UNIVERSITY of York

This is a repository copy of *Production and spectroscopic characterization of lytic polysaccharide monooxygenases*.

White Rose Research Online URL for this paper: <u>https://eprints.whiterose.ac.uk/139496/</u>

Version: Accepted Version

Book Section:

Hemsworth, Glyn R. orcid.org/0000-0002-8226-1380, Ciano, Luisa orcid.org/0000-0002-1667-0856, Davies, Gideon J. orcid.org/0000-0002-7343-776X et al. (1 more author) (2018) Production and spectroscopic characterization of lytic polysaccharide monooxygenases. In: Methods in Enzymology. Methods in enzymology . , pp. 63-90.

https://doi.org/10.1016/bs.mie.2018.10.014

Reuse

This article is distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs (CC BY-NC-ND) licence. This licence only allows you to download this work and share it with others as long as you credit the authors, but you can't change the article in any way or use it commercially. More information and the full terms of the licence here: https://creativecommons.org/licenses/

Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.



eprints@whiterose.ac.uk https://eprints.whiterose.ac.uk/

Production and Spectroscopic Characterisation of Lytic Polysaccharide 1

Monooxygenases 2

- Glyn R. Hemsworth^{1,2†}, Luisa Ciano^{3†}, Gideon J. Davies³, Paul H. Walton^{3*} 3
- 4

¹School of Molecular and Cellular Biology, Faculty of Biological Sciences, University of Leeds, Leeds LS2 9JT, United Kingdom

5 6 7 8 9 ²Astbury Centre for Structural Molecular Biology, University of Leeds, Leeds LS2 9JT, United Kingdom ³Department of Chemistry, University of York, York YO10 5DD, United Kingdom

10 *to whom correspondence should be addressed: paul.walton@york.ac.uk

11 ⁺these authors contributed equally to this chapter.

Contents 12

| 13 | 1. Introduction | 2 |
|----|---|-----|
| 14 | 2. Expression and Purification of LPMOs | 3 |
| 15 | 2.1 Recombinant expression and purification from the <i>E. coli</i> periplasm | 4 |
| 16 | 2.2 Recombinant expression and purification from the <i>E. coli</i> cytoplasm | 6 |
| 17 | 2.3 Recombinant expression and purification from yeast | 7 |
| 18 | 2.4 Recombinant expression and purification from fungi | 8 |
| 19 | 3. Electron Paramagnetic Resonance Spectroscopy | 9 |
| 20 | 3.1 EPR sample preparation for LPMOs | .10 |
| 21 | 3.2 Multi-frequency collection | .12 |
| 22 | 3.3 Simulation procedures | .13 |
| 23 | 6. Acknowledgements | .18 |
| 24 | 7. References | .18 |

Abstract 25

26 Lytic Polysaccharide Monooxygenases (LPMOs, also known as PMOs) are a recently 27 discovered family of enzymes that play a key role in the breakdown of polysaccharide 28 substrates. The ability of LPMOs to introduce chain breaks, using an oxidative mechanism, has 29 particularly attracted attention as the world seeks more cost-effective and environmentally 30 friendly ways of producing second generation biofuels for the future. LPMOs are mononuclear 31 copper dependent enzymes and have an unusual active site which includes the N-terminal 32 residue of the protein in the copper coordination sphere. This N-terminal histidine sidechain 33 is also methylated in fungal enzymes, the molecular reason for which is still a debated topic. 34 The production of these enzymes poses several challenges if we are to understand their 35 chemical mechanisms. Here we describe the methods that have been used in the field to 36 produce LPMOs and provide information on the workflows that we use for our Electron 37 Paramagnetic Resonance (EPR) Spectroscopy experiments. EPR has been a particularly 38 powerful tool in the study of these enzymes and our objective with this chapter is to provide 39 some helpful information for researchers for whom this technique might be daunting or theoretically difficult to access. 40

41 Keywords: LPMO, biofuel, bioinorganic chemistry, EPR, protein production

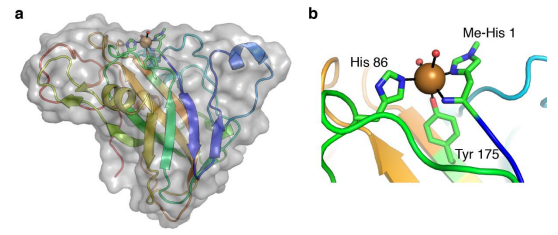
42 1. Introduction

43 The production of second generation biofuels from lignocellulosic feedstocks has been hotly 44 pursued for the last decade as a more sustainable fuel source in comparison to oil and first-45 generation biofuels (several reviews are available, e.g. (Merino and Cherry, 2007; Naik et al., 46 2010; Saini et al., 2015)). One of the major challenges towards the realisation of the cost-47 effective production of these fuels, however, has been the nature of the substrate, cellulose, 48 and its resistance to degradation (Himmel et al., 2007). The study of enzymes involved in the 49 breakdown of this substrate was pioneered by Reese and Mandels through the 1950s 50 (Mandels and Reese, 1957; Reese et al., 1950), 60s (Mandels and Reese, 1960; Mandels et al., 51 1962) and 70s (Mandels et al., 1971, 1974), and we continue to make new discoveries in this 52 area to this day (See recent reviews by Bischof et al. (2016) and Payne et al. (2015)). What is 53 now viewed as the "classical" model of enzymatic cellulose deconstruction requires the 54 combined action of essentially three distinct enzymatic activities to allow the liberation of 55 single glucose molecules from this challenging substrate (Reviewed in Payne et al. (2015)). 56 The glycoside hydrolases involved include endoglucanases, cellobiohydrolases and β -57 glucosidases, which synergise with one another to bring about this molecular conversion for 58 metabolism in nature. These naturally occurring enzymes, however, despite their efficiency 59 in degrading their substrates, have arguably been too slow to make the enzymatic conversion 60 of biomass cost effective in the biorefinery. As such, new enzymatic activities have been 61 sought after (as reviewed by Harris et al. (2014) and Horn et al. (2012)). In this context, Reese, 62 in his original 1950 publication, suggested that there was likely an additional activity required 63 to disrupt the crystalline structure of cellulose to allow glycoside hydrolases access to the 64 substrate (Reese et al., 1950). This activity is now thought to be provided by a family of 65 enzymes known as Lytic Polysaccharide Monooxygenases (LPMOs) (See Beeson et al. (2015), 66 Ciano et al. (2018) and Hemsworth et al. (2015) for recent reviews).

67

68 Prior to the demonstration of LPMO activity in 2010 (Vaaje-Kolstad et al., 2010), LPMOs were 69 defined in two families in the carbohydrate active enzymes database (www.cazy.org) 70 (Lombard et al., 2014) – Glycoside hydrolase-61 (GH61) and carbohydrate binding module-33 71 (CBM33) (See Morgenstern et al. (2014) for a review). In 2010, Harris et al. (2010) had 72 demonstrated that proteins classified as GH61s from Thielavia terrestris and Thermoascus 73 aurantiacus were able to reduce the enzyme load required for the depolymerisation of 74 cellulose 2-fold in a metal dependent manner. The biochemical basis for this became clear 75 when Vaaje-Kolstaad et al. showed that the structurally related protein, CBP21 from Serratia 76 marcescens (classified as a CBM33 in CAZy), was able to oxidatively induce chain breaks into 77 chitin in a reducing agent and O₂ dependent manner, thereby establishing CBM33s as LPMOs 78 for the first time (Vaaje-Kolstad et al., 2010). Subsequent work demonstrated that GH61s 79 were also oxidative enzymes and that they required a copper cofactor to function (Phillips et 80 al., 2011; Quinlan et al., 2011). LPMOs have now been discovered from diverse organisms 81 across the tree of life and have been implicated in the deconstruction of a range of 82 polysaccharides beyond cellulose and chitin (Agger et al., 2014; Bennati-Granier et al., 2015; 83 Borisova et al., 2015; Couturier et al., 2018; Frommhagen et al., 2015; Lo Leggio et al., 2015; 84 Simmons et al., 2017; Vu et al., 2014a). Due to their redox function, LPMOs are now defined 85 in CAZy as auxiliary activities (AA) (Levasseur et al., 2013) with six families currently classified - AA9, AA10, AA11, AA13, AA14 and AA15. 86

88 LPMOs are unusual in being monooxygenases which utilise only a single copper ion in their 89 active site (see Solomon et al. (2014) for a useful review on copper dependent enzymes). 90 Indeed, there have been recent suggestions that LPMOs may, under certain conditions, also 91 function as peroxygenases, but further work is still required to determine the true 92 physiological co-substrate for LPMOs (Bissaro et al., 2017; Kuusk et al., 2018). LPMOs bind 93 their single copper cofactor in a motif that has been dubbed the "histidine brace" (Quinlan et 94 al., 2011), an arrangement of two histidine residues, one of which is the N-terminal residue, 95 which coordinate the copper ion in a T-shaped geometry (Figure 1) (see reviews by Ciano et 96 al. (2018), Hemsworth et al. (2013a), Vaaje-Kolstad et al. (2017) and Vu & Ngo (2018)). In 97 addition, for fungal enzymes at least, there is a modification to the N-terminal histidine 98 sidechain in the form of τ -N-methylation which is only detected when proteins are expressed 99 in fungal hosts (Frandsen et al., 2016; Lo Leggio et al., 2015; Li et al., 2012; Quinlan et al., 100 2011; Vu et al., 2014a, 2014b). The importance of this modification is still debated, but it is 101 clear that LPMOs remain active in its absence when expressed in e.g. Pichia pastoris (Bennati-102 Granier et al., 2015; Bey et al., 2013; Couturier et al., 2018; Wu et al., 2013). The requirement 103 for the N-terminal histidine in the mature protein has dictated the strategies that have been 104 used to produce these enzymes which we will discuss in this chapter. We will also describe 105 our workflows for conducting Electron Paramagnetic Resonance (EPR) spectroscopy on these 106 enzymes. EPR has been a powerful tool in contributing towards unravelling the steps along 107 the reaction coordinate catalysed by LPMOs. EPR is a conceptually challenging technique, 108 however, which requires expertise to both successfully simulate the data and to interpret the 109 resulting spin Hamiltonian parameters in a meaningful way to gain insights into the electronic 110 aspects of the copper active site. We hope that the topics covered in this chapter will, 111 therefore, prove useful not only for those working on LPMOs, but more broadly for 112 researchers interested in studying diverse copper dependent enzymes.



- 113
- Figure 1. Example structure of an AA9 LPMO. a, overall structure of *Thermoascus aurantiacus* AA9 LPMO shown in cartoon coloured from N- (blue) to C-terminus (red). b, Close up view of
- the active site copper ion (orange sphere) coordinated by the histidine brace which includes
- 117 the N-terminal methylated histidine of the protein.
- 118

119 2. Expression and Purification of LPMOs

- 120 LPMOs have been identified from diverse organisms including bacteria (Vaaje-Kolstad et al.,
- 121 2005a, 2010), viruses (Chiu *et al.*, 2015), fungi (Karkehabadi *et al.*, 2008; Langston *et al.*, 2011;

122 Phillips et al., 2011; Quinlan et al., 2011; Saloheimo et al., 1997) and more recently higher 123 eukaryotes (Sabbadin et al., 2018). This can pose challenges towards the choice of expression 124 strategy in order to produce pure proteins to be studied. This choice is compounded by the 125 fact that these proteins are typically secreted in their native hosts, leading to the generation 126 of the N-terminal histidine integral to the active site (for reviews please see Hemsworth et al. 127 (2013a) and Vaaje-Kolstad et al. (2017)). Endogenously, LPMOs will often also contain 128 disulfide bonds, and eukaryotic family members can be further modified by N- and O-129 glycosylation, as well as methylation of the N-terminal histidine. The choice of expression 130 system must therefore be tailored towards the target enzyme and can have a significant 131 impact on the protein yield, and potentially on the activity of the enzyme though this has not 132 been investigated in great detail.

133 2.1 Recombinant expression and purification from the E. coli periplasm

134 Escherichia coli is the workhorse of the academic lab when it comes to protein production. Bacterial LPMOs of the AA10 class are therefore most commonly purified from this organism. 135 136 The most common approach has been to secrete the resultant protein to the periplasm 137 (Courtade et al., 2017; Forsberg et al., 2014a; Hemsworth et al., 2013b; Vaaje-Kolstad et al., 138 2005a, 2010). It is also possible, however, to express these enzymes in the cytoplasm using 139 specialised E. coli strains (Forsberg et al., 2011, 2014b; Gregory et al., 2016). These have 140 largely been our methods of choice for LPMO production and though fungal enzymes are 141 rarely produced using these strategies, we were able to express and purify an AA11 from 142 Aspergillus oryzae using a periplasmic secretion system in E. coli (Hemsworth et al., 2014).

143

144 In order to target the protein to the periplasm, we carefully design our constructs for cloning 145 into a pelB containing vector and typically use HiFi DNA Assembly (New England Biolabs), In-146 Fusion (Takara) or PIPE cloning (Klock and Lesley, 2009). This strategy allows the target gene 147 to be placed directly after the signal sequence without the addition of any residues from use 148 of restriction enzymes and negates the need to use site directed mutagenesis later to remove 149 any undesired residues. Others have found success using the CBP21 leader peptide as the 150 signal sequence (Crouch et al., 2016; Gardner et al., 2014), and one should always consider 151 screening other potential leader peptides in order to improve the chances of obtaining good 152 yields of soluble protein in the periplasm. In addition, careful thought needs to be put towards 153 the use of affinity tags using the periplasmic secretion strategy. An affinity tag would typically 154 be placed at the C-terminus in this case. We have tended to avoid using His-tags for fear that 155 they might interfere with copper binding and produce unwanted copper binding sites during 156 our EPR analyses and also, potentially, reactivity studies. If a His-tag is used to purify the 157 protein and not removed prior to such an analysis, then careful controls must be performed 158 as Cu binding to the tag cannot be excluded if the metal ion is added in excess. EPR can offer 159 a quick and powerful way to check the sample for the presence of any additional/adventitious 160 binding sites (see below). Other affinity tags can also be used of course, with a strep-tag recently used for the purification of the latest LPMO discovered in family AA15 providing a 161 162 prominent example (Sabbadin et al., 2018).

163

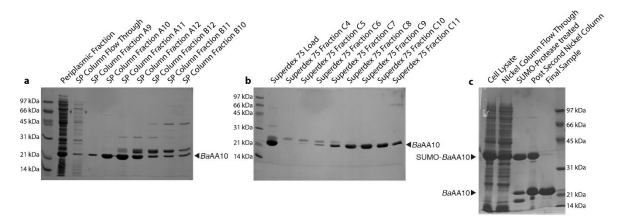
Once the desired constructs are obtained and confirmed as correct by Sanger sequencing, plasmids are transformed typically into BL21(DE3) *E. coli* cells as a first port of call. Other *E. coli* strains should also be considered. Transformants are plated out on selective LB agar plates and allowed to grow at 37 °C overnight. A 5 mL LB liquid culture is then inoculated using 168 a single colony the next day and the cells are allowed to grow at 37 °C in a shaker at 180 rpm 169 throughout the day. Once the culture is visibly turbid, 500 μ L is used to inoculate a 50 mL 170 culture which is left to grow overnight in the same conditions. The next day, 2 L baffled flasks 171 each containing 500 mL of LB are then inoculated using 5 mL each from the overnight culture and these are incubated at 37 °C, shaking at 180 rpm. The OD₆₀₀ is monitored closely and once 172 173 this has reached 0.6 the temperature is then typically cooled to 16 °C for about an hour. 174 Protein expression can then be induced by the addition of isopropyl β-D-1-175 thiogalactopyranoside (IPTG) to a final concentration of 1 mM and the cultures are left 176 overnight. The next day cells are harvested by centrifugation at 8,000 x g, the spent media discarded and cell pellets can be frozen at -80 °C until ready for subsequent use. 177

178

179 We use a modified periplasmic preparation protocol rather than the cold osmotic shock 180 protocol (Neu and Heppel, 1965) that is often used as we have found that we recovered better 181 yields of protein using this approach. Typically, cell paste is thoroughly resuspended in 3x 182 volumes of 20% sucrose (20 mM Tris can also be included to stabilise pH). To this, 40 µL of 10 183 mg/mL lysozyme is added for every gram of cell paste being used and the suspension is then left on ice for 1 hour. 60 µL of 1 M MgSO₄ is then added for every gram of cell paste being 184 185 used and the cells are left on ice for an additional 20 minutes. Cell debris is then harvested by 186 centrifugation at 10,000 x g for 20 minutes and the supernatant is removed as Fraction I to a 187 fresh container and kept on ice. The pellet is re-suspended in ice cold water and this is also 188 left on ice for a further 1 hour (some DNAse can be added at this point to try and reduce the 189 viscosity of the solution produced at the end of the process). The cell debris is again removed 190 by centrifugation at 10,000 x g for 20 minutes and the supernatant is removed as Fraction II. 191 Fraction I and II can be checked at this point to see whether the LPMO is abundantly present 192 in one fraction over the other, however, usually both fractions are combined for subsequent 193 purification.

194

195 If using an affinity tag, standard chromatographic approaches can be used for the purification 196 of LPMO. We typically perform purifications using FPLC instruments using an affinity 197 chromatography step for the initial purification and gel filtration for a final "polishing" step. 198 In the absence of a tag we would typically aim to purify via ion exchange chromatography 199 followed by gel filtration (see Figure 2a and b for example gels). We would recommend using 200 Good's buffers throughout any purification, these represent biologically compatible buffers 201 which should not interact significantly with biologically relevant cations such as copper (Good 202 et al., 1966). If attempting to measure copper binding to LPMO it is imperative that all buffers 203 are thoroughly de-metallated prior to the final step of the purification. This is typically 204 achieved by cycling the buffer through a 10 mL chelex column (Bio-Rad) for 24 hours before 205 being used in gel filtration. The protein should also be thoroughly treated with EDTA (typically 206 at 10 mM concentration or higher for at least an hour) to remove any metal that might already 207 be bound since LPMOs have very high affinity for copper. Indeed, it is probably this high 208 affinity for copper that resulted in low level activity, which could have been mistaken for 209 hydrolytic activity, during early studies on GH61s (Karlsson et al., 2001). The high affinity of 210 the enzyme for copper allows it to scavenge its cofactor even when present at very low 211 concentration and could also have contributed to the initial confusion over the identity of the 212 correct metal co-factor following the initial structure determinations for these enzymes 213 (Karkehabadi et al., 2008; Vaaje-Kolstad et al., 2005b, 2010).



215

Figure 2. Example gels for LPMO Purification strategies. **a**, gel showing initial purification of *Ba*AA10 from the periplasm using an SP ion exchange column. **b**, subsequent purification of BaAA10 using a Superdex 75 size exclusion column to yield the final sample (Hemsworth *et al.*, 2013b). **c**, SDS-PAGE gel showing stages from the purification of *Ba*AA10 using a SUMOtagged construct (Gregory *et al.*, 2016).

221

222 If an apo-enzyme is not desired, but rather a copper loaded sample is to be used for study, 223 prior to the final gel filtration step, CuCl₂ can be added in a slight excess (typically 1.5x) in 224 order to load the protein with copper prior to gel filtration. The sample is then typically passed 225 through a 16/600 Superdex 75 column (GE Healthcare) to remove any excess copper, giving 226 a highly purified, copper loaded sample ready for study. In order to store protein, we typically 227 concentrate to 10 mg/mL or more using a 3, 5 or 10 kDa Vivaspin column depending on the 228 size of the LPMO (avoid cellulose-based membranes concentrators). The sample is then flash 229 frozen in liquid nitrogen for storage at -80 °C.

230

Before discussing other strategies for LPMO production in *E. coli* and other organisms, there is another option for protein secretion to obtain mature LPMOs - to secrete the protein from gram positive bacteria into the culture medium (Nakagawa *et al.*, 2015). We have not used this approach ourselves, but this could be an excellent strategy for large scale LPMO production with industry often using gram-positive expression systems for the production of many stable and industrially-useful enzymes.

237 2.2 Recombinant expression and purification from the *E. coli* cytoplasm

238 Given that LPMOs are typically extracellular enzymes that contain disulfide bonds, the periplasmic expression protocol described above often represents the first port of call for 239 240 protein production (Courtade et al., 2017; Forsberg et al., 2014a; Hemsworth et al., 2013b; 241 Vaaje-Kolstad et al., 2005a, 2010). This is not the only option however, and there are reports 242 of LPMOs being expressed intracellularly using cleavable tags at the N-terminus to aid 243 purification (Forsberg et al., 2011, 2014b; Gregory et al., 2016). In order to achieve this, the 244 protease to be used to remove that tag must be carefully selected in order to leave the native 245 N-terminal histidine available for copper binding. Factor Xa (Forsberg et al., 2014b) and enterokinase are two such proteases. We prefer the use of a SUMO-tag (Champion-pET-246 247 SUMO, Invitrogen) which can be removed using SUMO-protease (Invitrogen) (Gregory et al., 248 2016). This method has the advantage that the SUMO-tag can act as a solubility enhancing 249 tag in addition to providing an N-terminal histidine tag to aid in the purification of the protein. 250 SUMO-protease recognises the tertiary structure of the SUMO domain and cleaves directly after a di-glycine motif which can be placed directly upstream of the N-terminal His coding region, so that cleavage results in the release of the native protein. Constructs can be carefully designed using the cloning strategies discussed above in order to achieve this without the use of restriction enzymes. This strategy was used to improve the yield of the AA10 from *Bacillus amyloliquefaciens* as much as 3-fold (Gregory *et al.*, 2016). While this protein lacked disulfide bonds which would have aided in the correct folding of the protein intracellularly, we have since used this strategy to express disulfide containing LPMOs intracellularly using specialised

- cells such as the NEB Express T7 cells (New England Biolabs) (Unpublished data).
- 259

The specific protocol for protein expression in *E. coli* is much the same as that discussed above with cells grown in shake flasks and the temperature typically being lowered to 16 or 20 °C upon IPTG induction to ensure correct protein folding. Cells are harvested the same way by centrifugation at 8,000 x g, with spent media being discarded and cell pellets frozen at -80 °C until ready for lysis.

265

266 Since the protein is now present in the cytoplasm, there is no need to perform the laborious 267 periplasmic prep protocol in order to release the protein from the cells. Cells are simply 268 resuspended in Buffer A (typically 50 mM HEPES pH 7, 250 mM NaCl, 30 mM imidazole) and 269 can be lysed by one's preferred method (typically sonication or French press is used in our 270 laboratories). The cell lysate can then be loaded directly onto a nickel resin to which the His6-271 SUMO-tagged LPMO will bind (Figure 2c). Protein is then released by the application of a 272 gradient from 0 to 100% buffer B (Buffer A + 300 mM imidazole) and 1.8 mL fractions are 273 collected across the gradient. Peak fractions containing His₆-SUMO-LPMO are combined 274 ready for tag removal. In order to prepare for tag cleavage, it is important to remove excess 275 imidazole and NaCl from the sample which can be detrimental to SUMO-protease function. 276 This can be achieved by buffer exchange on a 10 kDa MW cut-off concentrator, using a 277 desalting column or by dialysis. The protease buffer should also contain a reducing agent to 278 ensure optimal SUMO-protease activity since it is a cysteine protease. We, therefore, typically 279 buffer exchange our samples into 20 mM Tris pH 8, 5 mM β -mercaptoethanol. Small scale tag-280 cleavage reactions should be set up the first time that this strategy is used to try to obtain 281 optimal ratio of protease to SUMO-LPMO which can then be scaled up to the whole sample. 282 We typically perform tag removal at a SUMO-LPMO concentration of 1-2 mg/mL overnight at 283 4 °C.

284

Once the cleavage has been performed, the released tag, any uncleaved protein and the SUMO-protease are removed from the sample by passing it back down a nickel column ensuring that the flow through, which will contain the mature LPMO, is collected (Figure 2c). The differences in the LPMO's affinity for copper over nickel ensure that there is not transmetallation of the enzyme. The enzyme sample can then be concentrated, copper loaded or thoroughly EDTA treated as described above, before a final polishing step using a 16/600 Superdex 75 column (GE Healthcare) to yield the final sample (Figure 2c).

292 2.3 Recombinant expression and purification from yeast

For many LPMOs, expression in *E. coli* has not been possible and so laboratories have turned to other expression systems for challenging targets. *Pichia pastoris* is generally the most common yeast expression system used for protein production in academic laboratories and has been used for the production of several LPMOs (Kittl, Kracher, Burgstaller, Haltrich, & Ludwig, 2012; Bennati-Granier *et al.*, 2015; Tanghe *et al.*, 2015). While *P. pastoris* lacks the machinery required to perform the methylation of the N-terminal histidine found in fungal LPMOs, it maintains the ability to glycosylate proteins which may be important to ensure the correct folding of LPMOs from eukaryotic organisms.

301

302 The most common vector used for LPMO production in *P. pastoris* is the commercially 303 available pICZa (Invitrogen) (Bennati-Granier et al., 2015; Couturier et al., 2018; Kittl et al., 304 2012), although other vectors such as the pPpT4 vector can also be used (Tanghe et al., 2015). 305 These are shuttle vectors which allow all of the cloning to be performed in E. coli for speed 306 and ease before the vector is introduced into the *P. pastoris* host for protein expression. As 307 for expression in E. coli, similar factors need to be considered during construct design in P. 308 *pastoris*. The vectors contain signal sequences to direct the protein for secretion and so the 309 LPMO sequence must be carefully cloned such that the codon for the N-terminal histidine is 310 immediately placed after the cleavage site for the signal peptide. Alternatively, if the protein 311 is from a fungal source the native leader sequence of the LPMO gene can also be used rather 312 than the signal peptide available in the vector (Kittl et al., 2012; Tanghe et al., 2015).

313

314 The target gene is typically placed under the control of the AOX1 promoter in these vectors, 315 allowing induction of protein expression using methanol (Bennati-Granier et al., 2015; Couturier et al., 2018; Kittl et al., 2012). Cells are typically grown in defined media based on 316 317 the protocol provided by Invitrogen so we will not re-describe this here. Thanks to the use of 318 the signal sequence the protein is directed out of the cell into the media and so the protein 319 needs to be directly purified from here. Affinity tags can be incorporated at the C-terminus of 320 the protein to assist in the purification, but in many cases the protein has been purified 321 without the use of an affinity tag, for example Kittle et al (2012) used a multi-step purification 322 process involving hydrophobic chromatography, ion-exchange chromatography and finally 323 size-exclusion chromatography.

324

325 Copper loading procedures are much the same as those described above for LPMOs expressed 326 in *E. coli*, with the addition of a small excess of copper prior to the final gel-filtration step in 327 the purification. If the protein is to be used for crystallography, it is often necessary to 328 deglycosylate it. In our laboratories this step is typically achieved using Endoglycosidase H 329 (New England Biolabs) following the manufacturer's protocols. If, however, the protein is to 330 be used in assays, we typically do not deglycosylate the protein as this can affect both protein 331 stability and, potentially, protein-substrate interactions.

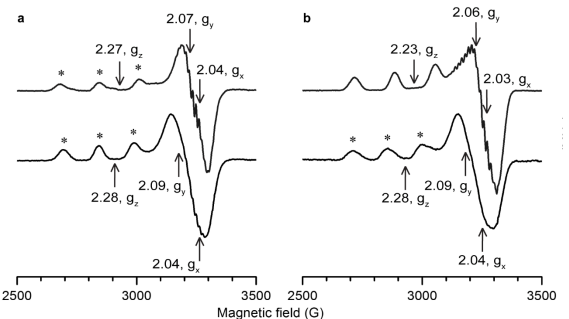
332 2.4 Recombinant expression and purification from fungi

333 The majority of LPMOs that have been studied to date are found in fungi and so there are 334 clear advantages to expressing these proteins in fungal hosts. These include correct insertion 335 of methylation on the N-terminal histidine sidechain as well as protein secretion into the 336 extracellular milieu following correct glycosylation and protein folding. Fungal recombinant 337 expression systems, however, are far less well established in academic laboratories. We have, 338 therefore, benefited greatly from our long-term collaboration with Novozymes who use a 339 highly specialised Aspergillus oryzae expression system for LPMO production on a large scale 340 (Frandsen et al., 2016; Lo Leggio et al., 2015; Quinlan et al., 2011). The high yields offered by 341 this system are able to meet the high sample demands for EPR (see below). These systems 342 are not readily available for use by others, however, and so examples of recombinant LPMO 343 expression from fungal sources are harder to find in the literature. Phillips et al (2011) early 344 on were able to purify *Neurospora crassa* LPMOs directly from secretomes when the fungus 345 was grown on cellulose. Since then, the Marletta lab and their colleagues have been able to 346 use a N. crassa expression system to produce both N. crassa LPMOs (Vu et al., 2014b) and 347 LPMOs from other fungi (Span et al., 2017). Using this approach, a knock-in system is used to 348 insert the LPMO gene downstream of a glyceraldehyde-3-phosphate dehydrogenase 349 promoter (GAP) (Bardiya and Shiu, 2007). The DNA is then incorporated into the N. crassa 350 genome by recombination and colonies in which the gene has been successfully incorporated 351 need to be screened for LPMO production. As for Pichia pastoris, the LPMO should be 352 secreted into the growth media allowing direct purification from the cleared media. Affinity 353 tags can be incorporated at the C-terminus of the protein to assist in this respect if they do 354 not interfere with subsequent planned experiments.

355

356 **3. Electron Paramagnetic Resonance Spectroscopy**

357 Once pure protein is obtained, a key technique that has been used in the field to gain insight 358 into the copper environment in LPMOs is Electron Paramagnetic Resonance (EPR) 359 spectroscopy. The fact that copper in its oxidised (+2) state (S=1/2, Kramers' doublet) has a single unpaired electron makes it EPR active. Therefore, by collecting EPR spectra from pure 360 361 LPMOs it has been possible to gain additional insight into the electronic state of the copper 362 centre in the resting protein. In addition, upon the inclusion of polysaccharide substrates in 363 the samples, it is sometimes possible to observe marked changes in the EPR spectra (Figure 364 3) (Bissaro et al., 2018; Borisova et al., 2015; Frandsen et al., 2016; Simmons et al., 2017), 365 suggesting significant changes occur at the copper active site in the presence of substrate, 366 which might be of key importance to activating oxygen and hence catalysis. In order to 367 interpret these spectra and understand what the changes in the EPR mean, it is necessary to 368 simulate the data. While there are algorithms that will attempt to simulate the data 369 automatically, these often fall short of a truly meaningful simulation. Optimal EPR simulation 370 is, therefore, still best performed by an expert EPR spectroscopist and requires considerable 371 patience and knowhow in order to obtain the best possible outcome. Here we describe the 372 steps taken to prepare samples for EPR analysis and provide information on how we interpret 373 our EPR spectra.



374

Figure 3: Example of changes observed in the X-band CW EPR spectra of an AA9 upon addition of substrate. **a**, *Ls*AA9A–G6 (top) and *Ls*AA9A with no cellohexaose (bottom) in low chloride conditions. **b**, *Ls*AA9A–G6 (top) and *Ls*AA9A with no cellohexaose (bottom) in high chloride conditions. ***** indicates signals from copper site where water is the exogenous ligand, other signals from species with chloride as exogenous ligand. Figure adapted from Frandsen *et al* (2016).

381 3.1 EPR sample preparation for LPMOs

382 Most EPR spectroscopy on LPMOs has been performed in frozen solution at X-band 383 frequencies (ca 9.3 GHz). At these frequencies, sample holders are typically quartz tubes with 384 roughly 3 to 5 mm internal diameters. The height of the resonator cavity will be of the order 385 of 10-15 mm, meaning that ideally a sample volume of ca 300-600 μ L is required. In reality, 386 by optimising the collection parameters, good EPR spectra can be obtained with much smaller 387 volumes, in the order of 80 to 120 µL with a concentration of at least 0.15 mM Cu. Albeit often 388 seen by biochemists as a highly demanding technique in terms of protein consumption, EPR 389 is non-destructive, and often samples can be fully recovered and re-used after the 390 experiments. In our hands, the repeated cooling/warming cycles that a sample typically 391 experiences during collection of EPR spectra do not lead to protein degradation. Obtaining a 392 signal with good signal-to-noise is a function of several experimental variables including the 393 capabilities of the EPR spectrometer, the temperature at which the spectra are collected, the 394 number of scans and the experimental settings. However, good spectra with excellent signal-395 to-noise ratios can be obtained with sample concentrations and volumes reported above at 396 100 to 170 K with acquisition settings reported in Table 1. Of particular note is the power 397 setting which must be at a level which does not cause saturation of the absorbance, especially 398 in concentrated samples. This level should be such that relaxation rates are greater than 399 absorption rates. For copper spectroscopy, relatively high power settings can be used because of the high relaxation rates associated with a metal centre. For detailed spectra, 400 401 then higher sample concentrations may be needed, possibly up to 1-2 mM concentration. 402 Great caution is needed at these concentrations however since face-to-face dimerization of 403 LPMOs may occur with the result that two nearby Cu centres will change their coordination 404 geometries or will lead to dipolar relaxation effects that perturb spin Hamiltonian values. Our 405 experience in this regard is that concentrations of ca 2 mM lead to spectral changes that could

406 be associated with protein dimerization.

- 407
- 408 Table 1: Standard acquisition parameters for frozen solution X-band EPR spectra of LPMOs.

| Centre field | 3000 G | |
|----------------------|---------------|--|
| Sweep | 2300 – 3700 G | |
| Modulation amplitude | 4 G | |
| Modulation frequency | 100 kHz | |
| Time constant | 163.8 msec | |
| Sweep time | 92 sec | |
| Conversion time | 90 msec | |
| Power | 5 – 10 mW | |
| Scans | 1-8 | |
| Temperature | 150 – 170 К | |

409

410

411 EPR spectra collected in aqueous solution often require the use of a glassing agent such as 412 glycerol or sucrose to minimise the formation of ice crystals. Such crystals can increase 413 microwave scattering, reduce the microwave cavity quality factor, cause changes in protein 414 structure, pH or ionic strength, or even shatter the microwave tube! For smaller tubes which 415 are used with high frequency EPR this can be a particular problem. Tube breakages for X-416 band EPR tubes are less common, especially if thick wall tubes are used. For work with LPMO 417 solutions, glycerol is typically the glassing agent of choice (10-50% v/v), although sucrose can 418 also be used (2 mg/ μ L). When combined with very rapid freezing then the resulting sample 419 usually forms a good glass. The cooling method needs to be carefully considered however in 420 that very rapid freezing is required to prevent entropic driven processes from occurring at the 421 metal centre. For instance, at room temperature the copper coordination may include the 422 three atoms of the ligating histidine brace along with a fourth exogenous ligand. Cooling 423 slowly to typical EPR collection temperatures may drive the formation of a five coordinate Cu 424 site in which a further water molecule is recruited into the coordination sphere. The result 425 would be that the EPR spectrum collected at low temperature is not representative of the 426 solution phase structure. Therefore, very rapid freezing is required to 'trap' the room 427 temperature species. This is best achieved by plunging the EPR sample tube into a cold liquid 428 (e.g. methanol in dry ice (Chaplin et al., 2016)) rather than into liquid nitrogen where — despite 429 the very low temperature of the liquid—the resulting bubbles of N₂ insulate the tube from 430 the liquid to prevent very rapid freezing. Notwithstanding the advantages of rapid cooling of 431 samples, the experimenter needs to be aware that any heterogeneity in protein structure, 432 where the interconversion is slower than the cooling timescale, may be trapped by the cooling 433 process, such that sample heterogeneity is increased, the result of which is a severe loss of 434 spectral resolution. It is not unusual for rapidly-cooled samples to have broader spectral 435 peaks that those samples cooled less rapidly (Guzzi et al., 2001). Necessarily, therefore, 436 different cooling techniques need to be explored. In our experience, a happy compromise 437 can be found for X-band LPMO samples when a sample at room temperature is plunged into 438 the stream of cooling N2 gas which cools the resonator. Also, on occasion for high 439 concentration samples, it has been possible to collect spectra of LPMO solutions at room 440 temperature. The slow tumbling rate of the protein in solution means that the spectrum is still anisotropic, from which it is possible to link the spin Hamiltonian parameters at roomtemperature with those at low temperature (unpublished data).

443

444 The type of glassing agent that is employed in these studies is an important consideration in 445 experimental design. The principal consideration is that glassing agents have a 'poly-ol' 446 structure which can mimic that of the natural polysaccharide substrate of the enzyme and 447 possibly interfere with the Cu active site by binding close to it. Accordingly, the dependency 448 of the spectrum on the concentration of glassing agent should be established in any 449 Unfortunately, an examination of the LPMO literature shows that such experiment. 450 dependency experiments are relatively rare and only appear to have been done in a handful 451 of cases (Sabbadin et al., 2018). When there is evidence to suggest that the glassing agent is 452 interfering with the Cu centre, then it is recommended changing the agent to another (e.g. 453 DMSO, Ficoll) or that the spectrum is collected without any glassing agent at all. The latter 454 method appears to work surprisingly well for LPMO samples (Bissaro et al., 2018; Borisova et 455 al., 2015; Chaplin et al., 2016; Forsberg et al., 2014b, 2014a, 2016).

456

457 One of the great advantages of EPR is that spectra can be obtained on a sample in any state 458 of matter: liquid, solid, slurry, frozen glass or even a gas. For LPMOs, of course, since the 459 substrates are usually insoluble polysaccharides, the interaction of the LPMO with its natural 460 substrate can be studied directly. Moreover, introducing a solid phase sample into a solution 461 to create a slurry, essentially removes the need for a glassing agent. Necessarily, the nature 462 of the solid phase means that protein binding may be slow or incomplete and it is advisable 463 to leave the slurry sample for some time before then acquiring the EPR spectra. In the 464 majority of cases, at least part of the protein binds very rapidly (Frandsen et al., 2016; 465 Simmons et al., 2017), but for some samples we have noticed that full binding can take up to 466 several hours and/or cannot be achieved, so would recommend that samples are left 467 incubating overnight in the absence of oxygen and/or a reducing agent. It is also important 468 that the introduced sample is isotropically distributed within the EPR tube, otherwise 469 alignment effects could complicate the spectrum. Beyond these considerations, it appears as 470 if solid state slurries of LPMO solutions with their insoluble substrates are perfectly good 471 samples for both CW and pulsed EPR studies (Bissaro et al., 2018; Borisova et al., 2015; 472 Frandsen et al., 2016; Simmons et al., 2017).

473 3.2 Multi-frequency collection

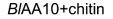
474 EPR spectra of LPMOs in the Cu(II) states give spin Hamiltonian parameters in which there is 475 either a typical 'type 2' Cu centre with large IA_zI values (ca 400-600 MHz) (Borisova et al., 476 2015; Chaplin et al., 2016; Couturier et al., 2018; Forsberg et al., 2014a, 2016; Frandsen et al., 477 2016; Hansson et al., 2017; Hemsworth et al., 2014; Quinlan et al., 2011; Sabbadin et al., 2018; 478 Simmons et al., 2017; Span et al., 2017) or a more rhombic Cu(II) centre that has a reduced 479 |A_z| value but large |A_x| and |A_y| values (Bissaro *et al.*, 2018; Forsberg *et al.*, 2014b; Gregory 480 et al., 2016; Hemsworth et al., 2013b). In both cases the perpendicular region of the spectrum cannot be unambiguously simulated from the X-band data alone. For this reason, if an in-481 depth analysis is required, collecting spectra at different frequencies is paramount. 482 483 Depending on the kind of information desired, one can decide to use lower (e.g. S-band) or 484 higher (e.g. Q-band) frequency than X-band, with the latter being usually the preferred 485 method. A couple of examples of this approach are present in the LPMO literature (Couturier 486 et al., 2018; Hansson et al., 2017), but it is easy to imagine that more and more will appear as

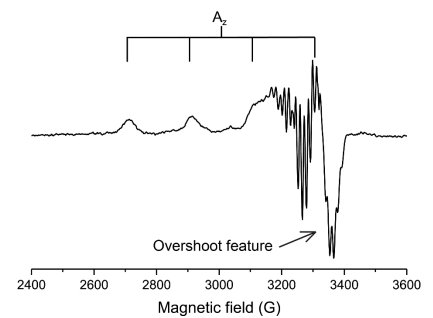
487 the fine details of the electronics at the Cu active site are investigated. For Q-band spectra, 488 guartz tubes of ca 2 mm in diameter are employed, while the samples are typically 15-30 µL 489 in volume at a concentration of copper of 1 mM or higher. The same, if not greater, caution 490 in optimising the acquisition settings as for X-band should be exercised when collecting Q-491 band spectra. In our experience, obtaining good quality Q-band spectra, especially if protein 492 concentration is an issue due to protein aggregation at high concentration or limited amount of protein available, is a non-trivial problem, but it can be sometimes overcome with patience 493 494 and trials of different settings. Typical settings used for our experiments include centre field 495 at 1150 mT, sweep width 150 mT, modulation width 0.6 – 1 mT, time constant 0.3 s, scan time 496 $1 - 2 \min$, power 0.5 - 1 mW.

497

507

498 Collecting CW Q-band spectra has a further benefit in that EPR spectra of LPMOs can 499 sometimes contain high field features (Bissaro et al., 2018). These features under simulation 500 (see below) are very hard to assign unambiguously as they could arise either from large 501 hyperfine coupling constants or from low principal g values. There are indeed cases in the 502 literature where such features have been simulated with low q values. It is, however, well 503 known in the wider EPR community that the high field signals may arise from a spectral 504 artefact known as an overshoot feature, associated with large |Az| hyperfine coupling values 505 (see figure 4). By collecting CW EPR spectra at different frequencies, the normal choice being 506 Q-band, this misinterpretation can be avoided.





508

Figure 4: X-band CW EPR spectrum of an AA10 from *Bacillus licheniformis, BI*AA10, after
addition of squid pen chitin, showing the overshoot feature associated with the large |A_z|
value (unpublished data).

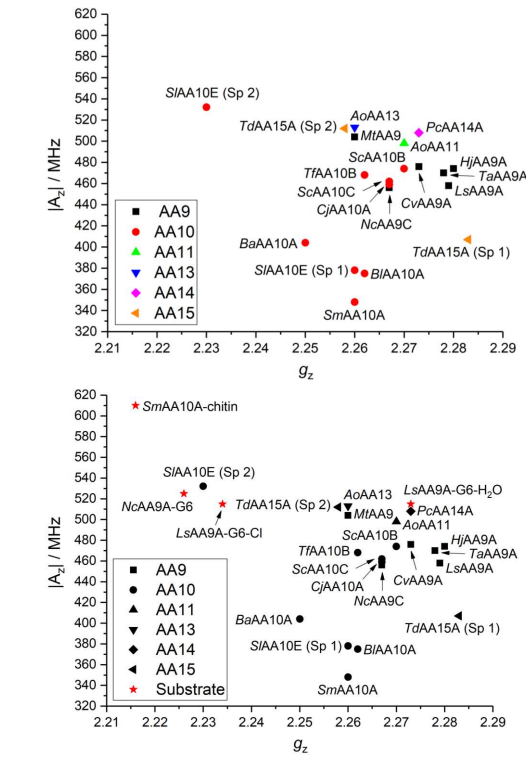
512

513 3.3 Simulation procedures

514 Once the CW spectra have been recorded, simulations have to be performed in order to gain

- 515 information about the copper active site. Several simulation packages are available to users,
- some of them developed by EPR instruments manufacturers (*e.g.* the WinEPR and SimFonia

517 suites (Bruker)), others made available by researchers in the field (e.g. SpinCount (The 518 Hendrich Metalloprotein Group) or EasySpin (Stoll and Schweiger, 2006)). Regardless of the 519 programme used to carry out the simulations, some general considerations can be done on 520 how to approach such a task, in particular for those who might not have encountered EPR 521 before. Determining the spin Hamiltonian parameters for the parallel region of the spectrum 522 is in general straightforward, as at least 3 of the 4 hyperfine peaks to the Cu can be clearly 523 observed between 2500 and 3100 G in the X-band spectra of LPMOs. Careful analysis of this 524 region of the spectrum can also reveal the presence of multiple species or adventitious 525 binding sites, if a second set of hyperfine peaks is present either as individual peaks or as 526 shoulders of the main component of the spectrum. Specifically, we would recommend 527 particular care if the LPMO carries an affinity tag, as highlighted in section 2.1. In these 528 circumstances, EPR provides a very powerful tool for the identification of second binding sites 529 in amounts as low as 10% of the main species. As mentioned above, most LPMOs show a type 530 2 copper active site according to the Peisach-Blumberg (P-B) classification (Peisach and 531 Blumberg, 1974), with SOMO mostly $d(x^2-y^2)$ in character and close to square planar 532 geometry. While the P-B classification is a useful 'first-step' guide to the general structure of 533 the copper ion in the active sites of LPMOs, we would caution against its in-depth use for the 534 assignment of coordinating atoms (e.g. determining between CuN₂O₂ and CuN₃O 535 coordination spheres), due to the multiple contributing factors to both g_z and $|A_z|$ values, the 536 most problematic of which is the unknown degree of covalency in the metal-ligand bonds. In 537 this regard, it is indeed possible to notice the high degree of variability in the published EPR 538 parameters of LPMOs in a P-B plot (Figure 5 and Table 2) even for proteins that, from a 539 structural point of view, show the same coordination environment (recently reviewed by 540 Ciano et al. (2018) and Vu and Ngo (2018)). Finally, in the context of the broad interpretation 541 of spin Hamiltonian parameters of LPMOs, there are reports of compressed trigonal 542 bipyramidal structures with $d(z^2)$ SOMOs, but this assignment is not in accord with the 543 spectral envelopes observed in all LPMO EPR spectra (Vu and Ngo, 2018).



546



Figure 5: Peisach-Blumberg plots of published EPR data for LPMOs (data and references reported in Table 2). For clarity and consistency, all enzymes have been named using the CAZy abbreviation, where two letters (in italics) define the organism the protein originates from, followed by the Auxiliary Activity family and the specific enzyme, if reported and more than one are present in the same organism (*e.g. Ls*AA9A). Top: P-B plot of LPMOs divided by family, according to the legend on the plot. Bottom: P-B plot as above, with the parameters for substrate-bound enzymes included; "G6" is used for cellohexaose.

Table 2: Table of g_z and $|A_z|$ LPMO EPR values published to date, used for the plots in Figure

557

| 5. | | | |
|------------------------------------|--------------------------------|--------------|------------------------|
| LPMO | Reference | gz | A _z / MHz |
| NcAA9C | Borisova <i>et al.,</i> 2015 | 2.267 | 456 |
| TaAA9A | Quinlan et al., 2011 | 2.278 | 470 |
| HjAA9A | Hansson <i>et al.,</i> 2017 | 2.280 | 474 |
| MtAA9 | Span <i>et al.,</i> 2017 | 2.260 | 504 |
| CvAA9A | Simmons et al. 2017 | 2.273 | 476 |
| <i>Ls</i> AA9A | Frandsen <i>et al.</i> , 2016 | 2.279 | 458 |
| BaAA10A | Hemsworth <i>et al.,</i> 2013b | 2.25 | 404 |
| CjAA10A | Forsberg et al., 2016 | 2.267 | 462 |
| SmAA10A | Forsberg et al., 2014b | 2.260 | 348 |
| ScAA10C | Forsberg et al., 2014b | 2.267 | 459 |
| B/AA10 | Forsberg et al., 2014b | 2.262 | 375 |
| <i>Tf</i> AA10B | Forsberg et al., 2014b | 2.262 | 468 |
| ScAA10B | Forsberg et al., 2014a | 2.270 | 474 |
| S/AA10E | Chaplin et al., 2016 | 2.260 (Sp 1) | 378 (Sp 1) |
| | | 2.230 (Sp 2) | 532 (Sp 2) |
| AoAA11 | Hemsworth et al., 2014 | 2.27 | 498 |
| <i>Ao</i> AA13 | Lo Leggio <i>et al.,</i> 2015 | 2.26 | 513 |
| PcAA14A | Couturier <i>et al.,</i> 2018 | 2.273 | 508 |
| TdAA15A | Sabbadin <i>et al.,</i> 2018 | 2.283 (Sp 1) | 407 (Sp 1) |
| | | 2.258 (Sp 2) | 512 (Sp 2) |
| NcAA9C-G6 | Borisova <i>et al.,</i> 2015 | 2.226 | 525 |
| <i>Ls</i> AA9A-G6-H ₂ O | Frandsen <i>et al.</i> , 2016 | 2.273 | 515 |
| LsAA9A-G6-Cl | Frandsen <i>et al.</i> , 2016 | 2.234 | 515 |
| SmAA10A-chitin | Bissaro et al., 2018 | 2.216 | 610 |

558

559

560 The perpendicular region of the spectrum, instead, is not as straightforward to interpret and 561 simulate, especially from X-band data alone. Here, the partial overlap of the q_x/q_y and $|A_x|/|A_y|$ 562 values and the usually broad appearance of the spectrum do not allow to unambiguously 563 determine the spin Hamiltonian parameters of the copper ion. Great care should therefore 564 be used in these circumstances. The problem can be overcome by simultaneous simulating 565 both X- and Q-band data, when multi-frequency collection of the spectra is a possibility. Simultaneous fits of the data from two different frequencies significantly narrows the sets of 566 567 possible spin Hamiltonian values, with the further advantage that artefacts arising from 568 overshoot features, whether present, can be all but eliminated.

569

Analysis of the superhyperfine (SHF) coupling, when visible, can provide insights into the coordination environment of the copper active site and possible changes triggered by the binding of substrate (Frandsen *et al.*, 2016). Simulation of the SHF coupling can be a very time consuming and, at times, frustrating challenge, but it can provide key information about the active site. In our experience, a reliable simulation of superhyperfine coupling patterns can take many days to perform accurately. Moreover, it is important to point out that the most direct and reliable method to obtain coupling values to coupled nuclei is the use of pulsed 577 EPR techniques such as ENDOR or HYSCORE (not discussed herein), in which the orientation-578 selective feature of the techniques allows some separation of the individual coupling 579 components of each coordinating atom. However, as pulsed techniques are not easily 580 accessible and can require high amount of protein, careful analysis of CW spectra can give 581 surprisingly accurate information about the type and number of nuclei coupled to the 582 unpaired electron and their coupling values. The risk of over-parameterising the problem 583 should be taken in account, and we would recommend to proceed in steps and add further 584 coupled nuclei only if the experimental data cannot be satisfactory simulated otherwise. 585 Simultaneous simulations of both first and second derivative spectra can be of great help in 586 establishing the coupling values, although usually only the largest component of the coupling, 587 when anisotropic, can be determined. If possible, isotopic labelling (e.g. ¹⁵N) should be 588 performed in order to restrict the set of possible values which would give good fit to the 589 experimental data.

590

591 Finally, some authors have used DFT calculations to augment their interpretations of EPR data 592 (Bissaro *et al.*, 2018). While useful in a broad context, there are well-known problems in using 593 DFT to give accurate representations of spin Hamiltonian parameters (Neese, 2009, 2017). 594 Indeed, these problems appear to be evident in DFT calculations performed on LPMOs. The 595 key difficulties stem from the issues associated with the inaccurate modelling of the Fermi 596 contact term (which, in turn, arises from configuration interaction of the SOMO with s-orbitals 597 and polarisation of core electrons at the copper ion), the relatively poor modelling of medium-598 distance electron-electron interactions, the important role of dispersion forces, and the 599 tendency of DFT calculations to overly apply covalency to metal-ligand bonding. In partial 600 mitigation of these problems, we would recommend using a basis set in which core functions 601 are enhanced (e.g. CP(PPP) (Neese, 2002)). For the latter issue, a functional should be 602 selected in which the degree of Hartree-Fock (HF) exchange has been manually increased, e.g. 603 B3LYP-D functional has 20% HF exchange set as default, but should be increased to ~38% with 604 the final value being determined by trial and error. However, as is evident in the preceding 605 sentence, this is necessarily an empirical procedure and our experience in this regard is that 606 the spin Hamiltonian parameters obtained from DFT calculations for LPMOs can be 607 significantly different from the experimental values. As such, great caution must be exercised 608 in using DFT-determined spin Hamiltonian parameters for in-depth interpretation of the 609 active site electronics, even when there appears to be good agreement with the experimental 610 values. Indeed, whether the results of DFT calculations on spin Hamiltonian parameters can 611 give useful absolute results at all in such a complicated system is debateable. Our advice in 612 this regard therefore is that trends in calculated parameters may be meaningful (when the 613 same basis sets, functionals and methods are used between systems), as might the absolute 614 signs of calculated hyperfine coupling values, but that further values should not be used as a 615 basis for in-depth analysis of the electronic structure of the SOMO without a critical 616 evaluation of the underlying issues. In this context, our further advice is that a ligand field 617 analysis in which many of the 'problematic' values are parameterised offers a more reliable 618 and suitably-caveated analysis of LPMO EPR spectra, from which appropriately framed 619 conclusions can be drawn.

621 6. Acknowledgements

- 622 GRH gratefully acknowledges support from a BBSRC (Biotechnology and Biological Sciences
- 623 Research Council) David Phillips Fellowship (grant # BB/N019970/1). LC, GJD and PHW also
- 624 gratefully acknowledge the support of the BBSRC under grant BB/L001926/1.

625 7. References

- 626 Agger, J.W., Isaksen, T., Várnai, A., Vidal-Melgosa, S., Willats, W.G.T., Ludwig, R., Horn, S.J.,
- 627 Eijsink, V.G.H., and Westereng, B. (2014). Discovery of LPMO activity on hemicelluloses
- shows the importance of oxidative processes in plant cell wall degradation. Proc. Natl. Acad.
 Sci. U. S. A. *111*, 6287–6292.
- Bardiya, N., and Shiu, P.K.T. (2007). Cyclosporin A-resistance based gene placement system
 for Neurospora crassa. Fungal Genet. Biol. 44, 307–314.
- Beeson, W.T., Vu, V. V, Span, E.A., Phillips, C.M., and Marletta, M.A. (2015). Cellulose
- 633 degradation by polysaccharide monooxygenases. Annu. Rev. Biochem. *84*, 923–946.
- Bennati-Granier, C., Garajova, S., Champion, C., Grisel, S., Haon, M., Zhou, S., Fanuel, M.,
- 635 Ropartz, D., Rogniaux, H., Gimbert, I., et al. (2015). Substrate specificity and regioselectivity
- 636 of fungal AA9 lytic polysaccharide monooxygenases secreted by Podospora anserina.
- 637 Biotechnol. Biofuels 8, 90.
- Bey, M., Zhou, S., Poidevin, L., Henrissat, B., Coutinho, P.M., Berrin, J.-G., and Sigoillot, J.-C.
- (2013). Cello-oligosaccharide oxidation reveals differences between two lytic polysaccharide
 monooxygenases (family GH61) from Podospora anserina. Appl. Environ. Microbiol. *79*,
 488–496.
- 642 Bischof, R.H., Ramoni, J., and Seiboth, B. (2016). Cellulases and beyond: the first 70 years of 643 the enzyme producer Trichoderma reesei. Microb. Cell Fact. *15*, 106.
- Bissaro, B., Røhr, A.K., Müller, G., Chylenski, P., Skaugen, M., Forsberg, Z., Horn, S.J., VaajeKolstad, G., and Eijsink, V.G.H. (2017). Oxidative cleavage of polysaccharides by monocopper
 enzymes depends on H2O2. Nat. Chem. Biol. *851*, 1.
- 647 Bissaro, B., Isaksen, I., Vaaje-Kolstad, G., Eijsink, V.G.H., and Røhr, Å.K. (2018). How a Lytic 648 Polysaccharide Monooxygenase Binds Crystalline Chitin. Biochemistry *57*, 1893–1906.
- 649 Borisova, A.S., Isaksen, T., Dimarogona, M., Kognole, A.A., Mathiesen, G., Várnai, A., Røhr,
- 650 Å.K., Payne, C.M., Sørlie, M., Sandgren, M., et al. (2015). Structural and Functional
- 651 Characterization of a Lytic Polysaccharide Monooxygenase with Broad Substrate Specificity.
- 652 J. Biol. Chem. *290,* 22955–22969.
- 653 Bruker EPR Simulation Suites. https://www.bruker.com/products/mr/epr/epr-654 software/simulation-suites/overview.html
- 655 Chaplin, A.K., Wilson, M.T., Hough, M.A., Svistunenko, D.A., Hemsworth, G.R., Walton, P.H.,
- 656 Vijgenboom, E., and Worrall, J.A.R. (2016). Heterogeneity in the Histidine-brace Copper
- 657 Coordination Sphere in Auxiliary Activity Family 10 (AA10) Lytic Polysaccharide
- 658 Monooxygenases. J. Biol. Chem. 291, 12838–12850.
- 659 Chiu, E., Hijnen, M., Bunker, R.D., Boudes, M., Rajendran, C., Aizel, K., Oliéric, V., Schulze-
- 660 Briese, C., Mitsuhashi, W., Young, V., et al. (2015). Structural basis for the enhancement of
- virulence by viral spindles and their in vivo crystallization. Proc. Natl. Acad. Sci. U. S. A. 112,

- 662 3973–3978.
- 663 Ciano, L., Davies, G.J., Tolman, W.B., and Walton, P.H. (2018). Bracing copper for the 664 catalytic oxidation of C–H bonds. Nat. Catal. *1*, 571–577.
- 665 Courtade, G., Wimmer, R., Røhr, A.K., Preims, M., Felice, A.K.G., Dimarogona, M., Vaaje-
- Kolstad, G., Sørlie, M., Sandgren, M., Ludwig, R., et al. (2016). Interactions of a fungal lytic
 polysaccharide monooxygenase with β-glucan substrates and cellobiose dehydrogenase.
 Proc. Natl. Acad. Sci. U. S. A. *113*, 5922–5927.
- 669 Courtade, G., Le, S.B., Sætrom, G.I., Brautaset, T., and Aachmann, F.L. (2017). A novel 670 expression system for lytic polysaccharide monooxygenases. Carbohydr. Res.
- 671 Couturier, M., Ladevèze, S., Sulzenbacher, G., Ciano, L., Fanuel, M., Moreau, C., Villares, A.,
- 672 Cathala, B., Chaspoul, F., Frandsen, K.E., et al. (2018). Lytic xylan oxidases from wood-decay
- 673 fungi unlock biomass degradation. Nat. Chem. Biol. 14, 306–310.
- 674 Crouch, L.I., Labourel, A., Walton, P.H., Davies, G.J., and Gilbert, H.J. (2016). The
- 675 Contribution of Non-catalytic Carbohydrate Binding Modules to the Activity of Lytic
- 676 Polysaccharide Monooxygenases. J. Biol. Chem. *291*, 7439–7449.
- 677 Forsberg, Z., Vaaje-Kolstad, G., Westereng, B., Bunæs, A.C., Stenstrøm, Y., MacKenzie, A.,
- 678 Sørlie, M., Horn, S.J., and Eijsink, V.G.H. (2011). Cleavage of cellulose by a CBM33 protein.
 679 Protein Sci. 20, 1479–1483.
- 680 Forsberg, Z., Mackenzie, A.K., Sørlie, M., Røhr, Å.K., Helland, R., Arvai, A.S., Vaaje-Kolstad,
- 681 G., and Eijsink, V.G.H. (2014a). Structural and functional characterization of a conserved pair
- of bacterial cellulose-oxidizing lytic polysaccharide monooxygenases. Proc. Natl. Acad. Sci.
- 683 U. S. A. *111*, 8446–8451.
- Forsberg, Z., Røhr, Å.K., Mekasha, S., Andersson, K.K., Eijsink, V.G.H., Vaaje-Kolstad, G., and
 Sørlie, M. (2014b). Comparative study of two chitin-active and two cellulose-active AA10-
- type lytic polysaccharide monooxygenases. Biochemistry *53*, 1647–1656.
- 687 Forsberg, Z., Nelson, C.E., Dalhus, B., Mekasha, S., Loose, J.S.M., Crouch, L.I., Røhr, A.K.,
- 688 Gardner, J.G., Eijsink, V.G.H., and Vaaje-Kolstad, G. (2016). Structural and Functional
 689 Analysis of a Lytic Polysaccharide Monooxygenase Important for Efficient Utilization of
- 690 Chitin in Cellvibrio japonicus. J. Biol. Chem. *291*, 7300–7312.
- 691 Frandsen, K.E.H., Simmons, T.J., Dupree, P., Poulsen, J.-C.N., Hemsworth, G.R., Ciano, L.,
- Johnston, E.M., Tovborg, M., Johansen, K.S., von Freiesleben, P., et al. (2016). The molecular
- basis of polysaccharide cleavage by lytic polysaccharide monooxygenases. Nat. Chem. Biol.*12*, 298–303.
- 695 Frommhagen, M., Sforza, S., Westphal, A.H., Visser, J., Hinz, S.W.A., Koetsier, M.J., van
- 696 Berkel, W.J.H., Gruppen, H., and Kabel, M.A. (2015). Discovery of the combined oxidative
- 697 cleavage of plant xylan and cellulose by a new fungal polysaccharide monooxygenase.
- 698 Biotechnol. Biofuels *8*, 101.
- 699 Gardner, J.G., Crouch, L., Labourel, A., Forsberg, Z., Bukhman, Y. V, Vaaje-Kolstad, G.,
- Gilbert, H.J., and Keating, D.H. (2014). Systems biology defines the biological significance of
- redox-active proteins during cellulose degradation in an aerobic bacterium. Mol. Microbiol.
- *94,* 1121–1133.
- 703 Good, N.E., Winget, G.D., Winter, W., Connolly, T.N., Izawa, S., and Singh, R.M. (1966).

- Hydrogen ion buffers for biological research. Biochemistry 5, 467–477.
- 705 Gregory, R.C., Hemsworth, G.R., Turkenburg, J.P., Hart, S.J., Walton, P.H., and Davies, G.J.
- 706 (2016). Activity, stability and 3-D structure of the Cu(ii) form of a chitin-active lytic
- polysaccharide monooxygenase from Bacillus amyloliquefaciens. Dalton Trans. 45, 16904–
 16912.
- Guzzi, R., Stirpe, A., Verbeet, M., and Sportelli, L. (2001). Structural heterogeneity of blue
 copper proteins: an EPR study of amicyanin and of wild-type and Cys3Ala/Cys26Ala mutant
 azurin. Eur. Biophys. J. *30*, 171–178.
- Hansson, H., Karkehabadi, S., Mikkelsen, N., Douglas, N.R., Kim, S., Lam, A., Kaper, T.,
- 713 Kelemen, B., Meier, K.K., Jones, S.M., et al. (2017). High-resolution structure of a lytic
- polysaccharide monooxygenase from Hypocrea jecorina reveals a predicted linker as anintegral part of the catalytic domain. J. Biol. Chem.
- 716 Harris, P. V, Welner, D., McFarland, K.C., Re, E., Navarro Poulsen, J.-C., Brown, K., Salbo, R.,
- 717 Ding, H., Vlasenko, E., Merino, S., et al. (2010). Stimulation of lignocellulosic biomass
- 718 hydrolysis by proteins of glycoside hydrolase family 61: structure and function of a large,
- r19 enigmatic family. Biochemistry *49*, 3305–3316.
- Harris, P. V, Xu, F., Kreel, N.E., Kang, C., and Fukuyama, S. (2014). New enzyme insights drive advances in commercial ethanol production. Curr. Opin. Chem. Biol. *19*, 162–170.
- Hemsworth, G.R., Davies, G.J., and Walton, P.H. (2013a). Recent insights into copper-
- containing lytic polysaccharide mono-oxygenases. Curr. Opin. Struct. Biol. 23, 660–668.
- Hemsworth, G.R., Taylor, E.J., Kim, R.Q., Gregory, R.C., Lewis, S.J., Turkenburg, J.P., Parkin,
- A., Davies, G.J., and Walton, P.H. (2013b). The copper active site of CBM33 polysaccharide
- 726 oxygenases. J. Am. Chem. Soc. 135, 6069–6077.
- 727 Hemsworth, G.R., Henrissat, B., Davies, G.J., and Walton, P.H. (2014). Discovery and
- characterization of a new family of lytic polysaccharide monooxygenases. Nat. Chem. Biol.*10*, 122–126.
- Hemsworth, G.R., Johnston, E.M., Davies, G.J., and Walton, P.H. (2015). Lytic Polysaccharide
 Monooxygenases in Biomass Conversion. Trends Biotechnol. *33*, 747–761.
- Himmel, M.E., Ding, S.-Y., Johnson, D.K., Adney, W.S., Nimlos, M.R., Brady, J.W., and Foust,
- T.D. (2007). Biomass recalcitrance: engineering plants and enzymes for biofuels production.
- 734 Sci. (New York, NY) *315*, 804–807.
- Horn, S.J., Vaaje-Kolstad, G., Westereng, B., and Eijsink, V.G. (2012). Novel enzymes for the
 degradation of cellulose. Biotechnol. Biofuels *5*, 45.
- Karkehabadi, S., Hansson, H., Kim, S., Piens, K., Mitchinson, C., and Sandgren, M. (2008). The
 first structure of a glycoside hydrolase family 61 member, Cel61B from Hypocrea jecorina, at
 1.6 A resolution. *383*, 144–154.
- 740 Karlsson, J., Saloheimo, M., Siika-Aho, M., Tenkanen, M., Penttilä, M., and Tjerneld, F.
- 741 (2001). Homologous expression and characterization of Cel61A (EG IV) of Trichoderma
- 742 reesei. Eur. J. Biochem. *268*, 6498–6507.
- 743 Kittl, R., Kracher, D., Burgstaller, D., Haltrich, D., and Ludwig, R. (2012). Production of four
- 744 Neurospora crassa lytic polysaccharide monooxygenases in Pichia pastoris monitored by a

- 745 fluorimetric assay. Biotechnol. Biofuels 5, 79.
- 746 Klock, H.E., and Lesley, S.A. (2009). The Polymerase Incomplete Primer Extension (PIPE)
- 747 method applied to high-throughput cloning and site-directed mutagenesis. Methods Mol.748 Biol. *498*, 91–103.
- 749 Kuusk, S., Bissaro, B., Kuusk, P., Forsberg, Z., Eijsink, V.G.H., Sørlie, M., and Väljamäe, P.
- (2018). Kinetics of H2O2-driven degradation of chitin by a bacterial lytic polysaccharide
 monooxygenase. J. Biol. Chem. *293*, 523–531.
- Langston, J.A., Shaghasi, T., Abbate, E., Xu, F., Vlasenko, E., and Sweeney, M.D. (2011).
- Oxidoreductive cellulose depolymerization by the enzymes cellobiose dehydrogenase and
 glycoside hydrolase 61. Appl. Environ. Microbiol. *77*, 7007–7015.
- Lo Leggio, L., Simmons, T.J., Poulsen, J.-C.N., Frandsen, K.E.H., Hemsworth, G.R., Stringer,
- 756 M.A., von Freiesleben, P., Tovborg, M., Johansen, K.S., De Maria, L., et al. (2015). Structure
- and boosting activity of a starch-degrading lytic polysaccharide monooxygenase. Nat.
- 758 Commun. *6*, 5961.
- 759 Levasseur, A., Drula, E., Lombard, V., Coutinho, P.M., and Henrissat, B. (2013). Expansion of
- the enzymatic repertoire of the CAZy database to integrate auxiliary redox enzymes.Biotechnol. Biofuels *6*, 41.
- Li, X., Beeson, W.T., Phillips, C.M., Marletta, M.A., and Cate, J.H.D. (2012). Structural basis
 for substrate targeting and catalysis by fungal polysaccharide monooxygenases. Structure
- 764 *20,* 1051–1061.
- Lombard, V., Golaconda Ramulu, H., Drula, E., Coutinho, P.M., and Henrissat, B. (2014). The
 carbohydrate-active enzymes database (CAZy) in 2013. Nucleic Acids Res. *42*, D490-5.
- Mandels, M., and Reese, E.T. (1957). Induction of cellulase in Trichoderma viride as
 influenced by carbon sources and metals. J. Bacteriol. *73*, 269–278.
- Mandels, M., and Reese, E.T. (1960). Induction of cellulase in fungi by cellobiose. J. Bacteriol. *79*, 816–826.
- Mandels, M., Parrish, F.W., and Reese, E.T. (1962). Sophorose as an inducer of cellulase in
 Trichoderma viride. J. Bacteriol. *83*, 400–408.
- Mandels, M., Weber, J., and Parizek, R. (1971). Enhanced cellulase production by a mutant
 of Trichoderma viride. Appl. Microbiol. *21*, 152–154.
- Mandels, M., Hontz, L., Nystrom, J., and Lee R Lynd, I.B. (1974). Enzymatic hydrolysis of
 waste cellulose. Biotechnol. Bioeng. *16*, 1471–1493.
- Merino, S.T., and Cherry, J. (2007). Progress and challenges in enzyme development for
 biomass utilization. Adv. Biochem. Eng. Biotechnol. *108*, 95–120.
- 779 Morgenstern, I., Powlowski, J., and Tsang, A. (2014). Fungal cellulose degradation by
- 780 oxidative enzymes: from dysfunctional GH61 family to powerful lytic polysaccharide
- 781 monooxygenase family. Brief. Funct. Genomics *13*, 471–481.
- Naik, S.N., Goud, V. V, Rout, P.K., and Dalai, A.K. (2010). Production of first and second
 generation biofuels: a comprehensive review. Renew. Sustain. Energy Rev. *14*, 578–597.
- 784 Nakagawa, Y.S., Kudo, M., Loose, J.S.M., Ishikawa, T., Totani, K., Eijsink, V.G.H., and Vaaje-
- 785 Kolstad, G. (2015). A small lytic polysaccharide monooxygenase from Streptomyces griseus

- 786 targeting α and β -chitin. FEBS J. *282*, 1065–1079.
- Neese, F. (2002). Prediction and interpretation of the57Fe isomer shift in Mössbauer spectra
 by density functional theory. Inorganica Chim. Acta *337*, 181–192.
- 789 Neese, F. (2009). Prediction of molecular properties and molecular spectroscopy with
- density functional theory: From fundamental theory to exchange-coupling. Coord. Chem.Rev. 253, 526–563.
- Neese, F. (2017). Quantum Chemistry and EPR Parameters. In EMagRes, (Chichester, UK:
 John Wiley & Sons, Ltd), pp. 1–22.
- Neu, H.C., and Heppel, L.A. (1965). The release of enzymes from Escherichia coli by osmotic
 shock and during the formation of spheroplasts. J. Biol. Chem. *240*, 3685–3692.
- Payne, C.M., Knott, B.C., Mayes, H.B., Hansson, H., Himmel, M.E., Sandgren, M., Ståhlberg,
 J., and Beckham, G.T. (2015). Fungal cellulases. Chem. Rev. *115*, 1308–1448.
- 798 Peisach, J., and Blumberg, W.E. (1974). Structural implications derived from the analysis of
- electron paramagnetic resonance spectra of natural and artificial copper proteins. Arch.Biochem. Biophys. *165*, 691–708.
- 801 Phillips, C.M., Beeson, W.T., Cate, J.H., and Marletta, M.A. (2011). Cellobiose dehydrogenase
- and a copper-dependent polysaccharide monooxygenase potentiate cellulose degradation
- 803 by Neurospora crassa. ACS Chem. Biol. *6*, 1399–1406.
- Quinlan, R.J., Sweeney, M.D., Lo Leggio, L., Otten, H., Poulsen, J.-C.N., Johansen, K.S., Krogh,
- K.B.R.M., Jørgensen, C.I., Tovborg, M., Anthonsen, A., et al. (2011). Insights into the
- 806 oxidative degradation of cellulose by a copper metalloenzyme that exploits biomass
- 807 components. Proc. Natl. Acad. Sci. U. S. A. *108*, 15079–15084.
- Reese, E.T., Siu, R.G.H., and Levinson, H.S. (1950). The biological degradation of soluble
 cellulose derivatives and its relationship to the mechanism of cellulose hydrolysis. J.
- 810 Bacteriol. *59*, 485–497.
- 811 Sabbadin, F., Hemsworth, G.R., Ciano, L., Henrissat, B., Dupree, P., Tryfona, T., Marques,
- 812 R.D.S., Sweeney, S.T., Besser, K., Elias, L., et al. (2018). An ancient family of lytic
- 813 polysaccharide monooxygenases with roles in arthropod development and biomass
- 814 digestion. Nat. Commun. 9, 756.
- 815 Saini, J.K., Saini, R., and Tewari, L. (2015). Lignocellulosic agriculture wastes as biomass
- 816 feedstocks for second-generation bioethanol production: concepts and recent
- 817 developments. 3 Biotech *5*, 337–353.
- Saloheimo, M., Nakari-Setälä, T., Tenkanen, M., and Penttilä, M. (1997). cDNA cloning of a
 Trichoderma reesei cellulase and demonstration of endoglucanase activity by expression in
- 820 yeast. Eur. J. Biochem. 249, 584–591.
- 821 Simmons, T.J., Frandsen, K.E.H., Ciano, L., Tryfona, T., Lenfant, N., Poulsen, J.C., Wilson,
- L.F.L., Tandrup, T., Tovborg, M., Schnorr, K., et al. (2017). Structural and electronic
- 823 determinants of lytic polysaccharide monooxygenase reactivity on polysaccharide
- substrates. Nat. Commun. 8, 1064.
- Solomon, E.I., Heppner, D.E., Johnston, E.M., Ginsbach, J.W., Cirera, J., Qayyum, M., Kieber-
- 826 Emmons, M.T., Kjaergaard, C.H., Hadt, R.G., and Tian, L. (2014). Copper active sites in

- biology. Chem. Rev. 114, 3659–3853.
- Span, E.A., Suess, D.L.M., Deller, M.C., Britt, R.D., and Marletta, M.A. (2017). The Role of the
- Secondary Coordination Sphere in a Fungal Polysaccharide Monooxygenase. ACS Chem. Biol. *12*, 1095–1103.
- Stoll, S., and Schweiger, A. (2006). EasySpin, a comprehensive software package for spectral
 simulation and analysis in EPR. J. Magn. Reson. *178*, 42–55.
- 833 Tanghe, M., Danneels, B., Camattari, A., Glieder, A., Vandenberghe, I., Devreese, B., Stals, I.,
- and Desmet, T. (2015). Recombinant Expression of Trichoderma reesei Cel61A in Pichia
- 835 pastoris: Optimizing Yield and N-terminal Processing. Mol. Biotechnol.
- 836 The Hendrich Metalloprotein Group Spincount.
- 837 http://www.chem.cmu.edu/groups/hendrich/facilities/index.html
- Vaaje-Kolstad, G., Horn, S.J., van Aalten, D.M.F., Synstad, B., and Eijsink, V.G.H. (2005a). The
- 839 non-catalytic chitin-binding protein CBP21 from Serratia marcescens is essential for chitin
- 840 degradation. J. Biol. Chem. 280, 28492–28497.
- 841 Vaaje-Kolstad, G., Houston, D.R., Riemen, A.H.K., Eijsink, V.G.H., and van Aalten, D.M.F.
- (2005b). Crystal structure and binding properties of the Serratia marcescens chitin-binding
 protein CBP21. J. Biol. Chem. *280*, 11313–11319.
- Vaaje-Kolstad, G., Westereng, B., Horn, S.J., Liu, Z., Zhai, H., Sørlie, M., and Eijsink, V.G.H.
 (2010). An oxidative enzyme boosting the enzymatic conversion of recalcitrant
 polysaccharides. Sci. (New York, NY) *330*, 219–222.
- Vaaje-Kolstad, G., Forsberg, Z., Loose, J.S., Bissaro, B., and Eijsink, V.G. (2017). Structural
 diversity of lytic polysaccharide monooxygenases. Curr. Opin. Struct. Biol. 44, 67–76.
- Vu, V. V., and Ngo, S.T. (2018). Copper active site in polysaccharide monooxygenases. Coord.
 Chem. Rev. *368*, 134–157.
- Vu, V. V, Beeson, W.T., Span, E.A., Farquhar, E.R., and Marletta, M.A. (2014a). A family of
 starch-active polysaccharide monooxygenases. Proc. Natl. Acad. Sci. U. S. A. *111*, 13822–
 13827.
- 854 Vu, V. V, Beeson, W.T., Phillips, C.M., Cate, J.H.D., and Marletta, M.A. (2014b). Determinants
- of regioselective hydroxylation in the fungal polysaccharide monooxygenases. J. Am. Chem.
 Soc. *136*, 562–565.
- 857 Wu, M., Beckham, G.T., Larsson, A.M., Ishida, T., Kim, S., Payne, C.M., Himmel, M.E.,
- 858 Crowley, M.F., Horn, S.J., Westereng, B., et al. (2013). Crystal structure and computational
- 859 characterization of the lytic polysaccharide monooxygenase GH61D from the Basidiomycota
- 860 fungus Phanerochaete chrysosporium. J. Biol. Chem. 288, 12828–12839.
- 861