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#### Complex coacervation and compartmentalized conversion of prebiotically relevant metabolites

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### 9 Abstract

10 Metabolism and compartmentalization are two of life's most central elements. Constructing synthetic assemblies based on prebiotically relevant molecules that combine these elements can provide insight 11 12 into the requirements for the formation of life-like protocells from abiotic building blocks. In this work, 13 we show that a wide variety of small anionic metabolites have strong enough interactions with oligoarginine (R<sub>10</sub>) to form coacervate protocells through liquid-liquid phase separation. The stability of 14 15 the coacervates can be rationalized by the molecular structure of the metabolites, and we show that three 16 negative charges for carboxylates or two negative charges complemented with an unsaturated functional 17 group for phosphates and sulfates is sufficient for phase separation. We show that these metabolites 18 remain reactive in compartmentalized systems. Protometabolic reactions that lead to an increased 19 interaction with the oligopeptide can be exploited to induce the formation of coacervate protocells. The 20 resulting coacervates can localize other metabolites and enhance their conversion. Finally, reactions of

21 compartmentalized metabolites can also alter the 22 physicochemical properties of the coacervates and ultimately 23 lead to protocell dissolution if the reaction products decrease 24 the coacervate stability. These results reveal the intricate 25 interplay between (proto)metabolic reactions and coacervate 26 compartments, and show that coacervates are excellent 27 candidates for metabolically active protocells.



### 28 Introduction

29 Metabolism and compartmentalization are fundamental elements of life.<sup>1, 2</sup> Prebiotic routes to form and 30 convert central metabolites and ways to compartmentalize them are therefore of utmost relevance. Prebiotic reactions of common metabolites such as ATP, NADH and citric acid have been studied 31 32 extensively, as these molecules play a key role across the tree of life in reaction pathways such as the tricarboxylic acid (TCA) cycle, which releases energy to fuel various cellular processes.<sup>3-9</sup> However, their 33 34 concentrations on the early Earth were likely very low, slowing down these protometabolic reactions to 35 non-viable pace. Moreover, some of the developed reactions proceed under mutually incompatible 36 conditions of, for instance, pH, redox potential or light. Compartmentalization of reagents can increase 37 local concentrations and create distinct chemical microenvironments, and might have therefore been an important step in gaining sufficiently high reaction rates and selectivity at the origins of life.<sup>10-13</sup> 38

Compartmentalization through self-assembly of amphiphiles into vesicles has long been 39 considered as the general mechanism for the formation of the first protocells.<sup>14</sup> However, this 40 41 encapsulation was likely inefficient without the presence of active exchange of components by complex transport machineries, limiting the local concentrations that could be reached inside these protocells.<sup>15</sup> 42 Since the discovery that membraneless organelles in modern cells can form by liquid-liquid phase 43 separation (LLPS), the analogous formation of protocells by LLPS has received increasing interest.<sup>16-18</sup> 44 Multiple weak associative interactions, such as charge-charge and cation- $\pi$  interactions, between 45 molecules can drive LLPS to form (complex) coacervates, which are droplets of a solute-rich phase 46 47 dispersed in a dilute continuous phase.<sup>19, 20</sup> These coacervate protocells have been shown to spontaneously concentrate reagents and increase reaction rates, <sup>21, 22</sup> and recently it has been shown that 48 49 they can even be formed by prebiotically relevant mononucleotides and short homopeptides containing just five residues.<sup>23, 24</sup> Moreover, droplets made from peptides with prebiotically relevant low molecular 50 51 weight were found to be more effective at generating distinct pH microenvironments, accumulating RNA 52 and stabilizing its secondary structure, than coacervates formed by high molecular weight 53 polyelectrolytes.<sup>24</sup>

54 Since many key metabolites have multiple negative charges, we hypothesized that these 55 metabolites could similarly undergo complex coacervation to become compartmentalized inside 56 protocells, and that these protocells could be capable of sustaining metabolic reactions. While the large 57 majority of complex coacervates in literature are composed of long polymers, a few examples of complex 58 coacervates made of small metabolites have been reported. AMP and citric acid have been shown to 59 phase separate with oligoarginine and protamine, a 4.2 kDa protein containing 21 arginine residues, respectively.<sup>24, 25</sup> Additionally, pyrophosphate and triphosphate were found to phase separate with 60 61 lysozyme and histatin-5, both of which have a net charge of z = +6 or +7 at the investigated pH.<sup>26, 27</sup> 62 However, a systematic analysis of phase separation of prebiotic metabolites is still lacking, and the 63 metabolic activity of the coacervate protocells is unexplored.

64 Here, we show that anionic metabolites with at least three charges, or two when complemented 65 with additional cation- $\pi$  or  $\pi$ - $\pi$  interaction sites, are able to form complex coacervates with the short 66 peptide oligoarginine (R<sub>10</sub>). We compare various types of prebiotically relevant metabolites, including 67 carboxylic acids, phosphates and sulfates in their ability to form coacervates, and show that their stability varies significantly due to small differences in molecular structures, such as their degree of unsaturation, 68 69 hydrogen bonding ability and charge density. We show that these metabolites continue to undergo 70 metabolic reactions and that these reactions can control compartmentalization and vice versa: metabolite 71 conversion can give rise to active formation and growth of coacervate protocells, metabolite conversion 72 can be accelerated inside coacervate compartments, and finally, metabolite conversion inside 73 coacervates can alter the nature of the protocells, by creating products that affect the stability of the 74 coacervates. These observations reveal the intricate interplay between protometabolic reactions and 75 coacervate compartments, and show that coacervates are excellent candidates to compartmentalize 76 protometabolic networks.

77 Results and discussion

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#### 79 Metabolites with at least two negative charges form stable coacervates with oligoarginine

80 Inspired by the central importance of metabolism in living systems, we wondered if key prebiotic 81 metabolites could be compartmentalized by liquid-liquid phase separation, and if so, how propensity for 82 phase separation is linked to the molecular structure. To answer these questions, we investigated three 83 classes of metabolites and related small molecules: carboxylic acids, phosphates, and sulfite/sulfates. We 84 selected di- and tricarboxylic acids from the tricarboxylic acid (TCA) and 4-hydroxy-2-ketoglutarate (HKG)<sup>4</sup> 85 cycles, complemented with an aromatic tricarboxylic acid. Phosphates were selected that are part of the 86 cellular respiration, like ATP, which functions as biochemical energy currency, or prebiotic analogues, such 87 as triphosphate and trimetaphosphate (TMP). Finally, we selected sulfites and sulfates and some iron complexes, because of their possible relevance in protometabolic reactions on early Earth.<sup>28, 29</sup> For each 88 89 molecule, we determined its ability to phase separate with a short oligoarginine peptide ( $R_{10}$ ). 90 Oligoarginine was chosen as the model cation because it is known to form coacervates with small molecules like AMP and because arginine-containing peptides have been suggested to occur on the early 91 Earth.<sup>24, 30</sup> To keep the system as simple as possible, stock solutions were adjusted to a pH of  $7.0 \pm 0.3$ 92 93 with HCl and NaOH and no buffer was used, since excess ions present in the buffer can screen the 94 electrostatic interactions and mask a potential phase separation. We decided to compare the ability of 95 metabolites to form coacervates at equal concentration, so the concentration of all metabolites is fixed 96 at 5 mM and  $R_{10}$  at 5 mM of arginine residues.

97 We found that carboxylic acid metabolites with three charges (z = -3) were all able to form 98 coacervate droplets with R<sub>10</sub>, while monocarboxylic acids (MCA) and dicarboxylic acid (DCA) resulted in 99 homogenous solutions (Table S1). For phosphates and sulfates, a similar trend is seen, as monophosphate 100 (z = -2) did not phase separate with R<sub>10</sub>, whereas pyrophosphate (z = -3) did. AMP and NADH were an 101 exception and formed coacervates with R<sub>10</sub> even though both molecules only have two negative charges. 102 This can be explained by the cation- $\pi$  interactions of R<sub>10</sub> with the nucleotide bases, which strengthen the 103 overall intermolecular interactions between the metabolite and R<sub>10</sub>, and is in agreement with previous



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106 Figure 1. Critical salt concentrations of coacervate forming metabolites with oligoarginine. (a) Tricarboxylic acids 107 phase separate with R10 while monocarboxylic acid (MCA) and dicarboxylic acid (DCA) do not. (b) Phosphates 108 containing two negative charges (blue) accompanied by a nucleobase can form coacervates with R10. (c) Sulfates 109 with two negative charges form coacervates with R10 while sulfites do not. (d) Ferri- and ferrocyanide coacervates 110 have relatively high critical salt concentrations while not containing aromatic groups. Critical salt concentrations 111 were determined by turbidity measurements and error bars represent the standard deviation (n = 3). Empty regions 112 between brackets indicate a variety of groups (Table S1). Concentration of all molecules is 5 mM (monomer units 113 for R<sub>10</sub>) or 10 mM for sulfites and sulfates.

observations<sup>24</sup>. Interestingly, pyrosulfate (**8**) was also able to phase separate with  $R_{10}$ , while having only two negative charges and no additional interacting groups. We attribute this to the high charge density and number of  $\pi$  electrons, which enhance pyrosulfate's ability to phase separate. Bisulfite did not phase separate with  $R_{10}$ , which supports the hypothesis that the  $\pi$ -electrons in the oxygen sp<sup>2</sup> bond interacts with  $R_{10}$  and contribute to phase separation.

119 To quantify the tendency of small metabolites to phase separate with cationic peptides, such as 120 R<sub>10</sub>, we determined the critical salt concentration (CSC) for each coacervate by titrating the coacervate 121 suspension with NaCl (Figure 1, Table S1). Differences in stability between different coacervate can be caused by the number and strength of interaction sites of their components.<sup>20</sup> For example, the 122 123 guanidinium groups of R<sub>10</sub> form relatively strong charge-charge interactions with carboxylate groups 124 compared to the cation- $\pi$  interactions they form with aromatic groups. The number of charged interaction 125 sites will thus have a large effect on the overall coacervate stability, whereas an increase in cation- $\pi$ 126 interactions will give a small increase in stability, and the ability to form hydrogen bonds an even smaller 127 increase.<sup>20</sup> For the tricarboxylic acids, which all have three negatively charged groups, a higher CSC was found with an increasing number or strength of additional interaction sites (Figure 1a). For example, the 128 129 additional hydroxyl group of citrate (2) and  $\pi$  bond of trans-aconitate (3) increased the CSC by 55 and 65 130 mM, respectively, compared to coacervates formed by  $R_{10}$  and tricarballylate (1). Trimesate had the 131 highest CSC of all tricarboxylic acid coacervates, which can be explained by the presence of the benzene 132 ring that forms relatively strong cation- $\pi$  interactions with R<sub>10</sub> compared to the other additional 133 interactions.<sup>20</sup>

For phosphate metabolites, a similar trend was observed: the CSC is determined in the first place by the number of negative charges, and secondly by additional interactions such as cation- $\pi$  interactions (Figure 1b). The CSC of R<sub>10</sub>/triphosphate (PPPi) coacervates was 300 mM higher than R<sub>10</sub>/pyrophosphate (PPi) coacervates due to the extra negatively charged phosphate group, whereas the CSC of R<sub>10</sub>/ATP coacervates was 100 mM higher than that of R<sub>10</sub>/PPPi coacervates because of the additional adenosine. Moreover, the CSC of R<sub>10</sub>/NADH coacervates was nearly double that of R<sub>10</sub>/AMP coacervates, demonstrating a nucleoside group can greatly increase coacervate stability. We also included

141 trimetaphosphate (TMP), which is a cyclic version of triphosphate and contains three negative charges. 142 Interestingly,  $R_{10}$ /TMP coacervates have a significantly lower CSC than  $R_{10}$ /PPi coacervates (116.47 mM 143 compared to 296.52 mM, respectively), while both molecules have the same net charge (z = -3) and 144 interacting groups. We attribute this decrease in coacervate stability to the lower flexibility of TMP 145 compared to PPi, which reduces the number of dynamic interactions it can make with R<sub>10</sub>, an effect that has been observed in coacervates composed of polylysine and DNA.<sup>31</sup> When comparing the general 146 147 stability of coacervates composed of phosphates compared to carboxylates, it becomes clear that the 148 phosphate coacervates have higher CSCs at similar charge. This can possibly be explained by the higher 149 charge density of the phosphates, as the anionic groups are not separated by an alkyl spacer like in 150 tricarboxylic acids. An additional cause could come from their relative hardness (which manifests itself in a Hofmeister series), leading to a stronger hydration of phosphates.<sup>32</sup> Complex coacervation would 151 152 release some of the bound water molecules, making phase separation with phosphates more favorable.

In the sulfite and sulfates section, it was found that  $R_{10}$ /pyrosulfate (8) coacervates had a higher CSC than  $R_{10}/1,3$ -benzenedisulfate (7) coacervates, which was surprising, as 7 contains an extra benzene group that forms cation- $\pi$  interactions with  $R_{10}$  (Figure 1c). A possible reason for this could be the reduced flexibility of 7 compared to 8, though it should be noted that a CSC difference of lower than 10 mM could have other causes and a wider range of compounds with varying charge densities should be used to determine the effect of molecular flexibility on electrostatic phase separation.

159 Lastly, it is striking how high the measured CSC of ferri- and ferrocyanide coacervates was 160 compared other anions with three or four negative charges (Figure 1d). A reason for this could be that 161 ferri- and ferrocyanide structurally contain six negative charges, which are countered by the ferric or ferrous cation to reach a net charge state of -3 and -4, respectively. This means that R10 can effectively 162 163 form more interactions with the iron complexes than with pyrophosphate, for example. Additionally, the 164  $\pi$ -electrons of the cyanide ligands could contribute to the coacervate stability by interacting with the sp<sup>2</sup> 165 hybridized guanidinium group of arginine, while the lone pairs on the nitrogen can form hydrogen bonds. 166 All these factors make ferri- and ferrocyanide coacervates incredibly stable compared to other small 167 molecule coacervates.

168 Finally, it should be noted that the CSCs discussed here are determined from R<sub>10</sub>/metabolite 169 coacervates, and replacing R<sub>10</sub> with a shorter polycation such as R<sub>5</sub> would decrease the CSC of the coacervates or completely prevent LLPS from occurring.<sup>33</sup> In addition to R<sub>10</sub>, we also investigated the 170 171 stability of  $K_{10}$ /metabolite coacervates to determine the difference between arginine and lysine in 172 complex coacervation (Table S1). K<sub>10</sub> was not able to form coacervates with any of the tricarboxylic acids 173 or sulfates used in this study. The CSCs of all  $K_{10}$ /phosphate coacervates were significantly lower than 174 R<sub>10</sub>/phosphate coacervates and TMP, AMP and NADH did not phase separate with K<sub>10</sub> while they do with 175  $R_{10}$ . These results confirm the consensus that arginine has a higher propensity to phase separate as a 176 result of the stronger cation- $\pi$  and additional  $\pi$ - $\pi$  interactions of arginine's guanidinium group compared to the amine group of lysine.<sup>24, 34, 35</sup> Taken together, these results show how metabolites can form stable 177 178 coacervate protocells with a relatively short polycation.

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#### 180 Active formation of coacervates by aldol addition

181 We then asked if compartmentalization of these metabolites is compatible with their conversion by metabolic reactions, which is a prerequisite for a primitive metabolism in coacervate protocells. We 182 183 expect that metabolism and coacervate compartments are mutually dependent: metabolic reactions 184 could affect the coacervates by inducing phase separation or affecting their stability, but coacervates can also affect the rates of metabolism, analogous to the interplay between cellular processes and 185 membraneless organelles.<sup>36,37</sup> Here, we show that conversion of metabolites can induce phase separation, 186 187 that the conversion of metabolites can be enhanced inside coacervates, and that metabolic reactions 188 taking place inside coacervates can affect their stability and ultimately lead to their dissolution.

Metabolic reactions of the compounds shown in Figure 1 can drive phase separation if the product of the reaction has stronger interactions with another species, such as R<sub>10</sub>. Previous work has shown that the enzymatic conversion of ADP to ATP, which involves the introduction of an additional negative charge, can induce phase separation.<sup>38, 39</sup> Although the use of enzymes is far from prebiotically relevant, reactions involving the addition or removal of charged groups are ubiquitous in prebiotic pathways as well. Several reactions in the tricarboxylic acid (TCA) cycle and its prebiotic analogues introduce additional carboxylate

195 groups.<sup>4,5</sup> To determine whether such prebiotic reactions could give rise to active coacervate formation, we 196 investigated the aldol addition reaction of 90 mM of the diacid malonate (9) with 120 mM glyoxylate (10) to form the triacid 3-carboxymalate (3-CM, 4), shown schematically in Figure 2a.<sup>5</sup> 3-CM should be able to 197 undergo LLPS with 5 mM R<sub>10</sub>, according to our results in Figure 1. Analysis by <sup>1</sup>H-NMR spectroscopy (Figure 198 199 2b, SI Figure S1) shows formation of 3-CM after 143 hours in a 22.8% yield. Similar yields were obtained in 200 absence of  $R_{10}$  (SI Figure S2-3). Nucleation of coacervate droplets could be seen after 59 hours under the 201 microscope at a concentration of 11 mM 3-CM, after which the coacervate droplets continued to grow 202 (Figure 2c, Supplementary Movie 1). Figure 2d shows the increase in average droplet size over time, which resembles previous observations of actively growing coacervate droplets.<sup>39</sup> A similar result was obtained for 203 204 the aldol addition reaction between  $\alpha$ -ketoglutarate and glyoxylate, which forms the tricarboxylic acids 205 isocitroyl formate and aconitoyl formate (SI Figure S4-7). These results show that metabolite-containing 206 coacervate protocells can be formed through prebiotically relevant metabolic conversions.



209 Figure 2 Active formation of coacervates. (a) Aldol addition reaction between dianion malonate (9) and glyoxylate 210 (10) generates 3-carboxymalate (3-CM, 4). The trianionic product can undergo LLPS with R<sub>10</sub>. (b) Plot of 211 concentrations over time for the reaction of 90 mM disodium malonate with 120 mM sodium glyoxylate in the 212 presence of 5 mM R<sub>10</sub> as observed by <sup>1</sup>H-NMR spectroscopy with 10 mM 3-(trimethylsilyl)propionic-2,2,3,3-d<sub>4</sub> acid 213 sodium salt internal standard. (c) Brightfield microscopy images for the reaction of 90 mM disodium malonate with 214 120 mM sodium glyoxylate in the presence of 5 mM R<sub>10</sub>, showing the formation of coacervates as the reaction 215 progresses. (d) Analysis of droplet size over time. Droplet area was converted to radius and mean droplet radius was 216 calculated. Shaded areas indicate the standard deviation over all droplets in the frame (blue) and detection limit 217 resulting from the threshold in droplet size analysis script (grey).

### 218 Oxidation of NADH is enhanced inside coacervates

219 After the observation of coacervate formation by conversion of malonate to 3-CM, we investigated if the 220 newly formed protocells could facilitate reactions involving metabolites. Recently, it was shown that simple coacervates can increase the rate of an aldol reaction<sup>18</sup>, which inspired us to investigate the 221 222 influence of coacervates on a simple redox reaction. NADH was selected as reducing agent because of the prebiotic plausible synthesis and importance in metabolic networks,<sup>40, 41</sup> and the possibility to monitor it 223 224 by fluorescence microscopy and absorbance. NADH was sequestered in the R<sub>10</sub>/3-CM coacervates formed 225 in the previous section with a partitioning coefficient ( $K_p$ ) of 91 ± 4.6, meaning that the concentration 226 inside the coacervate droplets is nearly two orders of magnitude higher than the supernatant (Figure 3a). 227 We hypothesized that the locally increased concentrations inside coacervates can increase the rate of 228 NADH-mediated reduction reactions (Figure 3b). We studied the conversion of NADH to NAD $^+$  by 229 ferricyanide, an anionic oxidizing agent with a strong tendency to phase separate and a possible presence on prebiotic Earth.<sup>29</sup> The reaction was followed by measuring the absorbance at 340 nm (SI Figure S8), for 230 231 30 minutes after the addition of a stoichiometric amount of ferricyanide. A sample containing L-arginine 232 instead of R<sub>10</sub> was used as control reaction without coacervates present.

233 We observed a significantly and consistently faster decrease in absorbance at 340 nm for the 234 reaction containing R<sub>10</sub>/3-CM coacervates compared to the control reaction, indicating that NADH 235 oxidation by ferricyanide is enhanced by their accumulation in coacervate droplets. The initial reaction 236 rate  $(v_0)$  was determined from a linear fit to the first three minutes of the reaction after addition of 237 ferricyanide (SI Figure S9). The initial rate of oxidation of compartmentalized NADH was 1.4 times higher than that of NADH in the control reaction, 31.8 and 23.4  $\mu$ M \* min<sup>-1</sup>, respectively. The fact that 238 239 coacervates can increase the rate of redox reactions has been shown before, but has thus far been limited to enzyme driven systems.<sup>42</sup> To verify that the  $v_0$  increase is due to compartmentalization of NADH and 240 241 not a difference in pH between the coacervates and surrounding solution, the pH was measured over the 242 course of the reaction. For both conditions, the pH slightly dropped during the reaction, but the difference 243 between the phase separated and dilute reaction was never greater than 0.1 (SI Figure S10) These results 244 show that coacervates can sustain and increase NADH oxidation, a process crucial for electron transport

- in biology. This is important, as chemical activity would have been essential for protocells and raises the
  possibility that similar coacervates composed of other metabolites shown in Figure 1 or mixtures of
- 247 metabolites could have acted as microreactors in a prebiotic environment.



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249 Figure 3. Oxidation of NADH is enhanced by coacervate droplets. (a) Microscopy pictures of R<sub>10</sub>/3-CM coacervates 250 containing 1 mM NADH in brightfield (left) and excited with a 405 nm laser (right). (b) NADH oxidation by ferricyanide 251 inside coacervate droplets. (c) Oxidation of NADH by  $Fe(CN)_6^{3-}$  is increased in the presence of  $R_{10}/3$ -CM coacervates 252 (purple, filled) compared to the control reaction containing Arg/3-CM (black, open). (d) The initial rate ( $v_0$ ) at which 253 NADH is oxidized is increased in coacervate droplets. Absorbance was measured at 340 nm and a background 254 measurement containing 1mM NAD<sup>+</sup> and 2 mM ferrocyanide in R<sub>10</sub>/3-CM coacervates or Arg/3-CM was subtracted 255 from the measured data. Shaded regions and error bars represent the standard deviation (n = 3).  $v_{0}s$  are significantly 256 different (\*\*\*) in a two sample *t*-test (p < 0.001).

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### 258 Oxidative decarboxylation of α-keto acids affects stability of coacervate protocells

Finally, having found that coacervates can affect the rate of oxidation of NADH, we wondered whether, vice versa, protometabolic reactions could also affect the coacervate protocells. To this end, we investigated the oxidative decarboxylation of the  $\alpha$ -keto acids isocitroyl formate (**11**) and aconitoyl formate (**12**) to form isocitrate (**13**) and aconitate (**3**), by reacting with hydrogen peroxide (Figure 4a). This reaction leads to loss of a keto group (C=O), producing anions that are likely to form less stable coacervates with R<sub>10</sub> than their precursors (cf. Figure 1). In addition, the carboxylic acid products have a higher pK<sub>a</sub> than the  $\alpha$ -keto acid substrates and have a slightly lower net charge. And lastly, the oxidative decarboxylation produces CO<sub>2</sub>, which can dissolve in the aqueous solution and dissociate into bicarbonate and a proton, thereby lowering the pH and increasing the ionic strength of the solution. Since the stability and physicochemical properties of complex coacervates are highly dependent on ionic strength and pH of the solution, such a reaction could significantly affect the nature of the coacervate protocells.<sup>36,43,44</sup>

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272 Figure 4: Oxidative decarboxylation of isocitroyl formate and aconitoyl formate decreases coacervate stability. (a) 273 Isocitroyl formate (11) and aconitoyl formate (12) undergo oxidative decarboxylation by hydrogen peroxide (HOOH) 274 to form isocitrate (13) and aconitate (3), respectively. During the reaction, carbon dioxide is formed, which dissolves 275 in water and decreases the pH due to dissociation into bicarbonate and a proton. (b) Detail of <sup>1</sup>H-NMR spectra over 276 time of the reaction of 15.2 mM isocitroyl formate and 4.1 mM aconitoyl formate with 5.9 equivalents hydrogen 277 peroxide and 1.05 mM 3-(trimethylsilyl)propionic-2,2,3,3-d4 acid sodium salt internal standard, showing the 278 disappearance of **11** and **12** and the formation of **13** and **3**. The full <sup>1</sup>H-NMR spectra can be found in SI Figure S11. 279 (c) Decrease in critical salt concentration of coacervates made with 15.2 mM isocitroyl formate, 4.1 mM aconitoyl 280 formate and 5 mM R<sub>10</sub> upon oxidation by 5.9 eq hydrogen peroxide. Error bars represent the standard deviation. 281 Critical salt concentrations are significantly different (\*\*\*\*) in a T-test (P < 0.0001). (d) Decrease in turbidity over 282 time upon addition of 3.5 μL 10 wt% (5.9 eq) hydrogen peroxide to coacervates made of 15.2 mM isocitroyl formate, 283 4.1 mM aconitoyl formate and 5 mM R<sub>10</sub> in the presence of 235 mM sodium chloride. Blanks where the same volume 284 of Milli-Q water and 0.4 mM hydrochloric acid (resulting in a similar decrease in pH as for addition of hydrogen peroxide to water) were added as reference. The standard deviation is depicted as shading around the curves. Insert: 285 286 Coacervate droplets as observed by Brightfield microscopy.

287 We started by investigating the reaction of 15.2 mM isocitroyl formate and 4.1 mM aconitoyl 288 formate with 5.9 equivalents hydrogen peroxide in Milli-Q water by <sup>1</sup>H-NMR spectroscopy (Figure 4b), 289 and observed that the oxidation reaction is completed in less than one hour, having a similar rate in 290 absence and presence of  $R_{10}$  (SI Figure S12). As the reaction progressed, a significant decrease in pH was 291 measured, starting at 8.42 before the reaction, and decreasing to 6.10 at the end of the reaction (SI Table 292 S2). Control experiments in which an equal amount of hydrogen peroxide was added to Milli-Q water only 293 showed a decrease in pH from 6.28 to 5.66 (SI Table S3), indicating that the addition of hydrogen peroxide 294 alone is not the reason for the significant lowering of pH. Instead, the CO<sub>2</sub> produced during 295 decarboxylation of isocitroyl formate and aconitoyl formate partly dissolves and dissociates into 296 bicabonate, which causes the decrease in pH.

297 We then investigated what the effect of the reaction is on the stability of the coacervate 298 protocells. We measured the critical salt concentration of coacervates made of 15.2 mM isocitroyl 299 formate and 4.1 mM aconitoyl formate with 5 mM  $R_{10}$ , before and after the reaction with 5.9 equivalents 300 hydrogen peroxide (Figure 4c), and observed a significant decrease in CSC from 274.1 ± 3.2 mM to 230.5 301 ± 2.0 mM due to the oxidative decarboxylation reaction. Control experiments performed in 100 mM MOPS 302 buffer (pH 7.55) showed a difference in CSC of 11.8 mM (SI Figure S13), indicating that decreased 303 coacervate stability can be explained by a combination of decreased interaction strength of the reaction 304 products, isocitrate / aconitate with R<sub>10</sub>, and the concomitant decrease in pH and increase in ionic strength 305 due to bicarbonate formation. Such a decrease in CSC and change in composition of the coacervates is 306 expected to coincide with a change in physicochemical properties of the protocell, including viscosity and surface tension.<sup>45-47</sup> Moreover, at specific conditions, this protometabolic reaction could be used to 307 308 dissolve the compartments in which the reaction is taking place. To test this, we again prepared 309 coacervate samples made of 15.2 mM isocitroyl formate and 4.1 mM aconitoyl formate with 5 mM R<sub>10</sub>, 310 but now added 235 mM NaCl. This value was chosen to be in between de CSCs of the reagents and 311 products, so that reaction with 5.9 eq hydrogen peroxide would result in dissolution of the coacervates. 312 We followed the turbidity of the reaction over time by plate reader, and indeed observed a disappearance of turbidity after 5 minutes (Figure 4d), indicating that the coacervates had fully dissolved. Control experiments where an equivalent amount of Milli-Q water or 0.4 mM HCl (resulting, in pure water, in a similar pH drop as the hydrogen peroxide) was added, only showed a gradual decrease in turbidity due to settling of the coacervate droplets. This shows that metabolic reactions, such as the oxidative decarboxylation of isocitroyl formate and aconitoyl formate, can have a significant effect on the stability of the coacervate protocells, and can even drive their dissolution.

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# 320 Conclusions

321 We have shown that coacervate protocells can form by phase separation of a wide range of prebiotically 322 relevant anionic metabolites with R<sub>10</sub>. The difference in electrostatic stability of the metabolite coacervates 323 made with different anions can be explained by the number of interaction sites, such as charged or aromatic 324 groups, and the type of anion. This difference in stability can be exploited to actively form coacervates, for 325 example through the conversion of malonate and glyoxylate to 3-carboxymalate. During this reaction, we 326 observed nucleation and steady growth of coacervate droplets. These metabolite coacervates were able to 327 sustain, and even enhance reactions that were localized to the coacervate interior, such as the oxidation of 328 NADH. We found an increase in oxidation rate by 40%, which we attribute to the high local concentration of 329 NADH and ferricyanide inside the coacervates. Lastly, we have shown that protometabolic reactions can also 330 change physicochemical properties of coacervate protocells, and even decrease their stability such that they 331 dissolve. We used the oxidative decarboxylation of isocitroyl formate and aconitoyl formate by hydrogen 332 peroxide to illustrate this effect, and show that the combination of decreased interaction strength of the 333 reaction products and the decrease in pH due to carbonation of the solution leads to a sufficient lowering 334 of the stability to dissolve the coacervate droplets.

The fact that stable coacervate droplets can be formed from a short polycation and small metabolites with as few as two negative charges is significant, as it offers new insights for the development of protocells with more prebiotically plausible materials. Importantly, the coacervates can be made chemically active with protometabolic reactions that produce or consume droplet material, which opens up possibilities to use the latest insights in prebiotic chemistry to generate liquid

- 340 compartments and to link together compartmentalization and metabolism, two central elements of living
- 341 systems. A scenario in which coacervate protocells nucleate, grow and eventually dissolve with
- 342 progression of a reaction cycle would be the next step of protocell construction.
- 343

#### 344 **Conflicts of interest**

- 345 There are no conflicts to declare.
- 346

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# 444 Footnotes

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