Loop engineering of a thermostable GH10 xylanase to improve low-temperature catalytic performance for better synergistic biomass-degrading abilities

Reporter: Shuai You
Background

**Pretreatment**

- Acid or alkali
- Ammonia recovery infiltration (ARP)
- Ammonia fiber expansion (AFEX)
- Steam blasting

**Biomass**

Xylanase + Cellulase

Improve efficiency

Wide industrial applications

Background

- Improve digestibility
- Reduce the risk of gastrointestinal diseases

Enzyme inactivation at high temperature granulation stage

Low enzyme activity at body temperature

Xylanase Ratio

- 90%
- 10%

Other enzymes (xylanase and others)

Background

Molecular modified

The Loop closed to the active center plays an important role in enzyme catalysis.

Rational design

Clustal Omega, NCBI, SWISS-MODEL et al.

WT: XYL10C_ΔN
Aim

Efficient degradation of lignocellulose

Enhance enzyme catalytic activity at low temperature

Target xylanase
Molecular docking and MD simulation \(\rightarrow\) Identify the mutation site (43 sites) and the design of primers

Mutated gene

\(\text{PIC9}_\gamma\)

(PCR) \(\rightarrow\) (Electric transfer GS115) \(\rightarrow\) (DNS)
Fig 1. Enzymatic properties of the purified recombinant wild-type XYL10C_ΔN and its three mutants.

(A) The optimal pH of each enzyme.
(B) pH-activity profiles tested at the optimal temperature for each enzyme (90°C).
(C) pH stability profiles.
(D) Temperature-stability profiles ($T_{50}$).
(E) Thermograms determined by using DSC.
(F) Half-lives of wild-type XYL10C_ΔN and its mutants at 65 °C.
Results

Thermostability:
M137E > M137/N269G > WT > N269G
Results

Stabilizes the conformation of the substrate binding channel

Enhances the interaction between the enzyme and the substrate molecule
## Results

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Beechwood xylan</th>
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<tbody>
<tr>
<td></td>
<td>Enzyme</td>
</tr>
<tr>
<td>XYL10C_(\Delta N)</td>
<td>1.39 ± 0.13</td>
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<tr>
<td>M137E</td>
<td>0.99 ± 0.03</td>
</tr>
<tr>
<td>N269G</td>
<td>1.02 ± 0.12</td>
</tr>
<tr>
<td>M137E/N269G</td>
<td>0.85 ± 0.06</td>
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</tbody>
</table>
Fig 2. Time-course hydrolysis of mulberry bark and corn cob. Separate hydrolysis: 5 U each of cellulase or xylanase simultaneous hydrolysis: 50 U each of cellulase and xylanase; control: no enzyme added (square) to substrates for 12 h (A) Mulberry bark (C) Corn cob. The DS curve for the mulberry bark samples are shown. Enzyme loading: cellulase 50 U, xylanase 50 U. The changes in the dry weight of the mulberry bark during separate and simultaneous hydrolysis with cellulases and xylanase after 24 h are shown (B) Mulberry Bark (D) Corn cob.
Fig. 3. Change in dry weight in three lignocellulosic substrates during separate and simultaneous hydrolysis with cellulases and xylanase after 72 h. (A) corncob. (B) mulberry. (C) wheat bran. (A) Mulberry bark (B) Corn cob
Fig. 4. Electron microscopy images of the micro structure of the mulberry bark samples treated with different enzymes. (A) Buffer treatment for 24 h (B) M137E/N269G alone (C) Cellulase alone (D) mixture of M137E/N269G and cellulase.

Fig. 5. Electron microscopy images of the micro structure of the corn cob samples treated with different enzymes. (A) Buffer treatment for 24 h (B) M137E/N269G alone (C) Cellulase alone (D) mixture of M137E/N269G and cellulase.
Results

Fig. 6. The effect of xylanases in different additions on the volume of bread. (A) The wild-type XYL10C_ΔN with different additions. (B) The mutant M137E/N269G with different additions.
Fig. 7. The effect of xylanase level in different additions on the specific volume of bread. (A) The wild-type XYL10C_ΔN with different additions. (B) The mutant M137E/N269G with different additions.
Conclusion

1. All the enzyme maintained similar pH and temperature optimal but the mutants showed notably improved catalytic performance under low temperature.

2. Loop region plays a vital role in hydrolysis of substrates, providing a reference for thermostable GH 10 xylanases engineering in improving their biochemical characteristics.

3. The successful improvement of XYL10C_ΔN makes the mutant produced in this study a potential xylanase for industrial applications on the degradation of lignocellulosic substrates even in diverse conditions.
Thanks for listening!

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