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## 1 **Review Article:**

## 2 Xylan Degrading Enzymes from Fungal Sources

3 Authors:

## 4 N. Kirikyali and I.F Connerton\*

5 Division of Food Sciences, School of Biosciences, University of Nottingham, Sutton
6 Bonington Campus, Loughborough, LE12 5RD, UK.

7 \*Corresponding Author

8 Tel.: +44 115 95 16 161

9 Fax: +44 115 95 16 162

10 E-mail address: ian.connerton@nottingham.ac.uk

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## 12 Abstract

13 Fungi have the ability to degrade xylan as the major component of plant cell wall 14 hemicellulose. Fungi have evolved batteries of xylanolytic enzymes that concertedly act to depolymerise xylan backbones decorated with variable carbohydrate branches. As an 15 16 alternative to acid extraction in industrial processes the combination of endo-1,4-β-xylanase 17 and  $\beta$ -xylosidase can reduce xylan to xylose. However, unlike chemical extraction procedures enzyme systems can selectively hydrolyse  $\alpha$ -L-arabinofuranosyl, 4-O-methyl- $\alpha$ -D-18 19 glucuronopyranosyl, acetyl and phenolic branches, and therefore have the potential to 20 deconstruct hemicellulose whilst retaining desirable structural integrity and functionality. The 21 sources, structures and catalytic activities of fungal xylanolytic enzymes are reviewed and 22 discussed in the context of their biotechnological potential.

23 Key words: Hemicellulose, Xylan, B-xylosidase, Endo-xylanase, Xylose

#### 24 1. General Overview of Biological Processing

25 Biological processing of plant biomass has become a significant concern in the quest for 26 renewable energy. The initial focus of research has been the characterization and 27 quantification of lignocellulosic constituents within plants species. Consequently several 28 extraction methods leading to the hydrolysis of cellulose and hemicellulose fractions from 29 agricultural lignocellulosic biomass have been adopted, which include the use of concentrated 30 acids, alkali, hydrogen peroxide, steam explosion, hot water treatment, CO<sub>2</sub> explosion and 31 organic solvent treatments [1]. However the use of such harsh physical and chemical 32 treatments for hydrolysis can result in problems during post-treatment processes, especially if 33 the solubilised sugar products are be recovered as high value-added products for use in synthetic chemistry or in the food industry. Plant cell wall degrading enzymes of microbial 34 origin have therefore attracted industrial attention since they have the potential to replace 35 36 toxic chemical treatments in current use and create new functional ingredients. The catalytic 37 performance and formulation of functional enzyme cocktails for hydrolyzing plant biomass 38 together with associated methods for the efficient recovery of oligosaccharides has been the 39 subject of research to establish commercial feasibilities.

40 The field of bioconversion has evolved into a broad multidisciplinary research base that 41 focuses on the (i) enhancement of enzyme efficiency and specificity, (ii) genetic 42 improvement of organisms to produce tailored plant cell wall degrading enzymes, (iii) 43 engineering of strains that can efficiently utilize pentose sugars and (iv) ethanol tolerance. 44 The work contained in this review article summarises the current research into the utilization 45 of hemicellulose, in particular reports the molecular characteristics of xylanases and  $\beta$ -46 xylosidases from fungal sources, for industrial applications.

#### 47 **2. Composition of Plant Biomass**

48 Lignocellulosic materials are composed of cellulose, hemicellulose, pectin, proteins and an

aromatic polymer lignin. Cellulose is a generally a linear polymer constructed of D-glucose subunits associated by  $\beta$ -1,4 glycosidic bonds that have a propensity to form microfibrils. Long chains of fibrils of cellulose are interconnected by hydrogen bonding to make it crystalline in nature. Plant cellulose is often embedded within a matrix containing hemicellulose and lignin [2, 3].

Lignin is a heterogeneous complex of polyphenolic compounds synthesized from phenylpropanoid units such as p-coumaryl, coniferyl, syringyl and sinapyl alcohols that are connected by non-hydrolyzable C-C and aryl-ether linkages. In addition lignin is also interconnected to cellulose and hemicellulose to form a physical barrier and provide cell walls with structural integrity in order to resist microbial attack and oxidative stress [4, 5].

Hemicellulose has a heteropolymeric structure, which is comprised of a linear xylan 59 60 backbone with short lateral side chains of different sugars. Xylan is the most abundant 61 constituent of hemicellulose in annual plants and hardwood species, and is comprised of a 62 linear chain of D-xylopyranosyl residues linked by  $\beta$ -1,4 glycosidic bonds [1, 3, 4]. The 63 lateral side chains linked to xylan are formed from mannan, galactan and arabinan polymers 64 or singularly attached D-xylose, D-mannose, D-galactose, D-glucose, L-arabinose, α-L-65 arabinofuranosyl and 4-O-methyl-glucouronic, ferulic, acetic, p-coumaric and galacturonic 66 acids [6]. However, the constituents of side chains display large variations in composition among plant species. For example in softwood xylans are not acetylated and contain  $\alpha$ -1,3-67 68 glycosidic bonds at C-3 positions, whereas hardwood xylans have high levels of acetylation 69 and emerge as 0-acetyl-4-0-methylglucuronoxylan [3].

The ratio of each polymer within lignocellulose materials have been reported in the literature to vary between the sources of lignocellulose. The majority of the carbohydrate fraction is composed of cellulose and hemicellulose is the second most abundant constituent. Hydrolysis of hemicellulose into sugar monomers as well as cellulose would result in the consumption of 74 more than 60 % of plant materials into valuable products and result in reduced agricultural75 waste.

### 76 **3. Breakdown of Lignocellulose**

77 Xylanolytic and cellulolytic complexes have been identified and explored in saprophytic soil inhabiting fungi and bacteria as they require preformed organic compounds as energy sources 78 79 for cellular synthesis. Unlike other eukaryotes, fungi obtain nutritional requirements by secreting polymer-degrading enzymes, from their hyphal tips, to their surroundings and 80 81 absorb enzymatic breakdown products [7]. Moreover, competent fungal species exhibit two 82 types of extracellular catalytic systems, firstly a hydrolytic system whereby polysaccharides are hydrolysed and a second oxidative ligninolytic system opens phenyl rings for the 83 84 degradation of lignin [5,8]. However, not all fungi are able to synthesize ligninolytic enzymes. This specialized activity is reserved largely to basidiomycetous white rot fungi and 85 86 plant pathogenic/saprophytic fungi, including Armillaria mellea, Pleurotus ostreatus, 87 Phanerochaete chrysosporium, Echinodontium taxodii, Aspergillus sp., Fusarium sp., Ceriporiopsis subvermispora and Botrytis cinerea that have the ability to produce such 88 89 enzymes [8-10].

#### 90 **3.1 Lignin Biodegradation**

Lignin biodegradation is an oxidative process that utilises members of the phenol oxidase family of enzymes. These comprise of lignin peroxidase, manganese peroxidase and laccases, that collectively catalyze the oxidation of variable phenolic moieties. Laccases are of general significance because of their non-selective ability to catalyse the oxidation of a variety of substrates including diphenols, polyphenols, diamines, substituted phenols and aromatic amines [8]. The peroxidases of lignocellulosic fungi utilise hydrogen peroxide that is generated by the enzymes glyoxal oxidase and glucose-2-oxidase [11]. These enzymes have 98 been characterised in Basidiomycota group of white rot fungi including Strobilurus 99 ohshimae, Echinodontium toxodii, Ceriporiopsis subvermispora, Phanerochaete 100 chrysosporium, Trametes versicolor and Bjerkandera adusta which oxidise phenolic lignin 101 units to diffusible products that can be acquired through hyphal tips and then utilised by 102 intracellular catabolic pathways [8-11].

Once dissociated from lignin cellulose and hemicellulose are available to a wider range of competitive microbial species [12]. The ability to degrade plant materials efficiently arises from the multiple cellulolytic and xylanolytic isoenzymes, which have diverse biochemical properties that have evolved through genetic selection for substrate availability and competitive efficiency within microbial communities [13].

## 108 **3.2 Cellulose Biodegradation**

109 Cellulose degrading microorganisms play a key role in carbon cycle by reprocessing carbon 110 fixed by photosynthesis. Many filamentous fungi have been characterized to have a complete 111 cellulolytic system, with all the enzymes required to hydrolyse cellulose efficiently via 112 synergistic actions [14]. Cellulases comprise of three classes of hydrolytic enzymes that 113 hydrolyse  $\beta$ -1,4- glycosidic bonds: (i) Endoglucanases, (ii) Exoglucanases and (iii)  $\beta$ -114 glucosidases.

115 Endoglucanases, also known as endo-1,4-β-glucanase or endocellulase, are responsible for random cleavage of internal sites within cellulose fibres. Exoglucanases (synonyms: exo-1,4-116 117 β-D-glucanase, cellobiohydrolases and exocellulase) remove short oligosaccharides from the ends of glucan chains, which then allow  $\beta$ -glucosidase to hydrolyse cellooligosaccharide 118 119 chains and cellobiose into glucose [15,16]. In addition to the enzymes listed above novel 120 types of cellulase enzymes have been recently identified. Swollenin (SWOI) from T. reesei 121 has been described to disrupt cellulosic fibres without showing any hydrolytic activity. Their role is analogous to plant expansins, which are thought to promote accessibility to cellulases 122

123 that depolymerise cellulose fibrils [11,17].

#### 124 **3.3 Hemicellulose Biodegradation**

125 Unlike cellulose, hemicellulose has a more heterogenic structure that comprises of a linear 126 main chain of  $\beta$ -1,4 linked D-xylose backbone and short lateral side chains of different sugar 127 residues [18,19]. Complete depolymerisation of hemicellulose fractions, into xylose and other monosaccharide sugars, requires a compiled cooperative action of a diverse range of 128 129 enzymes. Enzymatic hydrolysis of hemicellulose commences with the removal of side chains 130 that block the sites where xylanases cleave the xylan backbone. Endo-1, 4- $\beta$ -xylanase 131 enzymes do not cleave the xylan backbone randomly but cleave the glycosidic bonds in a 132 selective manner depending on the chain length and degree of branching of substrate 133 molecules and the presence of its constituents [20]. The cleavage of the xylan backbone yields xylo-oligosaccharides and the final trimming is carried out by  $\beta$ -xylosidase that 134 hydrolyses the  $\beta$ -1,4 glycosidic linkages of short chain xylo-oligosaccharides and xylobiose 135 from the non-reducing termini to release xylose monomers [21,22]. Accessory side chains are 136 further removed by  $\alpha$ -glucuronidase, acetylxylan esterase, arabinase,  $\beta$ -mannosidase,  $\alpha$ -L-137 138 arabinofuranosidase and other hemicellulolytic esterases [6], as illustrated in Figure 1

#### 139 **4. Sources of Xylanolytic Complexes**

140 Xylanolytic and cellulolytic complexes have been identified and characterized in saprophytic 141 soil inhabiting organisms and plant pathogens. Hydrolysis of plant tissues has been reported 142 as a unifying characteristic of plant pathogens and rotting organisms since these set of 143 enzymes play a role during softening of the region of penetration and invasion of host cell 144 wall structures for obtaining nutrients [23].

However xylanases and β-xylosidases also originate from plants such as maize, potato tubers,
barley and ripening strawberry, plum and pear fruits in which they are thought to mobilize

147 polymers and remodel plant cell walls [18,23,24]. Glycoside hydrolases from plants tend to 148 be produced during ripening processes that cause fruit tissues to become softer and play a 149 crucial role in cell enlargement, germination and other physiological processes (i.e budding, 150 senescence, post-pollination processes). Sugar starvation in Arabidopsis has been reported to 151 induce the production of glycosyl hydrolases including  $\beta$ -galactosidase,  $\beta$ -xylosidase and  $\beta$ -152 glucosidase for the mobilisation of carbon storage reserves when photosynthesis is inhibited 153 [25]. Fewer glycoside hydrolases have been characterized from plant origins, although [18] 154 have reported the purification of a  $\beta$ -xylosidase from cell wall of maize during senescence and reported that the enzyme worked optimally at 37 °C and pH 4.5. Under optimal 155 156 conditions kinetic constants, K<sub>m</sub> and k<sub>cat</sub>, against the synthetic substrate p-nitrophenylxvloside were 2.5 mM and 6.5 sec<sup>-1</sup> respectively, which is relatively low compared to  $\beta$ -157 158 xylosidases of fungal origin. The purified enzyme provided promising results through 159 synergistic activity with Tricoderma reesei endoxylanase to increase xylose production by 94 160 % during the hydrolysis of corn stover.

161 Yeast species such as Sugiyamaella xylanicola [26], Spathaspora brasiliensis, Spathasopra 162 xylofermentans [27] and Candida materiae [28] have been isolated from rotting wood and 163 demonstrated to elaborate extracellular enzymes that include xylanolytic activities in the presence of xylan as a sole carbon source. The ability of these species to grow and release 164 165 xylose raised the idea of using these species during fermentation of five carbon substrates. Spathasopra species are known for their ability to ferment xylose producing ethanol and have 166 167 the potential to be used during bioethanol production by fermentation of agricultural 168 materials [27].

169 However, the filamentous fungi remain attractive sources of robust industrial enzymes since 170 they operate in extra-cellular environments and are readily recovered from fermentates to 171 ease downstream processing.  $\beta$ -xylosidases produced from yeasts, for example, are generally 172 cell-associated [29], whereas filamentous fungi tend to secrete the majority into the growth 173 medium.

174 Along with saprophytic and phytopathogenic mesophilic fungi, thermophilic fungi have 175 attracted attention from researchers as sources of thermostable enzymes. It is unusual for 176 fungi to breach the upper temperature boundaries of eukaryotes and produce enzymes which are stable at high temperatures to achieve faster reaction rates, however such fungi that can 177 withstand temperatures of 50 °C are of potential biotechnological use [29]. Thermophilic 178 179 fungal species were discovered a century ago from self-heating hay sacks. Exothermic 180 reactions associated with saprophytic/mesophilic microorganisms result in favourable 181 temperatures for thermophilic fungal spores to germinate and compete for resources at 182 temperatures above 40 °C [14]. The genome sequences of several thermophilic filamentous fungi have been determined. Mining these DNA sequences has established the presence of 183 184 redundant xylanolytic complexes that contain enzymes with the potential to operate at elevated temperatures, such as those reported from Talaromyces thermophilus, Myriococcum 185 186 thermophilum, Scytalidium thermophilum and Thermomyces lanuginosus [30-32].

187

## 5. Synergy between hemicellulose hydrolyzing enzymes

Due to the complexity of the hemicellulose structure, hemicellulase enzymes tend to act in 188 189 concert with auxiliary enzymes. Three types of synergic relationships have been observed: (i) 190 homo-synergy between main chain cleaving enzymes (i.e. Endo-1,4-\beta-xylanases); (ii) hetero-191 synergy between main chain cleaving and debranching enzymes (e.g. α-L-192 arabinofuranosidases) and (iii) anti-synergy whereby one enzyme inhibits the activity of 193 another enzyme regardless whether the primary activity is main chain cleavage or 194 debranching [3]. Interactions between selected hemicellulases have been carried out on 195 different substrates in various combinations in order to uncover the best heterosynergic 196 activity for optimum hydrolysis and for the formulation of commercial enzyme cocktails. The 197 composition of these cocktails require careful consideration because xylan degrading 198 enzymes, like other plant cell wall degrading enzymes, exhibit variation in structure and 199 substrate specificities. Xylanolytic activities can operate as domain components of 200 multifunctional enzymes that include carbohydrate binding modules of different molecular 201 sizes with varying abilities to permeate branched substrates to access the main xylan 202 backbone.

203 In several studies, incubation of endo-1,4-β-xylanases with birchwood xylan have been 204 reported to yield short chain xylooligosaccharides, with xylobiose and xylotriose 205 predominating and minor amounts of xylose [30,33-35]. Addition of B-xylosidase to 206 birchwood xylan releases negligible amounts of xylose due to the presence of a limited 207 number of non-reducing ends that are the substrate for the enzyme [36]. The presence of 208 branching hetero substituents may also be hindering exo-cutting β-xylosidase activity (Figure 209 1). However, the combination of endo-1,4- $\beta$ -xylanase and  $\beta$ -xylosidase has been confirmed 210 to enhance the release of reducing sugar from xylans up to 25 fold that of  $\beta$ -xylosidase treatment alone. The catalytic activity of xylanase, could be classified as a heterosynergic 211 relationship since the action of xylanase produces more unsubstituted non-reducing ends for 212 213  $\beta$ -xylosidase to attack, hence an effective increase in substrate concentration [3,36].

A combined treatment of  $\beta$ -xylosidase and arabinofuranosidase almost doubled xylose release from water extractable wheat arabinoxylan compared to  $\beta$ -xylosidase treatment alone. The removal of  $1\rightarrow 3$  linked arabinose from singly substituted xylopyranosyls near nonreducing ends provided access for  $\beta$ -xylosidase [37]. The ability to degrade water soluble wheat arabinoxylan into reducing monomers by a combination of endo-1,4- $\beta$ -xylanase and  $\beta$ xylosidase could be increased 2.5-fold by the addition of  $\alpha$ -L-arabinofuranosidase [37-42].

#### 220 6. Influence of Carbon Source on Xylanase and β-Xylosidase Production

Hydrolysis of plant polysaccharides is essential for fungal organisms to obtain energy and nutrients for growth. As a consequence filamentous fungi have evolved to secrete relatively large quantities of enzymes that degrade plant carbohydrates into fermentable sugars. Distinct substrates, including agricultural by-products such as wheat straw, corn cobs, sugar beet pulp, corn stalks and selected carbohydrates (i.e. birch wood xylan, oat spell xylan and beech wood xylan) are used for the induction of cellulolytic and hemicellulolytic enzymes under solid state cultivation.

228 Studies have been conducted in order to test the response of fungal organisms to different 229 carbon sources within culture media and investigate expression patterns of enzymes. Model 230 organisms like Neurospora crassa have been demonstrated to sense the presence of cellulose 231 in the environment and induce lignocellulolytic enzymes in response to the exposure of 232 particular carbon source available. Expression of cellulases and hemicellulases in response to 233 Avicel (crystalline cellulose) has been reported to induce 17 of 21 predicted cellulase and 11 234 of 19 predicted hemicellulase genes in Neurospora crassa genome on the basis of RNA 235 sequencing [43,44]. Cellulolytic enzymes are synthesized in association with hemilolytic 236 enzymes when fungal organisms are actively grown on cellulose containing media owing to 237 traces quantities of xylan that often exist in conjunction with commercial preparations of 238 cellulose that are sufficient to trigger basal transcription of hemicellulase encoding 239 components [45].

Expression patterns of hemilolytic enzymes in response to a range of carbon sources have been studied by several research groups with similar conclusions. Xylan as the sole carbon source will lead to the expression of endo-xylanase and  $\beta$ -xylosidase synthesis, as demonstrated in Penicillium purpurogenum [46], Trichoderma reesei [47], Scytalidium thermophilum [48,49]. Xylose residues released from xylobiose or xylan has been suggested to induce expression of xylanolytic complexes [20] but in some microorganisms the accumulation of xylose at high concentrations (45 g/L) has been observed to have a repressive effect on both xylanase and  $\beta$ -xylosidase expression, indicating that carbon catabolite repression can be triggered by co-catabolic products [47]. The presence of glucose severely reduces the levels of mycelial  $\beta$ -D-xylosidase achievable in Scytalidium thermophilum and Penicillium sclerotiorum, indicating that  $\beta$ -D-xylosidase synthesis is subject to carbon catabolite repression [48,50].

## 252 **7. Regulation of Hemicellulase Encoding Genes**

253 Filamentous fungi are thought to respond to short chain oligosaccharides that are initially 254 released from polymeric substrates, which include derivatives such as xylitol and arabinol 255 that have been considered to induce expression of xylanolytic enzymes [11,47]. Regulatory elements and binding sites of transcriptional activators have been identified in the promoter 256 257 regions of the genes encoding cellulolytic and xylanolytic enzymes. The transcriptional 258 activator, XlnR, regulates the xylanase-encoding genes of Aspergillus niger [51]. XlnR 259 contains a zinc binuclear cluster DNA binding domain that binds to the consensus sequence 260 5'-GGCTAR-3' with orthologues in other fungal species [52-55]. Transcriptome analysis of 261 wild type A. niger and a XlnR knockout strain revealed 25 genes to be positively regulated by XlnR that included genes of  $\beta$ -xylosidase, endoxylanase, arabinofuranohydrolase, xylose 262 263 reductase, cellulases and sugar transporters [56]. Klaubauf et al. [57] have further examined the secretome profiles of wild type and XlnR mutants ( $\Delta x lnR$ ) of Fusarium graminearum, 264 265 Magnaporthe oryzae, Trichoderma reesei, Aspergillus niger and Aspergillus nidulans in the 266 presence of 25 mM glucose, 25 mM xylose and 1% (w/v) beechwood xylan medium. This 267 study concluded that the regulation of core set of xylanolytic enzymes including GH11 endoxylanase, GH3 and GH43  $\beta$ -xylosidases,  $\alpha$ -glucouronidase were highly dependent on XlnR 268 transcriptional regulator but other side-chain cleaving enzymes such as  $\alpha$ -arabinofuranosidase 269

270 from Aspergillus niger and  $\alpha$ -galactosidase from Trichoderma reesei and Aspergillus 271 nidulans were unaffected by the deletion of XlnR.

272

Several studies have demonstrated that the expression of fungal plant cell wall degrading enzymes is subject to carbon catabolite repression [34,53,58,59]. The mechanism controlling the preferential utilization of substrates over alternative carbon sources is mediated by the wide-domain repressor CreA which alters the transcription of XlnR [34,53,58,60]. CreA binds the promotors of the genes encoding the xylanolytic enzymes and the activator XlnR to create a double lock mechanism in which CreA represses transcription of the activator and the target genes.

## 280 8. Biochemical Characterisation of Xylanolytic Complexes

281 The majority of fungal xylanases are single subunit proteins of varying molecular sizes that 282 generally fall within the range of 20-60 kDa with the exception being reports of dimeric 283 xylanases from Talaromyces emersonii with molecular weights of 131 and 181 kDa [14,61]. 284 Several eukaryotic endoxylanases occur as glycosylated enzymes in which carbohydrate 285 groups are covalently linked to the protein or are present in dissociable complexes [29]. Carbohydrate content of the three xylanases, Xa, XbI and XbII encoded by Talaromyces 286 287 byssochlamydoides have been reported as 36.6, 31.5 and 14.2 % respectively whilst carbohydrate content was not reported for the xylanases encoded by Talaromyces emersonii 288 [61]. It is proposed that the presence of carbohydrate moieties associated with xylanases 289 290 allow the proteins to tolerate higher temperatures. Xylanases of mesophilic origin such as 291 Aspergillus and Penicillium species, exhibit pH optima ranging from 3 to 7 and exhibit 292 optimal activities at temperature ranging from 40 to 60 °C [34]. It is, however, notable that a 293 few fungi of mesophilic origin produce xylanases with increased thermal stability, for 294 example the xylanase of Ceratocystis paradoxa, remains active at 80 °C for 1 hour [29].

Endoxylanases from thermophilic organisms typically exhibit pH optima ranging from 4.5 to
6.5 and temperature optima in the range of 55 to 65 °C.

297 Fungal  $\beta$ -xylosidases are often characterized as monomeric glycoproteins, however, some have been reported to comprise of more than one subunit such as Humicola insolens β-298 299 xylosidase, which is characterized as a heterodimeric protein of 68 and 17 kDa subunits that 300 could have arisen by post-translational cleavage [62]. The  $\beta$ -xylosidases of Neocallimastix 301 frontalis and Aspergillus pulverulentus have also been characterized as dimeric enzymes with 302 molecular masses 180 and 190 kDa respectively [50]. Zanoelo et al. [63] characterized a cell 303 wall bound  $\beta$ -D-xylosidase from Scytalidium thermophilum, which has a carbohydrate content of 12 % and a molecular weight of 45 kDa. Recombinant enzymes purified from 304 305 yeasts or filamentous fungi tend to show variation in molecular mass from the native enzyme 306 sources due to post-translational modifications. The  $\beta$ -xylosidases of Talaromyces emersonii 307 and Trichoderma reesei have been expressed in Aspergillus oryzae, and both recombinant 308 enzymes were estimated 15 kDa larger than their theoretical molecular masses [40]. The 309 methylotrophic yeast Pichia pastoris is commonly used for heterologous protein expression 310 studies tends to hyperglycosylate recombinant enzymes depending on the number of N-311 glycosylation sites within the target protein. Fungal glycosyl hydrolase family 3 β-312 xylosidases from Neurospora crassa and Aspergillus oryzae expressed in Pichia pastoris 313 were observed to be hyperglycosylated with predicted molecular masses of 81.8 and 84.7 314 kDa as compared with SDS-PAGE estimates between 120-180 kDa, 153-165 kDa 315 respectively [64,65]. Similarly a 65.6 kDa Phanerochaete chrysosporium β-xylosidase 316 expressed in P. pastoris to produce a protein 83 kDa mass as a result of post-translational 317 glycosylation.

318 In contrast to endoxylanases,  $\beta$ -xylosidases from mesophilic and thermophilic origin exhibit 319 optimal temperature ranges between 50 and 70 °C and their thermo-stability is highly variable depending on structural differences. The pH optima of fungal β-xylosidases from mesophilic and thermophilic origin differ with the formers ranging from pH 2.5-4 and the latter pH 5–7 [21,30,45,46,63-67].

323  $\beta$ -xylosidases have been reported to exhibit bifunctional activities against the synthetic 4-nitrophenyl-β-D-xylopyranoside 324 substrates (PNPX) and 4-nitrophenyl-α-L-325 arabinofuranoside (PNPA) [64,68]. The majority of  $\beta$ -xylosidases studied to date have been 326 characterised against PNPX but it should be noted that enzyme behaviour and performance in the natural environment or during industrial application may differ. For example, a V<sub>max</sub> of 327 1052  $\mu$ mol min<sup>-1</sup> mg<sup>-1</sup> was recorded for recombinant N. crassa  $\beta$ -xylosidase against PNPX as 328 compared with 10.2 µmol min<sup>-1</sup> mg<sup>-1</sup> against xylobiose with descending values for longer 329 chain xylooligosaccharide substrates [65]. Similarly Aspergillus oryzae KBN616 β-330 xylosidase exhibited greater catalytic efficiency (V<sub>max</sub> and k<sub>cat</sub>) against the synthetic substrate 331 PNPX rather than naturally occurring xylooligosaccharides, with a V<sub>max</sub> of 250 against PNPX 332 and 25.5 µmol min<sup>-1</sup> mg<sup>-1</sup> against xylobiose [64]. The structural configuration of the C-O-C 333 xylosidic bond between 1.4 xylose residues is markedly different to that presented by 334 xylopyranoside 4-nitrophenyl, which is likely to affect the positioning within the active site and 335 336 the rate of hydrolysis.

#### 337 9. Xylanase classification

338 The glycosyl hydrolases have been classified according to their amino acid sequence directed 339 structures and catalytic mechanisms [69-71]. These classifications are available in the 340 Carbohvdrate Active Enzymes database (http://www.cazy.org/Glycoside-(CAZv) 341 Hydrolases.html). Fungal xylanases (endo-1,4-β-xylanase, E.C.3.2.1.8) generally fall into two 342 families of the glycosyl hydrolases (GH) that were initially named F and G, and later 343 renamed as GH10 and 11 in the consolidated scheme [72]. However, enzymes with xylanase 344 activities associated with distinct catalytic domains can also be found in GH families 5, 7, 8 and 43. Exoxylanase activity (exo-1,  $4-\beta$ -xylanase, E.C.3.2.1.37) has also been reported from 345 346 the fungus Chaetomium thermophile [73].

#### 347 9.1 Glycoside Hydrolase Family 10

348 Family GH10 xylanases have a catalytic domain molecular mass of approximately 35 kDa with the possibility of translational fusions to variable accessory domains that can feature 349 alternative catalytic domains or carbohydrate binding modules. 350 The catalytic domain 351 structure is that of an eight-fold  $(\beta/\alpha)$  barrel, commonly referred to as the TIM-barrel fold 352 (Figure 2A). Structural studies of the binding of xylooligosaccharides indicate that GH10 xylanases contain five xylopyranose subsites A-E and that hydrolysis occurs between subsites 353 354 D and E [74]. Catalysis proceeds via a double displacement mechanism that retains the anomeric centre [75], using two glutamate residues located on the carboxy-terminal ends of 355 356 core  $\beta$ -strands 4 and 7 [76]. Comparative studies indicate family GH10 xylanases have higher 357 affinity for shorter linear  $\beta$ -1,4-xylooligosaccharides than family GH11, which has been 358 proposed to be as a consequence of smaller substrate binding sites [77]. This gives the family 359 GH10 enzymes greater flexibility with respect to the degree of polymerisation but restricts 360 the enzymes ability to cleave branched chain substrates. However, GH10 xylanase of the 361 thermophillic fungus Thermoascus aurantiacus has been reported to show 4-fold greater 362 activity against xylotriose in which the non-reducing moiety is linked to an arabinose sidechain, compared to the undecorated form of the oligosaccharide [78]. High resolution X-ray 363 364 crystallographic structures of GH10 xylanases from the theromophiles Thermoascus 365 aurantiacus and Thermomyces lanuginosus have enabled comparisons with enzymes derived 366 from mesophiles to assess the structural determinants that confer thermostability to the 367 enzymes. Thermoascus aurantiacus GH10 xylanases were noted to benefit from 368 improvements in hydrophobic packing, favourable interactions between charged side chains 369 with helix dipoles, the introduction of prolines at the N-termini of helices [79] and the 370 formation of salt bridges [80].

371 Structural and complementary mutational studies have highlighted that the N- and C-terminal 372 residues of GH10 xylanase from Aspergillus niger are disordered and could destabilise nonsubstrate bound monomers [81]. Removal of the five disordered residues located at the N-373 374 terminus of the protein, with or without the presence of C-terminal disordered leucine 375 (residue 302), resulted in a 2 to 4-fold increase in the half-life of the recombinant enzymes at 50°C. These data highlight a role for the protein termini in structural stability that had 376 377 previously been ascribed potential roles in oligomerisation and thermostability of a 378 thermophillic Bacillus ssp. family GH10 xylanase [82].

## 379 9.1 Glycoside Hydrolase Family 11

Family GH11 xylanases vary markedly in their biochemical characteristics in terms of pI, thermostability, pH profiles and catalytic properties [83]. However, they have a common core structure of approximately 20-25 kDa molecular mass, which is composed of two antiparallel  $\beta$ -sheets in the form of a  $\beta$ -jelly-roll. The overall jelly-roll structure is folded over to create an active site cleft. From the inner  $\beta$ -sheet two glutamate residues are orientated into the cleft, which represent the catalytic residues of a double displacement mechanism similar to that used by family GH10 xylanases (Figure 2B). The topology of GH11 xylanases have 387 been described to take the shape of a right hand, with the two  $\beta$ -sheets and  $\alpha$ -helix resembling the palm and fingers and two loop regions forming thumb and cord [84]. Several fungal 388 389 GH11 xylanases have been reported to contain disulphide bridges that could confer stability 390 to extra-cellular enzymes. Molecular dynamic studies indicate that the N-terminal regions of GH11 xvlanases initiate protein unfolding [85], and engineering an N-terminal disulfide bond 391 392 has been reported to enhance enzyme thermostability [86-88]. The xylanases from 393 Aspergillus niger [89], Aspergillus kawachii [90], and Scytalidium acidophilum [91] contain 394 disulfide bonds that connect the cord and to strand  $\beta 12$ , whereas the xylanases from 395 Paecilomyces varioti [92] and Thermomyces lanuginosus [93] disulfide bonds are located 396 between an  $\alpha$ -helix and strand  $\beta$ 11. Two disulfide bridges were observed in the family GH11 397 xylanase of the anaerobic ruminal fungus Neocallimastix patriciarum but the disulfide 398 located in the N-terminal region serves to tether the N-terminal located  $\alpha$ -helix to strand  $\beta$ 14 399 [94].

400 **10.** β-Xylosidase classification

401 The CAZy database places  $\beta$ -xylosidases into GH families 3, 8, 30, 39, 43, 52, 54, 116, 120 402 but  $\beta$ -xylosidases of fungal origin are restricted to families 3, 43 and 54. In addition, GH 403 family 31 refers to  $\alpha$ -xylosidase activities from Aspergillus niger and Aspergillus nidulans 404 that hydrolyse  $\alpha$ -D-xylosidic linkages

Glycoside families 3, 31 and 54 function by a double-displacement mechanism, whereas GH
406 43 family enzymes typically operate by inversion of the anomeric centre where aspartate and
glutamate are claimed to be the catalytic nucleophile base and proton donor respectively.

## 408 10.1 Glycoside Hydrolase Family 3

409 The CAZy database contains 4669 family GH3 protein sequences of which 585 are of 410 eukaryotic origin and exhibit the following catalytic activities:  $\beta$ -glucosidase (EC 3.2.1.21), 411 xylan 1,4-β-xylosidase (EC 3.2.1.37), N-acetylhexosaminidase (EC 3.2.1.52), glucan 1,3-β-412 glucosidase (EC 3.2.1.58), glucan 1,4-β-glucosidase (EC 3.2.1.74), exo-1,3-1,4-glucanase 413 (EC 3.2.1.-), α-L-arabinofuranosidase (EC 3.2.1.55), β-glucosylceramidase (EC 3.2.1.45), 414 isoprimeverose producing oligoxyloglucan hydrolase (EC 3.2.1.120), coniferin β-glucosidase 415 (EC 3.2.1.126) and β-N-acetylglucosaminide phosphorylases (EC 2.4.1.-).

Gene sequences encoding  $\beta$ -xylosidases classified within family GH3 have been reported 416 417 from several fungal sources including Aspergillus niger [95], Aspergillus oryzae [67], 418 Aspergillus japonicus [96], Aspergillus nidulans [97], Neurospora crassa [65], Humicola 419 insolens [98], Trichoderma reesei [47] and Talaromyces emersonii [99]. However, structural 420 information is only available for enzyme from Trichoderma reesei, for which small-angle X-421 ray scattering data has been collected and compared with the crystal structure of barley β-D-422 glucan exohydrolase [100]. These data suggest the Trichoderma reesei enzyme consists of 423 three domains as opposed to two characterised for the barley enzyme. The N-terminal domain consists of an  $(\alpha/\beta)_8$  TIM-barrel domain and the second an  $\alpha\beta\alpha$  sandwich with the third 424 425 remaining unclassified. Glycoside hydrolase family 3 enzymes perform catalytic reaction by 426 a double-displacement mechanism, in which two carboxylic acid residues located in the 427 active site are involved in the formation of a covalent glycosyl-enzyme intermediate. The GH 428 3 catalytic nucleophile, aspartate, is conserved across all family members and is located in the 429 N-terminal  $(\alpha/\beta)_8$  TIM barrel domain [101]. The catalytic residues of the family GH3  $\beta$ -430 xylosidases of Trichoderma reesei and Talaromyces emersonii were probed by carbodiimide-431 nucleophile modification, which resulted in complete inactivation of the enzymes suggesting 432 that carboxyl groups are required for catalysis, which is consistent with the candidate aspartate and glutamate residues identified from protein sequence comparisons [40]. 433

## 434 **10.2 Glycoside Hydrolase Family 43**

435 Glycoside hydrolase family 43 contains 312 of eukaryotic origin with the following enzyme

436 activities: β-xylosidase (EC 3.2.1.37), β-1,3-xylosidase (EC 3.2.1.-), α-L-arabinofuranosidase 437 (EC 3.2.1.55), arabinanase (EC 3.2.1.99), xylanase (EC 3.2.1.8) and galactan 1,3-β-438 galactosidase (EC 3.2.1.145), α-1,2-L-arabinofuranosidase (EC 3.2.1.-), exo-α-1,5-L-439 arabinofuranosidase (EC 3.2.1.-).

440 Fungal sources of family GH43 β-xylosidases include Penicillium herquei [102], 441 Cochliobolus carbonum [103], Aspergillus oryzae [104], Thermomyces lanuginosus [105] 442 and Paecilomyces thermophila [22]. Family GH43 β-xylosidases are predicted to have 443 molecular masses between 35-62 kDa that do not contain recognisable secretion signal 444 sequences, and are therefore likely to be cell associated enzymes.

445 To date structures for members have only been determined for β-xylosidases of bacterial origin. However, Ravenal et al [106] have modelled two GH43 enzymes from Penicillium 446 447 purpurogenum, a bifunctional  $\alpha$ -l-arabinofuranosidase/xylobiohydrolase and a  $\beta$ -xylosidase. 448 Like other members of the family the catalytic domains have a five blade  $\beta$ -propeller fold that 449 contains a two subsites and a funnel shaped active site [26,107]. During catalysis, the active 450 site harbours the substrate sugar molecule at the non-reducing end and the remaining sugar 451 backbone tends to be positioned at right angles to the enzyme tertiary structure. GH43 452 enzymes perform catalytic reaction by inversion of the anomeric centre. The catalytic 453 residues are believed to be formed by aspartate as a nucleophile base and glutamate as a 454 proton donor. The catalytic reaction is executed as a single displacement reaction in which 455 one carboxylate (in this case glutamate) protonates the substrate while a second (aspartate) 456 acts as a base to activate a nucleophilic water molecule, which results in a nucleophilic attack, cleavage of the glycosidic bond and inversion at the anomeric carbon. 457

## 458 **10.3 Glycoside Hydrolase Family 54 and Family 31**

459 Glycoside hydrolase family contains 37 enzymes of eukaryotic origin with  $\alpha$ -L-460 arabinofuranosidase (EC 3.2.1.55) and  $\beta$ -xylosidase (EC 3.2.1.37) activities. The majority of 461 the proteins classified within GH family 54 have  $\alpha$ -L-arabinofuranosidase activities but two Aspergillus awamori and Hypocrea koningii are 462 bifunctional with α-Lfrom 463 arabinofuranosidase/β-xylosidases activities against arabinose, xylobiose and arabinose 464 linked xylobiose [101]. There are no protein structures available for  $\beta$ -xylosidases however an  $\alpha$ -L-arabinofuranosidase (EC 3.2.1.55) structure has been determined from Aspergillus 465 466 kawachii [108]. The catalytic mechanism of GH Family 54 is considered to retain the anomeric carbon. 467

There are over 600 GH family 31 members of eukarvotic origin with  $\alpha$ -glucosidase (EC 468 3.2.1.20), a-1,3-glucosidase (EC 3.2.1.84), sucrase-isomaltase (EC 3.2.1.48), a-xylosidase 469 470 (EC 3.2.1.177),  $\alpha$ -glucan lyase (EC 4.2.2.13), isomaltsyltransferase (EC 2.4.1.-),  $\alpha$ -471 mannosidase (EC 3.2.1.24) and oligosaccharide  $\alpha$ -1,4-glucosyltransferase (EC 2.4.1.161) 472 activities. To date only one  $\alpha$ -xylosidase has been cloned and characterized from Asperillus 473 niger which is composed of 736 amino acids and exhibit activity against p-nitrophenyl- $\alpha$ -D-474 xyloside but not against p-nitrophenyl- $\beta$ -D-xyloside [108]. Based on  $\alpha$ -xylosidase structures 475 of bacterial origin, GH31 catalytic domains are  $(\beta/\alpha)_8$  barrel structures that operate by 476 retaining a mechanism with aspartate as the catalytic nucleophile and proton donor.

## 477 **11. Industrial Applications of Endo-xylanases and β-Xylosidases**

478 Despite the general cooperative action of xylanolytic enzymes, it is the endo-xylanases and  $\beta$ -479 xylosidases that are most often used to solubilize xylan in industrial processes. Industrial 480 applications include bread making, xylitol production, fruit juice clarification, processing of 481 pre-digested animal feed, bioethanol production and paper pulp processing [1,109].

482 Robust extraction methods are employed in the processing of wood to pulp and paper in order 483 to separate cellulose fibrils from lignocellulosic materials. These processes include the use of 484 concentrated acids, alkalis, hydrogen peroxide,  $CO_2$  explosion and organic solvent treatments 485 [1,20]. Finished white pulp is bleached with acid or toxic chlorine reagents at high 486 temperature and pressure, the by-products of which represent a threat to local ecosystems. To comply with environmental regulations the harsh chemical methods have been replaced with 487 488 alternative chlorine free bleaching such as hydrogen peroxide, ozone, sodium hypochlorite 489 and biobleaching [110]. Enzymatic treatments using endo-xylanase rich cocktails have been 490 used to enhance delignification. Hemicellulases are used in enzyme assisted bleaching to 491 soften and swell the fibre structure to enhance the efficiency of bleaching chemicals used in 492 later treatment stages [29,109]. However, it is essential for the enzyme preparation to be 493 completely free of cellulase activity, the presence of which would impair pulp quality. Most 494 fungal species co-express cellulolytic and xylanolytic to utilise plant cell wall materials. 495 However, a few strains of fungi have been reported to produce cellulase-free thermotolerant 496 xylanases that degrade the xylan of lignocellulosic components, and are suitable for 497 application in pulp bleaching bioprocesses: Aspergillus sp. 2M1 [111], Talaromyces 498 thermophiles stolk AX4 [112] and Thermomyces lanuginosus SSBP (formally known as 499 Humicola lanuginosa) [113].

500 Xylan rich agricultural wastes hydrolysed into its monosaccharide constituents via enzymatic treatments in a form of liquors containing 30 - 40 % xylose and 20 % other sugars (i.e. 501 502 arabinose, galactose, and mannose) are attractive renewable feedstocks for fermentations aimed to produce bio-based chemicals [114]. The most widely produced xylan derived 503 504 product is xylitol, which is a popular ingredient in oral care and chewing gum products. 505 Xylitol is a 5-carbon alcohol, produced by the chemical hydrogenation of xylose, that has 506 been shown to exhibit antimicrobial properties, and specifically to reduce the growth of 507 dental plaque and decrease the incidence of dental caries [115]. Xylitol is also used as an 508 artificial sweetener with less than quarter of the calories of glucose, but critically it does not 509 induce insulin release in humans, which makes it a suitable substitute for individuals with 510 diabetes [115, 116]. Xylose fermenting yeasts such as the Candida species C. maltosa, C.

511 tropicalis, C. guilliermondii and C. parapsilosis have been extensively studied for their 512 capability to ferment xylose into xylitol using xylose reductase. Microbial reduction of xylose enriched broth using Candida maltosa, was shown to yield 213 g  $L^{-1}$  xylitol from 250 g  $L^{-1}$ 513 514 xylose [117]. A combined environmentally low impact process has been reported for xylitol 515 bioconversion using hydrolysates of corncob and wheat bran (1:1 ratio) first generated by 516 solid state fermentation by Aspergillus terreus, and then utilised as feedstock for Candida 517 tropicalis to convert 75.14 % of the xylose to xylitol [118]. Genetic engineering methods 518 have also been used to create integrated xylitol production pathways in Saccharomyces 519 cerevisiae and Candida tropicalis [119,120]. Saccharomyces cerevisiae INVSc1 520 transformants co-expressing  $\beta$ -1,4-xylanase and  $\beta$ -xylosidase from Aspergillus terreus and 521 xylose reductase from Candida tropicalis was used in a semi-aerobic fermentation to produce 522 0.71 g xylitol/g xylan [119].

First generation bioethanol production was based on the fermentation of the easily 523 524 metabolisable sugar glucose, however much research has been carried out on the conversion of hemicellulose hydrolysate into second generation ethanol production [114]. Ethanol is 525 either used directly as a chemical or as an additive to gasoline up to 20 % by volume. The 526 527 blend of ethanol into gasoline is an improved substitute of methyl tertiary butyl ether (MTBE) that may provide cleaner combustion and reduce greenhouse gas emissions [1,5]. 528 529 Efficient production of bioethanol from lignocellulosic materials requires organisms that are 530 capable of the complete hydrolysis of agricultural waste materials and fermenting organisms 531 that convert all types of monosaccharide constituents into ethanol. Metabolic engineering 532 approaches have been employed in order to produce new strains with hyper-production 533 phenotypes for hemicellulase production and generate Saccharomyces cerevisiae and Pichia 534 stipitis strains capable of ethanolic pentose fermentation [120,121].

535 Xylooligosaccharides obtained by enzymatic hydrolysis of hemicellulose have been reported

as useful bioactive ingredients of food and health products. Similar to xylitol, xylooligosaccharides are moderately sweet compounds with no off-taste and no hazardous properties and can be applied to foods, juices and carbonated beverages. A few studies have reported that these short chain sugars variably support the growth of probiotic Lactobacillus and Bifidobacterium species promoting a number of health benefits such as the inhibition of pathogenic bacteria preventing gastro-intestinal infections and improved intestinal transit [122,123].

The use of exogenous hydrolytic enzymes in animal feed has significant potential to improve nutritive values and digestibility of feed. Commercially available enzymes were initially developed as silage additives in order to obtain pre-digested animal feed, however nowadays such enzymes are directly added into ruminant diets. These additives are approved to increase the metabolizable energy and subsequently lead to animal weight gain and improvements in milk production [124, 125].

549 Enzymatic clarification of fruit juices has become a popular process in juice production. The utilization of pectinases along with cellulases, xylanases and  $\beta$ -xylosidases for the 550 551 degradation of all polymeric carbohydrates found in fruit pulp can improve yields. Freshly 552 pressed juice is turbid and viscous, and without processing the carbohydrate pieces tend to 553 settle during storage. The hydrolysis of polysaccharides weakens residual cell walls that 554 results in the release of cell wall materials and a reduction in the water holding capacity, this 555 leads to efficient juice recovery with reduced turbidity and an increase in the levels of 556 reducing sugars [126]. For example, treatment of pineapple juice with Aspergillus niger 557 DFR-5 xylanase has been reported provide a yield of 71.3 % and a clarity value of 64.7 % [127]. Endo-xylanases and  $\beta$ -xylosidases are also used in the brewing industry on crushed 558 559 barley in order to solubilize arabinoxylans that can give beers a muddy appearance with 560 lower oligosaccharides. The hydrolysis of arabinoxylans in wort reduces the beer's turbidity

thus improving its appearance and viscosity [128,129].

562 Endo-xylanases and  $\beta$ -xylosidases are employed in the baking industry due to the abundance of hemicelluloses within cereals. Wheat flour is obtained by the milling process and consists 563 of starch and arabinoxylan. The addition of several carbohydrase enzymes such as amylases, 564 565 endo-xylanases and glucanases in post-milling processes tends to modify the quality of the flour. Endo-xylanases and  $\beta$ -xylosidases act on the arabinoxylan fraction of the dough to 566 567 liberate the water retained in the arabinoxylan that helps even distribution of water and rise in 568 viscosity [130]. The quality of bread obtained from enzyme treated dough was improved 569 regarding dough rise, bread shape, loaf volume and crumb structure [131]. However, it 570 should be noted that exogenous xylanases have to overcome an endogenous wheat xylanase 571 inhibitor protein to maximise their effects in bread making [132].

572

## 573 12. Concluding remarks

574 There are an increasing number of fungal xylanolytic enzymes either characterised or under study. These studies are prompted by the importance of hemicellulose as an abundant 575 576 carbohydrate in nature. Systematic approaches are likely to yield yet further enzymes and 577 connections as to how they interact with plants as symbionts, pathogens and saprophytes. Despite the abundance of hemicellulose, it is an underutilised resource, either as a renewable 578 579 bioenergy source or a source of complex chemicals. Biotechnological applications making 580 use of xylanolytic enzymes are increasing and will continue to do so as the concerted actions 581 of multi-enzyme systems become better understood.

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# 964 Figure Legends

965 Figure 1

966 A. Xylan polymer structure showing the actions of xylanolytic enzymes. The backbone of the 967 substrate is composed of 1,4- $\beta$  -linked xylose residues with branches of  $\alpha$ -arabinofuranose;  $\alpha$ 968 -4-O-methylglucuronic acid; ferulic or p-coumaric acids. B. Hydrolysis of xylobiose and 969 higher xylooligosaccharides by  $\beta$ -xylosidase.

970

971 Figure 2.

972 A. Structure of the family GH 10 xylanase from Thermoascus aurantiacus showing 973 TIM-barrel fold with the catalytic glutamate residues projecting in to the active site 974 cleft [80]. B. Structure of the family GH 11 xylanase from Trichoderma reesei 975 showing  $\beta$ -jellyroll fold with the catalytic glutamate residues projecting in to the 976 active site cleft [133].

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