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Catalytic Gels for a Prebiotically-Relevant Asymmetric Aldol Reaction in Water: From Organocatalyst Design to Hydrogel Discovery and Back Again

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ABSTRACT: This paper reports an investigation into organocatalytic hydrogels as prebiotically-relevant systems. Gels are interesting prebiotic reaction media, combining heterogeneous and homogeneous characteristics with a structurally-organised active ‘solid-like’ catalyst separated from the surrounding environment, yet in intimate contact with the solution phase and readily accessible via ‘liquid-like’ diffusion. A simple self-assembling glutamine amide derivative 1 was initially found to catalyse a model aldol reaction between cyclohexanone and 4-nitrobenzaldehyde, but did not maintain its gel structure during reaction. In this study, it was observed that compound 1 could react directly with the benzaldehyde to form a hydrogel in situ based on Schiff base 2 as a low-molecular-weight gelator (LMWG) This new dynamic gel is a rare example of a two-component self-assembled LMWG hydrogel and was fully characterised. It was demonstrated that glutamine amide 1 could select an optimal aldehyde component and preferentially assemble from mixtures. In the hunt for an organocatalyst, reductive conditions were applied to the Schiff base to yield secondary amine 3, which is also a highly effective hydrogelator at very low loadings with a high degree of nanoscale order. Most importantly, the hydrogel based on 3 catalysed the prebiotically-relevant aldol dimerisation of glycolaldehyde to give threonine and erythrose. In buffered conditions, this reaction gave excellent conversions, good diastereoselectivity and some enantioselectivity. Catalysis using the hydrogel of 3 was much better than using non-assembled 3 – demonstrating a clear benefit of self-assembly. The results suggest hydrogels offer a potential strategy by which prebiotic reactions can be promoted using simple prebiotically-plausible LMWGs, that can selectively assemble from complex mixtures. Such processes may have been of prebiotic importance.

Introduction

Gels are a colloidal state of matter in which a solid-like network is dispersed through a continuous liquid-like phase leading to solvent immobilisation.1 In the case of supramolecular gels, low-molecular-weight gelators (LMWGs) self-assemble via intermolecular non-covalent interactions into a nanoscale sample-spanning ‘solid-like’ network.2 Hydrogels have wide ranging potential applications, particularly as biomaterials3 and in environmental remediation.4 In some cases, gelation requires two distinct chemical components5 – such ‘two-component gels’ are tunable, but relatively rare, especially in the case of hydrogels that self-assemble from LMWGs in water. Examples include systems in which the two-components form a non-covalent complex that acts as the LMWG,6 or in which they come together to form the LMWG in a dynamic reaction, such as acylhydrazone formation.7 There are also examples of small molecules that undergo dynamic reactions like imine formation to form polymer network hydrogels,8 but such gels are more like polymer gels9 than those that self-assemble from LMWGs via non-covalent interactions. There has been interest in the development of gels for applications in catalysis.10 Gels benefit from a combination of heterogeneous and homogeneous characteristics, allowing the rapid diffusion of small molecules, like reagents and products, through the gel matrix, while potentially immobilising the catalyst by incorporation into, or interaction with, the gel nanofibres. Self-assembled organocatalytic gels have been developed,11 with a particular focus on systems in which catalysis is enhanced compared with the non-assembled organocatalyst. In landmark work, Escuder, Miravit and co-workers demonstrated the potential of gel networks to achieve a range of catalytic processes, including amino acid-mediated aldol reactions.12 They also developed self-sorted gel networks with orthogonal catalytic sites to achieve multi-step reactions in one system, demonstrating the potential of gels to act as systems for complex processes.13 Organocatalysis also plays a key role in the field of prebiotic chemistry, in which researchers aim to demonstrate mechanisms by which simple chemical building blocks, such as those present on the early earth, can be converted into the more complex chemical constituents of life.14 It is hypothesised that some of the dominant reactions of prebiotic chemistry are aldol condensations, which can result in the synthesis of sugars and complex carbohydrates.15 It has been argued that the first sugars arose from the formose reaction – an autocatalytic reaction of formaldehyde to
form glycolaldehyde, followed by aldol homologation of glycolaldehyde into higher sugars. Alternatively, it has been suggested by Sutherland that glycolaldehyde and higher sugars can be generated from formaldehyde by iterative Kiliani Fischer synthesis and amino oxazoline formation.

It was first reported that prebiotically-relevant amino acids, such as proline, could catalyse asymmetric aldol cyclisations in the early 1970s. In 2000, Barbas and List studied proline and other primary and secondary, cyclic and acyclic amino acids as general aldol catalysts proposing a six-membered enamine Zimmerman–Traxler transition state. In key work, Pizzarello and Weber showed the dimerisation of glycolaldehyde by \(\alpha,\alpha\)-disubstituted amino acids led to the formation of \(L\)-tetroses in 7\% ee. Similar studies by Breslow with stoichiometric quantities of amino acids showed the formation of \(D\)-glyceraldehyde with similar ee. Dipeptides have shown improved reaction selectivity and tetrose yield, and zinc–proline complexes can catalyse an aqueous aldol reaction in which a cocktail of higher carbohydrates was produced. Barbas demonstrated the catalysis of aldol reactions on activated nitrobenzaldehydes in water without the need for co-solvents by using proline modified with a hydrophobic group. Micellar assembly of the catalyst mediates the reaction by providing a hydrophobic environment for the organic reagents. This led Janda and co-workers to critically question what was really meant by performing organocatalysis “in water” – which remains a key question in the field. They also noted that adding an acid to limit general base catalysis enhanced enantioselectivity. Clarke and co-workers went on to apply this approach to prebiotically-relevant reactions, reporting the aldol condensation of TIPS-protected glycolaldehyde in water promoted by amino acid esters. They then reported aldol reactions on unprotected glycolaldehyde to generate threose and erythrose, and demonstrates a plausible prebiotic synthesis in water of 2-deoxy-\(D\)-ribose using amino acid esters and amino nitriles as organocatalysts.

Although gels have been studied to some extent in the field of organocatalysis as described above, only Escuder and co-workers’ have performed a prebiotically-relevant reaction, using a tripeptide organogelator to catalyse the reaction between TIPS-protected glycolaldehyde, yielding TIPS-protected threose and erythrose derivatives. In this research, the catalysis failed on the unprotected glycolaldehyde. The relative lack of development of self-assembled prebiotic organocatalytic gels is surprising given that such systems can form from simple, prebiotically-plausible molecular-scale building blocks. Furthermore, gels are known to play vital roles in life itself, for example, the cytoplasm interior of a cell has a gel-like structure. It has been suggested that gels may have been an effective medium for organisation and compartmentalisation prior to the evolution of protective membranes. Gels can, to some extent, control traffic in and out of themselves, and can immobilise larger objects, or interactive ones, effectively separating them from the surrounding environment. We therefore reason that hydrogels are fascinating potential prebiotic materials – the development of gels with prebiotic catalytic potential may provide insight into how gel assembly and catalysis can cooperate in the evolution of more complex yet better organised chemical systems. Indeed, there is considerable current interest in a ‘systems chemistry’ approach to prebiotic chemistry, in which holistic systems of multiple components collaborate in achieving emergent properties – we suggest self-assembled hydrogels may play an important, and under-recognised, role in such processes. With this in mind, we decided to embark on a programme of organocatalyst and hydrogelator discovery with the hope of gaining new insights relevant to both supramolecular gels and prebiotic organocatalysis (Scheme 1). We focus here on self-assembled gels as catalysts for a specific prebiotic process – unprotected aqueous glycolaldehyde dimerisation – a key step in the pathway leading to sugars.

Scheme 1. Outline synthesis of the new gels reported here and summary of their ability to catalyse aldol reactions.
**Results and Discussion**

**Glutamine Derivative 1 – Gelation**

Preliminary research, in which the behaviour of a number of modified amino acids such as compound 1 had been screened,

suggested such compounds had the potential to become involved in aldol catalysis. Compound 1 was synthesised by coupling dodecylamine and Boc-protected glutamine using EDC and DMAP, followed by acid-mediated removal of the Boc-protecting group (Scheme 2). The product was converted into free base form using NaOH (1 M) for 2–12 h.

![Scheme 2. Synthesis of glutamine amide 1.](image)

Initially a range of solvents were used to test the gelation of compound 1 at 3% wt/vol loading using a simple tube inversion test for gel formation. Partial gels (in which the gel partially prevented dropping of the sample on inversion) were formed in acetonitrile, tetrahydrofuran and toluene, and a full gel (resistant to tube inversion) in cyclohexane. Most importantly with regard to this study, a gel was formed in water. However, this was somewhat irreproducible, which we assigned to the relatively high solubility of compound 1 in water. Rather like in crystallisation events, this can make the nucleation of a gel network somewhat temperamental on application of a heat-cool cycle. We therefore did not characterise this gel in detail, but decided instead to move straight onto testing the proficiency of compound 1 as an organocatalyst.

**Glutamine Derivative 1 – Organocatalysis**

We tested compound 1 in a standard aldol reaction, which proceeds effectively and has easy-to-analyse products – the reaction between cyclohexanone and 4-nitrobenzaldehyde (Scheme 3). Unfortunately, the gel was not reproducible or stable enough to be used as a catalyst, but to provide some insight into organocatalysis, we probed the ability of compound 1 to catalyse this reaction in a heterogeneous ‘solution’ phase.

![Scheme 3. Aldol reaction investigated using organocatalyst 1.](image)

A solution of 4-nitrobenzaldehyde (1 eq) in cyclohexanone (10 eq) was added to a solution of the amide (0.1 eq) in water (20 mL) and a heterogeneous mixture formed (see ESI Section 5 for full experimental details). Reactions were analysed after 24 h, 48 h and 72 h – each time-point was a separate reaction because the heterogeneous nature of the reaction meant aliquot sampling did not give accurate representation, and each timepoint was analysed across triplicate repeat reactions. After the allotted time, the reaction mixture was extracted with dichloromethane (DCM) and solvent removed in vacuo. 'H NMR analysis of the crude products in deuterated chloroform (CDCl₃) was used to calculate the conversion and diastereomeric ratio (dr) (see ESI Figs. S7–8, Table S1).

The reaction proceeded with excellent conversions, up to 98% after 72 h with an antisy:n ratio of 1.3:1.0. The antisy:n ratio was higher at shorter reaction times (e.g. 2.1:1.0 at 24 h). Chiral HPLC was used to determine the enantioselectivity (ESI, Fig. S9), with results indicating that the antisy product was obtained with an ee of up to 32%, while the syn product was obtained with an ee up to 11% (ESI Figs. S10–S12, Table S2). However, these ee’s varied widely from reaction to reaction, reflective of the highly heterogeneous nature. The relatively low diastereo- and enantioselectivities suggest a relatively poorly organised transition state, as might be expected given just one amino acid is in the organocatalyst. Other studies have shown dipeptides can enhance selectivity as a result of the introduction of additional non-covalent interactions, although of course, in prebiotic terms, the catalyst is then more structurally complex, and less likely to spontaneously emerge from simple building blocks. It is also worth noting co-solvents, such as DMSO can be added to homogenise reactions to improve selectivity – for prebiotic relevance we chose to avoid this approach, but in terms of basic process optimisation we suggest this would be a useful approach.

Interestingly, on catalysis of the heterogeneous reaction between cyclohexanone and 4-nitrobenzaldehyde mediated by compound 1, we consistently observed that a stable sample-spanning gel was formed during the reaction (see Scheme 1). We reasoned that this gel was the result of interactions between the different components in the system, and set out to understand this in more detail.

**Schiff Base 2 – Two-Component Gelation**

To understand gel formation, we tested compound 1 on its own and with cyclohexanone and 4-nitrobenzaldehyde individually under the reaction conditions. Only in the presence of 4-nitrobenzaldehyde was a hydrogel consistently formed *in situ*. Mass spectrometry of the gel, and analysis of the dried residue by 'H NMR spectroscopy indicated quantitative conversion of catalyst 1 into Schiff base 2 (Scheme 4). There have been previous reports of two-component gels based on dynamic, reversible imine formation, but in all cases this leads to gels in organic solvents, whereas here the gel is forming in water. As noted in the Introduction, however, two-component hydrogels based on LMWGs that self-assemble through non-covalent interactions are relatively rare.

![Scheme 4. Reaction between glutamine amide 1 and 4-nitrobenzaldehyde to give compound 2 which assembles into gels in situ in water.](image)
component gels assembled at minimum gel concentrations (MGCs) as low as 0.5 mg/ml of glutamine amide 1 and 0.24 mg/ml (1 eq.) of 4-nitrobenzaldehyde, giving a total gelator loading of 0.074% wt/vol (ESI, Table S7). This is remarkable performance for a two-component LMWG hydrogel, and allows the system to be categorised as a ‘supergelator’ (i.e. gelation at loadings <0.1% wt/vol). The maximum gel concentration was a total gelator loading of 0.74% wt/vol - above this level, solubility problems meant that not all of the material dissolved prior to gel formation, and consequently, the resulting gels were not fully homogeneous.

We then characterised these gels in more detail. Firstly, we used simple tube inversion methodology to gain insight into the thermal stability of the gel (T_gel value) formed from a 1:1 mixture of components. As expected, on increasing the concentration, the T_gel increased from 61°C (0.15% wt/vol) to 83°C (0.74% wt/vol) as the sample-spanning network becomes more fully established at higher concentration (ESI, Table S8).

Obviously, the reaction to form the Schiff base requires 1:1 stoichiometry, but we were interested to know how much aldehyde would be required to assemble a gel. Tests of gelation using 1 mg of glutamine amide 1 indicated that gels would not form with <0.7 equivalents of 4-nitrobenzaldehyde (ESI, Table S6). However, once 0.7 equivalents were present, a gel resulted – clearly at this point, sufficient Schiff base is generated to support a sample-spanning gel-network (consistent with the MGC of the gelator described above). Gels were still formed by the Schiff base even in the presence of significant excesses of 4-nitrobenzaldehyde but, once >10 eq. were present the gels became less stable.

Further analysis of the 1:1 gel was performed using ^1'H NMR spectroscopy. In gel-phase NMR, mobile components can be detected, but the self-assembled ‘solid-like’ network has broadened peaks and is thus not observed. Spiking the gel with a mobile internal standard allows quantification of any mobile components in the gel. For the 1:1 gel, we did not observe Schiff base 2 in the ^1'H NMR spectrum, indicating it is indeed assembled into the ‘solid-like’ network (ESI, Fig. S34). Neither did we observe any mobile unreacted glutamine amide 1. However, we did observe ca. 0.3 eq. of mobile unreacted aldehyde. This suggests, in-line with the observations above, that ca. 0.7 equivalents of aldehyde are sufficient to cause the system to assemble into a gel. However, the remaining ‘excess’ unreacted glutamine amide 1 is also probably assembled within the gel as it is not observed as mobile in the NMR.

We then wanted to understand the gelation process. Given that glutamine derivative 1 shows some ability to assemble in its own right, and sometimes forms gels, we performed dynamic light scattering (DLS). It was evident based on light scattering, that compound 1 is indeed self-assembled (ESI, Fig. S35). It was not possible to provide an accurate size for these assemblies, because DLS can only do this for spherical systems. Interestingly, however, on addition of 4-nitrobenzaldehyde, the emergence of a new peak in the DLS (ESI, Figs. S36) clearly indicated that the self-assembly mode changes as the two components react with one another, consistent with the formation of a more effective self-assembled gel in situ. This suggests that glutamine amide 1 assembles into nanoscale objects, with limited ability to form an interactive 3D sample-spanning network. However, on reaction with 4-nitrobenzaldehyde, the solubility changes – decreasing in water – this will enhance fibre-fibre interactions, promoting the formation of a sample spanning network. In this way, the second component drives gelation. This model of self-assembly is reminiscent of the widely-explored Fmoc-peptide derivatives – in basic conditions the anionic carboxylate assembles into nanoscale objects (cylindrical micelles), and only on protonation does the lowering of solubility drive fibrillar gel network assembly.

The ability of the isolated and dried Schiff base derivative 2 to form hydrogels directly was also tested. Effective gels could not be directly formed from 2, probably because the solubility was too low for heating and/or sonication to fully dissolve the system and overcome the energy barrier to gelation. Once again, this is reminiscent of Fmoc-peptides, in which the self-assembling carboxylic acid gelator is difficult to assemble into gels directly, and is instead generated at a controlled rate in situ by protonation of the pre-assembled free carboxylate to assemble effective gels. We argue that for our new two-component Schiff base gelator 2, the reaction between amine 1 and the aldehyde delivers the LMWG into solution at an appropriate rate for it to then self-assemble into effective gels in situ.
the gelator. The aldehyde component is achiral and has UV-active absorption bands associated with the aromatic ring (ca. 280 nm), while the glutamine amide component is chiral, but only has an absorption band at ca. 220 nm associated with the chiral amide. The results (Fig. 1, top) indicated a small CD signal associated with each of the chromophores, demonstrating that within the gel, both experience a chiral environment, i.e., the achiral aromatic aldehyde is attached to the chiral glutamine amide unit, and thus experiences an induced CD effect. The CD signal was thermally responsive, as expected for a self-assembled system (Fig. 1, bottom). On heating from 20-40°C, the CD signal, somewhat unusually, increased in intensity – a result of the gel becoming slightly more transparent (limiting light scattering). On heating further from 40°C to 80°C, however, the CD band then decreased in intensity, as would be expected for the thermally-induced disassembly of a chiral gel nanostucture. At 80°C, the system shows no CD signal, indicative of complete disassembly into isolated molecules of 2.

Transmission and scanning electron microscopy (EM) provided further insight into the nanoscale morphologies of these gels. Although drying effects can be significant in sample preparation and impact on apparent morphologies, EM can still be useful for comparing related samples prepared in the same way. We used standard methods to image the gel samples (ESI, Section 3.1) that avoid any problems associated with ice crystal formation. For SEM imaging, we applied cryo-drying in attempt to minimise morphological reorganisation during the drying step. The sample with 1 mg/ml of 1 (total loading ca. 0.15% wt/vol) had a nanofibrillar morphology as typically observed for self-assembled gels, with nanofibre dimensions 55-110 nm (Fig. 2 left, Figs. S41-S42). However, on increasing the loading, the fibres became enlarged, and additional nodular aggregates also appeared to be present (Fig. 2 right, Figs. S43-45). It seems plausible that as the loading increases, excess material has a secondary aggregation mode. Alternatively, the nodules may be associated with less soluble material as the concentration of gelator is increased.

Rheological studies (see ESI) on the combination of glutamine amide 1 and 4-nitrobenzaldehyde using parallel plate geometry indicated that the self-assembled materials behave as gels with G’>G” (Figs. S37-S40). Interestingly, the gel exhibited self-healing properties being broken down by shear during injection, and then reforming *in situ* on standing (Fig. S46, Table S1). This gives this family of gelators potential biomaterials applications – for example in drug delivery.

With our interests in prebiotic chemistry, we then substituted 4-nitrobenzaldehyde for a simpler aldehyde, benzaldehyde, which is known to be a prebiotically-plausible building block. Pleasingly, benzaldehyde behaved in an analogous manner to 4-nitrobenzaldehyde, forming gels. In general, when testing aldehydes (ESI, Table S12), we found a preference for aromatic aldehydes over aliphatic aldehydes, but if electron donating groups were present on the ring, then gelation did not occur. This preference would reflect the inherent electrophilicity of the aldehyde, and the ability of the aromatic ring to self-assemble via π–π interactions, both preferred in electron poor systems. One fascinating property of two-component gels is the potential for one component to select its ideal partner from a mixture of possibilities. Obviously this has considerable relevance in a prebiotic setting, as it enables order to spontaneously emerge from a relatively complex set of inputs. We therefore tested the assembly of the gel based on glutamine amide 1 in the presence of benzaldehyde (1 eq.) and (4-hydroxy-3-methoxybenzaldehyde, vanillin, 1 eq.). Benzaldehyde is capable of supporting gelation (Fig. 3, left), but vanillin is not (Fig. 3, middle). Interestingly, the 1:1:1 glutamine amide:benzaldehyde:vanillin mixed system still formed a gel (Fig. 3, right), suggesting benzaldehyde can dominate. We then used 1H NMR analysis on this mixed gel to determine how much of each aldehyde was being incorporated into the ‘solid-like’ gel fibres. This indicated that the gel network incorporated 75% of the benzaldehyde, but only 29% of the vanillin (Fig. 3, Figs. S47-S48). There is therefore clear component selection for the preferred aldehyde that gives effective gelation. A majority of vanillin is left in the solution phase, while a majority of the benzaldehyde becomes assembled into the gel fibres, clearly demonstrating that gel assembly can select specific components from mixtures to assemble functional materials – a principle of prebiotic relevance.
In summary, in the hunt for a catalyst for the aldol reaction between cyclohexanone and 4-nitrobenzaldehyde, we discovered a highly effective two-component hydrogel that forms at low concentrations and can select preferred components from mixtures. Clearly there is very considerable scope for tuning the performance of this class of gels by varying: (i) aldehyde, (ii) amino acid, (iii) hydrophobic chain. Full detailed results of these structure-activity relationship studies will be reported elsewhere.

**Secondary Amine 3 – Gelation**

Having developed a highly effective new two-component hydrogel based on simple building blocks, we then wanted to achieve effective organocatalysis. We reasoned that simple reduction of Schiff base 2 would yield a secondary amine that may be an organocatalyst. Furthermore, given that the perturbation to molecular structure is relatively small, we believed it may also form a hydrogel. We opted to use Schiff base 2a, based on benzaldehyde, rather than the Schiff base based on 4-nitrobenzaldehyde, as it is simpler, and more prebiotically-relevant.

To synthesise the Schiff base in high yield, we made use of its gel-forming ability to drive the reaction to completion, giving us a straightforward reaction in water (Scheme 5). We suggest that gelation may, in the prebiotic world, have its gel-forming ability to drive the reaction to completion, giving us a straightforward reaction in water (Scheme 5).

After drying the gel product 2a, we reduced the imine using sodium borohydride in methanol to give desired amine 3 in good yield (Scheme 6). There has been interest in reductive amination, specifically to convert α-keto acids into amino acids, with a number of prebiotically-plausible approaches discussed in the literature. We were delighted to find that compound 3 is a highly effective hydrogelator (Fig. 3) – even more so than the two-component Schiff Base. It also formed gels in some organic solvents (Table S13). A heat/sonication/cool cycle was used to achieve consistent and fast (ca. 2 h) gel formation, with the MGC being just 0.03% wt/vol (Fig. S49). This is even lower than the precursor Schiff base, making this a highly potent LMWG. This suggests that the secondary amine may enhance non-covalent interactions between LMWGs.

![Figure 3. Benzylglutamine amide 3 hydrogel (loading 0.1% wt/vol) and SEM image of a dried sample of the gel.](image)

The T_{gel} value of this new gelator is 50°C at a concentration of 0.10% wt/vol. Scanning electron microscopy (SEM) imaging indicates the assembly of a fibrillar sample-spanning network with diameters of ca. 40 nm (Fig. 3). These fibres are smaller than those found in the two-component gel, consistent with the formation of a more stable gel at lower concentrations. It is also consistent with the fact that the visual appearance of the gel is very different to those observed for the two-component hydrogel, being much more optically transparent, as is typical of smaller nanoscale assemblies, which are less able to scatter the incident light.

![Figure 4. Circular dichroism (CD) spectra at different temperatures of the gel formed by compound 3 (0.1% wt/vol).](image)

The gel was studied by VT CD spectroscopy (Fig. 4). The CD signal was very different to that observed for the two-component system, with a much larger band (50 mdeg) at ca. 215 nm, associated with chiral organisation of the chiral amide. This compares with much smaller CD bands (ca. 5 mdeg) for the two-component system, and supports the view that secondary amine 3 is a better organised, more effective self-assembling hydrogel. The CD spectrum is thermally responsive; by 70°C no CD signal is observed, confirming it can be attributed to LMWG self-assembly.

Analysis of the gel by 1H NMR spectroscopy (ESI, Fig. S50) indicated 100% incorporation of gelator into the ‘solid-like’

Scheme 5. Synthesis of Schiff base 2a from benzaldehyde and glutamine amide 1, using gelation to drive the reaction to completion in water.

Scheme 6. Reduction of imine with sodium borohydride to yield secondary amine 3.
gel network with no NMR resonances associated with free 'mobile' gelator being observed – again consistent with the highly organised assembly of this system.

The rheology of the gel formed by 3 at a loading of 4% wt/vol (ESI, Figs. S51-S52) indicated an effective soft gel, with $G' > G''$, and a $G'$ value of ca. 200 Pa. The gel crossover point is at ca. 3% strain. For detailed comparison we also performed rheology on precursor Schiff base gel 2a, formed from the combination of glutamine amide and benzaldehyde, under exactly the same conditions (Figs. S39-S40). In this case, the gel had a $G'$ value of 780 Pa and a crossover point at ca. 8% strain. Clearly the precursor gel is somewhat stiffer than the reduced secondary amine version of the gel. This is in-line with its more opaque appearance, indicating a greater degree of crystalline solid-like behaviour. Conversely, the secondary amine gelator 3 gives transparent, more nanoscale gels, that self-assemble at exceptionally low loadings. We were therefore excited to explore the potential use of these gels in organocatalysis.

**Secondary Amine 3 - Organocatalysis**

Initially, we tested the ability of compound 3 to catalyse the model reaction between cyclohexane and 4-nitrobenzaldehyde in the solution phase, to benchmark the organocatalyst against glutamine amide 1. In water, the conversion was ca. 43% after 72 h, the anti:syn ratio was ca. 2:0:1, with the anti product being produced in a typical ee of 15% and the syn product in a typical ee of 7% (ESI, Fig. S3, Figs. S3-S22). On performing the reaction in buffered conditions at pH 7, although the conversion dropped, the ee's rose to 55% for the anti product and 17% for the syn product (ESI, Table S4, Figs. S23-S25). Once again, we suspect better outcomes could have been achieved by adding a co-solvent, as the system is quite heterogeneous, but reaction optimisation was not the goal of our research. On testing this reaction on the gel, once the reagents were applied to the top of the gel, it broke down. We reasoned that the cyclohexane acts as an organic solvent, dissolving compound 3 and thus disrupting the self-assembled gel network.

![Scheme 7. Dimerisation of glycolaldehyde to give threose and erythrose.](image)

We therefore decided to probe the ability of this amine to catalyse a prebiotically-relevant aldol reaction that is fully compatible with aqueous-phase reaction conditions – the dimerisation of glycolaldehyde (Scheme 7). The gel was prepared in the standard way (20 mg, 49.6 μmol in 5 mL solvent), and the dimer of glycolaldehyde (59 mg, 0.49 mmol) was dissolved in water (200 μL) then rapidly added (< 1 min in total) in small aliquots (20 x 10 μL additions) to the surface of the gel, ensuring dispersion over the gel surface (ESI, Section 6). Under these conditions, the gel was ca. pH 6 during reaction. The system was monitored for 24 h to see if the gel remained intact. Initial studies were promising with the gel being stable even after 48 h.

To analyse the products, a trapping and analysis methodology was developed based on the conversion of the products to diphenyl hydrazone derivatives. Four individual standards were prepared from L- and D-threose and L- and D-erythrose by adding diphenyl hydrazine in methanol with acetic acid and stirring for 1 h (ESI, Section 4). The products were purified by column chromatography and analysed by chiral HPLC. The four enantiomers could be separated using an IC Chiral-Pak column with Hexane:IPA (90:10) at 40°C (ESI, Fig. S26).

Analysis of the dimerization of glycolaldehyde on the catalytic hydrogel was then carried out. At the end of each reaction, the hydrogel was dehydrated *in vacuo*. To the residue, diphenyl hydrazine in methanol was added followed by 1-2 drops of acetic acid and the mixture left to stir at room temperature for 1 h. The solvent was removed and $^1$H NMR analysis performed on the crude hydrazone product mixture to determine the conversion of glycolaldehyde to the erythrose and threose (ESI, Fig. S27). Column chromatography removed excess diphenyl hydrazine and glycolaldehyde, and chiral HPLC was performed on the mix of threose and erythrose hydrazone products (Fig. S26).

In the crude $^1$H NMR (Fig. S27), the triplet at 6.46 ppm corresponds to the hydrazone proton of glycolaldehyde, while the doublets at 6.49 ppm and 6.52 ppm correspond to the hydrazone proton of threose and erythrose respectively. As these peaks each integrate to one proton, the integrations can be simply used to calculate conversion and threo:erythro diastereoselectivity. When using deionised water as the solvent for the hydrogel, the conversion of glycolaldehyde into erythrose and threose was ca. 10%, with a diastereoselectivity for threose over erythrose of ca. 2-7%. In water, the conversion of glycolaldehyde into threose and erythrose increased dramatically to 43-76% (Table 1, Figs. S31-S33). Once again, in the absence of catalyst, no threose or erythrose products were observed.

The reaction was then performed using pH 7 buffer (0.01 M phosphate buffered saline, PBS) as solvent to form the gel. This allows control of pH during the reaction – in the unbuffered reaction, the pH started at ca. 6 and fell slightly during the course of reaction. Pleasingly, the conversion of glycolaldehyde to threose and erythrose increased dramatically to 43-76% (Table 1, Figs. S31-S33). Once again, in the absence of catalyst, no threose or erythrose was formed. In addition, there was still significant selectivity for threose over erythrose (ca. 2:1). Chiral HPLC again indicated a small ee in favour of L-threose (6.5%), but a change in the favoured enantiomer of erythrose, with small ee in favour of of L-erythrose (2.5%).
For purposes of comparison, we performed the dimerisation of glycolaldehyde reaction using compound 3 as a catalyst in solution, rather than in the gel phase. To achieve this, we added the same amount of compound 3 into water but omitted the heat/cool gel-forming step. Fascinatingly, with conversions of only ca. 5% either in water or in buffer, and no measurable ee’s, the reaction was very significantly less successful than when applying the catalyst in the gel form. This suggests, in-line with some other literature reports, that the highly organised nanostructure of the gel plays a key role in helping achieve a better reaction outcome. We reason that the nanofibrillar solid-like gel network enhances reactivity and helps ensure effective contact between the organocatalyst and the aqueous phase reagents.

In summary, hydrogelator 3 is catalytically proficient in a prebiotically-relevant aldol reaction in the gel phase, and can achieve excellent levels of conversion, along with diastereoselectivity and some enantioselectivity – the first time this has been demonstrated for a self-assembled LMWG in an unprotected aqueous-phase prebiotic reaction.

Although the enantioselectivity is relatively low, it is in line with other reports for this reaction. Furthermore, mechanisms for enantio-enrichment are well-known in the pre-biotic literature. A number of these rely on preferential removal of one enantiomer, which is, at least in principle, possible in a chiral gel, where one enantiomer may preferentially interact with the gel network, becoming effectively isolated from the surroundings and hence enriched. Our reaction work-up and characterisation methodology did not allow such effects to be observed here, but we suggest it as a potential advantage of gels in prebiotic catalytic systems that is worth further study.

Conclusions and Future Perspectives

In summary, this investigation into organocatalytic gels used simple prebiotically-relevant building blocks and reactions. In this way, we reported a self-assembling glutamine amide derivative 1, capable of organocatalysis of the reaction between cyclohexanone and 4-nitrobenzaldehyde in ‘solution’, but which could not maintain its gel structure during reaction. In our studies of this simple reaction, we found that compound 1 could itself, on reaction with benzaldehydes, form an effective two-component Schiff base hydrogelator in situ. This versatile, dynamic two-component gel was characterised in some detail as two-component hydrogels remain rare.

In the ongoing hunt for an effective organocatalyst, we then applied reductive conditions to benzaldehyde-modified Schiff Base 2a, to yield secondary amine 3. This minimal perturbation to the molecular structure yielded a highly effective and well-organised hydrogelator, active at loadings as low as 0.03% wt/vol. Most importantly, this new gel was catalytically proficient for the prebiotically-relevant dimerisation of glycolaldehyde to give threose and erythrose. The reaction proceeded with good diastereoselectivity, some enantioselectivity, and when the reaction was performed in buffered conditions, excellent conversions. The reaction using the gel-phase catalyst was much more successful that when the catalyst was used in the solution phase, suggesting the well-organised nanoscale environment, bringing the catalytic LMWG into intimate contact with the solution phase reagents, is beneficial for organocatalysis.

This report combines catalyst design with gelator discovery (and vice versa) to generate simple new functional gels with great potential for a range of applications, as well as effective performance in aqueous phase unprotected prebiotic reaction processes.

In terms of the prebiotic relevance of this general approach, we would highlight that:

1. LMWGs such as these are based on the self-assembly of prebiotically-plausible small molecules.
2. Gelators are preferentially selected and assembled from mixtures of components.
3. The gel catalyses an unprotected prebiotically-relevant aldol reaction proposed in the synthesis of sugars, in water, with good yield with diastero/enantio-selectivity, which are comparable with other studies.
4. Self-assembly into a gel is required in this case for effective organocatalysis, indicating a potential role for self-assembly in enhancing activity in mixtures.
5. Gels can, at least in principle, provide a unique environment, separated from the bulk, yet fully accessible to small molecules that can diffuse in and out – in this way they can be considered like ‘reaction vessels’ or even ‘primitive cells’.

| Entry | Solvent | Conversion (%) | Crude NMR d.r | HPLC ee% |
|-------|---------|----------------|---------------|----------|
|       |         |                | Ery: Thr:     |          |
| 1     | Water   | 12             | 1.00 : 2.77   | Ery: 2% Thr: 2% |
| 2     | Water   | 10             | 1.00 : 2.90   | Ery: 2% Thr: 3% |
| 3     | Water   | 7              | 1.00 : 1.38   | Ery: 2% Thr: 7% |
| 4     | pH 7    | 68             | 1.00 : 2.28   | Ery: 3% Thr: 7% |
| 5     | pH 7    | 43             | 1.00 : 1.94   | Ery: 1% Thr: 6% |
| 6     | pH 7    | 76             | 1.00 : 1.95   | Ery: 4% Thr: 10% |

Table S5: Dimerization of glycolaldehyde with benzylglutamine amide data on the hydrogel over 48h
In future work, we intend to extend the scope of gel-phase organocatalysts, while continuing to explore minimal systems that are capable of forming self-assembled hydrogels. We hope in the future to demonstrate advantages of compartmentalisation within gels. We suggest that self-assembled gels are previously overlooked materials of potential interest in the ‘systems approach’, in which a number of components can collaborate, both in terms of self-assembly and reactivity, to achieve outcomes that may have been of prebiotic importance.

ASSOCIATED CONTENT

Supporting Information
Full characterisation of novel compounds, materials and methods, details of analytical techniques, additional spectroscopic and analytical data, HPLC traces. This material is available free of charge via the Internet at http://pubs.acs.org. Reference spectroscopic, characterisation, reaction and analytical data can be found at DOI: 10.15124/f5a57bed-5fda-40fc-9bfa-a089136a01e0.

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Author Contributions
The manuscript was written through contributions of all authors. KH carried out the all of the experimental work, except for the rheology on the two-component gel and component selection experiments, which were carried out by AKP.

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