organic chemistry · Chemical engineering · Hands-on training

Introduction
Continuous-flow chemistry has rapidly established itself as a go-to technology to carry out difficult-to-handle reagents and reaction conditions [1, 2]. Notable examples where flow has made an undeniable impact are photochemistry [3–5], electrochemistry [6–8], multiphase reactions [9–12] and handling of toxic, explosive or other hazardous reagents and intermediates [13–15, 16, 17]. Its value has been recognized by the industry as well and many companies are establishing small expert groups specialized in flow chemistry [18, 19]. Hence, there is a high demand for skilled personnel with at least a notion of flow chemistry. It is therefore obvious that flow chemistry courses should be implemented in the academic training of future generations of chemists and chemical engineers [20–29].

However, flow chemistry courses in the chemistry curriculum are actually conspicuous by their absence. Historically, undergraduate teaching laboratories have focused on the use of round-bottom flasks to carry out organic chemistry experiments. Although, it is important for undergraduates to learn to work with this conventional and widely-used labware, it is all too often the only reactor type that students see in their entire education. Herein, we demonstrate that a broader education with regard to reactor technologies should not require a complete rethinking of the classical curriculum nor should it be expensive.

Implementation of flow chemistry in both theoretical and practical courses should be a focus of every chemistry major. Indeed, some reactions are better carried out in a flow reactor. As an example, some of the typical synthetic organic chemistry experiments require long reaction times (several hours or days) which is not always advantageous in a busy teaching schedule. Such reactions can often be accelerated in flow reactors via a strategy called process intensification [30, 31]. This would allow students to carry out more experiments within the given time slot. Moreover, hazardous reactions can be carried out in flow without risk, e.g. by generating in...
situ small quantities of a toxic substance which is subsequently reacted away in a follow-up transformation. Such experiments provide students with fundamental insights into reaction safety and the development of safe reaction environments. Furthermore, students can get insight in the importance of fundamental transport phenomena, like mass and heat transport. This is especially important for multiphase reaction conditions (gas-liquid or liquid-liquid biphasic reactions) or for exothermic reactions, two notoriously challenging examples to execute and to scale in both academia and industry [32].

As many chemical engineering and chemistry students will pursue a career in industry, where continuous processes are often encountered, it is paramount that students are familiar with the basic principles of continuous manufacturing and flow chemistry. While organic chemistry often lingers with batch processing, chemical engineering often sticks to residence time distribution experiments or the determination of other reactor parameters. By combining organic synthesis with continuous-flow techniques, students from both chemistry and chemical engineering tracks can learn from each other’s disciplines, i.e. chemists acquire skills in continuous processing while engineers pick up the benefits for the synthesis of organic molecules. This will create a mutual understanding, fostering more fruitful collaborations in the future.

Herein, we disclose our experience with two flow chemistry experiments which have been implemented in the opening teaching laboratory course at Eindhoven University of Technology. One experiment involves the synthesis of hazardous diazonium salts which can be subsequently reacted away to yield a diazo dye (Solvent Yellow 7). A second experiment involves an organic-aqueous biphasic oxidation of thiols to disulfides and provides insight in segmented flow. Both experiments have been carried out by >400 students in the past 4 years and have been well received.

**Experiment 1: Synthesis of solvent yellow 7 in continuous flow**

The synthesis of Solvent Yellow 7 involves two distinct steps, including a diazotization to yield the corresponding diazonium compound and a subsequent diazo coupling reaction with phenol to generate the targeted diazo dye (Fig. 1a). This experiment is an example of multistep flow reaction sequences [33–35] and shows convincingly how to handle hazardous intermediates. The explosion-sensitive diazonium salts are generated in the first step and immediately reacted away in an electrophilic substitution reaction [36, 37]. Hence, at any given time, the total inventory of the hazardous intermediate is kept small alleviating the safety risks associated with this transformation. The first reaction takes place under acidic conditions, while the second reaction requires basic conditions. All starting materials were selected because of their low cost and relative low toxicity. The concentration of the different solutions is low ($C = 0.01$ M) which is advantageous both in terms of reduced chemical consumption and enhanced safety.

The setup is assembled by the students in pairs as shown schematically in Fig. 1b. The two reactors are made from perfluoroalkoxy alkane (PFA) capillaries (ID = 760 μm, volume 1 mL) according to a procedure described elsewhere (Fig. 1c) [38]. The reactors are connected to the syringe pump, to each other and to the outlet with standard microfluidic fittings. Once the setup is assembled, the different solutions are made with the proper concentrations. To gain time, the solutions can also be made by the supervising tutors. The solutions are taken up in 4 plastic syringes (BD Discardit II®), which are subsequently mounted onto a syringe pump. In addition, the concentrations of the reagents are calculated so that all syringes are pushed at the same flow rate. By doing so, only a single syringe pump is required per experiment, reducing further the cost associated with this experiment. Next, several samples are collected (1 mL/sample) for different residence times ($t_R = 15, 30, 45, 60, 90, 120, 150$ and $300$ s). Prior to each sample, students need to wait four residence times in order to ensure steady state data collection. Samples are subsequently diluted and the conversion is measured using a benchtop UV-vis spectrometer ($\lambda = 400$ nm).

Next, the students are asked to analyze their results. First, they need to calculate the kinetic constant. The reaction is first order in aniline, which allows us to use the following equation:

\[
\frac{dC}{dt} = kC
\]  

with $C$, the concentration [M] and $t$, the reaction time [s]. Integration of this equation leads to:

\[
\ln \left( \frac{1}{1-X} \right) = kt
\]

with

\[
X = \frac{C_0 - C_t}{C_0}
\]

The value of the rate constant ($k$) can be readily obtained by plotting the conversion in function of the residence time (Fig. 2a). Next, $\ln \left[1/(1-X)\right]$ is plotted in function of the residence time (Fig. 2b). The value for the rate constant can be subsequently derived from its slope, i.e. $k = 0.0045$ s$^{-1}$. While slight variations are obtained by different sets of students, the order of magnitude of the rate constant is the same for all.

Next, we ask the students to calculate the time needed to obtain 99% conversion of the starting material. By using equation 2 with $X = 0.99$, it can be easily understood that $t_R \geq 1024$ s.
Equation 2 can be reorganized to result into the following equation:

\[ X = 1 - e^{-kt} \]  

(4)
gives confidence to the students that their calculations were accurate.

In principle, the entire experiment requires about four hours of research time per pair of students, this includes setting up the reactor assembly, carrying out the experiments and performing the UV-vis measurements. Since this experiment is done by first year students in the first month of their academic track, we have decided to spread the experiment over two half days. The first day is mainly used to construct the setup and ensuring its proper performance, e.g. no leakage, chemicals mixed in right stoichiometry, calculating the required residence times and getting familiar with the analysis equipment. On the second day the data can be collected and handled. A detailed procedure can be found in the Supporting Information.

**Experiment 2: Dimerization of octanethiol in continuous flow**

A second experiment was introduced in our undergraduate teaching laboratory to showcase the advantages provided by microreactor technology for the execution of multiphase reaction conditions (Fig. 3). Upon merging two immiscible phases, a segmented flow regime is established in the capillary microreactor where the two phases are brought into close contact with each other [40]. Segmented flow provides fast mixing due so-called Taylor recirculation flow patterns, a well-defined interfacial area and reduced axial dispersion, allowing the reactor to operate as an ideal plug-flow reactor (Fig. 3b).

We selected the synthesis of disulfides via oxidative dimerization of thiols in flow, which has been developed previously in our group using photocatalytic [41–43] or electrochemical activation [44]. For the practical course, we wanted a robust, yet operationally simple procedure which avoided the use of mass flow controllers to dose air or oxygen into the liquid stream [45]. A biphasic reaction protocol was developed with 0.1 M octanethiol and 1 mol% I$_2$ in ethyl acetate and a 10% aqueous hydrogen peroxide solution (Fig. 3a). Both solutions are taken up in two syringes and mounted on a single syringe pump (Fig. 3c). Upon merging of the two solutions, a segmented flow regime can readily be observed. The reaction is completed in about 5 min residence time. After discarding the first four reactor volumes to reach steady state, the students are asked to collect sufficient material to isolate the disulfide (20 mL solution, to isolate 1 mmol of the targeted disulfide). This solution is subsequently washed with 0.2 M thiosulfate in a separatory funnel. Note that this is an exothermic reaction which consumes the excess of iodine and thus caution is required. Hence, the combined layers have to be gently shaken and pressure needs to be released from the separatory funnel. The organic phase is subsequently washed two more times with fresh 0.2 M thiosulfate solution. The organic phase is dried with magnesium sulfate and evaporated. The latter step can be preferably done in vacuo using a rotavapor or, since we have >100 students, evaporation can be done in a fume hood by heating the flask in a water bath. The reaction should in principle afford the corresponding disulfide in quantitative yield (276 mg of a transparent oil), but we observed sometimes yields >100% due to insufficient removal of the solvent. All experimental operations can be carried out by the students in four hours.

**Assessment**

This practical course is for some students their first experimental experience. Consequently, every practical experiment is preceded by an in-depth discussion of the mechanisms of
Students need to be able to follow the flow procedures accurately, including building up the flow setup, calculating the flow rates correctly to reach certain residence times and executing the recipes with accuracy.

2. They have to abide the safety regulations at all times.

3. They have to be able to accurately note down the made observations and the obtained results in the laboratory notebook.

The written report can be assessed more objectively on its completeness, e.g., accurate description of the observations, correct calculations and plots, adequate conclusion and perspective. We tolerate small mistakes (e.g., rate constant is off), especially when the students assess their performance accurately in the conclusion. Generally, the written reports on the flow experiments are of good to excellent quality. However, it is important to note that the flow experiments are the last ones in the practical course. Hence, the students have by that time already more experience in writing reports and have received ample feedback on previous reports.

Hazards

While microreactors can handle safely small amounts of hazardous chemicals, one should still be careful as the stock solutions and the collecting vessels contain still several mL of solution. Links to appropriate safety hazards data for all chemicals are included in the Supporting Information.

Conclusions

Two laboratory experiments that incorporate microflow techniques were developed for first year undergraduate students at the Eindhoven University of Technology. These experiments are embedded in a general Undergraduate Teaching Laboratory and introduces the concept of continuous-flow processing. It trains the students in a practical manner, which enables them to understand the theory behind the use of flow reactors. The students also get acquainted with building a reactor setup, how to calculate flow rates and convert those to reaction times, reaction stoichiometry and flow synthesis. Importantly, our experiments use cheap and easy-to-operate capillary microreactors and only a single syringe pump per experiment. We believe that these experiments can be readily implemented into both chemistry and chemical engineering curricula.

Acknowledgments

The execution of a practical course with >100 students per annum is without a doubt a formidable task effort at the department of Chemical Engineering and Chemistry at TU Eindhoven. We would like to thank ir. P. F. A. M. (Peter) Janssens, who is the Director of Education at the department of Chemistry and Chemical Engineering, for his support during the implementation of flow chemistry in the curriculum. Also, we would like to thank all the student assistants in the past five years who helped guiding and supervising this practical course. Finally, we like to thank Ms. Christa Schilders for the administrative tasks during the practicum.

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