

Structure-based mutagenesis of *Penicillium griseofulvum* xylanase using computational design

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ABSTRACT

Penicillium griseofulvum xylanase (PgXynA) belongs to family 11 glycoside hydrolase. It exhibits unique amino acid features but its three-dimensional structure is not known. Based upon the X-ray structure of *Penicillium funiculosum* xylanase (PfXynC), we generated a three-dimensional model of PgXynA by homology modeling. The native structure of PgXynA displayed the overall β -jelly roll folding common to family 11 xylanases with two large β -pleated sheets and a single α -helix that form a structure resembling a partially closed right hand. Although many features of PgXynA were very similar to previously described enzymes from this family, crucial differences were observed in the loop forming the “thumb” and at the edge of the binding cleft. The robustness of the xylanase was challenged by extensive *in silico*-based mutagenesis analysis targeting mutations retaining stereochemical and energetical control of the protein folding. On the basis of structural alignments, modeled three-dimensional structure, *in silico* mutations and docking analysis, we targeted several positions for the replacement of amino acids by site-directed mutagenesis to change substrate and inhibitor specificity, alter pH profile and improve overall catalytic activity. We demonstrated the crucial role played by Ser44_{PgXynA} and Ser129_{PgXynA}, two residues unique to PgXynA, in conferring distinct specificity to *P. griseofulvum* xylanase. We showed that the pH optimum of PgXynA could be shifted by -1 to $+0.5$ units by mutating Ser44_{PgXynA} to Asp and Asn, respectively. The S44D and S44N mutants showed only slight alteration in K_m and V_{max} whereas a S44A mutant lost both pH-dependence profile and activity.

We were able to produce PgXynA S129G mutants with acquired sensitivity to the Xylanase Inhibitor Protein, XIP-I. The replacement of Gln121_{PgXynA}, located at the start of the thumb, into an Arg residue resulted in an enzyme that possessed a higher catalytic activity.

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Key words: endo- β -1,4-xylanase; homology modeling; glycoside hydrolase family 11; *in silico* mutagenesis; docking; xylanase inhibitor; site-directed mutagenesis.

INTRODUCTION

Endo-1,4- β -xylanases (EC 3.2.1.8 [EC]) catalyze the hydrolysis of the 1,4- β -xylanosidic linkage in xylan, one of the most abundant plant cell wall polysaccharides in nature. Arabinoxylans (AXs) are often highly complex polysaccharides.¹ In cereals, the 1,4- β -xylopyranosidic backbone is mainly substituted with monomeric L-arabinofuranosyl and D-glucuronate units by 1,2- and/or 1,3- α -glycosidic linkages.² Although AXs are minor constituents, they are important for the functionality of cereals in biotechnological processes such as bread-making, gluten-starch separation, animal feed, paper manufacturing, and bioethanol production.³ Family 11 xylanases (<http://afmb.cnrs-mrs.fr/CAZY>) are highly specific, displaying exclusive substrate specificity toward D-xylose containing

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substrates and preference for insoluble polymeric substrates, and thus often preferred for technical applications. Therefore, a full understanding of glycoside hydrolase family 11 (GH11) properties is a key step toward the engineering of new xylanases for various applications. The family is highly homologous,⁴ displaying a perfectly conserved jelly-roll framework in which two large β -pleated sheets and one α -helix form a structure that resembles a partly closed right hand.^{5,6} The catalytic machinery is composed of two glutamic acid residues, which function as the nucleophile and acid/base catalyst, located in the middle of a cleft formed by the twisted part of the β sheet and the α helix. The cleft can accommodate up to seven β -1,4 xylopyranose units.⁷ Each xylose is accommodated in a subsite which is given a negative or positive number dependent on whether it binds the glycone or aglycone region of the substrate, respectively, with glycosidic bond hydrolysis occurring between the -1 and $+1$ subsites.⁸ To date, three-dimensional structures of 18 GH11 xylanases are available. Although GH11 enzymes show very closely related three-dimensional structures, xylanases display a wide panel of specificities with regards to pH activity, thermostability, thermophilicity, inhibition sensitivity.^{9–17} Optimum pH values are highly variable within this family and range between pH 2.0 and 4.6 for acidophilic xylanases and between pH 5.0 and pH 6.5 for the so-called “alkaline” xylanases.¹⁸ Structural alignment of GH11 xylanases showed strong correlation between the nature of the residue hydrogen bonded to the general acid/base catalyst and the pH optimum of these enzymes; it is Asn in “alkaline” xylanases and Asp in those with a more acidic pH optimum.^{4,19} Site-directed mutagenesis studies on acidophilic adaptation have confirmed the importance of residues located in the neighborhood of the glutamic acids, in particular in the thumb and at the edge of the cleft^{19–21}

The GH11 xylanase from *Penicillium griseofulvum*, PgXynA, is unique in that, unlike the xylanases characterized so far, it does not contain the strictly conserved Asp/Asn residue characteristic of the low or high pH optimum xylanases. PgXynA represents a unique example of a natural xylanase with a Ser residue at this position. We recently enzymatically characterized the enzyme substrate and inhibition specificity.²² Like many other carbohydrate-active enzymes, naturally occurring proteinaceous inhibitors of xylanases are present in plant.¹⁷ PgXynA is one of the rare examples of a fungal GH11 xylanase insensitive to the wheat xylanase inhibitor XIP-I²². The XIP-I strategy for inhibition of GH11 xylanases involves substrate-mimetic contacts and interactions occluding the active site.²³ We postulated that structural determinants around the “thumb” region of PgXynA was key to XIP-I resistance, although it remains to be demonstrated.^{22,23} These specific characteristics of PgXynA make it an attractive model to study structure–function relationships in the GH11 xylanase family.

Here, we have adopted a rational approach to creating suitably modified xylanase mutants by using computational design methods to guide structure-based site-directed mutagenesis. The theoretical model, presented in this work, was used as the basis for *in silico* screening for selecting mutants retaining parental folding. We experimentally evaluated the impact of mutations on novel enzymes with changed substrate and inhibitor specificity, altered pH profile, and improved overall catalytic activity.

MATERIALS AND METHODS

Materials

E. coli DH5 strain was used for DNA manipulation and BL21 strain (DE3) *pLys* for protein expression (Novagen, Merck Biosciences, Fontenay, France). High purity salt-free oligonucleotides were from Invitrogen (Cergy Pontoise, France), restriction endonucleases and DNA modifying enzymes were from Promega (Madison, WI), pGEX-1 λ T expression vector, GST glutathione agarose beads, and thrombin protease from Amersham Biosciences (GE healthcare, Orsay, France), BugBuster and benzonase from Novagen, lysosyme, β -D xylose, and dinitrosalicylic acid from Sigma (Lyon, France). Low viscosity wheat arabinoxylan (LVAX) and xylo-oligosaccharides (X2, X3, X4, X5, and X6) were from Megazyme International (Wicklow, Ireland). Purified wheat arabinoxylan substrates with different arabinose to xylose ratio were kindly provided by C. Courtin (Katholieke University Leuven, Belgium).

Bioinformatics

The signal peptide^{24,25} was predicted using the TargetP 1.1 server (<http://www.cbs.dtu.dk/services/TargetP/>). Sequences were retrieved with Blast²⁶ from the ebi website (<http://www.ebi.ac.uk/blast/>), using PgXynA as the query sequence.²⁷ 2D alignments were obtained using ClustalW²⁸ or T-Coffee²⁹ from the ebi (<http://www.ebi.ac.uk/Tools/sequence.html>) and 3D alignments using ESPRIPT³⁰ (<http://esript.ibcp.fr/ESPrIPT/ESPrIPT/>).

Computational homology modeling

Homology modeling was performed, using Modeler 6v2,³¹ with the B chain of the xylanase of *Penicillium funiculosum* (PfxYnC; ITE1)²³ as reference sequence and target template. Several models of the PgXynA were generated to satisfy the spatial restraints issued from the alignment with the target protein. The model with the best score function and the correct stereochemistry, as verified by PROCHECK,³² was selected and minor repositioning was carried out through short minimization.³³ The superimposition of the target structure and the

Table I
Oligonucleotides Used in the Construction of pGEX1λT/PgXynA Mutants

Name	Oligonucleotide sequence (5' → 3')
R7Tfor	ATCACCCAGAACGAG ACC GGAAACCAACGGCGGC
R7Trev	GCCGCCGTTGGTTC GGT CTCGTTCTGGGTG
F14Yfor	ACCAACGGCGGCTACT ACT ACTCTTTCTGGACC
F14Yrev	GGTCCAGAAAGAGTAG T AGTAGCCGCCGTTGG
S44Dfor	TGGAAGAATTGCGG GAT TTTCACCTCTGGCAAGGGC
S44Drev	CCCTTGCCAGAGGTG AAA TTCCCGCAATTCTCCAG
S44Nfor	TGGAAGAATTGCGG AAAT TTTCACCTCTGGCAAGGGC
S44Nrev	CCCTTGCCAGAGGTG AAA TTTCCGCAATTCTCCAG
S44Afor	TGGAAGAATTGCGG AGCT TTTCACCTCTGGCAAGGGC
S44Arev	CCCTTGCCAGAGGTG AAAG CTCCCGCAATTCTCCAG
Q121Rfor	ATCTACAAGCATACT CGG GTCACCCAGCCTTCG
Q121Rrev	CGAAGGCTGGTTG ACC CGAGTATGCTTGATG
S129Gfor	CAGCCTTCGATCATT GCG GATTCTAGCACCTTCGACCAG
S129Grev	CTGGTCGAAGGTGCT AGA AT GCC AATGATCGAAGGCTG
ΔD130for	CCTTCGATCATTTCGCTAGCACCTTCGACC
ΔD130	GTGGAAGGTGCTAGACGAAATGATCGAAGGC

Mutations are in bold underlined.

models was carried out using DaliLite³⁴ from the ebi website (<http://www.ebi.ac.uk/DaliLite/>).

In silico mutation analysis

In silico mutation analysis was performed to identify the maximum number of mutations that would still guarantee a correct folding of the protein. Since the N-terminal sequence of GH11 xylanases is crucial for the thermostability and thermophilicity of the enzymes,^{9–14} the residues 1–30 of the PgXynA mature protein were excluded from *in silico* mutations. Mutational changes were exclusively computed in the variable loop regions to maintain PgXynA topology and activity. The residues were mutated either into their analogues in the sequence of *Trichoderma reesei* xylanase II (TrXynII; 1XYO)⁶ or into other alternatives issued from Blast to produce a modeled chimera. The stereochemistry of the chimera was verified using PROCHECK and the side chains were optimized through a quick energy minimization.³³ The potential energy of the final protein was consistent with that of the native model and no steric clash was observed with the mutated residues.

Substrate docking

A xylohexaose was built by β-1,4 linking a nondistorted xylopentaose, extracted from the complex with a xylan binding domain of *Streptomyces lividans* (1MC9),³⁵ and a xylose moiety added at the reducing end. The xylohexaose was positioned in the catalytic groove of the PgXynA model based upon the positioning of the xylotriose in the active site of the xylanase of *Bacillus agaradhaerans* (1H4H).³⁶ The (φ ; ψ) dihedral angles of the sugar chain were slightly revisited to fit into the groove with respect to the (φ ; ψ) xylobiose map.³⁷ The complex was optimized

throughout a cascade of energy minimization and molecular dynamic runs using Charmm forcefield (newly acquired Discovery studio, Accelrys[®]).

Site-directed mutagenesis, expression, and purification

Mutations were introduced into the pGEX-1λT/PgXynA plasmid using the QuickChange[®] XL site-directed mutagenesis kit (Stratagene) following manufacturer's recommendations using a pair of overlapping complementary oligonucleotides for each mutation designed to contain the corresponding nucleotide changes. The primers used in this study are given in Table I. *E. coli* BL21 (*pLys*) strain was transformed with pGEX-1λT/PgXynA mutants and recombinant clones grown at 37°C in LB media containing 50 μg mL⁻¹ ampicillin and 34 μg mL⁻¹ chloramphenicol with shaking at 200 rpm. Induction of expression, bacterial lysis, purification, and thrombin cleavage of recombinant proteins was performed as previously described.²²

Protein assays, electrophoresis, and mass spectrometry analysis

The protein concentration was determined using the Bio-Rad protein assay kit with bovine serum albumin as the standard. SDS-PAGE was performed in 12% (w/v) polyacrylamide gel (Bio-Rad, Marnes-la-Coquette, France) using a Pharmacia LMW electrophoresis calibration kit. Native IEF was carried out at 4°C in the Bio-Rad gel system, using ampholine carriers of pH range 3.0–6.0 (Sigma) and pI standards ranging from 4.45 to 8.2 (Bio-Rad). Proteins were visualized either by Coomassie or silver staining. MALDI mass spectra were obtained on an Ettan Maldi-Tof Pro (GE Healthcare Uppsala,

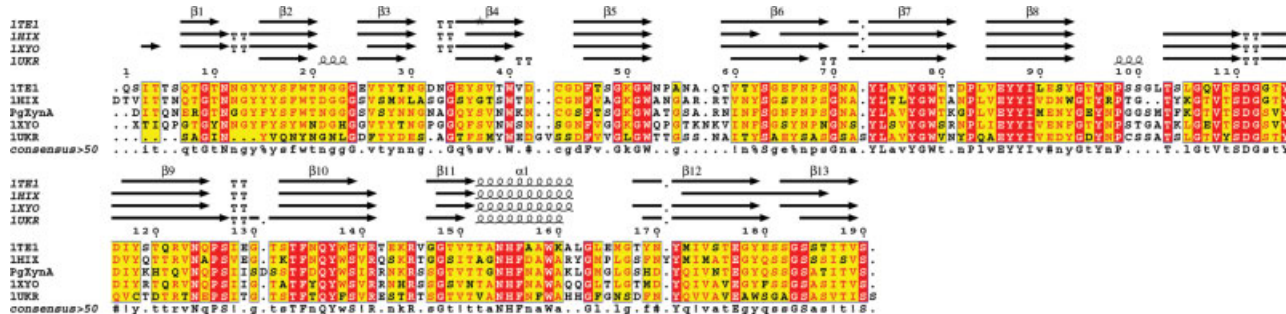


Figure 1

Structural alignment of xylanase sequences from *P. griseofulvum* (PgXynA),²⁷ *P. funiculosum* (ITE1),²³ *Streptomyces sp. S38* (IHIX),⁴³ *T. reesei* (IXYO)⁶ and *A. niger* (IUKR)⁴⁴ using ESPript. The secondary structural elements are shown above for the 3D solved structures. Strictly conserved residues are boxed in red and highly conserved ones colored in red.

Sweden) operating in positive linear mode with delayed extraction. The samples were treated as previously described.²²

Xylanase and inhibition activity assays

Purified xylanase activity was measured as described previously.³⁸ Briefly, 20 μL of enzyme (PgXynA wt or mutants) was mixed with 180 μL of substrate in McIlvaine's buffer pH 5.5 at 30°C for 5 min. The reaction was terminated by the addition of 300 μL dinitrosalicylic acid (DNS) reagent and boiled for 5 min. The reactions were cooled and centrifuged for 5 min at 13,000 rpm and 200 μL was transferred to a microtitre plate. The absorbance at 550 nm was measured relative to a xylose standard curve (0–180 $\mu\text{g mL}^{-1}$). One unit of xylanase activity was defined as the amount of protein that released 1 μmol of xylose/min at 30°C and pH 5.5. Optimal pH for xylanase activity was estimated using LVAX (10 mg mL^{-1}) in McIlvaine's buffer in a pH range of 2.4–7.5. For determination of apparent Michaelis-Menten constants, the initial velocities of the enzymes were measured at 30°C in McIlvaine's buffer, pH 5.5, with substrate concentrations ranging from 2 to 20 mg mL^{-1} . Because of the heterogeneous nature of polymeric substrates, their molar concentrations could not be calculated. Consequently, only an apparent value for the Michaelis constant, $K_{m(\text{app})}$, was determined. The kinetic parameters were estimated using weighted nonlinear squares regression analysis with the Grafit software (Biosoft, Cambridge, UK). Inhibition was detected by adding increasing molar equivalents of XIP-I³⁸ to the enzyme solution up to a maximum molar ratio of 30:1. The K_i value was determined at different concentrations of LVAX substrate (2–18 mg/mL) at 30°C and pH 5.5 in the presence of various amounts of XIP-I.

High performance anion exchange chromatography-pulsed amperometric detection

Products generated by hydrolysis of wheat arabinoxylans by wild type and mutated xylanases were analyzed by HPAEC (Dionex, Sunnyvale, CA) equipped with a Carbo-Pac PA-100 analytical column (250 \times 4 mm), a GP40 gradient pump and an AS3500 auto sampler. The hydrolysis was carried out using appropriate enzyme and substrate concentrations in McIlvaine's buffer pH 5.5. The enzymatic reaction was stopped by the addition of 1 M sodium potassium tartrate pH 11.5 and boiled for 5 min prior to injection (20 μL) on the HPAEC system. Elution was carried out as previously described.²² The effluent was monitored using an ED40 electrochemical detector.

RESULTS

Model of PgXynA and substrate docking

Since no crystal structure was available, a molecular model of PgXynA was generated using the *P. funiculosum* xylanase (PfxynC; ITE1)²³ as the template (Fig. 1). As expected, PgXynA exhibited the overall β -jelly-roll shape, typical of GH11 xylanases and reminiscent of a right hand, with the palm located between Thr152_{PgXynA} and Leu162_{PgXynA}, the cord by a loop positioned between Gly93_{PgXynA} and Lys104_{PgXynA} and the thumb by a hairpin loop located between Val122_{PgXynA} and Ser131_{PgXynA}. By analogy with other GH11 xylanases,^{5,6,23,39,40} the cleft is paved with strictly conserved aromatic residues that line up the active site with putative π stacking interactions.⁴¹ The carbonyl oxygens of the predicted catalytic residues, Glu85_{PgXynA} (catalytic nucleophile) and Glu177_{PgXynA} (catalytic acid-base), are located less than 6 Å apart in the model, which is consistent with the cata-

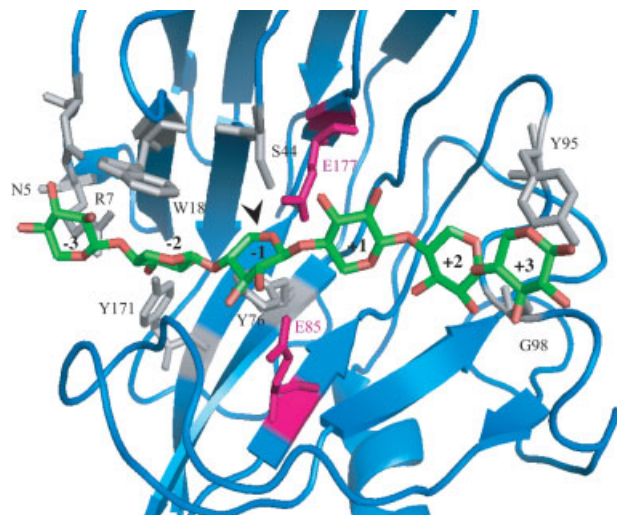


Figure 2

Close up view of the active site of PgXynA model structure, with the bound modeled xylohexaose. The residues predicted to participate in substrate binding through hydrogen bonding or van der Waals interactions are labeled. The xylohexaose molecule, modeled in the energetically most favorable conformation, is atom-sticked and atom-type colored in green. The distortion from 4C_1 to 2B_5 for the xylose moiety bound to subsite (-1) is indicated by an arrow.

lytic apparatus of a “retaining” glycoside hydrolase.^{39,40,42} The nucleophile belongs to a highly conserved cluster [V, I]-E-Y-Y [I, V], whereas the acid/base belongs to the less conserved [T, V]-E-G-Y motif.

The superimposition of the PgXynA model with that of the PfXynC 3D structure led to a root mean square deviations of 0.2 Å between the C α of 189 residues and a Z score of 38.1. The significant structural differences between PgXynA and PfXynC are mostly located between strand β_5 and β_6 (Ala52_{PgXynA}-Ser55_{PgXynA}) and within the thumb region (Ile127_{PgXynA}-Ser131_{PgXynA}) that includes the amino acid insertion of Asp130_{PgXynA}. Other differences concern amino acid side chains spread along the sequence defining the catalytic groove: Arg7_{PgXynA} versus Thr7_{PfXynC}, Phe14_{PgXynA} versus Tyr14_{PfXynC}, Ser44_{PgXynA} versus Asp44_{PfXynC}, Ala52_{PgXynA} versus Asn52_{PfXynC}, Met89_{PgXynA} versus Leu89_{PfXynC}, Gly98_{PgXynA} versus Ser98_{PfXynC}, Gly99_{PgXynA} versus Ser99_{PfXynC}, Met101_{PgXynA} versus Leu101_{PfXynC}, His119_{PgXynA} versus Thr119_{PfXynC}, Gln121_{PgXynA} versus Arg121_{PfXynC}, Ser129_{PgXynA} versus Gly129_{PfXynC} (Fig. 1). These residues differed in the length, charge and positioning of their 3D side-chains and were also distinct in other GH11 xylanases for example, xylanases from *Streptomyces* sp. S38 (1HIX),⁴³ *T. reesei* (TrXynII; 1XYO)⁶ and *Aspergillus niger* (1UKR)⁴⁴ (Fig. 1). Among those, Arg7_{PgXynA}, Phe14_{PgXynA}, Ser44_{PgXynA}, Gln121_{PgXynA}, and Asp130_{PgXynA} were targeted for experimental site-directed mutations (discussed later).

The 3D homology model was further challenged *in silico* to determine the robustness of the xylanase against mutations. A sequence variation of up to 16% was permitted to produce a xylanase retaining stereochemical and energetical control of protein folding. The permissive mutations are listed in “Supplementary Materials” and a xylanase chimera (not shown) was homology modeled based on TrXynII (XYO)⁶ that displayed most of the identified mutations (see MATERIAL AND METHODS).

To aid in the design of active site mutants, a modeled xylohexaose was docked in the active site of the PgXynA model, from subsites (-3) to (+3) (Fig. 2). The docking of the substrate revealed similar interactions with residues that contact the sugar rings in the experimentally solved structure of GH11 xylanases in complex with oligosaccharides and covalent inhibitors.^{5,6,23,36,39–44} From these studies, it can be suggested that Asn5_{PgXynA}, Arg7_{PgXynA}, and Trp18_{PgXynA} define the subsite (-3); Trp18_{PgXynA} stacks the xylose in subsite (-2), assisted by hydrogen bonding of Arg7_{PgXynA}, Tyr76_{PgXynA}, and Tyr171_{PgXynA}; Ser44_{PgXynA}, Pro125_{PgXynA}, and Glu177_{PgXynA} define subsite (-1). Ser44_{PgXynA} and Tyr72_{PgXynA} are engaged in hydrogen bonding of xylose in subsite (+1). The xylose in subsite (+2) appears hydrogen bonded to the carbonyl oxygen of Gly98_{PgXynA} and the reducing ring in subsite (+3) π -stacked to Tyr95_{PgXynA}. The PgXynA model is in accordance with previous kinetic analyses reporting the predominant production of xylo-triose from xylohexaose, suggesting that the specificity region of the xylanase spans about six xylose units.²² After several runs of molecular dynamic with no constraints, the xylose moiety located at subsite (-1) underwent a distortion from the 4C_1 chair conformation towards the boat 2B_5 geometry, as assessed by the measurements of the Pucker parameters.⁴⁵ This conformation allowed atoms C5, O5, C1, and C2 of the sugar in subsite (-1) to achieve coplanarity (Fig. 2). This is suggestive of a transition state conformation since a 2B_5 geometry was observed in the crystal structure of a xylo-triose bound to subsite (-1) of *B. agaradhaerens* xylanase³⁶ where it is essential to the formation of an oxocarbenium ion-like transition state, characteristic of the double-displacement catalytic mechanism. This distortion was also observed in the crystal structure of a 2-fluoro-xylose residue bound to subsite (-1) of *B. circulans* xylanase.³⁹ To summarize, the docking of a substrate analogue into the catalytic site of PgXynA highlighted Arg7_{PgXynA} and Ser44_{PgXynA} as relevant target for modifying substrate binding in GH11 xylanases.

Site-directed mutagenesis

Based upon sequence alignments and analysis of the PgXynA 3D model, mutations R7T, F14Y, S44A, S44D, S44N, Q121R, S129G, Δ D130 were introduced by site-directed mutagenesis and some combined in double

Table II

Specific Activities and Kinetic Parameters of wt and Mutant PgXynA Xylanases on Wheat Arabinoxylan (LVAX)

PgXynA	SA ^a (U mg ⁻¹)	K _{m(app)} (mg mL ⁻¹)	k _{cat} (s ⁻¹)	k _{cat} /K _{m(app)} (mL s ⁻¹ mg ⁻¹)	K _i (nM)
wt	1100 ± 84	3.8 ± 0.7	545 ± 29	143	N/A
R7T	1085 ± 71	11.9 ± 1.5	448 ± 25	38	N/A
F14Y	1226 ± 92	5.9 ± 0.8	773 ± 47	131	N/A
S44D	493 ± 31	4.4 ± 0.7	298 ± 17	66	N/A
S44N	954 ± 42	3.9 ± 0.4	454 ± 24	116	N/A
S44A	45 ± 3	4.6 ± 0.8	27 ± 2	6	N/A
Q121R	1464 ± 46	7.7 ± 1.5	1063 ± 86	138	N/A
S129G	764 ± 31	7.5 ± 0.9	484 ± 28	64	82
S129G/ΔD130	627 ± 40	6.4 ± 0.3	457 ± 8	71	3.9
ΔD130	1146 ± 34	5.7 ± 1.0	601 ± 42	105	N/A
S129G/S44D	95 ± 4	ND	ND	ND	ND
S129G/S44N	866 ± 22	6.5 ± 1.0	543 ± 45	83	90

N/A, not applicable; ND, not determined.

^aSpecific Activity at 30°C, pH 5.5 in McIlvaine's buffer, substrate concentration 10 mg ml⁻¹.

mutants. All mutant and wild-type (wt) xylanases were expressed in *E. coli*, purified as previously described and characterized in terms of molecular mass using mass spectrometry, pI and activity. The specific activity and kinetics parameters (*k*_{cat}, *K*_{m(app)}, *K*_i) of PgXynA xylanases were determined using wheat arabinoxylan (LVAX) as substrate (Table II).

Active site mutants

The R7T and F14Y mutants retained specific activity comparable to the wt-PgXynA (Table II). The activity of

the wild-type and Ser44 mutants was investigated at pH values between 2.4 and 7.5, using LVAX as substrate. The wt-PgXynA (pI 8.6), as also reported for PfXynC (pI 4.2), displayed over 50% activity in the pH range 4.5–6.5 with an optimum pH near 5.0 (Fig. 3). However, around 30% of activity was still observed at pH 7.3 for PgXynA whereas no activity was detected at this pH for PfXynC.²² Furthermore PgXynA retained full activity for 2 h in the pH range 4.0–7.5 at 30°C (not shown). In addition and in contrast to all GH11 “alkaline” xylanases characterized so far, PgXynA contains a Ser residue instead of the expected Asn residue adjacent to the acid-base catalyst (Fig. 2). This unique specificity of the *P. griseofulvum* xylanase A was investigated further by site-directed mutagenesis. The S44A mutation abolished the bell-shaped pH-activity profile of PgXynA whereas mutations to Asp or Asn resulted in a shift in the pH profile by ~0.5–1.0 pH units (Fig. 3). The S44D mutation shifted the activity to acidic pHs by ~1 unit, decreasing the optimum pH of PgXynA to 4.5 whereas the S44N mutation had the opposite effect, shifting the activity to alkaline pHs by ~0.5 with a pH optimum of 5.5. Moreover the S44D mutant had a broader pH profile retaining ~60% of its maximum activity at pH 3.0, where the wild-type enzyme is almost inactive (Fig. 3). This is consistent with previous mutagenesis studies of GH11 xylanases demonstrating correlation between the presence of an Asn or an Asp residue at position 44 (PgXynA numbering) and the pH optimum of the mutated enzymes. The substitutions led to an overall reduction in activity (S44N > S44D > S44A) mainly due to a reduction in *k*_{cat} whereas the apparent affinity remained unchanged (Table II).

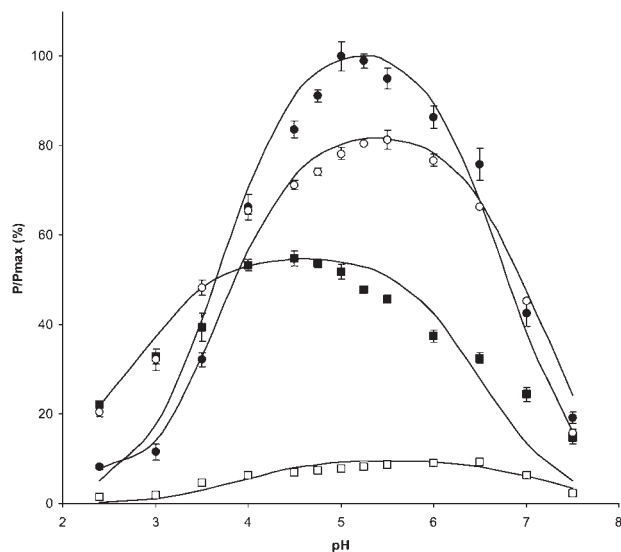


Figure 3

pH dependence of the wild-type (●) and mutant enzymes S44N (○), S44D (■), and S44A (□). The experimental data were fitted (solid lines) using the equation as described in de Lemos Esteves and coll.²¹ P represents the activity measured at various pH, and P_{max} the maximum activity observed for wild-type PgXynA at pH 5.0. The following pK_a values were used to fit the equation: wt, 3.7 and 6.7; S44D, 2.5 and 6.5; S44N, 3.5 and 7.2; S44A, 3.9 and 7.2.

“Thumb” mutants

Among the “thumb” mutants, the Q121R mutant showed a marked 33% increase in specific activity, mainly due to a two-fold increase in catalytic turnover (*k*_{cat}) (Table II). To evaluate the mode of action of the

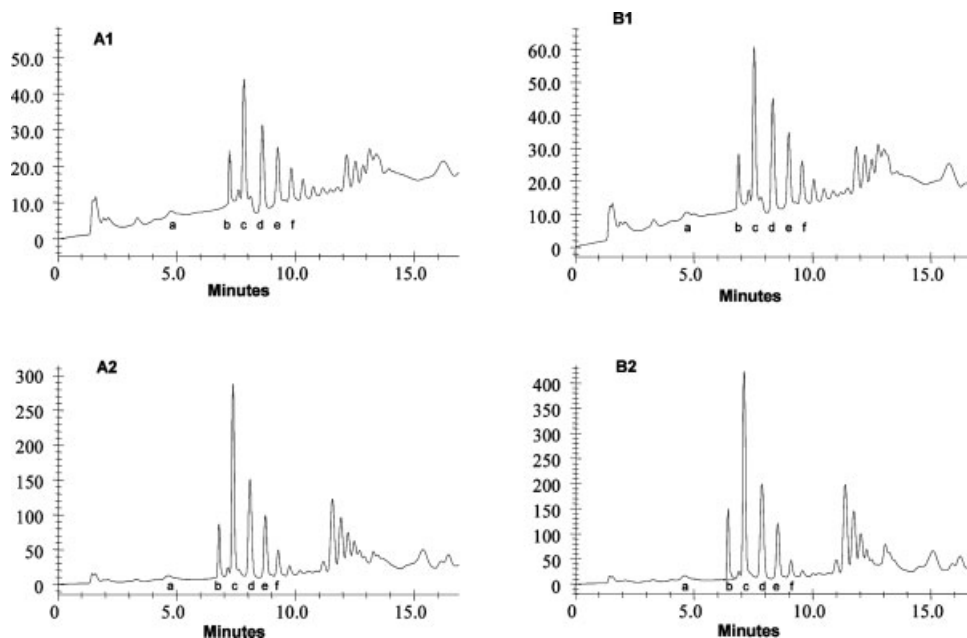


Figure 4

HPAEC analysis of wheat LVAX hydrolysis by PgXynA wt (A) and Q121R (B). PgXynA wt and mutant (20 nM) were incubated with 5 mg/mL wheat arabinoxylan in McIlvaine's buffer of pH 5.5 at 30°C. The reaction was stopped after 10 min (1) and 60 min (2), and subjected to HPAEC analysis: a, b, c, d, e, and f correspond to xylose, xylobiose, xylotriose, xylotetraose, xylopentaose, and xylohexaose, respectively.

mutant, the hydrolysis products generated on highly polymeric substrate were analyzed by high-performance anion exchange chromatography (HPAEC-PAD). The Q121R mutation did not alter the hydrolysis product profile of wheat arabinoxylan, that is, a mixture of xylo-oligosaccharides was generated during the initial stages of hydrolysis and as the reaction continued, the oligosaccharides were progressively degraded yielding principally xylobiose and xylotriose towards the end of the reaction, but an overall increase of released xylo-oligosaccharides products was observed (Fig. 4). The Q121R mutant was further tested on wheat arabinoxylan fractions of different arabinose/xylose ratio in order to test the effect of the substitution on the specific activity. The mutated enzyme showed highest increase in activity on low substituted xylan: A/X 0.36 > A/X 0.49 > A/X 0.68, with a maximum increase of up to 50% on A/X 0.36 compared to the wild-type (not shown). This is consistent with the generally accepted statement that GH11 xylanases preferentially cleave in unsubstituted regions of the AX backbone, whereas GH10 cleave in decorated regions, being less hampered by the presence of D-glucuronate and L-arabinofuranosyl substituents along the xylan backbone.^{2,42}

The wheat protein inhibitor XIP-I inhibits fungal but not bacterial GH11 xylanases. The structural basis for the inhibition came from the detailed analysis of the 3D structure of XIP-I in complex with PfxYnC.²³ Inhibition of the GH11 xylanase is mediated by the insertion of a

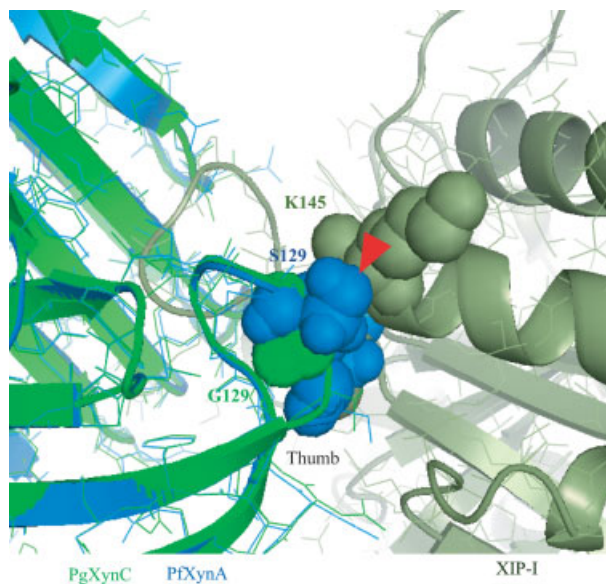


Figure 5

Evidence of steric hindrance between PgXynA and XIP-I. The inhibitor was extracted from its solved complex with PfxYnC (1TE1).²³ The backbone of PgXynA was superimposed onto the coordinates of the PfxYnC backbone, bound to XIP-I. PfxYnC and PgXynA are in green and blue, respectively whereas XIP-I is represented in light green. The Gly129_{PfxYnC}, Ser129_{PgXynA} and Lys145_{XIP-I} side-chains are in CPK spheres. The clash between Ser129_{PgXynA} and Lys145_{XIP-I} is indicated by an arrow.

XIP-I Π -shaped loop (148–153) into the enzyme active site but interactions with regions (thumb and palm) surrounding the entrance of the active site groove are also important recognition determinants for XIP-I, with the overall shape and amino acid composition of the thumb playing a significant role in inhibitor binding. As demonstrated earlier, the wt-PgXynA is insensitive to XIP-I²². In the present work, we showed that deletion of Asp130_{PgXynA}, protruding from the thumb region, did not alter inhibition sensitivity of the mutated enzyme whereas the S129G and the S129G/S44N substitutions conferred inhibition sensitivity with $K_i = 82$ nM and $K_i = 90$ nM, respectively (Table II and Fig. 5). The low specific activity of the S129G/S44D mutant prevented to study the effect of inhibition by XIP-I. However, a variant highly sensitive to the inhibitor was obtained when combining deletion of Asp130 with S129G mutation. The S129G/ Δ D130 mutant had a $K_i = 3.9$ nM, which is one of the lowest compared to tested natural enzymes.¹⁶ These results demonstrate that although deletion of Asp130_{PgXynA} is required to reach optimum inhibition level, the Asp130 insertion is not *per se* responsible for the lack of interaction with XIP-I and that Gly129_{PgXynA} plays a determinant role in the interaction with the inhibitor by ensuring correct positioning of the loop and preventing steric hindrance.

DISCUSSION

An enzyme in general is known for its specificity due to extreme sensitivity at crucial positions. Nevertheless, to maintain its folding and more importantly its function, an enzyme has to be robust to mutational events. Homology modeling with the target templates of PfXynC and TrXynII computed a native and a chimera model of PgXynA protein, respectively. The permissive level of *in silico* mutations reached 16%, confirming the predicted robustness of the GH11 xylanase folding.^{4,5}

In addition, bioinformatics, docking, and molecular dynamics were used to identify residues crucial for substrate binding, pH activity or motion flexibility. These predictions were validated through experimental site-directed mutagenesis. In particular, two residues highly conserved in other GH11 xylanases but different in PgXynA have been shown to be functionally significant. These are Ser44 and Ser129 (Fig. 1). Ser44 belongs to the highly conserved motif [GN/DFV/LxGKGW] where it replaces the conserved *Asp* or *Asn* residue found in “acidic” and “alkaline” GH11 xylanases, respectively.^{4,18–21,44,46} These neighboring residues of the acid/base catalyst determine the pH activity range of the enzyme following a “reverse protonation” mechanism dependent upon the electrostatic linkage of the groups involved in the interaction.⁴⁴ In the present study, we showed that we could drive the optimum pH activity of PgXynA toward the pH of the acidic or alkaline xylanases by mutating the Ser residue into an Asp or Asn

residue, respectively. However these mutations occurred at the expense of the activity since these groups may be involved in ground and/or transition state binding. In the PgXynA model structure with the modeled xylohexaose, the hydroxyl group of the Ser residue appears to be involved in bidentate hydrogen bonding of xylose residues in subsite (–1) and (+1), suggesting that it could participate in the distortion of the nonreducing xylose in (–1) towards its transition state geometry, thus leading to a boat geometry. Regarding the pH dependency of the enzymes, the side chain of the serine is significantly shorter than an aspartic acid or an asparagine residue so its participation in the protonation state of the neighboring catalytic Glu177 is jeopardized. The increased efficiency of the S44D mutant under more acidic conditions could be attributed to the formation of a shorter hydrogen bond between Asp44 side-chain and the acid/base catalyst Glu177 (Fig. 2). S44N has the same side-chain length as compared to its aspartate counterpart but does not contain any charge and thus cannot hydrogen bond Glu177. These results complement previous mutational studies where the Asp/Asn residues were substituted by their counterparts in other GH11 xylanases.^{4,18–21,44,46} Particularly relevant to the present work was the mutation of Asp to Ser in the acidic xylanase C from *Aspergillus kawachii*, leading to a shift in pH optimum from 2 to 4, thus slightly lower than the mutation to Asn which shifted the pH optimum of the mutant to 5.²⁰

The “thumb” region is known to play an important role in the interaction of GH11 xylanases with XIP-I, as first demonstrated by site-directed mutagenesis of xylanase from *A. niger*⁴⁷ and later on xylanases from *Fusarium graminearum*⁴⁸ and *Bacillus subtilis*.⁴⁹ In the present work, analysis of S129G, Δ D130, S129G/S44N, and S129G/ Δ D130 mutants showed that the insertion of Asp130 within the thumb region of PgXynA was not solely responsible for the lack of inhibition of the enzyme by XIP-I, as previously suggested.²² Deleting Asp130_{PgXynA} and substituting the adjacent Ser129_{PgXynA} by Gly, as it is in PfXynC, restored inhibition of PgXynA by XIP-I. In the model of PgXynA, the Asp130_{PgXynA} and Ser129_{PgXynA} side chains appear to clash against Asn147_{XIP-I} and Lys145_{XIP-I}, respectively, hampering interaction in that area (Fig. 5). Together, these results showed that the resistance of PgXynA to XIP-I is mostly due to the lack of flexibility of the loop, causing steric clashes between PgXynA and the inhibitor. A recent computational study of the thumb loop flexibility of a xylanase from *Thermobacillus xylanilitticus* using geometric path analysis indicated a motion amplitude of several Angströms.⁵⁰ High level of inhibition requires the loop to adopt an optimum conformation governed by the nature of the 129–130 residues in PgXynA.

Finally, we have improved the overall catalytic efficiency of some of the PgXynA mutants by mutating residues outside the core of the active site to avoid perturbing the exquisitely balanced network of complex interac-

tions that surround the catalytic groups. In particular the Q121R variant, located at the beginning of the “thumb”, showed higher specific activity on wheat arabinoxylan and retained product hydrolysis profile similar to the wild-type. Although Gln121_{PgXynA} is not seen to be involved in any interaction in the PgXynA model, substitution into an arginine leads to an extensive hydrogen bonding network with Gln124, Gln136, and the catalytic residue Glu85 (not shown). In addition, Arg121 appears to hydrogen bond the O₂ of xylose in subsite (-1) and O₄ and O₅ in subsite (+1), which could explain the enhanced affinity toward the substrate through stabilization of the ground state during catalysis (not shown).

CONCLUSIONS

In summary, we have devised a computational procedure to identify residues to be mutated with no change in the stereochemical and energetical control of the protein folding. Other mutations were suggested to modify substrate binding, pH activity or selective inhibition and experimental analysis confirmed that the procedure was successful in identifying xylanase mutants with novel specificity. This combination is a straightforward alternative to directed evolution and high throughput screening.⁵¹ By using molecular modeling, docking, *in silico* analysis and site-directed mutagenesis, we have generated knowledge to improve family 11 xylanases.

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