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Old Yellow Enzyme homologues in Mucor circinelloides: Expression profile and biotransformation

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(Article begins on next page)

1 Old Yellow Enzyme homologues in *Mucor circinelloides*: expression profile and biotransformation. 2 Alice Romagnolo^a, Federica Spina^a, Anna Poli^a, Sara Risso^a, Bianca Serito^a, Michele 3 Crotti^b, Daniela Monti^c, Elisabetta Brenna^b, Luisa Lanfranco^a and Giovanna Cristina 4 5 Varese^a* ^a Department of Life Sciences and Systems Biology, University of Turin, viale P. A. 6 7 Mattioli 25, 10125 Turin, Italy ^b Department of Chemistry, Materials and Chemical Engineering "G. Natta", 8 9 Politecnico di Milano, via L. Mancinelli 7, 20131 Milan, Italy ^c Istituto di Chimica del Riconoscimento Molecolare, CNR, Via M. Bianco 9, 10 11 20131 Milan, Italy *Corresponding author (G.C. Varese) Tel.: +39 0116705984; Fax: +39 0116705962 E-mail address: cristina.varese@unito.it Postal address: viale P.A. Mattioli 25, 10125 Turin, Italy 12 13 14 15 16 17

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21

22	Abstract
23	The reduction of C=C double bond, a key reaction in organic synthesis, is mostly
24	achieved by traditional chemical methods. Therefore, the search for enzymes capable of
25	performing this reaction is rapidly increasing. Old Yellow Enzymes (OYEs) are flavin-
26	dependent oxidoreductases, initially isolated from Saccharomyces pastorianus.
27	In this study, the presence and activation of putative OYE enzymes was investigated in
28	the filamentous fungus Mucor circinelloides, which was previously found to mediate
29	C=C reduction. Following an in silico approach, using S. pastorianus OYE1
30	amminoacidic sequence as template, ten putative genes were identified in the genome of
31	M. circinelloides. A phylogenetic analysis revealed a high homology of McOYE1-9
32	with OYE1-like proteins while McOYE10 showed similarity with thermophilic-like
33	OYEs.
34	The activation of meoyes was evaluated during the transformation of three different
35	model substrates. Cyclohexenone, α -methylcinnamaldehyde and methyl cinnamate were
36	completely reduced in few hours and the induction of gene expression, assessed by
37	qRT-PCR, was generally fast, suggesting a substrate-dependent activation. Eight genes
38	were activated in the tested conditions suggesting that they may encode for active
39	OYEs. Their expression over time correlated with C=C double bond reduction.
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45	Keywords
46	α ,β-unsaturated compounds, C=C double bond reduction, gene expression, <i>Mucor</i>
47	circinelloides, Old Yellow Enzymes.

1. Introduction

48

49 The reduction of C=C double bonds is a key reaction in organic chemistry but it is 50 usually carried out by metal catalysts with a strong impact on the technical and 51 economic feasibility of the process [1, 2]. For instance, Yang et al. [3] reported that 52 toxic traces of heavy metals remained in the reaction products and needed to be 53 removed before pharmaceutical use. 54 Since the major challenges of bulk and fine chemicals synthesis are the reduction of the 55 environmental impact and process costs, biocatalysis became one of the most intriguing 56 alternative to traditional processes. The use of microorganisms or their enzymes has 57 recently found room in the industrial production of pharmaceuticals, flavors, aromas, 58 etc. [2, 4]. The biological reduction of activated C=C double bonds may be carried out 59 by flavin-dependent oxidoreductases, namely ene reductases (ERs), belonging to the 60 Old Yellow Enzyme (OYE) family (EC 1.6.99.1) [5]. They catalyze the asymmetric 61 hydrogenation of C=C double bond conjugated with electron withdrawing groups 62 (EWGs) in the presence of NAD(P)H as cofactor [2]. In contrast with heavy metals, 63 which are capable of mediating *cis*-hydrogenation, OYEs can catalyze this reaction 64 trans-fashion with high stereo-selectivity [1]. 65 The reactions catalyzed by OYEs are very interesting and have strong application 66 outcomes. Robinson and Panaccione [6] showed the involvement of OYE homologues 67 involvement in the biosynthetic pathway of ergot alkaloids, commonly used to treat 68 disorders such as Alzheimer's disease, dementia, type 2 diabetes, and 69 hyperprolactinemia or to induce labor and reduce bleeding (lysergic acid-derived 70 drugs). OYE1 from S. pastorianus transformed methyl 2-hydroxymethylacrylate in (R)-71 3-hydroxy-2-methylpropanoate, known as "Roche-Ester", which is a chiral building 72 block for the synthesis of vitamins (vitamin E) [4]. Some fragrance compounds 73 (muscone), antibiotics (rapamycin), and natural products have been obtained by OYE-

74 mediated reduction [2]. The 12-oxophytodienoate reductase enzymes (OPRs, EC 75 1.3.1.42), OYE homologues from plants, are involved in the biosynthesis of jasmonic 76 acid, which is implicated in the regulation of plant responses to abiotic and biotic 77 stresses as well as plant growth and development [7]. Pentaerythritol tetranitrate 78 reductase (PETNR) from Enterobacter cloacae successfully degraded tri nitro toluene 79 (TNT) [8]. 80 OYEs have been ubiquitously described in yeasts, bacteria, animals and plants, and 81 recently in filamentous fungi [9]. Fungi are perfect candidates to set up biocatalysis 82 processes: they combine operative versatility to simple growth conditions and they are a 83 well-known enzymatic machinery [1, 2, 4, 10]. For instance, a homologue of OYE has 84 been discovered in Aspergillus fumigatus and Claviceps purpurea and associated to the 85 ergot biosynthesis [6, 11]. To date, most of the literature evidences focused on 86 Ascomycetes and Basidiomycetes [9, 12] but the presence of OYE homologue within 87 Zygomycota phylum has never been assessed. 88 Despite the potential application in several biotechnological fields, microorganisms and 89 enzymes are still scarcely used in manufacturing processes, mostly due to the lack of 90 suitable biocatalysts. Novel enzymatic activities with strong catalytic potential could be 91 achieved with traditional functional screening or advanced molecular approaches [2, 4]. 92 Genome-wide analysis is a useful tool to identify OYEs homologues among the 93 available fungal genomes. For instance, Nizam et al. [9] by analysing 60 Ascomycota 94 and Basidiomycota genomes identified 424 OYEs homologues and provided a first 95 classification of these enzymes within the fungal kingdom. They also explored the 96 evolutionary significance of fungal OYEs. Unfortunately, this data can be considered 97 just a first step, and the actual capability of strains to transform target compounds by 98 reducing C=C double bond need further validation.

In this work, we aimed to fill the lack of information about the occurrence of OYEs in fungi belonging to the Zygomycota phylum. *Mucor circinelloides* was selected due to its ability of converting several substrates [13]. Despite those interesting results, the enzymatic pattern responsible for the reactions has never been investigated before. The availability of *M. circinelloides* complete genome sequence (Joint Genome Institute, JGI: http://jgi.doe.gov) allowed a genome-mining approach to investigate the presence of putative OYEs homologues.

2. Results

2.1 Identification of putative OYEs in the genome of M. circinelloides

In order to identify OYE encoding genes in the filamentous fungus *M. circinelloides*, a BlastP analysis (Basic Local Alignment Search Tool, NCBI, USA) on the complete genome of *M. circinelloides* using *S. pastorianus* OYE1 as query was performed. Ten putative sequences were retrieved and named McOYE1-McOYE10 (Table 1). The 10 amino acid sequences and the amino acid sequence of OYE1 were aligned to evaluate sequence similarities (Table 1). Nine McOYEs showed a similarity with *S. pastorianus* OYE1 of about 40 % while McOYE10 showed a lower similarity (25.33 %; Table 1).

Table 1

Putative OYE homologues of *M. circinelloides* - McOYE1, McOYE2, McOYE3,

McOYE4, McOYE5, McOYE6, McOYE7, McOYE8, McOYE9 and McOYE10 - with

sequence ID according to JGI database and identity percentage with *S. pastorianus*OYE1.

McOYE	Sequence ID	ID matrix (%) with OYE1	ID matrix (%) with McOYE1	ID matrix (%) with McOYE2	ID matrix (%) with McOYE3	ID matrix (%) with McOYE4	ID matrix (%) with McOYE5	ID matrix (%) with McOYE6	ID matrix (%) with McOYE7	ID matrix (%) with McOYE8	ID matrix (%) with McOYE9	ID mat (%) w <i>McOYI</i>
1	160302	44.14	97.50	89.60	70.40	75.00	67.80	66.10	60.60	67.00	58.40	37.0
2	137297	43.99	89.60	97.50	69.70	75.30	68.70	66.10	61.10	67.00	59.50	36.40
3	177510	42.19	71.20	70.50	96.40	76.10	63.30	63.60	57.10	61.60	60.50	40.70
4	155592	41.30	73.40	73.70	73.60	100.00	65.30	65.70	57.80	64.80	60.10	41.60
5	110873	43.41	65.80	66.70	60.40	65.30	100.00	64.50	61.60	65.90	59.10	39.10
6	144573	42.30	64.40	64.30	61.10	64.90	64.60	100.00	67.70	66.10	5 58.20	38.20
7	153280	41.80	63.70	64.20	62.10	62.20	66.30	68.80	90.50	60.50	56.60	45.10
Q	76836	42.10	65.70	65.70	50.50	64.80	65.00	66.10	56.70	100.00	50.20	

134845 38.19 56.90 57.90 57.90 59.50 57.70 57.50 54.90 58.50 10 152500 35.90 35.20 39.20 41.60 39.10 40.30 48.40 41.60 25.33 120 121 122 Three conserved domains typical of OYEs were found in all the 10 sequences: the FMN 123 binding site, the active site and the substrate binding site (Supp. Fig. 1). 124 Specific primer pairs were designed on the nucleotide sequences of the 10 putative 125 mcoves (Supp. Table 1) and tested by conventional PCR on genomic DNA. Amplicons 126 of the expected size (about 200 bp) were obtained (Supp. Fig. 2). PCR products were 127 sequenced confirming the specificity of the primers pairs and the authenticity of the 128 DNA sequences. 129 A phylogenetic analysis was performed by implementing sequence data analyzed by 130 Nizam et al. [9], who divided OYEs proteins into three groups: Class I, Class II and 131 Class III. Nine out of 10 McOYEs clustered together in Class I showing a specie-132 specific clade whereas McOYE10 was located within Class II (Fig. 1). 133 2.2 Biotransformation of conventional substrates and gene expression 134 The expression profile of the 10 putative OYEs homologues was monitored on the RNA 135 extracted from the mycelium grown in liquid culture during the biotransformation of 136 three conventional substrates presenting different EWGs. For each substrate, data on 137 biotransformation and gene expression pattern are presented. Since the substrate was 138 dissolved in dimethyl sulfoxide (DMSO), mcoyes activation was also evaluated in the 139 presence of this solvent to exclude artifacts. None of the genes was activated in the 140 presence of DMSO (data not shown). The 10 genes showed a basal activity in the 141 absence of substrates (data not shown). 142 2.2.1 Cyclohexenone (CE) 143 M. circinelloides completely reduced the substrate CE into cyclohexanol within 24 h; 144 the reaction process is well known: first an OYE reduces the C=C double bond of CE

97.00

41.80

100.0

145 producing cyclohexanone, then the keto group is reduced by an alcohol dehydrogenase 146 (ADH) into cyclohexanol (Fig. 2 A) [13]. As shown in Fig. 3 A, the reaction began 30 147 min after the addition of CE to the medium and at 3.5 h the C=C double bond was 148 completely reduced producing cyclohexanone which was continuously converted in its 149 corresponding alcohol, cyclohexanol. 150 The transcripts level of the 10 mcove homologues was monitored both in the presence 151 and absence of CE at 30 min, 1 h, 2 h and 5 h (Fig. 4 A). With the exception of mcoye7 152 and mcove8 that did not show activation upon CE exposure (data not shown), all the 153 other genes were activated within the first two hours. In particular mcoye2, mcoye1 and 154 mcoye 10 displayed a fast and strong induction of gene expression: 730, 111 and 76 fold 155 compared to the control sample without CE at 1 h and at 30 min for mcoye10 (Fig. 4 A). 156 Mcoye4 and mcoye5 showed an activation of 30-50 fold, while for mcoye3, mcoye6 and 157 mcoye9 the induction of gene expression compared to the control sample remained 158 below 20 fold at the different time points. Noteworthy, expression levels of these genes 159 decreased to the control values after 5 h. 160 A clear relation between *mcoyes* expression profile and the biotransformation of CE 161 was observed (Fig. 3 A). Mcove2 transcript levels were strongly induced at the 162 beginning of the reductive process reaching the maximum at 1 h when 20 % of CE had 163 been converted into cyclohexanone. 164 2.2.2 α -Methylcinnamaldehyde (MCA) 165 M. circinelloides completely reduced the C=C double bond of MCA within 20 h. Both 166 the C=C double bond (OYE) and the aldehydic group (ADH) of the substrate were 167 reduced (Fig. 3 B). One hour after MCA addition, α-methylcinnamyl alcohol 168 represented 50 % of the substrates in the culture medium. The concentration of this first 169 product increased until 8 h and then dropped down before 20 h. The production rate of

170 the saturated alcohol was constant, starting from 2 h until 20 h, when it was the 171 remaining metabolite detected (Fig. 3 B). 172 According to literature data [2], OYEs are able to catalyze the reduction of the C=C 173 double bonds of unsaturated aldehydes, whereas they are usually inactive on allylic 174 alcohols. In the case of M. circinelloides, α -methylcinnamyl alcohol seemed to be the 175 intermediate of the conversion of MCA into the corresponding saturated alcohol. Thus, 176 α-methylcinnamyl alcohol was added directly to M. circinelloides cultures, and indeed 177 its conversion into α -methyldihydrocinnamyl alcohol was observed to be complete after 178 48 h. In order to elucidate this reduction pathway, the dideuterated α -methylcinnamyl 179 alcohol, showing two deuterium atoms linked to the carbon atom bearing the OH group, 180 was prepared (Figure 2 B). This compound was submitted to bioreduction with M. 181 circinelloides and a monodeuterated saturated alcohol was recovered. The formation of 182 this compound could be explained only admitting the formation of the unsaturated 183 aldehyde as an intermediate, because the two deuterium atoms should have been 184 preserved in the direct reduction of the starting allylic alcohol. The alcohol 185 dehydrogenases, which are present in the fermentation medium, catalyse the oxidation 186 of the allylic alcohol to the unsaturated aldehyde, which is easily reduced by ERs and 187 removed from the equilibrium. Then, the saturated aldehyde is further reduced by 188 ADHs to afford the corresponding saturated alcohol. The intermediate aldehydes did not 189 accumulate in the reaction medium and it was not possible to detect them during the 190 reaction course by GC/MS analysis. On the basis of these results, the reaction sequence 191 shown in Figure 2 C can be hypothesized for MCA. 192 As for CE, the transcripts level of the 10 putative *mcoyes* was monitored both in the 193 presence and absence of MCA at 30 min, 1 h, 2 h and 4 h (Fig. 4 B). Mcoye2 showed 194 the highest gene activation level with about 2,880 fold compared to the control sample, 195 followed by mcoye1 that displayed 1,860 fold induction (Fig. 4 B). An induction of

196 gene expression of about 500 fold was observed for mcoye4, mcoye5 and mcoye10. 197 Remarkably, for these 5 genes the highest activation was reached 1 h after the addition 198 of the substrate. For mcoye3 and mcoye6 the highest induction levels (138 and 243, 199 respectively) were observed at 2 h. Mcoye 9 displayed moderate gene activation (about 200 14 fold) only at 4 h, mcoye7 and mcoye8 did not show activation, as it was observed 201 with CE (data not shown). 202 In this case too, a relation between *mcoyes* gene expression activation and MCA 203 biotransformation was observed (Fig. 3 B). The strongest activation of mcovel in 204 presence of MCA was reached at 1 h, well before the beginning of the C=C reduction, 205 represented by the formation of the saturated alcohol. 206 2.2.3 Methyl cinnamate (MCI) The MCI substrate was completely reduced by M. circinelloides within 66 h; both the 207 208 C=C double bond and the ester group were reduced producing cinnamyl alcohol and 209 phenylpropanol. The exact reaction profile is unknown; however, the one reported in 210 Figure 2 D can be hypothesized on the basis of what has been observed for MCA, 211 starting from the enzymatic hydrolysis of the ester moiety followed by the biocatalysed 212 reduction of the COOH group to primary alcohol. As shown in Fig. 3 C, MCI decreased 213 slightly but constantly until 66 h, when all the substrate was transformed. The detected 214 amount of cinnamyl alcohol was never more than 20 %; also the level of 215 phenylpropanol remained low (< 20 %) until 9 h, after which the concentration reached 216 100 % within 66 h. 217 The transcripts level of the 10 putative *mcoyes* was analyzed in presence and absence of 218 MCI at 1, 2, 3 and 6 h (Fig. 4 C). *Mcove1* was the most induced gene (about 300 fold). 219 An activation of about 60 fold was observed for mcove2, mcove5, and mcove10, while 220 mcoye3, mcoye4, mcoye6, and mcoye9 showed 20 fold induction. Mcoye1 and mcoye6 221 displayed the fastest activation, with the maximum within the first 2 h; after that time

222 point, their transcripts level rapidly decreased. Mcove 10 showed a peak between 2 and 3 223 h (Fig. 4 C). Also in this case, neither mcove7 nor mcove8 were activated upon substrate 224 exposure (data not shown). 225 A relation between OYE activation and the biotransformation of MCI was observed 226 (Fig. 3 C). The transcription of *mcoye1* started early, when the substrate was still the 227 only detectable compound in the reaction mixture. 228 3. Discussion 229 Hydrogenation of C=C double bonds is an important reaction in several manufacturing 230 processes for the production of bulk and fine chemicals; researchers and industries are 231 moving towards more sustainable approaches as biocatalysis and in recent years, several 232 research groups have focused on the identification of OYEs homologues to be exploited 233 in different processes [1]. In the last few years the attention was given to OYEs from 234 filamentous fungi; nevertheless only few studies report their occurrence and their 235 physiological role in this group of organisms [9, 13]. 236 An *in silico* approach allowed to identify in the genome of the zygomycete fungus M. 237 circinelloides 10 gene sequences that shared similarity and conserved domains with 238 known OYEs. The presence of multiple OYE genes appears to be a common feature not 239 only among Ascomycetes and Basiomycetes [7, 12] but also in Zygomycetes: indeed 240 with a similar approach we found from 4 to 10 putative OYE sequences within some of 241 the completely sequenced genomes (Suppl. Table 2). 242 A phylogenetic analysis grouped the McOYEs in two classes: nine proteins (McOYE1-243 9) were placed in Class I, including most of the OYE1-like proteins [5, 9, 12], while 244 McOYE10 clustered with Class II. Genome sequence data allowed to hypothesize that a 245 number of Class I McOYEs are located within the same chromosome; this information 246 may suggest duplication events for some of these genes, as suggested by Corrochano et 247 al. [14]. Class II gathers OYEs originally identified from different thermophilic bacteria

248 [5, 9]; however, the recent work by Nizam et al. [9, 12] demonstrated that a number of 249 sequences, although not yet characterized, from filamentous fungi (Ascomycota and 250 Basidiomycota) also clusters within Class II. To the best of our knowledge, this is the 251 first report of an OYE homologue from a Zygomycota belonging to this Class. 252 The fungal enzymatic activity was analyzed in the presence of three different substrates 253 while previous works considered only one substrate or a series of compounds belonging 254 to the same chemical class [15]. M. circinelloides showed a strong enzymatic activity 255 being able to completely reduce the C=C double bond of the three substrates. CE was 256 converted very fast (3.5 h), followed by MCA (20 h) and MCI (66 h), suggesting an 257 increasing recalcitrance of the molecules. These results are in line with those obtained 258 by Gatti et al. [2], who demonstrated that the carbonyl moiety acts as a strong activator, 259 while the ester group is a weak EWG. Being able to convert compounds with different 260 EWGs, M. circinelloides was very versatile; during the biotransformation the EWG 261 influenced only the timing of the reaction; the ester group of MCI was the weakest 262 EWG as the reaction was accomplished in 66 h. 263 The reduction of α,β -unsaturated ketones has been extensively studied using either the 264 whole microorganism or the purified enzymes [5, 15, 16]. Generally CE is a well 265 reduced compound; in fact M. circinelloides completely reduced the C=C double bond 266 (100 %) in only 3.5 h. Comparable yields were achieved with other filamentous fungi: a 267 previous study, which examined 28 filamentous fungi for the reduction of three 268 different conventional compounds, showed that CE was the easiest to reduce for almost 269 all the fungi (96.4 %); in particular, 19 fungi completely reduced this molecule [13]. 270 Stueckler et al. [7] reported that purified OYE1 (S. pastorianus) reduced 92 % of CE 271 and purified YqjM (OYE from Bacillus subtilis) reduced 85 % of CE. 272 The reduction of α -substituted cinnamaldehydes is very important at industrial level [2]. 273 Aldehyde is considered a good EWG and MCA was completely reduced within 20 h;

274 Fardelone et al. [17] obtained comparable yields using a commercial strain of S. 275 cerevisiae in the biotransformation of cinnamaldehyde derivatives. Other authors 276 reported that MCA is not always an easily reduced compound. For instance, Goretti et 277 al. [18] analyzed different non conventional yeasts in the reduction of MCA and found 278 that only *Kazachstania spenceroum* was able to convert this substrate with a yield of 60 279 %. Romagnolo et al. [13] reported that, among 19 fungi tested, only two, belonging to 280 the *Mucor* genus were able to completely convert the C=C double bond of this 281 substrate. 282 The bioreduction of MCI and its derivatives is not frequently reported in the literature, 283 suggesting a possible recalcitrance of this molecule to OYE-mediated 284 biotransformation. A biotransformation study performed on 7 bacterial, yeast and plant 285 OYEs homologues showed a conversion rate of MCI < 1 % [19]. Therefore, the ability 286 of *M. circinelloides* to completely reduce MCI is remarkable, since unsaturated esters 287 with no other EWG are rarely converted by OYEs. 288 BlastP analysis, using OYE1 of S. pastorianus as query, allowed the identification of 10 289 putative genes coding for OYEs, confirmed by PCR amplification and sequencing. The 290 high versatility found in the reduction of different compounds by M. circinelloides may 291 depend on its enzymatic pattern and on the possibility to activate distinct genes 292 specifically in the presence of different molecules or in defined environmental 293 conditions. In a recent paper, Nizam et al. [9] performed a genome-wide analysis on 294 available genomes of filamentous fungi: 60 species were investigated leading to the 295 identification of 424 OYE homologues. Surprisingly, some species were shown to 296 possess up to 22 OYEs homologues in their genome, while, in other microorganisms the 297 number of OYEs homologues number was more exiguous: only two homologues are 298 present in S. cerevisiae, while there are four in Shewanella oneidensis [20, 21].

299 Gene activation upon exposure to CE and MCA was extremely high (i.e. up to 2,900 300 fold for mcove2 in presence of MCA) and occurred soon after substrate addition. Nizam 301 et al. [9, 12] monitored the expression profile of 6 OYEs homologues from the 302 Ascochyta rabiei in two different conditions reporting an increase of 80 fold in 303 transcript levels during plant infection and a weaker activation during oxidative stress. 304 Among the 10 genes identified in M. circinelloides, mcoye1 and mcoye2 showed the 305 highest degree of gene activation (70-2,900 fold), followed by mcoye4, mcoye5 and 306 mcove10 (20-800 fold). Mcove3, mcove6 and mcove9 were poorly activated, while 307 transcripts of mcoye7 and mcoye8 were never activated in each condition. On the basis 308 of these results it seems reasonable to conclude that 8 out of 10 putative OYEs 309 homologues are rapidly activated in response to the substrates addition. 310 A relation between the biotransformation of each substrate and the expression profile of 311 the eight putative OYEs homologues has been observed. Generally, the transcript levels 312 reached the maximum peak before the beginning of the C=C double bond reduction. For 313 example, during CE analysis, the maximum peak of expression of mcoye2 was reached 314 after 1 h when 20 % of substrate was reduced. 315 The biological role of these enzymes as well as their cell localization is still an open question. By in silico analysis Nizam et al. [9], found that the majority of the OYE 316 homologues were allegedly located in the cytoplasm and in the cytoskeleton, although 317 318 some of them were associated to other cell compartments such as nucleus, peroxisomes, 319 plasma membrane. Only three OYE seemed to be extracellular. A preliminary 320 experiment carried out on M. circinelloides during the biotransformation of CE, showed 321 that ene reductase activity was detected only in presence of cell debris indicating that 322 these enzymes may be intracellular (data not shown); further and deeper experiments 323 are needed to confirm this hypothesis.

- 324 Studies are in progress to analyze the secondary and tertiary structure of these enzymes
- by in silico approaches [22]. In order to purify and catalytically characterize McOYEs,
- efforts will concentrate on the production of the homologues of *M. circinelloides* by
- 327 heterologous expression systems.

4. Materials and methods

329 4.1 Fungal strain

328

- 330 Mucor circinelloides 277.49 was obtained from CBS (CBS-KNAW fungal biodiversity
- centre) and was selected due to its capability of reducing C=C double bonds [13]. The
- 332 strain is preserved as MUT 44 at the Mycotheca Universitatis Taurinensis (MUT),
- 333 Department of Life Sciences and Systems Biology, University of Turin.
- *4.2 Chemicals*
- 335 CE, MCA and MCI were purchased from Sigma-Aldrich. Stock solutions of 500 mM of
- each substrate were prepared in DMSO (Sigma-Aldrich).
- 337 (E)-2-methyl-3-phenylprop-2-en-1,1- d_2 -1-ol (dideuterated α -methylcinnamyl alcohol)
- was prepared by reduction of ethyl (*E*)-2-methyl-3-phenylacrylate (0.50 g, 2.6 mmol)
- with DIBAL-D (7.9 mmol, 0.7 M in toluene) in THF. After the usual work-up, the
- dideuterated compound was obtained (0.41 g, 2.3 mmol, 89 %). ¹H NMR (CDCl₃, 400
- 341 MHz): $\delta = 7.39 7.19$ (5H, m, aromatic hydrogens), 6.53 (1H, q, J = 1.5 Hz, CH=C),
- 342 1.91 (3H, d, J = 1.5 Hz, CH3); GC-MS (EI) $t_R = 14.1$ min: m/z (%) = 150 (M⁺, 92), 107
- 343 (68), 91 (100). 2-Methyl-3-phenylpropan-1-d-1-ol (monodeuterated α -
- methyldihydrocinnamyl alcohol) was isolated from the reaction medium and
- 345 characterized by NMR and GC/MS analysis: 1 H NMR (CDCl₃, 400 MHz): $\delta = 7.37 -$
- 346 7.13 (5H, m, aromatic hydrogens), 3.45 (1H, m, *CH*DOH), 2.75 (1H, dd J = 13.5 and
- 347 6.4 Hz, CHHPh), 2.43 (1H, dd J = 13.5 and 8.0 Hz, CHHPh), 1.97 (1H, m, CHCH₃),
- 348 0.92 (3H, s, CH_3); GC-MS (EI) $t_R = 12.6 \text{ min: m/z}$ (%) = 151 (M⁺, 10), 133 (23), 118
- 349 (27), 91 (100).

350 4.3 Genome mining and phylogenetic analyses 351 BlastP analysis was performed on the complete genome of M. circinelloides strain 352 277.49 (Joint Genome Institute, JGI: http://jgi.doe.gov) using the sequence of OYE1 of 353 Saccharomyces pastorianus (UniProtKB accession no. Q02899) as query. Primer pairs 354 for qRT-PCR assays were designed by using Primer 3 (http://primer3.ut.ee/) (Supp. 355 Table 1). Total genomic DNA was extracted from the mycelium grown in MEA liquid 356 medium (20 g/l glucose, 20 g/l malt extract, 2 g/l peptone) for 24 h using the CTAB 357 method [23]. Oligonucleotides were tested by conventional PCR on genomic DNA. The 358 PCR mixture included distilled water, PCR buffer (10 X), 1 mM deoxynucleotide 359 triphosphates (dNTPs), 10 mM of each primer, 0.5 U of DNA polymerase (Tag DNA 360 polymerase, Qiagen) and 100 ng of genomic DNA in a total volume of 20 µl. 361 Amplifications were performed using a T100 Thermal Cycler (BIORAD). For the 362 validation of mcoye1 F-R, mcoye2 F-R, mcoye3 F-R, mcoye5 F-R, mcoye6 F-R, mcoye9 363 F-R e mcoye 10 F-R, the amplification protocol was as follows: 95 °C (5 min), 34 cycles 364 of 95 °C (40 sec), 60 °C (50 sec) and 72 °C (50 sec), 72 °C (8 min). For the detection of 365 mcoye4 F-R, mcoye7 F-R e mcoye8 F-R the amplification protocol was as follows: 95 366 °C (5 min), 34 cycles 95 °C (40 sec), 56 °C (50 sec) and 72 °C (50 sec), 72 °C (8 min). 367 PCR products were loaded on a 1.5 % agarose electrophoresis gel stained with ethidium 368 bromide; the molecular weight marker used was the GelPilot 1 kb Plus Ladder (cat. no. 369 239095, Qiagen). Products were purified and sequenced at Macrogen (The 370 Netherlands). Newly generated sequences were analyzed using Sequencher 5.4 (Gene 371 Code Corporation). 372 To perform the phylogenetic analyses, over 400 OYEs aminoacidic sequences of fungi 373 were aligned with MUSCLE (http://www.ebi.ac.uk/Tools/msa/muscle/) using default 374 conditions for gap openings and gap extension penalties and trimmed by TrimAl (v 1.2) 375 (http://trimal.cgenomics.org) with the AUTOMATED 1 setting. The analysis was

376 performed using two approaches. First, a phylogenetic tree was derived by Bayesian 377 Inference (BI) implemented in MrBayes (v 3.2.2) (http://mrbayes.sourceforge.net) 378 under a mixed amino acid substitution model. The alignment was run over 10 million 379 generations with two independent runs each containing four Markov Chains Monte 380 Carlo (MCMC) and sampling frequency of every 300 iterations. The first 2,500 trees 381 were discarded as "burn-in" (25 %). Using the Sumt function of MrBayes a consensus 382 tree was generated and posterior probabilities were estimated. In a second approach, 383 Maximum Likelihood (ML) was performed using RAxML GUI (v 1.5 b) [24] with 384 WAG+I+G model. Statistical reliability was determined by Bootstrap analysis. All the 385 phylogenetic trees were visualized using FigTree (v 1.4) 386 (http://tree.bio.ed.ac.uk/software/figtree). 387 4.4 Biotransformation by whole cell system 388 A conidia suspension of *M. circinelloides* was made from pre-growth mycelium in 389 MEA solid medium (same composition of MEA liquid with the addition of 20 g/l of 390 agar). 10⁶ conidia were inoculated in 100 ml flasks containing 40 ml of MEA liquid 391 medium. Flasks were incubated at 25 °C in agitation. After 2 days, substrates were 392 added (5 mM final concentration), each cultural line was run in triplicate. In addition, 393 biotic controls (in absence of substrates) were set up. 394 According to previous results (unpublished data), the conversion of CE, MCA and MCI 395 was followed for 24 h, 48 h and 7 d, respectively. Every 2 h, 1 ml of broth and 100 mg 396 of biomass were collected to perform chemical analysis and RNA extraction. 397 respectively. The mycelium was frozen in liquid nitrogen and stored at - 80 °C until the 398 analysis. 399 At any collection time point, pH and glucose content were measured. The concentration 400 of reducing sugars was obtained following the reaction with 3,5-dinitrosalycilic acid 401 assay (DNS) [25], using a modified protocol as described by Spina et al. [26]. At each

- 402 time point and at the end of the experiment, fungal biomasses were separated from the
- 403 culture medium by filtration and dried in oven at 60 °C for 24 h to calculate the dry
- 404 weight.
- 405 *4.5 Chemical analyses*
- Samples taken at the different time points were extracted by two-phase separation using
- 407 0.4 ml of methyl t-butyl ether (MTBE) as solvent; the organic phase was dried over
- anhydrous Na₂SO₄ and analyzed by GC/MS.
- 409 GC/MS analyses were performed on an Agilent HP 6890 gas chromatograph equipped
- with a 5973 mass detector and an HP-5-MS column (30 m \times 0.25 mm \times 0.25 μ m,
- 411 Agilent), employing the following temperature program: 60 °C (1 min) / 6 °C min⁻¹ /
- 412 $150 \,^{\circ}\text{C} \, (1 \,^{\circ}\text{min}) / 12 \,^{\circ}\text{C} \,^{\circ}\text{min}^{-1} / 280 \,^{\circ}\text{C} \, (5 \,^{\circ}\text{min})$. The end products of the
- biotransformations were identified by GC/MS analysis, using authentic commercial
- samples as reference compounds: i) cyclohexenone $t_R = 5.40 \text{ min } m/z \text{ 96 (M}^+, 33), 81$
- 415 (19), 68 (100); cyclohexanone $t_R = 4.65 \text{ min } m/z 98 \text{ (M}^+, 47), 83 (13), 55 (100);$
- 416 cyclohexanol $t_R = 4.45 \text{ min } m/z \ 100 \ (M^+, 2), 82 \ (35), 57 \ (100), ii) \alpha$
- 417 methylcinnamaldehyde $t_R = 14.7 \text{ min } m/z \ 146 \ (M^+, 64), 145 \ (100), 117 \ (79), 91 \ (43);$
- 418 α -methylcinnamyl alcohol $t_R = 15.5 \text{ min } m/z 148 (M^+, 50), 115 (63), 91 (100); <math>\alpha$ -
- 419 methyldihydrocinnamyl alcohol $t_R = 13.7 \text{ min } m/z 150 \text{ (M}^+, 12), 117 \text{ (62)}, 91 \text{ (100)}; iii)$
- 420 methyl cinnamate $t_R = 16.03 \text{ min } m/z \ 162 \ (\text{M}^+, 58), \ 131 \ (100), \ 103 \ (72); \ \text{cinnamyl}$
- 421 alcohol $t_R = 12.80 \text{ min } m/z \ 134 \ (M^+, 53), \ 115 \ (65), \ 92 \ (100)$; phenylpropanol $t_R = 12.36$
- 422 min m/z 136 (M⁺, 21), 117 (100), 91 (84).
- 4.6 RNA extraction, first strand cDNA synthesis and quantitative Real-Time PCR
- 424 experiments
- The extraction of RNA was performed from about 100 mg of fungal biomass using the
- 426 RNeasy Plant Mini Kit (Qiagen). Quantity and quality of RNA samples were checked
- spectrophotometrically (Tecan Infinite 200, i-control software). After DNase treatment

428	(TURBO DNA-free, Ambion), RNA quality has been tested again and for all the
429	samples, the ratios of absorbance 260/280 were between 1.8 and 2.2. Subsequently they
430	were processed to obtain cDNA with the use of the Super-Script II Reverse
431	Transcriptase (Invitrogen), following instructions.
432	qRT-PCR were performed with an iCycler iQTM Real-Time PCR Detection System
433	(BIORAD); reactions were carried out in a final volume of 15 μ l by using iTaq
434	Universal SYBR GREEN Supermix (BIORAD), specific primers (3 μM ; Table 1) and
435	cDNA . For the detection of mcoye1, mcoye2, mcoye3, mcoye5, mcoye6, mcoye9 and
436	mcoye10, the amplification protocol was as follows: 95 °C (1.5 min), 40 cycles of 95 °C
437	(15 sec), 60 °C (30 sec) and 72 °C (50 sec), 72 °C (8 min). For the detection of mcoye4,
438	mcoye7 and mcoye8 the amplification protocol was as follows: 95 °C (1.5 min), 40
439	cycles 95 °C (15 sec), 56 °C (30 sec) and 72 °C (50 sec), 72 °C (8 min). The <i>M</i> .
440	circinelloides β-actin encoding gene was used as internal control [27]. The relative
441	expression was calculated using the $2^{-\Delta\Delta Ct}$ method [28]. One-way ANOVA and Tukey's
442	tests (p \leq 0.05) were performed to assess the statistical significance of the gene
443	expression data (IBM SPSS Statistics for Macintosh, Version 22.0).
444	4.7 Availability of materials and data
445	Authors confirm that all relevant data are included in the article and its supplementary
446	information file.
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451	References

- 452 [1] Stuermer, R., Hauer, B., Hall, M. & Faber, K. Asymmetric bioreduction of activated
- 453 C=C bonds using enoate reductases from the old yellow enzyme family. Curr. Opin.
- 454 Chem. Biol. 11, 203–213 (2007).
- 455 [2] Gatti, F. G., Parmeggiani, F. & Sacchetti, A. Synthetic strategies based on C=C
- 456 bioreductions for the preparation of biologically active molecules. In: Synthetic
- methods for biologically active molecules: exploiting the potential of bioreductions, 1st
- ed., Wiley WHC, Weinheim; 2013.
- 459 [3] Yang, J. W., Fonseca, M. T. H., Vignola, N. & List, B. Metal-free, organocatalytic
- asymmetric transfer hydrogenation of α,β -unsaturated aldehydes. Angew. Chem. Intern.
- 461 Ed. 44, 108–110 (2005).
- 462 [4] Soetaert, W. & Vandamme, E. J. The scope and impact of industrial biotechnology.
- In: Industrial biotechnology: sustainable growth and economic success, 1st ed. Wiley-
- 464 VHC, Weinheim; 2010.
- 465 [5] Toogood, H. S., Gardiner, J. M. & Scrutton, N. S. Biocatalytic reductions and
- chemical versatility of the old yellow enzyme family of flavoprotein oxidoreductases.
- 467 ChemCatChem 2, 892-914 (2010).
- 468 [6] Robinson, S. L. & Panaccione, D. G. Diversification in ergot alkaloids in natural and
- 469 modified fungi. Toxins 7, 201-218 (2015).
- 470 [7] Stueckler, C., Reiter, T. C., Baudendistel, N. & Faber, K. Nicotinamide-indipendent
- asymmetric bioreduction of C=C bonds via disproportionation of enones catalyzed by
- 472 enoate reductases. Tetrahedron 66, 663-667 (2010).
- 473 [8] Huang, M., Hu, H., Ma, L., Zhou, Q., Yu, L. & Zheng, S. Carbon-carbon double-
- bond reductases in nature. Drug. Metab. Rev. 46, 362-378 (2015).
- 475 [9] Nizam, S., Verma, S., Borah, N. N., Gazara, R. K. & Verma, P. K. (a)
- 476 Comprehensive genome-wide analysis reveals different classes of enigmatic old yellow
- 477 enzyme in fungi. Sci. Rep. 4, 4013 (2014).

- 478 [10] Brandl, J. & Andersen, M. R. Current state of genome-scale modelling in
- 479 filamentous fungi. Biotechnol. Lett. 37, 1131-1139 (2015).
- 480 [11] Cheng, J. Z., Coyle, C. M., Panaccione, D. G. & O'Connor, S. E. A role for Old
- 481 Yellow Enzyme in ergot alkaloids biosynthesis. J. Am. Chem. Soc. 132, 1776-1777
- 482 (2010).
- 483 [12] Nizam, S., Gazara, R. K., Verma, S., Singh, K. & Verma, P. K. Comparative
- 484 structural modeling of six Old Yellow Enzymes (OYEs) from the necrotrophic fungus
- 485 Ascochyta rabiei: insight into novel OYE Classes with differences in cofactor binding,
- organization of active site residues and stereopreferences. Plos One 9 (2014).
- 487 [13] Romagnolo, A., Spina, F., Brenna, E., Crotti, M., Parmeggiani, F. & Varese, G. C.
- 488 Identification of fungal ene-reductase activity by means of a functional screening. Fung.
- 489 Biol. 119, 487-493 (2015).
- 490 [14] Corrochano, L. M., Kuo, A., Marcel-Houben, M., Polaino, S., Salamov, A., et al.
- Expansion of signal transduction pathways in fungal by extensive genome duplication.
- 492 Curr. boil. 26, 1577-1584 (2016).
- 493 [15] Zagozda, M. & Plenkiewicz, J. Enantioselective reduction of α,β-unsaturated
- ketones by Geotrichum candidum, Mortierella isabellina and Rhodotorula rubra yeast.
- 495 Tetrahedron: Asymm. 17, 1958-1962 (2006).
- 496 [16] Hall, M., Stueckler, C., Hauer, C., Stuermer, R., Friedrich, T., Breuer, M., Kroutil,
- W. & Faber, K. Asymmetric bioreduction of activated C=C bonds using Zymomonas
- 498 mobilis NCR enoate reductase and old yellow enzymes OYE 1-3 from yeasts. Eur. J.
- 499 Org. Chem. 1511-1516 (2008).
- 500 [17] Fardelone, L. C., Rodrigues, A. R. & Moran, P. J. S. Baker's yeast mediated
- asymmetric reduction of cinnamaldehyde derivatives. J. Mol. Catal. B: Enz. 29, 41-45
- 502 (2003).

- 503 [18] Goretti, M., Ponzoni, C., Caselli, E., Marchegiani, E., Cramarossa, M. R.,
- 504 Turchetti, B., Forti, L. & Buzzini, P. Bioreduction of α,β-unsaturated ketones and
- aldehydes by non-conventional yeast (NCY) whole-cell. Biores. Technol. 102, 3993-
- 506 3998 (2011).
- 507 [19] Tasnadi, G., Winkler, C. K., Clay, D., Sultana, N., Fabian, W. M. F., Hall, M.,
- 508 Ditrich, K. & Faber, K. A substrate-driven approach to determine reactivities of α,β-
- 509 unsaturated carboxylic esters towards asymmetric bioreduction. Chem. Eur. J. 18,
- 510 10362-10367 (2012).
- 511 [20] Niino, Y. S., Chakraborty, S., Brown, B. J. & Massey, V. A new Old Yellow
- Enzyme of Saccharomyces cerevisiae. J. Biol. Chem. 27, 1983-1991 (1995).
- 513 [21] Brigé, A., Van Den Hemel, D., Carpentier, W., De Smet, L. & Van Beeumen, J. J.
- 514 Comparative characterization and expression analysis of the four Old Yellow Enzyme
- 515 homologues from *Shewanella oneidensis* indicate differences in physiological function.
- 516 Biochem. J. 394, 335-344 (2006).
- 517 [22] Catucci, G., Romagnolo, A., Spina, F., Varese, G. C., Gilardi, G. & Di Nardo, G.
- Enzyme-substrate matching in biocatalysis: *in silico* studies to predict substrate
- preference of ten putative ene-reductases from *Mucor circinelloides* MUT 44. J. Mol.
- 520 Catal. B: Enz. 131, 94-100 (2016).
- 521 [23] Gardes, M. & Bruns, T. D. ITS Primer with Enhanced Specificity for
- Basidiomycetes-Application to the Identification of Mycorrhizae and Rusts. Mol. Ecol.
- 523 2, 113-118 (1993).
- 524 [24] Stamatakis, A. RAxML Version 8: A tool for Phylogenetic Analysis and Post-
- Analysis of Large Phylogenies. Bioinformatics 30, 1312-1313 (2014).
- 526 [25] Miller, G. L. Use of DNS reagent for the measurement of reducing sugar. Anal.
- 527 Chem. 31, 426-428 (1959).

- 528 [26] Spina, F., Fidaleo, M., Nanni, A., Romagnolo, A. & Varese, G. C. Fungal Laccases
- 529 production using tomato-based medium: a factorial design approach. Environ. Eng.
- 530 Manag. J. 14, 1743-1750 (2015).
- 531 [27] Lee, S. C., Li, A., Calo, S. & Heitman, J. Calcineurin plays key roles in the
- 532 dimorphic transition and virulence of the human pathogenic zygomycete Mucor
- 533 *circinelloides*. Plos. Path. 9, 1-20 (2013).
- [28] Livak, K. J. & Schmittgen, T. D. Analysis of relative gene expression data using
- realtime quantitative PCR and the 2^-ΔΔCt method. Methods 25, 402–408 (2001).

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- 539 Author Contribution Statement
- 540 **A.R., F.S.:** wrote the manuscript
- **A.P.:** performed phylogenetic analysis
- **A.R., F.S., S.R., B.S.:** performed lab experiments
- 543 L.L.: gene expression experimental design
- 544 M.C., D.M., E.B.: performed chemical analysis and data curation
- 545 **G.C.V:** project administration and supervision
- All authors reviewed the manuscript.

547

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- expression analyses.
- The authors declare no competing financial interests.

Figure and figure legends

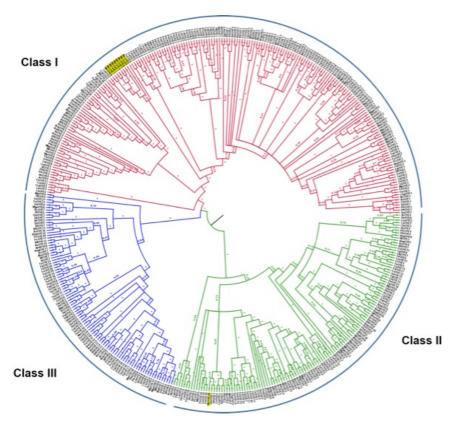


Figure 1. Evolutionary relationship of deduced OYE proteins based on Bayesian inference analysis of the structure-based amino acid sequence alignment. The numbers at the nodes indicates Bayesian posterior probabilities. The phylogenetic tree was implemented from Nizam et al., 2014 [9].

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 α -methyldihydrocinnamaldehyde α -methyldihydrocinnamyl alcohol

Figure 2. Reaction profiles of (A) CE, (B, C) MCA and (D) MCI biotransformations.

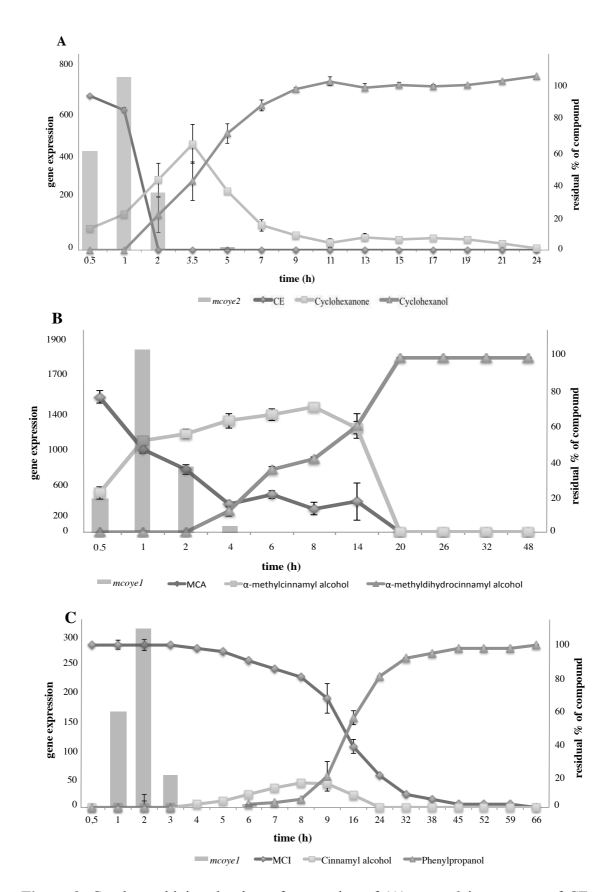


Figure 3. Graph combining the data of expression of (A) *mcoye2* in presence of CE (bars) with the biotransformation data of CE (lines); (B) *mcoye1* in presence of MCA (bars) with the biotransformation data of MCA (lines); (C) *mcoye1* in presence of MCI

- 567 (bars) with the biotransformation data of MCI (lines). Data are the averages \pm standard
- deviations (error bars) of the results of at least three different biological replicates.

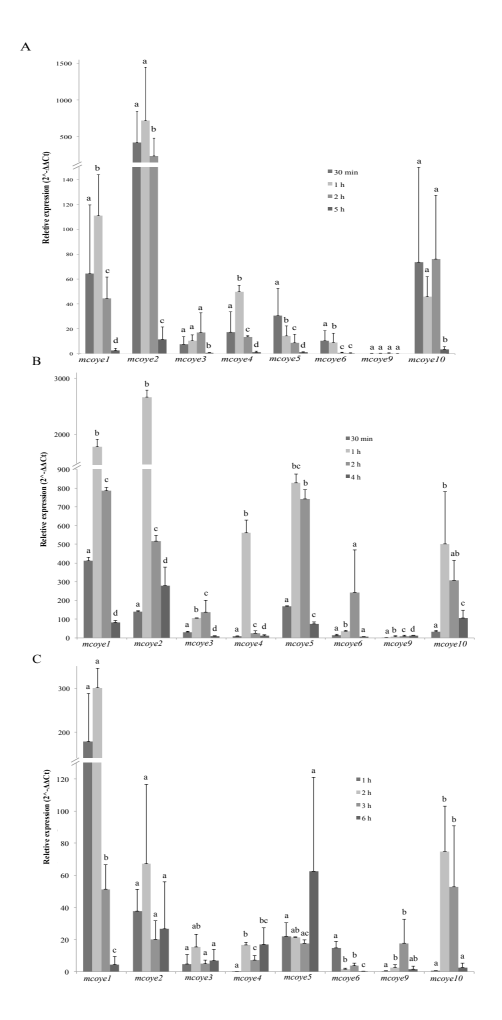


Figure 4. Gene expression of OYE homologues in presence of (A) CE, (B) MCA and CC MCI during the time course experiments. The relative gene expression was calculated with the $2^{-\Delta\Delta Ct}$ method according to Livak & Schmittgen [27] using the β -actin as housekeeping gene [26] and the control (non treated) as reference sample. Different letters indicate statistically significant difference (p < 0.05, ANOVA and Tukey's tests) for each gene at the different time points.