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1 **Effects of hemicelluloses on dehydrogenative polymerization of monolignols with cationic**
2 **cell wall-bound peroxidase**

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9
10 **Abstract**

11 To elucidate the influence of polysaccharides on hardwood lignification, dehydrogenative
12 polymerization of monolignols, coniferyl alcohol (CA) and sinapyl alcohol (SA), was attempted
13 with recombinant cationic cell wall-bound peroxidase (rCWPO-C) and horseradish peroxidase
14 (HRP) in measurement cells of a quartz crystal microbalance with dissipation (QCM-D). Hardwood
15 cellulose nanofibers were anchored; hemicelluloses, xylan, partially acetylated xylan (AcXY),
16 galactoglucomannan, and xyloglucan, and the enzymes were subsequently adsorbed onto the QCM-
17 D sensor surface, enabling fabrication of artificial polysaccharide matrices. The largest amount of
18 rCWPO-C is found to be adsorbed onto AcXY among all the polysaccharides, which affords the
19 largest amount and size of spherical dehydrogenation polymers (DHPs) from both CA and SA. In
20 contrast, no DHP and a small amount of DHPs are formed from SA and CA, respectively, by HRP
21 catalysis in all of the polysaccharide matrices. This study demonstrates important functions of a real
22 tree-derived peroxidase, rCWPO-C, and AcXY for hardwood lignification.

23
24 **Keywords:** Dehydrogenation polymer; sinapyl alcohol; cationic cell wall-bound peroxidase;
25 hemicellulose; partially acetylated xylan.

26 **Abbreviations:** AcXY, partially acetylated WXY; CA, coniferyl alcohol; CNFs, hardwood
27 cellulose nanofibers; DHP, dehydrogenation polymers; 2,6-DMP: 2,6-dimethoxyphenol; GGM,
28 galactoglucomannan; HRP, horseradish peroxidase; QCM-D, a quartz crystal microbalance with
29 dissipation; rCWPO-C, recombinant cationic cell wall-bound peroxidase; SA, sinapyl alcohol;
30 WXY, a water-soluble fraction of commercial beech xylan; XG, xyloglucan.

31

32 **Introduction**

33 Lignin is one of the major wood cell wall components, and its formation, termed lignification, occurs
34 through radical couplings between monolignols, coniferyl alcohol (CA), sinapyl alcohol (SA), *p*-
35 coumaryl alcohol (Ralph, Brunow, & Boerjan, 2007), and/or oligolignols. Cell wall formation in
36 trees starts with the deposition of polysaccharides, cellulose and hemicelluloses on the plasma
37 membrane to form a polysaccharide matrix. Subsequently, lignification occurs in the matrix.
38 Therefore, the matrix has been proposed to affect lignification with respect to the amount formed
39 and the lignin structure. The influence of hemicelluloses on lignification has been elucidated from
40 the spatial arrangement of cell wall components (Donaldson, 1994; Terashima, Yoshida, Hafrén,
41 Fukushima, & Westermark, 2012; Warinowski et al., 2016; Wi, Singh, Lee, & Kim, 2005) and lignin
42 structures in lignin-carbohydrate complexes (LCCs) isolated from wood. However, the latter
43 investigations have suggested contradictory functions of hemicelluloses for lignification. One
44 investigation has reported that xylan promotes the *b-O-4'* linkage as a predominant interunitary
45 linkage of lignin (Giummarella, Zhang, Henriksson, & Lawoko, 2016), but another study has
46 reported that glucomannan facilitates this linkage more than xylan (Du, Gellerstedt, & Li, 2013; Du
47 et al., 2014). These opposite results should be attributed to the fact that the investigations do not
48 directly address the lignification process.

49 To overcome the experimental drawback, dehydrogenative polymerization of CA was attempted
50 as an artificial lignification by endwise polymerization (Saake, Argyropoulos, Beinhoff, & Faix,
51 1996) with horseradish peroxidase (HRP) as a catalyst in an artificial polysaccharide matrix, which

52 was fabricated by depositing commercial beech xylan on bacterial cellulose (BC) (Li et al., 2015).
53 As a result, the xylan yielded larger amounts of dehydrogenation polymers (DHPs) with a higher
54 frequency of β -O-4' linkage than cellulose. By using BC-galactoglucomannan (GGM) and BC-
55 xyloglucan (XG) matrices, GGM was found to inhibit polymerization, while XG contributed to DHP
56 formation with condensed structures such as lignin in primary cell walls (Lyu et al., 2021). However,
57 these investigations still had a drawback in that the HRP used is not a tree enzyme and hardly
58 oxidizes SA (Aoyama et al., 2002; Veitch, 2004). Thus, HRP is unsuitable for the elucidation of
59 lignification in hardwood derived from CA and SA.

60 This study aims to elucidate hardwood lignification from DHP formation in artificial
61 polysaccharide matrices by using a real tree enzyme. It has been reported that native glucuronoxylan
62 in hardwood is partially acetylated with the degree of substitution (DS) ranging from 0.40–0.75
63 (Pawar, Koutaniemi, Tenkanen, & Mellerowicz, 2013; Qaseem & Wu, 2020; Teleman, Lundqvist,
64 Tjerneld, Stålbrand, & Dahlman, 2000; Teleman, Tenkanen, Jacobs, & Dahlman, 2002). Based on
65 this fact, we hypothesize that this acetyl (Ac) group should play an important role in hardwood
66 lignification. However, little work has been conducted to investigate it, although many have been
67 done for unsubstituted glucuronoxylan (Li et al., 2015; Pawar et al., 2017). In this study, a partially
68 acetylated xylan is firstly prepared as a model of native hardwood xylan from a commercially
69 available xylan, because this commercial xylan, extracted from beech with an alkaline aqueous
70 solution, does not contain Ac group. Then, the water-soluble fractions of the commercial xylan and
71 partially acetylated xylan are separately subjected to the fabrication of polysaccharide matrices. If
72 there is a difference in the DHP formation between these polysaccharide matrices, our hypothesis
73 would be verified.

74 As mentioned above, HRP is not proper to verify this, and therefore, a cationic cell wall-bound
75 peroxidase (CWPO-C) is focused on. This enzyme was discovered in poplar (*Populus alba* L.)
76 (Aoyama et al., 2002; Sasaki, Nishida, Tsutsumi, & Kondo, 2004) and is able to oxidize not only
77 CA but also SA and polymeric lignin (Aoyama et al., 2002; Sasaki et al., 2004). However, it is

78 difficult to isolate a large quantity of CWPO-C from hardwood, and thus, recombinant CWPO-C
79 (rCWPO-C) (Shigeto, Itoh, Tsutsumi, & Kondo, 2012) is used in this study.

80 This study adopts a quartz crystal microbalance with dissipation (QCM-D) to analyze the
81 interactions between lignification-involving materials and to elucidate DHP formation (Elschner,
82 Adam, Lesny, Joseph, & Fischer, 2022; Wang, Qian, Roman, Glasser, & Esker, 2013) because
83 QCM-D enables measurements using a small amount of samples. Accordingly, artificial
84 polysaccharide matrices are fabricated on the QCM-D sensor with hardwood-derived cellulose and
85 hemicelluloses (the partially acetylated and unsubstituted xylans, GGM, and XG) prior to the real-
86 time monitoring of DHP formation catalyzed by rCWPO-C and HRP as a reference.

87

88 **2. Materials and Methods**

89 ***2.1. Water-soluble fraction of commercial beech xylan (WXY) and its partially acetylated xylan*** 90 ***(AcXY)***

91 Commercial beech xylan (SERVA Electrophoresis GmbH, Heidelberg, Germany) was dissolved in
92 water at room temperature (25 °C) for 1 day, and the suspension was centrifuged to obtain a water-
93 soluble fraction as the supernatant. The water-soluble fraction was delignified with sodium chlorite
94 according to a modified Wise method and freeze-dried to yield WXY. The sugar constituents of
95 WXY were 76.8% xylose, 7.3% glucose, 12.8% glucuronic acid, 0.3% Klason lignin, and 0.7% acid
96 soluble lignin (Lyu et al., 2021).

97 WXY (5 g) was suspended in *N,N*-dimethylacetamide (50 mL) with stirring at 120 °C for 16
98 h, and then LiCl (3.76 g) was added to the suspension. The mixture was stirred at 80 °C for
99 approximately 16 h until WXY was completely dissolved. Methane sulfonic acid (2.5 mL) and
100 acetic anhydride (32.5 mL) were added to the solution, and the mixture was stirred at 80 °C for 16
101 h. The resultant solution was slowly poured into distilled water. The precipitate was collected by
102 filtration, rinsed with distilled water, and then dried *in vacuo* overnight to yield fully acetylated
103 WXY with a yield of 78% based on the theoretical output value.

104 The fully acetylated WXY (1 g) was dissolved in chloroform (100 mL) with stirring at room
105 temperature. Sodium methoxide (2.4 g) was then added to the solution and stirred at room
106 temperature for 16 h. The solution was neutralized with acetic acid and then dialyzed against
107 distilled water for 3 days. The water was changed every 4 h during the daytime. The dialyzed
108 suspension was evaporated to the solid and dried *in vacuo* at 50 °C for 2 days. The resultant solid
109 (0.22 g) was suspended again in 50 mL of water for 16 h, and the supernatant was collected by
110 centrifugation. It was lyophilized to give the water-soluble fraction (AcXY). The AcXY formed
111 65% of the total weight of the partially deacetylated derivative of fully acetylated WXY.

112 The degree of substitution (DS) of fully acetylated WXY and AcXY was determined by a
113 titration method (Taira et al., 2020) and FT-IR measurements (see Fig. S1 for details of the DS
114 determination). ¹H-/¹³C-NMR spectra for WXY and AcXY were measured in deuterated
115 dimethylsulfoxide (DMSO-*d*₆) or deuterium oxide using a 500 MHz spectrometer (Bruker
116 AVANCE Neo, Billerica, United States). The chemical shift of the deuterated solvent was used as
117 a reference ($\delta = 2.49$ ppm) for ¹H-NMR. The molar mass distributions of WXY and AcXY were
118 measured by using a size exclusion chromatography (SEC) system (see Supporting Information for
119 the detail conditions).

120

121 **2.2. Other polysaccharides**

122 A hydrogel of cellulose nanofibers (CNFs) derived from hardwood kraft pulp was provided by
123 Hokuetsu Corporation (Tokyo, Japan). The gel was diluted to approximately 1/5 consistency and
124 dialyzed against distilled water for 3 days. The dialyzed CNF suspension was concentrated to 0.74
125 g/L by evaporation. The composition of the CNFs was 72.5% glucose, 16.8% xylose, and 0.5%
126 Klason lignin (Lyu et al., 2021).

127 Crude glucomannan was extracted and purified according to a previous report (Lyu et al., 2021).
128 The obtained glucomannan (GGM) was composed of 13.7% glucose, 39.4% galactose, 41.1%
129 mannose, 0.1% Klason lignin, and 0.9% acid soluble lignin.

130 Xyloglucan (XG) extracted from tamarind seed was purchased from Megazyme (Ireland) and
131 used as received. XG was composed of 47.9% glucose, 24.9% xylose, 1.8% arabinose, 18.5%
132 galactose, 0.9% Klason lignin, and 1.2% acid-soluble lignin (Lyu et al., 2021).

133

134 **2.3. Enzyme activity measurement**

135 rCWPO-C preparation and purification procedures were conducted using the same method as
136 reported previously (Shigeto et al., 2012). HRP was purchased from FUJIFILM Wako Pure
137 Chemical Industries (Osaka, Japan) and used as received. The enzyme activity of rCWPO-C and
138 HRP (Shigeto et al., 2012) was measured at 25 °C on the basis of the oxidation of guaiacol and 2,6-
139 dimethoxyphenol (2,6-DMP) in Tris-HCl (50 mM, pH 7.5) buffered solution and phosphate
140 buffered saline (PBS, pH 6.1, 0.01 M phosphate containing 0.8 w/v% NaCl and 0.02 w/v% KCl),
141 respectively. The absorbance of the enzymatic oxidation products (i.e., tetraguaiacol and
142 coerulignone from guaiacol and 2,6-DMP, respectively) were monitored at a wavelength of 470 nm
143 and 469 nm, respectively, using a UV-vis absorption spectrophotometer (U-3310, HITACHI High-
144 Tech Science, Kyoto, Japan).

145 Enzyme activity (A U/mg) based on the generation of the product within 1 min after H₂O₂
146 addition was calculated by using the following equation (Shigeto et al., 2012; George, 1953;
147 Wariishi et al., 1992):

$$148 \quad A = \frac{(\Delta E_1 - \Delta E_2) \times v}{\varepsilon \times c \times v_0}$$

149 where ΔE_1 is the rate of increase in absorbance (per min) of the reaction solution, ΔE_2 is the rate of
150 increase in the absorbance (per min) of the blank solution, v is the total volume of the reaction
151 mixture, c is the enzyme concentration, and v_0 is the volume of the added enzyme solution. ε is the
152 absorptivity of the oxidation product and is equal to 26.6 L/mmol/cm for tetraguaiacol (George,
153 1953) and 49.6 L/mmol/cm for coerulignone (Wariishi, Valli, & Gold, 1992).

154

155 **2.4. QCM-D measurements**

156 A CNF-coated QCM-D sensor was prepared according to our previous report (Lyu et al., 2021). The
157 frequency change (Δf) of the sensor before and after CNF deposition was measured using a QCM-
158 D (Q-sense AB, Biolin Scientific, Västra Frölunda, Sweden). The dried amount of CNFs deposited
159 onto the sensor was calculated by using the Sauerbrey equation (Sauerbrey, 1959):

160
$$\Delta m = C(\Delta f/n)$$

161 where Δm (ng/cm²) is the weight change, C is the mass sensitivity constant of the sensor (-17.7
162 ng/cm²/Hz), and n is the overtone number ($n = 5$ in this study). For a Δm of less than 4000 ng/cm²,
163 the CNF coating was further repeated until Δm exceeded 4000 ng/cm², at which point the CNFs
164 theoretically covered the whole surface of the sensor as a monolayer (Lyu et al., 2021; Sugiyama,
165 Vuong, & Chanzy, 1991).

166 The dried CNF-coated sensor was set in a measurement cell and conditioned with a flow of
167 Mill-Q water. Then, a hemicellulose solution at 1 mg/mL was introduced into the cell for 2 h. Mill-
168 Q water was subsequently introduced for 15 min to remove unbound hemicellulose. All QCM-D
169 measurements were performed at 25 °C, and the flow rate for all solutions was controlled to be 50
170 μ L/min by using a peristaltic pump (ISM795C, Iamatec™, Fisher Scientific, Sweden). The changes
171 in the Δf and dissipation factor (ΔD) of the sensor were monitored during measurements. The
172 dissipation factor (D) is defined by the following equation (Dixon, 2008):

173
$$D = \frac{E_{diss}}{2\pi E_{stored}}$$

174 where E_{diss} is the dissipated energy and E_{stored} is the stored energy during one oscillation.

175 After the hemicellulose adsorption process, Mill-Q water in the cell was exchanged with Tris-
176 HCl or PBS buffered solutions. Then, an enzyme buffered solution (rCWPO-C in a Tris-HCl
177 buffered solution or HRP in a PBS buffered solution) at a concentration of 5 U/mL for guaiacol was
178 introduced at 50 μ L/min for 30 min. After the adsorption experiment, the sensor surface was rinsed

179 with the buffered solution for 15 min. This enzyme adsorption was also conducted on the water-
180 swelled CNF-coated sensor without hemicelluloses.

181 CA and SA were synthesized according to the same method as reported previously (Quideau
182 & Ralph, 1992). After enzyme adsorption, a monolignol (2.7 mM) buffered solution containing
183 H₂O₂ (2.7 mM) was introduced into the cell for 1 h. The sensor surface was successively rinsed with
184 the corresponding buffer solution for 15 min and Milli-Q water for 30 min and subjected to the
185 following observation. As a reference, monolignol adsorption was conducted for a CNF-coated
186 sensor without H₂O₂ after enzyme adsorption. Each measurement was repeated at least two times
187 until the same trend profiles were obtained.

188

189 ***2.5. Atomic force microscope (AFM) imaging***

190 The sensors used for DHP formation were dried under a N₂ flow and observed using AFM (SPA-
191 400 AFM, HITACHI High-Tech Science, Kyoto, Japan). Images of the sensor surfaces were taken
192 in tapping mode using a SI-DF-40 cantilever (HITACHI High-Tech Science, Kyoto, Japan) and
193 expressed as shape images. The mean surface roughness (Ra) over a scan area of 2.5 × 2.5 μm² was
194 estimated by using Gwyddion software.

195

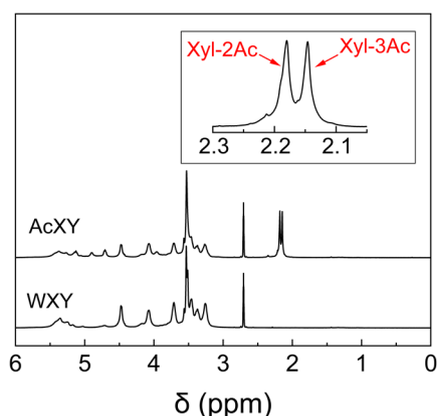
196 **3. Results and Discussion**

197 ***3.1. Preparation and characterization of AcXY***

198 WXY was fully acetylated under homogeneous conditions and then deacetylated using sodium
199 methoxide to yield partially acetylated xylan with a DS of 0.50 (see Fig. S1 for the DS
200 determination), which is close to the DS of native xylan in hardwood (Pawar et al., 2013; Teleman
201 et al., 2000; Teleman et al., 2002). Its water-soluble fraction was collected to yield AcXY with a
202 similar DS to that obtained before fractionation. The size exclusion chromatograms of WXY and
203 AcXY are shown in Fig. S2 and Table S1. As WXY shows a bimodal chromatogram (Lyu et al.,

204 2021), AcXY also shows a similar chromatogram, but the peak in the low molar mass range was
205 slightly shifted toward the lower molar mass region than that of WXY.

206 The distribution of acetyl (Ac) groups in AcXY was analyzed by using ^1H - and ^{13}C -NMR (Fig.
207 1 and Fig. S3, respectively). The ^1H -NMR spectrum shows chemical shifts at 2.17 and 2.15 ppm,
208 which are assigned to the methyl protons of the Ac group at the C-2 position (Xyl-2Ac) and C-3
209 position (Xyl-3Ac), respectively (Zhong, Cui, & Ye, 2017), and their integral ratio is approximately
210 1:1. No chemical shift for the Ac groups at both the C-2 and C-3 positions (Xyl-2,3Ac) was observed.
211 These results suggest that the Ac group is located at either the C-2 or C-3 position in each
212 anhydroxylose unit of AcXY. Thus, a partially acetylated xylan (i.e., AcXY) was successfully
213 prepared as the model of native hardwood xylan for the following investigations.



214
215 **Fig. 1.** ^1H -NMR spectra for AcXY and WXY in $\text{DMSO-}d_6$. An expanded spectrum of AcXY shows
216 the assignments for the methyl protons of the Ac groups.

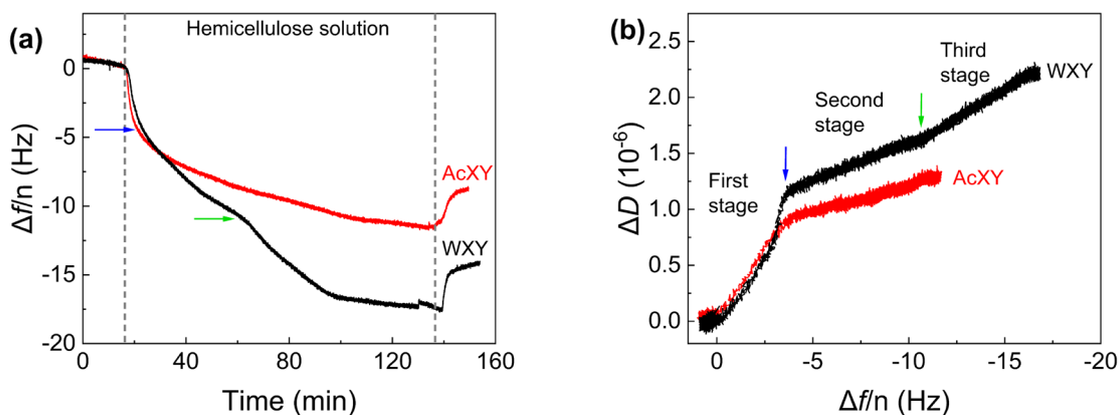
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218

219 **3.2 Hemicellulose adsorption on CNFs**

220 In our previous report (Lyu et al., 2021), hemicellulose adsorption on cellulose was monitored using
221 a QCM-D, and the adsorption amount followed the order of XG > GGM > WXY. In this study, the
222 AcXY adsorption on CNFs was further investigated by using the QCM-D. It should be noted that
223 the CNFs used in this study were derived from hardwood kraft pulp, thus, originally containing a
224 small amount of xylan (*ca.* 17%).

225 Figure 2 shows two types of QCM-D profiles for AcXY: $\Delta f/n$ versus elution time (Fig. 2a) and
226 ΔD versus $\Delta f/n$ (ΔD - $\Delta f/n$ plot, Fig. 2b), where the profiles for WXY are depicted as a reference
227 (Lyu et al., 2021). $\Delta f/n$ exhibits the adsorption amount of AcXY and WXY on the CNF-coated
228 sensor, and a smaller $\Delta f/n$ corresponds to a larger adsorbed amount. The ΔD - $\Delta f/n$ plot reflects the
229 viscoelastic change in the adsorbed layer on the sensor; a steep curve indicates that the surface layer
230 becomes viscous or soft, whereas a gradually increasing curve indicates that the surface becomes
231 elastic or rigid (Littunen, Mai-Gisoni, Seppälä, & Master, 2017).



232

233 **Fig. 2.** QCM-D profiles for frequency change ($\Delta f/n$) as a function of elution time (a) and dissipation
234 change (ΔD) as a function of $\Delta f/n$ (b) during adsorption of AcXY and WXY on the CNF-coated
235 sensor. The blue and green arrows show the border for each stage.

236

237 As shown in Fig. 2b, the ΔD - $\Delta f/n$ profile for AcXY can be divided into two stages: the first
238 stage ranges from 0 to -4 Hz, and the second stage ranges from -4 to -11 Hz. On the other hand, the

239 profile for WXY can be divided into three stages: the first and second stages are located in the same
240 ranges as those for AcXY, but the third stage appears from -11 to -17 Hz.

241 In the first stage, the $\Delta f/n$ profiles for AcXY and WXY (Fig. 2a) are almost identical, with $\Delta f/n$
242 showing a rapid decrease, indicating the rapid adsorption of AcXY and WXY onto the CNF surface.
243 Concomitantly, ΔD also rapidly increases, suggesting that the sensor surfaces become soft due to
244 the adsorption of hydrated AcXY and WXY. In the second stage, the $\Delta f/n$ for AcXY and WXY
245 gradually decreases, and the slope of the ΔD - $\Delta f/n$ plot becomes less steep than that in the first stage,
246 indicating that the surface becomes relatively rigid. This phenomenon can be attributed to the release
247 of bound water from hydrated AcXY and WXY to achieve further adsorption of the hemicelluloses
248 on CNFs (Farooq et al., 2020). Furthermore, the rate of decrease of $\Delta f/n$ for WXY in the second
249 stage is faster than that for AcXY. Such rapid adsorption of WXY can be promoted via much
250 hydrogen bonding interactions with CNFs compared with AcXY.

251 At the third stage of WXY, $\Delta f/n$ shows a rapid decrease followed by a gradually decrease. The
252 ΔD - $\Delta f/n$ plot for WXY also changes to a steeper slope than that in the second stage. These results
253 suggest that the formed layer becomes soft through a different mode of adsorption. It has been
254 reported that xylan without substituents tends to self-associate (Eronen, Österberg, Heikkinen,
255 Tenkanen, & Laine, 2011; Kabel, van den Borne, Vincken, Voragen, & Schols, 2007). Thus, the
256 softer layer formed in the third stage can be attributed to the self-association of WXY, resulting in
257 a maximum adsorption on CNFs. In contrast, the total amount of AcXY adsorbed on CNFs is smaller
258 than that for WXY because the Ac group in AcXY can inhibit self-association.

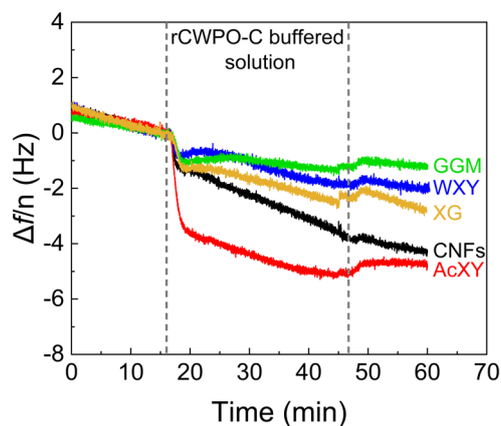
259

260 ***3.3. Enzyme activity and adsorption on polysaccharides***

261 The oxidation activities of rCWPO-C and HRP were determined with guaiacol and 2,6-DMP as
262 model substrates for CA and SA, respectively. rCWPO-C shows a high activity of 432 U/mg for
263 2,6-DMP, which is approximately three times higher than the activity of 148 U/mg for guaiacol. On
264 the other hand, HRP shows an extremely low activity of 177 U/mg for 2,6-DMP, which is only 0.35

265 times the activity of 503 U/mg for guaiacol. Therefore, rCWPO-C is confirmed to have a high ability
266 to oxidize syringyl nuclei (Aoyama et al., 2002; Shigeto et al., 2012).

267 The enzyme solution at 5 U/mL for guaiacol oxidation was flowed into the QCM-D sensor coated
268 with polysaccharides, and the time course for the enzyme adsorption was monitored by using QCM-
269 D. In this measurement, the CNFs were used as a reference material to evaluate the effect of each
270 hemicellulose on rCWPO-C adsorption. rCWPO-C shows the largest adsorption amount on AcXY
271 among the polysaccharides used in this study (Fig. 3), whereas the adsorption amount of HRP on
272 AcXY is less than that on CNFs (Fig. S2). rCWPO-C is more hydrophobic than HRP (Aoyama et
273 al., 2002), and the acetylation of xylan increases the hydrophobicity (Busse-Wicher et al., 2014).
274 Therefore, it is considered that rCWPO-C adsorption on AcXY can be enhanced via hydrophobic
275 interactions.



276
277 **Fig. 3.** $\Delta f/n$ profiles as a function of elution time for rCWPO-C adsorption on polysaccharide-coated
278 sensors.

279

280 3.4. DHP formation in artificial wood cell wall polysaccharide matrices

281 DHP formation from CA was attempted in HRP-adsorbed QCM-D sensors coated with CNFs by
282 flowing CA solution with/without H_2O_2 (Wang et al., 2013). As shown in Figs. 4a and -e, there is a
283 difference of 7 Hz between the maximum absolute values for $\Delta f/n$ ($|\Delta f/n|$) in the presence and
284 absence of H_2O_2 . After the QCM-D measurement with H_2O_2 , the AFM image of the sensor surface

285 (Fig. 5b) shows the presence of small particles on the fibrous CNFs, indicating DHP formation from
286 CA on the CNF-coated sensor by HRP catalysis. Furthermore, for the cases with H₂O₂, all the
287 maximum $|\Delta f/n|$ values for the other polysaccharide-coated sensors are higher than that for the CNF-
288 coated sensor without H₂O₂, suggesting that more DHPs are formed. In particular, the largest $|\Delta f/n|$
289 value is recorded for the AcXY-coated sensor, where DHP particles are also observed by AFM (Fig.
290 5c). These results imply that AcXY facilitates DHP formation from CA. On the other hand, when
291 the SA solution is flowed (Figs. 4b,f), all $|\Delta f/n|$ values are almost constant regardless of the presence
292 or absence of H₂O₂ and the type of polysaccharides used for coating the sensors. Considering the
293 decrease in $\Delta f/n$ at the beginning of the SA flow, DHP is not formed from SA due to the low activity
294 of HRP for syringyl nuclei (Wang et al., 2013), and only SA adsorption proceeds on the sensor
295 surface.

296 For CA flow into the rCWPO-C-adsorbed sensor coated with CNFs (Figs. 4c,e), there is a
297 larger difference of 58 Hz between the maximum $|\Delta f/n|$ in the presence and absence of H₂O₂ than
298 that for HRP-adsorbed sensors, suggesting the superior DHP formation of rCWPO-C from CA
299 compared to that of HRP. The AFM images of the resultant sensor surfaces (Figs. 5d,e) clearly show
300 the DHP particles formed and adsorbed along the fibers. The highest $|\Delta f/n|$ in the presence of H₂O₂
301 is recorded on the AcXY-coated surface among all polysaccharides. This can be explained by not
302 only the higher adsorption amount of rCWPO-C on AcXY than on other polysaccharides but also
303 the hydrophobicity of AcXY.

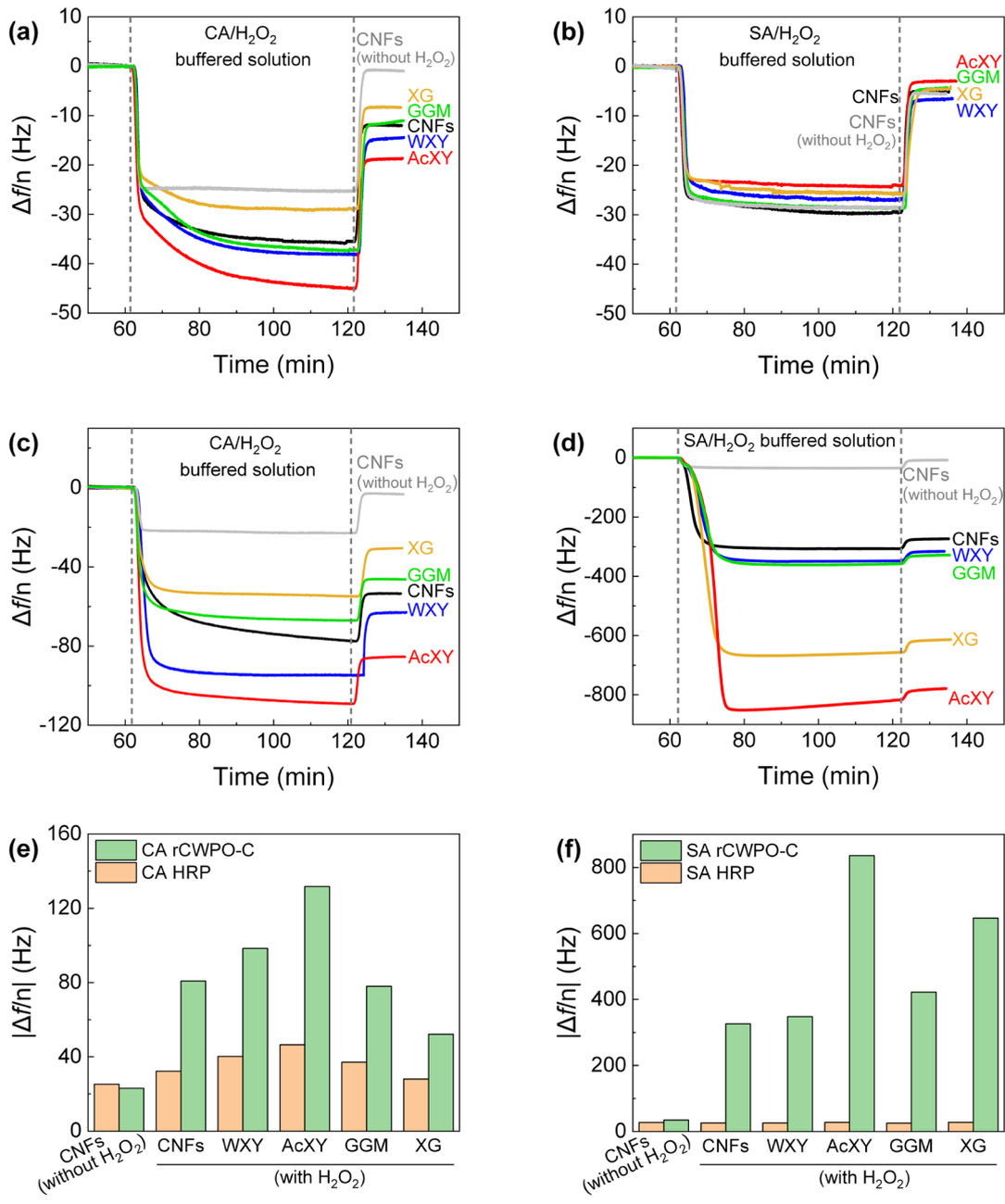
304 When the SA/H₂O₂ solution is flowed (Figs. 4d,f), the maximum $|\Delta f/n|$ values in all
305 polysaccharide matrices are 3–12 times larger than those obtained from the CA/H₂O₂ solution in
306 the corresponding matrices because of the superior oxidation activity of rCWPO-C for syringyl
307 nuclei compared to that for guaiacyl nuclei (Aoyama et al., 2002; Shigeto et al., 2012). In particular,
308 the AcXY-coated sensor exhibits the highest $|\Delta f/n|$, which is six times higher than that for CA. The
309 AFM image of the sensor (Fig. 5g) shows many particles with larger diameters of several hundred
310 nanometers, and the particles completely cover the sensor surfaces. The formation of “sphere”

311 nanoparticles of lignin and DHP from monolignols has been also confirmed in previous research
312 (Terashima et al., 2012; Chao et al., 2013). Although the mechanism to form globular lignin in wood
313 cell walls has not been clarified yet, Terashima et al. assumed that it would be a micellar aggregate
314 of oligolignols folded at the β -O-4 bond with their phenolic ends on the outer part of the aggregate.
315 The formation of globular lignin in the early stages of lignification is a very important phenomenon,
316 and the detailed mechanism is a subject for our further investigation.

317 As shown in Figs. 5f, g and Fig. S3 of AFM images of the sensors coated with other
318 polysaccharides, this study clearly demonstrates that rCWPO-C catalyzes the formation of a large
319 amount and size of spherical DHPs from SA in all polysaccharide matrices. Furthermore, this
320 tendency is the most significant for AcXY among all polysaccharides. These results suggest the Ac
321 group in xylan plays an important role in hardwood lignification, verifying our hypothesis.

322 For other polysaccharides, XG was found to promote DHP formation from SA catalyzed by
323 rCWPO-C followed by AcXY, as shown in Fig. 4d, although WXY and GGM are comparable to
324 CNFs. Thus, XG should be an important polysaccharide for lignification in the primary cell walls
325 of hardwood. On the other hand, WXY promotes DHP formation from CA catalyzed by rCWPO-C
326 and HRP compared with XG, GGM, and CNFs, implying that the xylan backbone itself may also
327 be related to lignification.

328 DHP formation from CA in unsubstituted polysaccharide matrices has been examined in our
329 previous studies using HRP, which revealed significant effects of WXY and XG on the amount and
330 structure of the generated DHPs (Li et al., 2015, Lyu et al., 2021). However, this is the first study to
331 demonstrate the significance of Ac groups on present in hardwood xylan through DHP formation
332 from both CA and SA catalyzed by a real tree enzyme, rCWPO-C. As galactoglucomannan in
333 softwood also contains a small amount of Ac group (Willför, Sundberg, Tenkanen, & Holmbom,
334 2008), its function will be further investigated to understand softwood lignification.



335

336 **Fig. 4.** $\Delta f/n$ profiles as a function of elution time during the flow of CA (a) and SA (b) buffered

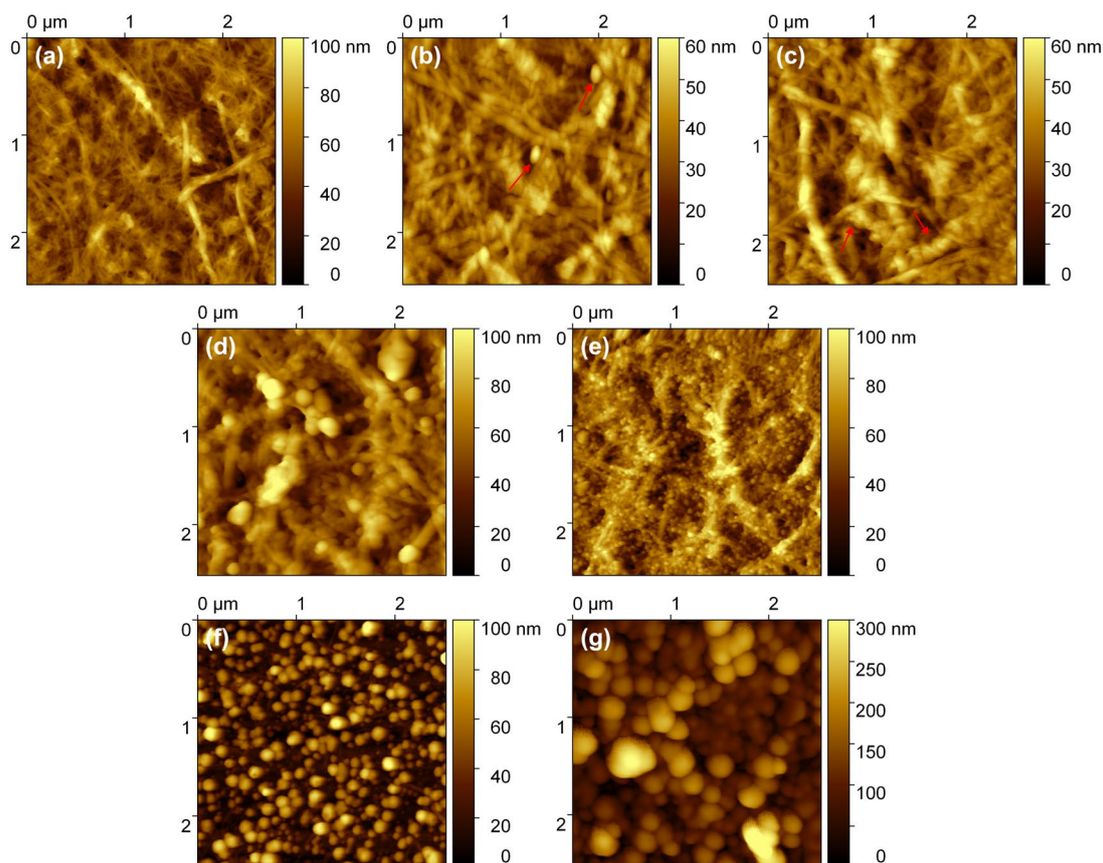
337 solutions on a HRP-adsorbed sensor and during the flow of CA (c) and SA (d) buffered solutions

338 on a rCWPO-C-adsorbed sensor. $|\Delta f/n|$ values at the smallest $\Delta f/n$ during flowing CA (e) and SA (f).

339 The sensors were coated with polysaccharides, CNFs, WXY, AcXY, GGM and XG, and the

340 measurements were performed with/without H₂O₂.

341



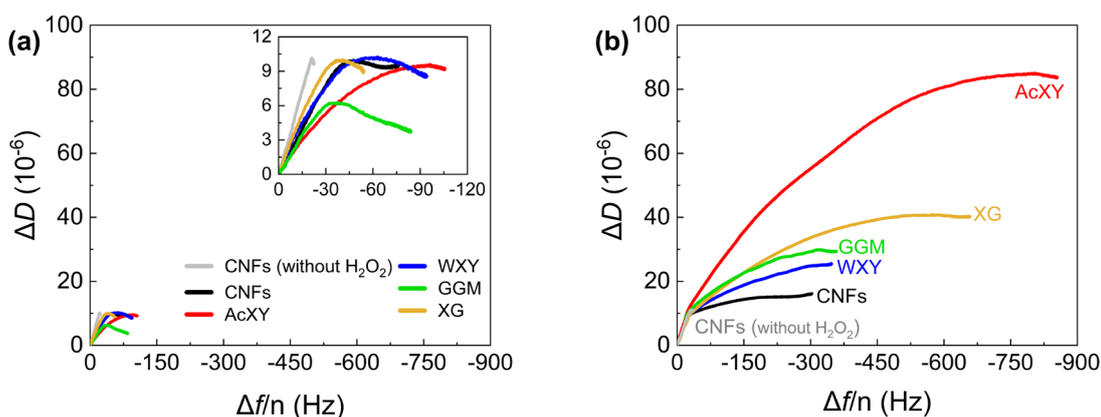
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343 **Fig. 5.** AFM images of QCM-D sensor surfaces. (a) CNF-coated sensor. (b) CNF- and (c) AcXY-
 344 coated HRP-absorbed sensors after flowing CA/H₂O₂, where the formed DHP particles are indicated
 345 by arrows. (d) CNF- and (e) AcXY-coated rCWPO-C-absorbed sensors after flowing CA/H₂O₂; (f)
 346 CNF- and (g) AcXY-coated rCWPO-C-absorbed sensors after flowing SA/H₂O₂.

347

348 Figure 6 shows the ΔD - $\Delta f/n$ profiles obtained during DHP formation by rCWPO-C catalysis.
 349 For CA flow (Fig. 6a), the profile without H₂O₂ is steeper than other profiles with H₂O₂, suggesting
 350 that the sensor surfaces become rigid upon DHP formation with H₂O₂. This can be attributed to
 351 expulsion of bound water on the polysaccharide matrix, induced by the formation of hydrophobic
 352 DHPs. In particular, the sensor surface with AcXY is more rigid than that with other polysaccharides
 353 because the amount of DHP formed in the AcXY matrix is the largest. On the other hand, for SA
 354 flow (Fig. 6b), the slopes for all ΔD - $\Delta f/n$ profiles are almost identical at the initial stage from 0 to -
 355 25 Hz of $\Delta f/n$, and then, the slopes become less steep at the latter stage. These profiles suggest that

356 SA adsorption occurs first, and then, the surface becomes rigid as DHP formation proceeds. At the
 357 final stage, the surfaces become more rigid, possibly because the DHP densely accumulates onto
 358 the sensor. This tendency is more pronounced in the AcXY-coated sensor than in other
 359 polysaccharide-coated sensors.



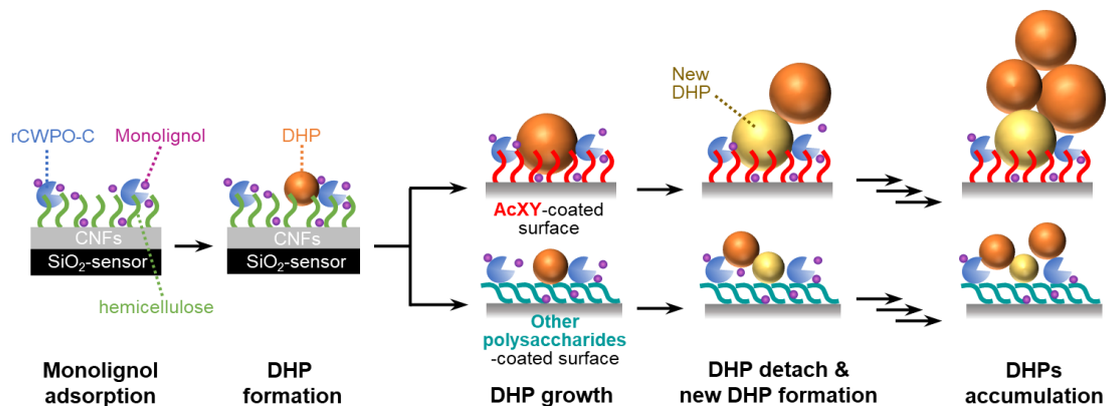
360

361 **Fig. 6.** ΔD vs. $\Delta f/n$ plots for flowing CA (a) and SA (b) buffered solutions with/without H_2O_2 on
 362 rCWPO-C-adsorbed sensors coated with polysaccharides.

363

364 Based on the above experimental findings, a DHP formation mechanism in the peroxidase-
 365 polysaccharide matrix (Fig. 7) is proposed as follows: First, considering that rCWPO-C is adsorbed
 366 onto the sensor surface coated with polysaccharides, DHP formation should occur on only the
 367 surface. After the formed DHP grows into a spherical particle with a diameter of hundreds of
 368 nanometers, it can become detached from the matrix surface. Concomitantly, a new formation of
 369 another DHP particle begins at this site. These formed particles associate with each other through
 370 several interactions, such as hydrophobic and hydrogen bonding interactions (Uraki et al., 2012).
 371 This process occurs successively. Finally, the DHP particles accumulate, as shown in the AFM
 372 images (Fig. 5). When the matrix consists of AcXY, it adsorbs a large amount of rCWPO-C through
 373 hydrophobic interactions, leading to the formation of a large quantity of DHPs. Furthermore, AcXY
 374 also holds the DHP particles through hydrophobic interactions for a longer time than other
 375 polysaccharides, which enables the DHP particles to grow to a larger size. Consequently, both the

376 quantity and particle size of the formed DHPs are the largest on AcXY among those on all
377 polysaccharides.



379 **Fig. 7.** Proposed mechanism for DHP formation in rCWPO-C-adsorbed polysaccharide matrices.

380

381 **4. Conclusion**

382 The components in wood cell walls are well known, but the interactions among them and their
383 cooperative functions on lignification have not been clarified. In this study, dehydrogenative
384 polymerization of monolignols catalyzed by a poplar-derived peroxidase, rCWPO-C, in artificial
385 polysaccharide matrices is real-time monitored by using QCM-D. It is found that DHP formation is
386 more facilitated when the matrix contains AcXY than WXY. This result supports our hypothesis
387 that Ac group in native glucuronoxylan would play an important role in hardwood lignification. In
388 addition, a mechanism for the lignification process in hardwood is proposed based on the large
389 amount of rCWPO-C adsorption on AcXY and the development of DHP particles assisted by AcXY.
390 These investigations would be expanded to elucidate the lignification process in softwood because
391 it also contains partially acetylated galactoglucomannan.

392

393 **CRedit authorship contribution statement**

394 **Yan Lyu:** Investigation, Formal analysis, Data curation, Writing – original draft. **Shiori Suzuki:**

395 Supervision, Writing – original draft & review & editing. **Hiroki Nagano:** Investigation, Formal

396 analysis, Resource (AcXY). **Kengo Shigetomi:** Supervision, Writing – review & editing. **Tutaka**
397 **Tamai:** Resource (GGM & XG). **Yuji Tsutsumi:** Resource (rCWPO-C). **Yasumitsu Uraki:**
398 Conceptualization, Supervision, Project administration, Funding acquisition, Writing – review &
399 editing.

400

401 **Conflict of interest disclosure**

402 The authors declare no competing interest.

403

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407

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