

Cite this: *Chem. Sci.*, 2024, 15, 7092

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## Constitutional adaptation to $pK_a$ modulation by remote ester hydrolysis†

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The mechanisms through which environmental conditions affect the expression of interconnected species is a key step to comprehending the principles underlying complex chemical processes. In Nature, chemical modifications triggered by the environment have a major impact on the structure and function of biomolecules and regulate different reaction pathways. Yet, minimalistic artificial systems implementing related adaptation behaviours remain barely explored. The hydrolysis of amino acid methyl esters to their corresponding amino acids leads to a drastic change in  $pK_a$  (ca. 7 and 9, respectively) that protonates the free amino group at physiological conditions. Dynamic covalent libraries (DCvLs) based on amino acid methyl esters and aldehydes respond to such hydrolysis and lead to constitutional adaptation. Each of the libraries studied experiences a DCvL conversion allowing for constituent selection due to the silencing of the zwitterionic amino acids towards imine formation. The selective action of different enzymes on the DCvLs results in states with simplified constitutional distributions and transient chirality. When additional components (*i.e.*, scavengers) that are not affected by hydrolysis are introduced into the dynamic libraries, the amino acid methyl ester hydrolysis induces the up-regulation of the constituents made of these scavenging components. In these systems, the constituent distribution is resolved from a scrambled mixture of imines to a state characterized by the predominance of a single aldimine. Remarkably, although the final libraries display higher “simplicity”, the different transient states present an increased complexity that allows for the emergence of organized structures [micelle formation] and distributions [up-regulation of two antagonistic constituents]. This reactive site inhibition by a remote chemical modification resembles the scenario found in some enzymes for the regulation of their activity through proximal post-translational modifications.

Received 23rd February 2024  
Accepted 9th April 2024

DOI: 10.1039/d4sc01288g

rsc.li/chemical-science

## Introduction

Cross-talks between amino acid residues located at different positions in proteins control essential processes like molecular recognition and catalysis.<sup>1</sup> In addition, selective post-translational modifications (PTMs) on amino acid sidechains offer a powerful biological toolkit to expand the polypeptide functional repertoire.<sup>2–4</sup> These chemical modifications, occurring in well-defined microenvironments, commonly lead to large perturbations in the  $pK_a$  of surrounding ionizable groups that govern the function of biomolecules.<sup>5</sup> For instance, the phosphorylation of a serine residue induces changes in the  $pK_a$  of a nearby histidine sidechain and modulate the reactivity of cofilin proteins (Fig. 1A).<sup>6</sup> Therefore, chemical systems that can adapt to analogous remote transformations deserve to be

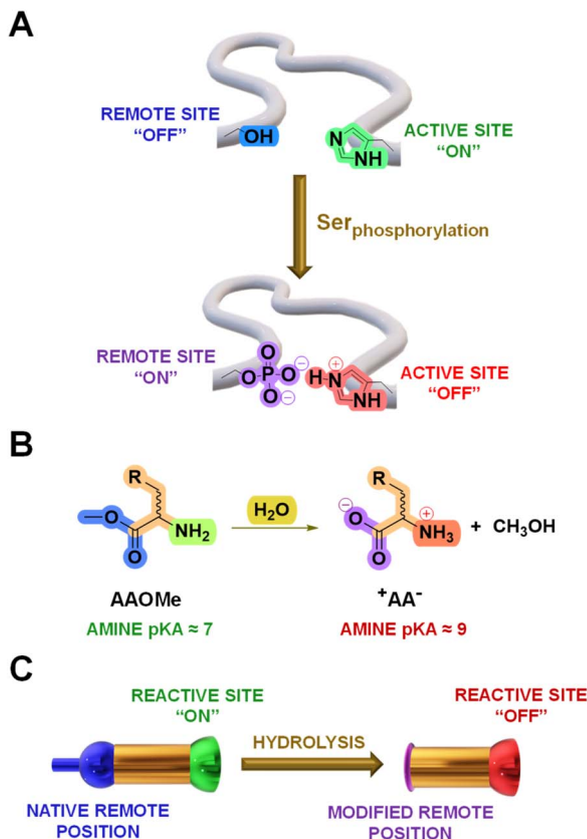
thoroughly explored as they might enlighten the mode-of-action underlying such biological processes.

An extensive amount of studies discussing adaptive systems rely on Constitutional Dynamic Chemistry (CDC) at either the molecular (Dynamic Covalent Chemistry, DCvC) or supramolecular level (Dynamic Non-Covalent Chemistry, DNCvC).<sup>7</sup> In DCvC, the reversible covalent bonds involved allow for error-checking and rearrangement processes under thermodynamic control.<sup>8–11</sup> Hence, the distribution of diversified Dynamic Covalent Libraries (DCvLs) can be directed to a certain state upon exposure to external stimuli or chemical effectors.<sup>12,13</sup> For example, the sorting of “messy” DCvLs has been attained using both physical (precipitation,<sup>14</sup> distillation,<sup>15</sup> adsorption<sup>16</sup>) and chemical (oxidation,<sup>17</sup> templates,<sup>18</sup> catalysis,<sup>19</sup> coupled enzymatic reactions<sup>20</sup>) methods. In a similar manner, Constitutional Dynamic Networks (CDNs) are composed of products (*i.e.*, constituents) constitutionally linked by exchangeable building blocks (*i.e.*, components) that adapt their distribution through agonistic and antagonistic relationships.<sup>21,22</sup> Substantial research efforts have been devoted to build-up relatively simple CDNs mimicking complex biological behaviours.<sup>23–25</sup> Related dynamic networks mediate the communication between

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† Electronic supplementary information (ESI) available: Experimental section, main text support figures and tables, copies of <sup>1</sup>H, <sup>13</sup>C NMR, and HRMS spectra. See DOI: <https://doi.org/10.1039/d4sc01288g>





**Fig. 1** (A) Representation of a decrease in activity at the catalytic site of a protein by a PTM (serine phosphorylation) accompanied by a change in the  $pK_a$  of a nearby residue (histidine protonation). (B) Hydrolysis of AAOMe (terminal  $-NH_2$  site reactive in imination) to yield  $^+AA^-$  (terminal  $-N^+H_3$  site silent in imination) and a molecule of  $CH_3OH$ . (C) Representation of the proposed remote modification leading to the inhibition of the components of DCvLs (colour code has been maintained for comparison).

biomolecules (e.g., DNA, RNA, proteins), leading to programmed reaction patterns that regulate key intracellular processes like signal transduction and enzymatic activity.<sup>26</sup> However, the design of DCvLs, and their underlying CDNs, able to undergo structural and/or constitutional adaptation in response to  $pK_a$  variations triggered by remote chemical modifications remains unexplored.

The hydrolysis of amino acid methyl esters (AAOMe) to their corresponding zwitterionic amino acids ( $^+AA^-$ ) leads to a drastic  $pK_a$  change in the remote  $-NH_2$  group, resulting in its protonation at neutral pH (predicted  $N_\alpha$   $pK_a$  values of ca. 9 and 7, respectively; Fig. 1B and Table S1†).<sup>27</sup> Thus, we envisaged that DCvLs based on the condensation of AAOMe and water-soluble aldehydes should experience constitutional adaptation in response to the hydrolysis of the remote ester group since the resulting zwitterionic  $^+AA^-$  entity should be “poisoned” towards imine formation through reactive site protonation (Fig. 1C). After first devising the most reactive components towards imination, we optimized the conditions for the hydrolysis of the AAOMe ester group. Then, the time evolution of the DCvLs was investigated, observing that they resulted in states of higher “simplicity” (simplification of the

output of the system)<sup>28</sup> through transient states of higher complexity [enzymatic selection/transient chirality/sorting] and diversity [increased number of components/constituents]. Furthermore, such constituent selection induced emerging behaviours in the final simplified states of the libraries: micelle formation and antagonistic sorting (see below).

## Results and discussion

### Component screening

The first step was to select the reaction components by screening a series of water-soluble reactive aldehydes and different amines. Imines are generally disfavoured in water due to their propensity to hydrolyse back into the reactants.<sup>29</sup> Nevertheless, a rational design of the system (increased aldehyde electrophilicity and amine nucleophilicity) can assist to overcome this issue and foster relatively high yields.<sup>30</sup>

The imine yields for the reaction between different aldehydes, namely **A1–6**, and **ArgOMe** were evaluated at room temperature and  $pD = 7.0$  (PBS 50 mM).  $^1H$  NMR signal integration after 1 h of equilibration showed that the highest imine formation was observed when using **A1**, likely as a result of the strong electron-withdrawing capacity of the two sulfonate groups (Fig. 2A).

In terms of the amine screening, the studied  $\alpha$ -AAOMe gave imine yields of ca. 30% (Fig. 2B). In contrast, the yield attained when using  $\beta$ -AlaOMe as a nucleophile was 9%, in good agreement with the higher  $pK_a$  calculated for this derivative (Table S1†). Several commercially available amines were also assayed to select a nucleophile with similar reactivity to that of the  $\alpha$ -AAOMe. The most reactive amine was found to be **B1** which gave a remarkable imine yield of 67%, assigned to its low degree of protonation arising from the presence of a positive charge in close proximity.<sup>31</sup> A similar scenario was found in the case of **B2**, but with a lower imine formation (25% abundance). The amines presenting  $pK_a$  values between 8.7–9.3 (viz., **B3**, **B5**, **B6**, and **B7**) resulted in imine yields of ca. 15%. As predicted, amine **B4** ( $pK_a = 10.2$ ) did not react with aldehyde **A1** as its amino group is completely protonated at  $pD = 7.0$ . To increase the imine yields, the reaction between **A1** and **ArgOMe** was performed at higher concentrations maintaining a 1 : 1 ratio between components. The reaction at 80 mM gave 65% of imine **A1ArgOMe** (Fig. S1†). These results indicated that the low specificity of imine formation in water can be overcome at high concentrations and with adequate substrate selection, paving the way for the study of dynamic covalent reactions and libraries (see sections below).

### Hydrolysis of AAOMe

Next, we decided to investigate the ester hydrolysis of AAOMe. It is well established that AAOMe are hydrolysed to the corresponding amino acids ( $^+AA^-$ ) under basic conditions.<sup>32</sup> However, the pH must be optimized to attain the best conditions in terms of hydrolysis rates as well as imination efficiency and selection. Although higher pH values would favour imine formation and ester hydrolysis, undesired aldimines with  $^+AA^-$  would also be formed at high pH (Table S2†). Thereby, three



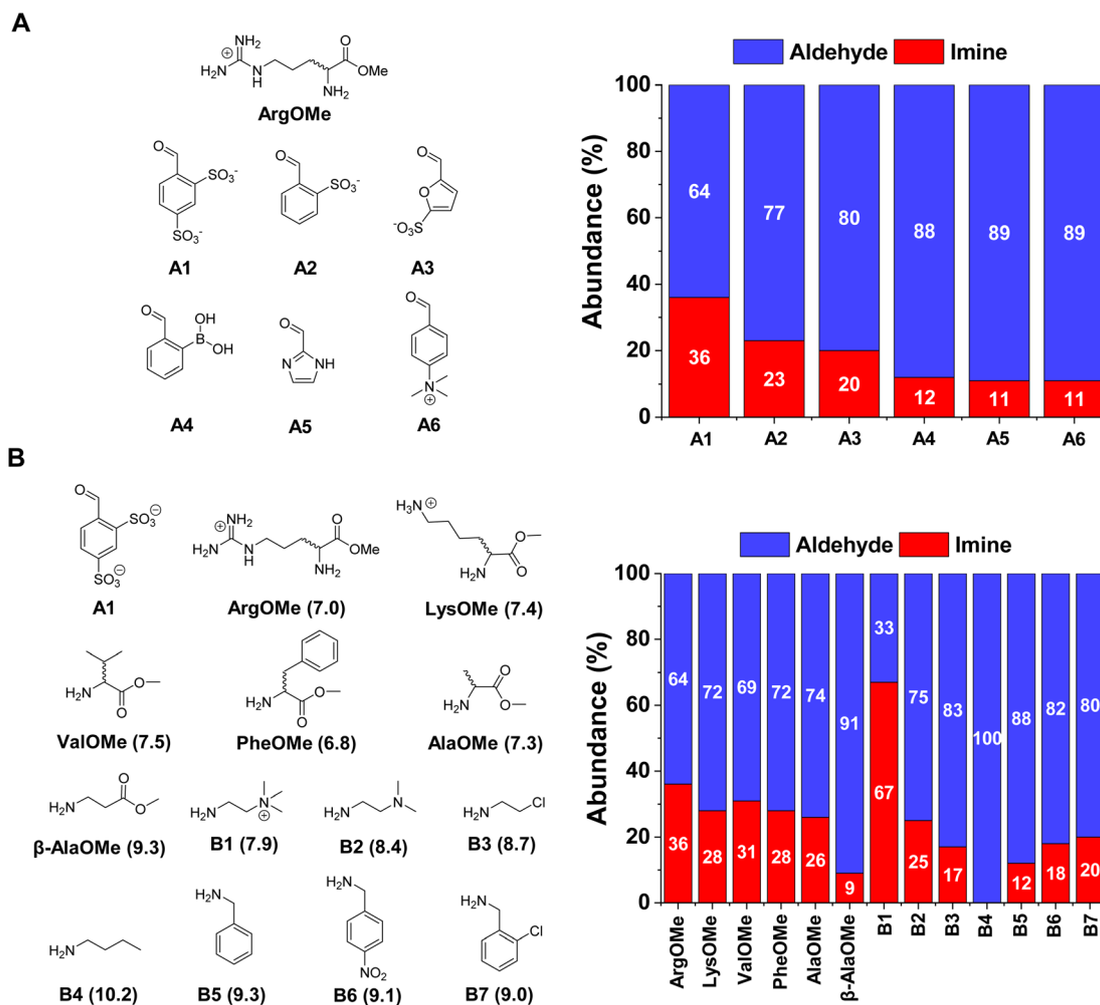


Fig. 2 (A) Aldehyde screening using ArgOMe as the model amine. (B) Amine screening using best-performing A1 as aldehyde. Left: Chemical structures of the different reagents employed. Right: Component abundances obtained after equilibration (1 h at room temperature). The hydrate detected was included in the aldehyde composition. Amine  $pK_a$  values (calculated using MarvinSketch) have been included in brackets. All abundances were calculated using  $^1H$  NMR spectroscopy (500 MHz,  $D_2O$ ,  $pD = 7.0$ , PBS buffer 50 mM, 293 K). Initial concentration of reagents 5 mM each in all cases. Error in abundance: 5%.

experiments were run at  $pD$  values ranging from 6.2–7.8 to reveal the effect on the aforementioned variables.<sup>33</sup> The reaction course could be easily monitored by  $^1H$  NMR spectroscopy, observing the disappearance of the signals assigned to ArgOMe and the appearance of those of  $^+Arg^-$  and methanol (Fig. S2<sup>†</sup>). As expected, the fastest hydrolysis rates were obtained under basic conditions, with the reaction at  $pD = 7.8$  being slightly favoured over the one at  $pD = 7.0$  (Fig. S3 and Table S3<sup>†</sup>). Notwithstanding, the  $-N_{\alpha/\beta}H_2$  degree of protonation calculated revealed that, at  $pD = 7.8$ , the  $^+Arg^-$  inhibition towards imination would not be complete, as a significant amount of the reactive site would be in its nucleophilic non-protonated form (10%; see Table S2<sup>†</sup>). Separated reaction of ArgOMe and  $^+Arg^-$  with A1 at these different  $pD$  values corroborated this assumption (Table S4<sup>†</sup>). Although the highest imine yields were obtained at  $pD = 7.8$  (44%), the undesired reaction between  $^+Arg^-$  and A1 was also detected (3% yield of  $A1^+Arg^-$ ). On the other hand, slightly lower  $A1ArgOMe$  yields (36%) were noted at

$pD = 7.0$ , but with complete silencing of  $A1^+Arg^-$  (entry 2, Table S4<sup>†</sup>). Therefore, the optimal  $pD$  was found to be 7.0.

We then evaluated the effect of the substrate structure on the hydrolysis rate, both in terms of the sidechain nature and the distance between the carbonyl unit and the amino group (*i.e.*,  $\alpha$ - or  $\beta$ -AAOMe). Results suggest that  $\beta$ -AAOMe hydrolyses at slower rates than the  $\alpha$ -derivatives. For instance, the hydrolysis of  $\beta$ -AlaOMe was much slower than that of AlaOMe, as shown by the yields of 16 and 44% observed after 48 h, respectively (Fig. S4 and Table S5<sup>†</sup>). In terms of the sidechain effect, the  $\alpha$ -AAOMe presenting the highest hydrophathy,<sup>34</sup> namely ValOMe, was barely hydrolysed after 48 h ( $\approx 20\%$   $CH_3OH$  yield; Fig. S4<sup>†</sup>). This could be related to a less efficient attack of water molecules on the ester group because of the hydrophobic environment. LysOMe presented the fastest hydrolysis rate, and the other  $\alpha$ -AAOMe (*viz.* ArgOMe and PheOMe) gave similar profiles (Table S5<sup>†</sup>). Thus, LysOMe was selected for the subsequent experiments (see sections below).



Previous reports proved that the hydrolysis of esters in water could be catalysed by imidazole-derivatives and by enzymes with esterase activity.<sup>35,36</sup> Acetylcholine esterase (**AChE**) is a conserved enzyme that presents superior activity in the hydrolysis of acetylcholine to choline (Fig. S5A†). The active site is composed of two well-defined functionalities: the esteratic site (containing a serine residue which cleaves the ester bonds) and the peripheral site (anionic groups that fix the substrate in the optimal position through supramolecular interactions).<sup>37</sup> We envisaged that **AChE** could selectively catalyse the hydrolysis of the **AAOMe** that contained a positively charged unit in the sidechain, namely **ArgOMe** and **LysOMe**, as they present a relatively similar chemical structure to that of acetylcholine (Fig. S5B†). The model hydrolysis of **ArgOMe** was then studied in the presence of imidazole and **AChE**. Whereas the imidazole-based approach was not successful even at high base concentrations (200 mol%), up to a 3.2-fold increase in hydrolysis rate could be observed with increasing catalytic amounts of **AChE** (Fig. S6 and Table S6†). As expected, the catalytic activity of **AChE** was notably dependent on the structure of the **AAOMe**, since it only accelerated the hydrolysis of **ArgOMe** and **LysOMe** (Fig. S7 and Table S7†). In view of these results, two additional **AAOMe** hydrolases were tested: **trypsin** and **chymotrypsin**. **Trypsin** was found to selectively catalyse the depletion of **LysOMe**, in a similar manner to that of **AChE**, but with faster rates (Fig. S8 and Table S8†). In contrast, **chymotrypsin** showed excellent selectivity for the catalytic hydrolysis of **PheOMe** (Fig. S8 and Table S8†).

Considering the chiral nature of the enzymatic active sites, we also surmised that **AChE** could preferentially catalyse the hydrolysis of one of the enantiomers of **LysOMe**. Indeed, the rate of hydrolysis of (*D*)**LysOMe** was not significantly affected by the enzyme (Fig. S9 and Table S9†). On the other hand, the use of 0.01 mol% of **AChE** induced complete hydrolysis of (*L*)**LysOMe** in less than 16 h.

Therefore, **AChE**, **trypsin** and **chymotrypsin** are suitable catalysts for the selective kinetic simplification of complex mixtures of **AAOMe** (*vide infra* DCvL1), as well as for challenging enantioselective kinetic resolutions of **AAOMe** racemic mixtures (*vide infra* DCvL3).

### Constitutional adaptation to AAOMe hydrolysis in a dynamic covalent reaction

Several reports show how peptide-based supramolecular systems are able to assemble/disassemble in response to ester/anhydride hydrolysis, as a result of solubility differences between the generated carboxylate groups and the parent units.<sup>38–41</sup> However, the change in the reactivity of vicinal functionalities in response to such ester hydrolysis (*e.g.*, potential decrease in the nucleophilicity of surrounding groups by supramolecular cross-talks, see biological example in Fig. 1A) has been quite overlooked, as well as its implementation in dynamic covalent libraries.

Once the reaction conditions were optimized (see sections above), the effect of **AAOMe** hydrolysis on the imine yields was monitored for the reaction between **AAOMe** and **A1**. Considering that the  $-N_{\alpha/\beta}H_2$  group of the hydrolysis product  $^+AA^-$  is

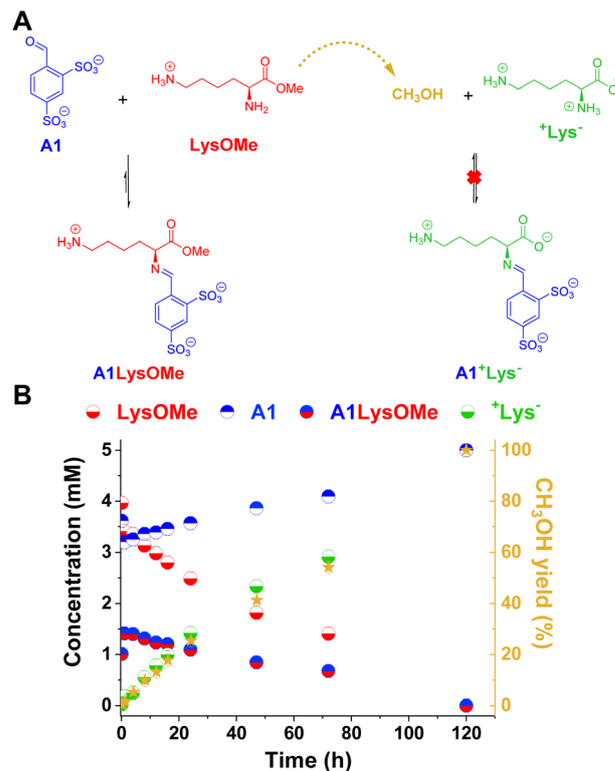


Fig. 3 (A) Chemical representation of the **LysOMe** reaction with **A1** influenced by the hydrolysis of **LysOMe** to produce  $^+Lys^-$ . (B) Concentration profile for the different species detected in solution over 120 h of reaction. The right Y-axis corresponds to the yield of methanol (dark yellow stars) calculated. Concentration and yields determined by  $^1H$  NMR spectroscopy (500 MHz,  $D_2O$ , pD = 7.0, PBS buffer 50 mM, 293 K). The hydrate detected was included in the aldehyde concentration (mM). Initial concentration of reagents: 5 mM. Error bars correspond to 5% of the measurement.

protonated, this component should not react with the released aldehyde (see entry 2 in Table S4†), so that the final composition of the system is expected to consist predominantly of free **A1** and  $^+AA^-$ .

The reaction between **LysOMe** and **A1** (5 mM each) at room temperature in the absence of enzyme was monitored for 120 h using  $^1H$  NMR spectroscopy (Fig. 3). The aldimine **A1LysOMe** was formed quite fast, reaching its maximum concentration of *ca.* 1.5 mM (30% yield) after 1 h of reaction. However, its concentration progressively decreased over time, as the hydrolysis of **LysOMe** occurred (see  $CH_3OH$  yield in Fig. 3B). As predicted, the final component distribution was  $^+Lys^-$  (*ca.* 5 mM) and **A1** (*ca.* 5 mM), with aldimine **A1<sup>+</sup>Lys<sup>-</sup>** not being detected. Subsequently, the adaptive response – and emergence of new properties – in more complex DCvLs was investigated (see following sections).

### Dynamic covalent libraries (DCvLs)

**Enzymatic action on 3-component [1 × 2] libraries.** A competition experiment based on **A1** : **LysOMe** : **PheOMe** gave after 1 h a mixture of 2.9 mM **A1**, 4.1 mM **LysOMe**, and 3.8 mM **PheOMe**, together with 1.2 mM **A1LysOMe** and 0.7 mM **A1PheOMe** (DCvL1, Fig. 4A, B and S10†).



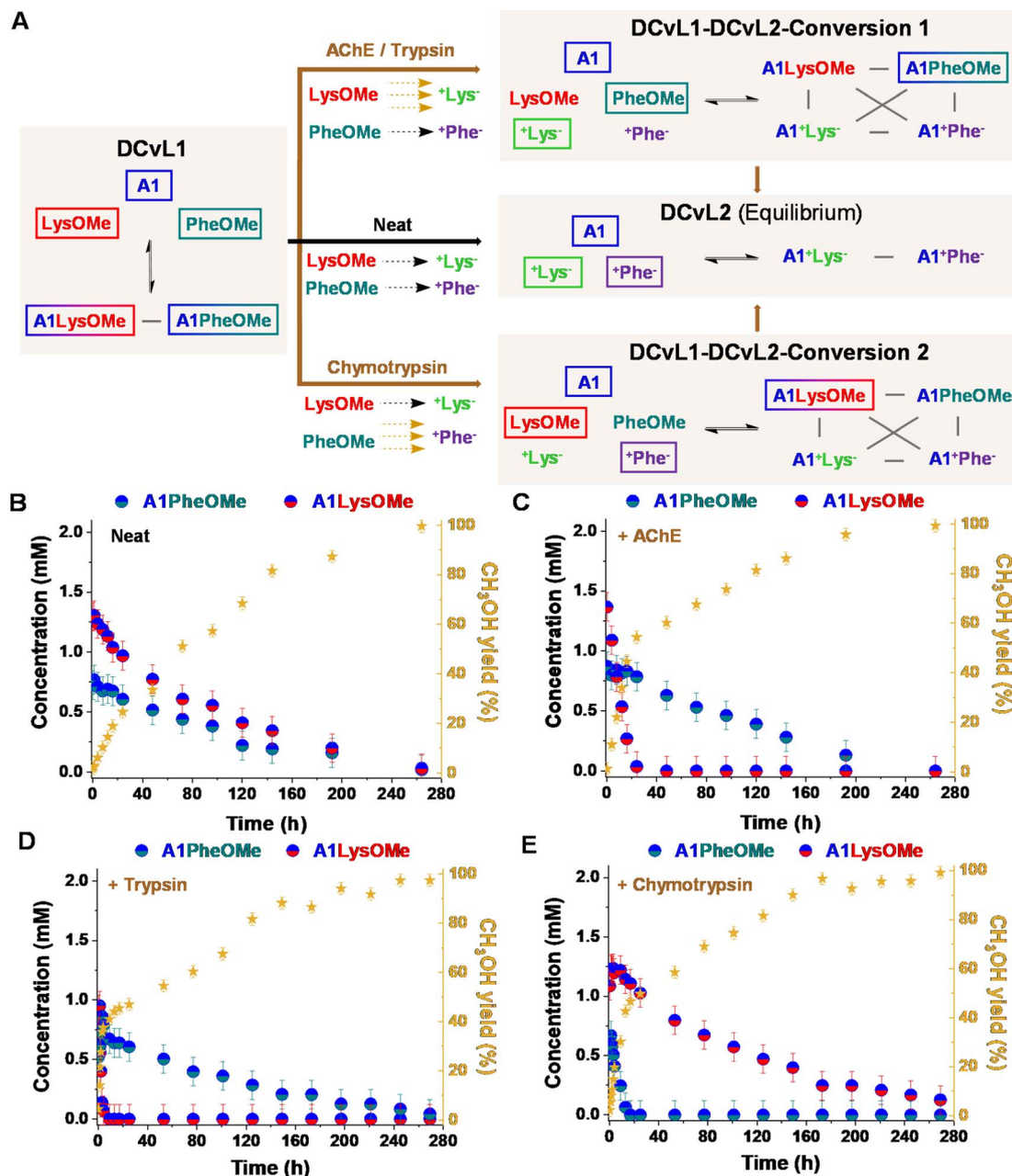


Fig. 4 (A) Reaction between A1, LysOMe, and PheOMe in the absence (black path) and presence of AChE, trypsin and chymotrypsin (brown paths, below). The DCvL1-DCvL2-Conversion 1/2 (5 components, 4 constituents) from DCvL1 (3 components, 2 constituents) to DCvL2 (3 components, 2 constituents) have also been introduced. Predominant compounds (>0.7 mM) are marked with a coloured rectangle. Grey lines correspond to antagonistic relationships; the brown and black arrows represent the time evolution of DCvL1 to reach DCvL2, in the presence and absence of enzymes, respectively. Concentration profile for the different constituents detected in solution over 264 h of reaction in the absence (B) and presence of (C) AChE, (D) trypsin, and (E) chymotrypsin. Concentration and yields determined by <sup>1</sup>H NMR spectroscopy (500 MHz, D<sub>2</sub>O, pD = 7.0, PBS buffer 50 mM, 293 K). Initial concentration of reagents: 5 mM.

The hydrolysis of both AAOMe to their corresponding <sup>+</sup>AA<sup>-</sup> derivatives converted DCvL1 (3 components: A1:LysOMe:PheOMe, 2 constituents: A1LysOMe:A1PheOMe) into the DCvL2 (3 components: A1:<sup>+</sup>Lys<sup>-</sup>:<sup>+</sup>Phe<sup>-</sup>, 2 constituents: A1<sup>+</sup>Lys<sup>-</sup>:A1<sup>+</sup>Phe<sup>-</sup>) through a transient state of higher constitutional complexity (5 components: A1:LysOMe:PheOMe:<sup>+</sup>Lys<sup>-</sup>:<sup>+</sup>Phe<sup>-</sup>, 4 constituents: A1LysOMe:A1PheOMe:<sup>+</sup>Lys<sup>-</sup>:<sup>+</sup>Phe<sup>-</sup>). After 264 h of reaction, the system reached the equilibrium characterized by

the absence of imines and the presence of A1, <sup>+</sup>Lys<sup>-</sup>, and <sup>+</sup>Phe<sup>-</sup> (ca. 5 mM each, Fig. S10 and S11<sup>†</sup>). In the absence of an enzyme, a scrambled mixture of aldimines was always observed due to the subtly similar hydrolysis rates for LysOMe and PheOMe, with the concentration of constituent A1LysOMe being subtly higher than that of A1PheOMe (Fig. 4B and S4; see also Table S5<sup>†</sup>). However, when 0.01% of AChE was added to the mixture, a different scenario was found since this enzyme was able to selectively



catalyse the hydrolysis of **LysOMe** (Fig. S7 and Table S7<sup>†</sup>). **A1LysOMe** completely vanished within the first 24 h, because of the very efficient hydrolysis of **LysOMe** to <sup>+</sup>**Lys**<sup>−</sup> catalysed by the enzyme. The concentration of these species after 20 h was of *ca.* 0.1 mM and 4.9 mM, respectively (Fig. S12 and S13<sup>†</sup>). Once the hydrolysis of **LysOMe** was completed, a drastic negative change in the slope **CH<sub>3</sub>OH** yield was observed, accompanied by a minor up-regulation of **A1PheOMe** (Fig. 4C).<sup>42</sup> The fast and selective consumption of **LysOMe** resulted in a product distribution unattainable in the absence of enzyme, with **A1PheOMe** being the predominant aldimine constituent during the **DCvL1-DCvL2-Conversion 1**. **DCvL1** was also subjected to **trypsin** and **chymotrypsin** to study the distinct adaptive behaviour (*i.e.*, change in sorting selectivity) of this dynamic system. In the presence of **trypsin** (Fig. 4D), the new **DCvL1-DCvL2-Conversion 1** species distribution after 5 h was of **LysOMe** (<0.1 mM), **PheOMe** (3.8 mM), **A1LysOMe** (<0.1 mM), **A1PheOMe** (0.7 mM), <sup>+</sup>**Lys**<sup>−</sup> (5 mM), and <sup>+</sup>**Phe**<sup>−</sup> (0.4 mM), stressing that the only aldimine constituent during this higher-in-complexity state was **A1PheOMe** (Fig. S14<sup>†</sup>). Alternatively, **chymotrypsin** accelerated the hydrolysis of **PheOMe** (Fig. 4E), as evidenced by the predominance of the imine **A1LysOMe** (*ca.* 1 mM after 24 h) during the transient **DCvL1-DCvL2-Conversion 2** (Fig. 4A and E; see also Fig. S15<sup>†</sup>). With time, the imino-species that were not affected by the enzymes were also depleted, reaching in all cases the final **DCvL2** equilibrium state after *ca.* 264 h.

One notes that the lifetime of the sorted distribution might be modulated with the enzyme concentration. The **DCvL1** was thus studied in the presence of varying amounts of **chymotrypsin**. Whereas the use of 0.01% of the enzyme resulted in a transient sorting (**A1LysOMe** predominant aldimine) lasting for 250 h (Fig. S16B<sup>†</sup>), the introduction of 0.006 and 0.003% **chymotrypsin** resulted in sorted states for 240 h and 204 h, respectively (Fig. S16B and C<sup>†</sup>). Alternatively, when 0.03% of the enzyme was added, the **A1PheOMe** imine vanished in less than 4 h and the sorting was observed for 260 h (Fig. S16D<sup>†</sup>). These results inferred that the selectivity and lifetime of the sorted transient states can be easily controlled by the nature and the amount of enzyme introduced.

We also evaluated the possibility to build-up a similar library but based on a more challenging process: the selective chiral resolution of a racemic mixture induced by a remote modification. **AChE** was found to catalyse the hydrolysis of (*L*)**LysOMe** but it barely affected the hydrolysis rate of (*D*)**LysOMe** (Fig. S8 and Table S8<sup>†</sup>). The **DCvL3** using **A1** : (*L*)**LysOMe** : (*D*)**LysOMe** as components was then assayed (5 mM each, Fig. 5A). In this case, <sup>1</sup>H NMR analysis of the reaction course was less useful as it did not allow for differentiation between each pair of enantiomeric components, not even in the presence of water-soluble chiral shifting reagent [(*R*)-1,2-diaminopropane-*N,N,N',N'*-tetraacetato]samarate(III).<sup>43</sup> Yet, one can estimate the abundance of each species considering the rates of hydrolysis for each separated

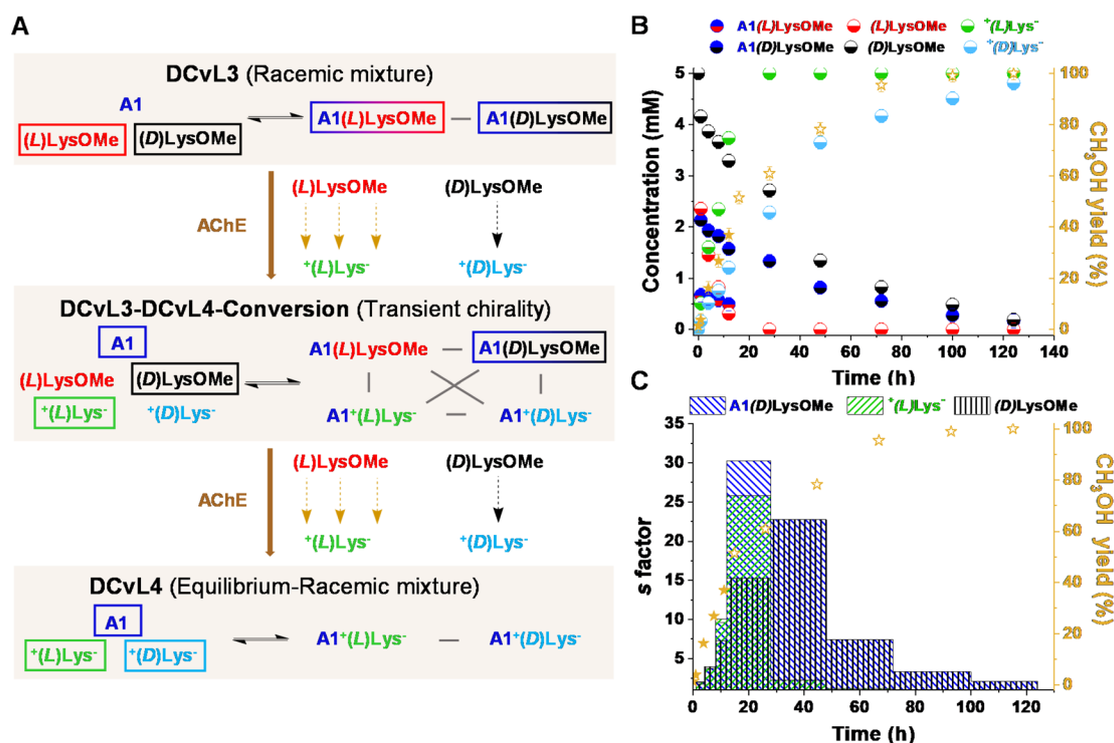


Fig. 5 (A) Reaction between (*D*)**LysOMe**, (*L*)**LysOMe** and **A1** in the presence of **AChE** (0.01 mol%). **DCvL3-DCvL4-Conversion** (5 components, 4 constituents) from **DCvL3** (3 components, 2 constituents) to **DCvL4** (3 components, 2 constituents) has also been introduced. Predominant compounds (>1.5 mM) are marked with a coloured rectangle. Grey lines correspond to antagonistic relationships; the brown arrows represent the time evolution of **DCvL3** to reach **DCvL4**. (B) Concentration profile for the different constituents detected in solution over 140 h. Concentration and yields determined by <sup>1</sup>H NMR spectroscopy (500 MHz, D<sub>2</sub>O, pD = 7.0, PBS buffer 50 mM, 293 K). Initial concentration of reagents: 5 mM. (C) Time evolution of the *s* factors calculated for **A1(D)LysOMe**, <sup>+</sup>(*L*)**Lys**<sup>−</sup>, and (*D*)**LysOMe** over 126 h.



reaction and the yield of  $\text{CH}_3\text{OH}$  monitored for the kinetic resolution of the racemic mixture (see ESI† for more details). The time evolution for the different species revealed the conversion of the DCvL3 to DCvL4 (Fig. 5A). After 16 h of reaction, a severe deceleration in  $\text{CH}_3\text{OH}$  production rate was observed, indicating that all (*L*)LysOMe had been consumed (Fig. S17†). At this point, the composition of the system was: (*L*)LysOMe (<0.1 mM), (*D*)LysOMe (2.5 mM), A1(*L*)LysOMe (<0.1 mM), A1(*D*)LysOMe (1.3 mM),  $^+(\textit{L})\text{Lys}^-$  (5 mM), and  $^+(\textit{D})\text{Lys}^-$  (2.1 mM) (Fig. 5B). Therefore, this higher-in-complexity DCvL3-DCvL4-Conversion (5 components, 4 constituents) presented remarkable transient chirality,<sup>44</sup> with the highest *s* factor of *ca.* 30 being attained after 24 h (Fig. 5C).<sup>45,46</sup> After 120 h, the complete consumption of both enantiomers of LysOMe was achieved, with the DCvL reaching its equilibrium state (DCvL4): a racemic mixture of  $^+(\textit{L})\text{Lys}^-$  and  $^+(\textit{D})\text{Lys}^-$  together with A1 (5 mM each; Fig. S18†). These results emphasize how the selective action of different enzymes on DCvLs can drastically change their constitution, allowing for transient states of higher constitutional complexity with sorted distributions.

#### Chemically-driven selection in $[1 \times 2]$ , $[1 \times 3]$ and $[2 \times 2]$ DCvLs

$[1 \times 2]$  DCvL. In the systems described above, the ester hydrolysis gave rise to the unreactive  $^+\text{AA}^-$  component and the up-regulation of free aldehyde A1. It can be noted that if the competition experiments are performed in the presence of an amino component (B2) which can scavenge A1, the concentration of the resulting imine (A1B2) should be amplified with time as the silencing of the competing component AAOMe occurs by its conversion into  $^+\text{AA}^-$ . Initially, DCvL5 (reaction between A1, B2, and LysOMe; Fig. 6A and S19A†) was studied at 5 mM concentration for each reagent. Nevertheless, only a minor increase in the concentration of A1B2 was observed with time as the component LysOMe was hydrolysed to the inhibited  $^+\text{Lys}^-$  (Fig. S19B and S20†). Based on previous results (Fig. S1†), we evaluated the AAOMe hydrolysis and imination reactions of A1 and AAOMe/ $^+\text{AA}^-$  at 80 mM. At such high concentrations, the kinetic profiles for the hydrolysis of AAOMe were comparable to the ones observed at 5 mM (Fig. S21 and Table S10 vs. Fig. S4 and Table S5†), but the reaction between  $^+\text{AA}^-$  and A1 could not be completely inhibited at 80 mM as some A1 $^+\text{AA}^-$  aldimine was observed (*ca.* 5–10%, see Table S11†).

DCvL5 was then assessed at 80 mM to see the effect on both selection and activity. As expected, a more pronounced up-regulation of A1B2 was noted, with a concentration increase of 17 mM after 400 h (Fig. 6B; see also Fig. S19B† for comparison). During the hydrolysis of LysOMe, the system experienced a conversion from DCvL5 to DCvL6, with the corresponding increase in complexity during the DCvL5-DCvL6-Conversion: from 3 to 4 components and from 2 to 3 constituents (Fig. 6A).<sup>28,47,48</sup> The final composition of DCvL6 was: 52 mM A1B2, 26 mM A1, 3 mM A1 $^+\text{Lys}^-$ , 77 mM  $^+\text{Lys}^-$ , and 28 mM B2 (Fig. S22 and S23†). Remarkably, as the LysOMe hydrolysis occurred, the system evolved from a scrambled mixture of two constituents in DCvL5 (*ca.* 35 mM of A1B2 and A1LysOMe) to a sorted outcome in DCvL6 containing A1B2 as the main aldimine constituent.<sup>49–51</sup>

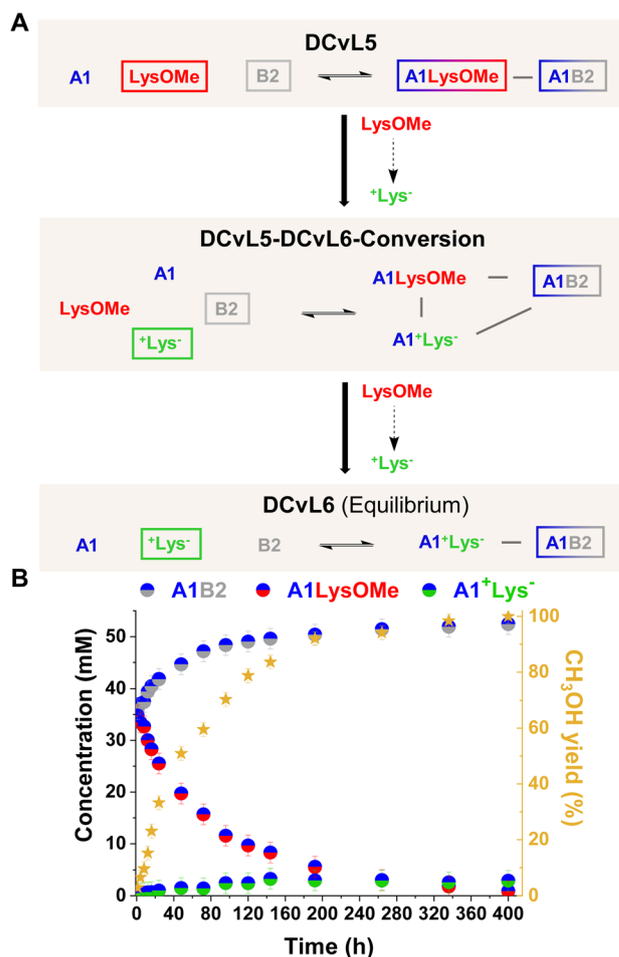
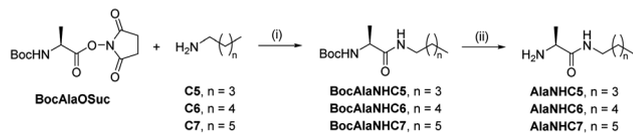


Fig. 6 (A) Reaction between A1, LysOMe, and B2. DCvL5-DCvL6-Conversion (5 components, 4 constituents) from DCvL5 (3 components, 2 constituents) to DCvL6 (3 components, 2 constituents) has also been represented. Predominant compounds (>35 mM) are marked with a coloured rectangle. Grey lines correspond to antagonistic relationships; the black arrows represent the time evolution of DCvL5 to reach DCvL6. (B) Concentration profile for the different constituents detected in solution over 400 h of reaction. Concentration and yields determined by  $^1\text{H}$  NMR spectroscopy (500 MHz,  $\text{D}_2\text{O}$ ,  $\text{pD} = 7.0$ , PBS buffer 500 mM, 293 K). Initial concentration of reagents: 80 mM.

This chemically-driven adaptation in DCvL5 was further studied in the absence of buffer (*i.e.* neat  $\text{D}_2\text{O}$ ). The presence of the quite basic component B2 led to an initial  $\text{pD}$  value of 8.3, resulting in slightly higher imine abundances than in the buffered system at  $\text{pD}$  7.0 (*ca.* 40 mM and 35 mM, respectively; see Fig. 6B and S24†). The LysOMe to  $^+\text{Lys}^-$  conversion triggered a similar constitutional rearrangement to that observed in the presence of buffer, but at subtly slower rates in this case. The equilibrium state of DCvL6 was characterized by the absence of A1LysOMe and the predominance of A1B2 (*ca.* 60 mM). During the DCvL5-DCvL6-Conversion, the  $\text{pD}$  of the system dropped 1.2 units as a result of the ester hydrolysis, reaching a final  $\text{pD}$  value of 7.1 (Fig. S25†). Thus, such noteworthy decrease in  $\text{pD}$  values regulated the expression of the A1 $^+\text{Lys}^-$  constituent (in terms of imine formation) and allowed for a sorted state with higher fidelity in DCvL6.





Scheme 1 Two-step synthesis of amphiphilic pseudopeptidic compounds AlaNHcX. (i) DCM, r.t., 16 h. (ii) TFA, DCM, r.t., 3 h.

$[1 \times 3]$  DCvL. We decided to study the behaviour of related  $[1 \times 3]$  systems on addition of a further adaptive step by combining the up-regulation of the unaffected constituent to the generation of a macroscopic property. To this end, a new family of amphiphilic pseudopeptidic compounds with different chain lengths (AlaNHcX, Scheme 1) was synthesized following reported procedures.<sup>52</sup> These amines were selected since their aldimine derivatives formed upon reaction with a charged aldehyde (e.g., A1) should present a prominent tendency to self-assemble in water and form micelles.<sup>53</sup> <sup>1</sup>H NMR at 80 mM indicated yields of ca. 50% for the imines A1AlaNHcX (see Table S12<sup>†</sup>). These values, despite being slightly lower than those of AAOMe under the same conditions, showed that the pseudopeptides were suitable for competition experiments. Moreover, the concentration of the aldimines A1AlaNHcX was not altered over time, indicating that the amide bond was stable to hydrolysis for at least 1 month.

The self-assembling properties of the different compounds were then evaluated by fluorescence emission spectroscopy using Nile Red (NR) as a hydrophobic probe.<sup>54</sup> SDS was used as a control for the fluorescence emission values of a system containing micelles. The parent amines AlaNHcX barely formed micelles even at high concentrations (80 mM, Fig. S26<sup>†</sup>). In contrast, aldimine derivatives were found active in micellization. For instance, A1AlaNHc7 seemed to self-assemble into micellar structures, as evidenced by the intense fluorescence response observed (Fig. S27<sup>†</sup>). The CMC for A1AlaNHc7 was in the 7–12 mM range. DLS analyses of the A1AlaNHc7 sample at 12 mM revealed the presence of micelles with an average diameter of 7.9 nm (Fig. S28<sup>†</sup>).

The hydrolysis reactions occurring in the DCvL7 (A1, LysOMe, ArgOMe and AlaNHc7; 1 eq. each) should result in the decrease of A1LysOMe and A1ArgOMe together with an up-regulation of A1AlaNHc7 over time (Fig. 7A). When the concentration of A1AlaNHc7 increases from below to above CMC, the spontaneous formation of micelles (A1AlaNHc7-mic) should be noted in the solution. Reaction between A1 : LysOMe : ArgOMe : AlaNHc7 (35 mM each) resulted in the DCvL7 with a constituent concentration of 10, 9, and 6 mM respectively for A1LysOMe, A1ArgOMe, and A1AlaNHc7. Since the concentration of A1AlaNHc7 was below the CMC, the relative fluorescence response observed was of 0.3, in good agreement with the absence of micelles.

The hydrolysis of ArgOMe and LysOMe to their corresponding <sup>+</sup>AA<sup>-</sup> released A1 into the medium, as the hydrolysed amino acids were silent towards imination due to protonation. AlaNHc7 scavenged the generated aldehyde and led to an amplification of A1AlaNHc7. After 90 h of reaction, the

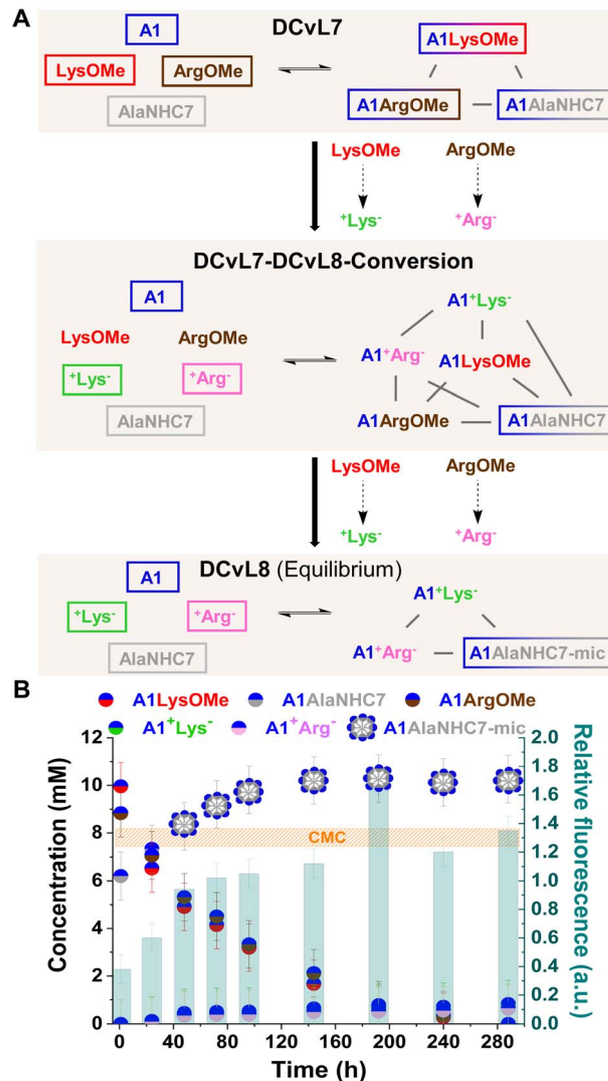


Fig. 7 (A) Reaction between A1, LysOMe, ArgOMe, and AlaNHc7. DCvL7-DCvL8-Conversion (6 components, 5 constituents) from DCvL7 (4 components, 3 constituents) to DCvL8 (4 components, 3 constituents) has also been represented. Predominant compounds (>5 mM) are marked with a coloured rectangle. Grey lines correspond to antagonistic relationships; the black arrows represent the time evolution of DCvL7 to reach DCvL8. (B) Concentration profile for the different constituents detected in solution over 290 h of reaction. Concentrations determined by <sup>1</sup>H NMR spectroscopy (500 MHz, D<sub>2</sub>O, pD = 7.0, PBS buffer 500 mM, 293 K). Initial concentration of reagents: 35 mM. CMC determined by fluorescence spectroscopy.

concentration of A1AlaNHc7 was of ca. 10 mM, with a relative fluorescence emission >1 (Fig. 7B). Therefore, A1AlaNHc7 assemblies (namely A1AlaNHc7-mic) were present in the solution. Such high relative fluorescence values were maintained over time, indicating that the micelles were stable under the conditions assayed. DLS analyses for the final sample revealed the presence of micelles with a hydrodynamic diameter of 7.2 nm. The observed DCvL7-DCvL8-Conversion presented a higher complexity (6 components, 5 constituents) than the initial and final states (DCvL7 and DCvL8, respectively, both with 4 components and 3 constituents). The equilibrium



distribution of the DCvL8 was of 10 mM A1AlaNHCH7-mic, 1 mM A1<sup>+</sup>Lys<sup>-</sup>, 1 mM A1<sup>+</sup>Arg<sup>-</sup>, 25 mM A1, 34 mM <sup>+</sup>Lys<sup>-</sup> and 34 mM <sup>+</sup>Arg<sup>-</sup> (Fig. S29 and S30<sup>†</sup>). Therefore, DCvL7 experiences a triple-state adaptation: DCvL conversion + selective constituent amplification + formation of self-assembled dynamic micelles; resulting in the emergence of a property neither contained in the components nor in the initial library.

[2 × 2] DCvL. Finally, the adaptive behaviour of a [2 × 2] DCvL was also studied. As reported, [2 × 2] CDNs respond to constitutional relationships between species upon addition of

certain effectors.<sup>55–57</sup> The up-regulation of the most thermodynamically-favoured constituent of a network (*e.g.*, AB) is accompanied by the amplification of its agonistic – and least fitted – A'B' constituent, at expenses of the antagonistic AB' and A'B constituents.<sup>7,26</sup> We surmised that building-up a [2 × 2] library in which one of the competing amines is silenced by hydrolysis should result in a simultaneous increase in concentration of the two antagonistic constituents that are not hydrolysed.

The initial distribution of DCvL9, built-up from LysOMe, B2, A1 and A3 (Fig. 8A), revealed that the aldimines derived from A1 were present in higher concentrations (*ca.* 35 mM) than those of A3 (*ca.* 20 mM) as a result of its higher reactivity (see Fig. 2A and B). As the LysOMe ester hydrolysis proceeded, DCvL9 converted into DCvL10 through the higher-in-complexity DCvL9-DCvL10-Conversion, as seen from the <sup>1</sup>H NMR spectra by the appearance of <sup>+</sup>Lys<sup>-</sup> signals. This hydrolysed component barely reacted with the aldehydes, resulting in the up-regulation of A1B2 and A3B2. Despite their antagonistic relationship – both constituents contain B2 as building block –, an increase in concentration of *ca.* 15 and 10 mM was noted for A1B2 and A3B2, respectively (Fig. 8B). The species distribution after 264 h of reaction was: 50 mM A1B2, 28 mM A3B2, 3 mM A1<sup>+</sup>Lys<sup>-</sup>, < 1 mM A3<sup>+</sup>Lys<sup>-</sup>, 29 mM A1, 48 mM A3, 77 mM <sup>+</sup>Lys<sup>-</sup> and 2 mM B2 (Fig. S31 and S32<sup>†</sup>). Thus, the DCvL conversion, triggered by ester hydrolysis, led to the constitutionally-disfavoured up-regulation of two antagonistic constituents in DCvL10 (see Fig. 8A). This behaviour resulted from the high degree of protonation of the competing amine (*i.e.*, <sup>+</sup>Lys<sup>-</sup>) that stimulated the scavenging of A1 and A3 by B2.

## Conclusions

The hydrolysis of different AAOMe constituents to their corresponding <sup>+</sup>AA<sup>-</sup> in a series of DCvLs affects imine formation with several aldehydes. This remote chemical modification increases the protonation degree of the N<sub>α/β</sub> amino group due to the lower pK<sub>a</sub> of the hydrolysed species, thus hampering imine formation.

Monitoring the reaction between LysOMe and A1 showed the expected depletion of A1LysOMe as <sup>+</sup>Lys<sup>-</sup> was produced, with the aldimine A1<sup>+</sup>Lys<sup>-</sup> not being observed due to the protonation of <sup>+</sup>Lys<sup>-</sup> (inhibited component). The kinetic simplification of two dynamic libraries (DCvL1 and DCvL3) in the presence of different enzymes (*i.e.*, AChE, trypsin and chymotrypsin) was attained, with the catalysed remote chemical modification leading to selective silencing of components, sorted states and transient chirality. In [1 × 2] and [1 × 3] DCvLs, containing a component that was not hydrolysed (*i.e.*, the scavenger B2 or AlaNHCH7), the hydrolysis of the AAOMe resulted in the up-regulation of the constituents made of these components. Such DCvLs experienced a chemically-driven constitutional sorting, evolving from an almost statistical distribution (2 and 3 aldimines for DCvL5 and DCvL7, respectively) to a state characterized by the predominance of a single imine. In addition, the DCvL7 displayed a triple-state adaptation based on DCvL conversion, selective constituent up-regulation and supramolecular self-assembly with emergence of a macroscopic property

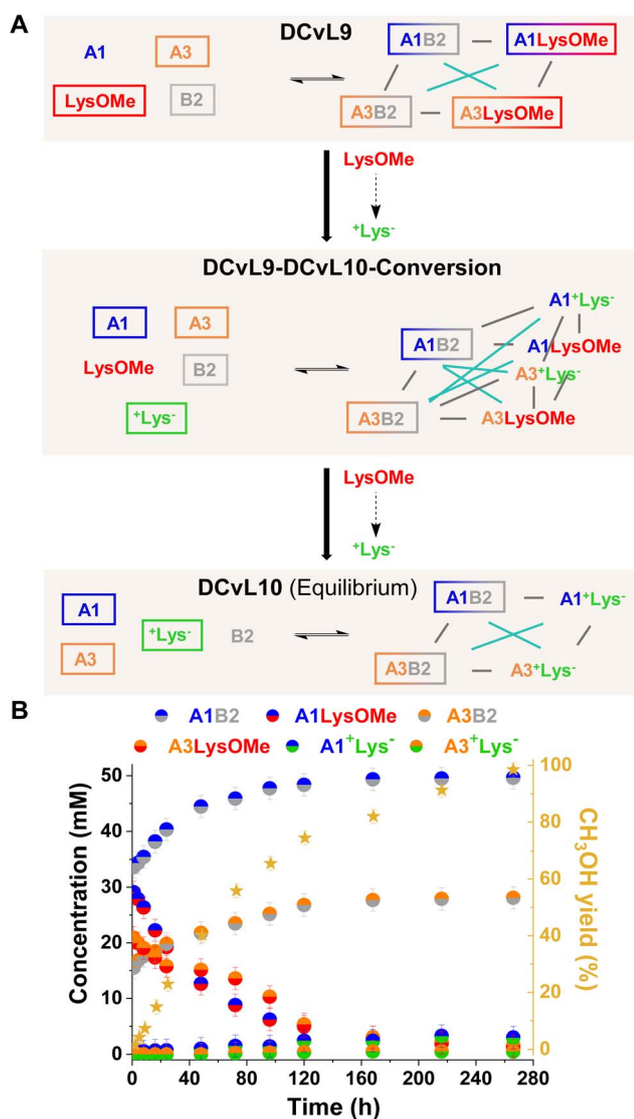


Fig. 8 (A) Reaction between A1, A3, LysOMe, and B2. DCvL9-DCvL10-Conversion (5 components, 6 constituents) from DCvL9 (4 components, 4 constituents) to DCvL10 (4 components, 4 constituents) has also been represented. Predominant compounds (>15 mM) are marked with a coloured rectangle. Grey and cyan lines correspond to antagonistic and agonistic relationships, respectively; the black arrows represent the time evolution of DCvL9 to reach DCvL10. (B) Concentration profile for the different constituents detected in solution over 264 h of reaction. Concentration and yields determined by <sup>1</sup>H NMR spectroscopy (500 MHz, D<sub>2</sub>O, pD = 7.0, PBS buffer 500 mM, 293 K). Initial concentration of reagents: 80 mM.



that was not present in the initial library: the generation of micelles. A  $[2 \times 2]$  library was also constructed in which one of the amino components was selectively poisoned by hydrolysis. The conversion from DCvL9 to DCvL10 resulted in a concentration increase for two constituents with an antagonistic relationship. Thus, all the DCvLs studied present transient states of higher constitutional complexity that adapt and evolve with time to states of higher “simplicity”. In addition, the interplay between the action of different enzymes and the libraries dictates the selectivity and the lifetime of the transient sorted states (up-regulated constituent and/or chirality).

In a broader sense, such adaptive responses in simple dynamic networks offer an evolutionarily relevant scenario were a remote side-reaction triggered by the environment (*i.e.*, solvent) amplifies the population of the “surviving” constituents at the expense of the “poisoned” ones which undergo the detrimental chemical modification. These results will pave the way for the design of related dynamic systems of higher functionality with emerging functions. Furthermore, the observed silencing process based on reactive site protonation showcases the mode-of-action found in some proteins for the regulation of their activity through specific PTMs that lead to  $pK_a$  modulations. Therefore, this work will also contribute to our knowledge on sophisticated biological mechanisms governed by supra-molecular cross-talks between remote sites.

## Data availability

Data available upon request from the authors.

## Author contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript. Conceptualization: F. E., J. M. L. Data curation: F. E., T. R. Formal analysis: F. E. Funding acquisition: J. M. L. Investigation: T. R. (supporting), F. E. (lead). Methodology: F. E. Project administration: J. M. L. Resources: J. M. L. Supervision: F. E., J. M. L. Validation: F. E. Visualization: F. E. Writing original draft: F. E. Writing – review and editing: T. R. (supporting), F. E., J. M. L.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

This work was supported by the ERC (Advanced Research Grant SUPRADAPT 290585), the University of Strasbourg Institute for Advanced Study (USIAS), and Fondation Jean-Marie Lehn. F. E. acknowledges Fundación Ramón Areces for a postdoctoral fellowship. T. R. thanks the CSC Graduate School funded by the French National Research Agency (CSC-IGS ANR-17-EURE-0016) for a PhD fellowship. The authors thank Dr J. L. Schmitt, Dr A. Osypenko, Dr Z. Yang, Dr S. Jung, B. Kozibroda, M. J. Aguilera, and C. Antheaume for helpful discussions.

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