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Continuous flow synthesis of cyclobutenes *via* lithium ynolates†

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Batch reactions that involve the generation of highly reactive species require a cryogenic temperature, complicated manipulations by chemists, and higher amounts of reagents, resulting in energy wastage and high costs. In this study, we developed a continuous flow synthesis of functionalised cyclobutenes, where the first step was the flash generation of short-lived lithium ynolates. Lithium ynolates were generated by the reaction of α,α,α -tribromomethyl ketones and *n*-butyllithium at 30 °C in 2.5 s and transferred to the next reactor before decomposition. The optimal reaction time (2.5 s) and temperature (30 °C) were determined *via* in-line Raman spectroscopy. The one-flow process involved three steps: the generation of lithium ynolates, the [2 + 2] cycloaddition reaction with α,β -unsaturated esters, and acetylation of the resulting unstable lithium enolates. These reactions were mediated by several reactive chemical species such as lithium ynolates, ketenes, and lithium enolates. Our green, flash flow approach to generating ynolate anions does not require cryogenic conditions and is highly reproducible and scalable, making it suitable for practical applications.

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Green foundation

1. We present a greener protocol to generate ynolate anions in flow. This protocol eliminates the need for cryogenic conditions and is highly reproducible and scalable. We also present a greener synthetic method of highly functionalized cyclobutenes. This three-step, one-flow process affords cyclobutenes without complicated manipulations by chemists, thus saving manpower.
2. In the optimised reaction conditions, we reduced the consumption of organolithium reagents by half. Moreover, the mild reaction temperature of 30 °C and an extremely short residence time of 23.1 s, which are applied by using the flow reactor system, contributed to energy savings.
3. In the future, this flow synthesis will be applied for the process synthesis of pharmaceutically important cyclobutenes.

Introduction

Flow reactions play a central role in modern synthetic chemistry and engineering.^{1,2} Flow microreactors, in particular, provide precise control over the reactivity of short-lived intermediates. In addition, continuous flow systems facilitate the

integration of several reactions involving highly reactive intermediates without isolation within a few minutes.^{3–6} This could be attributed to the advantageous mechanical features of flow microreactors. First, the continuous stream for a precise reaction time facilitates optimal mass transfer (Fig. 1a). Flow reactions can allow a reaction time of milliseconds, which is unachievable for batch systems. Second, their small dimensions permit precise temperature control and fast mixing, facilitating the desired reactions without side reactions (Fig. 1b). Consequently, continuous flow systems have empowered fast reactions *via* highly reactive chemical species such as organolithium, as demonstrated in the field of flash chemistry.⁷ In certain cases, flow reactions eliminate the necessity of cryogenic conditions, which are typically required in batch reactions. Furthermore, continuous flow synthesis ensures increased safety, high scalability, and

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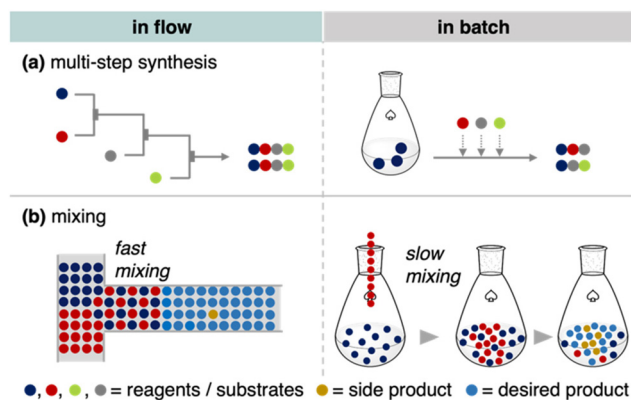


Fig. 1 Mechanical features in continuous flow and batch systems: (a) mass transfer in multi-step synthesis, (b) fast mixing in a flow system and slow mixing in a batch system.

reproducibility, broadening the synthetic applications of short-lived species.^{8,9}

Lithium ynolates are highly reactive species.¹⁰ Except for a few isolable ynolates, most of them have not been detected and quantified.^{11,12} Consequently, controlling their reactivity and employing them for organic synthesis remain a significant challenge.^{13–15} We previously developed a one-pot synthesis of cyclobutenes **4** via the generation of lithium ynolates. The synthesis involved three reactions: (1) the generation of lithium ynolates **2** from α,α,α -tribromomethyl ketones **1** and *tert*-butyl lithium (*t*-BuLi); (2) the [2 + 2] cycloaddition of lithium ynolates **2** with α,β -unsaturated esters via reactive ketene intermediates; (3) the acetylation of the resulting lithium enolates **3**

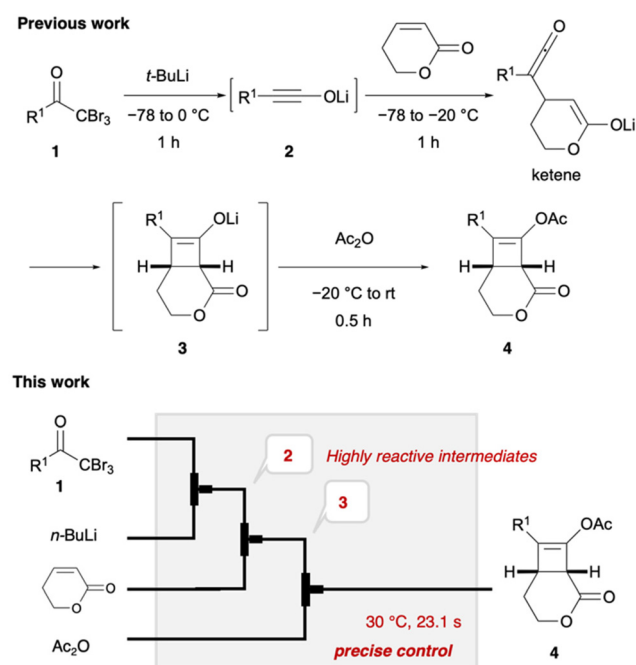


Fig. 2 Continuous synthesis of cyclobutenes in batch (previous work) and flow (this work) systems.

(Fig. 2).¹⁶ As alkoxide polymerises the α,β -unsaturated esters, the first step of the reaction was conducted under alkoxide-free conditions. This reaction features wide substrate scope of ynolates. However, this procedure has several drawbacks: cryogenic conditions, complicated operation, and temperature changes from -78 to 0 °C during every step. Additionally, precise temperature regulation is essential, as any deviation could lead to the rapid decomposition of two unstable intermediates, ynolates and cyclobutenes. This is a common problem in the generation and handling of lithium ynolates.^{17,18} Furthermore, excess consumption of *t*-BuLi at cryogenic temperatures hampers its application in the scalable and sustainable synthesis of cyclobutenes. Considering the advantages of flow chemistry and our previous success in flow synthesis, we envisioned that a continuous flow system would solve these drawbacks and improve the consumption of excess reagents and energy.¹⁹ Shindo's group previously utilised a microflow reactor to generate lithium ynolates from α,α -dibromo esters with precise control of the reaction temperature.^{20–22} However, the flash generation of lithium ynolates from α,α,α -tribromomethyl ketones has never been reported. Therefore, in this study, we developed an original flow protocol for the flash generation of lithium ynolates in a flow system and integrated it with the flow synthesis of cyclobutenes. Cyclobutenes are an important structural motif in small-molecule drug candidates. However, their green synthetic methods are limited.^{23,24} Our microflow synthesis is green in terms of safety, scalability, and productivity.

Results and discussion

Optimisation of reagents and solvents in batch for flow synthesis

We sought a less pyrophoric lithium reagent than *t*-BuLi for the lithium–halogen (Li–X) exchange of **1a**, as *t*-BuLi is highly pyrophoric and may produce bubbles due to the formation of volatile isobutene.^{9,25} We tested other organolithium reagents for the synthesis of **4aa** in batch (Table 1). Entry 1 lists the pre-

Table 1 Optimisation of organolithium reagents and solvents in batch for flow synthesis

Entry	Solvent	BuLi (equiv.)	Yield of 4aa (%)
1 ^a	Et ₂ O	<i>t</i> -BuLi (8)	70
2	THF	<i>t</i> -BuLi (8)	30
3	THF	<i>s</i> -BuLi (4)	13
4	THF	<i>n</i> -BuLi (4)	46

^a Reaction temperature for acetylation: -20 °C to rt.

viously optimised conditions.¹⁶ The use of Et₂O produces LiBr precipitate, which blocks the flow stream. Next, THF was used to prevent precipitation, which afforded **4aa** in 30% yield (entry 2). The use of *t*-BuLi or *s*-BuLi in THF decreased the yield of **4aa** due to side reactions of the organolithium reagents with THF (entries 2 and 3). In contrast, *n*-BuLi improved the yield of **4aa** to 46% (entry 4). Although this yield was lower than the previously reported yield of 70%, we considered it sufficient at this stage prior to further flow optimisation.

Monitoring the formation of lithium ynolates

The formation of lithium ynolates was monitored in real time in a flow system (Fig. 3a).^{26–29} The characteristic band of the lithium ynolates at $\sim 2100\text{ cm}^{-1}$ can be monitored using the ReactRaman instrument.¹¹ Hence, we integrated the ReactRaman instrument into the flow system (Fig. 3a).

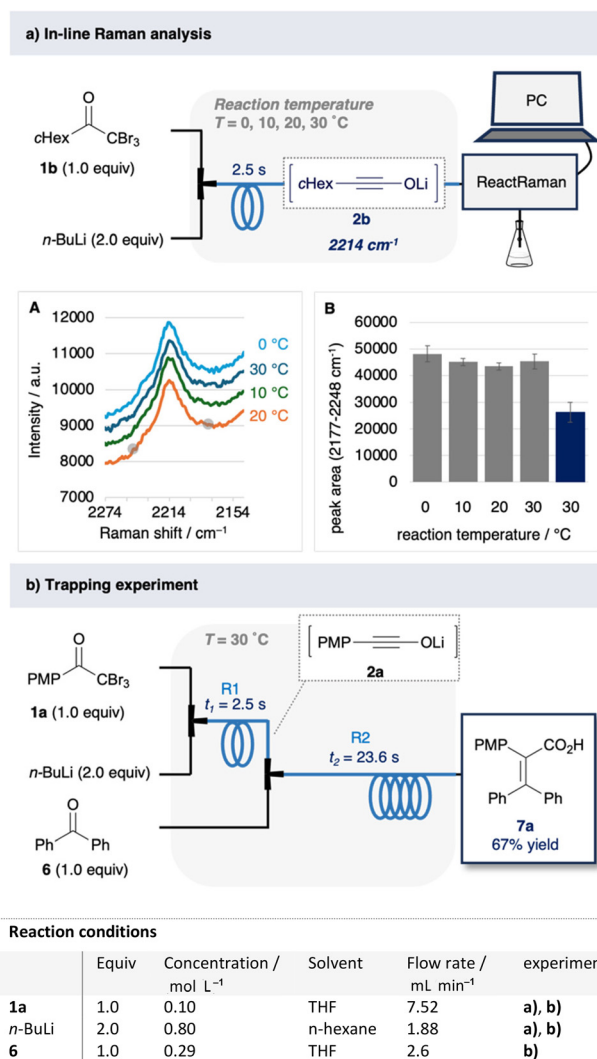


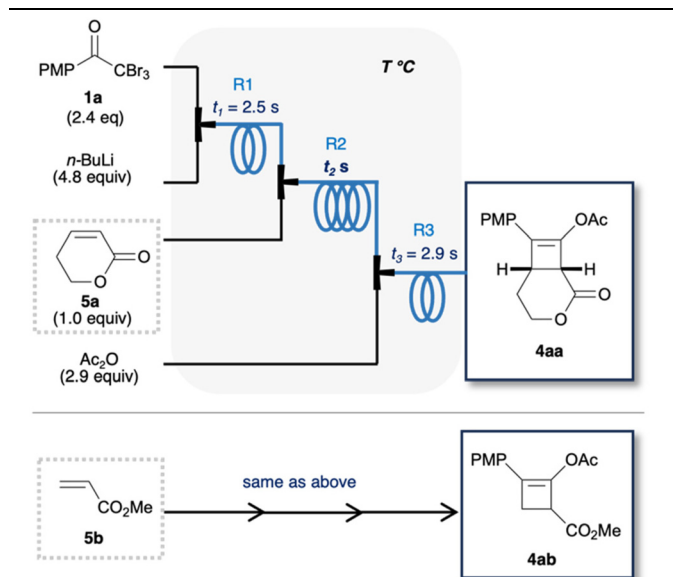
Fig. 3 (a) In-line Raman analysis of **2b**. A: Observed Raman spectra, residence time = 2.5 s, B: peak area ($2177\text{--}2248\text{ cm}^{-1}$), residence time = 2.5 s (grey bar), 22.5 s (blue bar). (b) Trapping experiment.

Lithium ynolates are formed *via* rapid reactions starting from double Li–X exchange.¹⁶ Therefore, to achieve rapid mixing, we employed a total flow rate of 9.4 mL min^{-1} and a mixer with an inner diameter of $250\text{ }\mu\text{m}$. The reaction mixture was monitored at $30\text{ }^{\circ}\text{C}$ after a residence time of 2.5 s. However, the monitoring experiment using **1a** was hampered by the saturation of the Raman spectra. Therefore, we used α,α,α -tribromomethyl cyclohexyl ketone (**1b**) as the substrate. Compound **1b** exhibits a characteristic Raman peak at 2214 cm^{-1} only when mixed with *n*-BuLi (Fig. S4 and Table S4†). This Raman band was identified as the alkynyl moiety of lithium ynoate **2b**, which is consistent with previously reported IR stretching frequencies [ESI 4.2†]. The Raman peak intensity remained unchanged in the temperature range of $0\text{--}30\text{ }^{\circ}\text{C}$ (Fig. 3a (A, B)). This indicates that a consistent amount of ynoate **2b** was produced in this temperature range. In contrast, the prolonged residence time of 22.5 s at $30\text{ }^{\circ}\text{C}$ reduced the Raman peak intensity (Fig. 3a and b, blue bar). Overall, these results indicate that a significant amount of ynoate was produced at room temperature at a mixing time of 2.5 s, and longer mixing times than 2.5 s may lead to ynoate decomposition. Furthermore, as a quantitative reference, we trapped the ynoate **2a** with a known electrophile (Fig. 3b). Ynoate **2a** reacts with benzophenone (**6**) to afford a stable alkene **7a**.^{20,30} We conducted this reaction in the flow system. The generated ynoate **2a** was trapped by **6** to afford alkene **7a** in 67% yield. That is, **2a** was obtained in 67% yield, at least in the first step.

Optimisation of the [2 + 2] cycloaddition of ynoate **2a** in flow synthesis

Next, we examined the continuous synthesis of cyclobutenes **4** (Table 2) in the flow system. This synthesis involves the formation of lithium ynoate **2a** from **1a**, the subsequent [2 + 2] cycloaddition with **5a** or **5b**, and acetylation. To optimise the conditions for the second step, the residence time (t_1) in the first reactor loop (R1) was fixed at 2.5 s. In the second reactor loop (R2), the residence time (t_2) range was set from 5.9 to 23.6 s, based on prior in-line Raman analysis that had demonstrated significant decomposition of **2b** at 22.5 s (*vide supra*). Thus, the various t_2 s in R2 and the various reaction temperatures (T) in all of the reactor loops were screened in the continuous process. Screening with **5a** demonstrated that a residence time between 17.7 and 23.6 s in R2 at $20\text{--}30\text{ }^{\circ}\text{C}$ afforded the best yield of **4aa** (74–75%) (Table 2a). Screening with **5b** showed that a residence time of 17.7 s in R2 at $30\text{ }^{\circ}\text{C}$ resulted in the highest yield of **4ab** (63%) (Table 2b). Besides, the lower the reaction temperature, the lower the yield of **4aa** or **4ab**.³¹ Hence, the [2 + 2] cycloaddition proceeds smoothly at approximately room temperature within the time scale. Moreover, this result demonstrates the potential of flash chemistry in preventing side reactions. Although the equivalents of ynolates and residence time (t_3) in the third reaction were re-examined, the initially set parameters afforded the best results (Tables S2 and S3†). Finally, a t_2 of 17.7 s in R2 and $30\text{ }^{\circ}\text{C}$ were selected as the optimised reaction conditions for generality.



Table 2 Screening of temperature and residence time (t_2) to synthesise cyclobutenes **4aa** and **4ab**. NMR yields of (a) **4aa** and (b) **4ab**

Reaction conditions: **1a** (2.4 equiv., $c_1 = 0.10 \text{ mol L}^{-1}$, $v_1 = 7.52 \text{ mL min}^{-1}$, THF); $n\text{-BuLi}$ (4.8 equiv., $c_2 = 0.80 \text{ mol L}^{-1}$, $v_2 = 1.88 \text{ mL min}^{-1}$, $n\text{-hexane}$); **5** (1.0 equiv., $c_3 = 0.12 \text{ mol L}^{-1}$, $v_3 = 2.6 \text{ mL min}^{-1}$, THF); Ac_2O (2.9 equiv., $c_4 = 0.23 \text{ mol L}^{-1}$, $v_4 = 4.0 \text{ mL min}^{-1}$, THF). $t_1 = 2.5 \text{ s}$, $t_3 = 2.9 \text{ s}$.

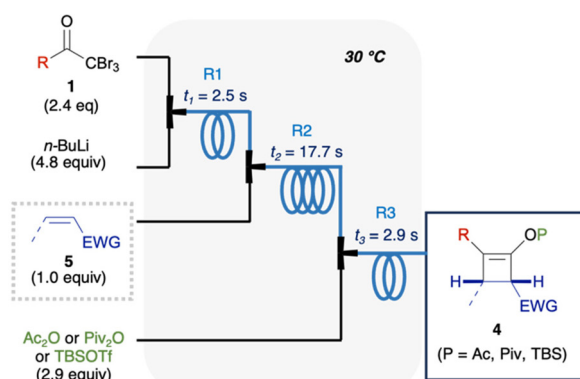
(a)		t_2 (s)			
Yield of 4aa (%)		5.9	11.8	17.7	23.6
T (°C)	30	72	71	74 (61)	74
	20	67	70	74	75
	10	64	67	68	N.D.

(b)		t_2 (s)			
Yield of 4ab (%)		5.9	11.8	17.7	23.6
T (°C)	30	43	56	63 (70)	62
	20	28	46	53	54
	10	22	30	41	47

Yields of **4** from the batch reaction are shown in the parentheses.

Substrate scope

The optimised reaction conditions were applied to the flow synthesis of 10 cyclobutenes (Fig. 4). Cyclohexyl (**1b**) and isobutyl (**1c**) ketones afforded **4ba** and **4ca** in 85% and 76% yields, respectively. Compared to the batch process, the flow process afforded higher yields of **4** with improved reproducibility. On the other hand, the use of **5b** afforded **4bb** and **4db** in lower yields (30–48%) than 64% yield in the batch process. It is considered that sterically less hindered **5b** involved undesired side reactions in this flow system. Similarly, the use of 2,5-dihydrofuranone afforded a low yield of **4ac** (27%). A plausible side reaction is considered to be the over-reaction of



Optimised reaction conditions

	Equiv	Concentration / mol L ⁻¹	Solvent	Flow rate / mL min ⁻¹
1	2.4	0.10	THF	7.52
$n\text{-BuLi}$	4.8	0.80	$n\text{-hexane}$	1.88
5	1.0	0.12	THF	2.6
Ac_2O or Piv_2O or TBSOTf	2.9	0.23	THF	4.0

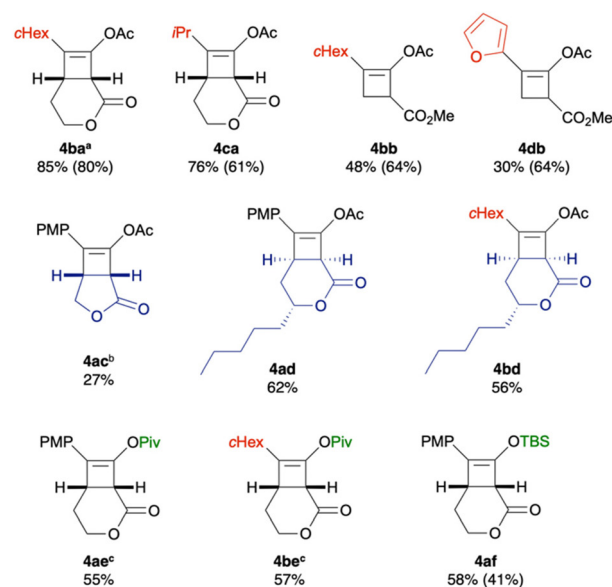
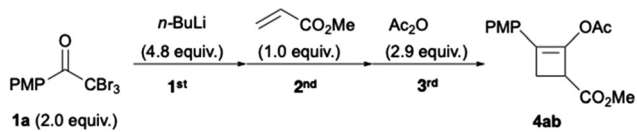


Fig. 4 Substrate scope. ^1H NMR yields of cyclobutenes **4** in flow synthesis, with batch yields in the parentheses. For the flow synthesis of **4ba**, **4ac**, **4ae**, and **4be**, few parameters of the optimised reaction conditions were changed as follows; ^a $T = 0^\circ\text{C}$, $t_3 = 1.5 \text{ s}$. ^b $T = 20^\circ\text{C}$. ^cConcentration of **5** = 0.10 mol L^{-1} .

ketene or enolate intermediates with ynolates. The other plausible side reaction is considered to be 1,2-addition³² of ynolates with **5**, because some side products from this reaction were observed in unsuccessful experiments (ESI, S10†). [2 + 2] cycloaddition with massoia lactone proceeded diastereoselectively to afford **4ad** and **4bd** in 62% and 56% yields, respectively. Relative stereochemistry of **4ad** was confirmed by X-ray crystallography. Trapping with pivalic anhydride or *t*-butyldimethylsilyl trifluoromethanesulfonate (TBSOTf) afforded pivalate ester **4ae** and **4be**, or silyl enol ether **4af**, respectively, in moderate yields (55–58%).

Table 3 Flow vs. batch synthesis of **4ab**

						
Reaction time/temperature						
Entry	Reactor	1st	2nd	3rd	Yield of 4ab (%)	STY (g h ⁻¹ L ⁻¹)
1	Batch ^a	1 h/−78 °C	1 h/−78 to −20 °C	0.5 h/−78 to −20 °C	70	0.98
2	Batch	60 s/30 °C	60 s/30 °C	60 s/30 °C	16	14
3	Flow	2.5 s/30 °C	17.7 s/30 °C	2.9 s/30 °C	63	7.2 × 10 ²

^a *t*-BuLi (8.0 equiv.) was used in a previous study instead of *n*-BuLi. Et₂O was used instead of THF.

Flow vs. batch synthesis and scale up

Finally, the flow conditions were compared with the corresponding batch conditions (Table 3). We used flow conditions (entry 3) for batch synthesis (entry 2), albeit with a slightly prolonged reaction time. Thus a low yield of **4ab** (16%) was afforded. When the batch synthesis was conducted at −78 °C, we observed a moderate yield (70%) of **4ab**, which is comparable to the flow synthesis yield (63%; entries 1 and 3). Unlike the batch systems that require cryogenic temperatures to stabilise the reactive and short-lived ynolates, the flow system did not need cryogenic temperatures. Furthermore, the space-time yields (STYs)^{33,34} of the flow system were over 50 times that of those in the batch system (Table 3, entries 1–3). This is attributed to the short residence time of 23.1 s over three steps in the flow system. This result is favourable for the scale up of this green synthesis. Hence, we used 3.12 mmol methyl acrylate (Fig. 5) as 0.12 M THF solution (26 mL), with a 0.1 M THF solution of **1a** (75.2 mL), 0.80 M *n*-hexane solution of *n*-BuLi (18.8 mL), and 0.23 M THF solution of acetic anhydride (40 mL). A flow reaction for 10 min afforded a 67% yield of **4aa** (607 mg) with a high STY of 8.1 × 10² g L⁻¹ h⁻¹. This demonstrates the utility of the flow protocol for the large-scale synthesis of highly functionalised cyclobutene **4aa**, which is impossible in a batch system. The advantages over the corresponding batch reaction, including the avoidance of cryogenic temperatures and reduction in reaction time, were effective for a straightforward and safe scale up. Moreover, the use of

n-BuLi reduced the consumption of organolithium reagents by half, which minimised wastage and lowered the synthetic cost. Our results should facilitate the development of a safe industrial process to synthesise cyclobutenes.

Conclusions

In this study, we achieved the continuous flow synthesis of cyclobutenes, starting from the flash generation of lithium ynolates. The synthesis is mediated by highly reactive species such as lithium ynolates, ketenes, and lithium enolates, which require complicated manipulations in batch. Nevertheless, we established a protocol that allows precise control of the reactivity without complicated temperature control. Non-cryogenic conditions shortened the total reaction time to 23.1 s, which prevented the decomposition of the reactive intermediates. First, we monitored the generation and degradation of lithium ynolates using in-line Raman analysis in the residence time range of 2.5–22.5 s at 30 °C to identify the best protocol for the continuous synthesis. The large-scale synthesis of cyclobutene **4aa** with the optimised conditions afforded a high STY (8.1 × 10² g L⁻¹ h⁻¹). Unlike batch protocols, the flow protocol uses a lower amount of safer reagents, is easy to scale up, conserves energy as it does not require cryogenic temperatures, and saves time and space. The mild reaction temperature of 30 °C (nearly room temperature) and an extremely short residence time of 23.1 s contributed to energy savings. Overall, this established flow protocol is highly sustainable. Notably, this facile three-step, one-flow process affords functionalised cyclobutenes without complicated manipulations by chemists, thus saving manpower. Our results improve the chemistry of lithium ynolates and enhance their utility in green chemistry by providing a simple and sustainable protocol.

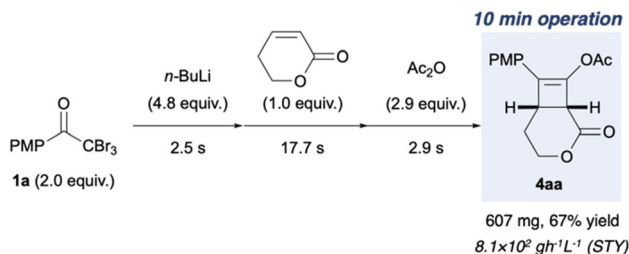


Fig. 5 Large-scale flow synthesis of **4aa**. The reactions were performed under the optimised reaction conditions shown in Fig. 4, and the reaction mixture was collected for 10 min.

Data availability

The data supporting this article have been included in the ESI.†



Conflicts of interest

There are no conflicts to declare.

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