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# Sustainable and fast synthesis of functionalized quinoxalines promoted by natural deep eutectic solvents (NADESs)†

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Functionalized quinoxalines, embedding acid sensitive protecting groups, can be prepared with a fast (5 min) and high yielding (>90%) protocol using a choline chloride/water NADES at room temperature without any additional activation. The products obtained after solvent extraction do not require any further purification. The NADES used can be recycled up to six times maintaining the same efficiency level.

#### Introduction

The quinoxaline unit is embedded in a notable number of compounds showing manifold practical applications. In the field of materials chemistry, quinoxaline derivatives show liquid crystal properties,<sup>1</sup> are known as corrosion inhibitors,<sup>2</sup> and find wide use in photovoltaic devices.<sup>3</sup> On the other hand, these heterocyclic systems are also currently employed as anticancer agents,4 for their recognized antiviral and antitubercular activities,5 as well as for their many other important biomedical applications (Fig. 1).6 Over the years, remarkable interest in these quinoxaline-containing compounds has resulted in the development of a high number of protocols, which often feature a limited level of sustainability, such as those using transition metal catalyzed reactions.7 A survey of the existing procedures available for the synthesis of quinoxalines reveals that the condensation of 1,2-diamino compounds with 1,2-dicarbonyl derivatives represents the most straightforward approach to obtaining these target products.8 This simple reaction can occur even under uncatalyzed conditions in water,9 or ethanol, 10 but is limited to the utilization of poorly functionalized substrates or requires harsh conditions (Scheme 1). Better results can be generally obtained by catalyzed or promoted processes working under acidic homogeneous or

heterogeneous conditions. Evaluating the nature of various acids which can be possibly used for this reaction, acetic acid appears as the most advantageous one mainly considering its sustainability features. However, several processes entailing the use of acetic acid often require relatively high temperatures or prolonged reaction times, which considerably curtail the sustainability features of the corresponding synthetic protocol. A protic ionic liquid obtained mixing DABCO, acetic acid and water (1:1:3) has been recently used for the synthesis of quinoxalines showing some advantages over traditional procedures but a relatively high temperature (80 °C, 20–40 min) is still required for a proper reaction. In an attempt to minimize these drawbacks, we turned our attention to the use of

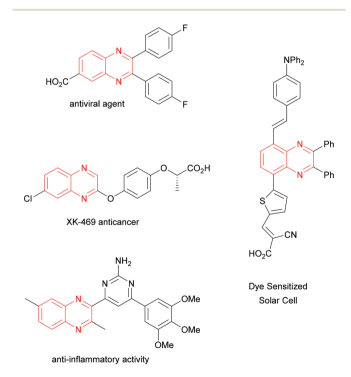


Fig. 1 Bioactive compounds featuring the quinoxaline ring.

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$$R^{1}$$
  $R^{2}$   $R^{3}$   $R^{4}$   $R^{4}$   $R^{4}$   $R^{2}$   $R^{3}$   $R^{4}$   $R^{4}$   $R^{2}$ 

a) H<sub>2</sub>O, no catalyst, rt or 230 °C, 30 to 60 min. Poor yields with diaryl 1,2-diketones and functionalized substrates.9

- b) EtOH, grinding, rt or reflux, 10 to 30 min No functionalized substrates. 10
- c) AcOH, EtOH or neat, rt (sonication) or 60-80 °C, 2 to 12 h. High temperature, long reaction time. 11
- d) This work: NADES (ChCl/water, 1:3.3), 10 M, rt, 5 min.
  - Short reaction time Functionalized substrates.
  - No purification.
     NADES recycled.
     Low E-factor.

Scheme 1 Synthetic protocols for the preparation of quinoxalines.

deep eutectic solvents (DESs), which have been recognized as formidable green media in several processes because of their low toxicity and high biodegradability. 13

A DES is defined as a mixture of compounds showing a significant depression of the melting point compared to those of the single pure components. The eutectic point represents the composition ratio at which the minimum value of the melting point is recorded. This DES composition is normally employed for synthetic and practical purposes. Additionally, DESs feature a low viscosity, a high thermal stability and a low vapor pressure, which make them ideal candidates to replace volatile organic solvents (VOCs) in several processes. Particularly, the peculiar activating properties of DESs have been demonstrated when used as selective extracting solvents from lignin and other biomass matrixes.14 Well documented synthetic applications of DESs can be highlighted in multicomponent and asymmetric catalyzed processes, 15 metal and biocatalyzed reactions, 16 and polymer and materials science. 17 According to their composition, the chemical features of DESs can be suitably tuned to provide basic, neutral or acidic environments. From a sustainable standpoint, natural deep eutectic solvents (NADESs) are of particular interest since their components are polar compounds featuring high biodegradability and biocompatibility. 18 These NADESs are usually binary mixtures of hydrogen-bond acceptors (HBA) and hydrogen-bond donors (HBD) and the more popular HBA is certainly choline chloride (ChCl), which also finds widespread use as a nutritional additive.19 Acetic acid, urea, lactic acid and water are employed as HBD together with ChCl in the preparation of various NADESs that feature different properties. Water represents a special case of HBD components in NADESs since it has been recently demonstrated that ChCl/water mixtures in the range of 1:3-4 ratios behave as a real NADES rather than a concentrated water solution of the ammonium salt.20 As a matter of fact ChCl/ water in a ratio of 1:3.33 remains liquid even at -80 °C.<sup>21</sup> In this paper we report for the first time a fast, clean and efficient synthesis of functionalized quinoxalines by condensation of 1,2-dicarbonyl derivatives with 1,2-diamino compounds promoted just by a ChCl/water NADES in 5 min at room temperature.

## Results and discussion

The stability of ChCl-based DESs appears to be due to the extended network of hydrogen bonding and van der Waals interactions involving ionic atoms and also ammonium methyl groups. In most NADESs, the activation of reactants toward the condensation is therefore supposed to conveniently occur by their embedding in this network and lowering the energy of the transition state.<sup>22</sup> The first trial using ChCl/ AcOH on the model reactants 1a and 2a gave the expected quinoxaline 3a after 5 min stirring in high yield (Table 1, entry 1). Comparable results were obtained using the combination of ChCl/lactic acid but the time required for a complete conversion was notably longer (Table 1, entry 2). A related behavior was observed employing the very popular DES ChCl/urea, which gave a complete conversion of the reactants after 1 h but required heating at 45 °C (Table 1, entry 3). Interestingly, the outstanding results achieved with ChCl/AcOH were maintained using ChCl/water, which introduced a notable improvement in the sustainability features of the protocol (Table 1, entry 4). In this context, it should be observed that ammonium halides in methanol or acetonitrile solutions have been used in the synthesis of quinoxalines but under heating or prolonged reaction times. 23 The crucial activating effect exerted by a DES was highlighted by running the reaction under neat conditions without any further activation. In this instance, compound 3a was obtained in only 50% yield together with a large amount of unreacted materials (Table 1, entry 5). In order to reduce the amount of NADES used in this process, the concentration was progressively increased by tenfold (10 M) without any significant reduction in the chemical yield (Table 1, entries 6 and 7). A further increase in the concentration up to

Table 1 Optimization studies

Entry	DES (molar ratio)	Conc.	Time (min)	Yield (%)
1	ChCl/AcOH (1:2)	0.5 M	5	95
2	ChCl/lactic acid (1:2)	0.5 M	70	92
$3^b$	ChCl/urea (1:2)	0.5 M	60	93
4	ChCl/water (1:3.3)	0.5 M	5	95
5 <sup>c</sup>	None	_	360	50
6	ChCl/water (1:3.3)	1.0 M	5	93
7	ChCl/water (1:3.3)	10 M	5	94
8	ChCl/water $(1:3.3)$	15 M	90	92

<sup>a</sup> Reaction conditions: 1a (0.3 mmol) and 2a (0.3 mmol) dissolved in DES at room temperature. <sup>b</sup> Reaction at 45 °C. <sup>c</sup> Reaction under neat conditions.

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15 M caused a notable increase in the reaction time, although the yield of the obtained product was still excellent (Table 1, entry 8).

The optimized reaction conditions were then employed to evaluate the substrate scope for the preparation of various functionalized quinoxalines 3 (Table 2). Unsymmetrical 1,2-diketones bearing an aryl framework and an alkyl framework were initially chosen for the condensation with symmetrical 1,2-diaminobenzenes leading to quinoxalines 3a-f in excellent yields.

The same pattern has been obviously observed using symmetrical 1,2-dicarbonyls with unsymmetrical 1,2-diaminobenzenes for the preparation of compounds 3g and m. As

Table 2 Substrate scope for the synthesis of quinoxalines 3 a

<sup>a</sup> Reaction conditions: diketone 1 (0.5 mmol) and diamine 2 (0.5 mmol) stirred in ChCl/water (1:3.3), 10 M for 5 min. <sup>b</sup> Reaction carried out on 5 mmol of reactants.

expected, the condensation of a couple of unsymmetrical reactants was devoid of any regioselectivity leading to an equimolar mixture of products 3k and 3k' although in excellent overall yield. The reaction of benzil with 1,2-diaminobenzene also conveniently affords product 31, showing that our procedure offers a better performance over that employing just water for the coupling reaction. $^{9a}$  At this point our goal was to demonstrate the peculiar mildness of this protocol especially when compared to procedures based on the utilization of acidbased catalysts or promoters. For this reason, a series of dicarbonyl reactants embedding acid sensitive groups, such as acetals, which are popular protecting groups of alcohols and ketones, were tested with our method. Products 3n and o containing a tetrahydropyranyl group and a 1,3-dioxolanyl group are obtained in very high yields without any trace of compounds arising from cleavage of these protecting groups. A comparison with the same reaction carried out in water shows that in spite of the neutral conditions, compound 3n is obtained in only 65% yield together with a notable amount of various byproducts (Scheme 2). In all the examples reported, the quinoxaline is obtained in pure form after extraction from the NADES with EtOAc, drying and solvent evaporation.

**Scheme 2** Comparative results using a NADES and water for the synthesis of **3n**.

Scheme 3 Green metric calculations (PMI and E-factor) for selected reactions.

**PMI** = (0.741 g + 0.541 g + 0.606 g + 4.961 g + 0.250 g) / 1.046 g = 6.78

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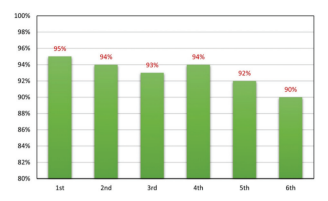


Fig. 2 Recycling of the NADES used in the synthesis of quinoxalines.

Repeating the reaction on a 5 mmol scale for the synthesis of compounds **3a** and **3e** has revealed an increase in the recorded chemical yield making this procedure rather appealing for large scale applications. For the preparation of these two compounds, green metrics such as the PMI (process mass intensity) and *E*-factor (environmental factor)<sup>24</sup> have been calculated in scaled up reactions, giving interesting values as reported in Scheme 3.

Although ChCl is a rather inexpensive material, recycling of NADES has been pursued with the aim to reduce the waste produced in our process (Fig. 2). The NADES recovered after solvent extraction by vacuum evaporation is freed of residual ethyl acetate, and then reused for a further six reactions always leading to the corresponding product 3a in good yield.

# Conclusions

Synthetic approaches to quinoxalines, although simple, are often unsuitable for the preparation of compounds embedding labile protecting groups or related functional groups. Even protocols carried out under neutral conditions show a limited substrate scope when tested with functionalized substrates. The utilization of the NADES, ChCl/water (1:3.3), provides a formidable green medium enabling the fast activation of reactants, and easy recovery of products, which are pure enough to avoid any further purification for practical use. This procedure has been used for the preparation of simple as well as functionalized quinoxalines embedding acetal protecting groups, which are totally unaffected by the reaction conditions. Thus, the devised protocol offers a cheap and competitive method from a sustainable standpoint for the preparation of quinoxalines.

#### **Author contributions**

G. L. carried out the experimental work. A. P. and M. P. conceptualized and designed the experiments.

#### Conflicts of interest

There are no conflicts to declare.

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