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Continuous and convergent access to vicinyl amino alcohols†

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Five active pharmaceutical ingredients (APIs) containing the vicinyl amino alcohol moiety were synthesized using a convergent chemical assembly system. The continuous system is composed of four flow reaction modules: biphasic oxidation, Corey–Chaykovsky epoxidation, phenol alkylation, and epoxide aminolysis. Judicious choice of reagents and module order allowed for two classes of β -amino alcohols, aryl and aryloxy, to be synthesized in good (27–69%) overall yields.

The chemical synthesis of active pharmaceutical ingredients (APIs) is generally accomplished *via* a linear batch approach where a single process produces a single compound. Flow chemistry is a conceptual advance that allows for greater control over reaction conditions¹ and the ability to combine several sequential flow reactors to achieve multi-step processes.^{2–4} Chemical assembly systems (CAS)⁵ are based on telescoping sequential robust reaction units that are linked together in a non-iterative fashion. Multi-step syntheses of a wide range of compounds of similar⁶ or unique⁵ structural cores can be accessed in a divergent fashion (Fig. 1). Here, we report on the application of CAS to convergent syntheses, where two structural variations of a key epoxide intermediate can be trapped in an aminolysis reaction to produce two distinct classes of β -amino alcohols, including five APIs.

A variety of biologically active natural products, chiral auxiliaries, ligands, and APIs for the treatment of hypertension or as bronchodilators contain β -amino alcohols as an important structural core.^{7–9} Synthetic approaches to β -amino alcohols abound,¹⁰ including amino acid reduction,^{8a} coupling reactions, α -functionalization,¹¹ amino-hydroxylation, and epoxide ring opening.¹² Two classes of pharmaceutically relevant compounds, aryl and aryloxy vicinyl amino

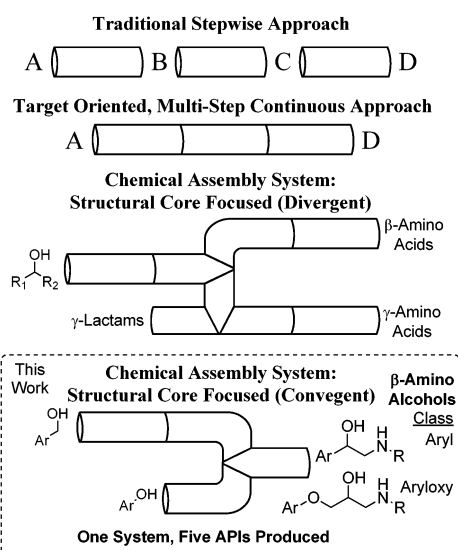


Fig. 1 Comparison of concepts for the chemical synthesis of small molecules.

alcohols, should be accessible *via* convergent epoxidations followed by a final ring opening with an amine (Fig. 2).

Aryl epoxides should be accessible *via* a one-to-two step oxidation/Corey–Chaykovsky process. Employing robust and selective biphasic alcohol-oxidation conditions we developed earlier,⁵ the first transformation is readily achieved and thus a continuous Corey–Chaykovsky epoxidation needed to be developed. Using benzaldehyde as a model system, a 0.5 M toluene solution was

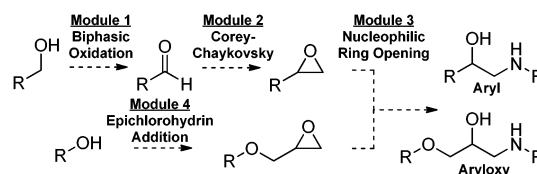


Fig. 2 Convergent synthesis of aryl and aryloxy vicinyl amino alcohols *via* a concluding epoxide ring opening.

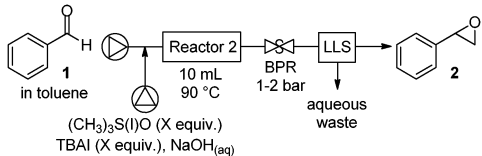
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† Electronic supplementary information (ESI) available: Characterization data, full experimental procedures, copies of ¹H and ¹³C NMR spectra of all new compounds. See DOI: 10.1039/c5cc06093a

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Table 1 Optimization of Corey–Chaykovsky reaction (module 2) in flow^a


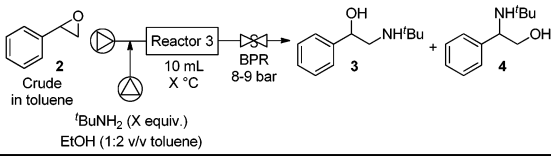
Entry	Me ₃ SOI ^b	NaOH ^b	TBAI ^b	Residence time (min)	Conversion of 1 (%)
1	1.4	1.4	0.1	24	95
2	1.7	1.7	0.1	24	97
3	2.0	2.0	0.1	24	96
4	1.7	1.7	0.1	40	96
5	1.7	1.7	0.1	10	88
6 ^c	1.7	1.7	0.1	24	85
7	1.7	1.7	0.0	24	79

^a Pump A: benzaldehyde (**1**) in toluene (0.5 M), flow rate: 0.083 mL min^{−1}. Pump B: (CH₃)₃S(I)O (1.7 equiv., 0.21 M), NaOH (1.7 equiv.), TBAI (0.1 equiv.) in water, flow rate: 0.33 mL min^{−1}, reactor size: 10 mL, pressure: 1–2 bar. ^b Equivalents with respect to benzaldehyde. ^c Reaction run at 60 °C. BPR: back pressure regulator, LLS: liquid/liquid separator. For full experimental details, see ESI.

mixed with an aqueous solution of trimethylsulfonium iodide ((CH₃)₃S(I)O), sodium hydroxide (NaOH), and a substoichiometric amount of phase transfer catalyst (tetrabutylammonium iodide (TBAI)). The resulting biphasic solution was passed through a 10 mL reactor held at 90 °C at a pressure of 2 bar. Clean conversions to styrene oxide were observed with (CH₃)₃S(I)O and NaOH in excess (1.4–2 equiv., Table 1 entries 1–3) after 24 minutes. Shorter reaction times, temperature, and the absence of TBAI resulted in decreased conversion (entries 5–7).

Using optimized conditions (97%, entry 2), we next turned to the amine ring-opening. While epoxide opening traditionally employed Lewis acids (LA), the elevated pressures and temperatures during microwave¹³ and flow¹² transformations allow for clean reactions without LA catalysts. We aimed to combine epoxidation and ring opening by modifying established flow conditions.¹² Thus, the crude toluene solution containing styrene oxide was mixed with ^tBuNH₂ in toluene/ethanol (2:1). With three equivalents of amine, good conversion to the desired regioisomer was observed after 50 minutes at 150 °C (Table 2, entry 1). Increasing the amount of amine to seven equivalents resulted in the highest regioselectivity at essentially full conversion (entry 3). Decreasing both the temperature and time resulted in lower conversions (entries 4 and 5). No product resulting from the bis-alkylation of the amine was observed under these conditions.

Modules 2 and 3 were connected with the help of a modified Jensen liquid–liquid separator,¹⁴ allowing for the toluene phase exiting the epoxidation to be used directly in the ring opening reaction. Thus, after a total residence time of 74 minutes, benzaldehyde was transformed into 2-(*tert*-butylamino)-1-phenylethanol **3** in 57% yield following off-line crystallization. A range of haloarenes reacted with isopropyl or *tert*-butyl amine without intermediate purification using the modular set-up. Moderate to good yields of the desired regioisomer were observed following product crystallization (Fig. 3). Tulobuterol, a β₂-adrenergic agonist used to treat asthma,¹⁵ was targeted. After passing 2-chloro benzyl alcohol through modules 1, 2, and

Table 2 Optimization of aminolysis (module 3) in flow^a


Entry	^t BuNH ₂ (equiv.)	Residence time (min)	Temp. (°C)	Ratio ^b 3 : 4 : 2
1	3	50	150	68 : 18 : 14
2	5	50	150	76 : 13 : 11
3	7	50	150	83 : 13 : 4
4	5	50	130	64 : 13 : 23
5	5	20	150	62 : 12 : 26

^a Crude styrene oxide (**2**, 0.33 M in toluene), ^tBuNH₂ in toluene/ethanol (1:0.5), flow rates each 0.2 mL min^{−1}. For full experimental details, see ESI. ^b As determined by ¹H NMR. BPR: back pressure regulator.

3 – with separators between modules 1/2 and 2/3 – the API was obtained in 27% yield (Fig. 3).

A number of hypertension medications such as propranolol, alprenolol, and metoprolol¹⁶ contain the aryloxy β-amino alcohol motif. These targets were selected to expand the scope of the modular flow synthesis of vicinyl amino alcohols. Typical phenol alkylations require a large excess of epichlorohydrin equivalents¹⁷ in the presence of an inorganic base.¹⁸ However, homogeneous solutions would facilitate the development of a flow process. A biphasic solution, obtained by mixing aqueous sodium hydroxide solution with a phenol in neat epichlorohydrin, should provide the desired aryloxy epoxide (Table 3).

The sodium hydroxide solution (1.95 M, 1.3 equiv.) is combined with phenol in three equivalents of neat epichlorohydrin *via* a T-mixer and passed through a reactor (1 h reaction time at 35 °C). Two major alkylation products (**10**, **11**) were observed in roughly a 2:1 ratio, with the desired epoxide **10** obtained in 56% (Table 3, entry 1). Addition of substoichiometric amounts of PTC resulted in an increased conversion of the intermediate halohydrin (**11**) into the desired product (entry 2). At a reaction temperature of 45 °C, **10** was obtained in 86% yield while no intermediate halohydrin was observed (entry 3). At higher temperatures, a diol was formed by sodium hydroxide mediated ring opening of the epoxide.¹⁹ Reducing the excess of epichlorohydrin to two equivalents (entry 5, final conditions) had no effect, while a further reduction of the reagent to one equivalent yielded sizable amounts

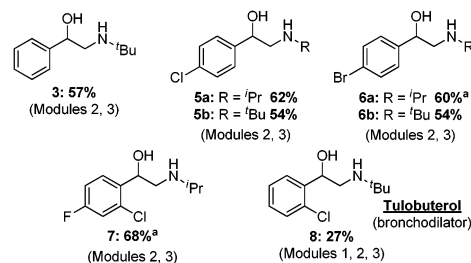
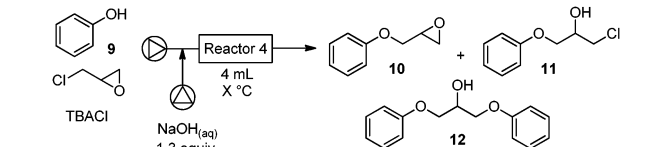


Fig. 3 Aryl vicinyl amino alcohols produced by continuous synthesis. ^a Residence time of Corey–Chaykovsky reaction is 40 min. See ESI† for full experimental details.



Table 3 Optimization of the phenol alkylation in flow^a


Entry	Epichlorohydrin (equiv.)	Temp. (°C)	TBACl (equiv.)	Residence time (min)	Yield (%)		
					10	11	12
1	3	35	—	60	56	24	—
2	3	35	0.1	60	68	15	—
3	3	45	0.1	60	86	—	—
4	3	45	0.1	46	80	8	—
5	2	45	0.1	60	85	—	—
6	1	45	0.1	60	65	—	21

^a Pump A: phenol, TBACl and epichlorohydrin, pump B: 1.95 M NaOH (1.3 equiv.), reactor size: 4 mL. For full experimental details, see ESI.

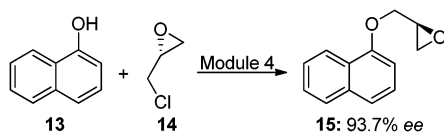
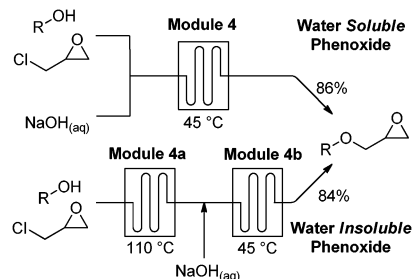
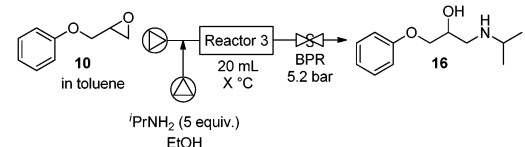
of dimer **12** (entry 6). Shortening the residence time in the reactor to 46 minutes resulted in incomplete conversion (entry 4).

Using this method, chiral aryloxy epoxides can be generated efficiently. When enantiopure epichlorohydrin is reacted using the conditions described above, the reaction proceeds with 93.7% ee (Fig. 4).

Module 4 depends on the solubility of the intermediate aryloxy in the biphasic solution. In cases where a precipitate forms, as occurred during the synthesis of metoprolol **20**, module 4 can split into a two-stage alkylation.^{20–22} Phenol was efficiently alkylated at 110 °C after 60 minutes when epichlorohydrin served as solvent (3.1 M, 2 equiv.). By mixing the resulting solution with aqueous NaOH at 45 °C, 84% of the desired epoxide could be obtained after only 30 minutes (Fig. 5).¹⁹

With a reliable process for the procurement of epoxide **10** in hand, the aminolysis module was connected (Table 2). Alkyl epoxides proved more active than aryl epoxides such that the ring opening proceeded at lower temperatures and shorter reaction times (Table 4). The epoxide solution was converted in just 20 min when mixed with five equivalents of isopropylamine in ethanol at 120 °C (entry 4). The reaction proceeds efficiently at 100 °C, but requires longer residence times (entry 2). Only one regioisomer and no amine dialkylation product were observed.

Finally, modules 3 and 4 were combined to yield the final sequence. The biphasic solution exiting module 4 is mixed with toluene. The toluene phase is removed using a membrane-based separator¹⁴ and directly fed into module 3. Amino alcohol **16** was obtained in 48% yield from phenol, following off-line crystallization (Fig. 6). Several active pharmaceutical ingredients were prepared based on a judicious choice of starting materials. Racemic and chiral aryloxy β-amino alcohols can be prepared

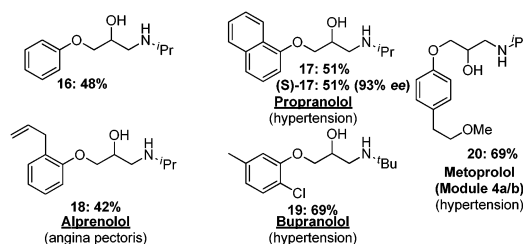
**Fig. 4** Synthesis of (S)-2-((naphthalen-1-yloxy)methyl)oxirane using enantiopure (R)-epichlorohydrin.**Fig. 5** To accommodate water-soluble and insoluble phenoxides, two variants of module 4, were implemented.**Table 4** Optimization of aminolysis in flow^a


Entry	T (°C)	Residence time (min)	Conversion (%)
1	80	30	78
2	100	30	100
3	100	20	85
4	120	20	100

^a Pump A: epoxide in toluene, pump B: isopropylamine (5 equiv.) in EtOH, reactor size: 20 mL, BPR: 5.2 bar. For full experimental details, see ESI. BPR: back pressure regulator.

including propranolol (**17**, 51%),²³ used to treat hypertension,²⁴ alprenolol (**18**, 42%), an angina pectoris medication,²⁵ and the hypertension drugs bupranolol (**19**, 69%),²⁶ as well as metoprolol (**20**, 69%).

Described is the development of a convergent chemical assembly system consisting of four modules that reliably produces aryl and aryloxy β-amino alcohols in good yields. Modular flow units include biphasic oxidations,⁵ Corey–Chaykovsky epoxidations, aryloxy alkylations, and epoxide aminolysis.¹² Aqueous in-line workup using liquid–liquid extractors resulted in continuous processes without intermediate purification. Starting from phenols or benzyl alcohols five active pharmaceutical ingredients for the treatment of hypertension, agina pectoris, and bronchodilation were produced. Enantiopure aryloxy β-amino alcohols can be prepared by starting from R/S epichlorohydrin.

**Fig. 6** Overview over aryloxy vicinyl amino alcohols produced by continuous synthesis.

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