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Bilirubin oxidase based enzymatic air-breathing cathode: Operation under pristine and contaminated conditions

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\textbf{A B S T R A C T}

The performance of bilirubin oxidase (BOx) based air breathing cathode was constantly monitored over 45 days. The effect of electrolyte composition on the cathode oxygen reduction reaction (ORR) output was investigated. Particularly, deactivation of the electrocatalytic activity of the enzyme in phosphate buffer saline (PBS) solution and in activated sludge (AS) was evaluated. The greatest drop in current density was observed during the first 3 days of constant operation with a decrease of ~60 $\mu$A cm$^{-2}$ day$^{-1}$. The rate of decrease slowed to ~10 $\mu$A cm$^{-2}$ day$^{-1}$ (day 3 to 9) and then to ~1.5 $\mu$A cm$^{-2}$ day$^{-1}$ thereafter (day 9 to 45). Despite the constant decrease in output, the BOx cathode generated residual current after 45 days operations with an open circuit potential (OCP) of 475 mV vs. Ag/AgCl. Enzyme deactivation was also studied in AS to simulate an environment close to the real waste operation with pollutants, solid particles and bacteria. The presence of low-molecular weight soluble contaminants was identified as the main reason for an immediate enzymatic deactivation within few hours of cathode operation. The presence of solid particles and bacteria does not affect the natural degradation of the enzyme.

\textbf{1. Introduction}

Enzymes are large, generally proteinaceous, biological molecules that catalyze specific reactions transforming substrates into products. Like all catalysts, they can increase reaction rates by decreasing the reaction activation energies. Enzymes are widely used by the scientific community mainly due to their selectivity and specificity [1]. One set of enzymes, belonging to the oxidoreductases family, can reduce oxygen into water performing the “so called” oxygen reduction reaction (ORR) [2–3]. The steps involved in the reduction mechanism of oxygen have been previously described in details [4,5].

Multi-copper oxidases (MCOs) are among the enzymes able to reduce oxygen to water [6]. The most known and heavily explored MCOs are ascorbate oxidase [7], laccase [8] and bilirubin oxidase (BOx) [9]. These enzymes have been demonstrated to perform ORR when immobilized on the surface of solid supports and extensively used as cathodes in biofuel cells [10–12]. The electrons necessary for ORR are transferred from the electrode to the enzyme directly without the utilization of electron carriers, such as mediators [13–16].

Particularly for the cathode, it has been shown that BOx has a higher ORR kinetic and durability compared to laccase and ascorbate oxidase [17].

Indeed, new concept of biological/enzymatic based electrocatalyst are actually developed because it has been previously reported that traditional inorganic ORR electrocatalyst based on carbonaceous materials [18–19], platinum [20–21] and non-platinum [22–23] group metals have very low activity toward ORR at neutral pH [23–24]. At contrary, high electrocatalytic activity in oxygen reduction at neutral pH has been demonstrate multiple times for MCOs [25–29]. Differences between Pt and BOx electrochemical activity have been previously shown [30,31,32]. In fact, at pH 7.2–7.3, BOx catalyst demonstrated higher open circuit potential (OCP) compared to platinum (~200 mV), due to the lower activation losses of the enzymatically catalyzed ORR, and generated much higher current densities than Pt, up to 500 $\mu$A cm$^{-2}$ at 0 mV vs Ag/AgCl [30].

The research on the use of enzymes as electrode components has generally focused on and limited to enzymatic fuel cells [33–34], where enzymes have been utilized at both the anode and cathode [35]. BOx-based enzymatic electrodes for applications in biofuel cells have been also deeply studied recently [36–40]. Enzymatic catalysis has been also explored in the design of biosensors due to the specific selectivity of the selected enzyme [41–43]. Lately, enzymatic cathodes have been utilized successfully in microbial fuel cells (MFCs), designing a hybrid biofuel cell with a microbial anode and enzymatic cathode [30,44–46]. Schatzle et al. [44] used a laccase cathode in a double chamber
MFC configuration, generating high power densities. Higgins et al. [45] separated the anodic and cathodic chambers with a solid polymeric membrane in their hybrid biofuel cell, with Shewanella oneidensis MR-1 at the anode and laccase at the cathode achieving a maximum power density of roughly 200 mW cm⁻² (25 W m⁻³) [45]. Bilirubin oxidase based cathode was also used in a hybrid biological/enzymatic fuel cell, with encapsulated S. oneidensis MR-1 at the anode, in order to increase the current output for powering marine sensors [46].

In that case, the hybrid fuel cell was working in marine environment [46]. At last, a single chamber membraneless MFC in a hybrid configuration with microbial mixed culture at the anode and bilirubin oxidase at the cathode was demonstrated [30], where both anode and the cathode were exposed to the same solution [30]. Despite the high output achieved, all of the above studies lack a durability tests that remained limited in hours/day time and consequently need to be addressed and reported.

Despite the superb efficiency toward ORR in neutral environment few doubts concerning applicability for long terms operational period can be raised. In fact, enzymes are subject to natural degradation over time [47], moreover, the exposure to various inorganic and organic soluble pollutants and solid matters (abiotic particles and mixed culture bacteria) can further decrease the activity of the enzymes. Usually enzyme activity on electrodes is studied in a short period of time identified in hours or few days during which the maximum activity is expressed and measured [12,17,25,30]. To the best of our knowledge, durability studies in continuous mode (more than 7 days) using MCO enzymes are missing in the current literature. Additionally, enzyme activity is mainly tested in buffer solutions [47–48], where no pollutants or suspended solids with abiotic or biotic nature has been present.

In order to evaluate the activity of BOx in a complex industrial media, in this work, the performance of a gas-diffusion bilirubin oxidase based cathode was tested continuously over time as a single electrode in two different electrolytes. Current generation of the BOx cathode was monitored in phosphate buffer saline (PBS) solution, which is the most popular electrolyte in biofuel cells and PBS with the addition of activated sludge (PBS + AS), AS was specifically selected due to the simultaneous presence of dissolved contaminants, solid non-biological particles (suspended solids (1 µm to 1 cm), colloids (10 nm to 1 µm)) and mixed consortium of various types of bacteria. Moreover, the presence of AS could simulate working conditions of MFC systems for wastewater purification. In order to separate the effects specifically due to bacteria, solid particles and dissolved inorganic pollutants, additional experiments were run with: i) an autoclaved AS + PBS (no active bacteria); ii) autoclaved AS + PBS with an enzyme encapsulated in silica-gel matrix as a physical barrier for contact with solid particles; and iii) mixed culture bacteria, “washed out” from AS, in PBS.

2. Materials and methodology

2.1. Cathode preparation

A bilirubin oxidase-based cathode was prepared as previously described [10]. Briefly, carbon cloth wet proof (30 wt.% PTFE, Fuel Cell Earth) was used as current collector and teflonized carbon black (loading 60 mg cm⁻², Vulcan XR 72 with 35% PTFE), referred as XC35, was hydraulically pressed on it (500 psi) and playing the role of gas-diffusion layer (GDL). Isopropanol (loading 40 µl cm⁻²) was then added to insert a hydrophobic/hydrophilic gradient from the outside of the GDL to the internal side facing the catalytic layer. On the inner side of the GDL, a multi-walled carbon nanotube paper (MWBP, Buckeye Composite) was placed and all together was pressed at 500 psi for 5 min. Bilirubin oxidase from Myrothecium verrucaria in quantity of 10 mg was dissolved in 1 ml of PBS (50 mM, pH 7.5), then deposited on the MWBP and left over night (at least 16 h) at 4°C to allow the enzyme immobilization [10]. Additionally, in the specified case, bilirubin oxidase cathode has been also immobilized using silica encapsulation technique in order to form a physical barrier for avoiding direct contact with solids [46]. The cathode geometric area, exposed to the solution, was 2.25 cm²

A control abiotic cathode has been prepared following the same protocol without the last step involving the enzyme addition and immobilization.

2.2. Cell configurations and test conditions

Single chamber glass MFC was modified with a lateral hole where the cathode was screwed (Fig. 1) [49]. The chamber was filled with 100 ml of different electrolytes. The solutions were made of: i) 50 mM phosphate buffer saline (PBS) solution, pH 7.5 (Fig. 1.a); ii) activated sludge (AS, 50 ml) and 50 ml of 50 mM PBS, pH 7.5 (named as PBS + AS) (Fig. 1.b); iii) autoclaved activated sludge (P, 50 ml) and 50 ml PBS, pH 7.5 (named as PBS + P) (Fig. 1.c); and iv) washed bacteria and suspended solids from 50 ml of activated sludge (B) and 50 ml PBS, pH 7.5 (named as PBS + B) (Fig. 1.d).

In order to discriminate the effect of dissolved pollutants from the effect of the solid non-biological particles, the cathodes have been encapsulated with a protective layer of silica to avoid the direct contact and interaction of solid particles with the enzyme. It has been shown previously that the silica layer is permeable to soluble compounds [46] and thus allowing studying the influence of the pollutants dissolved in the electrolyte.

Bacteria were collected from the AS by a centrifugation at 10,000 rpm for 10 min of 50 ml of activated sludge. The supernatant has been removed and PBS has been added. This procedure has been repeated 3 times in order to wash the bacteria from possible soluble pollutants.

In all cases, PBS contained 0.1 M KCl for increasing the conductivity. All the resulting solutions had similar pH corresponding to 7.5 ± 0.15 and remained similar during the experimental time.

2.3. Electrochemical measurements

The electrochemical measurements were performed in three stages: i) 1 h open circuit potential (OCP); ii) linear sweep voltammetry from the OCP to 0 mV (vs Ag/AgCl) at 0.2 mV s⁻¹ scan rate [49]; and iii) 22.5 h of chronoamperometry at 300 mA (vs. Ag/AgCl). All measurements were done using a potentiostat (Gamry 300). Three-electrode configuration was used for the electrochemical tests, as shown in Fig. 1 [50]. Platinum net, with comparable surface area to the working electrode, was used as counter electrode [49]. Saturated Ag/AgCl was used as the reference electrode and the BOx cathode was used as the working electrode. A homemade Luggin capillary was used to reduce the ohmic resistance [49].

2.4. Water analysis

For the elemental composition analysis, water samples were firstly homogenized and then filtered using 0.45 μm filter. For ICP-OES analysis, the samples were acidified using Ultra High Purity nitric acid (UHP HNO₃) by adding 2 drops for 100 ml sample. The samples were diluted to acceptable range within the calibration standards using 2% nitric acid. Samples were transferred into ICP-OES autosampler tubes and then placed for the auto sampler setup. The instrument optics was optimized using Hg and Mn view alignment for the wavelengths. The system was then calibrated using a blank and three point calibration standards diluted sequentially. Samples were then analyzed after the calibration curve was validated using QC check solutions. The data obtained were then validated, verified, and exported.

For the ions analysis, samples were analyzed using Ion Chromatography (IC). None acidified filtered samples (0.45 μm filter) were used in this analysis. The samples were diluted to acceptable range within the calibration standards using 18 MΩ water. Samples were then
transferred into IC autosampler tubes and placed in their locations for the auto sampler setup. The system was then calibrated using a blank and three point calibration standards diluted sequentially. Samples were then analyzed after the calibration curve was validated using QC check solutions. The data obtained were then validated, verified, and exported.

3. Results and discussion

3.1. BOx electrochemical response in PBS solution

The continuous operation of the BOx-based gas-diffusion cathode was first studied in PBS (pH 7.5), as a reference test with the goal of evaluating the operation of this cathode in the absence of interferences from soluble pollutants, bacteria and suspended solids. Fig. 2.a shows the current densities generated by the cathode over 45 days of a constant polarization at 300 mV vs. Ag/AgCl. A continuous decrease in the current produced by BOx-cathode was recorded over time. The current density/time dependence can be separated into three different successive regions based on the rate of current decrease that can be considered linear for each distinct region. Region I comprises the first three days of operation where a sharp decrease in the generated current was observed. The current density decreased at a rate of approximately 60 μA cm$^{-2}$ day$^{-1}$, starting from a value of ~350 μA cm$^{-2}$, and decreasing to ~175 μA cm$^{-2}$ during this initial period. Region II includes the performance from days 3 to 9, when an average current decay of ~10 μA cm$^{-2}$ day$^{-1}$ was detected. Region III comprised between days 9 and 45 where the decrease in the current densities was continuous with a rate of ~1.5 μA cm$^{-2}$ day$^{-1}$. Similar trend was achieved by BOx cathode encapsulated with silica indicating that the silica encapsulation did not have effect on the current output performance (results not showed). A linear decrease in the current density of BOx-based bio-cathode was also observed from Mano et al. who studied the operation of wired-BOx cathode [17]. Mano et al. reported loss of around 50% of cathode activity in 20 mM PBS (pH 7.4) with 0.15 M NaCl, at 37.5 °C, in 6 days, which was similar to the current decrease observed in this study [17].

To study the cathode degradation, potentiodynamic polarization curves were carried out each day after 1 h at open circuit conditions. Fig. 2.b shows polarization curves at selected days, representing the three regions with different rates of current decrease. Particularly, it can be noticed that while the OCP decreased slowly, the limiting current instead had a fast drop, especially in the initial days, stabilizing over time.
No cathodic current has been generated at 300 mV vs. Ag/AgCl in absence of BOx (control abiotic cathode). The OCP of the control electrode appeared to be 246 mV vs. Ag/AgCl therefore ORR on this electrode at potentials higher than OCP could not be performed. It has to be noted that the OCP of the BOx containing cathode has never reached the OCP of the control electrode (475 mV vs Ag/AgCl) indicating (residual or partial) BOx activity along the 45 days of operation. Singh et al. measured the electrocatalytic activity of adsorbed BOx along with the stiffness of the protein layer and concluded that structural rearrangements and water loss are the primary culprits in activity loss in these cathodes. It was also established that this effect is not a result of enzyme attachment but rather to the applied potential [47]. The latter can explain the fast current drop during the initial stages of the cathode operation.

3.2. BOx electrochemical response in PBS + AS

Continuous operational analysis has been also carried out introducing 50% of activated sludge, naturally containing pollutants, suspended solid matters and mixed culture of bacteria. This aspect is important for examining the effect of additional abiotic and biotic matters on the long-term operation of the BOx cathode. The same polarization set of experiments was carried out using the configuration shown in Fig. 1b.

The decrease in performance of the BOx cathode exposed to the activated sludge followed the same general trend as observed for the BOx cathode in clean PBS, but the current densities generated in presence of activated sludge were significantly lower (Fig. 3.a) indicating an immediate inhibition of the enzyme. In fact, at day 3, the operational characteristics of the cathode immersed in the PBS + AS electrolyte were already similar to the operational characteristics of the same cathode exposed only to PBS for 45 days. In the presence of AS, the first current drop was almost linear until day 3 with a rate of approximately 246 mV vs. Ag/AgCl indicating (residual or partial) BOx activity along the 45 days of operation. Singh et al. measured the electrocatalytic activity of adsorbed BOx along with the stiffness of the protein layer and concluded that structural rearrangements and water loss are the primary culprits in activity loss in these cathodes. It was also established that this effect is not a result of enzyme attachment but rather to the applied potential [47]. The latter can explain the fast current drop during the initial stages of the cathode operation.

3.3. Electrochemical characterization carried out combining chronoamperometry, polarization curves and open circuit potential

A comparison between the current recorded during the chronoamperometry study and the current generated during the potentiodynamic polarization of the electrode was carried out. Moreover, the changes in OCPs were monitored over the 45 days experiment (Fig. 4). The currents obtained during the polarization curves at 0 mV and 300 mV vs. Ag/AgCl were used for the comparison. Fig. 4.a shows the behavior of the BOx cathode in PBS while Fig. 4.b shows the trend of the cathode operation in PBS + AS. As it can be seen, the current obtained from the polarization curves is always higher than the one from the chronoamperometry, despite the fact that they follow similar pattern (Fig. 4). In the case of PBS, the OCP slightly decreased over time from 520 mV to 475 mV (Fig. 4.a). Moreover, it can be noticed that the OCP remained above 500 mV till day 32. To the best of our knowledge, this high cathode potential has never been recorded for any cathode working continuously at neutral conditions for 45 days. The high OCP value can be explained by the preserved BOx activity along the 45 days (Fig. 4.a). At the same time the BOx cathode in AS solution had initially high OCP close to 500 mV but the value dropped constantly until reaching 375 mV at day 45 (Fig. 4.b). Looking at the shape of the polarization curves (Figs. 2.b and 3.b) and the potential at which the diffusion limitation starts, it can be concluded that the gas-diffusion layer of the cathodes does not change over time. In fact, the slope of I/E is decreasing in Figs. 2.b and 3.b confirming that is the catalyst, mainly quantity and activity, responsible for this phenomenon. In other words, it is not a problem of mass transfer (like oxygen diffusion through the GDL or at the gas/liquid/solid interface) because the first part of the curves between OCP and 300 mV/AgAgCl are the one changing over time. In addition, it has to be pointed out that the OCP of the enzymatic cathodes never reached the low value OCP of the abiotic control electrode, which is another indication of enzyme activity along the course of the experiment.

Since the OCP indicates the existence of active enzyme and enzymatically catalyzed ORR and at the same time the current obtained from the polarization curves show preserved structure of the GDL, we can assume that the observed decrease in the cathode performance over time (Fig. 4) is due to a decreasing quantity of active enzyme units, which decrease occurs with much faster rate in presence of AS. The decrease of the cathode enzyme quantity/activity regardless of the solution can be explained by: i) enzyme natural deactivation; ii) leaching or release of enzyme from the surface to the bulk (non-covalent immobilization); iii) H2O accumulation that stops the gaseous oxygen (substrate accessibility). Our results indicate strong and fast inhibition effect of the activated sludge content on the enzymatic activity. In
presence of AS, the additional possibilities for decayed cathode performance are i) enzyme deactivation by soluble inhibitors and ii) indirect deactivation of enzyme by bacterial attachment on electrode surface. The presence of a variety of compounds in the AS (especially oxidizable compounds) can cause mixed potential at the cathode, which apparently leads to a decrease in the OCP of the electrode over time. Also, the variation of pH toward alkaline values could impact negatively the OCP but, as mentioned before, the pH of the bulk remained constant along the duration of the experiment not ensuring that the local pH at the electrode surface is not different if microorganisms colonized the surface [51–52]. The OCP values may be considered as a good indicator of enzyme activity and long-term electrode operation.

3.4. BOx electrochemical response as a function of pollutants, solids and bacteria presence

Activated sludge is a complex matrix, which contains not only dissolved organic and inorganic compounds but also bacteria and solid non-biological particles. In order to discriminate the effects of the dissolved compounds in the AS from those of suspended solids and bacteria, three additional experiments were performed (Fig. 5). The first experiment used autoclaved activated sludge mixed with PBS as electrolyte to eliminate the effects of the bacterial activity and study the influence of both, the soluble pollutants and the solid particles in the sludge (named as PBS + P). The second experiment used, BOx enzyme encapsulated in a silica gel matrix on the electrode surface. This test was conducted in PBS + AS (named as PBS + P encaps). The silica gel plays the role of a physical barrier and prevents the interaction of the enzyme with the solid particles and bacteria present in the AS and allows the soluble pollutants to penetrate the silica matrix and interact with the enzyme. Thus only the contribution of soluble pollutants can be studied. Encapsulation of biological specimens in silica matrix has been demonstrated to prolong the activity of the biological species and at the same time allows the penetration of water and ions [10]. The third experiment used a re-suspension of mixed culture bacteria and solid particles in a phosphate buffer (named as PBS + B). The latter allows monitoring the influence of bacteria on the cathode performance.

The current generated by the BOx cathode in activated sludge (PBS + AS) and autoclaved activated sludge with inactivated bacteria (PBS + P) was identical. Furthermore, the utilization of silica encapsulation (PBS + P encaps) for protecting the enzyme from a direct contact with the solid particles and bacteria did not change the current output that resulted similar to PBS + AS and PBS + P. This indicates that the presence of bacteria along with the solid particles in the electrolyte did not have a negative effect on the cathodes activity and the reason for the decreasing over time performance of the enzymatic electrodes was more probably due to soluble pollutants present in the AS.

The decay of current densities over time for the BOx cathode exposed only to PBS and to bacteria in presence of suspended solids (PBS + B) were also similar but higher than the rest of the experiments carried out (Fig. 5). The latter confirmed that neither bacteria nor the present solid particles in the AS negatively affected the current output of the BOx cathode. The reduction in the overall enzyme activity by the autoclaved activated sludge occurred at virtually the same rate as by live activated sludge as a result of the dissolved pollutants in both. Analysis of the autoclaved activated sludge + PBS solution were carried out to determine the elemental and ion content (Table 1).

![Fig. 4. OCP, current from the chronoamperometry at 300 mV vs. Ag/AgCl and current from the polarization curves at 300 mV and 0 mV for the BOx cathode in PBS (a) and PBS + AS (b).](image)

![Fig. 5. BOx cathode performance (current density) over 9 days when polarized with constant potential of 0.3 V vs. Ag/AgCl using PBS (red), PBS + AS (blue), PBS + P (pollutants, black), PBS + P with encapsulated BOx cathode (gray) and PBS + B (“washed” bacteria, green) as electrolyte. Only one replicate is shown in the figure.](image)

As expected, many different moieties were identified in the PBS + AS solution (Table 1). High concentrations of phosphate, potassium and chloride were due to the PBS + KCl solution added. Other detected anions were contributed by the activated sludge, including nitrate, sulfate and fluoride (Table 1). The presence of sulfate in an activated sludge has been previously reported [53–56], and it was suggested to be one reason for the deactivation of platinum cathodes utilized in MFC for wastewater treatment and electricity production [55–56]. It is also well known that the presence of halogens impacts the activity of mult copper oxidases [17]. The negative effect of halogens is far more pronounced when laccase is used [17]. The order of inhibitory effect of halogens on the enzymatic cathode has been found to be the following: F⁻ > Cl⁻ > Br⁻ [17]. Fluoride anion was reported to...
completely inhibit laccase activity, while the inhibition effect of chloride was less pronounced [17]. It has been widely discussed that the strong inhibition effect of $F^-$ is due to attachment of this anion to the T2/T3 center, blocking the internal electron transfer from the T1 center to the TNC [17,57–58]. It was suggested that contrary to fluoride, $Cl^-$ binds to the T1 center blocking the access to the substrate-binding pocket and suppressing electron transfer carried out through the utilization of mediators, but when direct electron transfer occurs, the influence of the chloride is small. Negligible influence of $Cl^-$ in the range of 0–1 M on the activity of BOx was demonstrated by Mano et al. [17]. The concentration of $Cl^-$ added in the electrolyte in our experiment was 0.1 M, proposing that the $Cl^-$ most likely is not the ion decreasing the cathodes output. Mano et al. also observed that 0.5 M fluoride addition decreased the current density of BOx cathode about 30%, and about 90% was lost at fluoride concentrations above 1 M [17]. The concentration of $F^-$ in AS used in the current study was determined to be 0.9 M. This concentration could impose huge negative effect on the BOx activity as previously demonstrated [17]. Recently, it has been shown that also hydrogen peroxide has a negative influence on the BOX performance [59–60]. Detailed investigation needs to be done to study the effect of each pollutant on the activity of BOX and the designed enzymatic cathode. At the end of the experiments, macroscopic images of the cathode surfaces exposed to the electrolyte were taken (Fig. 6). As expected, no biofilm formation was detected on the BOX cathode exposed to PBS for 45 days (Fig. 6.a) and to the autoclaved AS for 9 days (Fig. 6.d). In fact in both cases, the color remained black as for the origin cathode. Biofilms clearly had developed after 45 days in contact with the electrolyte containing active AS (Fig. 6.b) and also after 9 days on the BOx cathode in contact with resting bacterial cells and suspended solids (Fig. 6.c). Despite all these macroscopic deposits, there were no phenomena of transportation limitations observed as indicated by the polarization curves carried out and as we already discusses, no negative effect was occurring due to the presence of bacteria and solid particles.

4. Conclusion

Continuous operation of bilirubin oxidase cathode was investigated by placing the cathode in “clean” conditions and exposed to activated sludge. The cathodes in PBS only and in PBS supplemented with mixed culture of bacteria demonstrated higher performance than the cathodes in contact with soluble pollutants, naturally present in the activated sludge. OCP higher than 500 mV vs. Ag/AgCl over 32 days have been observed. The results demonstrated that the lost in the enzyme catalytic activity was due to the presence of dissolved inhibitors. Biofilm formation on the cathode and the presence of suspended particles did not affect the OCP rate. Generally, the OCP of the BOx cathode was a very significant parameter, and it may be used as an indicator of the enzyme activity.

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