Bioprospecting Fungal Glycosyl Hydrolases: Multi-locus Phylogenetic Analysis and Enzyme Activity Profiling for Enhanced Biomass Valorization

Meenal Rastogi
Amity University

Smriti Shrivastava (shrivastava.smriti@gmail.com)
Amity University
https://orcid.org/0000-0002-5382-5378

Pratyoosh Shukla
Banaras Hindu University Department of Bio-Technology

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Abstract

Lignocelluloses comprise of celluloses and hemicelluloses which can be effectively depolymerized to obtain fermentable sugars using diverse microbial enzymes, for subsequent conversion to various value-added products. Present study reports the bioprospecting of industrially significant microorganisms and their characterization to attain xylanases with high catalytic efficiency. Four potential xylanolytic fungi were identified through distinct primary and secondary screening process of 294 isolates from samples containing plant degrades. Morphological characterization and multigene analysis (ITS rDNA, 18S rDNA, nLSU rDNA, β-tubulin and actin gene) confirmed them Aspergillus niger AUMS56, Aspergillus tubingensis AUMS60, Aspergillus niger AUMS64 and Aspergillus fumigatus AUKEMS24 and their crude xylanase activities through submerged fermentation using comcob were 18.9, 32.29, 30.68 and 15.82 U ml\(^{-1}\), respectively. AUMS60 and AUMS64 have highest catalytic activity of 1429 U g\(^{-1}\) and 1243 U g\(^{-1}\), respectively, all having pH and temperature optima of 6.0 and 60°C respectively, where AUMS60 produced single xylanase (Xyn60; 36 kDa) and AUMS64 secreted 2 probable isozymes (Xyn64A and Xyn64B; 33.4 and 19.8 kDa). Maximum saccharification efficiency of AUMS60 and AUMS64 were 51.1% (13 h) and 52.2% (24 h) showing enhanced catalytic activity with various cations. Present research reports potential xylanases from indigenous fungi, providing opportunity for development of bio-catalysts concoction (novelty established) for enhanced saccharification of complex lignocelluloses finding specific industrial applications for production of value-added components.

Key Points

- Crude activity of Xyn60 and Xyn64A, B as 1429 U g\(^{-1}\) and 1243 U g\(^{-1}\)
- Physico-chemical characterization of Xyn60, Xyn64A and Xyn64B
- Saccharification efficiency of Xyn60, Xyn64A and Xyn64B (all above 50%)

Introduction

Growing concerns over global warming and hike in fuel prices has necessitated usage of lignocellulosic biomass (most abundant natural biopolymer) as a renewable and eco-friendly source of current fuel platform. There is a strong impetus worldwide, for usage of biofuels that are economical and sustainable (Rastogi and Shrivastava 2017). Holocelluloses (cellulose and all of the hemicellulose) forms major fraction of lignocelluloses along with lignin which can be ideally hydrolyzed to monosaccharides for effective conversion to various value-added products. Traditional thermochemical methods utilized for degradation of lignocelluloses typically generate toxic by-products, therefore emphasis is laid on enzymatic saccharification considering the cost and yield in the biorefineries (Varghese et al. 2017; Sunkar et al. 2020). Lignocellulosic biomass is naturally degraded by diverse glycosyl hydrolases (GHs) producing microorganisms that can be utilized for several industrial as well as biotechnological applications. Cellulases, xylanases, lipases, pectinases, proteases and ligninolytic enzymes are few of the enzymes being utilized for the improvement of microbial and enzymatic processes for degradation of lignocelluloses (Rastogi and Shrivastava 2020).

Xylanases are ubiquitously found in nature produced by a variety of organisms but fungi secrete much higher levels of xylanases than other microbes in addition to several auxiliary enzymes required for the degradation of the substituted xylan (Polizeli et al. 2005). Xylanase cleaves β-1,4 glycosidic backbone of xylan (major hemicellulosic fraction) generally releasing xylose and xylobiose such as xylo-oligosaccharides as end products (Shrivastava 2020). In the last few decades, microbial xylanolytic enzymes have demonstrated a tremendous biotechnological potential in various fields apart from bioenergy production, including food for improving dough elasticity, animal husbandry for boosting the weight of chicks, clarifying juices for degumming of fibers, bread making, pre-bleaching of Kraft pulp, deinking of
waste newspapers and wine making (Chadha et al. 2019; Rastogi and Shrivastava 2020). Enzymatic saccharification of lignocellulosic biomass is a feasible alternative to physico-chemical treatment but its usage is limited owing to the high cost of enzymes. On the other hand, a vast array of indigenous microorganisms can be exploited for production of desirable glycosyl hydrolases via submerged (SmF) or solid-state fermentation (SSF) by substituting purified xylan with lignocellulosic biomass as substrate for providing the necessary nutrients for the microbial growth and the induction of the enzymatic production (Sunkar et al. 2020). SmF system accounts nearly for 90% of total xylanase produced worldwide (Polizeli et al. 2005) and is considered more flexible for allowing large-scale fermentations (Taddia et al. 2020).

Filamentous fungi, such as *Aspergillus*, easily adapt for cultivation on solid substrates due to simulation of lignocelluloses to their natural habitat and hence deliberated as good producers of hydrolytic enzymes for holocellulose degradation (Dias et al. 2018). Presently, the usage of acidic and alkaline xylanases in industries is compromised due to the environmental impact and high energy costs while neutral and weak acidic enzymes with high shelf life are gaining popularity due to reduced energy and expenditure input for their production and functions (Guo et al. 2012). At industrial scale, thermostable enzymes (operating at 45-100°C) not only improve enzyme robustness and enhance higher mass transfer and reaction rates decreasing the amount of enzyme required, but also reduce viscosity to increase the solubility of reactants and products, reduce risk of contamination and improve hydrolytic performance due to long half-lives at high temperatures (Bhalla et al. 2013; Bibra et al. 2018). Therefore, scientists are always exploring new microbial sources having higher activities with novel characteristics to improve efficiency for biomass valorization. Although a plethora of cellulo-xylanolytic strains have been reported, they are far from fulfilling our needs.

Present investigation reports bioprospecting of potential microorganisms for production of industrially significant xylanases with high catalytic activities, multi-locus phylogenetic analysis of selective organisms and partial profiling of xylanase enzyme activities for enhanced biomass valorization. These organisms were isolated from diverse waste residues rich in plant debris, through screening on enrichment media and characterized through morphological/physiological characteristics and multigene analysis (internal transcribed spacer (ITS) rDNA, 18S small subunit rDNA, 28S large subunit (nLSU) rDNA, β-tubulin and actin gene sequences). Two isolates selected for further studies AUMS60 (*Aspergillus tubingensis*) and AUMS64 (*Aspergillus niger*) were investigated on production and profiling of potential extracellular xylanase from them namely Xyn60, Xyn64A and Xyn64B. Profiling of physico-chemical characteristics of these enzymes suggested their industrial significance. Effect of metal ions, pH and temperature optima and stability suggests great catalytic potential of these enzymes.

**Materials And Methods**

**Agro-residues, chemicals and sample collection**

Agro-residues obtained from local farms were washed thoroughly, followed by oven drying and grinding to obtain particles of uniform size 1 mm, to be used as carbon source during screening and fermentation process. Xylose (Sigma, USA), Oat spelt xylan, birchwood xylan, carboxymethyl cellulose (CMC), bovine serum albumin (BSA) and dinitrosalicylic acid (DNSA) were procured from HiMedia Laboratories Ltd. Mumbai, India. All the other media, chemicals, salts and reagents used were of analytical grade (Sigma-Aldrich, St. Louis, MO, USA; HiMedia, Mumbai, India; and Merck & Co., Inc.). Optical compound microscope (Make-Magnus MLX-M) was used for microscopic analysis.

Samples were collected aseptically in soil rich in decaying plant matter and garbage waste from different locations of Noida and Delhi, India such as Okhla Bird Sanctuary (28.5514° N, 77.3185° E), Challera (28.5545° N, 77.3372° E) and Tajpur Pahari (28.4948° N, 77.3059° E) and stored at 4°C until further study.
Isolation and screening of GHs producing microorganisms

Serially diluted samples inoculated on nutrient agar (NA) and potato dextrose agar (PDA, 50 µg ml⁻¹ ampicillin) plates, were incubated at 20-60°C for up to 72 h for isolating indigenous glycoside hydrolase producing strains. Isolates were sub-cultured and stored at 4°C and as glycerol stocks. Czapek-Dox mineral salt agar medium and Remazol brilliant blue xylan supplemented medium was utilized for screening of cellulolytic (Teather and Wood, 1982) and xylanolytic (Shrivastava et al. 2011) microorganisms, respectively.

Screening through enrichment media and determination of xylanase and endoglucanase activity

Enrichment medium (Potato dextrose broth supplemented with 1% corn cob; agricultural residue; pH 5.5) was inoculated with pre-inoculum (10⁶ spores/ml). Extracellular enzymes were produced through medium containing (g l⁻¹) yeast extract (5), KH₂PO₄ (1) and MgSO₄.7H₂O (0.5) supplemented with 2% (w/v) corn cob (pH 5.5) at 40°C in a rotary shaker at 100 rpm for 10-12 days, with enzyme activity of cell free extract determined at interval of 24 h.

Xylanase activity was determined by monitoring the production of reducing sugars by 3,5-dinitrosalicylic acid (DNS) method (Miller 1959) with minor modifications using xylose as standard according to Bailey et al. (1992). One unit of xylanase activity was defined as the amount of enzyme that catalyzes the release of 1 µmol of xylose equivalent per minute under defined assay conditions. CMCase activity was determined according to Ghose (1987), with 1% CMC as substrate. One unit of CMCase activity was defined as the amount of enzyme that releases 1 µmol of glucose per ml per minute.

Identification of fungal isolates

Xylanolytic and cellulolytic microbial strains were examined microscopically using Lactophenol cotton blue (Leck 1999) and Grams stain (Bartholomew and Mittwer 1952) according to standard procedures. Four potential fungal strains with high titers of xylanase were identified based upon the ITS region, 18S rRNA, 28S rRNA, β-tubulin and actin gene sequencing at commercial facility provided by Chromous Biotech Pvt. Ltd., and Eurofins Genomics India Pvt. Ltd., Bengaluru, India. Genomic DNA was extracted from the selected fungal isolates according to Al-Samarrai and Schmid (2000) with slight modifications and used for amplification of ITS region, 18S rRNA, nLSU rRNA, β-tubulin and actin genes using primer pairs ITS1/ITS4 (White et al.1990), n-SSU-0817/nu-SSU-1536 (Borneman and Hartin 2000), LROR/LR7 (Vilgalys and Hester 1990; Rehner and Samuels 1994), BENA1/BENA3 (Davolos et al. 2012) and ACT-512F/ACT-783R (Carbone and Kohn 1999), respectively. All nucleotide sequences of the genes were submitted to NCBI GenBank database. Similarity search was carried out for each gene sequence using online BLAST program (http://www.ncbi.nlm.nih.gov/) and analyzed (Table 1) using the ClustalW algorithm available in molecular evolutionary genetic analysis (MEGA X) software with default parameters (Kumar et al. 2018). The phylogenetic tree of the ITS region was generated by employing the maximum likelihood method using Tamura-Nei model (Tamura and Nei 1993) and the evolutionary distance was inferred using the bootstrap method (n = 1000 bootstrap replications).

Phylogenetic analysis

Multigene phylogeny construction through alignment of relevant sequences and trimmed matrices construction of ITS and nLSU as well as ITS, nLSU and β-tubulin sequences was done through MEGAX. A Bayesian interference (BI) analysis was conducted via MrBayes v3.2.2 (Ronquist et al. 2012) using a GTR + I + Γ model (General Time Reversible model) for 3000000 generations with sampling every 100 generations. The settings were as follows: nst = 6, rates = invgamma, MCMC heated chain set with nchains = 4 and temp = 0.2, ngen = 3000000, samplefreq = 100. Two independent analysis with four chains each (one cold and three heated) were run until the average standard deviation of the split frequencies dropped below 0.01. The initial 25% generations of MCMC sampling were discarded as burn-in.
Refinement of the phylogenetic tree was used for estimating BI posterior probability (BIPP) values (Zheng et al. 2020). The tree was viewed in FigTree v1.4 (Rambaut 2012).

**Enzyme production, partial purification and molecular characterization**

Enzyme production from strains AUMS60 and AUMS64 was carried out as mentioned above with fermentation media supplemented with 3% (w/v) corn cob (pH 7); at 40°C on a rotary shaker (110 rpm) for 7 days. Filtered cell free extract was concentrated 10-fold via ultrafiltration membrane with a molecular weight cutoff of 10-kDa (Millipore TFE system, Bedford, MA, USA). Xylanase activity of all samples were determined as mentioned previously and protein quantification was done through Lowry's method using BSA as standard (Lowry et al. 1951).

Electrophoretic analysis of crude and partially purified samples was done through native and SDS PAGE (Ornstein and Davis 1964, Laemmli 1970, Shrivastava et al. 2013) with protein bands developed through silver staining (Blum et al. 1987) documented through Gel documentation system (Bio-Rad Laboratories, Hercules, CA, USA). Zymography was carried out with method of Royer and Nakas (1990) with slight modification. Reagents were procured from Biorad (Bio-Rad Laboratories, Hercules, CA, USA) and Himedia (HiMedia Laboratories Pvt Ltd., Mumbai, India).

**Biochemical characterization of partially purified enzymes**

Effect of pH on partially purified enzymes from AUMS60 and AUMS64 was studied at pH 5-8 at 40°C, 50°C and 60°C (Sodium acetate and Tris chloride buffer, 0.05 mol l⁻¹). Stability at pH 5 and 6 were studied for 120 h at 40°C. Temperature optima was determined by studying catalytic efficiency from 30°C to 90°C. Thermostability was studied at 40°C, 50°C and 60°C. Effect of selective metal ions and chemicals (NaCl, KCl, CaCl₂, CuSO₄, MgSO₄, FeSO₄, EDTA and SDS) in dose of 1 and 10 mM was studied by incubation at 50°C for 30 min with respective analyte.

Effect of enzyme dose (20-100 U g⁻¹ of substrate) was studied on saccharification of purified substrate (1% (w/v) birchwood xylan prepared in Acetate buffer, pH 6.0) upto 24 h with incubation at 40°C in rotary shaker at varying speed of 100-140 rpm.

**Results**

**Screening and selection of GHs producing biocatalysts**

Out of 294 indigenous bacterial and fungal strains isolated from samples at temperature ranging from 20-60°C, 46% (135) and 12% (35) were cellulolytic and xylanolytic respectively (Fig. 1a). Among all glycoside hydrolase positive strains 69% were bacteria and 31% were fungi with most of them having optimum growth temperature of 40-50°C (Fig. 1b and 1c), complimenting the site of sample collection.

Hydrolytic zone on substrate supplemented medium ranged from 0.1 to 4 cm (Fig. 1d). Approximately 40%, 11-18% and 15% of cellulolytic isolates showed hydrolysis zone ranging from <0.5 cm, 0.6-2.0 cm and >2.0 cm, respectively. In case of xylanolytic isolates 45.7%, 11-20% and 23% species exhibited hydrolysis zone <1 cm, 1.1-2 cm, and > 2 cm, respectively. All isolates with hydrolytic zone greater than 2 cm were selected for further screening through enrichment media. Four fungal strains identified as most potential glycoside hydrolyase producers exhibiting crude xylanase (X) and cellulase activity (C) were AUMS56 (X=18.9 and C=0.28 U ml⁻¹), AUMS60 (X=28.58 and C=0.44 U ml⁻¹), AUMS64 (X=24.86 and C=0.67 U ml⁻¹) and AUKEMS24 (X=15.82 U ml⁻¹). The strains were deposited in National Collection of Industrial Microorganisms (NCIM), National Chemical Laboratory (NCL), Pune, India with the accession numbers, NCIM1422 (AUMS56), NCIM1423 (AUMS60), NCIM1424 (AUMS64) and NCIM1425 (AUKEMS24).
Morphological characterization and identification of fungal strains by ITS region rDNA sequencing

All four selected isolates were spore forming filamentous fungi, observed through morphological examination. BLAST analysis for ITS region suggested that strains AUMS56, AUMS60, AUMS64 and AUKEMS24 showed maximum sequence identity to *Aspergillus niger* T4 (MG833314.1, 99.68%), *Aspergillus tubingensis* CNU081066 (JF411067.1, 99.6%), *Aspergillus niger* Z4 (MH892847.1, 98.97%) and *Aspergillus fumigatus* IHEM-18963 (KP131566.1, 98.23%), respectively. For the amplified 18S region, the closest sequence matches for AUMS56, AUMS60, AUMS64 and AUKEMS24 were to *A. niger* KY (EU184861.1, 98.5%), *A. niger* ETYB-13 (GQ903338.1, 99%), *A. niger* CBS-554.65 (NG_065763.1, 99.8%) and *A. fumigatus* ATCC-13073 (LC485158.1, 99.4%), respectively. For the amplified nLSU region, the closest sequence match for AUMS56 was to *A. terreus* (GU362936.1, 96.2%), AUMS60 and AUMS64 was to *A. niger* PVF1 (MK273554.1) with 96.1% and 98.03% maximum identity, respectively and AUKEMS24 showed 90.8% identity with *A. fumigatus* nHF-01 (MN190286.1).

For the amplified β-tubulin region, the closest sequence matches for AUMS56 and AUKEMS24 were to *A. terreus* (FR775342.1, 88.4%) and *A. fumigatus* (AY048754.1, 87.6%), respectively while AUMS60 and AUMS64 showed maximum identity of 97.3% and 91.1% to *A. pulverulentus* (HE984408.1). For the amplified actin region, the closest sequence match for AUMS56 was to *A. campestris* IBT-28561 (XM_024835858.1, 80.6%), AUMS60 and AUMS64 was to *A. niger* PL5-2B (MF422182.1) with 90.14% and 85.9% identity, respectively and AUKEMS24 was to *A. fumigatus* (AF076692.1, 98.9%).

The strains were designated based upon the ITS sequence similarity results of BLAST. Their corresponding accession numbers are mentioned in Table 1.

**Phylogenetic analysis**

Maximum likelihood phylogenetic tree, generated using ITS region of 40 related sequences through MEGA X and Tamura-Nei model computed the evolutionary distances. The bootstrap consensus tree inferred from 1000 replicates (Felsenstein 1985) was taken to represent the evolutionary history of the taxa analyzed as shown in Supplementary Fig. S1. Strains AUMS56 and AUMS64 formed a sister node to *Aspergillus niger* while AUMS60 formed sister node to *Aspergillus tubingensis*. AUKEMS24 formed a separate clade with sister node to *Aspergillus fumigatus*. Multigene Bayesian analysis involved sequences of the ITS and nLSU rDNA genes from 39 strains. The aligned dataset consisted of 575 and 860 nucleotides from the ITS and nLSU rDNA gene sequences, respectively. Convergence was assumed as an average standard deviation of split frequencies of 0.004545 was achieved following 3000000 generations. From the generated phylogenetic tree (Fig. 2), representative isolates of *Aspergillus* species clustered together forming 5 phylogroups. Isolates AUMS60 and AUMS64 formed lineage with *Aspergillus tubingensis* and *Aspergillus niger* in the first phylogroup with 100% Bayesian posterior probability support. The well supported sister lineage to this phylogroup was the *A. niger* branch consisting of nine strains. Isolate AUMS56 showed similarity to both *A. tubingensis* and *A. niger* lineages while AUKEMS24 formed discrete lineage with *Aspergillus fumigatus*, clearly showing deviation from *A. tubingensis* and *A. niger* groups with 100% posterior probability support. The species *A. flavus*, *A. keveii* and *A. sydowii* all branched as sister lineages forming distinct clades and therefore treated as separate phylogroups. Results obtained from multigene Bayesian analysis involving sequences of the ITS, nLSU rDNA and β-tubulin genes (data not included) from 23 strains were similar to those obtained from multigene analysis of ITS and nLSU rDNA genes.

**Characterization of partially purified xylanase from AUMS60 and AUMS64**

Crude xylanase activity of *A. tubingensis* AUMS60 and *A. niger* AUMS64 was further enhanced to 32.29 U ml⁻¹ (specific activity, 19.5 U mg⁻¹) and 30.68 U ml⁻¹ (specific activity, 18.4 U mg⁻¹), respectively, upon increasing the concentration of corn cob (though not optimized). Xylanase activity of ultraltrate (Amicon 10 kDa cut off filter) obtained from cell free crude extract of AUMS60 and AUMS64 was observed as 316.46 (specific activity, 37.2 U mg⁻¹) and 305.14 U ml⁻¹.
(specific activity, 27.1 U mg⁻¹), respectively. Zymography suggested single xylanase from AUMS60 and two isoforms from AUMS64. Xylanase have been designated as Xyn60, Xyn64A and Xyn64B having molecular mass of approximately 36, 33.4 and 19.8 kDa, respectively (Fig. 3).

Temperature optima for AUMS60 xylanase was determined as 60°C, while xylanases from AUMS64 showed almost same activity at 60°C and 70°C (Fig. 4a). Although all xylanases in the study significantly lost their activity at 80°C and above. pH optima for all the enzymes at varied temperature range was determined as 6.0 (Fig. 4b).

Stability of partially purified xylanases determined at pH 5 and 6 for 120 hours showed 90% retention in activity till 96 h at pH 6 while 70% activity at pH 5 for AUMS60. Xylanases from AUMS64 were also comparatively stable (>70% residual activity) at both pH 5 and 6 (Fig. 4c).

All xylanases, Xyn60, Xyn64A and Xyn64B worked at pH 6.0, were stable at 40°C for more than 120 h, with considerable loss at 50°C and almost complete loss in activity at 60°C (Fig. 4c and 4d). Although temperature optima for all xylanases was 60°C, they were highly unstable at that temperature. A comparative analysis of xylanases exhibiting differing pH and thermal stability over a wide range reported in previous studies from *Aspergillus* species is shown in Table 2.

Study of metal ions and chemicals on xylanases suggested Na⁺¹ and K⁺¹ as significant enhancer for Xyn60, Xyn64A and Xyn64B. Cu⁺² acted as activator at low dose and inhibitor at high dose for overall Xyn64A and Xyn64B; while inhibitor at all dose for Xyn60. MgSO₄ at lower dose acted as moderate enhancer for all Xyn60, Xyn64A and Xyn64B and inhibitor for Xyn64A and Xyn64B at high dose. CaCl₂ and FeSO₄ had varied effects at different dose on Xyn60, Xyn64A and Xyn64B. EDTA and SDS showed inhibitory effect on activity for Xyn60, Xyn64A and Xyn64B at all dose conditions (complete data detailed as Table 3).

Study of saccharification efficiency of all xylanases observed in our investigation suggested maximum saccharification with enzyme dose of 100 U g⁻¹ of substrate at 140 rpm, showing 51.1% at 13 h and 52.2% at 24 h incubation for AUMS60 and AUMS64, respectively (Fig. 5).

**Discussion**

Extensive isolation and screening procedure based on various identification techniques suggested AUMS6 (Aspergillus niger), AUMS60 (Aspergillus tubingensis), AUMS64 (Aspergillus niger) and AUKEMS24 (Aspergillus fumigatus) as most potential extracellular xylanase producers from all indigenous isolates taken for study. Complete identification of these organisms to species level was done through multigene analysis. AUMS60 and AUMS64 were found to be most ideal extracellular xylanase producers among 294 strains studied and formed a discrete clade with *Aspergillus tubingensis* and *Aspergillus niger* with 100% Bayesian Inference posterior probability. Multigene analysis could not be carried out by involving the sequences of 18S rDNA and actin genes because of lack of availability of resources in the database. AUMS60, AUMS64 and AUMS65 were both cellulolytic and xylanolytic, whereas AUKEMS24 was only xylanolytic.

A cocktail of biocatalysts producing cellulases and xylanases simultaneously alludes to be an ideal approach towards the saccharification of complex lignocelluloses to obtain simpler sugars in biorefineries. AUMS60 and AUMS64 could produce xylanase with crude extract activity of 19.69 and 17.32 U ml⁻¹, respectively over 3 days of incubation using corncob as substrate. Though a significant xylanase production was obtained within 3 days, it persistently increased reaching maximum xylanase activity of 28.58 U ml⁻¹ (1429 U g⁻¹) and 24.86 U ml⁻¹ (1243 U g⁻¹) for AUMS60 and AUMS64, respectively over 10 days. The activity was further enhanced to 32.29 U ml⁻¹ (AUMS60) and 30.68 U ml⁻¹
(AUMS64) in only 7 days upon changing parameters such as pH and concentration of corn cob (though not optimized). All these xylanases had pH optima as 6.0 and temperature optima as 60°C and were stable at both pH 5.0 and 6.0 for more than 120 h, but completely lost temperature tolerance even at 50°C after 48 h of incubation.

Corn cob has significant amount of xylan content as compared to numerous other industrially significant agricultural residues (wheat bran, rice husk etc), thereby making it a potential substrate for xylanase production (Knob et al. 2014). Moreover, the abundance and easy availability of corn cob in India facilitated for the screening of xylanolytic microbes. Bandikari et al. (2014) reported xylanase activity of 810.7 U g⁻¹ produced from Trichoderma koeningi (30°C/7 days) from corn cob. Streptomyces sp. MSC702 produced xylanase with activity of 6.23 U ml⁻¹ at 50°C incubation for 48 h (Singh et al. 2012). Xylanase production using corn cob alone or in combination with other substrates has also been reported for A. niger (10.5 U ml⁻¹) and A. flavus (11.92 U ml⁻¹) (de Alencar Guimaraes et al. 2013), Aspergillus flavus (65 U ml⁻¹) (Chen et al. 2019) and most recently Penicillium purpurogenum (84.61 U ml⁻¹, using delignified corn cob) (Sunkar et al. 2020).

Two isozymes of xylanase were secreted by AUMS64 (Xyn64A and Xyn64B) and one xylanase (Xyn60) by AUMS60 that were comparable in size to various other xylanases as in A. flavus (20.2 kDa) (Chen et al. 2019), A. terreus S9 (33 kDa) (Sharma et al. 2018), A. fumigatus (20 kDa) (Lin et al. 2017) and A. japonicus PJ01 (20-34 kDa) (Li et al. 2015). Several xylanases of this molecular mass have been successfully cloned and expressed in both bacteria and yeasts including E. coli, Saccharomyces cerevisiae and Pichia stipitis (Alokika and Singh 2019). Hence, our enzymes hold great potential for production of genetically modified organisms with enhanced saccharification efficiencies for complex lignocelluloses. Different fungal strains such as Penicillium ramulosum N1, Aspergillus sp., Clostridium and Streptomyces sp., A. fumigatus SK1 and N2 (Lin et al. 2017) have been reported for producing multiple forms of xylanase. These isozymic forms may have different biochemical properties and specific activity which may impart stability over broad range of pH and temperature. Microbial isolates producing isozymes with different yield and xylan hydrolysis efficiency are considered more suitable for industrial use (Singh et al. 2020).

The optimum pH of enzymes obtained in this study is similar to that observed for xylanases from A. clavatus (Pasin et al. 2020), A. niger (Suleman et al. 2016) and A. fumigatus (Carvalho et al. 2015; Lin et al. 2017). The optimal temperature is in agreement with xylanases produced by A. niger Gyx086 (Wang et al. 2019), Aspergillus cf. tubingensis LAMAI 31 (Dos Santos et al. 2016), A. fumigatus (Carvalho et al. 2015), A. niger and A. flavus (de Alencar Guimaraes et al. 2013) that presented optimal temperatures ranging from 50-60°C. The enzymes obtained in this study showed pH stability at both pH 5 and 6 over long periods of time (more than 72 hours), which is highly significant for their use in industrial processes operating for longer durations. Moreover, the enzymes work perfectly around pH 6, which is the optimal pH for many industrial processes. Enzymes from AUMS60 and AUMS64 showed high temperature stability at 40°C (greater than 120 h) and could also retain about 80% activity for 4 h and 90% activity for 24 h at 50°C, respectively. The results were more encouraging as compared to various other studies as mentioned in Table 2. Even if the pH stability is slightly less, the storage temperature for our enzymes (40°C) is higher than most of the studies (0-25°C) which indicate their application at even high temperatures.

As reported through previous studies, cations have enhancing or inhibitory action on enzyme catalysis, thus was observed for xylanases in our study with sodium and potassium resulting in 80-100% increase in residual enzyme activity and cupric ion showing potential inhibition. This study is ideally performed with purified enzymes and have been reported for xylanase from A. terreus S9, which was strongly inhibited in the presence of (10 mM) Hg²⁺ and Cu²⁺ while Mg²⁺, Fe²⁺, Co²⁺ and EDTA caused slight to moderate inhibition, but Ca²⁺, Mn²⁺ and K⁺ promoted the activity (Sharma et al. 2018). Xylanases from A. clavatus NRRL1 were found to show halotolerance, exhibiting a relative activity of 110% in the presence of 10 mM NaCl and maintaining more than 90% relative activity over the NaCl range of
Activity of partially purified xylanase from *Aspergillus oryzae* LC1 increased in presence of Fe\(^{2+}\), Mg\(^{2+}\), Mn\(^{2+}\), Co\(^{2+}\) and Ag\(^{2+}\) but inhibited by CuSO\(_4\), HgCl\(_2\), ZnCl\(_2\) and EDTA (Bhardwaj et al. 2019).

The presence of xylan intertwined with cellulosic component in the lignocellulosic biomass creates resistance for efficient conversion of biomass into fermentable sugars. Therefore, we evaluated the performance of the xylanase enzymes obtained from AUMS60 and AUMS64 for depolymerization of birchwood xylan. In our study, the saccharification efficiency of AUMS60 and AUMS64 xylanases was found to be 51.1% (4.84 mg ml\(^{-1}\) reducing sugars) and 52.2% (4.95 mg ml\(^{-1}\) reducing sugars) in 13 and 24 hours, respectively when used at an enzyme dosage of 100 Units g\(^{-1}\) substrate. A xylanase from *S. variabilis* MAB3 was reported to achieve maximum saccharification efficiency of 51.1% from birchwood xylan in 72 hours by Sanjivkumar et al. (2018). *Pleurotus ostreatus* xylanase (rXyn162) resulted in the release of 88.4 mg l\(^{-1}\) reducing sugars from oat spelt xylan (Zhuo et al. 2018). In another study by Tu et al. (2019), the levels of xylo-oligosaccharides released via depolymerization of wheat-arabinoxylan, reached 451.4 mg l\(^{-1}\) within 12 hours of incubation using xylanase from *Termoascus crustaceus* JCM12803. In view of the previous findings, the enzymes used in our study accomplish better saccharification of birchwood xylan in much less time even while using low dosages of enzyme. Moreover, we have used enzymes that are partially purified (one-step purification). Most of the previous studies have utilized enzymes that have been purified to homogeneity, which again increases the cost of the enzymatic saccharification process. In order to develop a cost effective and efficient process for enzymatic saccharification of biomass, maximal fermentable sugars need to be released while maintaining minimum expenditures on the cost of enzymes as well as on the hydrolysis time.

There is a surge of interest in the investigation for xylanolytic microorganism for their use in various industries. Two potent fungal strains of *Aspergillus* sp. in this study produced significant levels of glycosyl hydrolases on raw corn cob as substrate and the xylanases produced were of high catalytic strength. Profiling of physico-chemical characteristics of these enzymes namely Xyn60, Xyn64A and Xyn64B suggested their industrial significance. Effect of metal ions, pH and temperature optima and stability, catalytic strength of the enzymes provides great scope for further studying enzymatic cocktail for enhanced biomass valorization that can be highly beneficial since utilizing these microbes not only helps in waste reduction and management but also leads to development of cost-effective processes for production of enzymes and successive value-added components. To the best of our knowledge we report that the enzymes in our study hydrolyzed birchwood xylan with higher saccharification efficiency than most of the previous findings. Based on this work, process optimization (e.g. enzymatic hydrolysis of substrate, dosages of AUMS60 and AUMS64 xylanases) may be ensued to improve the transformation rates of hemicellulose from varied lignocelluloses. Our findings suggest that these enzymes are better candidates for production of reducing sugar from complex polysaccharides, enabling their use in the various industries.

**Declarations**

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**Availability of data and material**

All data generated or analyzed during this study are included in this article and its supplementary information files.

**Authors’ contributions**
MR conducted experiments, analyzed data and wrote the manuscript. SS contributed in supervision, funding acquisition, review and editing. PS advised for final correspondence of the manuscript. All authors read and approved the manuscript.

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**Compliance with ethical standards**

**Conflict of interest**

The authors declare that there are no conflicts of interest.

**Ethical statement**

This article does not contain any studies with human participants or animals performed by any of the authors.

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| Sl. No. | Species                | Strain      | GenBank accession number |
|--------|------------------------|-------------|--------------------------|
|        |                        |             | ITS                      |
| 1      | *Aspergillus niger*    | AUMS56      | MT802125.1               |
|        |                        |             | LSU                      |
| 2      | *Aspergillus tubingensis* | AUMS60    | MN153521.1               |
|        |                        |             | β-tubulin                |
| 3      | *Aspergillus niger*    | AUMS64      | MN153517.1               |
|        |                        |             | actin                    |
| 4      | *Aspergillus fumigatus* | AUKEMS24   | MN153510.1               |
|        |                        |             | 18S rRNA                 |
| 5      | *Aspergillus fumigatus* | CBS 542.75 | MH860951.1               |
|        |                        |             |                         |
| 6      | *Aspergillus fumigatus* | CBS 123.59 | MH857830.1               |
|        |                        |             |                         |
| 7      | *Aspergillus niger*    | CBS 111.55  | MH857397.1               |
|        |                        |             | AB201359.1               |
| 8      | *Aspergillus fumigatus* | CBS 386.75 | MH860929.1               |
|        |                        |             |                         |
| 9      | *Aspergillus fumigatus* | CBS 158.71 | MH860046.1               |
|        |                        |             |                         |
| 10     | *Aspergillus tubingensis* | CBS 116.36 | MH855725.1               |
|        |                        |             |                         |
| 11     | *Aspergillus tubingensis* | CBS 103.12 | MH854604.1               |
|        |                        |             |                         |
| 12     | *Aspergillus tubingensis* | CBS 425.65 | MH858651.1               |
|        |                        |             |                         |
| 13     | *Aspergillus tubingensis* | CBS 107.55 | FJ629367.1               |
|        |                        |             |                         |
| 14     | *Aspergillus tubingensis* | CBS 130.52 | FJ629359.1               |
|        |                        |             |                         |
| 15     | *Aspergillus niger*    | CBS 554.65  | FJ629337.1               |
|        |                        |             |                         |
| 16     | *Aspergillus niger*    | LBM 134     | MK457457.1               |
|        |                        |             |                         |
| 17     | *Aspergillus niger*    | ATCC 16688  | AY373852.1               |
|        |                        |             |                         |
| 18     | *Aspergillus niger*    | ATCC 16404  | AY939787.1               |
|        |                        |             |                         |
| 19     | *Aspergillus spelaeus* | IG136       | MG976683.1               |
|        |                        |             |                         |
| 20     | *Aspergillus keveii*   | CBS 209.92  | MH862354.1               |
|        |                        |             |                         |
|   | Aspergillus Species        | Strain   | Accession 1 | Accession 2 | Accession 3 | Accession 4 |
|---|---------------------------|----------|-------------|-------------|-------------|-------------|
| 21| *fumigatus* SZJ01         | MK240371.1 | MK240374.1 | -           | -           | MK240373.1  |
| 22| *sydowii* ASAU-1          | KJ524907.1 | KJ524908.1 | KJ524906.1  | -           | -           |
| 23| *terreus* RCBC_1002       | GU362937.1 | GU362936.1 | -           | -           | -           |
| 24| *fumigatus* ATCC 16907    | AY214446.1 | AY216670.1 | -           | -           | -           |
| 25| *fumigatus* CBS 126847    | MH864332.1 | MH875778.1 | -           | -           | -           |
| 26| *fumigatus* CBS 130584    | MH865795.1 | MH877225.1 | -           | -           | -           |
| 27| *fumigatus* CBS 129448    | MH865339.1 | MH876789.1 | -           | -           | -           |
| 28| *fumigatus* CBS 120.53    | MH857125.1 | MH868658.1 | AY685156.1  | -           | -           |
| 29| *fumigatus* CBS 114.55    | MH857399.1 | MH868939.1 | -           | -           | -           |
| 30| *niger* CBS 263.65        | MH858562.1 | MH870199.1 | -           | -           | -           |
| 31| *niger* CBS 105.47        | MH856174.1 | MH867698.1 | -           | -           | -           |
| 32| *niger* CBS 134.54        | MH857268.1 | MH868802.1 | -           | -           | -           |
| 33| *niger* ANG-1             | MH266204.1 | MH423841.1 | -           | -           | -           |
| 34| *niger* T4                | MG833314.1 | -           | -           | -           | -           |
| 35| *niger* ATCC 9029         | KU729033.1 | KU729117.1 | JF450851.1  | -           | -           |
| 36| *niger* CBS 117.36        | MH855726.1 | MH867238.1 | -           | -           | -           |
| 37| *niger* C5334             | AF109327.1 | AF109343.1 | -           | -           | -           |
| 38| *niger* CSR3              | MF187478.1 | MF187479.1 | -           | -           | -           |
| 39| *flavus* V5F-13           | HQ395774.1 | HQ395773.1 | -           | -           | -           |
| 40| *flavus* CHS1             | MF599088.1 | MF590165.1 | -           | -           | -           |
| 41| *tubingensis* CNU081066   | JF411067.1 | -           | -           | -           | -           |
| 42| *niger* Z4                | MH892847.1 | -           | -           | -           | -           |
|   | Aspergillus fumigatus | IHEM 18963 | KP131566.1 | - | - | - | - |
|---|----------------------|------------|------------|---|---|---|---|
Table 2

Comparison of characteristics of xylanases from other microorganisms

| Substrate | Microorganism | pH stability | Temperature stability | Xylanase activity | References |
|-----------|---------------|--------------|-----------------------|-------------------|------------|
| 3% (w/v) Corn cob (SmF) | A. tubingensis AUMS60 | >70% (96 h at 40°C) at pH 5.0 - 6.0 | Stable at 40°C (120 h) | 32.29 U ml⁻¹ | This study |
| | A. niger AUMS64 | >70% (72 h at 40°C) at pH 5.0 - 6.0 | Stable at 40°C (120 h) | >82% (4 h at 50°C) | 30.68 U ml⁻¹ |
| Wheat bran 0.5% (w/v) + Corncob 0.5% (w/v) (SmF) | Aspergillus niger | >95% (1 h at 0°C) at all pH | >90% (24 h at 50°C) | 10.5 U ml⁻¹ | de Alencar Guimaraes et al. 2013 |
| | Aspergillus flavus | decrease of 10% at pH 3.0 – 5.5; 100% at alkaline pH (1 h at 0°C) | half-life > 75 min at 50°C | 11.92 U ml⁻¹ |
| 3% (w/v) Sugarcane Bagasse (SmF) | A. fumigatus M51 | >70% at pH 4 to 9 (24 h at 25°C) | >60% (1h at 50°C) | 35.6 U ml⁻¹ | Carvalho et al. 2015 |
| | A. fumigatus U2370 | >61% (1h at 50°C) | >50% at 50°C | 28.5 U ml⁻¹ |
| 1% (w/v) Birchwood xylan (SmF) | Aspergillus cf. tubingensis LAMAI 31 | Stable at pH 3.6 to 7.0 (24 h at 40°C) | Stable at 40 - 50°C (1 h) | 49.41 U ml⁻¹ | Dos Santos et al. 2016 |
| 0.1% (w/v) Beechwood xylan (SmF) | Aspergillus niger (KP874102.1) | 100% at pH 2.0 to 5.0 (1 h) | >79% (60 min at 37°C) | 229 U | Uday et al. 2017 |
| 5% (w/v) Wheat straw | A. niger Gyx086 | Not available | >70% at 50°C (72 h) | 6.19 ± 0.27 U ml⁻¹ | Wang et al. 2019 |
| 2.67% (w/v) Wheat bran (SmF) | A. terreus S9 | >75% at pH 6 to 10.0 (90 min) | >75% at 60-80°C (90 min) | 116 U ml⁻¹ (optimized) | Sharma et al. 2018 |
| 3% (w/v) Corn cob (SmF) | Aspergillus flavus | >70% at pH 3.5 to 10.5 (30 min) | >90% at 50°C (30 min) | 65 U ml⁻¹ | Chen et al. 2019 |
| 1% (w/v) Rice straw (SmF) | Aspergillus oryzae LC1 | >84% at pH 5 (12 h); >50% at pH 7 and 9 (8 h) | >50% at 50°C (12 h) | 1245 IU ml⁻¹ (optimized) | Bhardwaj et al. 2019 |
| 1% (w/v) Paper sludge, 0.5% (w/v) sugarcane bagasse | Aspergillus clavatus NRRL1 | >85% at pH 3 to 8 (24 h) | 80% and 100% at 50°C (1 h) | Not available | Pasin et al. 2020 |
| Waste from flour industry (SmF) | Aspergillus niger NRRL3 | ≥80% at pH 3 to 9 (48 h at 20°C) | >80% at 40°C (96 h) | 14.5 U ml⁻¹ (After optimization-138.3 U ml⁻¹) | Taddia et al. 2020 |

SmF: Submerged Fermentation, SSF: Solid state fermentation
Table 3
Effect of selective metal ions and reagents on xylanase activity

| Metal ions | Concentration (mM) | AUMS60 Relative activity (%) | AUMS64 Relative activity (%) |
|------------|--------------------|-----------------------------|-----------------------------|
| Control    | 0                  | 100                         | 100                         |
| NaCl       | 1                  | 200.24                      | 178.78                      |
|            | 10                 | 185.20                      | 182.14                      |
| KCl        | 1                  | 196.34                      | 179.71                      |
|            | 10                 | 198.36                      | 179.33                      |
| MgSO₄      | 1                  | 125.69                      | 124.83                      |
|            | 10                 | 124.31                      | 89.20                       |
| CuSO₄      | 1                  | 87.39                       | 110.85                      |
|            | 10                 | 1.82                        | 1.52                        |
| EDTA       | 1                  | 73.63                       | 71.58                       |
|            | 10                 | 79.89                       | 56.82                       |
| CaCl₂      | 1                  | 115.35                      | 125.80                      |
|            | 10                 | 65.34                       | 57.82                       |
| FeSO₄      | 1                  | 89.50                       | 82.43                       |
|            | 10                 | 124.83                      | 116.51                      |
| SDS*       | 1                  | 74.53                       | 22.61                       |
|            | 10                 | 74.37                       | 38.71                       |

*Incubation for 15 min

Figures
Figure 2

Phylogenetic tree derived from BI analysis based on the combined ITS and LSU rDNA data set, depicting the relationships of strains AUMS56, AUMS60, AUMS64 and AUKEMS24 with closely related taxa. Significant Bayesian posterior probability (>50%) are shown on the respective branches. Bar, 0.007 changes per position.

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- SupplementaryInformationRevised.pdf