ENZYMATIC DEGRADATION OF AZO DYES USING PEROXIDASE IMMOBILIZED ONTO COMMERCIAL CARRIERS WITH EPOXY GROUPS

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ABSTRACT. The treatment of colored wastewater has been paid attention in the recent years because of the high amount of dyes, mostly carcinogenic, discharged into the water. Enzymatic degradation shows several advantages such as: shorter processing time, reduction in sludge volume and ease of controlling the process. In this work, seven different industrial dyes were tested as substrates for horseradish peroxidase. The native enzyme was covalently immobilized onto two types of epoxy-supports, different in spacer length. Among the tested dyes the highest substrate conversions were achieved for Amido Black 10 (AB10). The highest recovered activity was obtained when the epoxy-activated support with longer spacer arm was used. After ten reuse cycles for the degradation of AB10, the covalently bound peroxidase preserved about 80% of the initial activity.

Keywords: horseradish peroxidase, biocatalysis, ReliZyme™ support, covalent immobilization, dye degradation, environment

INTRODUCTION

The use of enzymes for treatment of waste waters containing dyes was intensively studied, based on the capacity of enzymes to modify the original structure of dyes and thus obtaining compounds that can be degraded or eliminated more easily [1,2,3]. A wide range of enzymes, mainly oxidoreductases, have been proven to be excellent catalysts in the wastewater purification process, due to the mild reaction conditions, the ability to transform complex substances and higher reaction rates compared to chemical catalysts [2,3,4]. Oxidoreductases are able to catalyze degradation of phenols, anilines, benzidines, chlorophenols and various heterocyclic aromatic substances [5].

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Several factors can influence the degradation of residual dyes, such as the concentration and structure of dye, the presence of other compounds, pH and temperature of the effluent [6]. When enzymes are employed for the elimination of pollutants from waste waters, the properties of the biocatalyst must be properly tuned in order to ensure the highest efficiency, considering all these factors. Plant peroxidases have been extensively studied as a useful tool for bioremediation of industrial waste. Particularly, horseradish peroxidase (HRP) is a widely used catalyst in enzymatic reactions [7]. HRP proved important advantages including high activity and selectivity, high resistance to inhibition by various compounds over a broad concentration range, as well as high operability and reliability in different treatment conditions [5]. Studies have shown that in the presence of H2O2 HRP can effectively precipitate a wide variety of organic compounds, aromatic compounds, and recalcitrant contaminants, such as dyes and phenolic compounds. HRP has been extensively used in wastewater treatments as it offers long lifetime, stability and also retention of the enzymatic activity over a broad range of temperature and pH [8].

Recently it was found that chlorophenols, labeled as “priority pollutants” by the US Environmental Protection Agency, can be removed from contaminated water by HRP. Like other native enzymes, HRP has several drawbacks such as low stability, short lifetime, high susceptibility to inactivation, and difficulty in re-using [9]. Immobilization proved to be a viable solution to overcome these drawbacks, improving the activity, stability, and recovery of the enzyme. Moreover, immobilized enzymes could be broadly applied to various reaction environments and harsh conditions [9].

The immobilization of HRP was carried out using several supports: modified chitosan, polymeric acrylamide matrix, alginate, glycidyl methacrylate copolymers, glutaraldehyde-activated aminopropyl glass beads, magnetite, ion exchange resins, and various polymers [4,11,12,13].

Immobilization by covalent binding is considered appropriate for obtaining solid-phase biocatalysts with high operational stability [14]. Alongside the physical properties of the support, the structure and reactivity of the active group attached to the carrier is essential for a successful immobilization. Epoxy activated supports are very stable, can be stored for a long time and can be applied for immobilization of the enzymes to be used in the laboratory or at industrial scale [15]. Immobilization of enzymes on supports activated with epoxy groups is achieved by covalent bonding, thus increasing the stability of the enzyme, and diminishing the possible adverse effects of organic solvents, formed 9 products or temperature on the activity of the enzyme [16,17]. Covalent bonds are formed between the support and nucleophilic groups from the enzyme structure, such as amino, hydroxy or thiol groups. By covalent immobilization on such supports, the conformational changes are greatly diminished [16].
There are reports of immobilization on epoxy-activated polyacrylamide beads, like as Eupergi C [18]. Another possibility is to use a long-chained cross-linker holding epoxy groups. A ZnO nanowires/macroporous SiO$_2$ nanocomposite was used as support for HRP immobilization by in situ cross-linking, with diethylene glycol diglycidyl ether as cross-linker [19].

Although there are several studies on HRP immobilization, the utilization of commercial epoxy-functionalized methacrylate carriers was not yet reported. Compared to many other support materials, these carriers have the advantage of reproducible properties, large-scale availability, and convenient price. The aim of the present study was to develop a protocol for the covalent immobilization of HRP on epoxy-functionalized beads and to use the obtained biocatalysts for the degradation of industrial azo dyes. The effects of temperature and pH on the enzyme activity, as well as the operational stability have been evaluated for both free and immobilized HRP, using AB10 dye as model substrate. The results may provide new opportunities for the application of immobilized enzymes in the field of environmental protection, allowing the efficient oxidative degradation of residual dyes.

RESULTS AND DISCUSSION

The selectivity of HRP for commercial azo dyes

The selectivity of the HRP was tested using the native enzyme and six substrates, at 25°C and two pH values, 3.5 and 6.0. The chemical structures of the selected dyes are presented in Figure 1S (Supplementary material). The results obtained after 24 h of reaction, shown in Figure 1, indicate that at pH 3.5 the conversions were lower compared to those obtained at pH 6.0, except for the Neutral gray dye. At pH 6, the highest conversions values (up to 80%) were obtained for Amido Black 10 (AB 10) and Acid Orange 7 (AO 7) dyes.

Considering these results, the AB10 dye was used as substrate in the following experiments.

The effect of hydrogen peroxide concentration

Recent studies have shown that for HRP there is an optimal hydrogen peroxide/dye molar ratio [2]. However, addition of excess hydrogen peroxide, after conversion has reached optimal value, favored the dye degradation process. A possible explanation for this phenomenon might be that addition of excess H$_2$O$_2$ at the beginning of the process leads to formation of a larger amount of intermediate products which inhibit the peroxidase activity, or even excess hydrogen peroxide can inhibit the activity of the enzyme [2].
As hydrogen peroxide acts as co-substrate, three concentrations were tested. The results, presented in Figure 2, show that at concentrations higher than 1.43 mM there were no changes in activity, therefore in the subsequent studies the H₂O₂ concentration was set at 1.43 mM.

**Figure 2.** The effect of hydrogen peroxide amount on the decrease of absorbance at 611 nm in time, using AB10 dye as substrate
Immobilization of HRP onto ReliZyme™ supports

The practical interest on peroxidases, illustrated by the increasing number of publications in the past years, reveals that the research towards stabilization of this enzyme still presents scientific importance. Among the available immobilization methods, a simple and rapid covalent binding was selected. Considering that the biocatalyst was designed to be used in aqueous solutions, the covalent binding could also avoid the leaching of the enzyme from the support.

Two commercial ReliZyme™ supports, with active epoxy groups and different chain length and structure of the spacer arm, were selected (Table 1). The immobilization was performed in a single step, by mixing the enzyme solution with the support for 24 h. The activity of the immobilized HRP was determined by using ABTS as substrate, as described in the experimental part. The results, presented in Table 1, indicate high protein loading values (>90%), for both support types. The specific activity and the reproducibility were higher when the epoxy-amino HFA 403 support was used. Probably, the longer spacer arm favors multipoint covalent attachment of the enzyme to the epoxy groups of the support. However, considering the high immobilization efficiencies, the characterization studies were performed for both biocatalysts and the results were compared to the native enzyme.

Table 1. Protein loading and recovered activity of HRP immobilized onto epoxy- ReliZyme™ supports

| Type of the solid support | Protein loading [%] | Specific activity [U g protein⁻¹] |
|--------------------------|---------------------|----------------------------------|
| EP 403                   | 93.5                | 21.24 ± 2.36                     |
| HFA 403                  | 99.6                | 37.24 ± 0.78                     |

Comparative characterization of the native and immobilized horseradish peroxidase

Modification of the pH profile by immobilization

pH is one of the most important factors influencing the enzyme activity, because at extreme pH values the activity can be inhibited [20]. Often, for native enzymes the highest activity is in a relatively narrow pH range, and this shortcoming can be overcome by immobilizing the enzyme [20,21]. The reactions
were carried out at 25 °C, in the presence of 1.43 mM H₂O₂, for 30 minutes, under stirring at 300 rpm. Compared to the native enzyme that showed the optimal pH at 5, the activity of the immobilized biocatalyst was improved over the whole tested pH range (Figure 3).

![Figure 3](image)

**Figure 3.** Effect of pH on the decolorization of Amido Black 10 dye by native (black color), and immobilized HRP on HFA 403 (green color) and EP 403 (turquoise color) carriers. Reaction conditions: 25°C, 30 min, 1.43 mM H₂O₂, 300 rpm

The highest activity was obtained for the HRP-HFA biocatalyst, in the pH range 4-7.

An improvement of the pH stability of HRP by immobilization was also observed by Bilal et al., when calcium alginate was used as support and glutaraldehyde as linker [23]. The activity of the immobilized HRP on calcium alginate was at least 10% higher in the range 7-9 compared to the native enzyme. Mohameda et al. immobilized HRP onto activated wool support and Wool-HRP showed broad optimum pH at 7.0–8.0 [24].

**Effect of temperature on the stability of immobilized peroxidase**

The stability of the enzymes decreases with increasing temperature, the enzyme becoming inactive at high temperatures, due to the irreversible modification of the tertiary structure. Several studies concluded that thermal inactivation can be limited by enzyme immobilization [20]. To evaluate the effect of temperature on the enzymatic activity, the native and immobilized enzymes were incubated for 1h at 25°C, 35°C, 45°C and 55°C, in buffer solution pH 5.0, then cooled at 25°C when the model substrate (AB10) was added. The assay was carried out in the presence of 1.43 mM H₂O₂, for 30 minutes at 300 rpm.
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Figure 4. Thermal stability of native (black color) and immobilized horseradish peroxidase (on HFA 403, green color; on EP 403, turquoise color). The degradation of Amido Black 10 dye was carried out at 25°C, 30 min, 300 rpm, in presence of 1.43 mM H$_2$O$_2$

The results (Figure 4) indicate up to 60% activity loss for the native HRP with increasing temperature in the studied range, while the enzyme immobilized on both HFA 403 and EP 403 demonstrated excellent thermostability (only 6.1% and 18.4 % decrease, respectively). Among the two preparations, the HRP immobilized on HFA 403 showed superior thermal stability. Compared to ~84% recovered activity value for decoloring of Acid Black dye after incubation at 55°C, reported by Sun et al. for HRP immobilized onto ZnO nanowires/macroporous SiO$_2$ and diethylene glycol diglycidyl ether as cross-linker [19], the 93.8% value obtained in this study indicates a higher thermal stability after immobilization, which could be explained by the multipoint attachment of the cross-linked HRP, improving the rigidification of the protein and protect it from denaturation [19].

Operational stability of the immobilized enzyme

To increase the operational stability of the enzymes, in order to be reused in several production cycles, is one of the main objectives of immobilization. Provided that the immobilized enzyme does not lose its activity even after prolonged operational period, it can be employed in both continuous and discontinuous processes and the costs become lower [22].

The degradation of Amido Black 10 by HRP immobilized on HFA-403 epoxy-activated beads was studied in repeated batch reaction cycles, considering the activity of the first batch as reference. After ten cycles the relative activity
remained about 80% (Figure 5), an excellent value for an enzyme working in aqueous environment. Compared to previous literature reports concerning the reuse of HRP immobilized by different methods (Table 2), the operational stability of our preparate was higher, indicating the efficiency of the selected immobilization support and method.

![Figure 5. Stability of the peroxidase immobilized on HFA 403 epoxy beads, in repeated batches of Amido Black 10 degradation at 35°C, 30 min, 1.43 mM H₂O₂, 300 rpm.](image)

**Table 2.** Comparative evaluation of several immobilized horseradish peroxidases, based on the recovered activities after repeated reaction cycles

| Immobilization method   | Support           | Linker           | Reaction cycles | Recovered activity [%] | Ref. |
|-------------------------|-------------------|------------------|-----------------|------------------------|------|
| Encapsulation           | chitosan beads    | -                | 6               | 64.9                   | [25] |
| Covalent binding        | calcium-alginate  | GA*              | 7               | 40.0                   | [23] |
| Covalent binding        | activated wool    | -                | 7               | 50.0                   | [24] |
| Covalent binding        | chitosan beads    | hydrazine hydrate| 6               | 65.8                   | [25] |
| Covalent binding        | cashew gum polysacch. | GA      | 9               | 50.0                   | [27] |
| Covalent binding        | nano-composite    | epoxy cross-linker| 12              | 71.1                   | [19] |
| Covalent binding        | ReliZyme™ HFA 403 | -                | 10              | 76.1                   | This study |

*glutaraldehyde
CONCLUSIONS

Horseradish peroxidase was successfully immobilized by covalent bonding, on two ReliZyme™ supports containing active epoxy groups. The immobilization on the HFA 403 support proved to be more advantageous, allowing probably a multipoint linkage. By immobilization, both the stability and the activity of the enzyme were improved over a wider range of pH, particularly in the pH range 5-6, which represents the typical pH of waters contaminated with dyes. In the basic domain, the enzymatic activity decreased significantly, but even at pH 11 the immobilized enzyme was still active, unlike the native enzyme which was completely inactivated.

The thermostability of the enzyme was considerably enhanced by immobilization. In the case of peroxidase immobilized on HFA 403, 90% of the initial enzymatic activity was recovered after incubation for 30 minutes at 55°C, while the native enzyme lost ~60% activity in the same conditions.

The selected immobilization method and support were also able to provide a stable end robust enzyme, suitable for several utilization cycles at high residual activity. The immobilized enzyme was successfully reused in batch decolorization processes, the activity remaining at 80% of the initial value after 10 cycles of use.

EXPERIMENTAL SECTION

Chemicals

Amido Black 10 B (AB10), Acid Orange (AO7), Acid Red (AR), Direct Green (DG), Methylene Blue (MB), Neutral Gray (NG) dyes were purchased from Merck (Germany) and Loba-Feinchemie (Austria). Sodium hydrogen phosphate, sodium dihydrogen phosphate (used to prepare the buffer solutions), hydrogen peroxide 35%, citric acid, boric acid, trisodium phosphate, and 2,2'-azino-di (3-ethylbenzothiazoline-6-sulfonate) (ABTS) were from Sigma-Aldrich Chemie GmbH (Germany). The horseradish peroxidase (HRP) used in the immobilization process and as reference native enzyme was also purchased from Merck. Both commercial supports utilized for covalent immobilization of HRP were products of Resindion S.r.l (Italy), a subsidiary of Mitsubishi Chemical: ReliZyme™ HFA 403 and ReliZyme™ EP 403 (polymethacrylate beads with oxiranyl active groups which differ by the length and structure of the spacer, oxirane content min. 30 μmol/g wet, average pore diameter 40-60 nm).
Determination of peroxidase activity using ABTS

The activity of the immobilized HRP was tested using a known method with ABTS as substrate [28]. 20 μL native peroxidase solution of (1 mg/mL), or 10 mg immobilized peroxidase, were added to 100 μL of 3 mM ABTS, followed by addition of 1498 μL citrate-phosphate buffer solution pH 5 and 40 μL H2O2. The absorbance was measured at 414 nm, after 5 minutes incubation at 25 °C, using a Jasco V-530 UV/vis Spectrophotometer (JASCO, Japan).

Effect of hydrogen peroxide concentration

The influence of the co-substrate concentration (H2O2) was studied in the 0.21-2.86 mM range using AB10 as substrate. The decrease of the absorbance at 611 nm was continuously measured for 3500 s at 25 °C, using a Jasco V-530 UV/vis Spectrophotometer.

Immobilization by covalent binding on epoxy supports

900 μl of peroxidase solution (1.5 mg / mL) in 50 mM phosphate buffer pH 6.5 was added to 100 mg epoxy-activated support. The mixture was incubated for 24 hours at 1000 rpm and 25°C. The immobilized peroxidase was separated by filtration. At the end of the immobilization, the resin was washed 3 times with 20mM phosphate buffer, pH 6.5, and stored at 4°C until subsequent use.

pH profile of the immobilized peroxidase

The effect of pH on the activity of immobilized HRP (5 mg / mL) was investigated in the range of pH 4.0-12.0, using a wide pH buffer solution containing citric acid, boric acid and trisodium phosphate buffer [29]. The pH profile of native HRP (20 μL,1 mg / mL) was determined in the same conditions, as reference. The activity assay was performed as described in previous section.

Degradation of azo dyes by immobilized HRP

Six different dyes were used to determine the decolorization percentage subsequent to oxidation with immobilized HRP at two pH values, 3.5 and 6. The dye solutions were at 10 mg/mL concentration, and the experiments were carried out in different pH buffers (1 mL buffer) at room temperature after 3 h, 12 h and 24 h, respectively. The decrease of the absorbance was monitored by using a Jasco V-530 UV/vis Spectrophotometer.
Effect of temperature on the activity of immobilized HRP

The native (20 μL native enzyme 1 mg / mL) immobilized (5 mg / mL) enzyme was incubated for 2 hours in buffer, pH 5.0 in the absence of the substrate at different temperatures, in the range of 25°C to 55°C, followed by cooling on ice for 10 minutes and AB10 substrate was added and the decrease of the absorbance was monitored by using a Jasco V-530 UV/vis Spectrophotometer after 30 min.

Repeated use of the biocatalyst

The immobilized HRP activity was determined after repeated use of the biocatalyst at 30°C for 2 hours buffer, pH 5, with Amido black 10 as substrate. After each cycle, the immobilized enzyme was separated by filtration and washed three times with 50 mM phosphate buffer pH 6.5. A new reaction cycle was performed by adding the same amount of buffer and substrate. The decolorization percentage achieved with the reused biocatalyst was determined as previously described, and the resulting value was compared to the first cycle (set as 100%).

SUPPLEMENTARY MATERIALS can be obtained by request from the authors.

REFERENCES

1. S. Nouren; H.N. Bhatti; Biochem. Eng. J., 2015, 95, 9-19.
2. F. Gholami-Borujeni; A. H. Mahvi; S. Naseri; M. A. Faramarzi; R. Nabizadeh; M. Alimohammadi; Res. J. Chem. Environ., 2011, 15(2), 217-220.
3. J. T. Chacko; K. Subramaniam; Int. J. Environ. Sci., 2011, 1(6), 1250-1256.
4. M. Monier; D.M. Ayad; Y. SAA Wei; Int. J. Biol. Macromol., 2010, 46, 324-330.
5. M. Matto; Q. Husain; Ecotoxicol. Environ. Saf., 2009, 72, 965-971.
6. R. G. Saratale; G. D. Saratale; J. S. Chang; S.P. Gowindvar; J. Taiwan. Inst. Chem. Eng., 2011, 42, 138-157.
7. G.R. Lopes; D.C.G.A. Pinto; A.M.S. Silva; RSC Adv., 2014, 4, 37244-37265.
8. L. Y. Juna; L. S. Yon; N.M. Mubarak; C. H. Bing; S. Pan; M. K. Danquah; E.C. Abdullah; M. Khalid; J. Environ. Chem. Eng., 2019, 7, 1-14.
9. X. Xie; P. Luo; J. Han; T. Chen; Y. Wang; Y. Cai; Q. Liu; Enzyme Microb. Technol., 2019, 122, 26-35
10. B. Yu; H. Cheng; W. Zhuang; C. J. Zhu; J. Wu; H. Niu; D. Liu; Y. Chen; H. Ying; Process Biochem., 2019, 79, 40-48.
11. S. Venkata Mohan; K. Krishna Prasad; N. Chandrasekhar Rao; P.N. Sarma; Chemosphere, 2005, 58, 1097-1100.
12. O. Prodanović; M. Prokopijević; D. Spasojević; Z. Stojanović; K. Radotić; Z.D. Knežević-Jugović; R. Prodanović, Appl. Biochem. Biotechnol., 2012, 168, 1288-1301.
13. J. L. Gómez; A. Bódalo; E. Gómez; J. Bastida; A. M. Hidalgo; M. Gómez; Enzime. Microb. Technol., 2006, 39, 1016-1022.
14. Z. Zhao; M. C. Zhou; R. L. Liu; Catalysts, 2019, 9 (647), 1-15.
15. C. Mateo; O. Abian; R. F. Lafuente; J. M. Guisan; Enzyme Microb. Technol. 2000, 26, 509-515.
16. C. Mateo; V. Grazu; F. Lopez-Gallejo; R. Fernandez-Lafuente; Nat. Protoc., 2007, 2(5), 1022-1033.
17. C. Mateo; J. M. Palomo; G. Fernandez-Lorente; J. M Guisan; R. Fernandez-Lafuente, Enzym. Microb. Technol., 2000, 26, 509-515.
18. L. Pramparo; F. Stüber; J. Font; A. Fortuny; A. Fabregat; C. Bengoa; J. Hazardous Mat., 2010, 177, 990-1000.
19. H. Sun; X. Jin; N. Long; R. Zhang; Int. J. Biol. Macromol., 2017, 95, 1049-1055.
20. E. Biró; D. Budugan; A. Todea; F. Péter; S. Klébert; T. Feczkó; J. Mol. Catal. B Enzym., 2016, 123, 81-90.
21. B. Sahoo; S.K. Sahu; P. Pramanik; J. Mol. Catal. B: Enzym., 2011, 69, 95-102.
22. A. Cimporescu; A. Todea; V. Badea; C. Paul; F. Peter, Process Biochem., 2016, 51, 2076-2083.
23. M. Bilal; M. N. H. Iqbal; S. Z. H. Shah; H. Hu; W. Wang; X. Zhang; J. Environ. Manage., 2016, 183, 836-842.
24. S. A. Mohameda; A. A. Darwish, R.M.El-Shishtawy, Process Biochem., 2013, 48, 649–655
25. M. Bilal; H. M. N. Iqbal; H. Hu; W. Wang; X. Zhang; Sci. Total Environ., 2017, 575, 1352–1360.
26. M. Monier, D. M. Ayad; Y. Wei; A.A. Sarhan; Int. J. Biol. Macromol., 2010, 46(3), 324-330.
27. T. M. Silva, P. O. Santiago; L. L. A. Purcena, K. F. Fernandes, Mat. Sci. Eng. C-Mater, 2010, 30, 526–530.
28. H. Cai; X. Liu; J. Zou; J. Xiao; B. Yuan; F. Li; Q. Cheng; Chemosphere, 2018, 193, 833-839.
29. W.R. Carmody, J. Chem. Educ., 1961, 38, 559-560.