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# Tandem metalloenzymes gate plant cell entry by pathogenic fungi

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- 24
- 25

#### Abstract 26

Global food security is endangered by fungal phytopathogens causing devastating crop 27 production losses. Many of these pathogens use specialized appressoria cells to puncture plant 28 cuticles. Here, we unveil a pair of alcohol oxidase-peroxidase enzymes to be essential for 29 pathogenicity. Using *Colletotrichum orbiculare*, we show that the enzyme pair is co-secreted 30 by the fungus early during plant penetration, and that single and double mutants have impaired 31 penetration ability. Molecular modeling, biochemical and biophysical approaches revealed a 32 fine-tuned interplay between these metalloenzymes, which oxidize plant cuticular long-chain 33 34 alcohols into aldehydes. We show that the enzyme pair is involved in transcriptional regulation of genes necessary for host penetration. The identification of these infection-specific 35 metalloenzymes opens new avenues on the role of wax-derived compounds and the design of 36

- 37 oxidase-specific inhibitors for crop protection.
- 38

#### Short title: 39

Fungal metalloenzymes drive plant invasion. 40

#### **One Sentence Summary:** 41

- Fungal phytopathogens secrete tandem metalloenzymes that catalyze cuticle oxidation and 42
- drive plant cell entry. 43

# 44 Introduction

Fungal phytopathogens represent a serious threat to plant health (1) and global food 45 security (2). *Colletotrichum* and *Magnaporthe* species rank among the top 10 most devastating 46 fungal phytopathogens in the world and reduce crop yield by up to 30% (3). Despite being 47 separated by ca. 300 million years of evolution (4, 5), these fungi share remarkable similarities 48 in their infection strategy, notably the formation of a specialized cell dedicated to host 49 penetration called an appressorium (6-8) (Fig. 1A). This dome-shaped, darkly melanized cell 50 generates a high internal turgor and directs this mechanical pressure onto a needle-like 51 52 penetration peg (9-11), which emerges from a 2-500 nm pore at the appressorial base to puncture the plant's outer defensive barriers, namely the cuticle and epidermal cell wall. 53 Despite major advances in our understanding of the cellular processes preceding (6, 7, 11-14)54 and following (15, 16) plant cell entry, the (bio)chemical reactions occurring at this nanoscale 55 plant-fungus interface and their role in host penetration are not fully elucidated. 56

Copper radical oxidases (CROs) are enzymes with diverse substrate specificities, and 57 have been extensively studied since the late 1960s (17). Today, CROs include galactose 6-58 oxidases (GalOx) (17), glyoxal oxidases (18), and broad specificity primary alcohol oxidases 59 (AlcOx) (19, 20). Despite the detailed knowledge available on the enzymology and structure 60 of CROs, their biological function remains largely unknown. It has only been proposed that 61 glyoxal oxidases play a role in lignin degradation by fungal saprotrophs (18). During the course 62 of our previous work (19), we noted that genes encoding secreted AlcOx orthologs are 63 particularly widespread among phytopathogenic ascomycete fungi and are absent in plants. 64 Given that long-chain primary alcohols are components of the waxy cuticle of aerial plant 65 surfaces (21), we hypothesized that AlcOx could play a role in fungal pathogenesis. 66

In this study, we used a combination of 'omics analyses to unveil the pairing of AlcOx with a redox partner, namely heme-peroxidase, in some fungal plant pathogens of high agricultural importance. We used wet enzymology and molecular modeling to demonstrate and characterize the interplay between these metalloenzymes. Reverse genetics, live-cell imaging and fungal transcriptomics allowed us to probe the *in vivo* function of the enzyme pair, providing new molecular insights into the host penetration cascade.

### 73 **Results**

74

### 75 Discovery of the tandem Perox-AlcOx

studying the enzymology of *Colletotrichum* AlcOx enzymes for 76 While biotechnological applications (19, 22), we noticed the presence of a gene encoding a putative 77 peroxidase located adjacent to an AlcOx-encoding gene (Fig. 1B). To strengthen this initial 78 observation, we searched for *alcox* orthologs in 30 sequenced *Colletotrichum* genomes, which 79 revealed the near-ubiquitous presence of a putative peroxidase (hereafter called "Tandem 80 Peroxidase"). Interestingly, the perox and alcox genes were found in a head-to-head 81 arrangement (fig. S1), suggesting the presence of a bidirectional promoter for tight co-82 expression of the genes. Interestingly, these Tandem Peroxidases are never found in 83 combination with other types of CROs (fig. S1C) and both proteins encoded by the *perox-alcox* 84 pair are predicted to be secreted (Table S1). These observations aroused our interest because 85

it is known that CROs require activation by horseradish peroxidase (HRP) for maximum
activity *in vitro* (23, 24).

Our phylogenetic analysis of the peroxidase-catalase superfamily showed that the 88 Tandem Peroxidases cluster together in a sister clade within the under-explored ascomycete 89 Class II peroxidases (25) (fig. S2A). Furthermore, Tandem Peroxidases form a distinct clade 90 amongst the 333 Class II peroxidases found in *Colletotrichum* species (fig. S2B), suggestive 91 of neofunctionalization. A broad search for the co-occurrence of *perox* and *alcox* orthologs 92 93 across fungal genomes revealed that the pair is also present in *Magnaporthe* species, including the infamous causal agent of rice blast, *Magnaporthe oryzae* (syn. *Pyricularia oryzae*) (**Table** 94 S2). Mapping the occurrence of Perox-AlcOx protein pairs and their corresponding genomic 95 neighborhoods onto a phylogeny of representative pathogenic ascomycetes (Fig. 1B, fig. S3) 96 allowed us to conclude that the pair is present in most *Colletotrichum* species complexes for 97 which genome sequences are available and in *Magnaporthe spp.*, and that the head-to-head 98 organization of the pair is conserved in all these fungi, suggesting there is selection pressure to 99 retain the pairing and that it is has a critical role in the biology of these pathogens (Fig. 1B). 100

To further test the hypothesis of a functional linkage of the *perox* and *alcox* gene products, we parsed transcriptomic data available for those fungal species harboring the pair. Remarkably, this analysis revealed that both genes are always tightly co-transcribed at the appressorium stage in various pathosystems involving *Colletotrichum* species attacking maize, fruits, and model plants (26, 27), and in *M. oryzae* attacking rice and barley (28, 29) (**Fig. 1C and D**). In each case, the transcript levels are relatively low and detected within a narrow time window, which may explain why these genes were overlooked in previous studies.



Fig. 1. Genomic and transcriptomic analysis of the Perox-AlcOx pair. (A) The multistage plant 110 infection process of appressorium-forming fungi: appressorium (AP), penetration (PP), biotrophic (BP) 111 112 and necrotrophic (NP) phases. (B) Phylogenomic occurrence of the *perox-alcox* pair and consensus 113 genomic environment (within each clade) amongst pathogenic ascomycetes (black bold lettering indicates species for which transcriptomics data are shown in panel C, and grey lettering indicates 114 115 absence of the gene pair. Note that analogous tandem oxidases systems may occur in other fungi but 116 were possibly not detected due to the stringent sequence identity thresholds used to define AlcOx and Perox-encoding genes. Selected Colletotrichum species and associated consensus sequences are 117 representative of their respective species complexes; see fig. S3). (C) Time-course transcriptomic 118 analysis of the Tandem Peroxidases (Perox, green) and AlcOx (orange) encoding genes (gene accession 119 120 numbers in **Table S1**) during plant infection for different pathosystems (Myc., Mycelium) (26–29). Actual time points associated with each infection stages are provided in the Material and Methods 121 122 section. (**D**) Illustration of targeted plant hosts.

#### 123 Tandem Perox-AlcOx oxidize plant long chain alcohols

As a prelude to analyzing their biological function in vivo, we studied the substrate 124 specificity and enzyme interplay of the Perox-AlcOx pair in vitro (Fig. 2A). C. orbiculare was 125 selected as a model because it not only causes the economically important anthracnose disease 126 of cucurbits (e.g., melons, cucumber) but also has been used for decades as a model system for 127 studying fungal pathogenesis (30). Despite the notorious difficulties associated with 128 heterologous expression of such metalloenzymes, we successfully produced in the yeast Pichia 129 pastoris recombinant copper-radical AlcOx and heme-iron tandem peroxidase from C. 130 orbiculare (hereafter CorAlcOx and CorPerox, respectively). Similar to the previously studied 131 AlcOx orthologs from C. graminicola and C. gloeosporioides (19) (fig. S1A), CorAlcOx 132 oxidized both aromatic and long-chain aliphatic primary alcohols (fig. S4A). This finding 133 raises the possibility that fatty primary alcohols present in the cuticle of many plant species 134 (21), including cucumber, could be the native substrates of these enzymes. 135

On the other hand, CorPerox was confirmed to be a peroxidase, albeit with moderate 136 catalytic efficiency ( $k_{cat} = 1.52 \pm 0.02 \text{ s}^{-1}$ ,  $K_M^{H2O2} = 80 \pm 3 \mu M$ ,  $k_{cat}/K_M = 1.9 \times 10^4 \pm 0.1 \text{ s}^{-1} \text{ M}^{-1}$ 137 <sup>1</sup>) compared to the only previously characterized ascomycete Class II peroxidase (25), 138 commercial horseradish peroxidase (HRP) (31), and well-studied basidiomycete lignin-active 139 peroxidases (32)  $(k_{cat}/K_{M} = 10^{4} - 10^{7} \text{ s}^{-1} \text{ M}^{-1})$ . CorPerox was only active on low redox-potential 140 substrates and not on any of the substrates of canonical lignin-active peroxidases (fig. S4B), 141 and required the presence of calcium ions for stability (fig. S4C). These observations are in 142 agreement with structural predictions (fig. S5), which indicate the presence of two conserved 143 calcium ion binding sites, but absence of the manganese binding site and the surface-exposed 144 tryptophan involved in long-range electron transfer, which are two key features of lignin-active 145 146 peroxidases (33).

Despite its comparatively low peroxidase activity, CorPerox activates CorAlcOx for 147 oxidation of primary alcohols in a dose-dependent manner, and to a much greater extent than 148 the plant peroxidase HRP does (Fig. 2B). The pH optima of CorAlcOx and CorPerox were 149 markedly different (ca. 8 and 4, respectively, **fig. S4D**). Importantly, the pH of the environment 150 measured on cucumber cotyledons at the time of triggering appressorium penetration by C. 151 orbiculare was between 7.5 and 8.0. Taken together, these results indicate that AlcOx 152 activation is not dependent on highly efficient peroxidase activity. We also heterologously 153 produced the Perox-AlcOx pair from the rice blast pathogen *M. oryzae*, of which *Mor*AlcOx 154 was recently confirmed to be a primary alcohol oxidase (34). Here, we obtained an activation 155 profile for the MorPerox-MorAlcOx pair resembling that observed for the C. orbiculare pair 156 (fig. S4E cf. Fig. 2B). 157

Having determined optimal enzyme activation conditions, we then probed further the activity of the *Cor*Perox-*Cor*AlcOx pair on biologically relevant aliphatic alcohols. Despite challenges associated with substrate solubility in aqueous buffer, we clearly detected activity on hexadecan-1,16-diol and octadecan-1-ol, as well as on a crude preparation of waxes extracted from cucumber cotyledons (**Fig. 2C**). Importantly, such activity was detected only when both enzymes were present. Product analysis by gas chromatography unambiguously indicated that octadecan-1-ol was oxidized to the corresponding aldehyde (**fig. S4F**).

To obtain a deeper understanding of the activation of AlcOx by the Tandem Peroxidase, we analyzed electron transfer by electron paramagnetic resonance (EPR) spectroscopy (**Fig.** 

2D) and state-of-the art molecular modeling (Fig. 2E and fig. S6). Reduction in the EPR signal 167 of the inactive Cu(II)-non-radical form of CorAlcOx upon addition of CorPerox was 168 supportive of one-electron oxidation leading to the EPR silent, active Cu(II)-radical form (Fig. 169 2A and D). This change in electronic structure, observed in the absence of any substrate, 170 confirms the activity-independent activating role of CorPerox. It also indicates a close contact 171 between the enzymes during the activation process, which is concordant with our modeling 172 studies (see below) and with the sigmoidal, titration-like curves observed during activity assays 173 carried out in the presence of substrate (Fig. 2B). Further, CorAlcOx-CorPerox top 5 models 174 predicted by two independent computational techniques, PIPER (35) and AlphaFold2-175 Multimer (36) consistently placed CorPerox structures in front of the CorAlcOx active site 176 (fig. S6A). Refinement of these models with the all atom Monte Carlo software PELE (37), 177 which includes protein small rotations and translations followed by an exhaustive side-chain 178 prediction at the interface, resulted in a clear minimum (fig. S6B) where the heme group of 179 CorPerox is oriented towards the CorAlcOx copper ion (Fig. 2E and fig. S6C). Notably, the 180 binding surfaces of AlcOx orthologs are considerably more hydrophobic than that of GalOx 181 (fig. S7A and B), which is consistent with the idea that AlcOx may have evolved to interact 182 with the hydrophobic plant cuticle. Moreover, in the top model (Fig. 2E), there is a significant 183 decrease of solvent exposure of the CorAlcOx active site, defining a cavity between both 184 enzymes that could facilitate diffusion and binding of a long-chain alcohol substrate. C18 185 docking and PELE induced fit simulations confirm this point, revealing a pronounced local 186 minimum in which the alcoholic group of the substrate is well-positioned for catalysis (fig. 187 188 **S6D**).







192 Fig. 2. Biochemical and biophysical evidence for interplay between CorPerox and CorAlcOx. (A) Reaction mechanism of CROs showing activation of the resting, inactive form of the enzyme via 193 formation of a tyrosine radical, yielding the Cu(II)-radical active form. The latter will oxidize an alcohol 194 into the corresponding aldehyde followed by regeneration of the active form via the two-electron 195 reduction of O<sub>2</sub> into H<sub>2</sub>O<sub>2</sub>. (B) CorAlcOx oxidation rate of benzyl alcohol in presence of varying 196 amounts of HRP or CorPerox. (C) Activity of CorPerox-CorAlcOx on crude extract of cucumber waxes 197 and derived long-chain aliphatic alcohols, monitored via the production of chromogenic resorufin (RF), 198 product of the Perox-catalyzed oxidation of Amplex-Red by  $H_2O_2$ , the latter being the co-product of 199 200 AlcOx-catalyzed oxidation of primary alcohols into aldehydes. (D) EPR spectra of inactive CorAlcOx before (orange curve) and after mixture with CorPerox (black curve). EPR parameters of the Cu(II) 201 inactive form:  $g_z = 2.270$ ,  $A_z^{Cu} = 171 \ 10^{-4} \text{ cm}^{-1}$ ,  $g_x = 2.047$ ,  $A_x^{Cu} < 50 \ 10^{-4} \text{ cm}^{-1}$ ,  $g_y = 2.054$ ,  $A_y^{Cu} < 50 \ 10^{-5}$ 202 <sup>4</sup> cm<sup>-1</sup> and super-hyperfine coupling constant corresponding to two N-ligands  $A^{N} = 43 \ 10^{-4} \ cm^{-1}$ . (E) 203 204 Lowest energy CorAlcOx-CorPerox complex obtained by protein-protein modeling simulation with 205 PELE (see fig. S6 for more details). The copper atom is shown as a blue sphere, the heme group as 206 magenta sticks and calcium ions as purple spheres.

#### 207 The Perox-AlcOx pair gates plant penetration

To investigate the role of the Perox-AlcOx pair in plant infection, we isolated single and double gene deletion mutants of *C. orbiculare* (fig. S8). Inoculation of spore suspensions onto intact cucumber cotyledons showed that fewer and smaller lesions were formed by all the mutants compared to the wild-type strains (Fig. 3A and B). For instance, the proportion of lesions with a diameter > 4 mm fell from 95 % to < 20%. Furthermore, similar phenotypes were obtained for single and double mutants, suggesting that both oxidases are crucial for fungal pathogenicity.

Further investigations indicated that neither single nor double perox/alcox deletion 215 mutants were affected in mycelial growth (fig. S8A). Microscopy revealed that the significant 216 loss of pathogenicity of the mutants was due to a large decrease in the frequency of host 217 penetration (Fig. 3C). However, morphogenesis, cell wall melanization, and turgor build-up 218 219 within appressoria cells were virtually indistinguishable from those of the wild-type strain (fig. **S8B-D**). The next step in the infection process is the emergence of a needle-like penetration 220 peg through a pore in the basal cell wall of the appressorium (Fig. 1A), during which actin 221 222 assembly at the pore provides rigidity (38). Using a red fluorescent protein/actin-binding protein fusion (Lifeact-RFP), we found normal actin assembly at the appressorium pore for 223 both the single and double mutants inoculated onto cucumber cotyledons (fig. S8E and F). 224

Remarkably, the mutants could penetrate and form hyphae inside inert cellophane membranes (**fig. S9A-C**) and caused wild-type-like lesions when inoculated on mechanically wounded cucumber cotyledons (**Fig. S9D and E**), in contrast to the crippled invasive capacity observed on intact cotyledons (**Fig. 3C**). This points to a mechanism involving plant surface compounds, concordant with the catalytic activity of the Perox-AlcOx pair (**Fig. 2C**). These results collectively indicate that *Cor*AlcOx and *Cor*Perox play a crucial role during the early penetration stage, but are not involved in either appressorium or peg formation.

To further examine the function of the Perox-AlcOx pair during plant infection, we 232 attempted to localize the proteins by live-cell imaging of CorAlcOx-mCherry and CorPerox-233 GFP driven by their native promoters. Although the CorAlcOx-mCherry and CorPerox-GFP 234 235 complemented the defect in pathogenicity of the deletion mutants, fluorescence of CorAlcOxmCherry and CorPerox-GFP was not detectable during appressorium formation on cucumber 236 cotyledons (fig. S10A), suggesting that gene expression was too low, consistent with 237 transcriptomic data (Fig. 1C), or that the gene products were secreted and diffused away from 238 the penetration site. However, the constitutive overexpression of CorAlcOx-mCherry and 239 *Cor*Perox-GFP, driven by the *translation elongation factor (TEF)* promoter, revealed that both 240 proteins accumulated specifically at the appressorial penetration pore and that signal intensity 241 increased during penetration peg formation (Fig. 3D, fig. S10B and C, and Supplementary 242 Movie 1). This protein co-localization observed in vivo is consistent with gene co-expression 243 data (Fig. 1C) and rationalizes the co-operative activity demonstrated by biochemical assays 244 (Fig. 2). Remarkably, CorAlcOx-mCherry and CorPerox-GFP were detected at the plant 245 surface beneath detached appressoria, at the penetration site (Fig. 3E), suggesting that the 246 tandem metalloenzymes are secreted from appressoria into the plant epidermis. Strikingly, 247 CorAlcOx-mCherry and CorPerox-GFP were not detected at the penetration site on cellophane 248 membranes (fig. S10D), suggesting that interaction of the fungus with the plant surface triggers 249 local and specific recruitment of the tandem metalloenzymes to the pore. 250

To probe the presence of potential natural substrates of the fungal AlcOx in the neighborhood of the penetration site, we carried out a compositional analysis of waxes present at the surface of uninoculated cucumber cotyledons (**fig. S11**). This analysis showed that the extracted waxes are mainly composed of odd-numbered alkanes (C27-C33) and evennumbered long-chain primary alcohols (C24-C32). Fatty aldehydes were found only as traces. Thus, this experiment demonstrates that potential substrates of AlcOx represent a major part of the plant cuticular compounds while AlcOx reaction products are very minor components.

To further explore the role of Perox-AlcOx in the fungus-plant dialogue, we exposed the *C. orbiculare* single and double mutants to a product of the AlcOx, *viz.* the aliphatic long chain aldehyde *n*-octadecanal. For all gene deletion mutants, the addition of *n*-octadecanal partially restored appressorium penetration ability and lesion formation on cucumber leaves (**Fig. 4A and B**), suggesting that the role of the fungal Perox-AlcOx pair is to generate long chain aldehydes to prime the fungus for efficient plant infection.

Taken together, our results suggest that the role of the fungal Perox-AlcOx pair is to increase locally the concentration of long-chain aldehydes. These aliphatic compounds, members of the volatile organic compounds, are well-known to function as signal molecules (*39*), raising the possibility that the Perox-AlcOx pair generates signals to prime the fungus for efficient plant infection.

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To gain insights into the possible steps controlled by the Perox/AlcOx activities during 271 the early infection process, we performed comparative transcriptomic analyses of *C. orbiculare* 272 wild-type and double mutant strains at the appressorial stage on cucumber leaves and 273 cellophane (Fig. 5). Analysis of the differentially-expressed genes suggested that the Perox-274 AlcOx pair contributes to the regulation of a subset of 32 plant-inducible genes predicted to 275 encode small secreted proteins (SSPs), carbohydrate-active enzymes (CAZymes) and 276 membrane transporters (Table S6). SSPs and CAZymes are well-known fungal effectors 277 playing a key role in the molecular dialog with host plants. A phylogenetic analysis of 278 CAZymes (Table S6) present in this subset of genes revealed that the pair is required for the 279 up-regulation of genes encoding proteins directed towards the plant cell wall (PCW) and the 280 fungal cell wall (FCW) (Fig. 5). Among the former, we detected cellulose-active enzymes 281 known to display an enzymatic interplay - viz. the cello-oligosaccharide dehydrogenase AA7 282 and the lytic polysaccharide monooxygenase (LPMO) AA9 (40) – as well as pectin-active 283 enzymes from the PL1, PL3 and GH93 CAZy families. Regarding the FCW-targeting proteins, 284 we detected three proteins with carbohydrate binding modules (CBM50s; also called LysM 285 domains) that bind to chitin and function to evade recognition by host immune receptors during 286 infection (41). 287





Fig. 3. In vivo characterization of the role of the Perox-AlcOx pair during plant infection. A-B, 290 Infection phenotypes (A) and violin plot of the necrotic lesions size (B), at 5 dpi, for the wild-type (WT) 291 292 and perox/alcox deletion mutants of C. orbiculare on intact cucumber (Cucumis sativus) cotyledons. For each strain/plant combination, at least 60 inoculations were carried out. (C) Relative frequency 293 (WT set to 100%) of development of normal appressorium (AP) and penetrating hyphae (PH) by C. 294 orbiculare strains on cucumber cotyledons. Data are presented as average values (>300 appressoria for 295 296 each replicate, n = 3 independent biological replicates) and error bars show s.d. (**D-E**) Localization of the *C. orbiculare* tandem oxidases on cotyledons at 48 hpi, in the presence of appressoria cells (**D**) or 297 after detachment of the appressoria from the surface of the leaf (E). See movie S1 for a 3D view. In 298 299 panel **D**, white arrowheads indicate enzyme accumulation at the appressorium pore. In panel **E**, enlarged insets show a ring-like localization of the enzymes (scale bar = 1  $\mu$ m). In panels B-C, a one-tailed 300 independent *t*-test for each mutant vs WT was applied (\*\*\*P<0.001, \*\*\*\*P<0.0001). 301 302





Fig. 4. Effect of *n*-octadecanal on the pathogenicity of *perox/alcox* deletion mutants. The figure 305 306 shows the effect of the addition of 10 µM n-octadecanal (denoted as +C18) on necrotic lesions size 307 formed by C. orbiculare WT and mutant strains on cucumber cotyledons (24°C, 5 days) (A) and on the 308 formation of penetration hyphae (B). Control experiments, i.e. 1% ethanol without addition of noctadecanal, are denoted as Ref. In panel A, for each strain, we show one representative cotyledon 309 image of the infection phenotype and, below, a violin plot of lesions sizes based on 66 inoculation sites 310 per condition (carried out over 4 independent biological replicates). In panel **B**, data are presented as 311 312 mean percentages (>300 appressoria per replicate, n = 4 independent biological replicates) and error bars show s.d. In panels A and B, for each strain, a one-tailed independent t-test of +C18 vs Ref was 313 314 applied.



Figure 5. Loss of the Perox-AlcOx pair affects fungal gene expression during appressorium-316 mediated penetration of C. orbiculare. (A) Venn diagrams illustrating the differentially regulated (up-317 or down-regulated) genes between cucumber vs cellophane that are under control of the AlcOx-Perox 318 pair. **Top**: green circle indicates the number of downregulated genes (46 genes) in  $\Delta pair$  compared with 319 wild-type strain (WT), both incubated on cucumber; magenta circle denotes the number of genes (1608 320 genes) upregulated on cucumber vs cellophane in the WT strain. Bottom: Magenta circle indicates the 321 322 number of upregulated genes (86 genes) in *Apair* compared with WT, both incubated on cucumber; 323 green circle shows the number of downregulated genes (2887 genes) in the WT strain on cucumber vs cellophane. In the top and bottom Venn diagrams, the overlap indicates that 32 and 26 genes are 324 respectively upregulated or downregulated in the presence of the plant and also under control of the 325 AlcOx-Perox pair. (B) Class-wise distribution of the number of genes that were differentially expressed 326 327 in appressoria of the  $\Delta pair$  double mutant compared with WT on cucumber cotyledons. (C) Hierarchical clustering heatmap of differentially expressed genes encoding CAZymes and SSPs in the  $\Delta pair$  mutant 328 compared to WT on cucumber cotyledon. The expression values per gene were median normalized. 329 The columns were clustered by Euclidean distance. The expression levels of up- (magenta) and down-330 331 regulated (green) genes are shown as log2-transformed values. On the right-hand side of the figure, we show the presence of a predicted signal peptide (Y, Yes; N, No), phylogeny-based substrate specificity 332 predictions for CAZymes (PCW, plant cell wall; FCW, fungal cell wall; GAL, galactose-containing 333 compounds; Nd, not determined) and SSPs/CBM50s that are putative effectors (pale green circles) 334 335 predicted to localize in the plant cell (in the apoplast (A) and/or the cytoplasm (C)). See **Table S6** for 336 more details.

#### 338 **Discussion**

Research on plant invasion by appressorium-forming fungi teaches us that the 339 development of these specialized infection structures is an extremely complex, finely regulated 340 process. In this study, we have shown that fungal AlcOx, which are only encountered in 341 phytopathogens, have a very different biological function than the few distantly-related CROs 342 for which roles in morphogenesis (42, 43) or lignin degradation (18) were proposed. We 343 showed that the tandem Perox-AlcOx metalloenzymes are specifically deployed at the 344 initiation of penetration peg formation and their localization at the pore formed at the fungus-345 plant cell interface. Our results suggest that the oxidative action of Perox-AlcOx on long-chain 346 alcohols triggers a yet-to-be-elucidated biochemical cascade leading to penetration. Indeed, 347 functional complementation of the deletion mutants, which were defective in their ability to 348 puncture intact plant cuticles, by the product of the reaction suggests that cuticular alcohol 349 oxidation provides a chemical cue required for plant cell entry (Fig. 6). We speculate that 350 acquisition of the Perox-AlcOx pair provided the ecologically widespread Colletotrichum and 351 Magnaporthe species with an advantage, namely an in-house "locksmith" (i.e. the tandem 352 metalloenzymes) ensuring localized production of the aldehyde. This entry key is needed to 353 move to the next pathogenesis stage with the expression of PCW-degrading enzymes to 354 facilitate penetration, and of FCW-binding proteins and carbohydrate oxidases to evade host 355 immunity. For example, it has been suggested that, upon oxidation by fungal LPMOs and 356 oligosaccharide oxidases, plant oligosaccharides can no longer play their role as inducers of 357 host immune response (40, 44, 45). 358

It is interesting to speculate why Nature has evolved such a complex enzymatic system 359 relying on heme-iron peroxidase and elaborate copper radical chemistry when simpler, mono-360 enzyme systems could be employed for the same purpose (e.g. FAD-dependent oxidases active 361 on long-chain alcohols (46)). We show that the functional integrity of the secreted tandem 362 metalloenzymes system endows pathogenic fungi with fine control over oxidation reactions in 363 the extracellular space. We propose that the copper-radical center equips CROs with a "redox 364 switch" to turn on activity with high spatio-temporal resolution, when paired with a cognate 365 peroxidase (Fig. 6). More generally, our study suggests that fungi have evolved ways of 366 controlling oxidative reactions that are seemingly out of reach, i.e. beyond the bounds of the 367 fungal plasma membrane and cell wall, through tight genetic regulation and protein interplay 368 between secreted oxidoreductases. 369

In conclusion, using a combination of *in silico*, *in vitro* and *in vivo* approaches, we have 370 unveiled the existence of a natural redox partner, i.e. a heme-peroxidase, for AlcOx-type CROs, 371 and showed that the pair acts as a secreted virulence factor during early infection (Fig. 6). It is 372 noteworthy that the enzymology-driven approach pursued here was essential to bring to light 373 this unique mechanism, because low expression levels, fine temporal tuning, and highly 374 localized co-secretion of the Perox-AlcOx would have evaded classical 'omic approaches. The 375 376 specific occurrence of the Perox-AlcOx pair in most Colletotrichum and Magnaporthe species raises the possibility that functionally equivalent, coupled oxidative enzymatic mechanisms 377 may operate in other appressorium-forming fungal pathogens. We anticipate that the present 378 discovery will open new research avenues, notably on the role of wax-derived compounds in 379 380 the cascade of events leading to successful infection as well as on the development of oxidasespecific inhibitors as surface-acting, anti-penetrant drugs for crop protection. 381



386

Fig. 6. Schematic summary of the recruitment of the fungal Perox-AlcOx pair during early plant penetration and its proposed role in the induction of a biochemical cascade. The bottom panel is a zoom-in view illustrating the Perox-AlcOx reaction occurring at the fungus-plant interface and the triggered downstream cascade, highlighting the induction of genes encoding proteins targeting the plant cell wall (*pcw*) and fungal cell wall (*fcw*).

#### 393 Materials and Methods

394

#### 395 Chemicals and commercial enzymes

396 Most chemicals were purchased from Sigma (Darmstadt, Germany) or VWR (Fontenay-sous-397 Bois, France) unless stated otherwise. Molar concentrations of type II HRP (Sigma-Aldrich; 398 MW 33.89 kDa) were estimated by Bradford assay. *n*-octadecanal was purchased from TCI-399 Europe (Zwijndrecht, Belgium). All alcohol substrates stock solutions were prepared either in 400 H<sub>2</sub>O or in acetone and stored at -20°C. The concentration of H<sub>2</sub>O<sub>2</sub> stock solution was verified 401 at 240 nm ( $\epsilon^{240} = 43.6 \text{ M}^{-1}.\text{cm}^{-1}$ ).

402

# 403 **Bioinformatics**

Species tree of the 18 ascomycete genomes (shown in Fig. 1b) was constructed as previously 404 described (47). Core clusters containing only one protein-coding gene per species were 405 identified using FastOrtho (48) with the following parameters: 50% identity and 50% coverage. 406 Each cluster was aligned with MAFFT 7.221 (49), and curated alignments were concatenated 407 with Gblocks 0.91b (50). The tree was finally constructed with RAxML 7.7.2 (51) 408 (PROTGAMMAWAG model and 500 bootstrap). Phylogenetic analysis of AA5\_2 genes and 409 «standard» haem peroxidases (PFAM 00141, including Class II peroxidases) from 30 410 Colletotrichum genomes relied on 72 and 333 sequences, respectively (see fig. S2 legend and 411 **Table S3** for more details). The phylogenetic analysis of the peroxidase-catalase superfamily 412 relied on 150 sequences encompassing Class I (intracellular peroxidases), II (fungal secreted 413 peroxidases) and III (plant secreted peroxidases) peroxidases and from so-called Hybrid B 414 peroxidases, as previously reported (52). Subsequently to manual curation (removing signal 415 peptides), the sequences were first aligned using MAFFT-DASH (L-INS-i method) (53) and 416 the resulting multiple sequence alignment (MSA) was used to infer a phylogenetic tree using 417 RAxML (1000 bootstraps). The trees were then visualized in iTOL (54) and edited in 418 419 Illustrator<sup>®</sup>.

For the gene neighborhood survey, we retrieved 4 genes located upstream and downstream of each *Colletotrichum*'s AA5\_2 query (72 sequences). The resulting 569 genes were assigned to

422 78 different PFAM domains. The frequency of occurrence of a given type of domain in the 423 neighborhood of each AA5\_2 phylogenetic clade was then computed and visualized in Excel.

- 424 For interrogating the co-occurrence of both Perox and AlcOx-coding genes beyond
- *Colletotrichum* species two independent BLAST searches were run against the NCBI nonredundant database, using *Cor*AlcOx and *Cor*Perox as query sequences. 1,000 AlcOx-like (down to 37% sequence identity) and 1,000 Perox-like (25% sequence identity) sequences
- along with their corresponding source microorganism were retrieved. A cross-comparison of
  both lists of microorganisms, applying different sequence identity-based thresholds (60% for
  AlcOx and 30% for Perox) returned the list of species harboring both type of enzymes.
- 431 Transcriptomics data were retrieved from publicly available datasets (26–29). To normalize
- different dataset longitudinally, we expressed reported sampling time points as infection stages
- 433 in Fig. 1 as follows (hpi = hours post-infection): for *C. fructicola nara gc5/strawberry* (27),
- 434 AP (24 hpi), BP (72 hpi) and NP (144 hpi); for *C. orbiculare/N. benthamiana* (27), AP (24
- 435 hpi), BP (72 hpi) and NP (168 hpi); for *C. higginsianum/A. thaliana* (26), AP (22 hpi), BP (40 hpi) and NP (60 hpi); *C. argminicala/Maiza* (26), AP (22 hpi), BP (40 hpi) and NP (60 hpi);
- 436 hpi) and NP (60 hpi); *C. graminicola/Maize* (26), AP (22 hpi), BP (40 hpi) and NP (60 hpi);

437 *M. oryzae/Rice* (28), AP (8 hpi), BP (24 hpi) and NP (48 hpi); *M. oryzae/Barley* (29), AP (12
438 hpi), PP (24 hpi), BP (36 hpi) and NP (48 hpi).

439

## 440 Structure Prediction and Preparation

441 Surface hydrophobicity

Structural homology models were generated with AlphaFold (55) and surface hydrophobicity 442 of selected CROs was computed with the "protein-sol patches" online software (56). Average 443 hydrophobicity of the binding surface was determined as follows: using PyMOL 2.4, we 444 selected the residues constituting the entire binding surface of FgrGalOx, CgrAlcOx and 445 equivalent residues (based on MSA) in orthologous enzymes (5 GalOx and 11 AlcOx in total) 446 as well as those of 4 characterized hydrophobins (PDB 2N4O, 2LSH, 1R2M and 2FZ6) as 447 hydrophobic protein reference. Average hydrophobicity of the selected residues was computed 448 as a GRAVY index score (Kyte-Doolittle method) (57). 449

450 Preparation of models for docking experiments

We used Alphafold2 (55) to obtain a model for *Cor*AlcOx and *Cor*PerOx. To add the metals and cofactors to AlphaFold2 models, we performed a BLAST search in the Protein Data Bank proteins database and selected the PDB 2EIC (Sequence Identity 47%) to add the copper to *Cor*AlcOx, and the PDB 1MN2 (Sequence Identity 28%) to add the calcium ions and the heme group to *Cor*Perox. Subsequently, we prepared the systems with Schrödinger Protein Preparation Wizard (58) to determine the protonation states at pH 7 using PROPKA (59), and finally relax the systems performing a restrained minimization with convergence criteria for

- 458 heavy atoms to 0.30 Å using the OPLS\_2005 force field.
- 459 Protein-Protein Docking

For the generation of protein-protein poses (PPPs), we used PIPER (*35*) to generate 70,000 PPPs, which were clustered with the in-built clustering protocol using the default RMSD threshold of 9.0 Å and a minimum population of 10 poses per cluster. In parallel, we also used the recently developed Alphafold2 Multimer (*36*), which only uses sequence information, together with ion and cofactor placements, as described above.

### 465 *PPPs refinement with PELE*

We used the all atom Monte Carlo (MC) software PELE (Protein Energy Landscape 466 467 exploration), to map intermolecular interactions (37, 60). PELE follows a heuristic MC approach, generating new conformational proposals using vibration modes of the proteins with 468 translations and rotations of the ligand (CorPerox in this case), and relaxing the system with 469 structure prediction methods, so that the probability of acceptance in the Metropolis criterion 470 (61) remains high (60). Here, we refined the top five PIPER and five AlphaFold2-Multimer 471 models, using a simulation of 250 PELE steps with 256 computing cores (about 25 independent 472 trajectories per model). Each MC PELE step consists of a perturbation and a relaxation stage. 473 In the first one, a perturbation of the ligand, *Cor*AlcOx in this case, is first performed, including 474 random rotations of 0.01-0.04 rad and translations of 0.25-0.50 Å, and is followed by a 475 backbone perturbation of both proteins following the normal modes directions predicted by an 476 Anisotropic Network Model (ANM). In the second stage, the system is first relaxed by a high 477 478 resolution side chain prediction including all protein-protein interphase side chains, defined by

the region within 3 Å of any heavy atom of the other protein. Afterwards, a global minimization
is performed to relax the entire system, providing a final conformation and energy; this energy

is then used in a Metropolis importance sampling to accept/reject the MC step.

#### 482 *Exploring the substrate interactions*

The C18 substrate was created with Maestro 3D Builder, followed by a Glide docking (62) on 483 the lowest energy PPPs, as predicted with PELE. The docking grid was defined with a cubic 484 485 box of 30 Å centered in between the copper ion and Tyr120(OH) of the *Cor*AlcOx. Following the Glide rigid receptor docking, we performed a C18 induced fit rescoring simulation 486 involving 40 PELE steps (using 48 computing cores). Each PELE step consisted of a random 487 rotation of 0.01-0.04 rad and translation of 0.05-0.15 Å, using a spherically restrained search 488 space of radius 12 Å centered on the copper ion. The perturbation step was then followed by a 489 relaxation phase including side chain prediction (all side chains inside the spherical space) and 490 a full system minimization. 491

492

### 493 **DNA cloning and strain production**

DNA cloning and strain production of the AA5\_2 alcohol oxidases (AlcOx) from 494 Colletotrichum graminearum (CgrAlcOx, Genbank ID XM\_008096275.1, Uniprot ID 495 E3QHV8) was already carried out in previous study (19). The intron-free sequences of the 496 genes coding for the AlcOx from Colletotrichum orbiculare MAFF 240422 (CorAlcOx, 497 Genbank ID TDZ17043.1, Uniprot ID N4UTF2), the AlcOx from Magnaporthe oryzae 498 (MorAlcOx, Genbank ID XM\_003719321.1, Uniprot ID G4NG45), the Tandem Peroxidase 499 (Perox) from Colletotrichum orbiculare (CorPerox, Genbank ID TDZ17044.1, Uniprot ID 500 N4UUY4) and the Tandem Peroxidase from Magnaporthe oryzae (MorPerox, Genbank ID 501 XM\_003719322.1, Uniprot ID G4NG46) were synthesized after codon optimization for 502 expression in *P. pastoris* and inserted into a modified pPICZ $\alpha$ C vector using *Xho*I\* and *Not*I 503 restriction sites in frame with the  $\alpha$  secretion factor at N-terminus (i.e. without native signal 504 peptide) and with a  $(His)_6$ -tag at the C-terminus (without *c*-myc epitope) (Genewiz, Leipzig, 505 Germany). Transformation of competent P. pastoris X33, selection of zeocin-resistant 506 P. pastoris transformants screened for protein production was carried out as described by Haon 507 et al. (63). The best-producing transformants were conserved as glycerol stock at  $-80^{\circ}$ C. 508

509

### 510 Heterologous protein production in flasks

All proteins were first produced in 2 L Erlenmeyer flasks. To this end, single colonies of P. 511 pastoris X33 expressing each gene of interest were individually streaked on a YPD agar plate 512 containing Zeocin (100  $\mu$ g.mL<sup>-1</sup>) and incubated 3 days at 30°C. A single colony was then used 513 to inoculate 5 mL of YPD, in a 50 mL sterile Falcon tube and incubated during 5 h (30°C, 160 514 rpm). This pre-culture was used to inoculate at 0.2% (vol/vol) 500 mL of BMGY medium, in 515 a 2 L Erlenmeyer flask, incubated during approximately 16 h (30°C, 200 rpm) until the OD<sub>600</sub> 516 <sub>nm</sub> reached 4–6. The produced cellular biomass was then harvested by centrifugation (5 min, 517 16°C, 3,000 x g). For the AlcOx, the cell pellet was then resuspended in 100 mL BMMY 518 medium supplemented with methanol (1%, vol/vol) and CuSO<sub>4</sub> (500 µM). The culture was 519 incubated for 3 days (16°C, 200 rpm), with daily additions of methanol (1% added, vol/vol). 520 The Tandem Peroxidases production conditions were optimized and varied from the standard 521

- protocol as follows: the BMMY was supplemented with methanol (3% vol/vol), hemin (25  $\mu$ M) and CaCl<sub>2</sub> (2 mM). The culture was incubated for 3 days (20°C, 200 rpm), with daily additions of methanol (3%, vol/vol) and hemin (25 μM). Then, the extracellular medium was recovered by centrifugation (10 min, 4°C, 3,000 x *g*) and the supernatant filtrated on 0.45 μm membrane (Millipore, Massachusetts, USA) and stored at 4°C prior to purification.
- 527

## 528 Heterologous protein production in bioreactors

The upscaled production of CorPerox was carried out in 1.3 L and 7.5 L bioreactors (New 529 Brunswick BioFlo 115 fermentor, Eppendorf, Germany) as per the *P. pastoris* fermentation 530 process guidelines (Invitrogen) with the following optimizations: the glycerol fed-batch phase 531 was replaced by a sorbitol and methanol transition phase, besides 200 µM (1.3 L bioreactor) 532 and 150 µM (7.5 L bioreactor) of hemin were added to the methanol solution. CaCl<sub>2</sub> (10 mM 533 final) was added to the crude protein solution before being either directly purified or flash-534 frozen in liquid nitrogen and stored at -80°C. We verified that flash-freezing did not cause any 535 activity loss, for both AlcOx and Perox enzymes. 536

537

# 538 **Protein purification**

539 The filtered *Cor*AlcOx and *Mor*AlcOx crude supernatants were adjusted to pH 8.5, filtered on 540 0.22  $\mu$ m filters (Millipore, Molsheim, France), and purified by anion exchange 541 chromatography (DEAE) on a HiPrep FF 16/10 column (GE Healthcare, USA). Elution was 542 performed by applying a linear gradient from 0 to 500 mM NaCl (in Tris-HCl buffer 50 mM, 543 pH 8.5) over 20 column volumes, with a flow rate set to 5 mL.min<sup>-1</sup>.

- The filtered CorPerox and MorPerox culture supernatant was adjusted to pH 7.8 just before 544 purification and filtered on 0.22 µm filters (Millipore, Molsheim, France). Depending on the 545 volume to purify, the crude protein sample was either loaded on a His-Trap HP 5-mL column 546 (GE Healthcare, Buc, France) on a HisPrep FF 16/10 column (GE Healthcare) connected to an 547 ÄKTAxpress system (GE Healthcare) equilibrated with HEPES (10 mM, pH 8.0), NaCl (100 548 mM), CaCl<sub>2</sub> (2 mM) and imidazole (10 mM) buffer. Each (His)<sub>6</sub>-tagged recombinant enzyme 549 was eluted with HEPES (10 mM, pH 8.0), NaCl (100 mM), CaCl<sub>2</sub> (2 mM) and imidazole (500 550 mM) buffer. The Tandem Peroxidases were further purified by size exclusion chromatography, 551 using a HiLoad 26/600 Superdex 200 pg column (GE Healthcare) operated at 2.5 mL/min and 552 with a running buffer containing HEPES (10 mM, pH 8.0), NaCl (100 mM) and CaCl<sub>2</sub> (2 mM). 553 After SDS-PAGE analysis, fractions containing the recombinant enzyme were pooled, 554 concentrated and buffer exchanged in sodium phosphate (50 mM, pH 7.0) for the AlcOx or in 555 HEPES (10 mM, pH 8.0), NaCl (100 mM) and CaCl<sub>2</sub> (2 mM) buffer for the Tandem 556
- 557 Peroxidases.
- 558 Protein concentrations of *Cgr*AlcOx (52,337 Da,  $\epsilon^{280} = 101,215 \text{ M}^{-1}.\text{cm}^{-1}$ ), *Cor*AlcOx (52,317
- 559 Da,  $\epsilon^{280} = 92,735 \text{ M}^{-1}.\text{cm}^{-1}$ ), *Mor*AlcOx (62,894 Da,  $\epsilon^{280} = 90,020 \text{ M}^{-1}.\text{cm}^{-1}$ ), *Cor*Perox (26,137
- 560 Da,  $\epsilon^{280} = 21,345 \text{ M}^{-1}.\text{cm}^{-1}$ ), and *Mor*Perox (26,290 Da,  $\epsilon^{280} = 24,450 \text{ M}^{-1}.\text{cm}^{-1}$ ) were
- determined by the Bradford assay (64) using BSA as reference protein as well as by UV
- absorption at 280 nm using a Nanodrop ND-200 spectrophotometer (Thermo Fisher Scientific,
   Massachusetts, USA).
- 564

#### 567 Enzyme assays

For screening the substrate specificity of CRO-AlcOx enzymes, the alcohol substrates were 568 prepared in sodium phosphate buffer (50 mM, pH 7.0) in 96-well microplates and reactions 569 were initiated by the addition of a pre-mix of CRO-AlcOx (1 nM final concentration), HRP 570 (0.1 mg.mL<sup>-1</sup>) and 2,2'-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid (ABTS, 500 µM) in 571 sodium phosphate buffer (50 mM, pH 7.0). The tested substrates included D-glucose (50 mM 572 final concentration, D-Glc), D-galactose (50 mM, D-Gal), D-raffinose (50 mM, D-Raf), 573 xyloglucan (0.1% mM, XG), butan-1-ol (3 mM), butan-2-ol (3 mM), octan-1-ol (3 mM), 574 decan-1-ol (3 mM), 2,4-hexadiene-1-ol (3 mM, HD-OH), glycol aldehyde dimer (3 mM, 575 GAD), benzyl alcohol (3 mM, BnOH), 4-hydroxybenzyl alcohol (3 mM, p-OH BnOH), vanillic 576 alcohol (3 mM, Van-OH), syringic alcohol (3 mM, Syr-OH) and cinnamyl alcohol (3 mM, Cin-577 OH). The absorbance of the final reaction (100 µL total volume) was monitored at 414 nm 578 using a microplate spectrophotometer (TECAN), thermostated at 23°C. The 414 nm 579 absorbance allows to determine the concentration of ABTS cation radical over time (ABTS<sup>++</sup>, 580  $\epsilon^{414} = 31,100 \text{ M}^{-1}.\text{cm}^{-1}$ ), and in turn the rate of alcohol oxidation, considering a peroxidase 581 reaction stoichiometry for (H<sub>2</sub>O<sub>2</sub>:ABTS<sup>•+</sup>) of 1:2 and a CRO-AlcOx reaction stoichiometry for 582  $(alcohol:H_2O_2)$  of 1:1. 583

For screening the substrate specificity of *Cor*Perox, unless stated otherwise the enzyme (0.125 584 µM final) was prepared in citrate-phosphate buffer (50 mM, pH 4.0 to 7.0) in 96-well 585 microplates (for wavelength in the visible range) or in 1 mL Quartz cuvettes (for UV range), 586 in the presence of various substrates (vide infra). Reactions were initiated by the addition of 587 H<sub>2</sub>O<sub>2</sub> (100 µM final), incubated at 23°C, and monitored spectrophotometrically at the 588 wavelengths indicated below. The tested peroxidase substrates included: ABTS (500 µM) 589 converted into ABTS<sup>•+</sup> ( $\epsilon^{414}$  = 31,100 M<sup>-1</sup>.cm<sup>-1</sup>); 2,6-dimethoxyphenol (2,6-DMP, 500  $\mu$ M) 590 converted into hydrocoerulignone ( $\epsilon^{469} = 53,200 \text{ M}^{-1}.\text{cm}^{-1}$ ); guaiacol (500 µM) converted into 591 the final product tetraguaiacol ( $\epsilon^{470} = 26,600 \text{ M}^{-1}.\text{cm}^{-1}$ ); Reactive Black 5 (RB5, 100  $\mu$ M,  $\epsilon^{600}$ 592 = 20,000  $M^{-1}$ .cm<sup>-1</sup>) converted into non-chromogenic product RB5<sup>ox</sup>; veratryl alcohol (500  $\mu$ M) 593 converted into veratraldehyde ( $\varepsilon^{310} = 9,300 \text{ M}^{-1}.\text{cm}^{-1}$ ). For testing the manganese peroxidase 594 activity, CorPerox was mixed with Mn(II)SO<sub>4</sub> (1 mM final) in tartrate buffer (50 mM, pH 2.0

- activity, *Cor*Perox was mixed with Mn(II)SO<sub>4</sub> (1 mM final) in tartrate buffer (50 mM, pH 2.0 to 5.0) and the formation of Mn<sup>3+</sup>-tartrate complex upon addition of H<sub>2</sub>O<sub>2</sub> (100  $\mu$ M) was followed at 238 nm ( $\epsilon^{238} = 6,500 \text{ M}^{-1}.\text{cm}^{-1}$ ), as previously described (65).
- 598 All activities were expressed as Vi/E (s<sup>-1</sup>), i.e. the initial rate (Vi,  $\mu$ moles of H<sub>2</sub>O<sub>2</sub> consumed 599 per second) divided by the amount of enzyme (in  $\mu$ moles).
- 600 *Cor*Perox stability over time was carried out by monitoring the peroxidase activity of *Cor*Perox
- samples (50  $\mu$ M) stored in sodium acetate buffer (50 mM, pH 5.2), at 4°C, in the presence of
- varying concentrations of CaCl<sub>2</sub> (0-500 mM). The peroxidase activity of these samples was
- 603 measured as described above (final concentration of 0.5 μM *Cor*Perox), using ABTS (500 μM)
- and  $H_2O_2$  (100  $\mu$ M) as substrates, in citrate-phosphate buffer (50 mM, pH 4.0), at 23°C.
- Michaelis-Menten kinetic parameters of *Cor*Perox were determined by measuring the peroxidase initial rate, as described above, in the presence of ABTS (500  $\mu$ M) and varying

607 concentrations of  $H_2O_2$  (0-1200  $\mu$ M), in citrate-phosphate buffer (50 mM, pH 4.0), at 23°C. 608 Experimental data could be fit to the standard Michaelis-Menten equation (residual standard 609 error = 0.019).

- AlcOx activation by the peroxidases was assayed by monitoring changes in absorbance at 254 610 nm upon oxidation of benzyl alcohol (1.5 mM) into benzaldehyde by the CRO (10 nM final 611 concentration), in the presence of varying concentrations of peroxidase (0-1000 nM). Reactions 612 were carried out in sodium-phosphate buffer (50 mM, pH 7.0), at 23°C, in UV-transparent 613 cuvettes (1 mL reaction volume). The reactions were initiated by addition of the CRO and 614 vigorously mixed by pipetting up and down. The absorbance was measured using an Evolution 615 201 UV-Vis spectrophotometer (Thermo-Fisher). The concentration of benzaldehyde was 616 calculated as [Benzaldehyde]<sub>t</sub> =  $(Abs^{254 \text{ nm}}_{t} - Abs^{254 \text{ nm}}_{t0})/(\epsilon^{254}_{benzaldehyde}-\epsilon^{254}_{BnOH})$ , where 617  $\epsilon^{254}_{\text{benzaldehvde}} = 8,500 \text{ M}^{-1}.\text{cm}^{-1}$  and  $\epsilon^{254}_{\text{BnOH}} = 150 \text{ M}^{-1}.\text{cm}^{-1}$ . 618
- 619

#### 620 Gas chromatography (GC) analysis

Enzymatic reactions were carried out in 4 mL-clear borosilicate glass vials closed by screw 621 caps with PTFE septum (500 µL final reaction volume). CorAlcOx (2 µM final) was mixed 622 with CorPerox (2 µM) in sodium phosphate buffer (50 mM, pH 7.0). The reaction was initiated 623 by the addition of octadecanol (0.3 mg.mL<sup>-1</sup>, eq. 1.1 mM final) and the mixture was incubated 624 at 23°C at 190 rpm in an Innova 42R incubator (New Brunswick, USA), during 1 h. Following 625 a previously published protocol (22), the reaction mixture was then acidified by addition of 10 626 µL HCl (12 M). Products and possible remaining substrate were extracted by adding 500 µL 627 of hexane (containing 1 mM of internal standard dodecane), followed by shaking and 628 centrifugation for 5 min at 3,000 x g. The organic layer was transferred into a new vial and 629 analyzed with a GC-2010 Plus apparatus (Shimadzu, Japan) equipped with a flame ionization 630 detector (FID) and a DB-5 capillary column (30 m x 0.25 mm x 0.25 µm; Agilent). Nitrogen 631 (200 kPa) was used as carrier gas. The injector and detector temperatures were set at 250°C. 632 After injection (2 µL sample), the analytes were separated by applying the following 633 temperature program: step 1) from 65°C to 250°C over 9.25 min (i.e. 20°C/min); step 2) plateau 634 at 250°C for 6 min. For quantitation, standard curves of octadecanol, n-octadecanal and 635 octadecanoic acid were prepared by following the same procedure. 636

#### 638 Electron Paramagnetic Resonance (EPR)

EPR spectra were recorded on frozen solutions (120K) using a Bruker Elexsys E500 639 spectrometer operating at X-band equipped with a BVT 3000 digital temperature controller. 640 The following acquisition parameters were used: modulation frequency 100 kHz; modulation 641 amplitude 5 G; gain 87 dB; and microwave power, 20 mW. EPR spectra were simulated using 642 the EasySpin toolbox developed for Matlab (66). CorAlcOx (100 µM final), prepared in 643 sodium phosphate buffer (50 mM, pH 7.0), in the absence or presence of CorPerox (100 µM 644 final), was flash-frozen in liquid nitrogen and continuous-wave EPR spectra were recorded. 645 CorAlcOx and CorPerox were placed in contact for various amount of time (2.5 min, 15 min) 646 before flash-freezing the solution. Controls containing buffer only or the CorPerox were also 647 carried out. 648

649

### 650 Analysis of cuticular waxes from cucumber cotyledons

651 Cuticular waxes were extracted from 2-weeks old cotyledons by immersing 6 intact cotyledons 652 for 30 s in chloroform in a glass beaker. Chloroform was evaporated under a stream of nitrogen 653 gas and wax extracts were derivatized using N,O-Bis (trimethylsilyl)trifluoroacetamide 654 (BSTFA) and analyzed by gas chromatography coupled to mass spectrometry (GC-MS) as 655 previously described (67).

656

#### 657 Strains and media

658 Strain 104-T (MAFF240422) of *C. orbiculare* was used as the wild-type strain. All strains used 659 in this study are listed in **Table S4** (*13*, *38*, *68–70*). *C. orbiculare* strains were cultured on 3.9% 660 PDA (Nissui) at 24°C in darkness. For genetic manipulation, *Escherichia coli* DH5α-661 competent cells were maintained on Luria-Bertani (LB) agar at 37°C. For fungal 662 transformation, *Agrobacterium tumefaciens* C58C1 was maintained on LB agar at 28°C. 663 Transformations of *C. orbiculare* (*13*, *38*) were carried out as previously described.

664

### 665 Strain construction

Primers and plasmids used in this study are listed in **Table S5**. For construction of *CorAlcOx* deletion strains, 1.1-kb upstream and 1.0-kb downstream flanking sequences and a 1.0-kb fragment of the neomycin-resistance cassette was amplified with the respective primer pairs. For construction of *CorPerox* deletion strains, 1.1-kb upstream and downstream flanking sequences and a 1.4-kb fragment of the hygromycin-resistance cassette was amplified. These three fragments were inserted into linearized pPZP-PvuII using the In-Fusion HD cloning kit (Clontech). The same procedures were used for construction of *M. oryzae* gene deletion strains.

- 673 For construction of *CorAlcOx-mCherry* and *CorPerox-GFP* gene fusion, a 5.9-kb *CorAlcOx-*
- 674 *CorPerox* fragment containing 1.1-kb downstream flanking sequences was inserted into 675 linearized pPZP-PvuII-SUR, and the mCherry and GFP fragments were inserted.
- 676 For construction of *CorAlcOx-mCherry* overexpression strains, a 4.0-kb *CorAlcOx-mCherry*
- 677 fragment containing its 1.1-kb downstream flanking sequence was amplified from pPZP-678 AlcOx-mCherry-Perox-GFP-S, and fused to linearized pCAMSUR-TEF (71) containing the 679 translation elongation factor promoter of *Aureobasidium pullulans* (72). The same procedures
- 680 were used for construction of *CorPerox-GFP* overexpression strains.
- 681

# 682 Plant infection

Infection assays on detached cucumber leaves (*Cucumis sativus* L. 'Suyo') with conidial suspension ( $1 \times 10^5$  conidia/ml in distilled water) of *C. orbiculare* were performed as previously described (73). The inoculated leaves were incubated in a humid environment for 5 days at 24°C. For testing the effect of *n*-octadecanal supplementation, conidial suspensions ( $1 \times 10^5$ conidia/mL in 10 µM *n*-octadecanal dissolved in 1% ethanol or 1% ethanol as a control) were spotted onto detached cucumber leaves, and incubated in a humid environment for 5 days at 24°C.

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# 691 Microscopy

For observation of appressorium formation of *C. orbiculare*, a conidial suspension  $(5 \times 10^5$  conidia/ml) was placed on a multiwell glass slide or cover glass (Matsunami Glass),

respectively. Cells were incubated in a humid box for 24 h at 24°C in the dark. Observation of penetration hyphae on cucumber cotyledons and cellophane membranes were performed as previously described (*13*). Appressorial cytorrhysis assay was conducted based on a previous procedure (*68*).

A confocal laser scanning microscope LSM900 with Airyscan 2 (Carl Zeiss) equipped with a 698 Plan Apochromat 63×/1.4 Oil DIC objective (Carl Zeiss) was used to acquire confocal 699 microscopic images. Excitation/emission wavelengths were 488 nm/490-556 nm for GFP and 700 561 nm/565-630 nm for mCherry. Images were acquired and processed using ZEN Software 701 (Version 3.1; Carl Zeiss) and Imaris (Version 9.3.1; Bitplane). For detection of appressorial 702 actin assembly, cells were observed using a Zeiss Axio Imager M2 Upright microscope (Carl 703 Zeiss) equipped with a Plan Apochromat 100× oil immersion lens, an Axio Cam MRm digital 704 camera and excitation/barrier filter set of 595 nm/620 nm for RFP. Images were acquired using 705 Axiovision 4.8. Bright-field microscopy was performed using a Nikon ECLIPSE E600 706 microscope equipped with a 40× water immersion lens (Nikon) and an OLYMPUS DP74 707 digital camera system. 708

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#### 711 Microarray analysis

For sampling appressoria, the abaxial surface of cucumber cotyledons or cellophane 712 membranes (Wako Chemicals) were inoculated with 10 µL droplets or 10 mL, respectively, of 713 a conidial suspension  $(1 \times 10^6 \text{ conidia/ml})$ , and then incubated at 24°C in a humid box. After 714 16 h, the cellophane was frozen in liquid nitrogen. After 24 h, the lower epidermis of the 715 716 cotyledons was peeled off. All samples were ground in liquid nitrogen, and total RNA was prepared using the Maxwell RSC Plant RNA Kit (Promega) and the Agilent Plant RNA 717 Isolation MiniKit (Agilent Technologies). Microarray analyses were performed as described 718 previously (13) using the C. orbiculare  $(8 \times 60,000, 13,352)$  independent probes, Design ID: 719 060762) oligo microarray, according to the Agilent 60-mer Oligo Microarray Processing 720 Protocol (Agilent Technologies). The normalization condition: i) intensity-dependent Lowess 721 normalization; ii) data transformation, measurements less than 0.01 were set to 0.01; iii) per-722 chip 75<sup>th</sup>-percentile normalization of each array; and iv) the expression values per gene were 723 median normalized. The normalized data were subjected to a *t* test, with statistically significant 724 gene sets defined as those giving *P* values less than 0.05. The differentially-regulated genes 725 (fold change > 2 and P < 0.05) were selected and were used for further analysis. Functional 726 classification was based on the Gene Ontology (GO), protein families (Pfam) and C. orbiculare 727 genome information of CAZymes and SSPs. Note that to avoid redundancy in the count of up-728 and downregulated genes, all sequences considered as CAZymes were not counted in the 729 broader classes of Hydrolases/Transferases/Oxidoreductases. 730

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#### 983 Supplementary Materials:

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