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1 Introduction

There is an ever-growing incentive to move towards more sustainable and more efficient processes. Within the chemical industry, process intensification and the use of renewable feedstock can allow for major contributions to this end. The decrease in environmental impact can be further amplified by using less toxic and biodegradable reactants for producing chemicals. Surfactants account for a large share of the chemical industry. Their current total market size is \$39.69 billion, which is continuously expanding with approximately 2.5% each year.¹

Quaternary ammonium salts (QAS) are one of the primary surfactants, with an annual production estimated at millions of tonnes.^{2,3} These QASs are widely applied, in households as well as in various industries, with applications ranging from detergents³ to phase-transfer catalysts.^{3,4} Currently, the SARS-CoV-2 pandemic results in an increased demand in QASs, due

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Renewable dimethyl carbonate for tertiary amine quaternisation: kinetic measurements and process optimisation

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Quaternary ammonium salts (QAS) are an important part of the increasing surfactant market. Conventional production processes employ toxic alkyl halides in a Menshutkin reaction with a tertiary amine (DMDA). Dimethyl carbonate (DMC) can provide a renewable route, while also leading to more benign, and non-corrosive products. This work aims to use linear ramp-flow in a plug flow reactor (PFR), combined with inline ¹H NMR spectroscopy to determine reaction kinetics. These kinetics will be used to further optimise the production process with a computational model. Solvent effects were first studied in a batch reactor. Methanol (MeOH) was found most suitable as a solvent. Subsequently, the reaction kinetics were measured in a PFR set-up. The used ramp-flow was successfully validated with data from batch and steady-state experiments. Arrhenius parameters were determined with the ramp-flow method, which proved to be an accurate and efficient technique. The kinetic data was implemented in a computational model. After validation of the model with experimental data, it was employed to extrapolate this data and optimise the reaction. The optimum QAS productivity was predicted at 122 kg h⁻¹ L⁻¹, obtained at 270 °C, 0.25 min residence time, and a molar ratio of 1:2.5:10 (DMDA:DMC:MeOH). These conditions would provide significant intensification of the QAS production processes.

to their effectiveness as disinfectants.⁵ Their conventional production is performed according to the Menshutkin reaction, which was first published in 1890.⁶ Nowadays this reaction is still carried out almost exclusively in large batch reactors. As operation of this type of reactor is generally labour-intensive, costly, and not very scalable, progressing towards a continuous process seems paramount. Other challenges in the production process are introduced by the QAS's high viscosity, and the consecutive thermal degradation reaction. At higher temperatures, QASs are (depending on their configuration and anion) increasingly susceptible for decomposing according to Hofmann elimination.⁷

In the Menshutkin reaction, a tertiary amine reacts with an alkyl halide (predominately methyl chloride), as shown in Fig. 1.^{3,8} Under mild temperatures, between 60 and 100 °C, methyl chloride, benzyl chloride, dimethyl sulfate and diethyl sulfate are commonly employed to complete the quaternisation reaction.^{9,10} The reaction is well-known to occur according to an $S_N 2$ mechanism, in which the nature of the leaving group and the nucleophilic strength of the nitrogen base determine the rate of reaction.³ Although these alkylating agents allow for relatively high reaction rates, they are accompanied with high toxicities and environmental impacts.³ Dimethyl sulfate, *e.g.*, has a low LD₅₀ of 133 mg



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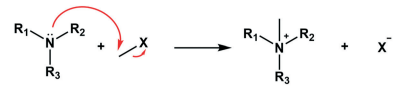


Fig. 1 The Menshutkin reaction of a tertiary amine with an alkyl halide according to an $S_N 2$ mechanism.^{3,6}

kg⁻¹,^{11–14} and is also known to be carcinogenic.¹⁵ As a result, their allowed concentrations in the final products is constrained to ppm levels to ensure safety, which complicates product purification.¹⁶

Dimethyl carbonate (DMC) has been posed as a highpotential renewable building block for various chemicals.^{17,18} It has negligible toxicity, is biodegradable, and can be produced sustainably from methanol (MeOH) and CO or CO₂ precursors.^{13,19,20} For its use as an alkylating agent in QAS production processes, several patents and articles have already been published.^{21–26} The resulting QAS features no corrosivity, making it eligible for specific demanding and specialty applications. In Fig. 2, an exemplary alkylation reaction with DMC is presented. Besides its high flammability, DMC is not hazardous, as there is no long-term toxicity and it has a high LD₅₀ at 5000 mg kg⁻¹.²⁰ Its high thermal stability up to 390 °C allows for high-temperature operation and reactant recovery.²⁷

The main disadvantage of DMC is its relatively low reaction rate compared to the aforementioned conventional reagents. For trialkyl amines, for example, the reaction rate constant with DMC at 125 °C (5.5×10^{-6} L mol⁻¹ s⁻¹) is similar to that of methyl chloride at room temperature (6.0×10^{-6} L mol⁻¹ s⁻¹).² However, as already stated by Friedli¹⁶ in 1990, it remains difficult to find literature relating reaction kinetics of particular amines. Often, tertiary amines with equally long side-chains are investigated. Previous studies on DMC alkylation rates in MeOH with various trialkyl amines have shown an activation energy of 79 kJ mol⁻¹ and rate

constants from 2.9×10^6 L mol⁻¹ s⁻¹ to 2.1×10^7 L mol⁻¹ s⁻¹ for temperatures of 115 to 155 °C.² For these trialkyl amines, the activation energy is independent for chain-lengths beyond three carbons.^{2,16}

Alkylation reactions with DMC are also highly susceptible to solvent effects.^{16,28} Reaction intermediates are known to be best stabilised by polar aprotic solvents, *e.g.*, acetonitrile, acetone, and propylene glycol.²⁹ Nevertheless, polar protic solvents such as MeOH, ethanol, and isopropanol are also typically employed for these types of reactions.^{2,16,28}

Another key advantage of using DMC as an alkylating agent, is the fact that it produces a QAS with an anion that can be readily exchanged according to the scheme in Fig. 3. This reaction results in the formation of carbon dioxide and MeOH which are both highly volatile, favouring the consecutive purification. Any organic or inorganic acid can facilitate this exchange, as long as their pK_a value is lower than methyl carbonic acid (pK_a 5.612).² Although ion exchange is also possible after the conventional alkylation process, it is very impractical as it is difficult to remove the resulting salts from the product.³⁰

Together with the side-chain length, the anion has a profound influence on the properties of the QAS and thus on its potential applications. Currently, about 500 different QASs are produced and commercially used.³¹ DMC-based QASs have the potential to be versatile platforms to produce a much wider range of QASs.

This research focuses on optimising the quaternisation of *N*, *N*-dimethyldecylamine (DMDA) with DMC (Fig. 2). This tertiary

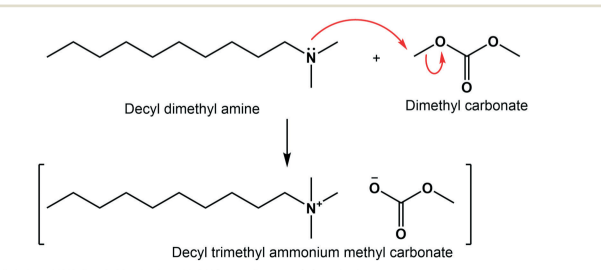


Fig. 2 The alkylation of N,N-dimethyldecylamine with DMC according to an S_N2 mechanism.



amine's kinetics have not been reported in literature, while it leads to industrially relevant products. First, solvent effects are investigated in a batch reactor, after which the most suitable solvent is used in kinetic experiments in flow. These experiments are carried out using linear ramp-flow under continuous analysis of the reactor outflow with Proton nuclear magnetic resonance (¹H NMR) spectroscopy. Similar methods have been applied in other studies.^{10,32-34} The kinetic parameters and solvent effects will then be used in a computational model of the flow reactor. This model is employed to extrapolate and optimise the operating conditions to maximise the yield and productivity of the reactor.

2 Methods

2.1 Materials

The materials used, their origin, and their purity are listed in Table 1. The DMDA was known to contain traces of other tertiary amines, some secondary amines, and very low amounts of unreacted primary amines. The solvents that are shown were used to determine solvent effects only in batch reactions and their purity was at least 99%. MeOH, however, was also used in the flow experiments. The main source of impurity in the solvents was water.

2.2 Solvent effects in batch

Batch experiments were performed to determine solvent effects on the alkylation rate of DMDA with excess DMC. The batch reactors consisted of a 1.5 mL glass vial, in which the liquid was contained by a membrane crimp cap. These reactors were filled with pre-prepared stock solutions, consisting of the concerning solvent, DMDA, DMC, and ¹H NMR standards. The ratio of the amine to DMC was varied between 1:5 and 1:10, while the ratio of the amine to solvent was varied between 1:0 (neat) and 1:15.

The vials were inserted in a heated mantle or an oil bath,
where a temperature was set ranging from 80 to 130 °C. For
each experiment, a series of six vials were filled with the
stock solution. One vial acted as a blank, while the others
were inserted into the heating source simultaneously, and
removed, one by one, at fixed time intervals. After their
removal, the vials were directly subjected to a thermal
quench in an ice bath. For the samples that showed
segregation, MeOH was added until a single phase was
obtained again. This dilution was accounted for by the
internal standard present in the mixture.

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The resulting time series was used to determine reaction rate constants for the various reaction mixtures and temperatures. These rate constants were determined by computing the conversion of the amine, X_A , according to eqn (1), which corresponds with a second-order kinetic equation. Here, the initial concentration of compound *i* ($C_{i,0}$), the reaction rate constant (k), and the residence time (τ) are the required input parameters. The consecutive fitting of the Arrhenius equation allowed for the determination of the desired kinetic parameters.

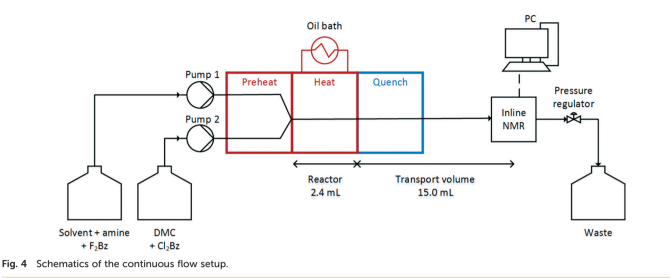
$$X_{\rm A} = \frac{\exp[(C_{\rm DMC,0} - C_{\rm A,0})k\tau] - 1}{\exp[(C_{\rm DMC0} - C_{\rm A,0})k\tau] - \frac{C_{\rm A,0}}{C_{\rm DMC,0}}}$$
(1)

2.3 Reaction kinetics in a plug flow reactor

The kinetic experiments in flow were performed in a PFR setup, of which a schematic is presented in Fig. 4. For reference, this set-up has also been used and described in previous work.¹⁰ Two Shimadzu CL 20 AD HPLC pumps were used to separately feed the two reactant mixtures through a pre-heat loop into the reactor. The pumps had been calibrated for their respective mixture. The first mixture consisted of the

Chemical	Туре	Supplier	Grade		
N,N-Dimethyldecylamine	Reactant	TCI	≥95% purity		
Dimethyl carbonate	Reactant	Sigma Aldrich	≥99% purity		
Methanol	Solvent	VWR Chemicals	Technical grade		
Ethanol	Solvent	Merck	Absolute grade		
Isopropanol	Solvent	VWR Chemicals	Technical grade		
Acetone	Solvent	Merck	Analysis grade		
Acetonitrile	Solvent	Biosolve	HPLC grade		
Dimethyl acetamide	Solvent	Alfa Aesar	≥99% purity		
Dimethyl sulfoxide	Solvent	Merck	Analysis grade		
1,4-Dichlorobenzene (Cl ₂ Bz)	¹ H NMR standard	Alfa Aesar	≥99% purity		
1,4-Difluorobenzene (F ₂ Bz)	¹ H NMR standard	Sigma Aldrich	≥99% purity		
Deuterated chloroform	¹ H NMR solvent	VWR Chemicals	≥99.80% purity		

 Table 1
 The used chemicals with their supplier and grade



solvent, DMDA, and a F_2Bz internal standard, while the second mixture contained DMC and a Cl_2Bz internal standard. This double internal standard approach allowed for determining individual compound concentration at the back-end of the reactor, and also to verify the flow and mixing ratio of the two inlet streams. Before each experiment, all solutions were stirred, and degassed with a helium purge.

The pre-heat loop consisted of two isolated 1/16'' stainless steel tubes (with an inner diameter of 1 mm), enclosed by HM-RD2011SO heat tracing from Mohr & Co, which was regulated with a thermocouple. The reactor was devised from a 1/8'' stainless steel tube (with a wall thickness of 0.711 mm for an inner diameter of 1.75 mm), with a length of 1 m for a total volume of 2.4 mL. Prior to the reactor volume, one of the pre-heat tubes was adapted to the 1/8'', in which the other 1/16'' tube was inserted. The resulting concentric tubes continued for 1 m, after which the inner tube was terminated, the inlet streams mixed, and the reaction volume started. The design allowed for inserting or retracting the inner tube, for a variable effective reactor volume. The entire 2 m 1/8'' tubing was fed through a 1/2'' stainless steel tube, connected to a LAUDA Proline P5 thermostat oil bath, serving as a heat exchanger. This ensured that the reactants were mixed at the desired operating temperature, and that the heat of reaction was removed. Isothermal operation could thus be assumed for the entire reaction volume. As this design made mixing at a precise spatial coordinate possible, the reactor volume can be considered very well-defined. The reactor system of concentric tubes has been schematically depicted in Fig. 5.

After the reactor, the tubing was immediately led through an ice bath to cool the reaction mixture to 0 °C. At this temperature, the reaction rate is negligible, and the reaction can thus be considered successfully quenched. Subsequently, the stainless steel tubing was converted to perfluoralkoxy alkane (which does not interfere with the spectrometer), and fed through a Magritek Spinsolve benchtop 43 spectrometer. After the ¹H NMR, a back-pressure regulator was installed to have a constant elevated pressure of 30 bar, which ensured a liquid phase for all compounds.

Kinetic experiments were conducted in a range from 130 to 160 °C. After equilibrating and reaching a steady-state in the

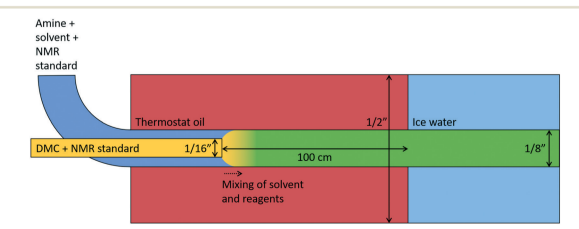


Fig. 5 Schematic representation of the reactor system of concentric tubes. The design allowed for a very well-defined and controlled reactor volume.

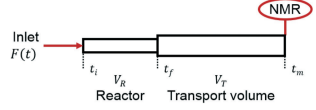


Fig. 6 Graphical representation of the relevant volumes and experimental times in the continuous flow setup.

reactor, a linear decreasing flow ramp was introduced. This resulted in modulation of the residence time. Therefore, a single experiment allowed for measuring reactor outlet concentrations at increasing reaction times. Both HPLC pumps were programmed, and although their ratio remained constant, total flow rates were varied between 0.20 and 1.60 mL min⁻¹. For each temperature, a separate experiment was carried out.

2.4 Ramp-flow and data processing

The ramp-flow method employs a linear decrease in the volumetric flow rate, resulting in an increasing residence time in the reactor, $\tau_{\rm R}$, over the duration of the experiment. Similar methods have been described with, *e.g.* model-based design of experiments, IR analysis, or HPLC analysis.^{35–38} In this study's method, the experimental time has to be converted into the actual residence time. The required processing has been described in previous work.¹⁰ One profound difference in this case, however, is the fact that no reaction occurs in the transport volume ($V_{\rm T}$). Fig. 6 shows a graphical representation of the set-up, where t_i , t_f , and t_m are the times at which a discrete element flowing through the reactor reaches its entrance, its exit, and the ¹H NMR sensor.

These times were used to compute the required residence times, according to eqn (2). The different situations in which a discrete element could be travelling through the set-up before, during, and after the flow ramp, are displayed in Fig. 7. The initial volumetric flow rate is represented by F_0 , while the final flow rate is shown as F_e .

$$\begin{aligned} \tau_{\rm R} &= t_{\rm f} - t_{\rm i} \\ \tau_{\rm T} &= t_{\rm m} - t_{\rm f} \end{aligned} \tag{2}$$

Accounting for the ramp of the flow, *a*, started at t_0 and ended at t_e , leads to the flow rate developing as a function of time as described in eqn (4). Under the assumption of plug-flow, and with the imposed flow ramp, the concerning reaction volumes can be expressed by eqn (3).

$$V_{\rm R} = \int_{t_{\rm i}}^{t_0} F_0 dt + \int_{t_0}^{t_{\rm f}} F_0 + at dt$$
(3)
$$V_{\rm T} = \int_{t_{\rm f}}^{t_{\rm m}} F_0 + at dt$$

$$t < t_0: F(t) = F_0$$

$$t_0 \le t < t_e: F(t) = F_0 - at$$

$$t > t_e: F(t) = F_0 - at_e = F_e$$
(4)

Eqn (3) can be integrated and substituted to acquire the desired set of equations for the unknown experimental times, t_i and t_f . These equations are shown in eqn (5), along with their associated boundary conditions. Subsequent substitution of these equations into eqn (2), results in calculation of the required residence times of the set-up.

$$t > t_{0} \wedge t_{i} \leq t < t_{f}: \quad \tau_{R} = t_{f} - t_{i}$$

$$t_{f} = -\frac{F_{0} + \sqrt{F_{0}^{2} - 2a\left(V_{T} - t_{m}\left(F_{0} + \frac{1}{2}at_{m}\right)\right)}}{a} \quad (5)$$

$$t_{i} = t_{f} + \frac{\frac{1}{2}at_{f}^{2} - V_{R}}{F_{0}}$$

Finally, a representative flow ramp and the resulting residence time profile in the reactor are shown in Fig. 8. Note that this residence time is based on ideal plug-flow behaviour, and is a mere estimation of the actual residence time.

2.5 ¹H NMR spectroscopy

For all kinetic measurements, ¹H NMR spectroscopy was used to determine the concentrations of the individual compounds in the reaction mixtures. The batch measurements were

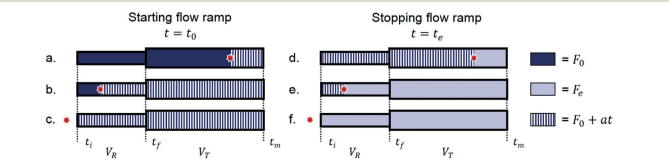


Fig. 7 An element travelling through the setup can be in six discrete situations.

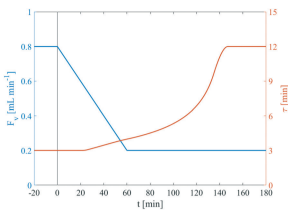


Fig. 8 Example of a flow ramp and the resulting residence time in the reactor for a reactor volume of 2.4 mL, and a transport volume of 15 mL.

carried out offline, in a Bruker-400 400 MHz NMR spectrometer. Prior to the analysis, 0.1 mL of sample was diluted with 0.4 mL of deuterated chloroform in a NMR tube, and subjected to 32 scans with a recycle delay of 2 s. Typical spectra before, and after the reaction are shown in Fig. 9c and d, respectively. The assigned protons are elaborated in Fig. 9a. The peak areas in the spectra were determined after an automated phase correction, an automatic third order Bernstein polynomial fit baselinecorrection, and one zero-filling step using the forward linear prediction method of Zhu–Bax. An internal standard allowed for the conversion from integrated area to the concentration of the respective species.

The analysis for the flow experiments was performed using a Magritek Spinsolve benchtop 43 NMR spectrometer (43 MHz). The ¹H NMR was calibrated and relative standard deviations lower than 1.5% were observed. As the reaction mixture was flowed through the NMR, *i.e.* measured in-line, the data acquisition was automated. At the sensor, every 6.4 s a spectrum was recorded with a 90° pulse angle. A powershim, executed before each experiment with a 0.5 mL min⁻¹ flow rate ensured the spectrometer was locked on the highest intensity peak (3.48 ppm, MeOH). During the experiments, the ¹H NMR was re-calibrated at 30 min intervals, to preclude any decay of the signal. The spectrometer could be operated with total flow rates of up to 2.5 mL min⁻¹, approximately. This maximum is dictated by the sensor volume and the acquisition time settings.

2.6 Reactor characterisation

In order to verify if plug-flow can be assumed for the setup, a reactor characterisation study was conducted. This study included a numerical verification, as well as experimental work. For the numerical characterisation, Reynolds and Péclet dimensionless numbers were used, as shown in eqn (6) and (7), where u and d represent the superficial velocity through, and the inner diameter of the tube, correspondingly.

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$$R_{\rm e} = \frac{ud}{v} \tag{6}$$

$$Pe = \frac{ud}{D}$$
(7)

It is generally accepted that Péclet numbers approaching zero indicate an ideally mixed system, while plug-flow is assumed from Péclet numbers exceeding 100.^{39,40} The diffusion coefficient, *D*, is computed according to the Wilke-Chang correlation, shown in eqn (8).^{41,42} A system of dilute DMC in MeOH is assumed, as its diffusion coefficient in MeOH is expected to be higher than that of the tertiary amine due to its smaller size. Furthermore, the dynamic viscosity, μ , of pure MeOH was calculated according to Xiang *et al.*,⁴³ and the characteristic length, *d*, is equal to 1.75 mm (the reactor's inner diameter). Finally, the association factor (*x*) has been reported to be 1.9.⁴⁴

$$D = 7.4 \times 10^{-14} \frac{T(xM_{\rm MeOH})^{\frac{1}{2}}}{\mu_{\rm MeOH} \overline{V}_{\rm DMC}^{0.6}}$$
(8)

Table 2 shows the resulting dimensionless numbers. For the operated range of temperatures, these have been calculated at the lowest, and highest flow rate. With Reynolds numbers not exceeding 112, all experiments exclusively operated in the laminar regime. Nevertheless, the Péclet numbers clearly demonstrate overall plug-flow behaviour, as only at the lowest temperature and flow, a value lower than 100 is observed.

The experimental characterisation was performed by means of a residence time distribution (RTD) analysis. The PFR set-up was subjected to a tracer step input, while keeping the total volumetric flow rate constant. Total flow rates between 0.15 and 2.00 mL min⁻¹ were applied with a MeOH solvent and DMDA as the traces. These RTD experiments were thus very representative of the kinetic experiments. The concentration of the tertiary amine was determined as elaborated in the ¹H NMR spectroscopy section. The resulting concentration profile was used to generate the corresponding *F*-curves, and to calculate the corresponding tanks-in-series (TIS), as well as the backmixing time.

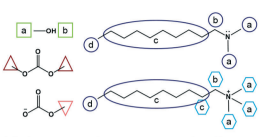
3 Modelling approach

3.1 Model structure

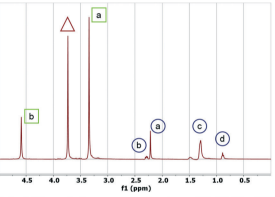
A computational model was built in close resemblance to the operated PFR. The model was employed to extrapolate the obtained experimental results into conditions that were infeasible in the PFR. It aims to represent the PFR, and it was validated using the experimental data. Thereafter, optimisation of the system's yield and productivity was carried out.

As the reactor characterisation has shown, the reactor operates according to non-ideal plug-flow. In this case, the

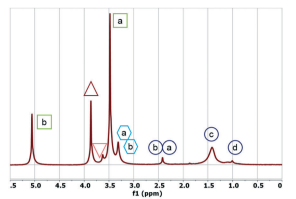
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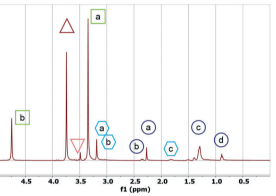
(a) Assignment of unique protons for all species present in the reaction mixture.



(c) Offline, before quaternisation, $^{1}\mathrm{H}$ NMR (400 MHz, none) δ 4.59 (s, 1H), 3.74 (s, 6H), 3.35 (s, 3H), 2.37 - 2.24 (m, 2H), 2.22 (s, 8H), 2.00 - 1.08 (m, 16H), 0.89 (m, 3H).



(b) Online, during quaternisation, ¹H NMR (43 MHz, none) δ 5.06 (s, 1H), 4.06 - 3.73 (s, 6H), 3.73 - 3.58 (s, 3H), 3.58 - 3.38 (s, 3H), 3.38 - 3.10 (m, 11H), 2.24 (m, 8H), 2.10 - 0.77 (m, 19H).



(d) Offline, after quaternisation, ¹H NMR (400 MHz, none) δ 4.75 (s, 1H), 3.74 (s, 6H), 3.49 (s, 3H), 3.35 (s, 3H), 3.19 (m, 11H), 2.44 – 2.31 (m, 2H), 2.27 (d, 6H), 1.82 (m, 1H), 2.00 – 1.08 (m, 16H), 0.88 (m, 3H).

Fig. 9 1 H NMR spectra of the quaternisation of DMDA with DMC in MeOH. b) shows a spectrum taken during flow reaction at 160 $^{\circ}$ C with a residence time of 6 minutes, reaching 60% amine conversion. c) and d) show spectra taken before and after batch reaction at 90 $^{\circ}$ C for 4 hours, also reaching 60% amine conversion. The line thickness for all spectra is the same, but the online spectrum has a lower signal-to-noise ratio.

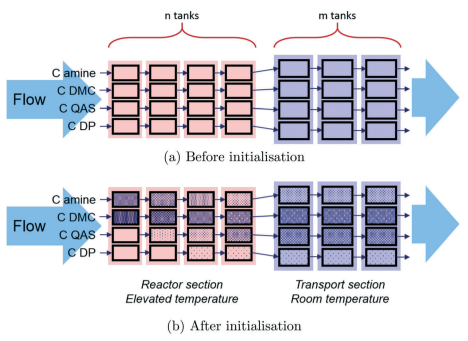
system can be approached by a dispersion model or a TIS model, which are equably suitable.³⁹ However, the TIS model does not require diffusion rate data, so this model is selected. The model was built in MATLAB, and solved by means of a ode45 ODE solver. Fig. 10 presents a schematic representation of the model. It consists of *n* tanks in $V_{\rm R}$, and *m* tanks in $V_{\rm T}$, depending on the operating conditions. Each compound has a mass balance for each cell, which communicates with the adjacent cells. The resulting balance

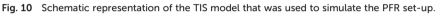
for a compound in tank i is shown in eqn (9). In accordance with the experiments, the reactor was considered to be operated at isothermal conditions.

$$\frac{\mathrm{d}C_i}{\mathrm{d}t} = \frac{F_{\mathrm{v}}}{V_{\mathrm{R}}} (C_{i-1} - C_i) N_{\mathrm{tanks}} - r_{\mathrm{tot}} \tag{9}$$

The overall reaction term (r_{tot}) consists of the QAS formation reaction, the degradation reaction, or both,

Table 2 The Reynolds and Péclet numbers at the used operating conditions						
Т	Low flow	High flow	Low Re	High Re	Low Pe	High Pe
[°C]	$[mL min^{-1}]$	$[mL min^{-1}]$	_	_	_	_
130	0.05	0.20	3	10	59	238
140	0.10	0.40	6	23	104	417
150	0.20	0.80	12	50	181	723
160	0.40	1.60	28	112	309	1235





depending on the species. All tanks were initialised at the respective initial concentrations, and at t = 0 the flow, heating, and reactions were started. Full convergence is obtained at equilibrium for all concentrations.

3.2 Parameter input

The actual PFR properties were used in the model ($V_{\rm R} = 2.4$ mL and $V_{\rm T} = 15$ mL). This ensured that the model could be validated with the experimental data. The reaction rate constants were computed based on the Arrhenius parameters that were determined with the ramp-flow experiments. For the degradation reaction, Arrhenius parameters from previous work were used.⁷

For the validation of the model, the input parameters were matched with the operating conditions. The flow rate was set between 0.05 and 1.60 mL min⁻¹, and the temperature was set between 130 and 160 °C. Resulting concentration and conversion profiles were compared with the respective experimental data.

After the validation was satisfactory, the operating conditions were expanded for the optimisation. Various molar ratios of DMDA:DMC:MeOH were varied, and a temperature of up to 300 °C was investigated. The flow rates were now set between 0.005 and 10 mL min⁻¹, corresponding with residence times of *ca.* 480 to 0.25 min. The main monitored design specifications of the model were the yield of QAS and the specific productivity of the reactor.

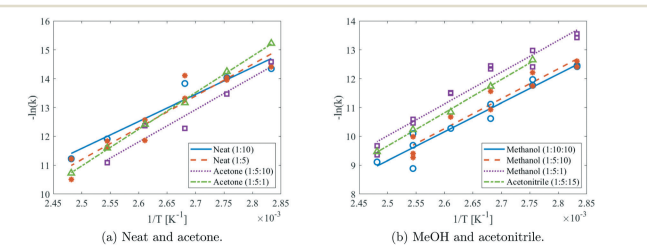


Fig. 11 Arrhenius plots of the batch experiments in different solvents and molar ratios.

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Table 3 Activation energies and frequency factors with 95% confidence bounds for different solvents and molar ratios

Reaction mixture DMDA:DMC:solvent	$E_{\rm a} \left[\rm kJ \ mol^{-1} \right]$	$-\ln k_0 []$	$k_0 \left[10^7 \text{ L mol}^{-1} \text{ s}^{-1} \right]$	<i>t</i> (<i>X</i> = 90%) at 150 °C [min]
MeOH (1:10:10)	83.2 ± 0.5	15.8 ± 0.1	0.760	16
MeOH (1:5:10)	86.3 ± 0.5	16.7 ± 0.2	1.80	26
MeOH (1:5:1)	91.6 ± 0.4	17.5 ± 0.1	4.06	32
Neat (1:10:0)	77.9 ± 0.8	11.8 ± 0.2	0.0139	133
Neat (1:5:0)	91.5 ± 0.6	16.3 ± 0.2	1.22	94
Acetone (1:5:10)	93.3 ± 0.9	17.4 ± 0.3	3.48	116
Acetone (1:5:1)	105.5 ± 0.1	20.7 ± 0.1	102	67
Acetonitrile (1:5:15)	95.8 ± 0.2	19.1 ± 0.1	20.6	42

4 Results and discussion

4.1 Solvent effects on quaternisation rate

Reaction kinetics of DMDA with DMC were determined in batch in MeOH, acetonitrile, acetone, and excess DMC. Isopropanol, ethanol, dimethyl acetamide, and dimethyl sulfoxide were also investigated, but their presence caused a side reaction with DMC. These side reactions were confirmed after further testing: DMC was added to all solvents, along with an internal standard and subjected to 130 °C for 28 hours. Side reactions were then indeed only observed in isopropanol, ethanol, dimethyl acetamide, and dimethyl sulfoxide, while the other solvents were confirmed to be stable. In these solvents, reaction rate constants of DMDA quaternisation were successfully determined in different ratios. A subsequent Arrhenius fit was performed and has been displayed in Fig. 11. Variations in concentrations were accounted for according to eqn (1) and this equation was used to fit the k values. As the graphs of different reactant ratios in the same solvent overlap, it appears that the second-order fit assumption is validated.

The Arrhenius parameters that resulted from the fit are presented in Table 3, along with their 95% confidence bounds. For reference purposes, the required reaction time to reach 90% conversion has also been computed. These indicate that MeOH is the preferable solvent, as it features the lowest required reaction times. Therefore, this solvent was selected for further use in the kinetic measurements in flow. Furthermore, at lower temperatures the neat production is outperforming the acetone system, while this is reversed at higher temperatures.

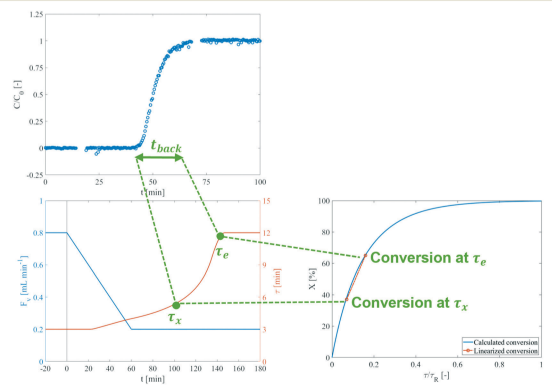


Fig. 12 An illustrative explanation of the check on the plug-flow assumption. If the change in conversion over the change in residence time from τ_x to τ_e , which correspond to the backmixing time t_{back} from the RTD, can assumed to be linear, the plug-flow assumption can be applied. The used plots are not from experiments at similar conditions, so the time axes do not match.

Table 4 The times used to estimate the change in conversion X, from τ_x to τ_e

$T [^{\circ}C]$	$\tau_{\rm e}$ [min]	$t_{\rm back} [{\rm min}]$	$\tau_{\rm x}$ [min]	$X(\tau_{\rm e})$ [%]	$X(\tau_{\rm x})$ [%]
130	48	192	20.9	67	40
140	24	96	10.2	65	37
145	12	48	5.2	52	28
150	12	48	5.2	62	36
155	6	24	2.6	49	26
160	6	24	3.4	49	32

Table 5 Operating conditions, fitted reaction rate constants, and confidence intervals for flow experiments in MeOH (130 to 160 °C)

<i>Т</i> [°С]	$ au_{ m i}$ [min]	$ au_{ m e}$ [min]	t _{ramp} [min]	Fitted k [10 ⁻⁴ mole per L s ⁻¹]	$\frac{R_{\mathrm{adj}}^2}{[]}$
130	12	48	240	1.31 ± 0.021	0.8718
140	6	24	120	2.94 ± 0.019	0.9780
145	3	12	60	3.73 ± 0.023	0.9869
150	3	12	60	4.76 ± 0.031	0.9845
155	1.5	6	30	7.21 ± 0.058	0.9856
160	1.5	6	60	8.01 ± 0.062	0.9946

4.2 Residence time distribution in the plug flow reactor

The RTD experiments resulted in *F*-curves for varying operating conditions within the intended range for kinetic experiments. Fig. 12 contains a representative *F*-curve, acquired at 0.35 mL min⁻¹. All curves showed a sharp breakthrough with a broadening tail, which is indicative of a slight deviation from ideal plug-flow.³⁹ The residence time at the midpoint of the *F*-curve was used to validate the set-up's total volume at 17.4 mL (of which 2.4 mL for the reactor section). A TIS model was fitted to the *F*-curves to determine the corresponding number of tanks. All except one of the RTD experiments (at 0.8 mL min⁻¹) resulted in a number of tanks greater than 50, at which plug-flow can be assumed.^{39,40}

Experimental validation of the plug-flow assumption was also achieved by estimating the change in conversion over the backmixing time, t_{back} . For an ideal PFR, this value would be 0. However, if the conversion can be approximated to be

linear for the concerning range, the distribution of residence times will result in an evenly-averaged instantaneous conversion. The backmixing time as a function of flow was fitted with respect to all RTD experiments, resulting in $t_{\text{back}} =$ $9.60/F_v = 9.60 \cdot \tau_e/V_R (R^2 = 0.831)$. The backmixing will be most severe at lower flow rates, *c.q.* the highest τ , at the end of the experiment (τ_e). The residence time of the fluid element that can be backmixed with the flow at τ_e , τ_{xy} is given by eqn (10).

$$t(\tau_{\rm x}) = t(\tau_{\rm e}) - t_{\rm back} \tag{10}$$

For all flow experiments, the resulting t_{back} , τ_{e} , and τ_{x} have been compiled in Table 4, along with the respective computed conversions. The change in conversion is the largest for the experiment at 140 °C, with other temperatures showing very similar ranges. Even the largest change in conversion is nearly linear, as shown in Fig. 12. Plug-flow can thus indeed be assumed for all conducted experiments.

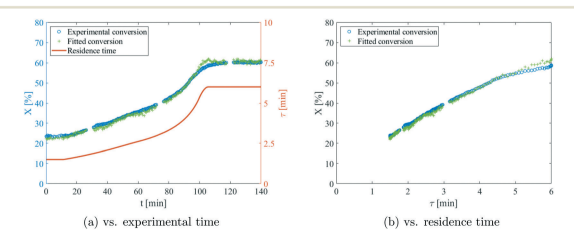


Fig. 13 Example of a flow experiment and its fit. This experiment was run at 160 °C and 1:5:10 ratio DMDA:DMC:MeOH. Gaps in the data are due to shimming.

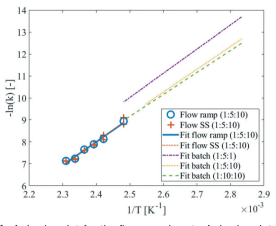


Fig. 14 Arrhenius plot for the flow experiments. Arrhenius plots from batch and steady-state flow data show good agreement. The molar ratios (DMDA:DMC:MeOH) provided in the legend.

4.3 Kinetic parameters in flow

The flow experiments with the linear ramp produced extensive sets of transient data for temperatures ranging from 130 to 160 °C (DMDA: DMC: MeOH = 1:5:10). These conditions and the residence times (Table 5) were selected based on the batch results and with respect to some physical limitations of the set-up. The data was fitted in accordance with eqn (1), resulting in conversion profiles. An example, obtained at 160 °C has been plotted in Fig. 13a. One key advantage of the ramp-flow method is distinctly illustrated here: a large amount of data points is acquired in a single experiment. The development of the experimental conversion clearly follows the trend of the residence time. Furthermore, the fitted conversion, based on the fitted reaction rate constant, is in agreement with the data. In Fig. 13b, the conversion (at 160 °C) has been plotted vs. the investigated residence time.

Subsequently, for each temperature a reaction rate constant was computed. These constants are presented in Table 5, and they are accompanied by the respective operating conditions, confidence intervals, and the R_{adj}^2 values. All experiments show high coefficients of determination, except for the *k* value at 130 °C. Nevertheless, no systemic error was observed in the data at this temperature. Arrhenius fitting then resulted in the

determination of the kinetic parameters for the 1:5:10 system. The computed activation energy is 88.5 ± 0.3 kJ mol⁻¹ and the frequency factor is $4.14 \pm 0.09 \times 10^7$ L mol⁻¹ s⁻¹. These values correspond well with the Arrhenius parameters in batch, presented earlier.

4.4 Method comparison

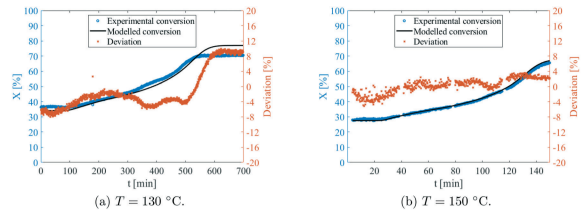
The linear ramp-flow method certainly has potential as a more efficient technique to acquire accurate kinetic data.¹⁰ However, the method is relatively novel and not very established yet. Therefore, validation of the method with respect to conventional methods, e.g. batch and steady-state flow, is paramount. Besides the reported batch and rampflow kinetic experiments, steady-state measurements were also conducted. In the Arrhenius plot in Fig. 14, the results acquired from batch, steady-state, and the ramp-flow method can be directly compared. The reaction constants of steadystate and ramp-flow are in very good agreement. And while the batch experiments were conducted at different temperatures, extrapolation of the fitted curve would evidently result in an acceptable fit of the flow data as well. Moreover, the kinetic parameters that were calculated from the steady-state data ($E_a = 91.1 \pm 0.1 \text{ kJ mol}^{-1}$ and $k_0 = 8.81 \pm$ 0.11×10^7 L mol⁻¹ s⁻¹), are also in accordance with those from the ramp-flow.

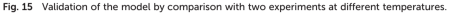
The comparison between the steady-state and the rampflow results has been presented in more detail in Table 6. For all experiments, the reaction rate constants and their relative difference with the corresponding ramp-flow values are shown. Steady-state reaction rate constants were determined at the upper flow rate, as well as at the lower flow rate. Deviations vary from 15.5% to -15.2%, with values as low as -0.7%. In general, the deviations are considered to be acceptable and not systemic. Ultimately, also based on the agreement with the batch results, the ramp-flow method is deemed to be validated.

As mentioned earlier, the ramp-flow method allows for the acquisition of a vast amount of transient data in a singular experiment. With the investigated operating conditions at least 150 data points were generated per run. Similar amounts of data are unfeasible with steady-state methods. Two steady-state equilibrations are required per data set with the ramp-flow method, in contrast to one equilibration for

 Table 6
 Comparison of the reaction rate constants determined with a flow ramp and at steady state flow. In brackets, the relative difference between the reaction rate constant from the flow ramp and the steady-state flow is shown

_	Ramp k	Steady-state k			
Temperature [°C]	$[10^{-4} \text{ L} mol^{-1} \text{ s}^{-1}]$	High flow $[10^{-4} L \text{ mol}^{-1} \text{ s}^{-1}]$	Low flow $[10^{-4} \text{ L mol}^{-1} \text{ s}^{-1}]$		
130	1.31 ± 0.021	$1.50 \pm 0.020 \ (15.5\%)$	$1.11 \pm 0.017 (-15.2\%)$		
140	2.94 ± 0.019	$3.05 \pm 0.018 (3.8\%)$	2.50 ± 0.022 (-14.9%)		
145	3.73 ± 0.023	3.80 ± 0.024 (1.8%)	3.90 ± 0.026 (4.5%)		
150	4.76 ± 0.031	$5.00 \pm 0.028 (5.0\%)$	$4.59 \pm 0.026 (-3.6\%)$		
155	7.21 ± 0.058	$7.30 \pm 0.049 (1.3\%)$	$7.53 \pm 0.052 (4.4\%)$		
160	8.01 ± 0.062	8.12 ± 0.069 (1.4%)	$7.96 \pm 0.066 (-0.7\%)$		





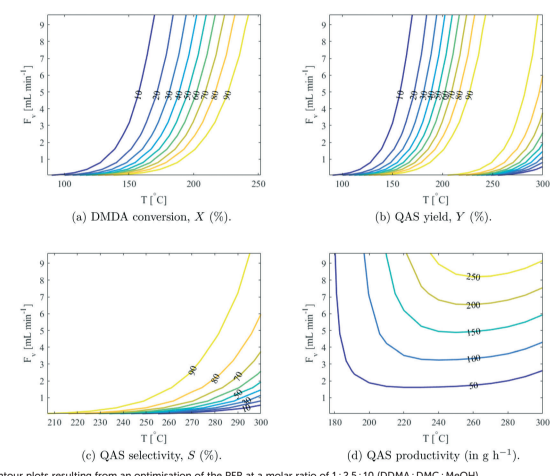


Fig. 16 Contour plots resulting from an optimisation of the PFR at a molar ratio of 1:2.5:10 (DDMA:DMC:MeOH).

every data point with the steady-state method. So even for a minimum amount of five data points, the time and material savings would be quite significant.

The comparison with batch experiments is less obvious. Approximately 150 mL of chemicals was used per experiment, which amounts to *ca.* 1 mL per data point. This could be further reduced by decreasing the transport volume to the spectrometer in particular. Nevertheless, this is still achievable in batch. However, the manual labour and time

required for preparing and analysing the batch samples is significantly higher. Especially at large-scale experiments that demand vast amounts of data, the ramp-flow method is profoundly superior.

4.5 Model validation

Prior to using the model for optimisation, it was first validated with experimental data. By settings the exact same

Table 7 Optimised reactor conditions for several molar ratios and reactor types (X is the conversion and S is the selectivity of the reaction). Molar ratio in DDMA: DMC: MeOH. Optimised variable in bold

Molar ratio	Reactor	t [min]	T [°C]	X [%]	S [%]	Productivity [g h ⁻¹]
1:2.5:10	PFR	0.25	270	99.71	97.51	293
1:2.5:10	PFR	0.25	220	64.67	99.91	194
1:2.5:10	PFR	48	150	100.00	99.90	1.5
1:5:10	PFR	0.25	250	99.72	99.20	251
1:5:10	PFR	0.25	220	84.21	99.90	213
1:5:10	PFR	48	140	99.36	99.96	0.1
1:10:10	PFR	0.25	260	99.04	98.65	132
1:10:10	PFR	0.25	220	96.99	99.91	86
1:10:10	PFR	48	150	99.28	99.90	0.7
1:2.5:10	CSTR	0.25	280	89.24	95.23	256
1:2.5:10	CSTR	0.25	210	39.97	99.93	120
1:5:10	CSTR	0.25	290	90.69	92.26	212
1:5:10	CSTR	0.25	210	36.33	99.93	92

operating conditions as the ramp-flow experiments, a direct comparison with the model could be made. In total, the results were compared at six different temperatures, in accordance to the ones shown in Tables 5 and 6. Two examples are shown in Fig. 15. A peculiar, yet acceptable deviation is observed at 130 °C, as can be seen in Fig. 15a. At lower times, the model predicts conversions that are a bit too high, though at higher times the model overshoots quite severely. No explanation was found for this remarkable observation which was absent at other temperatures that were used for the validation. In contrast, Fig. 15b, which was recorded at 150 °C, demonstrates excellent agreement. The deviation also appears to be random and not systemic at all. At different temperatures, very reasonable fitting was obtained as well. The model was thus considered validated, and is accurate within a range of approximately 10%. employed Consequently, it was to perform the aforementioned optimisation procedures.

4.6 Optimisation of a continuous process

The model provided insight into the optimum QAS production conditions, beyond the used PFR's limits. During the optimisation, the residence time was varied from 0.25 to 48 min (without a ramp), and the temperature ranged up to 300 °C. The resulting contour plots for all design specifications are presented in Fig. 16. The ratio between the reactants and MeOH was also varied. Lastly, the performance of a PFR was compared to a continuous stirred tank reactor (CSTR). The results have been summarised in Table 7.

The highest reactor productivity of 293 g h⁻¹ was computed at a molar ratio of 1:2.5:10 (DMDA:DMC:MeOH), 270 °C, and with a residence time of 0.25 min. Taking into account the reactor volume (2.4 mL), this translates to a specific productivity of 122 kg h⁻¹ L⁻¹. At these conditions a high conversion of 99.71%, and reasonable selectivity of 97.51% are reached. Conventional QAS production is carried out at approximately 120 °C, so this optimum poses quite a compelling increase in temperature. It demonstrates a clear opportunity for process intensification. It should be noted that the vapour pressure of MeOH is *ca.* 126 bar at this temperature, so an even higher pressure would be required. This is feasible in a PFR, especially since the high production rate allows for a small reactor volume.

Further temperature increase results in even higher required pressures, and lower selectivities. The reduced residence time this results in seems redundant and will certainly not outweigh the aforementioned drawbacks. For applications where a very high yield (over 99.9%) is desired the temperature has to be lowered to 150 °C. This does increase the reaction time to 48 min, however, and lowers the productivity to 1.5 g h⁻¹. Evidently, there can be an intermediate optimum, depending on the application.

Increasing the relative amount of DMC can also allow for a slight increase in selectivity. However, since this reduces the tertiary amine content of the reaction mixture, this leads to lower QAS production rates. The proposed optimum molar ratios could be further optimised by taking into account the technicals and economics of subsequent separation steps. Finally, Table 7 does allow for the comparison between production in a PFR *vs.* a CSTR. The results show that the PFR significantly outperforms the CSTR. This was expected, as the consecutive degradation reaction is accelerated here, due to the higher average QAS concentration.

5 Conclusions

Reaction kinetics of DMDA alkylation with DMC were successfully determined in various reactor systems. In batch, solvent effects were elucidated. MeOH was considered to be most eligible for industrial use, compared to acetonitrile, acetone, and solventless production. Isopropanol, ethanol, dimethyl acetamide, and dimethyl sulfoxide were found to be unsuitable because of side reactions.

A linear ramp-flow method was employed to measure the reaction kinetics in a PFR with in-line ¹H NMR. The Arrhenius parameters were computed (DMDA:DMC:MeOH = 1:5:10) at $E_a = 88.5 \pm 0.3$ kJ mol⁻¹ and $k_0 = 4.14 \pm 0.09 \times 10^7$

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L mol⁻¹ s⁻¹. The results were compared with the data in batch, and with conventional steady-state experiments. Consistent agreement was ascertained, so the method was considered valid. The ramp-flow method can allow for more efficient kinetic experiments. It saves time, material, and required labour costs compared to both batch, and steady-state techniques.

The PFR was modelled with a TIS model. The kinetic data from the ramp-flow method was used as input, along with degradation kinetic data. Using RTD studies and reactor characterisation, non-ideal plug-flow behavior was confirmed. Thereafter, the model was validated with data obtained in the PFR.

After validation, the model was employed to extrapolate the operating conditions from the experiments. This provided a platform to optimise the QAS production process, and productivity and yield were optimised. A maximum production rate of 122 kg $h^{-1} L^{-1}$ was predicted by the model at 270 °C, 0.25 min residence time, and a molar ratio of 1: 2.5:10. This shows considerable potential for the intensification of the process.

Conflicts of interest

There are no conflicts of interest to declare.

Note added after first publication

This article replaces the version published on 13th August 2021, which contained an error in the temperature in the Abstract.

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