Amrit Pal Kaur,1 Boguslaw P. Nocek,2 Xiaohui Xu,1 Michael J. Lowden,3 Juan Francisco Leyva,3 Peter J. Stogios,1 Hong Cui,1 Rosa Di Leo,1 Justin Powellski,3,4 Adrian Tsang3,5 and Alexei Savchenko1∗
1Department of Chemical Engineering and Applied Chemistry, University of Toronto, ON M5S 3E5, Canada.
2Structural Biology Center, Argonne National Laboratory, Argonne, IL 60439, USA.
3Centre for Structural and Functional Genomics, Departments of 4Chemistry and Biochemistry and 5Biology, Concordia University, Montreal, QC H4B 1R6, Canada.

Summary
The genome of the thermophilic fungus Scytalidium thermophilum (strain CBS 625.91) harbours a wide range of genes involved in carbohydrate degradation, including three genes, abf62A, abf62B and abf62C, predicted to encode glycoside hydrolase family 62 (GH62) enzymes. Transcriptome analysis showed that only abf62A and abf62C are actively expressed during growth on diverse substrates including straws from barley, alfalfa, triticale and canola. The abf62A and abf62C genes were expressed in Escherichia coli and the resulting recombinant proteins were characterized. Calcium-free crystal structures of Abf62C in apo and xylotriose bound forms were determined to 1.23 and 1.48 Å resolution respectively. Site-directed mutagenesis confirmed Asp55, Asp171 and Glu230 as catalytic triad residues, and revealed the critical role of non-catalytic residues Asp194, Trp229 and Tyr338 in positioning the scissile α-L-arabinofuranoside bond at the catalytic site. Further, the +2R substrate-binding site residues Tyr168 and Asn339, as well as the +2NR residue Tyr226, are involved in accommodating long-chain xylan polymers. Overall, our structural and functional analysis highlights characteristic differences between Abf62A and Abf62C, which represent divergent subgroups in the GH62 family.

Introduction
Plant-derived lignocellulosic biomass represents a major renewable energy resource, as well as a source of raw materials for production of bio-based products (Carroll and Somerville, 2009). However, its conversion into biofuels, fibres and other industrially important biomaterials is hampered by its complex structure, which requires appropriate catalysts to extract its constituents for industrial uses. In natural environments, filamentous fungi achieve conversion of lignocellulosic biomass through secretion of a plethora of diverse carbohydrate and lignin-degrading enzymes. Genome sequencing efforts have revealed that each filamentous fungus harbours 100 to 300 glycoside hydrolase (GH) protein-encoding genes that often include multiple members within a family. However, the number of characterized fungal GH family enzymes is relatively small compared with the numbers of sequenced fungal GH family genes. To better understand the bewildering diversity of these enzymes and their roles in degradation of complex substrates, detailed characterization of their molecular function and specificity is needed.

Arabinoxylan is a major component of the hemicellulose fraction of grasses, and is especially abundant in the endosperm wall of dietary grains such as wheat, triticale and oats (Henry, 1985). It is a heteropolysaccharide and consists of a main chain of β-1,4 linked D-xylopyranosyl sugar units with randomly distributed L-arabinose substituents. The arabinose substituents are linked through either α-1,2- or α-1,3- glycosidic bonds to xylose. Some xylose units of xylan may carry additional substituents such as 4-O-methyl glucuronic acid, acetyl
group or arabinose sugar esterified by coumaric or ferulic acids (de O Buanafina, 2009). These modifications in the xylan chain increase its complexity and can make it refractory to degradation.

Natural decomposition of arabinoxylan requires coordinated actions of endo-1,4-β-xylanases (EC 3.2.1.8), α-L-arabinofuranosidase (EC 3.2.1.55), α-glucuronidase (EC 3.2.1.139), acetyl (xylan) esterase (EC 3.1.1.72), ferulic acid esterase (EC 3.1.1.73) and β-xylosidase (EC 3.2.1.37) (de Vries et al., 2000; Sørensen et al., 2003). Combinations of such enzymes have been used to design ‘minimal’ enzyme cocktails for efficient arabinoxylan hydrolysis (Sørensen et al., 2007) for industrial applications. Among these, the α-L-arabinofuranosidases are the exo-acting enzymes that specifically remove L-arabinofuranose decorations from xylan or arabinan constituents of hemicellulose and are distributed in CAZY families GH3, GH43, GH51, GH54 and GH62 (www.CAZY.org).

Arabinoxylan arabinoxylanohydrolases belonging to the GH62 family specifically remove α-1,2- or α-1,3-linked L-arabinofuranose decorations from xylan (Kellett et al., 1990). Characterized fungal GH62 arabinoxylan arabinoxylanohydrolases that act on arabinoxylan include those from Aspergillus niger (Gielkens et al., 1997), Aspergillus tubingensis (Gielkens et al., 1997), Penicillium chrysogenum (Sakamoto et al., 2011), Coprinopsis cinerea (Hashimoto et al., 2011) and Penicillium funiculosum (De La Mare et al., 2013). However, other fungal GH62 hydrolases have also been reported to be active against branched arabinan and/or arabinogalactan, including enzymes from Aspergillus sojae (Kimura et al., 2000), Ustilago maydis and Podospora anserina (Siguier et al., 2014).

Based on the sequences listed at CAZY, the GH62 family is proposed to consist of two distinct subfamilies (Hashimoto et al., 2011; Siguier et al., 2014). Many sequenced fungal genomes, such as those of P. funiculosum (De La Mare et al., 2013) and Coprinopsis cinerea (Hashimoto et al., 2011), have been reported to carry at least two or more GH62 hydrolases which may either belong to the same or different subfamilies. Recently, we sequenced the genome of Scytalidium thermophilum (http://fungalgenomics.ca/), a thermophilic ascomycete with optimum growth temperatures nearing 50°C. This fungus is the dominant organism of mushroom compost (Wiegant, 1992; Straatsma et al., 1994) and is a source of thermostable enzymes (Guimarães et al., 2001; Zanoelo et al., 2004) with possible commercial applications. In this work, we have characterized the GH62 hydrolases from this fungus in terms of their induction patterns on biomass substrates, structure, biochemical properties and structure–function relationships.

Results

Sequence analysis and expression of S. thermophilum GH62 hydrolases

The genome sequence of the thermophilic fungus, S. thermophilum strain CBS 625.91 contains three genes – abf62A, abf62B and abf62C – predicted to encode secreted GH62 family arabinofuranosidases (Genbank accession numbers KJ545572, KJ545573 and KJ545574). Abf62A and Abf62B share 60% sequence identity between themselves, but only 34% and 36% sequence identity, respectively, with Abf62C (Table S1). A cladogram was constructed using GH62 sequences (Fig. 1) from various fungal genomes including S. thermophilum and was rooted at an out-group branch consisting of five distinct functionally and structurally characterized GH43 sequences: Arb43a from Cellvibrio japonicas [Protein Data Bank (PDB) i.d. 1GYD, (Nurizzo et al., 2002)], AbnB from Geobacillus stearothermophilus (2EXH, Brix et al., 2006), BaSAXHm2,3, Bacillus subtilis subsp. subtilis (3C7E, Vandermaarie et al., 2009), SaAraf43A from Streptomyces avermitilis (3AKF, Fujimoto et al., 2010) and AXHd3 from Humicola insolens (3ZXJ, McKee et al., 2012). Notably, despite significant variation in substrate specificities, the five representatives of GH43 enzyme cluster as one out-group, whereas the fungal GH62 enzymes are distributed into two phylogenetically distinct subfamilies i.e. GH62_1 and GH62_2 (Hashimoto et al., 2011; Siguier et al., 2014). For the GH62 enzymes of S. thermophilum sequences of Abf62A/Abf62B belongs to subfamily GH62_2, while Abf62C is a member of the GH62_1 subfamily.

It is not uncommon for fungal genomes to harbour more than one GH62 gene, and some such as C. cinerea (Stajich et al., 2010) and Myceliophthora thermophila (Berka et al., 2011) may feature multiple representatives of the same subfamily (Fig. 1). The existence of two different subfamilies and multiple members of the same subfamily suggests the possibility of functional diversity among various GH62 homologues that favours their coexistence in the course of evolution.

The sequences of all three S. thermophilum GH62 enzymes feature an N-terminal signal motif typical of extracellular fungal proteins. Abf62A is the only one of the three enzymes that includes a motif, at the C-terminal, similar to carbohydrate-binding module 1 (CBM-1) in addition to the core catalytic domain. The cellulose-binding properties of CBM-1-containing GH62 enzymes from C. cinerea and P. funiculosum have previously been reported (Hashimoto et al., 2011; De La Mare et al., 2013).

To probe the roles of abf62A, abf62B and abf62C in the degradation of different biomass substrates,
S. thermophilum cultures were grown in media supplemented with various polysaccharides, lignin, straws or wood pulps as carbon source (Berka et al., 2011). The expression of individual GH62 members was quantified by transcriptome analysis using ribonucleic acid sequencing (Fig. 2A). Robust expression of abf62A and abf62C was observed in S. thermophilum cultures grown on complex substrates such as straws from alfalfa, canola, barley and triticale, while only basal or no expression was detected for abf62B. The expression of abf62A was generally higher than that of abf62C, reaching up to fivefold higher level in culture on barley straw. Since expression of abf62B was minimal during growth on any of the selected substrates, we focused on functional and structural characterization of Abf62A and Abf62C.

Catalytic properties of Abf62A and Abf62C

For structural and functional characterization, Abf62A and Abf62C were produced in recombinant form in Escherichia coli and purified to homogeneity. DNA sequences encoding N-terminal signal peptides, corresponding to residues 1–30 of Abf62A and 1–18 of Abf62C were omitted during cloning, and proteins were produced with N-terminal polyhistidine tags. In addition, an Abf62A fragment designated Abf62A△CBM that corresponds to

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Fig. 2. Transcription and biochemical analyses of three GH62 enzymes of *S. thermophilum*.
A. RNA-Seq reads of gene transcripts of *abf62A*, *abf62C* and *abf62B* in *S. thermophilum* after growth on various complex substrates, as described in ‘Experimental procedures’.
B. Determination of optimum reaction pH and temperature for Abf62C, Abf62A and Abf62AΔCBM using wheat arabinoxylan as substrate with reaction conditions as described in supplemental Appendix S1 Experimental procedures.
C. Effect of divalent cation chelation (EDTA and EGTA, 2 mM each) and supplementation (Ca<sup>2+</sup>, Co<sup>2+</sup>, Mg<sup>2+</sup>, Mn<sup>2+</sup>, Ni<sup>2+</sup>, Cu<sup>2+</sup> and Zn<sup>2+</sup>; 2 mM chloride salt of each) on enzymatic activities of Abf62C, Abf62A and Abf62AΔCBM using wheat arabinoxylan as substrate.
the enzyme’s core catalytic domain (residues 30 to 322) was also produced.

Recombinant proteins were purified to homogeneity and tested for activity using pNP-α-L-arabinofuranoside. Since Abf62A, Abf62AΔCBM and Abf62C were all active against this generic substrate, activity was next tested against wheat arabinoxylan (a linear xylan backbone with α-1,2 and α-1,3 linked L-arabinofuranose), as well as branched sugar beet arabinan (which has terminal α-1,2 and α-1,3 L-arabinofuranose linked main chain) (Table 1). Abf62A and Abf62C are both active against these substrates but have no detectable activity against CM-linear 1,5-α-L-arabinofuranose, which contains exclusively non-terminal 1,5-α-L-arabinofuranose linkages, indicating that these enzymes lack endo-arabinanase activity. Since the enzymes are active against pNP-α-L-arabinofuranoside and branched arabinan as well as arabinoxylan, we refer to them as arabinofuranosidasises rather than arabinosylan arabino furanohydrolases (Kormelink et al., 1991).

Abf62A and Abf62C showed differences in relative activities on different substrates, as well as in kinetic parameters (Table 1, Fig. S1A). The \( k_{cat} \) and \( K_m \) determined for Abf62A on wheat arabinoxylan are both threefold higher than those determined for Abf62C. The specific activity on pNP-α-L-arabinofuranoside is about 10-fold higher for Abf62A versus Abf62C, and for sugar beet arabinan the specific activity of Abf62A is twice that of Abf62C. The activities of the Abf62AΔCBM fragment are consistent higher than those of the full-length enzyme. The temperature and pH optima for Abf62A and Abf62C were determined using reducing sugar assays with wheat arabinoxylan as substrates (Fig. 2B). The enzymes are optimally active at 50 °C and at pH ranges of 5.0–6.5 (Abf62A) and pH 5.5–7.0 (Abf62C). 

**Table 1.** Kinetic parameters of Abf62C and Abf62A wild type and variants on arabinan substrates.

| Protein          | High-viscosity wheat arabinoxylan | pNP-α-L-arabinofuranoside | Sugar beet arabinan |
|------------------|-----------------------------------|---------------------------|--------------------|
|                  | \( K_m \) \( mg \text{ ml}^{-1} \) | \( k_{cat} \) \( min^{-1} \) | Specific activity \( U \text{ mg}^{-1} \) | Specific activity \( U \text{ mg}^{-1} \) |
| Abf62C           | 3.66 ± 0.82                      | 16.55 ± 1.13              | 0.02               | 1.5 ± 0.38          |
| Abf62C_Y107A     | 7.56 ± 0.69                      | 14.88 ± 0.62              | 0.01               | N.A.               |
| Abf62C_Y168A     | 22.53 ± 1.59                     | 38.49 ± 2.27              | 0.01               | N.A.               |
| Abf62C_Y168T     | 6.02 ± 0.24                      | 14.81 ± 2.44              | N.D.               | N.A.               |
| Abf62C_Y226A     | 0.54 ± 0.29                      | 3.67 ± 0.23               | N.D.               | N.A.               |
| Abf62C_D194A     | N.D.                             | N.D.                      | 0.38               | N.A.               |
| Abf62C_W230A     | N.D.                             | N.D.                      | 0.35               | N.A.               |
| Abf62A           | 9.38 ± 1.98                      | 51.44 ± 5.02              | 0.24               | 2.8 ± 0.65         |
| Abf62A-ΔCBM      | 8.40 ± 2.02                      | 67.09 ± 7.14              | 0.86               | 7.3 ± 0.28         |

N.A., not analysed; N.D., no activity detected.

Structural characterization of Abf62C

The crystal structures of Abf62C in apo form and in complex with xylotriose were determined to 1.23 Å and 1.48 Å resolutions respectively. The Abf62C apo structure was determined using a selenomethionine substituted protein crystal by the single wavelength anomalous diffraction (SAD) method (Hendrickson, 1991), and it was then used as a search model to determine the structure of the enzyme’s core catalytic domain (residues 30 to 322) was also produced.
A comparative sequence analysis was undertaken (Fig. S2) to detect sequence conservation among various members of the two GH62 subfamilies, including structurally characterized representatives from the basidiomycete *Ustilago maydis* (UmAbf62A) and the ascomycete *Podospora anserina* (Siguier et al., 2014) points to three completely conserved residues Asp55, Asp171 and Glu230 as the catalytic triad of Abf62C. The disposition of these catalytic residues in the Abf62C structure (Fig. 3A and B) is similar to that of catalytic triads previously characterized in GH43 (Brüx et al., 2006) and GH62 (Siguier et al., 2014) enzyme family representatives. Interestingly, one of the phosphate molecules present in the Abf62C apo structure is bound in the active site cavity forming a network of hydrogen bonds with side chains of active site residues (Lys54, Arg259, His303 and Gln328) including that of the catalytic Glu230, thus suggesting the probable position of arabinose binding (Fig. S3A and B).

The Abf62C apo and xylotriose-bound structures superimposed with root-mean-square deviation of 0.24 Å over 324 C-alpha atoms (Fig. 3B), indicating minimal change in conformation upon substrate binding. However, detailed comparison of the apo and the xylotriose-bound Abf62C structures revealed the change in positions side chains of several residues (Fig. 4A and B) in response to binding to xylotriose, both towards the molecular surface (Tyr168, Trp229, Arg259 and Tyr338; Fig. 4A) and the catalytic core (Asp55, Asp171, Glu230 and His303; Fig. 4B). The most dramatic shifts were observed in Asp55 (2.7 Å and 2.2 Å of OD2 and OD1 atoms respectively), Tyr338 (1.7 Å shift of –OH group) and Glu230 (1.5 Å shift of atom OE2).

The recently characterized GH62 enzyme structures (Maehara et al., 2014; Siguier et al., 2014) contain a single calcium ion in the active site; a feature shared with previously characterized members of GH43 family as well (de Sanctis et al., 2010; Santos et al., 2014). Surprisingly, despite significant sequence similarity between *PaAbf62C* and *Abf62C* enzymes (64% identity, Table S1) and also the conserved calcium binding residue (H285 in *PaAbf62C* and His303 in *Abf62C*; Fig. 4C), the *Abf62C* structure does not harbour a calcium ion in its active site. This Abf62C feature is in line with the functional data presented above, which shows that Abf62C activity is not significantly affected by the presence of Ca\(^{2+}\) or chelating agents, suggesting structural and functional independence from divalent metal ion binding.

However, a 0.9 Å shift of the ND1 atom of His303 in Abf62C (Fig. 4C and D) in the xylotriose bound structure places this residue in a virtually identical position to the calcium-coordinating orientation observed in the structures of the previously characterized enzyme *PaAbf62C* (Fig. 4C) and *UmAbf62C* (Fig. 4D). The presence of a calcium ion in these two enzymes plays a structural role by restricting the movement of the histidine side chain.

### Table 2. Data and refinement statistics.

| Data collection statistics | Apo-Abf62C | Xylotriose-Abf62C |
|----------------------------|------------|-----------------|
| PDB ID                     | 4PVA       | 4PVI            |
| Space group                | P3\(\_\_\_\) | P3\(\_\_\_\)     |
| Unit cell (Å)              | a = 72.0,  | a = 49.6,       |
|                            | b = 72.0,  | b = 49.6,       |
|                            | c = 61.2,  | c = 232.6,      |
|                            | \(\alpha = \beta = 90^\circ\), | \(\gamma = 120^\circ\) |
| Resolution (Å)             | 27.5–1.23  | 23.7–1.48       |
| Wavelength (Å)             | 0.9794 Å   | 0.9794 Å        |
| Number of observed reflections | 461 208  | 112 129         |
| Number of unique reflections | 97 754 | 58 733          |
| Redundancy\(d\)            | 4.7 (1.8)  | 1.9 (1.6)       |
| \(R_{merge}\) (\%)         | 4.4 (41.8) | 2.5 (34.5)      |
| \(R_{merge}\) (\%)         | 4.9 (56.4) | 3.3 (46.6)      |
| Completeness\(b\) (\%)     | 94.7 (52.2)| 96.5 (96.4)     |
| I/\(\sigma\)               | 40.7 (1.80)| 22.4 (1.90)     |
| Phasing                     |             |                 |
| phasing method             | SAD        | MR              |
| Refinement statistics      |             |                 |
| \(R_{cryst}\) (\%)         | 14.40      | 13.02           |
| \(R_{free}\) (\%)          | 16.21      | 15.88           |
| protein residues           | 320        | 320             |
| xylotriose                 | 0          | 1               |
| solvent                    | 515        | 561             |
| RMSD from target values    |             |                 |
| bond lengths (Å)           | 0.013      | 0.011           |
| bond angles (deg)          | 1.90       | 1.51            |
| Average B factors (Å\(^2\))| 16.39      | 16.09           |
| protein                    | –          | 11.36           |
| Xylotriose                 | –          | 30.85           |
| H\(_2\)O                   | 31.34      | 30.85           |
| Ramachandran (\%)\(e\)    | 96.73/3.0  | 0.3             |

\(a\). \(R_{merge} = \sum_i \Sigma_{hkl} |I(hkli) – \langle I(hkli)\rangle| / \sum_i \Sigma_{hkl} |I(hkli)|\), where \(I(hkli)\) is the \(i\)th observation of reflection \(hkli\), and \(\langle I(hkli)\rangle\) is the weighted average intensity for all observations \(i\) of reflection \(hkli\).

\(R_{merge} = \sum_i \Sigma_{hkl} |I(hkli) – \langle I(hkli)\rangle| / \sum_i \Sigma_{hkl} |I(hkli)|\), and \(R_{merge} = \sum_i (1/(N-1)) \Sigma_{j \neq i} |I(hkli) – I(hkjl)| / \sum_i \Sigma_{hkl} \Sigma_{j \neq i} |I(hkjl)|\).

\(b\). Numbers in parentheses are values for the highest-resolution bin.

\(c\). As defined by MOLPROBITY (M.F. –the most favored/A.A additionally allowed/O. outlier).

\(d\). RMSD, root mean square deviation.

the Abf62C-xylotriose complex by the molecular replacement method (Vagin and Teplyakov, 2000). The statistics for both structures are presented in Table 2.

The Abf62C apo structure contains one polypeptide chain (residues 30 to 350) in the asymmetric unit. In addition, five phosphate ions and one glycerol molecule were modelled (Fig. 3A). The overall fold of Abf62C adopts the five-bladed \(\beta\)-propeller fold similar to the other representatives of the GH43_62_32_68 superfamily (PDB id 2EXH, Brüx et al., 2006; PDB id 4N2R, Siguier, et al., 2014; PDB id 1WMY, Maehara, et al., 2014). Each ‘blade’ in this fold consists of either four (Fig. 3A, ‘blade’ I, II, IV and III) or five (Fig. 3A, ‘blade’ V) \(\beta\)-strands forming antiparallel \(\beta\)-sheets that are interconnected through loops of variable lengths to form a funnel-like structure that encircles the central cavity, which houses the active site.
Apparent ly, substrate binding in Abf62C induces a conformational change of His303 from the preferred region in the apo structure to an energetically disallowed region of the Ramachandran plot for Abf62C-xylotriose (Fig. 4C and D).

**Xylotriose binding sub-site**

The xylotriose molecule occupies in a curved cylindrical sub-cavity (Fig. S4A and B) leading into the active site and is constrained by helix α8, (residue Tyr338, blade I), helix α6 (Tyr226 ‘blade’ IV), the loop connecting helix α5 with strand β11 (Tyr168, connects ‘blades’ III and IV), loop between α-2 and strand β-6 (Tyr107, ‘blade II’). The binding cavity involves many polar residues that are visualized in a long patch of acidic residues in an electrostatic representation (Fig. S3B). To define the structural basis for substrate recognition by Abf62C, the substrate binding cavity was divided into three sub-sites (Fig. 5A and B) using the xylotriose sugar ring nomenclature (McKee et al., 2012). Thus, sub-site +2R is where the reducing end of the xylotriose backbone binds, placing the scissile bond containing the central xylene ring at sub-site +1 (the active site) and the xylene at non-reducing end binds at sub-site +2NR. The type of interactions and H-bonding distances are listed in Table S3. The bound orientation of xylotriose at the Abf62C active site positions the +1 xylene ring relative to the catalytic triad so that its two hydroxyl groups (C2 and C3) are within 3 Å distance of the general base residue, Glu230. These observations support the functional data showing that Abf62C is active against both α-1,2 and α-1,3 L-arabinofuranosyl linkages.

**Probing the Abf62C active site residues by mutagenesis**

To test the individual roles of active site residues in substrate recognition and catalysis, corresponding Abf62C residues were individually substituted by alanine or other residues using site-directed mutagenesis (Table 1 and Table S4). The resulting variants were purified and tested for activity against wheat arabinoxylan and pNP-α-L-arabinofuranoside for comparison to the wild-type enzyme. As expected, replacement of the catalytic triad (D55A, D171A and E230A) and surrounding core residues such as Lys54, Tyr77, Arg259, Tyr338 and His303 by alanine renders Abf62C completely inactive on both substrates (Table S4). Many of these residues form hydrogen bonds with the phosphate ion trapped in the active site of the Abf62C apo structure (Fig. S3A and B) and suggest their equivalent participation in L-arabinose binding to GH62_2 subfamily enzymes (Siguier et al., 2014).

Next, we tested the Abf62C residues involved in interactions with xylotriose at all three sub-sites (Fig. 5A and...
B). At the +2R sub-site, respective substitution of Tyr107 or Tyr168 by alanine do not affect the specific activities in a significant way (Table 1 and Table S4). However, alanine substitution of Asn339 results in loss of activity on both arabinoxylan and pNP-α-L-arabinofuranoside (Table S4) supporting a role for this residue in substrate recognition as suggested by the structure (Fig. 5A and B). At the +1 sub-site, the individual substitution of residues Asp194, Trp229 and Tyr338 completely abrogates Abf62C enzymatic activity against wheat arabinoxylan (Table 1 and Table S4). Conversely, the D194A, W229F and W229A Abf62C variants display an obvious increase in otherwise barely detectable activity on pNP-α-L-arabinofuranoside. The replacement of Asp194 and Trp229 with smaller residues would create a larger opening around the catalytic Glu230 (Fig. 5A), which can better accommodate the p-nitrophenyl ring of pNP-α-L-arabinofuranoside. This also underlines non-catalytic roles of Asp194 and Trp229 in orienting the xylose backbone of the substrate. At the +2NR sub-site, Asp194 is also involved, together with Tyr226 in the orientation of the +2NR of xylotriose via a water molecule (Fig. 5B). This interaction is important as substitution of Tyr226 with alanine leads to complete loss of activity on wheat arabinoxylan (Table 1 and Table S4).

Overall, our mutagenesis data reveal the critical role of several Abf62C residues in the orientation of the substrate molecule in the active site. Our findings also underline the...
and linked arabinose at the end of the residues that are involved in interactions with xylose sequences including Abf62A indicates high conservation of the two enzymes. A sequence alignment of various fungal GH62 subfamily structure depicting substrate interactions between Abf62A and Abf62C in complex with xylotriose represents the first distinguish the functionality between these two enzymes. (Fig. S2) and structural features (Fig. 6A) that might discriminate the conserved histidine (His303 or His248) residues with glutamine residue of Abf62A (Gln207) ligand is replaced by a cysteine residue (Cys233; Fig. 4C and D and Fig. S4) although calcium-binding histidine residues are conserved in the two structures (His248 and His303) respectively. To probe individual roles of these residues in Abf62C and Abf62A catalytic activity, a series of mutations was designed (Table S4), and the variant enzymes were tested for activities on pNP-α-L-arabinofuranoside and wheat arabinoxylan substrates (Table S4). Substitution of aspartate or, to a lesser extent, with glycine. A replacement of the equivalent asparagine residue with glutamine in the case of the Streptomyces coelicolor enzyme (Maehara et al., 2014) is reported to increase its activity on longer chain substrates. Similarly, Abf62C Tyr168 appears to be conserved in members of subfamily GH62_1, whereas the equivalent position in members of the GH62_2 subfamily is occupied by threonine or alanine (Thr139 in Umabf62C and Thr148 in Abf62A, Fig. 6 and Fig. S2). Such a change in residue size may result in a wider cavity opening at the +2R site, thus facilitating binding of a longer or more substituted xylan backbone. Similarly, Abf62C residue Tyr226 at the +2NR sub-site is the least conserved among residues involved in the active site (Fig. 6A and B and Fig. S2). Notably, the bridged interactions between Abf62C Tyr226 and xylotriose (Fig. 5B) are compensated by non-equivalent direct hydrogen bonds between PaAbf62C Arg216 and cellotriose (Fig. 6A). Further, UmAbf62C and Abf62A feature alanine or asparagine residues, respectively, at the position equivalent to Tyr226 in Abf62C (Fig. 6B). These alternative residues of the GH62_2 subfamily have short side chains and therefore may not be able to interact with xylotriose or cellotriose chains due to their shorter side chains, and instead might contribute towards binding an additional xylose ring at +2NR ends.

Another distinguishing feature of Abf62C is that the position equivalent to a potential calcium-binding glutamine residue of Abf62A (Gln207) ligand is replaced by a cysteine residue (Cys233; Fig. 4C and D and Fig. S2) although calcium-binding histidine residues are conserved in the two structures (His248 and His303) respectively. To probe individual roles of these residues in Abf62C and Abf62A catalytic activity, a series of mutations was designed (Table S4), and the variant enzymes were tested for activities on pNP-α-L-arabinofuranoside and wheat arabinoxylan substrates (Table S4). Substitution of the conserved histidine (His303 or His248) residues with alanine in both Abf62C and Abf62A resulted in complete inactivation. The activities of Abf62A Q207A and Q207C variants, and the C233Q variant of Abf62C, were also dramatically abrogated compared with the wild type, highlighting potential functional significance of these residues. However, the loss of activity in the Abf62C C233Q variant
can be also attributed to steric clashes between the introduced glutamine’s side chain and those of Glu306 and Trp174, both of which are conserved in representatives of GH62_1 subfamily (Fig. S2). The equivalent positions in Abf62A are occupied by smaller residues, Asp238 and Thr154, which are conserved in representatives of GH62_2 subfamily.

Combined with the structural analysis, our mutagenesis studies suggest that the primary role of the Abf62C His303 residue is critical for catalysis but it may not involve the coordination of metal ion, unlike equivalent residues in calcium containing GH62 enzymes.

Discussion

Genes encoding a variety of GH43, GH51 and GH62 arabinofuranosidases are present in the genomes of many species of bacteria and fungi. These enzymes play important accessory roles in degrading arabinose-rich arabinan and arabinoxylan. The genome of the thermophilic fungus, S. thermophilum, features genes encoding two GH43, one GH51 and three GH62 enzymes (abf62A, abf62B and abf62C) (http://fungalgenomics.ca/) with potential arabinofuranosidase activities. Focussing on the GH62 family, we found that Abf62A and Abf62C were both expressed when the fungus was grown on a variety of complex substrates rich in arabinoxylan. These two enzymes represent the two subfamilies of GH62 enzymes, a combination of which is often found in fungal genomes containing multiple GH62 representatives.

To understand the significance of the coexpression of two S. thermophilum GH62 enzymes, we characterized their activities and obtained the crystal structure for Abf62C. During preparation of this manuscript, two other fungal GH62 enzymes, PaAbf62C from Podospora anserina and UmAbf62C from Ustilago maydis (Siguier et al., 2014), and one bacterial GH62 from S. coelicolor (Maehara et al., 2014) were also structurally characterized. The GH62 structures from U. maydis (Siguier et al., 2014) and S. coelicolor (PDB id 3WMY, Maehara, et al., 2014) represents the GH62_2 subfamily, whereas the
PaAbf62C enzyme structure from the GH62_1 subfamily was obtained in complex with cellobiose inhibitor. We were able to determine the structure of Abf62C in complex with a true substrate component, xylotriose, thus revealing for the first time the molecular framework involved in GH62 interactions with part of a substrate molecule.

A sequence and structural comparison of Abf62C with other GH62 enzymes highlighted the conservation of residues involved in positioning of the substrate arabinose moiety. Specifically, the residues proximal to the substrate’s scissile bond are invariably conserved between both GH62 subfamilies, with Abf62C showing a unique feature of Trp229 taking the place of an otherwise highly conserved phenylalanine residue. On the other hand, the residues participating in xylene binding at sub-sites +2R and +2NR show more variability that may reflect adaptability among GH62 enzymes to the diverse nature of xylan substrates.

Calcium binding is another important aspect of GH62 enzyme active sites. Several determined structures for GH43 and GH62 enzymes contain calcium ion in their catalytic pockets, anchored by a conserved histidine residue and a network of ordered water molecules. Some of the Ca^{2+} ion containing GH43 arabinanases, including BsaArb43b (de Sanctis et al., 2010) and TpABN from Thermotoga petrophila (Santos et al., 2014) are strongly inhibited by the presence of a chelating agent. In a recently proposed mechanism, the positively charged Ca^{2+} ion in the active site of these GH43 enzymes induces hyper-polarization of an adjacent histidine residue, which in turn affects the functional protonation states of the vicinal catalytic acid residues (Santos et al., 2014). However, the active site of another GH43 family representative, ARN2 features a sodium ion interacting with the corresponding histidine residue (Santos et al., 2014) rather than a calcium ion. In the case of this enzyme, the global change in the protein architecture was proposed to contribute to the retention of the histidine molecular rotameric conformation, thus sustaining its role in catalysis (Santos et al., 2014). Notably, GH43 arabinanases structures lacking Ca^{2+} are not inhibited by chelation (Santos et al., 2014).

According to our data, the Abf62C enzyme is not affected by the presence of chelators, and no metal ion is observed in the enzyme active site despite the presence of the conserved histidine (His303) residue. These findings place Abf62C in the same category as the GH43 arabinases mentioned above (Santos et al., 2014) in which the proper positioning of the substrate, and catalysis, is apparently dependent on conformational flexibility of the active site residues rather than on the presence of a metal ion. Thus, we suggest that in the case of Abf62C, the active site residues, including His303, are evolved to possess the necessary flexibility to achieve a catalytically active state in the absence of a metal ion. However, at high concentrations, the large-radius divalent cations, such as Zn^{2+} and Cu^{2+}, were observed to inhibit Abf62C activity, possibly through low affinity binding to the conserved histidine residue of the active site and altering the catalytic environment. Thus our data indicate that, similar to GH43 family proteins, the GH62 enzymes also demonstrate significant variation with respect to the involvement of the metal ions in the enzyme catalytic center.

The transcription profiles showed that S. thermophilum abf62C and abf62A are upregulated when cultured in plant-derived biomass, as compared with simple sugars. The coexpression of abf62C and abf62A, representing two phylogenetically distinct subtypes, under different growth conditions suggests that the concerted action of Abf62A and Abf62C enzymes provides S. thermophilum an edge in its natural habitat in decomposing plant-derived biomass.

In conclusion, GH62 enzymes play a prominent role in removing arabinose substituents from arabinoxylan, and thus decreasing the complexity of biomass substrates for further downstream processing. With the exponential growth of genomic data revealing a plethora of lignocellulolytic enzyme sequences, the challenge is to understand their synergetic and individual functions in the degradation of complex substrates. Microorganisms from thermophilic environments represent particularly attractive ecological niche of enzymes that can potentially carry out complete degradation of complex substrates in an industrial setting. In this respect, our data show that the GH62 family includes structurally diverse representatives that may offer unique biochemical properties that may be suitable for such applications.

**Experimental procedures**

**Transcriptome analysis**

Scytalidium thermophilum was cultured on different substrates as described (Berka et al., 2011). Total RNA was extracted from mycelia (Semova et al., 2006) at early growth phase, and sequencing was performed using the mRNA-Seq method of Illumina’s Solexa IG at the McGill University-Génomique Québec Innovation Centre. The RNA-Seq reads, 50 nucleotides in length, were mapped and analysed as described (Berka et al., 2011). Fragments per kilobase of transcript per million (mapped reads) values were calculated from the counts using the transcript lengths and the total number of mapped reads from each sample.

DNA manipulation, cloning and expression of abf62C and abf62A in E. coli

Complementary DNA of S. thermophilum was prepared as described (Semova et al., 2006). Deoxyribonucleic acid fragments containing coding sequences for functional domains of
Abf62A and Abf62C were amplified from double-stranded cDNA and cloned (Table S2) into an N-terminal histidine tag containing ligation independent cloning (LIC) based pET15b vector (Novagen). The cloned abf62C (aa30-aa350), abf62A (aa18-aa391), abf62A-ΔCBM (aa18-aa322) were expressed and purified (details in supplemental Appendix S1 Experimental procedures) from BL-21 cells (DE3) Gold strain (Strategene). The oligonucleotide primers used for mutagenesis were designed (Table S2) using the online QuikChange Primer Design tool from Agilent Technologies and the Stratagene XL protocol.

**Activity assays**

The optimal pH conditions for enzymatic activities were determined using 50 mM Britton-Robinson (BR) buffer in pH ranges 2.0–9.0 at 40°C. The same buffer at pH 6.0 was used to determine the optimal temperature using wheat arabinoxylan as substrate. Enzymatic reaction using as substrate pNP-α-L-arabinofuranoside (Sigma N3641) as substrate (1 mM substrate, 1 μg of protein in 50 μl reaction) was carried out at 40°C for 30 min in 50 mM BR buffer, pH 6.0. Reactions were terminated by the addition of 50 μl of 1 M Na2CO3 and p-nitrophenol (pNP) release was determined at 410 nm. One unit of enzyme activity is defined as the amount of enzyme that releases 1 μmol of pNP per min from pNP-α-L-arabinofuranoside under these conditions. Specific activities of Abf62A, Abf62A-ΔCBM and Abf62C were determined by measuring the release of reducing sugars using the Nelson-Somogyi method (Green et al., 1989) adapted to 96 well polymerase chain reaction plates using wheat arabinoxylan (high viscosity, P-WAXYH), sugar beet arabinan (P-ARAB) and CM-Linear 1,5-α-L-arabinan (Megazyme, P-CMLA).

Reaction conditions used to determine kinetic parameters are indicated in Fig. S1A, and the calculations were carried using Prism 5.0 (GraphPad Software, USA). One unit (U) of enzyme activity is defined as the amount of enzyme required to produce 1 μmol of product/min at 50°C at optimum pH. The effect of divalent cation (CaCl2, NiCl2, ZnCl2, MgCl2, CuCl2) and chelator (0.2 M EDTA and 0.2 M EGTA) on enzymatic activities of the GH62 enzymes were assessed in 100 mM HEPES (N-2-hydroxyethylpipерazине-2-ethane sulfonic acid) buffer (pH 7.0) at 50°C for 30 min using 0.2% wheat arabinoxylan.

**1H-NMR assay and product analysis**

1H-NMR experiments were carried out according to the methods described by (Sakamoto et al., 2011). The details of the 1H-NMR and HPAEC-PAD detection of released arabinose are discussed in supplemental Appendix S1 Experimental procedures.

**Sequence alignment and phylogenetic analysis**

ClustalW2 (Goujon et al., 2010) was used to carry out the multiple protein sequence alignment as well as to calculate the phylogenetic tree. ESPRiPT (Gouet et al., 1999) and FIGTREE (http://tree.bio.ed.ac.uk/software/figtree/) were used to visualize the sequence alignment and calculated tree respectively. The sequences used in alignments were obtained from National Center for Biotechnology Information and Joint Genome Institute genomic data websites.

**Crystallization**

Selenomethionine crystals of Abf62C were obtained by hanging drop method at 22°C, using 1 μl of 17 mg ml⁻¹ of Abf62C within the reservoir buffer solution (0.1 M Tris pH 8.5, 2.2 M KH2PO4, 12% glycerol). Abf62C co-crystals with xylotriose were obtained by pre-incubating native Abf62C (17 mg ml⁻¹) with xylotriose (2 mM at 4°C, overnight) by hanging drop method at 22°C in reservoir buffer (0.2 M KH2PO4, 20% PEG3350, 20 mM xylotriose) solution. Twenty percent Paratone-N was supplied to reservoir solution prior to flash freezing crystals into liquid nitrogen.

**Data collection and structure determination**

Crystallographic data of apo and xylotriose-incubated Abf62C were collected at the 19-ID beamline of the Structural Biology Center at the Advanced Photon Source (Argonne National Laboratory, Argonne, IL, USA) (Rosenbaum et al., 2006). Data were collected at a wavelength of 0.9794 Å, from the single crystals and were processed using HKL3000 (Minor et al., 2006). Data collection statistics are presented in Table 2. The structure of apo-Abf62C was determined using diffraction data obtained from a single (SeMet-labeled) crystal by the SAD method (Hendrickson, 1991). The hexagonal crystal contains one monomer of apo-Abf62C in the asymmetric unit. The SAD phasing, density modification and initial protein model building was accomplished in the HKL-3000 (Minor et al., 2006) software package integrated with SHELXD, SHELXE (Sheldrick, 2010), MOLPHARE (Otwinowski, 1991), DM (Cowtan, 1994), ARP/WARP (Langer et al., 2008), SOLVE (Terwilliger and Berendzen, 1999) and RESOLVE (Terwilliger, 2000). The structure of Abf62C with xylotriose was determined by molecular replacement using the structure of apo-Abf62C as a search model. Molecular replacement searches were performed using the MOLREP program of the CCP4 suite (Vagin and Teplyakov, 2000). Both models were rebuilt using the program COOT (Emsley et al., 2010) and refined with PHENIX (Adams et al., 2010) and REFMAC 5.5 (Murshudov et al., 1997). The translation/libration/screw (TLS) operators were automatically determined using the program PHENIX and added in the final round of the refinement. The final refinement statistics for all structures are presented in Table 2. Prior to deposition of the structure in the PDB, the quality of the structure was verified with the set of validation tools in the program COOT (Emsley et al., 2010), as well as PROCHECK (Laskowski et al., 1993) and MOLPROBITY (Lovell et al., 2003). Crystal packing analysis using PISA (Krissinel and Henrick, 2007) showed limited contacts between symmetry-related molecules, strongly suggesting that Abf62C monomer (the asymmetric unit content) represents a biologically relevant unit. Electrostatic potential surfaces were calculated using the APBS PrVOL plugin (Petrey and Honig, 2003).
PDB accession codes

The atomic coordinates of apo-Abf62C and Abf62C-xylotriose complex have been deposited in the Research Co-laboratory for Structural Bioinformatics Protein Data Bank under accession codes 4PVA and 4PVI respectively.

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Conflict of Interest

None declared.

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Supporting information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site:

Fig. S1. Biochemistry of GH62 enzymes.
A. Kinetics parameters of three GH62 enzymes of *S. thermophilum* on wheat arabinoxylan. Varying concentrations of wheat arabinoxylan (P-WAXYH) were used to determine the kinetics of Abf62C (0.5 μg of protein, 100 mM HEPES pH 7.0), Abf62A (0.5 μg of protein, 100 mM citrate buffer pH 5.0) and Abf62AΔCBM (0.5 μg of protein, 100 mM citrate buffer pH 5.0) at 50°C for 30 min.
B. 1H-NMR. 1H-NMR spectra of untreated (A–C) and pretreated with AFase (D–F) wheat arabinoxylan (P-WAXYL). Peaks are labelled based on assignments by Sakamoto *et al.* 2011 as follows: (1, 4) an arabinose residue bound to C-3 of a single-substituted xylose residue (5.357 ppm), (2) an arabinose residue bound to C-3 of a double-substituted xylose residue (5.240 ppm), (3) an arabinose residue bound to C-2 of a double-substituted xylose residue (5.188 ppm) and (5) an arabinose residue bound to C-2 of a single-substituted xylose residue (5.250 ppm). (A) substrate only, (B) Abf62C, (C) Abf62A, (D) pretreated substrate only, (E) Abf62C and (F) Abf62A. All spectra were recorded on a Varian VNMRS-500 MHz spectrometer at 30°C.
C. HPAEC-PAD. Product analysis of Abf62C (3.4 μg ml⁻¹) activity on wheat arabinoxylan (P-WAXYL; 0.2%) in Britton-Robinson buffer pH 5 (30 mM) at 40°C for 30 min. (A) A mixture of 0.016 mM arabinose and 0.016 mM xylose as standards. (B) Only arabinose is detected in the enzymatic reaction of Abf62C and arabinoxylan. Monosaccharides were detected using a Dionex ICS-500 HPLC equipped with a Carboxp PA20 analytical column (3 mm × 150 mm).

Fig. S2. Arabinose binding in Abf62C.
A. H-bonding network formed by the central phosphate in the active site of the apo Abf62C structure.
B. Arabinose binding interactions of *Um*Abf62C (magenta) and their equivalent residues in Abf62C (orange).

Fig. S3. Molecular surface of Abf62C.
A. Molecular surface of xylotriose bound Abf62C (grey). The active site residues lining the binding pocket are shown in sticks (orange).
B. Electrostatic surface of Abf62C displaying the xylotriose bound by a highly positively charged (red) surface extending from the catalytic core. Red indicates negative potential, white is neutral, blue shows positive potential and surfaces were contoured between -20 and +20 kB T/e, where kB is the Boltzmann constant, T is temperature and e is the electronic charge.

Fig. S4. Protein sequence alignment between the GH62 enzymes of selected fungi. The secondary structure of Abf62C (Subfamily 1) and *Um*Abf62C residues (Subfamily 2) are presented on the top and bottom of the alignment respectively. The two GH62 subfamilies and the key residues involved in active centre of Abf62C are marked. The alignment figure was prepared by Espript (http://esprit.ibcp.fr/ESPrity/ESPrity).

Table S1. Sequence and structural homologies between GH62 enzymes.

Table S2. Primers sequences used to amplify and mutate target DNA.

Table S3. H-bonds and stacking interactions between Abf62C protein residues with sugars of xylotriose. At sub-site +2R, Abf62C residue Tyr107 forms stacking interactions with the plane of the +2R xylose of xylotriose, while side chains of Asn338 form three hydrogen bonds with two hydroxyl groups of the sugar. The arrangement/location of the xylose ring at the +1 sub-site is imperative for the catalytic Glu230 of Abf62C to access the scissile bond. Furthermore, the +1 xylose ring is oriented at the sub-site by stacking with the aromatic side chain of Tyr339 and by forming multiple hydrogen bonds with active site residues, including side chains of the catalytic Glu230 (one to 2-OH and two to 3-OH), Trp229 (2-OH), Arg259 (2-OH) and Asp194 (3-OH). The hydroxyl groups (2-OH and 3-OH) of the +2NR xylose ring form hydrogen bonds with two water molecules, which are in turn oriented by interactions with Tyr226 and Asp194.

Table S4. Summary of site directed mutants of Abf62C and Abf62A.

Appendix S1. Experimental procedures.