The Effect of Deep Eutectic Solvents as Co-solvent on Organophosphorus Hydrolase Targeting Engineering Enzyme-catalyzed

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Abstract: Deep eutectic solvents (DESs) are systems formed from one or more compounds in a mixture form, to give a eutectic with a melting point much lower than either of the individual components. DESs have attracted considerable attention as green alternative solvents to conventional solvents because they are not only eco-friendly, non-toxic, and biodegradable organic compounds, but also low cost and easy to produce and share several features and properties. DESs not only have particular properties in comparison with traditional organic solvents, but also their combination with organic solvents may also show improved and desired properties. In this study, DESs were used as the solvent in buffer containing organophosphorus hydrolase enzyme, in order to optimize the enzyme reaction buffer. Different DESs such as reline (choline chloride 2:1 urea), ethaline (choline chloride 2:1 ethanol) and glyceline (choline chloride 2:1 glycerol) were selected as an adjuvant for OPH enzyme reaction buffer. Biochemical properties, thermal stability and half-life of the OPH enzyme were studied, accordingly. In the reline, a reduction in the activity of the enzyme and in ethaline and glyceline, an increase in the activity and the stability were observed when compared with the buffer. The highest activity and stability in ethaline and glyceline in molar fraction was 0.025 and 0.25, respectively.

Keywords: Deep eutectic solvents; Choline chloride; Organophosphorus hydrolase; Reline; Ethaline; Glyceline.

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1. Introduction

Since enzymes as catalysts are involved in biochemical reactions which speed up the rate of the reactions, therefore the selection of a suitable solvent is considered as a critical issue [1]. Recently, non-aqueous solvents have been used to promote enzymatic processes rather than aqueous solvents [2, 3]. Over the past two decades, engineering of enzymatic solvents and enzymes has been studied extensively. Determining factors that cause fundamental and influential enzymatic changes are the key elements in enzyme processes which include solvents and effective conditions for enzyme activity such as pH, aqueous activity, surfactants and ionic liquids etc., that are used to increase the efficiency of enzymatic processes [4, 5]. Therefore, these factors alter the nature of the reaction buffer dramatically. In the present research, we discuss the effect of deep eutectic solvents (DESs) as a novel class of solvents [6, 7]. The published reports indicate the potential of ionic liquids as solvents for enzymatic reactions. In
particular, in bioconversion processes compared with conventional organic solvents, enzymes in ionic liquids own higher activity, better stability and more selectivity which indicates the positive effect of these solvents on enzymatic reactions [8]. Bioactive reactions can occur in ionic liquids [9, 10], hence, the catalytic activity, stability and the structure of enzymes in DESs have been broadly investigated [11]. DESs are a group of ionic liquid [12] solvents composed of a mixture of cheap polar molecules (liquids or solids) which the melting temperature drops sharply when combined [6, 9, 13].

Solvent and solvent interactions in the solvent mixture are much more complex than pure solvents due to the possibility of preferential solvation and solvent-solvent interactions [14]. Therefore, DESs can be used as solvents in a mixture of binary or triple solvents to increase the efficiency of processes and to promote the physical and chemical properties of solvents [15].

Organophosphorus compounds (OPs) are among the class of highly toxic substances, which are used as pesticides and insecticides, as well as nerve agents. The widespread use of these compounds increases public concern due to its adverse effects on human health and the environment by contaminating the soil, sediments and groundwater [13, 16, 17]. Given the recent advances in the biological sciences, the use of microorganisms to destroy these toxic substances as an economical and constructive tool compared to other common approaches seems appropriate [18, 19]. The Organophosphorus hydrolase enzyme is capable of hydrolyzing various OPs [4, 13, 16]. However, the thermal stability is low and the enzyme's profile significantly limits its industrial applications [20].

Regarding the importance of OPH as an important and effective enzyme in the biological system, as well as the tendency to examine the enzymatic behavior in non-aqueous media and especially solvents with suitable environmental characteristics, the OPH enzyme’s behavior were investigated in three different media. The media contained three most commonly used solvents with DES, including reline (choline chloride 1:2 urea), ethaline (cholinechlorid 1:2 ethylen glycole) and glyceline (choline chlorid 1:2 urea) that This particular type of solvents will provide guidance to design and develop truly green solvents with both low-toxicity and high biodegradability [12, 14-16]. This research investigated these interactions in new solvent media (such as DESs) along with their mixtures containing the buffers. In each part of this research, studies on activity, stability and half-life performed concurrently so that the results are used in order to provide a comprehensive interpretation of the changes in the way the OPH enzyme works.

2. Materials and Methods

In the present study, the effect of DESs was evaluated on the OPH function in the presence of several mole fractions of reline, ethaline and glycelin. For this purpose, in the beginning, the activity of the OPH in different mole fractions was tested, followed by thermal stability and the half-life assays in order to optimize the function of the enzyme in the reaction buffer.

2.1. Material.

Ethylene glycol (≥ 99.8%), glycerol (≥ 99.5%, spectrophotometric grade) and urea (99%) were purchased from Merck. All solvents were dried by 3Å molecular sieve. The water content of the dried materials was determined by coulometric Karl Fisher titration yielding
<500 ppm residual water, choline chloride and paraoxon were obtained from Sigma-Aldrich, choline chloride was dried at 50°C in a vacuum oven for 24 h.

2.2. Synthesis of DESs.

In order to prepare Reline, Ethaline, and Glyceline, choline chloride were mixed with urea, ethylene glycol, and glycerol as hydrogen bond donor (HBD), respectively. The material mixed and stirred while heating (near 80°C) until a homogeneous, colorless liquid formed.

2.3. Protein expression and purification.

The recombinant Rosetta-gami containing the OPH enzyme-encoding gene obtained from previous research [21]. The bacterial strain cultured in a 5 ml culture medium containing 50 μg/ml Kanamycin for 16 h. In the next step, 100 μl of pre-culture was inoculated in 100 ml of fresh culture medium containing Kanamycin and incubated to OD600 0.8 at 37°C. IPTG as the inducer added with a final concentration of 1 mM and after 5 h, the cells collected by centrifugation at 9000 rpm for 5 min and washed with 50 mM Tris-HCl buffer (pH 8). After the addition of lysis buffer (50 mM Tris-HCl, 3% TritonX-100, 8 mM EDTA, 8% sucrose), sonication was performed on ice. In order to isolate the intact cells, a 4-min centrifugation performed at 2500 rpm and then the supernatant centrifuged for isolation of the inclusion bodies with a 12000 rpm for 15 min. The washing of inclusion bodies carried out with 50 mM Tris buffer (pH 8), 3 M urea, and 8% sucrose [22]. The obtained precipitate used for protein refolding by dissolving in 8 M urea and gradually decreasing the concentration of urea [23]. Finally, the concentration of refolded protein calculated by Bradford method.

2.4. Spectrophotometric measurements.

The buffer mixture was prepared with DESs in different mole fractions (± 0.01 mg). To investigate the paraoxon degradation reaction, the spectrophotometric measurements was performed by Biochrom WPA Biowave II+ in the wavelength range of 405 nm, which is the absorption spectrum of 4-nitrophenol (as the reaction product). Other components such as solvents, reactant agents and intermediate products showed no absorbance in this wavelength range.

2.5. OPH activity assay.

To investigate the effects of DESs on enzyme activity, paraoxon was analyzed for production of 4-nitrophenol in the presence of different mole fractions of DESs containing 0, 0.025, 0.05, 0.075, 0.1, 0.25 of ethaline, glyceline and reline buffer mixtures. In order to evaluate the enzyme activity, OPH (final concentration of 3.0 × 10⁻² mg/ml) and paraoxon (final concentration of 2 mM), were added to reaction buffer (pH 8) in final concentration of 1 ml and incubated at 37°C. Finally, the activity was measured at 405 nm, followed by the detection of p-nitrophenol, produced by enzymatic hydrolysis of parathion, which creates yellow color. The enzyme activity was assayed following increasing the mole fraction of DES in the mixture. The control sample included the enzyme in the presence of buffer reaction and the rest of the activities calculated according to the control sample. Before reading the ODs, the extinction coefficients of paraoxon degradation to 4-nitrophenol calculated in the total molar fraction of reline, ethaline and glyceline. These experiments were conducted at the ideally pH-controlled environment and were repeated three times. It is worth noting that, the
activity of the OPH in the buffer in the absence of DESs, was considered as the control in order to calculate the activity. Furthermore, paraoxon hydrolysis evaluated in the presence of all mole fractions and its amount deduced from the results of mole fractions to investigate the effect of DES on spontaneously hydrolysis of parathion.

2.6. Thermostability.

As demonstrated in the previous studies, thermal destruction of OPH enzyme occurs at 65°C after several minutes [24, 25]. The thermal stability assay was performed in the presence of various mole fractions of ethaline and glyceline DESs, at 65°C. The incubation mixture contained OPH with concentration of $3.0 \times 10^{-2}$ mg/ml along with paraoxon (final concentration of 2 mM), in a volume of 1 ml of the reaction mixture (pH 8). The mixture was incubated for 15 min in the presence of different mole fractions of the DESs at 65°C. The samples were cooled on ice followed by the addition of paraoxon in order to assay the enzymatic hydrolysis at 37°C. In order to calculate and compare the activity accurately, before and after the thermal shock, the enzyme activity at 37°C, in the presence of buffer and absence of DES was set as the control. On the other hand, the stability of the OPH enzyme at room temperature (25°C) was evaluated by incubation of the enzyme in mixtures of ethaline and glyceline with buffer. Samples were assayed for their activity at different time intervals (0, 24, 50, 96, 125 and 168 h), by addition of 2 mM of paraoxon in the reaction buffer. The activity was calculated to monitor the changes in activity relative to the starting time of the enzyme-catalyzed reaction ($t_0$). In addition, the half-life ($t_{1/2}$) of the OPH enzyme was measured at 25°C. The tests presented here were performed in triplicate in order to obtain valid results.

3. Results and Discussion

Deep eutectic solvents (DESs) have already gained widespread attention to expand their applications in different areas due to their environmental and economic prospects. Hence, many enzymatic reactions have been conducted in DESs aiming to increase the activity and stability of enzymes in catalytic biological reactions [11]. Since DESs are able to affect the profile of enzyme activity and stability [26], it was decided to study these changes by designing a set of experiments involving OPH enzyme. The purpose was to represent an optimal formulation, which significantly modifies the enzyme functionality.

3.1. Enzyme activity assay.

In this study, the ability of DESs in elevating the enzyme activity was studied by measuring the progress of the OPH-catalyzed reaction in aqueous buffer solution and different concentrations of DES. The obtained results of the enzyme activity in three mixture buffers of reline, ethaline and glyceline in different mole fractions are demonstrated in figure 1. Reduced activity was observed during the reline mole fraction enhancement. Reline buffer contains urea, affecting the protein structure which results in a continuous decrease in mole fraction of 0.1 to the extent that the enzyme almost lost its activity [27]. OPH activity elevated relative to the increase of the ethaline mole fraction in the mixture, compared to the enzyme activity in the buffer. The increase in the mole fraction of $x_{Et} = 0.075$ was more noticeable than the control sample. The maximum activity was reported in the mole fraction of 0.025; however, the enzyme almost lost its activity in the mole fraction of 0.25. Upon the increase of glyceline in the buffer to the mole fraction of 0.1, OPH activity increased accordingly. The maximum
activity was observed in the mole fraction of 0.025, followed by increasing the glyceline to the mole fraction of 0.25, which caused a decrease in activity to an extent that it reached slightly near the amount of the control sample. Due to the sharp decline in the activity in mole fraction of 0.25 and more, following up the amount of activity was impossible therefore, higher mole fractions were excluded from the experiments.

**Figure 1.** Effects on organophosphorus hydrolase activity in the presence of increasing mole fraction of reline [28], ethaline [29] and glyceline (GLY) as deep eutectic solvents (Retained enzyme activity was measured, 1mg/ml enzyme concentration and 0.1 Tris-HCl buffer, pH 8.0, 2 mM paraoxon, in the presence of deep eutectic solvents (≤x = 0.25)).

Upon analyzing results obtained from previous experiments, ethaline and glyceline were chosen for further analysis. Reline was dismissed due to its denaturing effect on enzyme. The observed changes in the buffer mixtures containing ethaline and glyceline can be due to the fact that: (a) One of the known properties of DESs is the formation of a hydrogen bond between the HBD species and the halogen ion present in chlorine chloride salt. The used DESs in this study contain chloride depletion anions. However, the molar concentration of these chlorides decreases during the preparation of DES by addition of 1:2 choline chloride to glycerol and ethylene glycol (HBD), which here the glycerol and ethylene glycol have a tendency to form hydrogen bonds with chloride, therefore the likelihood to leave a negative effect on OPH enzymes is greatly reduced [30]. b) The hydrophilic nature of DESs allows the water molecules to act differently, which confers a new function to the water molecules and this effect is observed by increased activity in the mole fraction of 0.025 of ethaline and glyceline. c) In the mole fraction of more than 0.025, the DES concentrations of ethaline and glyceline were increased tremendously which led to the declination in the enzyme activity. Viscosity is regarded as a vital factor in enzyme activity. Solvent viscosity affects rate of biocatalytic chemical reactions due to mass transfer limitations [31].

Given the increase in the amount of ethaline and glyceline in the water, the viscosity of the mixture elevates and the motility of enzyme and other components decreases exceedingly. In previous studies, it has been observed that the enzyme activity decreases relative to viscosity [28, 32], which here in this study, by increasing the viscosity and increasing the mole fraction of the ethaline and glyceline in the buffer, the ability of the enzyme to conduct the hydrolysis reaction decreases and therefore the enzyme activity reduces dramatically.
3.2. Thermostability of OPH.

Improvement in the thermal stability of the enzyme is one of the outstanding effects of DESs [33]. In this study, the mole fraction of ethaline and glyceline is the indicator used to assess the OPH activity changes. Obtained amounts of activities were calculated after deducing the control amounts in order to study and observe the thermostability of OPH against high temperature. A reduction of activity occurred following the increasing mole fraction of ethaline and glyceline. The enzyme exhibited higher activity in mole fraction of glyceline, compared to ethaline, which indicates the thermal stability of OPH, and it can be concluded that the thermostability of OPH enzyme increases in the presence of the mole fraction of rich solvents of ethaline and glyceline (Figure 2).

Figure 2. Effects on OPH activity in the presence of increasing mole fraction of ethaline and glyceline after 15 min in T = 60°C to calculate thermal stability. Retained enzyme activity was measured in 1mg/ml enzyme concentration and 0.1 Tris-HCl buffer, pH 8.0, 2 mM paraoxon, in the presence of deep eutectic solvents (≤x = 0.25).

Despite the activity decline in all mole fractions, this decrease in activity can be tracked down in high mole fractions of ethaline and glyceline. The results displayed in figures 1 and 2 show that in the mole fraction of 0.25 of ethaline, only 12% of the activity was reduced while the activity in the lower mole fraction, such as 0.025 with the maximum activity, decreased by 61% and it decreased even more in the control sample by 75%. Similarly, buffer containing glyceline, the activity reduction was observed in the mole fraction of 0.25 and 0.025, which decreased by 14% and 68%, respectively. It can be concluded that the enzyme's stability in high mole fractions of DESs at high temperatures is remarkably high. As mentioned earlier, increasing the DES concentration enhances the thermal stability of OPH, which this characteristic is due to proper solvation of the enzyme by DESs, thus protecting the enzyme against the temperature.

3.3. Half-life OPH.

In this section, the sustainability of OPH in mole fraction of ethaline and glyceline along with buffer was evaluated. These compounds are known to have an effect on the stability and half-life of the enzyme [26]. The OPH enzyme in the control sample lost its activity completely after a total of 168 h. The higher the concentration of ethaline and glyceline, the lower the amount of change occurs in the activity, so that it lost only 30% of its activity in the mixture of 0.25% of ethaline after 168 h. In the mixture of glyceline with buffer in the mole fraction of 0.25 after 168 h, the enzyme lost just 15% of its activity. These results indicate the high stability of OPH in the rich mole fractions of ethaline and glyceline, therefore lower amounts of change occur in the activity, so that OPH lost only 30% of its activity (Figures 3 and 4). The half-life
of the OPH is reported in table 1 and it increases in DES solutions. In addition, OPH adopts longer half-life in DES solutions containing glyceline compared to ethaline. As a result, according to the results of the previous studies on the OPH half-life improvement [34], by applying DES solution, a new formulation of enzyme reaction can be obtained in order to enhance the stability of the enzyme.

**Table 1.** Half-life (hour) of enzyme activity during prolonged incubation of OPH in various mole fraction of ethaline and glyceline with buffer, pH 8, T = 37°C.

| Storage condition (xDES in buffer) | Ethaline | Glyceline |
|-----------------------------------|----------|-----------|
| 0                                 | 60       | 60        |
| 0.025                             | 68       | 126       |
| 0.05                              | 80       | 128       |
| 0.075                             | 96       | 137       |
| 0.1                               | 165      | 169       |
| 0.25                              | >170     | >170      |

**Figure 3.** Relative activities of OPH in the presence of aqueous solutions containing glyceline mixtures and dried paraoxon. Constant amounts of enzyme concentration (1 mg/ml) and paraoxon (2 mM) were used, T=25°C.

**Figure 4.** Relative activities of OPH in the presence of aqueous solutions containing ethaline mixtures and paraoxon, Constant amounts of enzyme concentration (1 mg/ml) and paraoxon (2 mM) were used, T=25°C.
4. Conclusions

This study was conducted to evaluate the effect of DES on thermal stability and half-life of OPH enzyme in order to improve the enzyme function. In the lower mole fractions of ethaline and glyceline DESs with buffer, the activity of OPH increases, but as we move forward to higher DES concentrations, it is inversely proportional to activity and directly proportional to enzyme stability, but instead the stability of the OPH increases. This amount of increase in the activity and stability of the OPH in mixtures of glyceline is higher than the ethaline. It also demonstrates that DESs can leave a positive effect on the activity, but in the higher mole fractions of DESs due to an increase in factors such as viscosity, the activity drops. Nevertheless, the stability of the OPH enzyme in mole fractions of 0.25 and 0.1, in which the concentration of ethaline and glyceline is high, the half-life of the OPH enhances eminently. Since DESs are relatively easy to make, inexpensive, non-toxic and biodegradable, it seems that proper formulated DES solutions can be considered as an ideal solution in order to control the activity and stability of the OPH enzyme. With further investigations, this system can be used to develop enzyme protection and disinfectant systems for military and civilian applications.

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Conflicts of Interest

The authors declare no conflict of interest.

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