Increasing the Enzyme Loading Capacity of Porous Supports by a Layer-by-Layer Immobilization Strategy Using PEI as Glue

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Received: 12 June 2019; Accepted: 29 June 2019; Published: 29 June 2019

Abstract: A new strategy to increase the enzyme-loading capacity of porous supports was investigated. Lipase from Pseudomonas fluorescens (PFL) was immobilized on octyl-agarose (OA) beads and treated with polyethyleneimine (PEI). Then, PFL was immobilized on the previous PFL layer. Next, the biocatalyst was coated with PEI and a third layer of PFL was added. Sodium dodecyl sulfate polyacrylamide electrophoresis showed that the amount of PFL proportionally increased with each enzyme layer; however, the effects on biocatalyst activity were not as clear. Hydrolyzing 50 mM of triacetin at 25 °C, the activity of the three-layer biocatalyst was even lower than that of the bi-layer one; on the contrary its activity was higher when the activity was measured at 4 °C in the presence of 30% acetonitrile (that reduced the activity and thus the relevance of the substrate diffusion limitations). That is, the advantage of the multilayer formation depends on the specific activity of the enzyme and on the diffusion limitations of the substrate. When octyl agarose (OA)-PFL-PEI-PFL preparation was treated with glutaraldehyde, the activity was reduced, although the enzyme stability increased and the immobilization of the last PFL layer offered results similar to the one obtained using the three-layer preparation without glutaraldehyde modification (90%).

Keywords: biocatalyst volumetric activity; enzyme immobilization multilayers of enzyme; lipase from Pseudomonas fluorescens; polyethyleneimine

1. Introduction

Enzyme immobilization is a key step in the enzyme biocatalyst design [1]. Initially, its purpose was just to facilitate the reuse of these expensive biocatalysts. However, soon, the technique showed a great potential to improve many enzyme properties, such as stability, selectivity, specificity, purity, etc. [2–11]. The selection of the support is in many instances decisive for the final implementation of the biocatalysts [12]. Price is considered relevant in some instances, but that depends on the degree of enzyme improvement that can be achieved after immobilization and on other features of the support [11]. Their mechanical properties should be compatible with the reactor system. For instance, porous rigid supports like silicates are useful for fixed bed reactors, but they can be broken in stirred systems, while flexible supports are adequate for stirred systems but produce problems in fixed bed reactors [1]. The particle size should be adequate for the filters and catalysts retention equipment. Moreover, one very important feature is the enzyme loading capacity: if one can immobilize twice the amount of enzyme on one support than on another, the support price of the final catalyst is reduced by half [12].
In a similar way, large-size non-porous supports offer a very low specific area and low loading, unless going down to nanoscale, and in this case handling and price may become problematic [1,13–16], as standard filters will be inefficient to retain the biocatalyst. The impact of enzyme loading capacity of the support on the support selection could be reduced if somehow one could increase the loading capacity of the support. This could be achieved if one enzyme layer is immobilized over a previously immobilized enzyme layer, making it possible to load enzyme until the entry of new enzyme molecules is hindered by the reduction of the support pore size, or substrate diffusional limitations make inefficient the firstly immobilized enzyme molecules layer near to the support surface. Using a non-porous support, the only relevant substrate diffusion problem will be the diffusion from the upper enzyme layer to the lower enzyme layer in contact with the support, while using porous supports, the entry of the substrate in the particle can increase the diffusion problems. The enzyme activity may play an important role on the importance of these substrate diffusion problems, a poorly effective enzyme may be a good model [17], but we can mimic this by using experimental conditions where the enzyme activity decrease.

To reach this goal, it is necessary to “activate” the immobilized enzyme molecules surface in a way that allows the immobilization of a new layer of enzymes. This may be achieved using polyethylenimine (PEI) [18–21] (Figure 1). This polymer is multicationic, has primary, secondary and tertiary amino groups and has been broadly used in biocatalysis design, in some instances to coimmobilize enzymes [22,23] or enzymes and cofactors [24–26]. The only problem (with the second and further enzyme layers) of this new strategy is that, after immobilizing the enzyme via ion exchange on PEI, the new coating with PEI of the previously ionically exchanged enzyme could produce some enzyme leaching from the support by competition among the PEI coating the previously immobilized enzyme layer, the immobilized enzyme molecules layer and the new doses of free PEI.

Figure 1. Scheme of the multilayer immobilization strategy proposed in this paper. PEI: polyethylenimine; PFL: lipase from Pseudomonas fluorescens.

In this first approach, we have tried to build a three-layer biocatalyst using the lipase from Pseudomonas fluorescens (PFL). This enzyme is very popular in literature [27–29]. To simplify the study, we have immobilized a first layer of PFL on octyl agarose (OA) beads via interfacial activation, a kind
of lipase biocatalysts often submitted to the coating with PEI to prevent enzyme release under drastic conditions, thus improving the immobilized enzyme stability [30–32]. The lipase immobilization via interfacial activation on hydrophobic supports mimics the adsorption of lipases on substrate drops and involves the hydrophobic areas of the enzyme near to the active center. The adsorption is strong but the large size of the enzyme pocket and the layer of octyl groups leaves a space large enough to the substrates to reach the active center of the enzyme, even giving usually a lipase hyperactivation due to the involvement of the open form of the lipase in the immobilization [33,34].

Next, the immobilized enzyme was coated with PEI. Then, we immobilized a second layer of PFL on the PEI layer and studied if a new coating with PEI could produce the release of the ionically exchanged enzymes. If this leaching did not occur, a third layer of enzyme was immobilized. The effect of this layer-by-layer biocatalyst building on the biocatalysts volumetric activity was evaluated using different substrates and conditions. It may be expected that when specific activity of the enzyme is very high, the volumetric activity of the biocatalysts may not be increased by using this multilayer strategy due to the increase of substrate diffusional limitations (diffusion of substrate rate may be lower than enzyme substrate consumption rate) [35–38]. However, if the specific enzyme activity is not very high and substrate diffusion limitations are scarce, a continuous increase in enzyme activity may be expected when the number of enzyme layers increases.

2. Results and Discussion

2.1. Immobilization of PFL on octyl agarose

To take full advantage of this strategy, maximum PFL loading of octyl agarose must be utilized. Thus, Figure 2 shows the immobilization yields at different PFL loadings: 1 mg/g and 80 mg/g. While using 1 mg/g the immobilization is very rapid and almost total in a very short time, using 80 mg of PFL/g of support, the immobilization yield is only 73.5% after 2 h. This overloading was used in all further experiments to ensure full coating of the support with the enzyme and the successive layers.

This high enzyme loading increased the substrate diffusion limitations of p-nitrophenyl butyrate (pNPB); thus, while using low enzyme loadings an increment of the enzyme activity is observed upon immobilization (around 10%), at maximum loading, the enzyme activity even decreased, the immobilized enzyme has below 75% of the activity of the free enzyme. That may be visualized in Figure 2.

![Figure 2](image_url)

**Figure 2.** Immobilization courses of PFL on octyl agarose at different enzyme loadings ((A): 1 mg/g, (B): 80 mg/g). Experiments details as described in Methods. Circles: supernatant; triangles: suspension; squares: reference.

There are reports on the effect of enzyme loading and immobilization rate on the final immobilized enzyme stability [39,40]. To study if the use of overloaded PFL biocatalysts could have some effects on...
enzyme stability, both preparations were inactivated (Figure 3). The overloaded biocatalysts presented higher stability than the low loaded preparation, although part of this effect may be explained by an artifact caused by the diffusional limitations (the specific activity of the overloaded enzyme was 68% of that of the lowly loaded preparation), it looks that the stability of the overcrowded preparation was slightly higher under the inactivation conditions. That is, the use of overcrowded PFL preparations had even advantages in terms of enzyme stability.

Figure 3. Effect of enzyme loading in the inactivation courses of octyl PFL biocatalysts. Inactivation carried out in 50 mM Tris at pH 7 and 75 °C. Experiments details as described in Methods. Squares: highly loaded biocatalyst; circles: lowly loaded biocatalyst.

2.2. Coating of OA-PFL Biocatalysts with PEI

The PEI coating of the immobilized PFL produced a clear increment in the enzyme activity versus \( p\text{-NPB} \) (around 10%) (Table 1). This strategy has been used to stabilize other lipases immobilized on this support, and the positive effect on enzyme activity found here is an additional interesting feature [30–32]. This increase in activity is quite relevant considering that the overloaded biocatalyst already has diffusion limitations. A similar experiment using preparations with only 1 mg/g afforded an increment of enzyme activity of 30% (not shown).

Table 1. Mass activity of different PFL biocatalysts in hydrolysis of 1 mM \( p\text{-nitrophenyl butyrate (pNPB)} \) at 25 °C at pH 7. Experiments were performed as described in Methods. The loading of each layer as fairly similar, around 60–65 mg of protein/g of octyl agarose.

| PFL Preparations     | Activity versus \( p\text{-NPB} \) (U/g) |
|----------------------|----------------------------------------|
| OA-PFL               | 96.1 ± 1.0                             |
| OA-PFL-PEI           | 105.8 ± 1.2                            |
| OA-PFL-PEI-PFL       | 132.2 ± 1.1                            |
| OA-PFL-PEI-PFL-PEI   | 123.4 ± 1.0                            |
| OA-PFL-PEI-PFL-PEI-PFL | 107.7 ± 0.9                           |

Next, we analyzed the effect of the PEI coating in the stability of both, highly and lowly loaded biocatalysts. Figure 4 shows that the effect is not very relevant in contraposition with the results using other enzymes, where at least the overloaded enzyme biocatalysts increased its stability upon PEI coating [30,32]. This may be related to the very strong immobilization of the enzyme on octyl agarose [41], as the main reason for the PEI coating stabilization was associated to the prevention of the enzyme release [30,32].
with the components of the commercial powder, as it is able to become immobilized on octyl agarose way, the substrate can only circulate between the holes of the enzyme molecules in the upper layer. This scarce effect of the immobilization of the second layer of enzyme on the biocatalyst activity may be caused by an increase of the diffusion limitations, mainly affecting the enzyme layer immobilized via interfacial activation on the octyl layer [38]. The coating of this second PFL layer with PEI decreased the pNPB activity, which was only slightly higher than that of the first PFL-octyl layer. A further addition of PFL (again with around 80% immobilization yield) to get a three PFL layers biocatalyst decreased the pNPB final activity of the biocatalyst (Table 1).

The explanation for this effect on pNPB activity of the enzyme loading should be based on an increase of the substrate diffusion issue. PEI is a random coil polymer that permits the entry of the substrate without serious additional limitations. However, after the enzyme immobilization on the PEI layer, the substrate diffusion problems increased, as the protein layer really makes the entry of the substrate difficult. Moreover, the interactions of the rigid protein molecules with the PEI bed made that it lost mobility, promoting that the substrate diffusion become even more difficult. That way, the substrate can only circulate between the holes of the enzyme molecules in the upper layer increasing the tortuosity of the way the substrate must follow to reach the enzyme molecules located below [38]. This effect became very negative for the third layer of enzyme, when even the pore diameter of the support will be also reduced and the substrate diffusion limitations will be strongly increased. Now, the substrate will could not follow a straightforward way to reach the enzyme molecules in contact with the support, but should follow the tortuous route existing between enzyme molecules [38]. Figure 5 shows the sodium dodecyl sulfate –polyacrylamide gel electrophoresis study of the different biocatalysts, where the amount of protein proportionally increased with the number of enzyme layers. The band of more than 66 KD could be the result of covalent dimers of PFL formed by Maillard reaction with the components of the commercial powder, as it is able to become immobilized on octyl agarose at low ionic strength.

From these results, it can be seen that just two enzyme layers produced the most positive effects on the biocatalyst mass activity, as the third layer produce even a decrease in the activity. Thus, we did not try to immobilize new layers of enzyme and focused to look for reactions where the three-layer biocatalyst offers a real advantage. In the next section, the activity of the different biocatalysts was analyzed using triacetin, a substrate that allowed higher concentrations than pNPB.

2.3. Immobilization of Further PFL Layers on OA-PFL-PEI Biocatalyst

Immobilization of PFL on the OA-PFL-PEI biocatalyst is quite rapid (not shown results) and, in this instance, the final pNPB activity of the biocatalyst increased less than 40% compared to the activity of the free and previously immobilized enzyme; this occurred even though the amount of immobilized enzyme was similar to that of the first PFL layer (around 80%). This scarce effect of the immobilization of the second layer of enzyme on the biocatalyst activity may be caused by an increase of the diffusion limitations, mainly affecting the enzyme layer immobilized via interfacial activation on the octyl layer [38]. The coating of this second PFL layer with PEI decreased the pNPB activity, which was only slightly higher than that of the first PFL-octyl layer. A further addition of PFL (again with around 80% immobilization yield) to get a three PFL layers biocatalyst decreased the pNPB final activity of the biocatalyst (Table 1).

Figure 4. Effect of PEI coating of lowly (A) and highly (B) loaded PFL octyl biocatalysts. Inactivation carried out in 50 mM Tris at pH 7 and 75 °C. Other details are described in Methods section. Circles: unmodified preparations. Squares: PEI coated preparations.

Figure 5 shows the sodium dodecyl sulfate –polyacrylamide gel electrophoresis study of the di...
We checked the adsorption strength of PFL on the PEI bed, by incubating the biocatalysts in 1 M ammonium sulfate at pH 7, and we found that less than 1% of the enzyme molecules were released to the medium even at 37 °C. This showed the very strong immobilization of PFL on the PEI bed.

2.4. Activity of the Different PFL Biocatalysts versus Triacetin

Table 2 shows the activity of the different PFL preparations in triacetin hydrolysis at 25 °C. The activity of OA-PFL was decreased just by coating the enzyme with PEI. This decrease is very likely due to the high volumetric activity of the biocatalysts: there are substrate diffusion limitations even using 50 mM of substrate. In fact, the activity considering the enzyme load and comparing the preparation of 1 mg PFL/g of support, the activity of the overloaded preparation is around 70% of the expected one. As the PEI coating may reinforce these problems [38], an increase in the diffusion problems may be the responsible of this activity decrease instead of a direct negative effect of PEI on the enzyme properties versus this substrate. To investigate this, an immobilized OA-PFL preparation loaded with 1 mg enzyme/g of support was modified with PEI: the activity increased by a 10% versus triacetin (not shown results), confirming that in the case of the overloaded preparation the decrease in activity is mainly caused by diffusion limitations and not by enzyme inactivation.

Table 2. Mass activity of different PFL biocatalysts in hydrolysis of 50 mM triacetin at pH 5. Activity is given in micromoles of acetic acid released per minute and gram of biocatalysts. T1: 25 °C. T2: 4 °C in the presence of 30% acetonitrile (v/v). Experiments were carried out as described in Methods.

| PFL Preparations | Activity versus Triacetin (T1) (U/g) | Activity versus Triacetin (T2) (U/g) |
|------------------|--------------------------------------|--------------------------------------|
| OA-PFL           | 384.4 ± 2.3                          | 17.6 ± 0.3                           |
| OA-PFL-PEI       | 277.0 ± 1.8                          | 17.9 ± 0.4                           |
| OA-PFL-PEI-PFL   | 441.4 ± 3.2                          | 33.6 ± 1.1                           |
| OA-PFL-PEI-PFL-PEI | 793.6 ± 4.4                       | 76.7 ± 1.5                           |
| OA-PFL-PEI-PFL-PEI-PFL | 575.8 ± 3.3                      | 82.0 ± 1.4                           |

Figure 5. SDS-PAGE of the different immobilized PFL preparations. The experiments were performed as described in Methods (load was 80 mg/g per layer). Lane 1: Low molecular marker; Lane 2: Octyl Agarose-PFL, Lane 3, OA-PFL-PEI, Lane 4: OA-PFL-PEI-PFL, Lane 5: OA-PFL-PEI-PFL-PEI, Lane 6: OA-PFL-PEI-PFL-PEI-PFL.
Immobilization of a second layer of PFL increased the biocatalyst activity versus triacetin, and in this case, the coating of this second layer of PFL with PEI greatly increased the biocatalyst activity (almost doubling the total enzymatic activity). This difference with the effect of the first PEI coating may be based on the different immobilization strategy in the two enzyme layers: the first one was immobilized via interfacial activation, with the active center oriented towards the rigid support surface, while the second one was immobilized via ion exchange on a polymeric mobile and flexible bed. To check this, PFL was immobilized on a PEI coated support, and the treatment with PEI doubled the enzyme activity (not shown). A further last layer of PFL was again negative for the final volumetric activity of the biocatalysts, very likely because of a reinforcement of the diffusional limitations to the substrate [38], both from outside to inside in the pore of the support and from above to below through the three enzyme layers.

Trying to find conditions where the three layers of enzyme may have advantages, the temperature was decreased to 4 °C and 30% acetonitrile was added (this improved the selectivity of the reaction towards 1,2 diacetin [42]). These conditions decreased the biocatalyst activity from 384.4 to 17.6 U/g (Table 2), conditions where the substrate diffusion limitation may be greatly decreased (the relevant point is the substrate diffusion versus substrate conversion rates). The effect of the PEI modification of OA-PFL on the enzyme activity was even slightly positive (though at 25 °C the biocatalyst activity decreased). The second layer of PFL increased the activity by almost a twofold factor, and the same may be said of the modification with PEI. Under these reaction conditions, addition of the third layer of PFL produced a small increase of biocatalyst activity (from 76.7 to 82 U/g), lower that the values expected from the second PFL layer (which, considering the amount of immobilized protein, should be triple of the activity of the one PFL layer), but enough to increase the total volumetric activity of the biocatalyst. That is, when the enzyme specific activity was decreased and that way the substrate diffusional limitations were diminished, the positive effect of the multilayers on the biocatalyst activity increased. The limit of the interest of increasing the number of layers will depend on many factors: substrate, substrate concentration and experimental conditions.

2.5. Treatment of the Biocatalysts with Glutaraldehyde

Some authors propose the treatment with glutaraldehyde of enzymes immobilized via ion exchange to prevent the enzyme release if high ionic strength or drastic pH values are used (e.g., hydrolysis of concentrated oils solutions, where the pH may decrease below 4 in the presence of high concentrations of free fatty acids) [23,43–45].

Thus, in a last trial, the biocatalysts were treated with glutaraldehyde (Table 3). When the second layer was treated with glutaraldehyde, only 14 and 8% of the activity was lost using triacetin at 25 °C and at 4 °C in 30% acetonitrile, respectively. This shows that the treatment with glutaraldehyde of the second PFL layer do not harm significantly the PFL structure. However, the effect of PEI coating of the biocatalyst treated with glutaraldehyde was more negative: the biocatalyst lost more than 25 and almost 50% of its activity in triacetin hydrolysis at 25 °C or at 4 °C/30% acetonitrile, respectively. In the third PFL layer, the effect of glutaraldehyde treatment of the previous layers was also negative and the biggest drop in activity was observed in triacetin hydrolysis at 25 °C, where the biocatalyst lost about 28% of its activity. These negative effects of glutaraldehyde treatment in enzyme activity may be based on the PEI crosslinking that may increase the diffusion problems as the polymer will not be longer random coil [38].

An SDS-PAGE study showed that only the last PFL layer, not treated with glutaraldehyde, can be released from the biocatalyst, confirming the successful massive intermolecular crosslinking of the other two underlying enzyme layers (Figure 6). This treatment clearly prevents enzyme release from the support even under very harsh conditions (boiling in the presence of SDS).
Table 3. Effect of the glutaraldehyde treatment of the PFL biocatalysts on enzyme activity in the hydrolysis of 50 mM triacetin at pH 5. T1: 25 °C. T2: 4 °C in the presence of 30% acetonitrile (v/v). The data are given as relative activity, considering the activity of the respective no treated preparation as 100%. Experiments were performed as described in Methods.

| PFL Preparations          | Relative Activity versus Triacetin, (%) (T1) | Relative Activity versus Triacetin, (%) (T2) |
|---------------------------|----------------------------------------------|----------------------------------------------|
| OA-PFL-PEI-PFL-GA 1%      | 86.4 ± 1.9                                   | 91.9 ± 3.2                                   |
| OA-PFL-PEI-PFL-GA 1%-PEI  | 73.6 ± 3.1                                   | 51.2 ± 2.1                                   |
| OA-PFL-PEI-PFL-GA 1%-PEI-PFL | 72.5 ± 1.1                             | 90.4 ± 3.1                                   |

Figure 6. SDS-PAGE of the glutaraldehyde modified PFL preparations. The experiments were performed as described in Methods (load was 80 mg/g per layer). Lane 1: Low molecular marker; Lane 2: OA-PFL-PEI-PFL-Glutaraldehyde, Lane 3: OA-PFL-PEI-PFL-Glutarialdehyde-PEI, Lane 4: OA-PFL-PEI-PFL-Glutarialdehyde-PEI-PFL.

Figure 7 shows how the treatment of the OA-PFL-PEI with glutaraldehyde has a relevant stabilizing effect. Although the first inactivation steps are fairly similar, there was a fraction (accounting for 75% of the enzyme activity) that is much more stable than the unmodified biocatalysts. The stabilization can be produced by either intermolecular or intramolecular crosslinking. To check the importance of the intramolecular crosslinking, the lowly loaded preparation coated with PEI was modified with glutaraldehyde, and the resulting stabilization effect was negligible. That suggests that the main effect of the glutaraldehyde treatment in the overloaded preparations was caused by intermolecular crosslinking. In any case, the positive effect of the glutaraldehyde treatment was very clear in terms of stability, although it was not as positive in terms of activity. The final decision will depend on the necessity or not to improve enzyme stability and prevent enzyme release.
3. Materials and Methods

3.1. Materials

Lipase from *Pseudomonas fluorescens* (PFL, 0.132 mg protein/mg powder, determined by Bradford method [46]), polyethylenimine (PEI) (MW 25,000), triacetin, p-nitrophenyl butyrate (p-NPB), glutaraldehyde solution Grade II (25% in H₂O) and acetonitrile for HPLC (gradient grade, ≥ 99.9%) were purchased from Sigma Aldrich. Cl-4B octyl-Sepharose beads and low molecular weight (LMW) marker for SDS electrophoresis (14.4–97 kDa) were purchased from GE Healthcare. All other reagents were of analytical grade.

3.2. Methods

All experiments were performed by triplicate and the values are given as mean values and experimental errors.

3.2.1. Immobilization of PFL

Immobilization of PFL on Octyl Agarose (OA)

PFL was immobilized on octyl-agarose using 1mg/g and 80 mg/g (overload of the support capacity that is around 60mg/g). A proper mass of lipase was diluted in 5 mM sodium phosphate buffer (pH 7) and then added to the support in a ratio 1/10 or 1/15, for 1 mg/g and 80 mg/g, respectively. The immobilization was performed under mechanical stirring at room temperature during 2 hours. A reference suspension was utilized to check the effects of the immobilization conditions on the PFL activity; this was the enzyme under identical conditions and mixed with inert agarose beads. Samples of supernatant and suspension were periodically withdrawn and their activities were measured using the pNPB assay described below.

Coating of the Immobilized Enzyme with PEI

The immobilized PFL preparations were treated with 10% w/v PEI solution at pH 7 [47]. This step was conducted under gentle stirring at 4 °C during 18 hours. Subsequently, the coated biocatalyst was filtered and washed many times with distilled water.

Immobilization of PFL on PEI-coated Biocatalysts

PFL was immobilized on the overloaded biocatalyst and PEI coated offering 80 mg/g in 5 mM sodium phosphate at pH 7 during 2 hours at 25 °C. Then, the biocatalyst was filtered and washed with
an excess of distilled water. In some cases, the biocatalyst was treated with glutaraldehyde solution 1% (v/v) in 50 mM sodium phosphate at pH 7, for 1 hour [48–50].

3.2.2. Determination of Enzymatic Activities

Hydrolysis of p-nitrophenyl Butyrate (p-NPB)

In the hydrolysis of p-NPB, 50 µL of substrate (50 mM) was added to 2.5 mL of 25 mM sodium phosphate buffer at pH 7 and 25 °C. The reaction was started by adding 50–100 µL of the enzyme sample (solution or suspension) and the hydrolysis continued during 90 seconds, with the continuous liberation of the product, the p-nitrophenol, which was detected at 348 nm using a spectrophotometer with magnetic stirring and thermostatization system.

Hydrolysis of Triacetin

Hydrolysis of triacetin was performed using 50 mM of substrate at pH 5 to prevent acyl migration [42]. 0.1 g of wet biocatalyst was added to the substrate solution at 25 °C or, in some instances, at 4 °C in presence of 30% acetonitrile to reduce the enzyme activity [42]. The catalytic activity of the biocatalysts was determined by the production of 1,2 diacetin, at conversions degrees between 15–22%. This conversion was measured using a HPLC (Kromasil C18 column of 15 cm × 0.46 cm) with a flow rate of 1mL/min of 15% acetonitrile-85% water as mobile phase. The compounds were detected with a UV detector (Waters 486) at 230 nm and the retention times were about 4 min for 1,2 diacetin and 18 min for triacetin [47].

3.2.3. Electrophoresis SDS-PAGE

The SDS-PAGE analyses were carried out using 12% and 5% of polyacrylamide as resolution gel and concentration gel, respectively [51]. The immobilized PFL samples were diluted in a solution 10% (v/v) SDS, calculating a final protein concentration of 15 mg of protein per ml in the solution. The samples were boiled for 5 minutes, centrifuged, boiled again (5 minutes) and, finally, 15 µL of the samples were loaded in electrophoresis gel. 5 µL of Low Molecular Weight marker proteins (LMW-SDS Marker—14.4—97 kDa) were used as standard. The gels were stained with Coomassie brilliant blue.

3.2.4. Thermal Inactivation

Enzyme biocatalysts (solution for free enzyme or suspension for immobilized enzyme) were incubated in 50 mM Tris at pH 7 and incubated at the indicated temperature (immobilized enzyme was diluted 1:9, free enzyme was at 1mg/ml). A temperature that gave a reasonable inactivation rate was selected. Periodically, samples were taken and their activity versus pNPB was quantified considering 100% the activity of the initial preparation.

4. Conclusions

A biocatalyst with three layers of PFL has been built. This was based on the use of PEI as glue. Modification of the enzyme with PEI using lowly loaded enzyme preparations is positive for pNPB and triacetin hydrolysis, but using the highly loaded biocatalyst it is only clear when triacetin is hydrolyzed at 4 °C and in the presence of solvents but not at 25 °C, very likely due to the increase in the substrate diffusion limitations. The third layer of enzyme only increases the biocatalyst activity under conditions where the specific activity of the enzyme is not very high. That is, the number of multilayers that can be built with increase in the final biocatalyst activity depends on the activity that the biocatalyst expresses versus the specific substrate and under the specific conditions. This strategy could be extrapolated to other lipases, and also to other industrially important enzymes. For other enzymes, the initial immobilization strategy should be changed by other immobilization method (e.g., covalent), useful for most enzymes. These studies are in course in our laboratory.
Author Contributions: N.S.R., S.A.-P., and C.M.-S.: investigation, formal analysis, writing—original draft preparation, Y.L.: investigation, V.C.-C.: methodology, formal analysis, writing—review and editing, L.R.B.G.: methodology, funding acquisition, writing—review and editing, R.F.-L.: funding acquisition, supervision, formal analysis, methodology, conceptualization, writing—original draft preparation, review and editing.

Funding: This research was funded by MICIU grant number CTQ2017-86170-R (Spain). N. S. R. thanks to CNPq for a predoctoral fellowship (CNPq scholarship–Brazil).

Acknowledgments: Martinez (Novozymes Spain) is gratefully recognized by the donation of the enzymes.

Conflicts of Interest: The authors declare no conflict of interest.

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