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How Molecular Evolution Technologies can Provide Bespoke Industrial Enzymes: Application to Biofuels

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Abstract — How Molecular Evolution Technologies can Provide Bespoke Industrial Enzymes: Application to Biofuels — Enzymatic hydrolysis of lignocellulose is one of the major bottlenecks in the development of biological conversion of lignocellulosic biomass to biofuels. One of the most efficient organisms for the production of cellulolytic enzymes is the fungus Trichoderma reesei,
mainly thanks to its high secretion capacity. The conversion of cellulose to glucose involves three types of cellulases working in synergy: endoglucanases (EC 3.2.1.4) randomly cleave β-1,4 glycosidic linkages of cellulose, celllobiohydrolases (EC 3.2.1.91) attack cellulose chain ends to produce celllobiose dimers which are converted into glucose by the β-glucosidases (EC 3.2.1.21). Unexpectedly, the amount of β-glucosidase (BGL1) from T. reesei hyperproducing strains represents a very low percentage of the total secreted proteins. A suboptimal content of this enzyme limits the performance of commercial cellulase preparations as celllobiose represents the main inhibitor of the cellulolysis reaction by celllobiohydrolases. This bottleneck can be alleviated either by overexpressing the β-glucosidase in T. reesei or optimized its specific activity. After giving a brief overview of the main available technologies, this example will be used to illustrate the potential of directed evolution technologies to develop enzymes tailored to fit industrial needs. We describe the L-Shuffling™ strategy implemented with three parental genes originating from microbial biodiversity leading to identification of an efficient β-glucosidase showing a 242-fold increase in specific activity for the pNPGlc substrate compared to WT (Wild Type) Cel3a beta-glucosidase of T. reesei. After expression of the best improved β-glucosidase in T. reesei and secretion of a new enzymatic cocktail, improvement of the glucosidase activity allows a 4-fold decrease of cellulase loading for the saccharification of an industrial pretreated biomass compared to the parental cocktail.

INTRODUCTION

In the emerging low carbon economy, enzymes have become key tools to develop sustainable and clean industrial processes in major sectors such as bioenergy and chemicals manufacturing. This article opens a room for exploiting molecular evolution approaches in order to design ideal cocktails of enzymes for the development of biological conversion of lignocellulosic biomass to biofuels. Indeed, lignocellulosic biomass is an abundant and cost-efficient substrate but its breakdown into fermentable sugars remains one of the major challenges for the development of a sustainable biofuel production process.

The article will start with an overview of the context, the social and the economical issues of 2G ethanol production. Then, we will describe the directed evolution technologies that could be applied to overcome these issues and illustrate such approach by describing a L-Shuffling™ strategy implemented with three parental genes originating from microbial biodiversity leading to identification of an efficient β-glucosidase showing a 242-fold increase in specific activity for pNPGlc substrate compared to WT Cel3a beta-glucosidase of T. reesei. We will then describe the potential of the new generated secretome after expression of the best improved β-glucosidase in T. reesei.

1 CONTEXT, SOCIAL AND ECONOMIC ISSUES

Bioethanol represents a major part of biofuels and is to date nearly exclusively produced from sugar and starch containing crops. However, this first generation bioethanol cannot meet all needs and increasing production may lead to competition for arable land and increase pressure on world market prices for cereals and basic crops for food production (Fairley, 2011).

Therefore, growing efforts are undertaken to develop second generation biofuels which use the lignocellulosic, non-edible parts of the plant as substrate for biofuel production. Using this biomass source is also leading to substantially higher greenhouse gas emission savings than first generation bioethanol (60-80% savings when compared to CO₂ emissions from gasoline) (Edwards et al., 2007). The recent new EU directive on renewable energy includes a mandatory market share of 10% for renewable energy in the transport sector in 2020. The directive includes a minimum saving of 35% GHG (greenhouse gas) emissions for current biofuels and of 60% for production plants to be installed after 2017. These commitments pave the way for second generation biofuels (Directive 2009/28/EC on the promotion of the use of energy from renewable sources).

The classical second generation bioethanol production process comprises four major steps. First, a physico-chemical pretreatment, under acidic or alkaline conditions, destroys the rigid structure of the plant cell wall (for review, see Chundawat et al., 2011). This material can then be submitted to enzymatic hydrolysis in order to generate fermentable sugars, mainly glucose. These sugars can be converted into ethanol by yeast fermentation. Finally, the ethanol thus produced is recovered by distillation. The main issue for an industrial application of such process is its high cost compared to first generation bioethanol. It is anticipated that for a competitive production, the 2G ethanol production cost must be divided at least by two.
One of the most important cost-contributors in bioethanol production is enzymatic hydrolysis, and many efforts are aimed at decreasing the cost of this step (Margeot et al., 2009). Biotechnology has therefore become one of the key growth drivers for so-called “cellulosic bioethanol”, also referred to as “2G bioethanol”. Due to its capacity to produce large amounts of cellulose degrading enzymes, T. reesei has been extensively studied in various fields of white biotechnology, especially in biofuel production from lignocellulosic biomass (Kubicek et al., 2009). In parallel of studies dedicated to improvement of cellulolytic cocktails, other process schemes allowing for production of 2G ethanol have been proposed in order to lower its cost. In one of these suggested processes, the hydrolysis and fermentation steps are carried out in a single step (“Simultaneous Saccharification and Fermentation”, SSF, opposite to “Separate Hydrolysis and Fermentation”, SHF) (Olofsson et al., 2008). The advantages of such processes include the consumption by yeasts of sugars that would inhibit cellulases and the reduction of the risks of bacterial contamination during the fermentation step. However, inhibition of cellulases by ethanol and different temperature optima for yeast and cellulases remain problems to be solved. A promising alternative is the consolidated bioprocessing (CBP) which uses only a single organism expressing cellulolytic enzymes and fermenting the released sugars (Olson et al., 2012). The latter concept has yet to reach industrial maturity.

Whatever the process, conversion of lignocellulosic renewable resources into biofuels creates a crucial need for second generation enzymes with optimized characteristics, fine-tuned properties and enhanced performance.

2 MOLECULAR EVOLUTION

Directed evolution of proteins offers different alternatives for optimizing enzyme characteristics and performance. Protein engineering has first been performed by using techniques based on rational design and structural approaches. However, randomized methods rapidly proved to have a much broader scope of application than rational design for rapidly optimizing enzymes and other proteins (Illanes et al., 2012). Actually, randomized directed evolution technologies require no prior knowledge of the tridimensional structure of the target protein. The evolution of the enzymes is solely directed by the assays used for screening for variants with improved characteristics.

Therefore, randomized “directed evolution” technologies have had an increasing impact to address the need for enzyme optimization. “Directed evolution” mirrors dramatically accelerates what has long been made when breeding plants and animals to produce new crops and livestock in order to fulfill the needs of mankind.

The successful outcome of enzyme directed evolution experiments depends on three major points which include:

- the potential of evolution of the enzyme(s) to be engineered (the so-called “parental” enzymes);
- the method chosen for generating the pool of next-generations enzymes;
- the screening method(s) applied for identifying the enzyme variants having appropriate characteristics.

Until the end of the 90’s, mutagenesis was the most widely used tool for generating new variants of an enzyme. The most popular technique for introducing mutations along a gene sequence has long been error-prone PCR (Polymerase Chain Reaction) and is still widely employed (Song et al., 2012). However, it has rapidly been observed that mutagenesis-based technologies suffer severe limits such as the relatively limited sequence space that can be rapidly explored, and a potential of evolution which is limited by the number of beneficial mutations that a gene can cumulate between two selection events. Some of these advanced methods have recently been extensively reviewed elsewhere (Dougherty and Arnold, 2009; Brustad and Arnold, 2011). In order to overcome the limits of classical mutagenesis-based approaches and optimize directed evolution protocols. This review will therefore develop in detail two major technologies developed by Proteus, one called EvoSight™ based on random mutagenesis and the other called L-Shuffling™ based on gene shuffling.

2.1 Random Mutagenesis

The random mutagenesis EvoSight™ method was designed in order to rationalize the directed evolution strategies and make it possible to run enzyme optimization programs within timelines that are compatible with industrial requirements (Fourage et al., 2006; Chodorge et al., 2005).

EvoSight™ is a three-step method. A first step consists in experimentally assessing the “plasticity” (i.e., the ability of a protein to accept mutations with a limited loss of activity) of the specific protein to be engineered, by rapidly evaluating the frequency of functional variants in a series of libraries produced by random mutagenesis under increasing mutational loads. The data generated by this first step are then processed using a mathematical model in order:

- to determine the optimal mutation load for that particular system;
to statistically determine the minimum number of clones to be screened to ensure that improved variants will be detected.

These results enable to strongly reduce the size of the library to be screened. After screening, the experimental determination of the actual evolution rate of the system (i.e. the frequency of improved clones in the screened library) enables to determine the potential of evolution of the specific enzyme/function system considered. The EvoSight™ algorithm also enables to determine the number of independent benefic mutations for that particular system and therefore provides guidelines for defining further directed evolution successful strategies.

### 2.2 Gene Shuffling

Since the late 90’s, *in vitro* “sexual” evolution has been developed to allow for poolwise recombination of the parental genes (Tobin *et al.*, 2000). Poolwise recombination enables mutations from many parental genes to recombine in a single progeny, thus dramatically increasing the number of positive mutations that can be cumulated between two selection events and enabling a much broader sequence space to be explored. The term “gene shuffling” was coined to name this approach. It appeared rapidly that recombining related parental genes has dramatically accelerated the rate of evolution. This new scheme achieves a much larger evolutionary distance of the best clones from the parental genes than when using classical random mutagenesis. The first *in vitro* methods for poolwise recombination of parental genes developed, such as DNA shuffling (Stemmer, 1994) or SteP (Zhao *et al.*, 1998), represented a major breakthrough in protein engineering. They were based on polymerase chain reaction-like recombination. Although extremely powerful, these pioneering methods also suffered limitations. The kinetics of hybridization of the megaprimers formed during the PCR-like recombination rounds used to limit the size of the genes that can be efficiently shuffled. Because of the inherent properties of the randomized PCR-like recombination methods, the libraries thus generated frequently contained a significant proportion of non-functional mutants. As a consequence, larger size libraries generally had to be screened so that positive hits are not missed.

To overcome these limitations, a new gene shuffling technique was developed involving no polymerase, called L-Shuffling™. This ligation-based random recombination method (Dupret *et al.*, 2000; Ravot *et al.*, 2005) permits the recombination of parental genes without the use of any polymerase, therefore reducing the risk of unwanted mutations. In this technology, the random recombination is assured by the ligation, by means of a suitable DNA ligase, of the ends of fragments of the parental gene variants hybridized onto an assembling template (*Fig. 1*). This ligation-based process enables randomized recombination, maintaining and combining the DNA information of the parental genes and generating a high proportion of functional variants.

Benefits of L-Shuffling™ include the ability of engineering both long and short genes as well as genes with high or limited homology. Most shuffled proteins generated by L-Shuffling are functional, which enables quicker screening and faster release of new products.

The impact of L-Shuffling™ for optimizing the characteristics of biocatalysts is well documented (Fourage *et al.*, 2005, 2009) and not limited to the engineering of enzymatic activity. For instance, the technology has been applied for antibody affinity maturation (Chodorge *et al.*, 2008). In this paper, the authors demonstrate that the most improved variant, with a 22-fold affinity gain, emerged from the recombination-based approach and not from a cumulative process such as error-prone PCR. Moreover, an analysis of mutations preferentially selected in the recombined population demonstrated strong cooperative effects when tested in combination with other mutations but small, or even negative, effects on affinity when tested in isolation. This study also demonstrated that the inclusion of an L-Shuffling™ recombination step enabled the selection of novel combinations of mutations, and therefore the exploration of a broader sequence space, compared with a parallel strategy which omitted recombination. This result is a strong argument for the combinatorial, poolwise-based approaches to protein evolution when compared to iterative mutagenesis based approaches.
3 MOLECULAR EVOLUTION OF Cel3a
BETA-GLUCOSIDASE

3.1 Experimental Section

3.1.1 Materials

p-nitrophenyl β-D-glucopyranoside (pNPGlc) was purchased from Sigma. Plasmid pET26b + used for creating L-Shuffling libraries was supplied by Novagen.

3.1.2 Construction of Plasmid pET26cay

The cDNA encoding the three parental glucosidases (A, B and C for β-glucosidases from Chaetomium globosum, Trichoderma reesei and Neurospora crassa respectively) was amplified by PCR with the specific primers (see the specific primers below)

Amplification was performed with 30 cycles of 94°C for 1 min, 60°C for 45 s, and 72°C for 1.5 min, with Pfu DNA polymerase (Promega) using the following PCR mix:

- 2 µL DNA template at 20 ng/µL;
- 2 µL of primers at 10 pmol/µL;
- 4 µL of dNTPs at 5 mM each;
- 10 µL of 10× buffer;
- 1 µL Pfu at 2.5 U/µL;
- 79 µL H2O.

After digestion using NdeI and HindIII restriction nuclease, the 2.1-kb PCR products were cloned into the pET26b + multicopy vector using the corresponding restriction sites leading to pETcay construct. After validation by automated DNA sequencing with the T7 promoter and terminator primers, the resulting plasmid, named pETcayB, A and C respectively, were used to prepare the three L-Shuffling libraries as described previously (Ravot et al., 2005; Ayrinhac et al., 2011).

3.1.3 Screening of L-Shuffling™ Library

For the first round of evolution, E. coli MC1061(DE3) colonies expressing β-glucosidase variants were grown in 96-well microtiterplates at 37°C during 4 hours and 20°C during 20 hours after induction with 100 µM IPTG in 150 µL of Luria-Bertani (LB) medium complemented with 60 mg/mL of kanamycin. For the second and third rounds, cells were grown during 20 hours at 20°C without IPTG induction. After centrifugation (4 min at 4 000 g) and resuspension in 100 µL of 100 mM succinate buffer at pH 5.0 comprising 2.2 mM of pNPGlc (in a presence of 80 g/L of glucose for the third round), cells were incubated for 3 hours at 23°C. After this incubation, activities were estimated spectrophotometrically by measuring the release of p-nitrophenolate ion at 414 nm after adding one volume of Na2CO3. In each plate, the values were compared to the value obtained with cells expressing the WT-glucosidases tested in the same conditions.

3.1.4 Characterization of Glucosidase Variants

Growth conditions used for the characterization of the β-glucosidase variants were the same as the ones used during the screening of the second round of L-Shuffling™ (variants were expressed without IPTG induction. Under such conditions, the level of expression is only driven by the leakage of the promoter and no detectable expression of the variants was observed on SDS-PAGE). After centrifugation (5 min at 8 000 g), E. coli MC1061(DE3) clones expressing the improved glucosidase variants were resuspended in 0.8 mL of 100 mM succinate buffer at pH 5.0 and different amounts of resuspended cell pellets were incubated 1.5 h using saturating concentration of pNPGlc at 50°C.

3.2 Results

In order to evolve β-glucosidase activity, a first round of L-Shuffling™ was carried out using two parental sequences sharing 70% of amino acid identity. For microplate expression, induced cultures using 100 µM of IPTG were needed in order to detect β-glucosidase activity of T. reesei. It should be noted that activity of the second parental protein (protein A) was not detectable under these screening conditions. Around 20 000 clones were analyzed as described in the experimental section using 2.2 mM of p-nitrophenyl

Specific primers

SgeneA (5’- GGAATTCATATGCTGAGGCCGCGACTG G -3’) and
ASgeneA (5’- CCCAAGCTTCTAGGCGGTCAGGCTGCC -3’);
SgeneB (5’- GGAATTCATATGTTTGTACCTCCTGCAGGGA -3’) and
ASgeneB (5’- CCCAAGCTTCTACGCTACCGACAGAGTG -3’); and
SgeneC (5’- GGAATTCCATATGGCTGAGGAGAGGATCG -3’) and
ASgeneC (5’- CCCAAGCTTCTAGTGAGGAGAGGATCG -3’).
β-D-glucopyranoside (pNPGlc) as substrate. Among the selected improved clones, 16 were sequenced showing large sequence diversity even if a hot-spot was suspected in the N-terminal part of the protein. As expected, due to the chosen L-Shuffling™ strategy, the skeleton of the improved variants was based on the reference protein. Figure 2 shows a sequence alignment of the 3 best variants together with parental genes BGL1 and A. The figure highlights the fact that comparison of improved variants can lead to identification of potentially important amino acids. Here the Histidine (H) in position 225 from gene A seems more efficient than the glutamine (Q) in the original Bgl1 gene, as this amino acid is found in every improved mutant.

The clone (164A2) selected as the best performer among the three tested ones when expressed in *Escherichia coli* was also analyzed for its β-glucosidase activity improvement on pNPGlc when expressed in *Trichoderma reesei* (Fig. 3).

Using saturating concentration of substrate, kat value of the wild-type β-glucosidase was assessed. A strong improvement of the specific activity was observed (Fig. 4) whatever the expression system used was. Based on these results, a second round of L-Shuffling™ was performed using the sixteen sequences shown in Figure 3 and introducing the WT β-glucosidase C as an additional parental gene. The same screening strategy was applied except that IPTG induction was no longer necessary due to the strong improvement achieved during the first round. Around 20 000 clones were analyzed as described in the experimental design using 2.2 mM of pNPGlc as substrate. Among the selected improved
clones, we determined the DNA sequence of 14 clones. The sequences confirmed the presence of a hot-spot in the N-terminal part of the protein (Fig. 4) demonstrating that this protein region is of importance for activity.

The clone (100B11) selected as the best performer among the three tested ones when expressed in *Escherichia coli* (see Fig. 5) was also analyzed for its β-glucosidase activity improvement on pNPGlc when expressed in *Trichoderma reesei*.

As already observed for 164A2 clone, a strong improvement of the specific activity (Fig. 6) was observed compared to native Cel3A β-glucosidase, whichever expression system was used. An 11-fold improvement was reached when 100B11 gene was expressed in *T. reesei*. Surprisingly, despite the high β-glucosidase activity of protein C in the tested conditions (data not shown), no DNA fragment of that β-glucosidase C was found in the sequence of the three best performers, although other less improved variants resulting from the same round of shuffling contained fragments originating from β-glucosidase C (Fig. 7).

The molecular diversity of the variants selected during the second round of directed evolution was limited, hence suggesting that a plateau of molecular evolution has been reached. In order to further enhance the β-glucosidase activity, the introduction of additional molecular diversity would be required. In the case of the subsequent round of evolution which aimed at improving another parameter using a new screening assay, such diversity can be provided by the parental genes.

As native Cel3A β-glucosidase is inhibited by high concentrations of glucose, which is also a bottleneck for production of 2G ethanol through SHF process, a new screening assay was defined for the third round of L-Shuffling™, based on the same conditions than the ones used for the second round, except that 80 g/L of glucose were added during the activity test. In such reaction conditions, around 20 000 clones were analyzed and one clone (149G7, see Fig. 7 for sequence) was selected.
for a better resistance to high glucose concentrations compared to Cel3A (Fig. 6).

As shown in Figure 8, an improved residual activity in the presence of 80 g/L glucose was also observed compared to 100B11 (best performer identified during the second round) when 149G7 gene is expressed in *Escherichia coli* and under saturating concentrations of substrate.

However, SDS-PAGE analysis has demonstrated (data not shown) that the observed improvement of activity per amount of cells (see Fig. 8) may be due to a better expression of the recombinant glucosidase 149G7.

In conclusion, the protein encoded by clone 149G7 shows a lower inhibition by high glucose concentration than the best variant from the second round of L-Shuffling™ experiment (protein 100B11) while conserving a specific activity in the same range.

**4 SACCHARIFICATION OF LIGNOCELLULOSIC BIOMASS**

Secretomes produced by *T. reesei* strain TR3002 expressing 100B11 gene and *Trichoderma* strain CL847 expressing native Cel3A were used as cellulolytic cocktails for steam-exploded wheat straw hydrolysis (Fourage *et al*., 2010). Kinetics were determined in shake-flasks and bioreactor (Bio-Laffite) under stirring conditions (500 tr/min). The wheat straw was pretreated by steam explosion (19 bar, 3 min) previously soaked in 0.05 M H₂SO₄ for 15 hours. After filtration, the solid phase mainly containing cellulose and lignin was suspended at 10% Dry Matter (DM) in a total volume of 50 mL for shake-flask experiments and 2 L for bioreactor experiments of 1 M acetate buffer at pH 4.8 and 48°C. Protein concentration was determined using the Folin method (Lowry *et al*., 1951). Enzymatic cocktails were used at 5, 10 and 20 mg per g of DM and kinetics were carried out for 72 h or 96 h. Shake flasks were done in duplicates. Bioreactor experiments were not replicated but 2 independent samples were taken for each time point. Enzymatic activities were inactivated by thermal denaturation and, after centrifugation and filtration, the released glucose was titrated with a glucose analyzer using glucose oxidase method (Analox, UK). The results obtained are shown in Figure 9 (shake-flasks) and Figure 10 (bioreactor). Figure 9 shows that for any enzyme dose TR3002 secretome outperforms the reference CL847. The effect is more visible at high enzymes...
doses, suggesting that at 5 mg/g DM, other activities besides β-glucosidase become critical for hydrolysis. This effect is even more pronounced for doses below 5 mg/g DM (our unpublished results). In bioreactor (Fig. 10), steering allows a more efficient mixing and suspension of the substrate, resulting in faster kinetics. After 24 h, the maximum yield (60 g/L of glucose) was obtained when TR3002 secretome was used as biocatalyst, while after 72 h of incubation this maximum yield was not held using the same amount of CL847 secretome.
(20 mg of secretome/g of DM, Fig. 10a). This major improvement enabled a 4-fold decrease in secretome loading (5 mg of secretome/g of DM) without any loss in activity performance of hydrolysis of a steam exploded wheat straw in bioreactor (Fig. 10b).

CONCLUSION

The present results demonstrate that molecular evolution technologies combined with an efficient high throughput screening assay enable to rapidly identify strongly improved variants for industrial purposes. In this particular example, we demonstrate how an evolved secretome resulting from in vitro recombination processes enables to lower the cost-contribution of the saccharification step in biofuel manufacturing cost. Despite no structural information was available for Cel3A β-glucosidase from Trichoderma reesei and no β-glucosidase activity was demonstrated for the two other parental proteins A and C using in this approach, enzyme variants showing up to 242-fold improvement in enzyme variants showing up to 242-fold improvement in cellulase activity is 11 times higher than in the secretome from the unmodified strain. This new secretome enables a 4-fold decrease in cellulase loading without any loss in hydrolysis performance for degradation of a steam exploded wheat straw in bioreactors.

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