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Exploiting cyclodextrins as artificial chaperones to enhance enzyme protection through supramolecular engineering[†]

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We report a method of enzyme stabilisation exploiting the artificial protein chaperone properties of β -cyclodextrin (β -CD) covalently embedded in an ultrathin organosilica layer. Putative interaction points of this artificial chaperone system with the surface of the selected enzyme were studied *in silico* using a protein energy land-scape exploration simulation algorithm. We show that this enzyme shielding method allows for drastic enhancement of enzyme stability under thermal and chemical stress conditions, along with broadening the optimal temperature range of the biocatalyst. The presence of the β -CD macrocycle within the protective layer supports protein refolding after treatment with a surfactant.

Introduction

Molecular chaperones are a family of multidomain proteins involved in the protein production machinery.¹⁻⁴ They support nascent proteins to achieve functional folding. They are also involved in unfolding misfolded proteins or folded proteins sentenced for proteolysis.^{5,6} Chaperones are also used to enable or increase yields of recombinant protein expression.^{7–9} In addition, they substantially increase protein stability under thermal stress conditions.^{10–12} Inspired by natural chaperones,



In our effort to stabilise enzymes and produce robust nanobiocatalysts, we worked on the concept of supramolecular enzyme engineering. It refers to a concept of enzyme supramolecular modification without manipulating the protein sequence. It aims at meticulously controlling the enzyme nano-environment. Applying this approach, we developed a method to protect enzymes immobilised at the surface of silica nanoparticles by growing, in a thickness-controlled fashion (down to nanometre precision), a protective organosilica layer.²⁷ The protection of the enzyme was shown to arise from a large set of supramolecular interactions established between the protein surface and the organosilica layer.²⁸ These interactions may also be optimised in order to engineer the active site of the protected enzyme and to switch a nonenantioselective enzyme into a selective one.²⁹

Herein, we report a method of enzyme immobilisation and supramolecular organosilica shielding. This method exploits

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Fig. 1 Schematic representation of chaperone-stabilised enzyme embedded in an organosilica layer.

the protein chaperone properties of a novel cyclodextrin building block, Fig. 1. The enzyme selected to establish the proof of concept of this method, a lipase from the *Bacillus* genus, when shielded in CD-containing organosilica, exhibits not only higher thermal stability but also an improved capacity to refold after chaotropic treatment. We expect this method to be versatile in that it can be applied to a range of enzymes for which thermal stability is an issue. This method can eventually contribute to improve the efficiency of industrial biocatalytic processes.

Results

As model enzyme, we used a lipase enzyme (Lip_{MRD9}) from the Bacillus genus (WP_034624255.1), identified by sequencebased metagenomic bioprospecting (see ESI[†] for details). First, we studied the possible CD-binding sites at the surface of this model protein. To that end, we employed the SiteFinder PELE protocol. The simulation encompassed 191 000 sampling steps, comprehensively exploring the protein surface. Fig. 2A illustrates the energy profile (created with Matplotlib library)³⁰ of the Lip_{MRD9}-CD interaction, where we monitored the ligand solvent accessible surface area (SASA) versus the protein-CD binding energy. Notably, few binding poses reached energies of $-60 \text{ kcal mol}^{-1}$, indicative of a robust interaction between the CD and the protein's surface. Furthermore, the lowest binding energies align with the minimum ligand SASA and minimum total energies of the system, reinforcing the concept of heightened affinity in that specific protein region. A focused analysis was then performed on those poses with the lowest binding energy, based on filtering the CD orientation where secondary alcohols are required to be oriented towards the protein surface.²¹ This was justified by the fact that the CD building block produced is modified at the primary rim; this is expected to drastically reduce its ability to bind to the protein surface via the primary rim. Out of the 25 steps with a binding energy lower than -50 kcal mol⁻¹, 9 conformed to the correct orientation. Within these 9 steps, we identified three distinct binding sites (Fig. 2B).



Fig. 2 Results for the first round of SiteFinder PELE protocol simulations. Solvent Accessible Surface Area (SASA) *vs.* binding energy (kcal mol^{-1}) for the whole simulation (A). The energy profiles were created with the Matplotlib library. Representation of the top three CD binding sites identified during the SiteFinder PELE simulation; residues inside the CDs cavity and catalytic serine are shown in orange (B).

After identifying the most promising three CD binding sites, a subsequent round of the SiteFinder PELE protocol was initiated. In this phase, the protein bound to the three CDs was considered the receptor, while an additional CD molecule was designated as the ligand. This strategy aimed to comprehensively explore the protein's surface with the three top binding sites already occupied, shedding light on potential tendencies of CDs to bind proteins that were already engaged with other CDs. As depicted in the scatter plot in Fig. 3A, just a few simulation steps managed to surpass the -50 kcal mol⁻¹ barrier. However, upon closer examination of the poses exhibiting binding energy below $-40 \text{ kcal mol}^{-1}$, a total of 44 structures, we identified 17 accurately positioned, ultimately leading to the selection of 5 distinct binding sites. Consequently, a total of 8 binding sites were now discernible (Fig. 3B). For each of the identified CD binding sites, a structural analysis was conducted, and the results are presented in Table S1 (ESI[†]). As depicted in the table, every pose exhibits a



Fig. 3 Results for the second round of SiteFinder PELE protocol simulations. Solvent accessible surface area (SASA) *vs.* binding energy (kcal mol⁻¹) for the whole simulation of the protein (A). The energy profiles were created with the Matplotlib library. Visualisation of the Lip_{MRD9} receptor bound to 8 CDs, showcasing the binding sites identified in this subsequent simulation round in cyan, the previously selected in green, and catalytic serine is shown in orange (B).

minimum of 5 hydrogen bonds between the protein and the ligand, among other interactions. The residues found within the CD cavities comprise Tyr, Leu, Asn and Ile. Three representative poses showcasing these interactions are shown in Fig. 2B. This set of results collectively supported the hypothesis of CDs binding at the protein surface through synergistic hydrogen bonding and hydrophobic inclusion.

Considering that shield formation occurs *via* organosilane self-sorting at the protein surface followed by polycondensation,^{27,31} we decided to produce a CD-derivative that can be used as a layer building block, *i.e.* bearing a triethoxysilane moiety. To that end, native β -CD was reacted with 3-isocyanatopropyltrimethoxysilane in dimethylformamide (80 °C); Fig. 4. The reaction product, CD-TES, was characterised by means of nuclear magnetic resonance (¹H and ¹³C), Fourier transform infrared spectroscopy and mass spectrometry. Collectively, the results confirmed the successful car-



Fig. 4 Chemical synthesis of a CD-TES, a β -CD derivative bearing triethoxysilane functions, $n \approx 1.5$ as estimated by ¹H NMR.

bamate bond formation with an average of 1.5 moiety per CD macrocycle (Fig. S2–5, ESI†).

Our model enzyme, Lip_{MRD9} (specific activity 19.3 U mg⁻¹) was immobilised at the surface of amino-modified SPs (average diameter 290 nm ± 20 nm) following a procedure previously published.²⁷ The SPs were further reacted with a mixture of tetra-ethyl-orthosilicate (TEOS), aminopropyl-triethoxysilane (APTES) and CD-TES to yield SPs with immobilised Lip_{MRD9} and shielded within a CD-containing organosilica layer, hereafter referred to as SP-Lip_{MRD9}-OS_{CD}. Reference particles, SP-Lip_{MRD9}-OS_{REF}, were produced by omitting the addition of CD-TES. The particles produced were characterised by scanning electron microscopy (SEM); Fig. 5. The micrographs showed, for SP-Lip_{MRD9}-OS_{CD}, a close-to-linear increase of the particles' diameter at a rate of *ca.* 6.5 nm h⁻¹; Fig. 5E.

It was accompanied by a moderate increase in surface roughness over the reaction duration, without impact on the polydispersity index which remained of 0.005. The layer growth kinetics for SP-Lip_{MRD9}-OS_{REF} followed a similar trend confirming the lack of relevant effect of CD-TES on the layer growth kinetics; Fig. 5F.

Next, we measured the biocatalytic activity of the immobilised enzymes using an established spectrophotometric assay of *p*-nitrophenyl butyrate hydrolysis; Fig. 6. The specific activity of SP-Lip_{MRD9} was measured to be 14.0 U mg⁻¹. For SP-Lip_{MRD9}-OS_{CD}, an activity increase to 127% and 133% (normalized to the specific activity of SP-Lip_{MRD9}) was measured for layer thickness values of 4.2 and 8.1 nm, respectively. When the layer was thicker at 10.9 and 14.5 nm, the activity slightly decreased to 118% and 97%. A similar trend was found for SP-Lip_{MRD9}-OS_{REF}; this is in good agreement with results obtained with other enzymes such as β -galactosidase or esterase.²⁷⁻²⁹

Next, we measured the influence of the artificial chaperone building block on the enzyme resistance to thermal stress conditions (Fig. 7A). To that end, the nanobiocatalysts produced were incubated at 50 °C for increasing durations. The soluble enzyme showed a decrease of activity to values of 62.1, 28.8 and 7.8% after 10, 20 and 30 min of reaction. This corresponds to a half-life of *ca.* 13 min. SP-Lip_{MRD9} and SP-Lip_{MRD9}-OS_{REF} displayed similar trends, with a slightly higher thermal stability than the soluble enzyme, with half-life values of 15 and 17 min, respectively. The stability of SP-Lip_{MRD9}-OS_{CD},



Fig. 5 Scanning electron micrographs of SP-Lip_{MRD9}-OS_{CD} measured after 30 (A), 60 (B), 90 (C) and 120 (D) min of layer polycondensation reaction. Layer thickness measured *via* a statistical analysis carried out on at least 100 SPs measured by SEM for SP-Lip_{MRD9}-OS_{CD} (E) and SP-Lip_{MRD9}-OS_{REF} (F). Scale bars represent 200 nm. Error bars represent standard deviation measured on at least 100 nanoparticles.



Fig. 6 Enzymatic activity measured for SP-Lip_{MRD9}-OS_{CD} (A) and SP-Lip_{MRD9}-OS_{REF} (B) values are normalised with the activity measured in the absence of shielding layer; error bars represent standard deviation measured on triplicates.

however, is markedly higher, with values of activity of 93.3, 76.8, 53.6% after 10, 20 and 30 min of incubation. This corresponds to a half-life of *ca.* 32 min and represent an increase of 246% when compared to the soluble enzyme. This consistent increase in enzyme stability may be directly attributed to the presence of the β -CD macrocycle in the protective layer. A stronger set of interactions between the surface of the protective shell and that of the protein may contribute to limit the



Fig. 7 Enzymatic activity measured at 50 °C during 60 min incubation for SP-Lip_{MRD9}-OS_{CD} (green), SP-Lip_{MRD9}-OS_{REF} (blue), SP-Lip_{MRD9} (orange), and Lip_{MRD9} (red) (A). Temperature profile for SP-Lip_{MRD9}-OS_{CD} (green) and Lip_{MRD9} (red) (B). Error bars represent standard deviation measured on triplicates.

detrimental thermodynamics effects of temperature on the protein conformation. Further, the biocatalytic activity the nanobiocatalysts produced was also studied at increasing reaction temperatures (Fig. 7B). The soluble enzyme, Lip_{MRD9}, showed an optimum activity at 30 °C and activity values over 80% in the range of 25–35 °C. The engineered counterpart, SP-Lip_{MRD9}-OS_{CD}, showed a maximum activity value at 40 °C and maintained more than 80% of activity over a broad range of 25–55 °C. The shift in the optimal temperature value and the relevant broadening of the temperature profile may be attributed to enzyme interactions with the support matrix. Collectively, these results bring compelling evidence of the stabilising effect of β -CD in the engineered organosilica layer.

Thermal and chemical protein denaturation are taking place following different molecular mechanisms involving different thermodynamic pathways.³² The encouraging results obtained with thermal stabilisation prompted us to test the stabilisation effect of the chaperone-based protective layer against chemical stress conditions. First, we tested the influence of an anionic surfactant, namely sodium dodecyl sulphate (SDS). Lip_{MRD9}, SP-Lip_{MRD9}-OS_{REF}, and SP-Lip_{MRD9}-OS_{CD} were incubated for 20 min in a buffered solution containing SDS (1%) prior to activity measurements; results are shown in Fig. 8A. The activity of Lip_{MRD9} dropped to 3%; activity values measured for SP-Lip_{MRD9}-OS_{REF} and SP-Lip_{MRD9}-OS_{CD} were consistently higher, with values of 15% and 21%, respectively. This demonstrated a moderate positive impact of the β -CD on the protein stability. Additionally, the possibility to refold the enzyme after SDS treatment was tested. To that end, Lip_{MRD9}-OS_{REF} and SP-Lip_{MRD9}-OS_{CD} were submitted to centrifugation and resuspended in Tris-HCl buffer. For the soluble enzyme, the surfactant was removed by dialysis against the same buffer. Activity measurements showed that while Lip_{MRD9} recovered only 20% of activity, SP-Lip_{MRD9}-OS_{REF} and SP-Lip_{MRD9}-OS_{CD} showed significantly better results with 68 and 84%, respectively. This effect was previously observed for soluble cyclodextrins and a carbonic anhydrase enzyme.¹⁸ It was attributed to the capacity of the cyclodextrin macrocycle to



Fig. 8 Enzymatic activity measured after SDS (1%) treatment for Lip_{MRD9} , SP- Lip_{MRD9} -OS_{REF}, SP- Lip_{MRD9} -OS_{CD} (white bars), and after further dialysis (red, blue, and green bars, respectively) (A). Enzymatic activity measured after urea (6 M) treatment for Lip_{MRD9} , SP- Lip_{MRD9} -OS_{REF}, and SP- Lip_{MRD9} -OS_{CD} (B); error bars represent standard deviation measured on triplicates.

strip away surfactant molecule. In the present work, it is more likely that the enzyme recovered its three-dimensional structure owing to the appropriate positioning of the CD moieties in the organosilica matrix. This effect of protein imprinting in organosilica layers was previously demonstrated with viruses.³¹

Next, we studied the denaturing effect of urea on Lip_{MRD9} and engineered counterparts by measuring the enzymatic activity after 20 minutes incubation with urea solution (6 M); as shown in Fig. 8B, SP-Lip_{MRD9}-OS_{REF}, and SP-Lip_{MRD9}-OS_{CD} were able to retain 16 and 27% activity after treatment, respectively, while Lip_{MRD9} could only retain 7% of its initial activity. This set of results suggest that the organosilica layer surrounding the enzyme supports its refolding as strongly suggested by the activity recovery values measured. While it is expected that the soluble enzyme may undergo irreversible aggregation and that immobilisation may prevent that, the significantly enhanced recovery for SP-Lip_{MRD9}-OS_{CD} brings additional evidence of the stabilising effect of β -CD within the protective layer.

Conclusions

In summary, we have demonstrated that the use of a cyclodextrin tri-alkoxysilane derivative in the formulation of an enzyme *via* organosilica shielding provides the immobilised biocatalyst with improved stability. *In silico* modelling using a Monte Carlo-based simulation algorithm allowed the identification of putative binding sites, where the macrocycle interacts with the protein through synergistic hydrogen bonding and hydrophobic inclusion. Once immobilised at the surface of SPs and shielded in a CD-containing organosilica layer, the model enzyme exhibited thermal stability markedly higher than its counterparts. Additionally, the range of temperature at which the enzyme is active was broadened. When subjected to chaotropic stress conditions, supramolecularly engineered enzymes exhibited moderately higher stability. The protein's ability to refold in an active state, however, was markedly enhanced. This increased stability suggests that supramolecularly engineered enzymes with artificial chaperone systems could withstand harsh chemical and physical operational conditions. This is expected to provide a longer shelf life and potential for broader applications in biotechnology.

Materials and methods

Chemicals

Tetraethyl orthosilicate (TEOS, \geq 99%), (3-aminopropyl)triethoxysilane (APTES, \geq 98%), 3-(triethoxysilyl)-propylisocyanate (95%), Trizma® hydrochloride (Tris–HCl, \geq 98%), ammonium hydroxide (ACS grade, 28–30%), ethanol (ACS grade, anhydrous), 2-(*N*-morpholino)ethanesulfonic acid monohydrate (MES, \geq 99.0%) and ammonium bicarbonate (\geq 98%) were purchased from Sigma-Aldrich (Switzerland). β -Cyclodextrin (97%) was purchased from BLD Pharmatech (Germany). Dimethylformamide (DMF, anhydrous, 99.9%) and PierceTM BCA protein assay kit were purchased from Thermo Fisher Scientific (Switzerland).

Buffer solutions

Buffer solutions were prepared with nanopure water. ABC buffer: 10 mM ammonium bicarbonate, 1.5 mM MgCl₂, pH 7.8. MES buffer: 10 mM MES, 1.5 mM MgCl₂, pH 7.5. Tris buffer: 50 mM Tris-HCl, pH 8.5.

Characterisation

Activity assays and bioconjugation reactions were performed and monitored using a Synergy H1 spectrometer (BioTek, Switzerland) into a 96-well plate (Microplate 96 Well Half Area). Nanopure water (resistivity $\geq 18 \text{ M}\Omega$ cm) was produced with a Millipore Synergy® purification system (Merck). Particles were imaged using a Zeiss SUPRA® 40VP scanning electron microscope (SEM). ¹H and ¹³C NMR (400 and 100 MHz respectively) were recorded on Bruker Avance-III HD spectrometer in DMSO-d₆ or MeOD using residual solvent signals as the internal standard. Infrared (IR) spectra were collected in attenuated total reflection (ATR) using a single reflection diamond ATR and Agilent Cary 630 FTIR Spectrometer. High resolution mass spectra were obtained with Bruker MaXis 4G spectrometer (Electrospray ionisation). For SEM imaging, a solution of bare SPs, SP-Lip_{MRD9}-OS_{CD} and SP-Lip_{MRD9}-OS_{REF} in nanopure water was prepared and spread on a silicon substrate. The samples were allowed to dry at 20 °C under atmospheric conditions and were subsequently sputtercoated with gold for 15 seconds at 20 mA. Secondary electron micrographs were acquired using the InLens mode with an accelerating voltage of 10 kV at a magnification of 150 000×. Particle sizes were measured on the acquired micrographs using the ImageJ free software. 100 measurements, at least, were performed for each sample (Tables S3 and S4, ESI[†]). Statistical analysis of SEM micrographs on SP-Lip_{MRD9}-OS_{REF} is shown in Fig. S6 (ESI[†]).

Modelling and simulations

PELE (Protein Energy Landscape Exploration) is a Monte Carlo-based simulation algorithm that unfolds in two

phases.³³ In the initial phase, the ligand undergoes random translation and rotation, while the protein backbone is perturbed using a normal mode method based on an anisotropic network model (ANM).³⁴ Moving to the second phase, optimisation of all rotamers from the ligand and its surrounding side chains occurs through a rotamer library, followed by an overall structural minimisation. Subsequently, acceptance or rejection of the new conformation is determined according to the Metropolis criterion. PELE can be used to study a large number of biological problems, including mapping molecular mechanism,³⁵ enzyme engineering,^{36,37} or binding site finder.³⁸ This last case is the one of interest in this work. The Site Finder protocol in PELE is a tool that conducts a global exploration across the protein surface to identify an appropriate binding site for a specified ligand. For each processor, a protein-ligand replica simulation is created, where the ligand is placed randomly and explores a defined area of the protein's surface. Note that the random position depends on previously positioned ligands, aiming at exploring novel surface areas; with enough processors (>32), the whole surface will be explored. For the best bound poses, those with lower binding energy and correct positioning (secondary alcohols pointing to the protein surface) were selected and studied. In some cases, a residue was found inside the cavity of the CD macrocycle. The structure of the protein used in this work, Lip_{MRD9}, was generated using AlphaFold2,39 and prepared with the protein preparation wizard, Schrödinger (United States).⁴⁰ Then the structure was set up for the site finder protocol using 192 processors, for a total of 191 000 Monte Carlos steps.

CD-TES synthesis

The cyclodextrin derivative was synthesised according to the procedure of Yilmaz *et al.* with slight adaptations.⁴¹ Cyclodextrin was dried overnight at 70 °C prior to the reaction. 2 mmol of cyclodextrin was dissolved in 20 mL dry DMF. 2 equivalents (4 mmol) of 3-(triethoxysilyl)-propylisocyanate were added and the reaction mixture was stirred for 12 hours at 80 °C. The mixture was cooled to room temperature and precipitated with acetone. The precipitate, CD-TES, was washed several times with acetone, collected and dried under vacuum.

Lip_{MRD9} immobilisation and shielding

Extensive details regarding the sequence-based metagenomic bioprospecting and the source and production of Lip_{MRD9} are provided in the ESI section.[†] Silica nanoparticles (SPs) were produced following a method described by Cumbo *et al.*³¹ inspired by the Stöber method.⁴² The nanoparticles synthesised were diluted in 400 mL H₂O to obtain a solution of [SPs] = 3.2 mg mL⁻¹. APTES (260 µL) was added to the previous solution, and the reaction mixture was stirred at 20 °C, 400 rpm for 90 minutes. Subsequently, the suspension was centrifuged at 4000 rpm for 15 minutes, and the supernatant was removed. The particles were resuspended in nanopure H₂O (50 mL) and submitted to ultrasonic treatment. This washing step was repeated twice. The particles produced, SP-NH₂, were then resuspended in water (50 mL) and stored at 4 °C. To a

suspension of SP-NH₂ (10 mL, 3.2 mg mL⁻¹) was added glutaraldehyde (25%, 40 μ L). The reaction mixture was stirred at 20 °C, 400 rpm for 30 min. Then, the suspension was washed twice with nanopure water, resuspended using an ultrasonic bath and resuspended in 5 mL MES buffer to yield SP-Glu (6.4 mg mL⁻¹). To a suspension of SP-Glu (2 mL, 6.4 mg mL⁻¹) was added 2 mL (in MES buffer) Lip_{MRD9} (final concentration of 73.6 μ g mL⁻¹) to reach [SP-Glu] = 3.2 mg mL⁻¹. The resulting mixture was stirred at 20 °C, 400 rpm for 1 h to produce SP-Lip_{MRD9}. The nanoparticles were centrifuged at 700 rcf for 5 minutes. A BCA assay was performed on the supernatant collected after enzyme immobilisation, which showed that 81% of the enzyme was immobilised at the surface of SPs; A summary of the results is presented in Table S2 (ESI[†]).

SP-Lip_{MRD9} (1.8 mL, 3.2 mg mL⁻¹) was incubated with TEOS (4 μ L) and CD-TES (13 mg) in 10 mM ABC buffer with 1.5 mM MgCl₂ at 10 °C, 400 rpm for 1 h. Subsequently, APTES (1.6 μ L) was added to the reaction media. The resulting mixture was stirred at 10 °C, 400 rpm for 120 minutes. Aliquots of 400 μ L were collected every 30 minutes. The resulting nanoparticles were centrifuged at 700 rcf for 5 minutes, washed and resuspended in 400 μ L Tris–HCl buffer ([SP-Lip_{MRD9}-OS_{CD}] = 3.2 mg mL⁻¹). A control experiment was conducted by replacing β -CD derivative with a volume of 2 μ L of TEOS and following the same shielding procedure as previously described ([SP-Lip_{MRD9}-OS_{REF}] = 3.2 mg mL⁻¹). The resulting suspension was allowed to cure at 20 °C for 18 hours for further activity recovery and stored at 4 °C.

Lip_{MRD9} activity assay

Lip_{MRD9} activity was measured spectrophotometrically using *p*-nitrophenyl butyrate (*p*-NPB) as substrate at 410 nm. The reaction mixture contained immobilised/soluble Lip_{MRD9} (3 μ g mL⁻¹), 1 mM *p*-NPB in isopropanol and 50 mM Tris–HCl (pH 8.5).

Author contributions

AFK, MF, SSS and PS conceptualised the work. AFK carried out enzymatic assays. SAN carried out the SP synthesis and microscopic characterisation. SSS and PS contributed to biocatalytic results' analyses. ARUP synthesised and characterised CD-TES. RMT and VG were in charge of *in silico* modelling and interactions studies. PV contributed by enzyme bioprospecting, synthesis, expression, production, and characterisation. PS and SSS supervised the project. The manuscript was written through the contributions of all authors who gave their approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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