

Supplementary material for:

Characterization of *Zalaria obscura* Y1223 hydrolases: Implications for lignocellulose conversion

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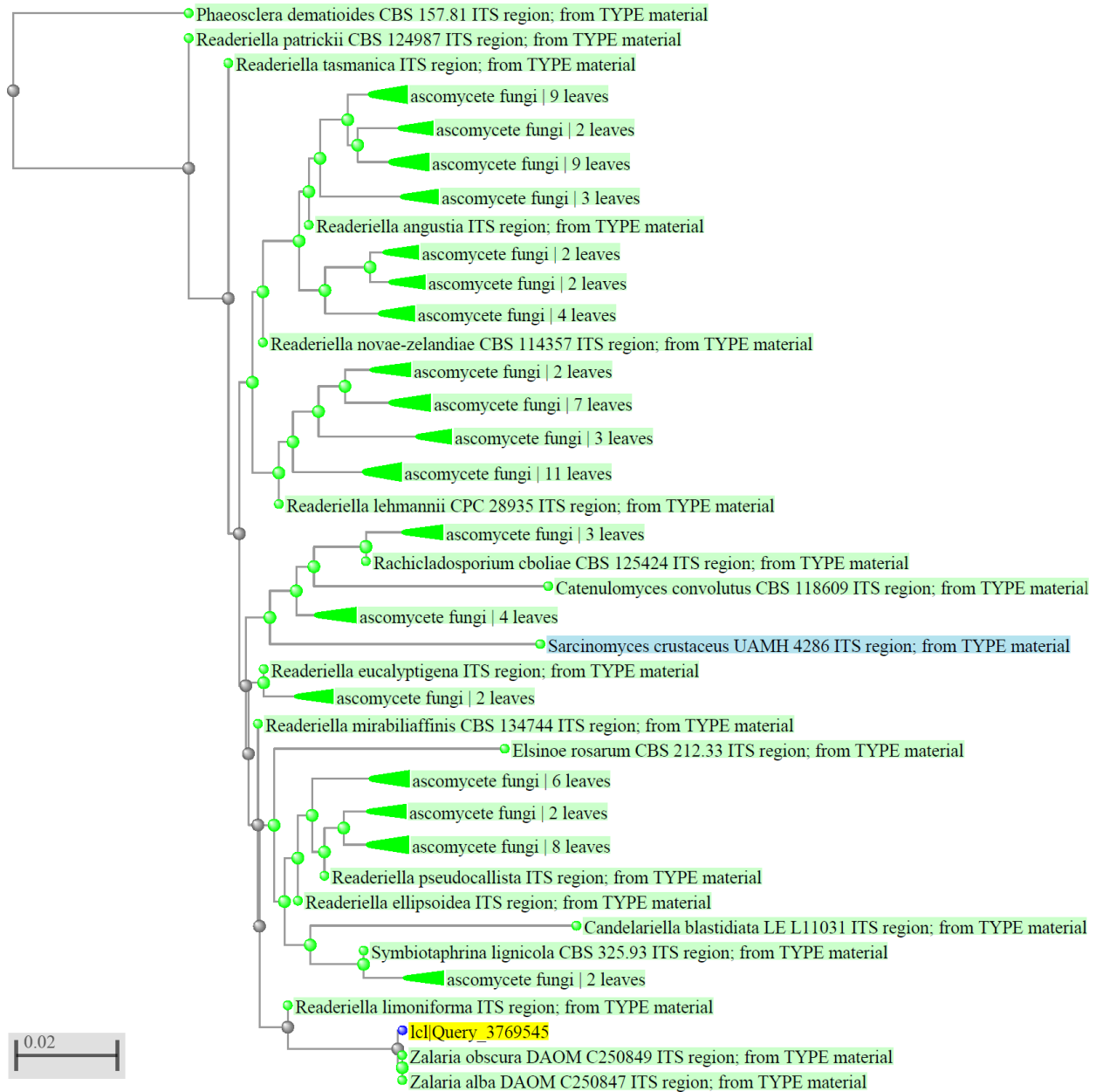


Fig. S1. Phylogenetic tree comparing ribosomal ITS sequences, to confirm taxonomic classification of *Z. obscura* Y1223 strain

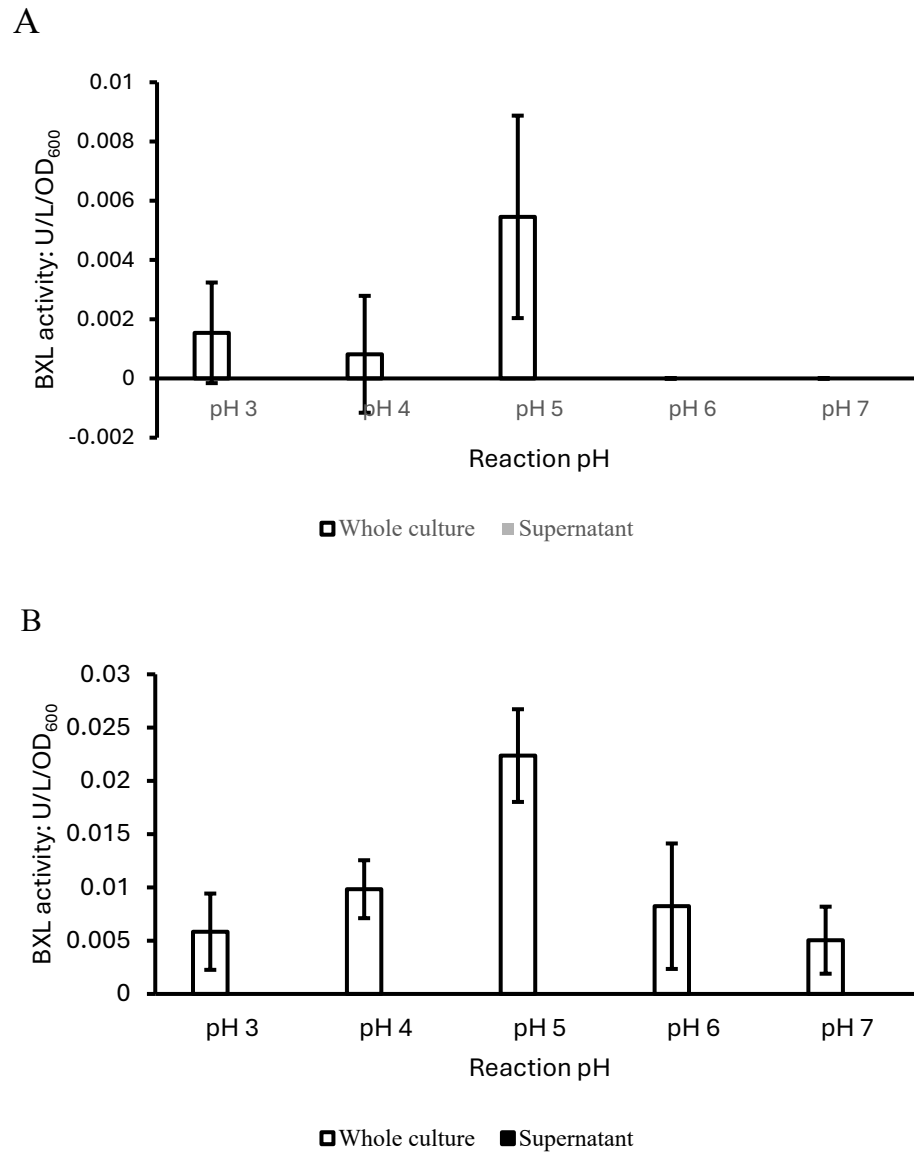


Fig. S2: *p*-Nitrophenyl (pNP) based activity assays performed at variable pH levels using citric-phosphate buffer pH 3-7. **A.** β -glucosidase (BGL) activity of *Z. obscura* Y1223 at variable pH levels. **B.** β -xylosidase (BXL) activity of *Z. obscura* Y1223 at variable pH levels.

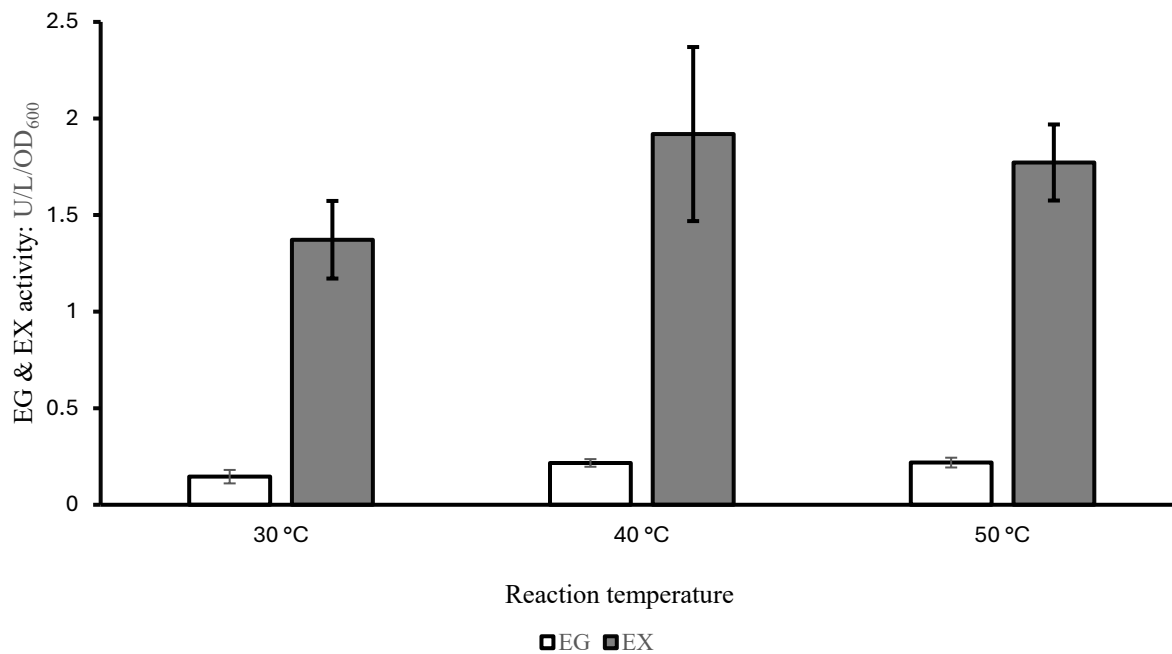


Fig. S3: Endoglucanase (EG) and endoxylanase (EX) activity of *Z. obscura* Y1223 cell free supernatant extracted from SC media supplemented with oat bran. EG and EX activity is shown using variable reaction temperatures of 30 – 50 °C.

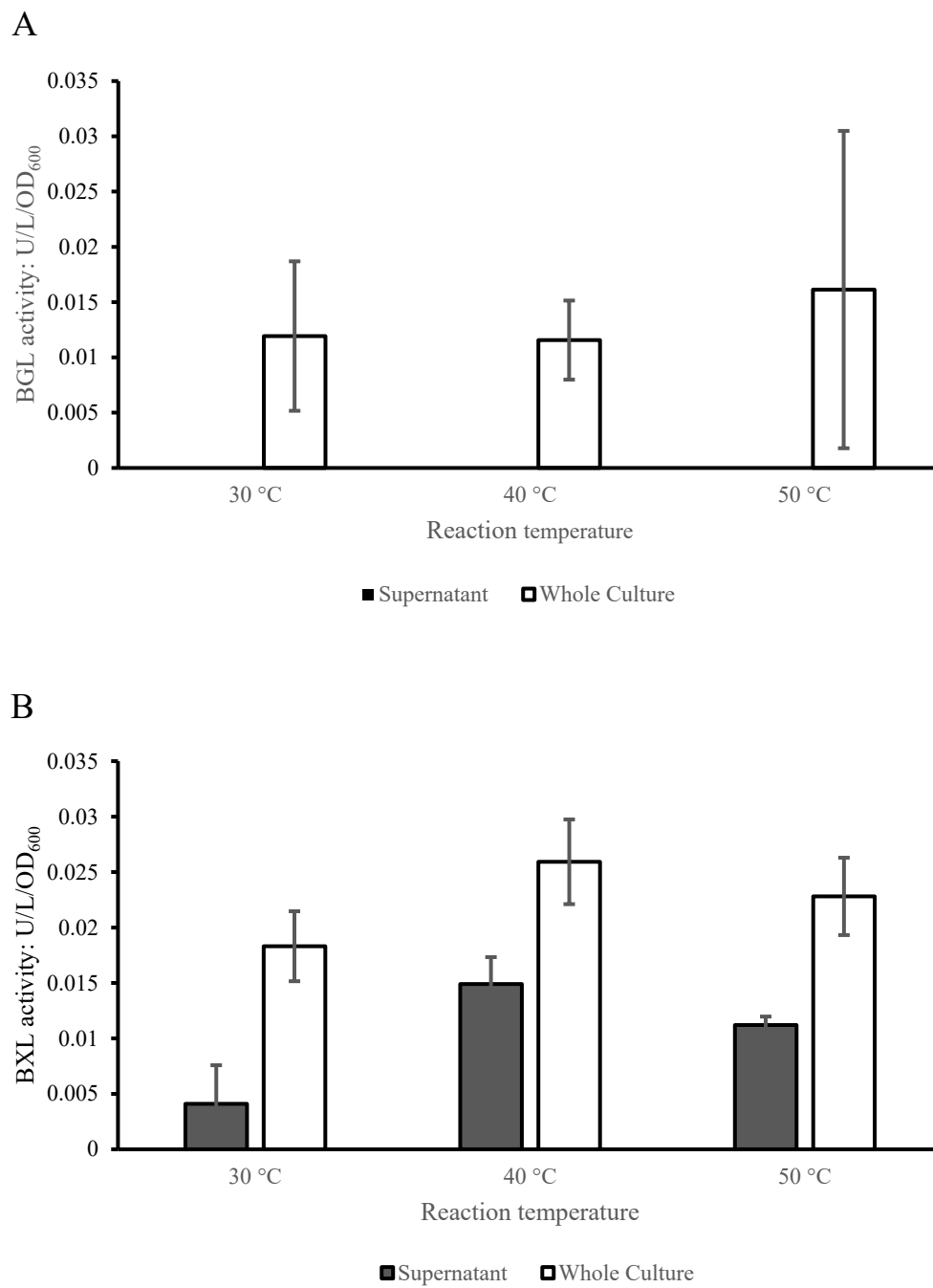


Fig. S4: *p*-Nitrophenyl (pNP) based activity assays performed at pH 5 and variable temperatures. **A.** β -glucosidase (BGL) activity of *Z. obscura* Y1223 at variable temperature. **B.** β -xylosidase (BXL) activity of *Z. obscura* Y1223 at variable temperature.