Co-immobilization and stabilization of xylanase, β-xylosidase and α-L-arabinofuranosidase from *Penicillium janczewskii* for arabinoxylan hydrolysis

César Rafael Fanchini Terrasan a,*, Lara Trobo-Maseda a, Sonia Moreno-Pérez a, Eleonora Cano Carmona b, Benedix Costa Pessela c, José Manuel Guisan a

a Departamento de Biocatálisis, Instituto de Catálisis y Petroleoquímica (ICP), Consejo Superior de Investigaciones Científicas (CSIC), Campus Universidad Autónoma de Madrid (UAM), Cantoblanco, 28049 Madrid, Spain

b Biochemistry and Microbiology Department, Biosciences Institute, Univ. Estadual Paulista—UNESP, PO 199, 13506-900 Rio Claro, SP, Brazil

c Departamento de Biotecnología y Microbiología de Alimentos, Instituto de Investigación en Ciencias de los Alimentos (CIAL), Consejo Superior de Investigaciones Científicas (CSIC), Campus Universidad Autónoma de Madrid (UAM), Cantoblanco, 28049 Madrid, Spain

A R T I C L E   I N F O

Article history:
Received 30 November 2015
Received in revised form 23 February 2016
Accepted 25 February 2016
Available online 3 March 2016

Keywords:
*Penicillium janczewskii*
Xylanase
β-xylosidase
α-L-arabinofuranosidase
Enzyme co-immobilization
Xylan hydrolysis

A B S T R A C T

Differently activated agarose-based supports were evaluated for co-immobilization of a crude extract from *Penicillium janczewskii* containing xylanase, β-xylosidase and α-L-arabinofuranosidase activities. Adequately selecting support and immobilization conditions (8 h, using agarose with 10% crosslinking) increased enzyme levels substantially, mainly in relation to the xylanase (2-fold). A coating with dextrans aldehyde MW 6000 Da, partially oxidized, covalently attached the enzymes to the support. Optimum activity was verified in the pH range 2–4, and at 50, 65 and 80 °C for the xylanase, α-L-arabinofuranosidase and β-xylosidase, respectively. The xylanase was highly thermostable retaining more than 70% of activity even after 24 h incubation at 60 and 70 °C; and at 80 °C its half-life was 1.7 h. The half-lives of the β-xylosidase and α-L-arabinofuranosidase at 50 °C were 2.3 and 3.8 h, respectively. The co-immobilization of the enzymes on a single support give raise to a functional multi-enzymatic biocatalyst acting in the complete hydrolysis of different and complex substrates such as oat spelt and wheat arabinoxylans, with xyllose yield higher than 40%. The xylanase and the α-L-arabinofuranosidase presented high stability retaining 86.6 and 88.0% of activity after 10 reuse cycles.

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1. Introduction

Xylan is the second most abundant biopolymer in plant cell walls and the main hemicellulosic polysaccharide. It is composed of a β-(1 → 4) d-xylopyranosyl backbone substituted at various degrees by side chain residues such as glucopyranosyl, 4-O-methyl-d-glucurono-pyranosyl, α-L-arabinofuranosyl, acetyl, feruloyl, and/or p-coumaroyl [1]. The precise composition of the polymer is strongly dependent on plant species and tissue. For instance, hard wood xylans often have d-glucuronic acid attached to their backbone, whereas L-arabinose is the most common branch in cereal xylans [2].

Given the diversity of xylan structures, their complete and efficient hydrolysis involves the synergistic action of main chain degrading enzymes, including *endo*-β-1,4-xylanases (EC 3.2.1.8) and *β*-d-xylosidases (EC 3.2.1.37), and side chain cleaving enzymes, including *α*-d-arabinofuranosidases (EC 3.2.1.55), *α*-glucuronidases (EC 3.2.1.139), acetyl xylan esterase (EC 3.1.1.72), and feruloyl esterases (EC 3.1.1.73). Endo-β-1,4-xylanase and *β*-d-xylosidase are the main enzymes responsible for the degradation of the polymer: xylanases cleave the internal β-(1 → 4) bonds in the xylan backbone, liberating different chain-length-(substituted) xyloligosaccharides, and β-xylosidases are exoglycosidases that release xyllose from the non-reducing ends of these xyloligosaccharides. β-xylosidases are critical for the systems since they carry the greatest work load in terms of number of glycosidic bonds cleaved, as well as in relieving product inhibition of xylanases [3]. Among other accessory enzymes, α-L-arabinofuranosidases are

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* Corresponding author at: Departamento de Biocatálisis, Instituto de Catálisis y Petroleoquímica (ICP), Consejo Superior de Investigaciones Científicas (CSIC), Campus Universidad Autónoma de Madrid (UAM), Cantoblanco, Calle de Marie Curie, 28049 Madrid, Spain.

E-mail addresses: cesarterrasan@hotmail.com, cesarterrasan@gmail.com (C.R. Fanchini Terrasan).

http://dx.doi.org/10.1016/j.procbio.2016.02.014
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exo-type enzymes that catalyze the cleavage of the terminal α-L-arabinofuranosyl residues from arabinosylated substrates. Xylan hydrolysis is not aleatory, i.e., the degree of substitution in xylan influence the products of hydrolysis for xylanases [4]. In this sense, accessory enzymes such as α-L-arabinofuranosidases are important since the removal of side-chain residues from xylan backbone may have a synergistic effect with the other xylanolytic enzymes [5] and also the difference in substrate specificity among different xylanases has important implications in the deconstruction of xylan [4].

Enzyme immobilization poses as a possibility to improve the characteristics of an enzyme in terms of stability and catalysis, as well as for process improvement allowing the reuse of the biocatalyst for many operational cycles [6]. Immobilization of more than one enzyme on the same support, however, is especially challenging, as it has to preserve the catalytic activity of all enzymes involved in the system and ideally improve their stability [7]. Many xylanolytic enzymes have been individually immobilized by different methods; and some studies have investigated the co-immobilization of two xylanolytic enzymes [8]. In this sense, the xylanase and β-xylanosidase from Talaromyces thermophillus were co-immobilized on chitosan and employed for the hydrolysis of oat spellet xylan, demonstrating the synergistic action of both enzymes by increasing the saccharification of the substrate [9]. In another study, co-immobilization of recombinant xylanase and α-L-arabinofuranosidase ongoloyxyl agarose was evaluated through different approaches in the hydrolysis of arabinoylan [10]. The effect of xylanase, β-xylanosidase and α-L-arabinofuranosidase from Aspergillus oryzae in the decomposition of arabinoylan was verified using the soluble enzymes in the moromi mash during soy sauce fermentation [11], nevertheless, co-immobilization of three xylanolytic enzymes acting cooperatively in the complete hydrolysis of complex substrates has not been reported to date.

This way, the aims of this work were to establish a protocol for simultaneous co-immobilization of the xylanase, β-xylanosidase and α-L-arabinofuranosidase from Penicillium janczewskii present in the crude extracellular extract, as well as improve the stabilization of the immobilized enzymes via post-immobilization techniques. After that, the immobilized enzymes were biochemically characterized and evaluated in the hydrolysis of arabinoxylans.

2. Materials and methods

2.1. Materials

Agarose with 4, 6 and 10% of cross-linking BCL were purchased from Agarose Bead Technologies (Madrid, Spain). p-nitrophenyl β-D-xylopyranoside (pNPX), glycidosil, potassium tetraborate tetrahydrate, sodium borohydride, sodium periodate, ethylenediamine, glutaraldehyde, Leuconostoc spp. dextran (MW 6000–100,000), polyethyleneimine (PEI, MW 1300), oat spellet and beechnwood xylans were obtained from Sigma-Aldrich Co (St. Louis, MO). β-xylene Assay Kit, xylole, p-nitrophenyl α-L-arabinofuranoside (pNP4ara) and low viscosity wheat arabinoxylans were from Megazyme ( Wicklow, Ireland). All reagents were of analytical grade.

2.2. Methods

2.2.1. Microorganism, enzyme production and preparation of enzyme extract

P. janczewskii (CRM 1348) is deposited in The Central of Microbial Resources, CMR-UNESP, Brazil. The microorganism was maintained on Vogel solid medium [12] and liquid cultures were prepared in the same medium with brewer’s spent grain as substrate, under optimized conditions for xylanolytic enzymes production [13]. After cultivation, the mycelium was removed by vacuum filtration and the culture filtrate was centrifuged (10,000g, 4°C, 15 min). The supernatant was dialyzed overnight against distilled water, 0.025 M sodium acetate buffer pH 5.0 or 0.025 M sodium phosphate buffer pH 7.0 before immobilization. A sample of the supernatant was also treated with 0.01 M sodium periodate for 1.5 h in order to oxidize sugar moieties of the enzymes and then dialyzed against 0.025 M sodium phosphate buffer pH 7.0.

2.2.2. SDS-PAGE

A sample containing 50 µg of protein prepared from the extracellular extract obtained under optimized conditions for xylanase production (medium with oat spellet xylan, pH 6.5, 7 days of cultivation, 30°C) was applied to SDS-PAGE performed in 8–18% (w/v) gradient gels, according to Laemmli [14]. The resolved protein bands were visualized after staining with 0.1% Coomassie brilliant blue R-250 dissolved in methanol, acetic acid, and distilled water (4:1:5 v/v/v). Standard proteins (Sigma) were phosphorylase b (97 kDa), bovine serum albumin (66 kDa), ovalbumin (45 kDa), carbonic anhydrase (29 kDa), trypsin inhibitor (20 kDa), and α-lactalbumin (14.2 kDa).

2.2.3. Enzyme assays

Xylanase activity was determined according to Bailey et al. [15] with 1% (w/v) beechwood xylan prepared in 0.05 M sodium acetate buffer pH 5.0 (before determining optimum pH) or pH 4.0 (after determining optimum pH) and appropriately diluted enzyme solution. Reducing sugars were quantified with DNS acid reagent [16], β-xylanosidase and α-L-arabinofuranosidase activities were determined in a reaction mixture containing, respectively, 3 mM NPX and pNP4ara prepared in 0.05 M sodium acetate buffer pH 5.0 (before determining optimum pH) or pH 4.0 (after determining optimum pH) and appropriately diluted enzyme solution to 1 mL final volume. Reactions were stopped by adding 1 mL of a saturated potassium tetraborate solution and the absorbance was measured at 405 nm [17]. One unit of activity was defined as the amount of enzyme required to release 1 µmol of product equivalent per min in the assay conditions at 25°C.

2.2.4. Preparation of support

Monoaminoethyl-N-4aminoethyl (MANAE)-agarose, was prepared as described elsewhere [18] and the glutaraldehyde-agarose support was prepared from MANAE agarose [19, 20]. Briefly described, we used 10 g of MANAE in 20 mL of 0.5 or 15% (v/v) glutaraldehyde solution prepared in 0.2 M phosphate buffer pH 7.0. The suspensions were kept under mild stirring at 25°C for 1 h and the cases of the supports activated with 0.5 or 15% (v/v) glutaraldehyde, respectively. This treatment permitted to fully modify the primary amino groups of the support with one or two glutaraldehyde molecules, respectively [19]. After that, the supports were filtered and washed exhaustively with 0.025 M sodium phosphate buffer and then with distilled water. Glyoxyl-agarose was prepared with the maximal activation degree, as previously described [21]. Polyethyleneimine (PEI) and dextran sulfate [22] and heterofunctional amino-glyoxyl and amino-epoxide [23,24] supports were prepared as described elsewhere.

The supports were initially prepared using agarose with 4% crosslinking, MANAE and 0.5% (w/v) glutaraldehyde supports were further prepared using agarose with 6 and 10% crosslinking. Activated supports were stored at 4°C and, before use, washed with incubation buffer according to immobilization condition.

2.2.5. Enzyme immobilization

Immobilizations were performed by suspending 1:10 (w/v) the activated supports in the dialyzed/diluted enzyme solution. Buffers
for each immobilization were: 0.025 M sodium phosphate buffer pH 7.0 for MANAE, PEI, glutaraldehyde, amino-epoxide and amino-glyoxyxyl supports, 0.05 M sodium acetate buffer pH 5.0 for dextran sulfate support, 0.1 M sodium bicarbonate buffer pH 10.0 for glyoxyxyl support (enzyme solution dialyzed against distilled water was 2-fold diluted in the buffer). Immobilizations were carried out under gentle agitation at 25 °C for 4 h or overnight incubation. During immobilization, samples of the suspensions and the supernatants were withdrawn and enzyme activities were measured. Proteins were measured during immobilization time-course and for immobilization for different periods.

2.2.6. Post-immobilization techniques

Dextran-coated derivatives: a mass of 0.5 g of the uncoated glutaraldehyde (Glut) derivative (immobilization carried out by 8 h) was added to 5 mL of aldehyde-dextran suspensions, pH adjusted to 7.0. Dextran with MW of 1500, 6000, 25,000 and 75,000 Da completely oxidized were initially evaluated [25]. Dextran with MW of 6000 with 20 and 40% degree of oxidation were further evaluated. The suspensions were generally agitated overnight, the derivative was then re-suspended in sodium borate buffer pH 8.5 or sodium phosphate buffer pH 7.0 and reduced by adding 1 mg/mL sodium borohydride. The suspension was gently agitated for 30 min, washed abundantly with water and vacuum filtered.

PEI-coated derivative: a mass of 0.5 g of the uncoated Glut derivative (immobilization carried out by 8 h) was added to 5 mL of 5% (w/v) PEI MW 1300 Da solution, pH was adjusted to 7.0. The suspension was gently agitated overnight, washed abundantly with water and vacuum filtered.

Glutaraldehyde cross-linked derivative: a mass of 1.0 g of the uncoated Glut derivative (immobilization carried out by 8 h) was added to 10 mL of 0.5% (w/v) glutaraldehyde solution pH adjusted to 7.0. The suspensions were gently agitated for 30 min, washed abundantly with water and vacuum filtered. A mass of 0.5 g of this derivative was directly reduced with 1 mg/mL sodium borohydride and 0.5 g was previously coated with dextran, as described above, and then reduced with 1 mg/mL sodium borohydride. The suspensions were washed abundantly with water and vacuum filtered.

2.2.7. Immobilization parameters

Immobilization yield was defined as the ratio between the activities (or protein) in the supernatant compared to the activity (or protein) in the control. Expressed activity was defined as the ratio of the activity in the final suspension after the immobilization process and the initial enzyme activity.

2.2.8. Evaluation of the attachment between enzyme and support

The glutaraldehyde agarose 10 BCL derivative (before and after coating with aldehyde dextran) was incubated in 0.005 M sodium phosphate buffer pH 7.0 with 0.5 M NaCl at 25 °C. After 1 h, enzyme activities were analyzed in the supernatant, as previously described.

2.2.9. Derivative characterization

2.2.9.1. Thermal stability. A mass of 0.1 g of the Glut derivative coated with dextran MW 6000 and 40% oxidation degree was suspended in 1.0 mL of 0.05 M acetate buffer pH 5.0 and incubated at 50 °C. In all cases, samples of the suspension were withdrawn at several intervals and the activity was assayed as previously described. Residual activity was calculated as the ratio between activity at a given time and the activity in the beginning of incubation (regarded as 100%).

2.2.9.2. pH stability. A mass of 0.1 g of the Glut derivative coated with dextran MW 6000 at 40% degree of oxidation was suspended in 1.0 mL of 0.05 M glycine HCl buffer pH 3.0, 0.05 M sodium acetate buffer pH 4.0 and 5.0, and 0.05 M sodium phosphate buffer pH 7.0. The suspension was incubated at 50 °C and after 4 h residual activity was assayed. Initial activities before incubation were regarded as 100%.

2.2.9.3. Optima pH and temperature. Optimum pH was determined by assaying enzyme activities of the Glut derivative coated with dextran MW 6000 at 40% degree of oxidation at 25 °C at various pH from 2.0 to 8.0. The following buffers were utilized: 0.05 M glycine-HCl pH 2.0 and 3.0, 0.05 M sodium acetate pH 4.0 and 5.0, 0.05 M sodium phosphate pH 6.2 and 7.0, and 0.05 M Tris HCl pH 8.0. Optimum temperature was determined by assaying enzyme activities at temperatures ranging from 40 to 85 °C, with 5 °C intervals, in 0.05 M sodium acetate buffer pH 5.0.

2.2.10. Hydrolysis of arabinoxylans

The hydrolysis of 0.5% (w/v) oat spelt xylan (arabinose residues ≤10%, glucose residues ≤15%, xylose residues ≥70%) and low viscosity wheat arabinoxylan (38/62 arabinose:xylose relation) were carried using the Glut derivative coated with dextran MW 6000 at 40% degree of oxidation. The reactions were carried out in 0.05 M sodium acetate buffer pH 4.0 at 40 °C. Collected samples were filtered and the adequately diluted supernatant was analyzed for xylose.

2.2.11. Reuse assay

Successive hydrolysis cycles of 0.5% (w/v) wheat arabinoxylan prepared in 0.05 M sodium acetate buffer pH 4.0 were performed using the Glut derivative coated with dextran MW 6000 at 40% degree of oxidation, at 1:10 proportion (w/v). Each cycle was carried out at 40 °C for 1 h under magnetic stirring (300 rpm). At the end of the cycles, the derivative was filtered, washed with 0.05 M sodium acetate buffer pH 4.0 and new substrate was added for a new reaction round. Samples of the supernatant were withdrawn during the first, the fifth and the tenth cycles, filtered and analyzed for reducing sugars. After the fifth and the tenth cycles the derivative was suspended in the washing buffer (1:10, w/v) and residual activities were measured, as previously described (activities before the first cycle were regarded as 100%).

2.2.12. Analytical methods

Protein concentration was determined with the modified Bradford’s method, with bovine serum albumin as standard [26]. Reducing sugars were quantified with DNS acid reagent, with xylose as standard [16]. Xylose was quantified using the enzymatic D-xylose Assay Kit. The first reaction involves the conversion of the α- to the β-anomeric form of D-xylose catalyzed by xylose mutarotase. The β-D-xylose was then oxidized by NAD+ to D-xyonic acid in the presence of β-xylose dehydrogenase. The amount of NADH formed in the reaction is stoichiometric with the amount of D-xylose. NADH was measured by the increase in absorbance at 340 nm (ε = 6300 M⁻¹ cm⁻¹). Samples of hydrolyzed oat spelt and wheat arabinoxylans were adequately diluted and analyzed according to the supplier instruction, in duplicate, and expressed as mean value. Xylose yield was calculated using 0.88 as the conversion factor of pentose to equivalent xylan, as below:

\[
\text{Xylose yield} (\%) = \frac{\text{xylose released (g)}}{\text{initial xylan (g)}} \times 0.88 \times 100
\]

3. Results and discussion

3.1. Enzyme immobilization

P. janczewskii was isolated from soil of the Brazilian Rainforest [27] and characterized as an excellent producer of xylanolytic
Table 1
Co-immobilization of xylanase, β-xylosidase and α-L-arabinofuranosidase from P. janczewskii on different agarose-based supports.

<table>
<thead>
<tr>
<th>Derivative</th>
<th>Condition/strategy</th>
<th>Xylanase</th>
<th>β-xylosidase</th>
<th>α-L-arabinofuranosidase</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Yield (%)</td>
<td>Expressed activity (%)</td>
<td>Yield (%)</td>
<td>Expressed activity (%)</td>
</tr>
<tr>
<td>MANAE</td>
<td>pH 7, 4h</td>
<td>53.4</td>
<td>20.0</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>pH 8.5, overnight</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Glutaraldehyde 0.5%</td>
<td>pH 7, 24h, sodium periodate treatment</td>
<td>100</td>
<td>167.5</td>
<td>100</td>
</tr>
<tr>
<td>Glutaraldehyde 10%</td>
<td>pH 7, 4h</td>
<td>63.2</td>
<td>20.8</td>
<td>100</td>
</tr>
<tr>
<td>PEI 1300 0.5%</td>
<td>pH 7, overnight</td>
<td>100</td>
<td>1.2</td>
<td>100</td>
</tr>
</tbody>
</table>

Enzyme activities were assayed in 0.05 M sodium acetate buffer pH 5.0 at 25 °C. Initial loads: xylanase—64 U; β-xylosidase—23 mU; α-L-arabinofuranosidase—34 mU; proteins—5.3 mg. Supports were prepared using agarose with 4% crosslinking.

Table 2
Co-immobilization of xylanase, β-xylosidase and α-L-arabinofuranosidase from P. janczewskii on different agarose beads MANAE and Glut supports.

<table>
<thead>
<tr>
<th>Derivative</th>
<th>Agarose beads (% CL)</th>
<th>Expressed activity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Xylanase</td>
<td>β-xylosidase</td>
</tr>
<tr>
<td>MANAE</td>
<td>4</td>
<td>20.0</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>14.6</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>14.0</td>
</tr>
<tr>
<td>Glut</td>
<td>4</td>
<td>20.8</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>23.1</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>26.1</td>
</tr>
</tbody>
</table>

Immobilization was carried out in 0.025 M sodium phosphate buffer pH 7.0 for 4 h at 25 °C. Enzyme activities were assayed in 0.05 M sodium acetate buffer pH 5.0 at 25 °C. Initial loads: xylanase—64 U; β-xylosidase—23 mU; α-L-arabinofuranosidase—34 mU; proteins—5.3 mg. CL—crosslinking. MANAE—monooaminoethyl-N-aminooethyl, Glut—0.5% (w/v) glutaraldehyde activated support.

Fig. 1. SDS-PAGE (8–18%) of the crude extracellular extract from P. janczewskii. Lane 1: standard proteins, phosphorylase b (94 kDa), bovine serum albumin (67 kDa), ovalbumin (43 kDa), carbonic anhydrase (30 kDa), trypsin inhibitor (20 kDa) and α-lactalbumin (14.4 kDa); Lane 2: crude extracellular extract (50 μg). Modified from [28].

enzymes in the absence of cellulases [13]. Production of xylanase, β-xylosidase and α-L-arabinofuranosidase has been previously optimized [28,29] but other enzymes from the xylanolytic complex may also be produced by this fungal strain since SDS-PAGE revealed the presence of several extracellular proteins (Fig. 1) [28]. Due to their potential application in many bioprocesses, several activation protocols, techniques and strategies were evaluated in order to optimize the concomitant immobilization of the crude extracellular xylanase, β-xylosidase and α-L-arabinofuranosidase on agarose-based supports, which have been widely used for immobilizing different enzymes [30].

In general, good results were observed in relation to the β-xylosidase immobilization, although the xylanase and the α-L-arabinofuranosidase were hardly immobilized on the supports (Table 1). Good balance for the immobilization of the three enzymes was obtained with ionic immobilization on MANAE agarose. The use of pH 8.5 and more prolonged immobilization period did not improve the activities on this support. When the enzymes were previously treated with sodium periodate to oxidize saccharide moieties of the proteins, the three enzymes were easily attached to the anionic MANAE support. This fact indicates that the presence of carbohydrates may be blocking the access of the enzymes to the reactive groups of the support or the formed aldehyde can readily react with these groups forming imine or Schiff bases. The removal of glycosylation was good in terms of immobilization; nevertheless, the enzymes became unstable even during mild conditions of enzymatic assays (not shown). The use of MANAE agarose cross-linked with 0.5% (w/v) glutaraldehyde also resulted in good enzyme immobilization. When this support was activated with higher glutaraldehyde concentration (10%, w/v), which results in the formation of glutaraldehyde dimers, improved immobilization yield of the xylanase was observed, probable consequence of the higher glutaraldehyde concentration, which allowed more attachments between enzyme and support. Nevertheless, these attachments may have been excessive, causing distortion in the xylanase structure resulting in the very low expressed activity. Expressed activities of the β-xylosidase and α-L-arabinofuranosidase increased by increasing glutaraldehyde concentration that may be related to the fact that these enzymes usually have higher MW than the xylanases or correspond to multimeric enzymes [31,32]. In this case, the higher glutaraldehyde concentration provided more attachments stabilizing the structure of the immobilized enzyme or provided enough attachments to stabilize all enzyme subunits, thus resulting in the higher expressed activity. Immobilization of enzymes on glutaraldehyde-activated supports has been largely used on supports previously activated with amine groups. Immobilization is promoted through a two-step mechanism: in a first step the enzyme is adsorbed on the support via an anionic exchange mechanism and then, the covalent immobilization occurs [20,33].

The enzymes could not be immobilized on PEI (Table 1), dextran sulfate and glyoxyl activated supports (not shown). In the case of glyoxyl support, the enzymes probably present low stability in pH
Table 3
Co-immobilization of xylanase, β-xylosidase and α-1-arabinofuranosidase from P. janczewskii for different periods on Glut agarose 10% crosslinking.

<table>
<thead>
<tr>
<th>Incubation (h)</th>
<th>Protein yield (%)</th>
<th>Xylanase</th>
<th>β-xylosidase</th>
<th>α-1-arabinofuranosidase</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Yield (%)</td>
<td>Expressed activity (%)</td>
<td>Yield (%)</td>
</tr>
<tr>
<td>4</td>
<td>78.1</td>
<td>79.8</td>
<td>26.1</td>
<td>94.3</td>
</tr>
<tr>
<td>8</td>
<td>83.1</td>
<td>80.1</td>
<td>41.8</td>
<td>100</td>
</tr>
<tr>
<td>16</td>
<td>84.3</td>
<td>81.7</td>
<td>11.2</td>
<td>100</td>
</tr>
</tbody>
</table>

Immobilization was carried out in 0.025 M sodium phosphate buffer pH 7 to 25 °C. Enzyme activities were assayed in 0.05 M sodium acetate buffer pH 5.0 at 25 °C. Initial loads: xylanase—64 U; β-xylosidase—23 mU; α-1-arabinofuranosidase—34 mU; proteins—5.3 mg.

Table 4
Post-immobilization strategies evaluated on co-immobilized xylanase, β-xylosidase and α-1-arabinofuranosidase from P. janczewskii on Glut agarose 10% crosslinking.

<table>
<thead>
<tr>
<th>Step</th>
<th>First coating</th>
<th>Sodium borohydride reduction</th>
<th>Second coating</th>
<th>Sodium borohydride reduction</th>
<th>Residual activity (%)a</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PEI</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>Xylanase</td>
</tr>
<tr>
<td></td>
<td>Dextran</td>
<td>*</td>
<td>−</td>
<td>−</td>
<td>66.9</td>
</tr>
<tr>
<td></td>
<td>Glutaraldehyde</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>39.6</td>
</tr>
<tr>
<td></td>
<td></td>
<td>*</td>
<td>−</td>
<td>Dextran</td>
<td>33.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>13.8</td>
</tr>
</tbody>
</table>

Immobilization was carried out in 0.025 M sodium phosphate buffer pH 7 to 25 °C. Enzyme activities were assayed in 0.05 M sodium acetate buffer pH 5.0 at 25 °C. Step present (+) or absent (−).a In relation to the expressed activities of the derivative prepared by 8 h incubation (Table 3).

Fig. 2. Time-course of total protein, xylanase, β-xylosidase and α-1-arabinofuranosidase from P. janczewskii co-immobilization on Glut agarose 10% crosslinking. Immobilization was carried out in 0.025 M sodium phosphate buffer pH 7.0 at 25 °C. Enzymatic activities were assayed in 0.05 M sodium acetate buffer pH 5.0 at 25 °C. Initial protein (5.3 mg), and activities (xylanase—64 U; β-xylosidase—23 mU; α-1-arabinofuranosidase—34 mU) in the supernatant were regarded as 100%. Relative (%) protein (+), and xylanase (▲), β-xylosidase (●) and α-1-arabinofuranosidase (■) activities.

Fig. 3. Thermostability of xylanase, β-xylosidase and α-1-arabinofuranosidase from P. janczewskii co-immobilized on MANAE-Glut derivative coated with MW 6000 and 40,000 dextrans (completely oxidized). Immobilization was carried out in 0.05 M sodium phosphate buffer pH 7.0 at 30 °C. Enzymatic activities were assayed in 0.05 M sodium acetate buffer pH 5.0 at 25 °C. Initial activities were regarded as 100%. Full symbols: derivative coated with dextran MW 6000. Empty symbols: derivative coated with dextran MW 40,000. Residual (%) xylanase (▲), β-xylosidase (●) and α-1-arabinofuranosidase (■) activities.

10. a required condition for optimal immobilization in this support [21]. In the case of heterofunctional amino-glyoxyl and aminooxipoxide supports (not shown) the enzymes were not immobilized, remaining in the supernatant even after pH increase or long-term incubation.

Considering the previous results, a comparison between the immobilization on MANAE and 0.5% (w/v) glutaraldehyde (further called Glut agarose) prepared with different agarose beads, i.e., 4, 6 and 10% of crosslinking, was carried out (Table 2). The expressed activity gradually increased by increasing the agarose crosslinking in the Glut derivative, whereas in the MANAE derivative the expressed activity decreased by increasing agarose crosslinking from 4 to 6 or 10%. The low crosslinking level in the 4 BCL agarose results in larger pores, which contain higher concentration of reactive groups for the co-immobilization of the enzymes in the MANAE support. The higher quantity of weak ionic interactions were not enough to distort enzyme structure resulting in the higher expressed activities. In opposition, the lower degree of derivatization with glutaraldehyde and the consequent fewer covalent bonds in the 10 BCL agarose resulted in less distortion to enzyme structures and consequent higher expressed activities. The Glut derivative using 10% cross-linked agarose was that which resulted in the highest expressed activity levels, corresponding to 26.1, 100 and 42.5% for the xylanase, β-xylosidase and α-1-arabinofuranosidase, respectively.

Based on previous observations, immobilization on the Glut support was followed by measuring the enzyme activities and protein in the supernatant during 16 h (Fig. 2). The enzymes were quickly immobilized on the support, i.e., after 30 min most of the enzyme
activities had disappeared from the supernatant. After this period, immobilization proceeded at lower rate, with exception of the xylanase whose activity remained in the supernatant (≈20%) for the subsequent period. Immobilization of proteins also proceeded at lower rate and after 16 h, 20% of the proteins still remained in the supernatant.

In sequence, three derivatives were prepared by carrying out immobilization for 4, 8 and 16 h in order to verify the enzyme activities in the derivatives obtained after these three immobilization periods (Table 3). As previously observed in Fig. 2, immobilization yield in relation to protein and enzymatic activities increased by increasing the immobilization period, nevertheless the best result in terms of expressed activities was obtained with the derivative prepared by proceeding immobilization during 8 h (Table 3). These results indicated that enzyme–support reaction for more prolonged periods may be causing distortion of the enzyme structures, resulting in activity loss. Thus, the selected support for the co-immobilization of the xylanase, β-xylanidase and α-L-arabinofuranosidase from P. janczewskii was agarose 10% crosslinking activated with MANAE cross-linked with 0.5% (v/v) glutaraldehyde and by carrying out immobilization at pH 7.0 for 8 h.

In order to obtain covalent bonds from Schiff base formed between protein lysines and aldehyde groups of glutaraldehyde, the support was incubated in pH 8.5 and reduced with sodium borohydride. The pH increase and reduction in itself negatively influenced the enzymatic activities, i.e., after the procedure, xylanase and β-xylanidase activities were 50 and 90% reduced, respectively, and no α-L-arabinofuranosidase activity was detected on the support. Due to these results, the non-reduced Glut derivative was submitted to post-immobilization techniques (Table 4 null, i.e., coating with PEI or dextran–aldehyde (and further reduced), and it was also submitted to a second round of cross-linking with glutaraldehyde (further reduced or not). Coating with the ionic polymer PEI had a strong negative influence and the three enzyme activities were totally depleted from the biocatalyst. Dextran-aldehyde coating and the maintenance of pH 7.0 during borohydride reduction resulted in residual activity corresponded to 66.9, 57.0 and 55.5% of those previously observed for the xylanase, β-xylanidase and α-L-arabinofuranosidase, respectively. When the derivative was submitted to a second round of glutaraldehyde crosslinking it presented residual activity of 39.6, 21.7 and 47.8% before, and 33.0, 7.7 and 25.9% after borohydride reduction, for the xylanase, β-xylanidase and α-L-arabinofuranosidase, respectively. If this latter derivative was coated with dextran–aldehyde and then reduced, the residual activity was even lower, corre-
The thermostability of xylanase, β-xylosidase and α-1-arabinofuranosidase from *P. janczewskii* co-immobilized on Glut derivative coated with dextran MW 6000 at 40% degree of oxidation. Incubation was carried out in 0.05 M glycine HCl buffer pH 3.0 at 50 °C for the β-xylosidase and α-1-arabinofuranosidase, and at 80 °C for the xylanase. Enzymatic activities were assayed in 0.05 M sodium acetate buffer pH 4.0 at 25 °C. Initial enzyme activities were regarded as 100%. Residual (%) xylanase (▲), β-xylosidase (●) and α-1-arabinofuranosidase (■) activities.

**Fig. 6.** Thermostability of xylanase, β-xylosidase and α-1-arabinofuranosidase from *P. janczewskii* co-immobilized on Glut derivative coated with dextran MW 6000 at 40% degree of oxidation. Incubation was carried out in 0.05 M glycine HCl buffer pH 3.0 at 50 °C for the β-xylosidase and α-1-arabinofuranosidase, and at 80 °C for the xylanase. Enzymatic activities were assayed in 0.05 M sodium acetate buffer pH 4.0 at 25 °C. Initial enzyme activities were regarded as 100%. Residual (%) xylanase (▲), β-xylosidase (●) and α-1-arabinofuranosidase (■) activities.

**Table 6.** Hydrolysis of wheat and oat spelt arabinoxylans by multienzymatic derivative of xylanolytic enzymes from *P. janczewskii*.

<table>
<thead>
<tr>
<th>Time (h)</th>
<th>Xylose yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>WAX</td>
</tr>
<tr>
<td>0.25</td>
<td>0.8</td>
</tr>
<tr>
<td>0.5</td>
<td>1.4</td>
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<tr>
<td>1</td>
<td>2.3</td>
</tr>
<tr>
<td>2</td>
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<tr>
<td>24</td>
<td>28.9</td>
</tr>
<tr>
<td>48</td>
<td>41.0</td>
</tr>
<tr>
<td>72</td>
<td>43.5</td>
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</tbody>
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Hydrolysis was carried out by incubating 0.05 g of Glut derivative coated with dextran MW 6000 at 40% degree of oxidation with 10 ml of 0.5% (w/v) substrate prepared in 0.05 M sodium acetate buffer pH 4.0 at 40 °C. WAX—wheat arabinoxylan; OSX—oat spelt xylan.

In the subsequent step, dextrans of different MW (6000, 40,000 and 100,000 Da; 100% degree of oxidation) were evaluated for coating the Glut derivative in order to improve enzyme activity and stabilization (Table 5). Coating the derivative with dextran MW 100,000 negatively influenced the immobilized enzymes and their activities were greatly reduced. Coating with dextran MW 6000 resulted in similar xylanase and β-xylosidase activities and half of the α-1-arabinofuranosidase activity was observed in relation to the derivative coated with dextran MW 40,000. A previous stability study, however, demonstrated that the xylanase was more thermostable in the derivative coated with dextran MW 6000 than with dextran MW 40,000 (as further presented). In this sense, derivatives coated with dextran MW 6000 at 20 and 40% degree of oxidation (DO) were prepared and the expressed activities were compared to those from the derivative coated with the completely oxidized dextran. Among them, the lowest enzyme levels were observed in the derivative coated with dextran at 20% DO. Similar levels of xylanase and β-xylosidase were observed in the derivatives coated with dextrans at 40 and 100% DO, while the α-1-arabinofuranosidase activity was substantially higher in the derivative coated with dextran at 40% DO (Table 5).

### 3.2. Derivative characterization

After selecting the most adequate support and optimizing immobilization conditions, the stability of the immobilized enzymes in the Glut derivative coated with dextrans MW 6000 and 40,000 Da (completely oxidized) was initially evaluated (Fig. 3). The immobilized xylanase was highly stabilized, retaining 80% of activity in the case of the derivative coated with dextran MW 6000, and around 60% in the derivative coated with dextran MW 40,000. Similar activity was observed even after 24 h incubation (shown up to 8 h). The β-xylosidase coated with dextran MW 6000 was only a little more stable than that coated with dextran 40,000; while for the α-1-arabinofuranosidase no differences were observed.

When the stability of the xylanase in the derivatives coated with dextran MW 6000 at different DO was evaluated at different pH and 50 °C (Fig. 4), it was observed that coating the biocatalyst surface with this polymer stabilized the enzymes in all pH. Among the
dextran oxidized at different degrees, the derivative coated with dextran at 20% DO presented the lowest stability, intermediate levels were verified with dextran at 100% DO and the highest stability was observed with the 40% DO, especially in the pH range from 3.0 to 5.0, in which more than 85% of the activity was retained.

Dextran coating as a post-immobilization technique has been widely used in enzyme technology for many different enzymes. When dextran at low degree of oxidation is used only some covalent attachments are formed between enzyme and the polymer, and the residual sugar chains remain attached rendering non-natural and large sugar moiety for the enzyme, i.e., the chemical glycosylation of a protein [36]. On the other hand, when the dextran is highly oxidized, it results in the formation of many covalent attachments between enzyme and support rigidifying enzyme structure, improving stabilization with consequent thermostabilization [25].

In this sense, some stabilization was achieved by glycosylation of the enzymes as observed for the xylanase from P. janczewskii, although it was not stabilized in a wide pH range. Additional covalent attachments rendered more stability to the enzyme at different pH, but when excessive it results in loss of activity as previously observed in Table 5. Thus, the best results were obtained with partially oxidized dextran, which corresponds to a mix of both glycosylation and an intermediate level of covalent attachments, rendering the highest stabilization to the co-immobilized enzymes.

Considering the stability of the immobilized xylanase in different pH and also that the derivative coated with dextran at 40% DO expressed good levels of the three enzyme activities, the derivative coated with dextran MW 6000 at 40% DO was the most promising, being further characterized and applied for arabinobioxylans hydrolysis.

When the enzymatic reactions were carried out at different conditions, the immobilized xylanase and α-1-α-arabinofuranosidase presented optimum activity in pH between 2.0 and 4.0, and the β-xylanosidase in pH 3.0 (Fig. 5a). The immobilized xylanase, α-1-α-arabinofuranosidase and β-xylanosidase presented optimum activity at 50, 65 and 80 °C, respectively (Fig. 5b). A shift in the pH for optimum activity is observed since the free xylanase presented optimum activity at pH 5.0 and both free β-xylanosidase and α-1-α-arabinofuranosidase were optimally active at pH 4.0. The optimum temperature of the xylanase was similar to that observed for the free-enzyme, while for the α-1-α-arabinofuranosidase and β-xylanosidase the optimum temperature was increased by 5 °C in relation to those previously observed for the free enzymes [28,29].

The xylanase was highly thermostable retaining 83.1 and 70% of activity even after 24 h incubation at 60 and 70 °C, respectively [not shown]. At 80 °C, its half-life was 1.7 h (Fig. 6). The β-xylanosidase and the α-1-α-arabinofuranosidase were very stable up to 40 °C, retaining 53 and 100% of the activity after 24 h incubation (not shown). At 50 °C, the half-lives of the β-xylanosidase and α-1-α-arabinofuranosidase were 2.3 and 3.8 h (Fig. 6). The immobilized α-1-α-arabinofuranosidase is therefore 25-fold stabilized in relation to its free counterpart [29]. The higher thermostability of the xylanase may be associated to its possible monomeric structure and lower MW that resulted in more attachments to the support and a more adequate coating by the dextran.

3.3. Hydrolysis of arabinobioxylans

The synergistic action of xylanolytic enzymes, divided into homoeosnergy, occurring between main-chain cleaving enzymes, and heterosnergy, occurring between main-chain cleaving and debranching enzymes, has been reported by using free- [37] and at a lesser extent by using immobilized enzymes [8]. In this hydrolysis study a low viscosity wheat arabinoxylan (WAX) and oat spelt xylan (OSX) were used. WAX is a highly arabinosylated polymer containing 41% arabinose and 59% xylose residues, according to the supplier. OSX holds a complex structure containing 81.4% xylose, 9.7% arabinose, 3.4% glucose, 1.2% galactose and 4.3% uronic acid residues [38]. The presence of substituents intensely limits the action of endo-xylanase, hampering the complete degradation of the polymers into their monomers. The functionality of the multienzymatic derivative was evaluated by hydrolyzing these substrates in pH 4.0 at 40 °C during 72 h (Fig. 7). As previously observed, at these conditions the xylanase and the α-1-α-arabinofuranosidase were very active, and the β-xylanosidase operate with 65% of its maximum activity: at this temperature the enzymes do not operate with maximum activity, nevertheless, they present high stability to operate for prolonged cycles. The prepared biocatalyst was active and the similar hydrolysis profiles revealed that they equally degraded both substrates, independent of structure and composition. After 8 h, the reaction velocity is reduced probably due to the inhibitory effect of xylose or arabinose on enzyme activities since the enzymes were stable in the reaction conditions. The maximum xylose yield verified after 72 h corresponded to 43.5 and 42.8% for WAX and OSX, respectively (Table 6).

The application of a biocatalyst in industry requires the stability of immobilized enzymes through many operational cycles. After consecutive cycles of wheat arabinoxylan hydrolysis, the
xylanase and the α-1,2-arabinofuranosidase were very stable retaining 86.6 and 88.0% of the activity even after 10 reuse cycles. The β-xylanase activity decreased to 78.7 and 47% after five and ten cycles, respectively (Fig. 8a). The product release decreased to 80 and 60% after five and ten cycles, respectively, that may be mainly related to the reduction in the β-xylanase activity during the cycles (Fig. 8b).

4. Conclusions

The simultaneous co-immobilization of the crude xylanase, β-xylanase and α-1,2-arabinofuranosidase from P. janczewskii was not such a simple procedure, but by adequately selecting the support and by optimizing parameters such as agarose cross-linking and immobilization period, the amount of each enzyme in the derivative can be considerably increased, allowing the preparation of a multienzymatic biocatalyst acting cooperatively in the complete degradation of complex substrates. By optimizing conditions, the expressed activity of the xylanase, the most difficult enzyme to be immobilized, in the glutaraldehyde derivative can be two-fold higher than initially verified. By optimizing the xylan coating, the activity and the stability of the enzymes could also be substantially increased. The mildly activated glutaraldehyde derivative coated with low molecular weight partially oxidized xylan is the most promising, presenting interesting properties, and offering advantages over the use of free enzymes. It is possible to obtain a very stable derivative, with improved temperature, pH and operational stability. Furthermore, the use of this multi-enzymatic derivative leads to the direct formation of xyllose monosaccharide from different arabinoxylans, therefore representing an encouraging alternative for degradation of hemicelluloses since it can be successively reused thereby reducing the need for new enzyme loads and consequently reducing the process cost.

References

