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1 **Distribution of methionine sulfoxide reductases in fungi and conservation of the free-**
2 **methionine-*R*-sulfoxide reductase in multicellular eukaryotes**

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5

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9

10 **Running title:** Methionine sulfoxide reductases in fungi

11

12 **Keywords:** fungi, genome, horizontal gene transfer, methionine sulfoxide, methionine sulfoxide
13 reductase, protein oxidation, thiol oxidoreductase.

14

15 **Highlights:**

- 16 • Free and protein-bound methionine can be oxidized into methionine sulfoxide (MetO).
- 17 • Methionine sulfoxide reductases (Msr) reduce MetO in most organisms.
- 18 • Sequence characterization and phylogenomics revealed strong conservation of Msr in fungi.
- 19 • fRMs_r is widely conserved in unicellular and multicellular fungi.
- 20 • Some *msr* genes were acquired from bacteria via horizontal gene transfers.

21

22 **Abstract**

23 Methionine, either as a free amino acid or included in proteins, can be oxidized into methionine sulfoxide
24 (MetO), which exists as *R* and *S* diastereomers. Almost all characterized organisms possess thiol-
25 oxidoreductases named methionine sulfoxide reductase (Msr) enzymes to reduce MetO back to Met.
26 MsrA and MsrB reduce the *S* and *R* diastereomers of MetO, respectively, with strict stereospecificity
27 and are found in almost all organisms. Another type of thiol-oxidoreductase, the free-methionine-*R*-
28 sulfoxide reductase (fRMsr), identified so far in prokaryotes and a few unicellular eukaryotes, reduces
29 the *R* MetO diastereomer of the free amino acid. Moreover, some bacteria possess molybdenum-
30 containing enzymes that reduce MetO, either in the free or protein-bound forms. All these Msrs play
31 important roles in the protection of organisms against oxidative stress. Fungi are heterotrophic
32 eukaryotes that colonize all niches on Earth and play fundamental functions, in organic matter recycling,
33 as symbionts, or as pathogens of numerous organisms. However, our knowledge on fungal Msrs is still
34 limited. Here, we performed a survey of *msr* genes in almost 700 genomes across the fungal kingdom.
35 We show that most fungi possess one gene coding for each type of methionine sulfoxide reductase:
36 MsrA, MsrB, and fRMsr. However, several fungi living in anaerobic environments or as obligate
37 intracellular parasites were devoid of *msr* genes. Sequence inspection and phylogenetic analyses allowed
38 us to identify non-canonical sequences with potentially novel enzymatic properties. Finally, we identified
39 several occurrences of *msr* horizontal gene transfer from bacteria to fungi.

40

41 **1. Introduction**

42 Life in the presence of dioxygen necessarily exposes the biological systems to oxidant
43 molecules. Due to its high reactivity, dioxygen can be converted to reactive oxygen species (ROS),
44 which play key roles in physiological and pathological contexts through redox modifications of
45 macromolecules [1]. Met, either as a free amino acid or as a residue included in a protein, can be oxidized
46 into Met sulfoxide (MetO) by the addition of an oxygen atom to the sulfur of the lateral chain. MetO
47 exists as diastereomer *R* or *S* (Met-*R*-O and Met-*S*-O, respectively) [2], and can be reduced back to Met
48 by the action of oxidoreductases called methionine sulfoxide reductases (Msr). The two main types are
49 the (seleno)thiol-containing MsrA and MsrB, which display strict stereoselectivities toward the *S*- and
50 the *R*-diastereomer of MetO, respectively [3–5]. Whereas MsrA reduces free MetO and protein-bound
51 MetO with similar catalytic efficiencies, MsrB generally only reduces efficiently the protein-bound
52 MetO [6]. Another enzyme, fRMs (free methionine-(R)-sulfoxide reductase) specifically reduces the
53 free form of Met-*R*-O [7–9]. Some bacteria also possess several molybdoenzymes that reduce
54 exclusively the free MetO [10,11] or both the free and protein-bound MetO [12–15]. Despite the lack of
55 sequence and structure similarities, MsrA, MsrB and fRMs generally catalyze the reduction of MetO
56 using a similar 3-steps-mechanism [9,16] : i) a ‘catalytic’ Cys (or, less frequently, a selenocysteine, Sec)
57 reduces the target MetO and is converted into a sulfenic (or selenic) acid [17], ii) an internal ‘resolving’
58 Cys reduces it through the formation of an intramolecular disulfide bond, and finally iii) the oxidized
59 Msr is regenerated through disulfide exchange with a thioredoxin [18]. Variations of this mechanism
60 exist, with some Msrs devoid of any resolving Cys, in which the sulfenic acid is directly reduced by an
61 external reducer [19–21]. Alternatively, in other Msrs, a disulfide exchange occurs with a second
62 internal resolving Cys before regeneration by the thioredoxin [7,22,23]. Genome mining analyses
63 indicated that *msrA* and *msrB* genes originated from early prokaryotes and are now present in all
64 organisms across the tree of life, with few exceptions [24–26]. Most organisms have few *msr* genes,
65 generally one of each type [24–26], but this number is generally higher in plants (e.g. up to 5 *msrA* and
66 9 *msrB* genes in *Arabidopsis thaliana* [25]). Moreover, some bacteria encode a bifunctional MsrA/MsrB
67 fusion and some others lack a MsrB, but no organism of any kind was so far described with only a MsrB
68 [24,26]. Finally, only very few organisms do not possess any *msr* gene at all, such as a few
69 endosymbiotic or obligatory parasitic bacteria and some archaea [24,26]. fRMs were so far only
70 reported in bacteria and a few unicellular eukaryotes [7–9]. Most interestingly, the only eukaryote for
71 which the absence of *msr* gene was observed is the fungus *Encephalitozoon cuniculi*, a microsporidium
72 having a remarkably reduced genome (~2.9 Mb) and living as intracellular parasite of mammals [24,27].
73 The conservation of typical Msrs and MsrBs in almost all known organisms argues for the critical role
74 of MetO reduction in cellular metabolism, and numerous studies showed that Msrs are involved in the
75 protection against oxidative stress and the regulation of protein functions. Schematically, Msr protective
76 roles against oxidative injuries occur through two main functions: i) the repair of oxidized proteins, and

77 ii) an antioxidant function through ROS scavenging by cyclic Met oxidation and reduction. Moreover,
78 the reversible Met oxidation was shown to act as a post-translational modification responsible for the
79 activation of enzymes and transcription factors or the regulation of protein-protein interactions [28–30].
80 The role of fRMsrs has been far less studied, but it is suspected to have an antioxidant function by
81 reducing free MetO and maintaining the pool of Met available for protein synthesis and for the
82 production of sulfur-containing metabolites [8,31]. Overall, these functions were well established in
83 animals, bacteria and plants, and have recently been reviewed [32–36].

84 Fungi are heterotrophic eukaryotes that colonize virtually all niches on Earth and exist as
85 unicellular or multicellular organisms. They can have numerous lifestyles, either as free living
86 organisms playing key roles in organic and inorganic matter cycling, or as symbionts or pathogens with
87 crucial impacts on plant and animal health [37]. As any other organism living in aerobic conditions and
88 exposed to environmental constraints, fungi are exposed to oxidative constraints and protein oxidation
89 [38]. However, the effects of Met oxidation and the roles of Msrs were largely overlooked in these
90 eukaryotes. The Msr system was mainly characterized in the yeast *Saccharomyces cerevisiae*, which
91 possesses one Msr of each type [6,8]. In *S. cerevisiae*, MsrA is located in the cytosol, MsrB is found in
92 mitochondria and in the cytosol and fRMsrs is located both in the cytosol and the nucleus [39,40]. Genetic
93 manipulations have shown that these Msrs are involved in the protection against oxidative stress and in
94 maintaining the yeast lifespan [6,8,41,42]. Consistently, the overexpression of *msrA* in the
95 basidiomycete *Pleurotus ostreatus* and of *msrB* in the yeast *Schizosaccharomyces pombe* increased the
96 resistance to oxidative constraints [43,44]. Moreover, the filamentous fungus *Aspergillus nidulans*
97 possesses one *msrA* and one *msrB* genes, and the deletion of one or both genes increased the sensitivity
98 of the fungus to oxidative treatments [45]. Few proteins with oxidized Met residues have been
99 characterized in fungi, but remarkably interesting findings were obtained. For instance, in *S. cerevisiae*,
100 the reversible oxidation of Met regulates the oligomerization state of the ataxin-2 protein and the activity
101 of the co-chaperone Fes1 [40,46]. Moreover, Met oxidation enhanced the activity of an α-galactosidase
102 in *Trichoderma reesei* [47]. Finally, in *A. nidulans*, it was demonstrated that the nuclear localization of
103 the nitrate-responsive transcription factor NirA was actively regulated through cyclic Met oxidation
104 [48]. These data indicate that, as for other organisms, fungal Msrs certainly play key roles under many
105 conditions of oxidative stress, such as biotic interactions or abiotic constraints.

106 In this study, we searched for *msr* genes in about 700 fungal genomes. We show that the great
107 majority of fungi have one *msrA* and one *msrB* genes. Moreover, we identified *fRmsr* genes in almost
108 all the fungi we analyzed and thereby demonstrate that the enzyme is conserved in these multicellular
109 eukaryotes. Finally, using a phylogenetic analysis and a close inspection of the sequence features, we
110 identified fungal Msrs with unusual sequence characteristics and uncovered horizontal gene transfers
111 from bacteria to fungi.

112 **2. Material and methods**

113 **2.1. Search for Msr homologs in fungi.**

114 The protein sequence of MsrA (Uniprot accession # C8Z745), MsrB (Uniprot accession # P25566) and
115 fRMsr (Uniprot accession # P36088) from *S. cerevisiae*, MsrP (Uniprot accession # P76342) and BisC
116 (Uniprot accession # P20099) from *Escherichia coli*, TorZ (Uniprot accession # P44798) from
117 *Haemophilus influenzae*, and DorA (Uniprot accession # Q57366) from *Rhodobacter sphaeroides* were
118 used as BLASTP and TBLASTN [49] queries to identify *msr* genes in 683 genomes available in the
119 MycoCosm database [50] (<https://mycocosm.jgi.doe.gov/mycocosm/home>).

120

121 **2.2. List of all explored fungal genomes.**

122 *Aaosphaeria arxii* CBS 175.79 v1.0; *Amniculicola lignicola* CBS 123094 v1.0; *Ampelomyces quisqualis*
123 *HMLAC05119* v1.0; *Aplosporella prunicola* CBS 121.167 v1.0; *Aulographum hederae* v2.0; *Bimuria*
124 *novae-zelandiae* CBS 107.79 v1.0; *Byssothecium circinans* CBS 675.92 v1.0; *Cercospora zae-maydis*
125 v1.0; *Clathrospora elynae* CBS 161.51 v1.0; *Cucurbitaria berberidis* CBS 394.84 v1.0; *Decorospora*
126 *gaudefroyi* v1.0; *Delitschia confertaspora* ATCC 74209 v1.0; *Delphinella strobiligena* CBS 735.71
127 v1.0; *Didymella exigua* CBS 183.55 v1.0; *Dissocoicum aciculare* v1.0; *Dothidotthia symphoricarpi*
128 v1.0; *Elsinoe ampelina* CECT 20119 v1.0; *Eremomyces bilateralis* CBS 781.70 v1.0; *Hortaea*
129 *acidophila* CBS 113389 v1.0; *Karstenula rhodostoma* CBS 690.94 v1.0; *Lentithecium fluviatile* v1.0;
130 *Lindgomyces ingoldianus* ATCC 200398 v1.0; *Lineolata rhizophorae* ATCC 16933 v1.0; *Lizonia*
131 *empirigonia* CBS 542.76 v1.0; *Lophiostoma macrostomum* v1.0; *Lophiotrema nucula* CBS 627.86 v1.0;
132 *Lophium mytilinum* CBS 269.34 v1.0; *Macroventuria anomochaeta* CBS 525.71 v1.0; *Massarina*
133 *eburnea* CBS 473.64 v1.0; *Massariosphaeria phaeospora* CBS 611.86 v1.0; *Melanomma pulvis-pyrius*
134 v1.0; *Microthyrium microscopicum* CBS 115976 v1.0; *Myriangium duriae* CBS 260.36 v1.0;
135 *Mytilinidion resinicola* CBS 304.34 v1.0; *Ophiobolus disseminans* CBS 113818 v1.0; *Patellaria atrata*
136 v1.0; *Phoma tracheiphila* IPT5 v1.0; *Piedraia hortae* CBS 480.64 v1.1; *Pleomassaria siparia* v1.0;
137 *Polychaeton citri* v1.0; *Polyplosphaeria fusca* CBS 125425 v1.0; *Pseudovirgaria hyperparasitica* CBS
138 121739 v1.0; *Rhizodiscina lignyota* CBS 133067 v1.0; *Saccharata proteae* CBS 121410 v1.0;
139 *Setomelanomma holmii* CBS 110217 v1.0; *Sporormia fimetaria* v1.0; *Teratosphaeria nubilosa* CBS
140 116005 v1.0; *Tothia fuscella* CBS 130266 v1.0; *Trematosphaeria pertusa* CBS 122368 v1.0;
141 *Trichodelitschia bisporula* CBS 262.69 v1.0; *Verruculina enalia* CBS 304.66 v1.0; *Viridothelium virens*
142 v1.0; *Westerdykella ornata* CBS 379.55 v1.0; *Zasmidium cellare* ATCC 36951 v1.0; *Zopfia rhizophila*
143 v1.0 [51]; *Acaromyces ingoldii* MCA 4198 v1.0; *Ceraceosorus guamensis* MCA 4658 v1.0; *Jaminaea*
144 sp. MCA 5214 v1.0; *Meira miltonrushii* MCA 3882 v1.0; *Pseudomicrostroma glucosiphilum* gen et sp.
145 nov. MCA 4718 v1.0; *Testicularia cyperi* MCA 3645 v1.0; *Tilletiopsis washingtonensis* MCA 4186 v1.0;

146 *Violaceomyces palustris* SA 807 v1.0 [52]; *Acidomyces richmondensis* BFW; *Acidomyces*
147 *richmondensis* BFW [53]; *Acremonium chrysogenum* ATCC 11550 [54]; *Agaricus bisporus* var *bisporus*
148 (*H97*) v2.0; *Agaricus bisporus* var. *burnettii* JB137-S8; *Gigaspora rosea* v1.0; *Rhizophagus*
149 *cerebriforme* DAOM 227022 v1.0; *Rhizophagus diaphanus* v1.0 [55]; *Amanita muscaria* Koide v1.0;
150 *Gymnopus luxurians* v1.0; *Hebeloma cylindrosporum* h7 v2.0; *Hydnomerulius pinastri* v2.0;
151 *Hypholoma sublateritium* v1.0; *Laccaria amethystina* LaAM-08-1 v2.0; *Paxillus adelphus* Ve08.2h10
152 v2.0; *Paxillus involutus* ATCC 200175 v1.0; *Piloderma croceum* F 1598 v1.0; *Pisolithus microcarpus*
153 441 v1.0; *Pisolithus tinctorius* Marx 270 v1.0; *Plicaturopsis crispa* v1.0; *Scleroderma citrinum* Foug A
154 v1.0; *Sebacina vermifera* MAFF 305830 v1.0; *Sphaerobolus stellatus* v1.0; *Suillus luteus* UH-Slu-Lm8-
155 n1 v3.0; *Tulasnella calospora* AL13/4D v1.0; *Oidiodendron maius* Zn v1.0 [56]; *Amanita thiersii*
156 Skay4041 v1.0 [57]; *Ambrosiozyma philentoma* NRRL Y-7523; *Candida boidinii* NRRL Y-2332;
157 *Citeromyces matritensis* NRRL Y-2407; *Nakazawaea wickerhamii* NRRL Y-2563; *Peterozyma xylosa*
158 NRRL Y-12939; *Saccharomycopsis capsularis* NRRL Y-17639; *Saturnispora disporea* NRRL Y-1447
159 [58]; *Amorphotheca resinae* v1.0; *Meliniomyces bicolor* E v2.0; *Meliniomyces variabilis* F v1.0;
160 *Rhizoscyphus ericae* UAMH 7357 v1.0 [59]; *Anaeromyces robustus* v1.0; *Neocallimastix californiae* G1
161 v1.0; *Piromyces finnis* v3.0; *Piromyces* sp. E2 v1.0 [60]; *Antonospora locustae* HM-2013 [61];
162 *Armillaria cepistipes* B5; *Armillaria gallica* 21-2 v1.0; *Armillaria ostoyae* C18/9; *Armillaria solidipes*
163 28-4 v1.0 [62]; *Armillaria mellea* DSM 3731 [63]; *Arthrobotrys oligospora* ATCC 24927 [64];
164 *Ascobolus immersus* RN42 v1.0; *Choiromyces venosus* 120613-1 v1.0; *Morchella importuna*
165 CCBAS932 v1.0; *Terfezia boudieri* ATCC MYA-4762 v1.1; *Tuber aestivum* var. *urcinatum* v1.0; *Tuber*
166 *borchii* Tbo3840 v1.0; *Tuber magnatum* v1.0 [65]; *Ascochyta rabiei* ArDII [66]; *Ascocoryne sarcoides*
167 NRRL50072 [67]; *Ascodesmis nigricans* CBS 389.68 v1.0 [68]; *Aspergillus aculeatinus* CBS 121060
168 v1.0; *Aspergillus brunneoviolaceus* CBS 621.78 v1.0; *Aspergillus costaricensis* CBS 115574 v1.0;
169 *Aspergillus ellipticus* CBS 707.79 v1.0; *Aspergillus eucalypticola* CBS 122712 v1.0; *Aspergillus*
170 *fijiensis* CBS 313.89 v1.0; *Aspergillus heteromorphus* CBS 117.55 v1.0; *Aspergillus homomorphus* CBS
171 101889 v1.0; *Aspergillus ibericus* CBS 121593 v1.0; *Aspergillus indologenus* CBS 114.80 v1.0;
172 *Aspergillus japonicus* CBS 114.51 v1.0; *Aspergillus neoniger* CBS 115656 v1.0; *Aspergillus niger*
173 (*lacticoffeatus*) CBS 101883 v1.0; *Aspergillus niger* (*phoenicis* Corda) Thom ATCC 13157 v1.0;
174 *Aspergillus niger* NRRL3; *Aspergillus niger* van Tieghem ATCC 13496 v1.0; *Aspergillus piperis* CBS
175 112811 v1.0; *Aspergillus saccharolyticus* JOP 1030-1 v1.0; *Aspergillus sclerotioricarbonarius* CBS
176 121057 v1.0; *Aspergillus sclerotioriger* CBS115572 v1.0; *Aspergillus uvarum* CBS 121591 v1.0;
177 *Aspergillus vadensis* CBS 113365 v1.0; *Aspergillus violaceofuscus* CBS 115571 v1.0; *Aspergillus*
178 *welwitschiae* CBS139.54b v1.0 [69]; *Aspergillus aculeatus* ATCC16872 v1.1; *Aspergillus brasiliensis*
179 v1.0; *Aspergillus carbonarius* ITEM 5010 v3; *Aspergillus glaucus* v1.0; *Aspergillus luchuensis* CBS
180 106.47 v1.0; *Aspergillus sydowii* CBS 593.65 v1.0; *Aspergillus tubingensis* v1.0; *Aspergillus versicolor*
181 v1.0; *Aspergillus wentii* v1.0; *Aspergillus zonatus* v1.0; *Penicillium chrysogenum* v1.0 [70]; *Aspergillus*
182 *bombycis* NRRL 26010 [71]; *Aspergillus calidoustus* [72]; *Aspergillus campestris* IBT 28561 v1.0;

183 *Aspergillus candidus* CBS 102.13 v1.0; *Aspergillus novofumigatus* IBT 16806 v1.0; *Aspergillus*
184 *ochraceoroseus* IBT 24754 v1.0; *Aspergillus steynii* IBT 23096 v1.0; *Aspergillus taichungensis* IBT
185 19404 v1.0; *Aspergillus albertensis* v1.0; *Aspergillus alliaceus* CBS 536.65 v1.0; *Aspergillus*
186 *arachidicola* v1.0; *Aspergillus avenaceus* IBT 18842 v1.0; *Aspergillus bertholletius* IBT 29228 v1.0;
187 *Aspergillus caelatus* CBS 763.97 v1.0; *Aspergillus coremiiformis* CBS 553.77 v1.0; *Aspergillus leporis*
188 CBS 151.66 v1.0; *Aspergillus minisclerotigenes* CBS 117635 v1.0; *Aspergillus nomius* IBT 12657 v1.0;
189 *Aspergillus novoparasiticus* CBS 126849 v1.0; *Aspergillus parasiticus* CBS 117618 v1.0; *Aspergillus*
190 *parvisclerotigenus* CBS 121.62 v1.0; *Aspergillus pseudocaelatus* CBS 117616 v1.0; *Aspergillus*
191 *pseudonomius* CBS 119388 v1.0; *Aspergillus pseudotamarii* CBS 117625 v1.0; *Aspergillus sergii* CBS
192 130017 v1.0; *Aspergillus tamarii* CBS 117626 v1.0; *Aspergillus transmontanensis* CBS 130015 v1.0
193 [73]; *Aspergillus clavatus* NRRL 1 from AspGD; *Aspergillus flavus* NRRL3357; *Aspergillus nidulans*;
194 *Aspergillus oryzae* RIB40; *Aspergillus terreus* NIH 2624; *Neosartorya fischeri* NRRL 181 [74];
195 *Aspergillus cristatus* GZAAS20.1005 [75]; *Aspergillus fumigatus* A1163 [76]; *Aspergillus fumigatus*
196 Af293 from AspGD [77]; *Aspergillus kawachii* IFO 4308 [78]; *Aspergillus niger* ATCC 1015 v4.0 [79];
197 *Aspergillus niger* CBS 513.88 [80]; *Aspergillus nomius* NRRL 13137; *Aspergillus ochraceoroseus*
198 SRRNC1432; *Aspergillus rambellii* SRRNC1468 [81]; *Aspergillus udagawae* IFM 46973 [82];
199 *Aureobasidium pullulans* var. *melanogenum* CBS 110374; *Aureobasidium pullulans* var. *namibiae* CBS
200 147.97; *Aureobasidium pullulans* var. *pullulans* EXF-150; *Aureobasidium pullulans* var. *subglaciale*
201 EXF-2481 [83]; *Auricularia subglabra* v2.0; *Coniophora puteana* v1.0; *Dichomitus squalens* LYAD-
202 421 SS1 v1.0; *Fomitiporia mediterranea* v1.0; *Fomitopsis pinicola* FP-58527 SS1 v3.0; *Gloeophyllum*
203 *trabeum* v1.0; *Punctularia strigosozonata* v1.0; *Stereum hirsutum* FP-91666 SS1 v1.0; *Trametes*
204 *versicolor* v1.0; *Wolfiporia cocos* MD-104 SS10 v1.0; *Dacryopinax primogenitus* DJM 731 SSP1 v1.0;
205 *Tremella mesenterica* Fries v1.0 [84]; *Auriculariopsis ampla* NL-1724 v1.0 [85]; *Baudoinia*
206 *compniacensis* UAMH 10762 (4089826) v1.0; *Cochliobolus heterostrophus* C4 v1.0; *Cochliobolus*
207 *heterostrophus* C5 v2.0; *Cochliobolus lunatus* m118 v2.0; *Cochliobolus sativus* ND90Pr v1.0;
208 *Hysterium pulicare*; *Rhytidhysteron rufulum*; *Septoria musiva* SO2202 v1.0; *Septoria populincola* v1.0;
209 *Setosphaeria turcica* Et28A v2.0 [86]; *Beauveria bassiana* ARSEF 2860 [87]; *Bjerkandera adusta* v1.0;
210 *Ganoderma* sp. 10597 SS1 v1.0; *Phlebia brevispora* HHB-7030 SS6 v1.0; *Neofusicoccum parvum*
211 UCRNP2; *Eutypa lata* UCREL1; *Phaeoacremonium aleophilum* UCRPA7 [88]; *Blastobotrys* (Arxula)
212 *adeninivorans* [89]; *Blastomyces dermatitidis* SLH14081 [90]; *Blumeria graminis* f. sp. *hordei* DH14;
213 *Blumeria graminis* f. sp. *hordei* Race1 [91]; *Blumeria graminis* f. sp. *tritici* 96224 [92]; *Botryobasidium*
214 *botryosum* v1.0; *Galerina marginata* v1.0; *Jaapia argillacea* v1.0; *Pleurotus ostreatus* PC15 v2.0;
215 *Ascoidea rubescens* NRRL Y17699 v1.0; *Babjeviella inositovora* NRRL Y-12698 v1.0; *Candida*
216 *arabinofermentans* NRRL YB-2248 v1.0; *Candida tanzawaensis* NRRL Y-17324 v1.0; *Cyberlindnera*
217 *jadinii* NRRL Y-1542 v1.0; *Hanseniaspora valbyensis* NRRL Y-1626 v1.1; *Hyphopichia burtonii* NRRL
218 Y-1933 v1.0; *Lipomyces starkeyi* NRRL Y-11557 v1.0; *Metschnikowia bicuspidata* NRRL YB-4993 v1.0;
219 *Nadsonia fulvescens* var. *elongata* DSM 6958 v1.0; *Ogataea polymorpha* NCYC 495 leu1.1 v2.0;

220 *Pachysolen tannophilus* NRRL Y-2460 v1.2; *Pichia membranifaciens* v2.0; *Tortispora caseinolytica* Y-
221 17796 v1.0; *Wickerhamomyces anomalus* NRRL Y-366-8 v1.0; *Saitoella complicata* NRRL Y-17804 v1.0
222 [93]; *Botryosphaeria dothidea* [94]; *Botrytis cinerea* v1.0 [95]; *Byssochlamys spectabilis* No. 5 [96];
223 *Candida albicans* SC5314 [97]; *Candida tenuis* NRRL Y-1498 v1.0; *Spathaspora passalidarum* NRRL
224 Y-27907 v2.0 [98]; *Capronia coronata* CBS 617.96; *Capronia epimyces* CBS 606.96; *Capronia*
225 *semiimmersa* CBS27337; *Cladophialophora bantiana* CBS 173.52; *Cladophialophora carrionii* CBS
226 160.54; *Cladophialophora immunda* CBS83496; *Cladophialophora psammophila* CBS 110553;
227 *Cladophialophora yegresii* CBS 114405; *Cyphellophora europaea* CBS 101466; *Exophiala*
228 *aquamarina* CBS 119918; *Exophiala mesophila* CBS40295; *Exophiala oligosperma* CBS72588;
229 *Exophiala sideris* CBS121828; *Exophiala spinifera* CBS89968; *Exophiala xenobiotica* CBS118157;
230 *Fonsecaea multimorphosa* CBS 102226; *Fonsecaea pedrosoi* CBS 271.37; *Coniosporium apollinis* CBS
231 100218; *Verruconis gallopava* [99]; *Cenococcum geophilum* 1.58 v2.0; *Glonium stellatum* CBS 207.34
232 v1.0; *Lepidopterella palustris* v1.0 [100]; *Ceriporiopsis (Gelatoporia) subvermispora* B [101];
233 *Chaetomium globosum* v1.0 [102]; *Chaetomium thermophilum* var *thermophilum* DSM 1495 [103];
234 *Cladonia grayi* Cgr/DA2myc/ss v2.0 [104]; *Cladosporium fulvum* v1.0; *Dothistroma septosporum*
235 NZE10 v1.0 [105]; *Cladosporium sphaerospermum* UM 843 [106]; *Clavispora lusitaniae* ATCC 42720;
236 *Lodderomyces elongisporus* NRRL YB-4239; *Meyerozyma guilliermondii* ATCC 6260 [107];
237 *Coccidioides immitis* RS ; *Coccidioides posadasii* C735 delta SOWgp; *Histoplasma capsulatum* NAm1;
238 *Uncinocarpus reesii* UAMH 1704 [108]; *Cochliobolus carbonum* 26-R-13 v1.0; *Cochliobolus*
239 *miyabeanus* ATCC 44560 v1.0; *Cochliobolus victoriae* FI3 v1.0 [109]; *Colletotrichum chlorophytii*
240 NTL11 [110]; *Colletotrichum fioriniae* PJ7 [111]; *Colletotrichum graminicola* M1.001 [112];
241 *Colletotrichum higginsianum* IMI 349063 [113]; *Colletotrichum incanum* MAFF 238712 [114];
242 *Colletotrichum nymphaeae* SA-01; *Colletotrichum salicis* CBS607.94; *Colletotrichum simmondsii*
243 CBS122122; *Trichoderma gamsii* T6085 [115]; *Colletotrichum orbiculare* 104-T [116]; *Colletotrichum*
244 *orchidophilum* IMI 309357 [117]; *Colletotrichum tofieldiae* 0861 [118]; *Conidiobolus coronatus*
245 NRRL28638 v1.0; *Coemansia reversa* NRRL 1564 v1.0; *Gonapodya prolifera* v1.0 [119]; *Coniochaeta*
246 *lignaria* NRRL 30616 v1.0 [120]; *Coniochaeta* sp. 2T2.1 v1.0 [121]; *Coniophora olivacea* MUCL
247 20566 v1.0 [122]; *Coprinellus micaceus* FP101781 v2.0; *Coprinopsis marcescibilis* CBS121175 v1.0;
248 *Crucibulum laeve* CBS 166.37 v1.0; *Dendrothele bispora* CBS 962.96 v1.0; *Helocybe sulcata*
249 OMC1185 v1.0; *Peniophora* sp. CONTA v1.0; *Pluteus cervinus* NL-1719 v1.0; *Polyporus arcularius*
250 v1.0; *Pterula gracilis* CBS309.79 v1.0 [123]; *Coprinopsis cinerea* [124]; *Coprinopsis cinerea*
251 AmutBmut pab1-1 v1.0 [125]; *Cordyceps militaris* CM01 [126]; *Corynespora cassiicola* CCP v1.0
252 [127]; *Cronartium quercuum* f. sp. *fusiforme* G11 v1.0 [128]; *Cryphonectria parasitica* EP155 v2.0
253 [129]; *Cryptococcus curvatus* ATCC 20509 v1.0; *Cryptococcus terricola* JCM 24523 v1.0 [130];
254 *Cryptococcus neoformans* var *neoformans* JEC21 [131]; *Cryptococcus neoformans* var. *grubii* H99
255 [132]; *Cylindrobasidium torrendii* FP15055 v1.0; *Fistulina hepatica* v1.0 [133]; *Cystobasidium*
256 *minutum* MCA 4210 v1.0 [134]; *Daedalea quercina* v1.0; *Exidia glandulosa* v1.0; *Fibularhizoctonia* sp.

257 CBS 109695 v1.0; *Laetiporus sulphureus* var. *sulphureus* v1.0; *Neolentinus lepideus* v1.0; *Peniophora*
258 sp. v1.0; *Sistotremastrum niveocremeum* HHB9708 ss-1 1.0; *Sistotremastrum sueicum* v1.0; *Calocera*
259 *cornea* v1.0; *Calocera viscosa* v1.0 [135]; *Daldinia eschscholtzii* EC12 v1.0; *Hypoxyton* sp. CI-4A v1.0;
260 *Hypoxyton* sp. CO27-5 v1.0; *Hypoxyton* sp. EC38 v3.0 [136]; *Debaryomyces hansenii* [137]; *Dekkera*
261 *bruxellensis* CBS 2499 v2.0 [138]; *Dentipellis* sp. KUC8613 v1.0 [139]; *Dichomitus squalens*
262 CBS463.89 v1.0; *Dichomitus squalens* CBS464.89 v1.0; *Dichomitus squalens* OM18370.1 v1.0 [140];
263 *Encephalitozoon cuniculi* GB-M1 [141]; *Encephalitozoon hellem* ATCC 50504; *Encephalitozoon*
264 *romaleae* SJ-2008 [142]; *Encephalitozoon intestinalis* ATCC 50506 [143]; *Endocarpon pusillum*
265 Z07020 [144]; *Endogone* sp FLAS 59071; *Jimgerdemannia flammicorona* AD002; *Jimgerdemannia*
266 *flammicorona* GMNB39; *Jimgerdemannia lactiflua* OSC166217 [145]; *Enterocytozoon bieneusi* H348
267 [146]; *Eremothecium gossypii* ATCC 10895 [147]; *Erysiphe necator* c [148]; *Eurotium rubrum* v1.0
268 [149]; *Exophiala dermatitidis* UT8656 [150]; *Fibroporia radiculosa* TFFH 294 [151]; *Fonsecaea*
269 *monophora* CBS 269.37 [152]; *Fonsecaea nubica* CBS 269.64 [153]; *Fusarium fujikuroi* IMI 58289
270 [154]; *Fusarium graminearum* v1.0 [155]; *Fusarium oxysporum* f. sp. *conglutinans* race 2 54008
271 (*PHW808*); *Fusarium oxysporum* f. sp. *cubense* tropical race 4 54006 (*I15*); *Fusarium oxysporum* f. sp.
272 *lycopersici* MN25 (*FoMN25*) NRRL 54003; *Fusarium oxysporum* f. sp. *radicis-lycopersici* 26381
273 (*CL57*); *Fusarium oxysporum* f. sp. *raphani* 54005; *Fusarium oxysporum* f. sp. *vasinfectum* 25433
274 (*Cotton*); *Fusarium oxysporum* Fo47; *Fusarium oxysporum* NRRL 32931 [156]; *Fusarium oxysporum*
275 f. sp. *lycopersici* 4287 v2; *Fusarium verticillioides* 7600 v2 [157]; *Fusarium oxysporum* f. sp. *melonis*
276 (*FoMelon*) NRRL 26406 [158]; *Fusarium oxysporum* f. sp. *pisi* HDV247 [159]; *Fusarium*
277 *pseudograminearum* CS3096 [160]; *Gaeumannomyces graminis* var. *tritici* R3-111a-1;
278 *Magnaporthe* *poae* ATCC 64411 [161]; *Glarea lozoyensis* ATCC 20868 [162]; *Grosmannia*
279 *clavigera* kw1407 [163]; *Gymnopus androsaceus* JB14 v1.0; *Chalara longipes* BDJ v1.0 [164];
280 *Heterobasidion annosum* v2.0 [165]; *Homolaphlyctis polyrhiza* JEL142 v1.0 [166]; *Hortaea werneckii*
281 EXF-2000 M0 v1.0 [167]; *Ilyonectria* sp. v1.0 [168]; *Kazachstania africana* CBS 2517; *Torulaspora*
282 *delbrueckii* CBS 1146 [169]; *Kluyveromyces lactis*; *Yarrowia lipolytica* (strain CLIB122) [170];
283 *Kuraishia capsulata* CBS 1993 [171]; *Laccaria bicolor* v2.0 [172]; *Lentinula edodes* B17 v1.1 [173];
284 *Lentinula edodes* W1-26 v1.0 [174]; *Lentinus tigrinus* ALCF2SS1-6 v1.0; *Lentinus tigrinus* ALCF2SS1-
285 7 v1.0 [175]; *Leptosphaeria maculans* [176]; *Leucoagaricus gongylophorus* Ac12 [177];
286 *Leucosporidiella creatinivora* 62-1032 v1.0; *Kockovaella imperatae* NRRL Y-17943 v1.0; *Naematella*
287 *encephala* UCDFST 68-887.2 v1.0; *Clochesyomyces aquaticus* v1.0; *Pseudomassariella vexata* CBS
288 129021 v1.0; *Protomyces lactucae* debilis 12-1054 v1.0; *Lobosporangium transversale* NRRL 3116
289 v1.0; *Absidia repens* NRRL 1336 v1.0; *Hesseltinella vesiculosa* NRRL3301 v2.0; *Rhizopus microsporus*
290 var. *microsporus* ATCC52813 v1.0; *Syncephalastrum racemosum* NRRL 2496 v1.0; *Basidiobolus*
291 *meristosporus* CBS 931.73 v1.0; *Linderina pennispora* ATCC 12442 v1.0; *Catenaria anguillulae* PL171
292 v2.0; *Rhizoclostratum globosum* JEL800 v1.0 [178]; *Lichtheimia corymbifera* JMRC:FSU:9682 [179];
293 *Macrophomina phaseolina* MS6 [180]; *Magnaporthe oryzae* 70-15 v3.0 [181]; *Malassezia globosa*

294 [182]; *Malassezia sympodialis* ATCC 42132 [183]; *Marssonina brunnea* f. sp. *multigermtubi* MB_m1
295 [184]; *Melampsora larici-populina* v2.0; *Puccinia graminis* f. sp. *tritici* v2.0 [185]; *Melampsora lini*
296 CH5 [186]; *Metarhizium acridum* CQMa 102; [187]; *Metarhizium robertsii* ARSEF 23; [188];
297 *Metschnikowia bicuspidata* single-cell v1.0; *Dimargaris crystalligena* RSA 468 single-cell v1.0;
298 *Piptocephalis cylindrospora* RSA 2659 single-cell v3.0; *Syncephalis pseudoplumigaleata* Benny S71-1
299 single-cell v1.0; *Thamnocephalis sphaerospora* RSA 1356 single-cell v1.0; *Blyttiomycetes helicus* single-
300 cell v1.0; *Caulochytrium protostelioides* ATCC 52028 v1.0; *Rozella allomycis* CSF55 single-cell v1.0
301 [189]; *Metschnikowia fructicola* 277 [190]; *Microbotryum lychnidis-dioicae* pIA1 Lamole [191];
302 *Microdochium bolleyi* J235TASD1 v1.0 [192]; *Microsporum canis* CBS 113480; *Trichophyton rubrum*
303 CBS 118892 [193]; *Mitosporidium daphniae* UGP3 [194]; *Mixia osmundae* IAM 14324 v1.0; *Tilletiaria*
304 *anomala* UBC 951 v1.0 [195]; *Moesziomyces aphidis* DSM 70725 [196]; *Monacrosporium haptotylum*
305 CBS 200.50 [197]; *Moniliophthora perniciosa* FA553 [198]; *Morchella importuna* SCYDJ1-A1 v1.0
306 [199]; *Mortierella elongata* AG-77 v2.0 [200]; *Mucor endophyticus*; *Mucor fuscus*; *Mucor lanceolatus*;
307 *Mucor racemosus* [201]; *Mucor lusitanicus* (*circinelloides*) MU402 v1.0 [202]; *Mucor lusitanicus*
308 CBS277.49 v2.0; *Phycomyces blakesleeanus* NRRL1555 v2.0 [203]; *Myceliophthora thermophila*
309 (*Sporotrichum thermophile*) v2.0; *Thielavia terrestris* v2.0 [204]; *Mycosphaerella graminicola* v2.0
310 [205]; *Nakaseomyces bacillisporus* CBS 7720; *Nakaseomyces delphensis* CBS 2170 [206]; *Nectria*
311 *haematococca* v2.0 [207]; *Nematocida parisii* ERTm1 [208]; *Neolecta irregularis* DAH-1 v1.0 [209];
312 *Neonectria ditissima* R09/05 [210]; *Neurospora crassa* FGSC 73 trp-3 v1.0 [211]; *Neurospora crassa*
313 OR74A v2.0 [212]; *Neurospora tetrasperma* FGSC 2508 mat A v2.0; *Neurospora tetrasperma* FGSC
314 2509 mat a v1.0 [213]; *Nosema ceranae* BRL01 [214]; *Obba rivulosa* 3A-2 v1.0 [215]; *Omphalotus*
315 *olearius* [216]; *Ophiostoma novo-ulmi* subsp. *novo-ulmi* H327 [217]; *Ophiostoma piceae* UAMH 11346
316 [218]; *Orpinomyces* sp. [219]; *Paecilomyces niveus* CO7 v1.0 [220]; *Paecilomyces variotii* CBS 101075
317 v1.0; *Paecilomyces variotii* CBS144490 HYG1 v1.0 [221]; *Paracoccidioides brasiliensis* Pb03;
318 *Paracoccidioides brasiliensis* Pb18 [222]; *Paraconiothyrium sporulosum* AP3s5-JAC2a v1.0;
319 *Pyrenophaeta* sp. DS3sAY3a v1.0; *Stagonospora* sp. SRC1lsM3a v1.0; *Alternaria alternata* SRC1lrK2f
320 v1.0 [223]; *Penicillium antarcticum* IBT 31811; *Penicillium coprophilum* IBT 31321; *Penicillium*
321 *decumbens* IBT 11843; *Penicillium flavigenum* IBT 14082; *Penicillium nalgiovense* FM193;
322 *Penicillium polonicum* IBT 4502; *Penicillium solitum* IBT 29525; *Penicillium steckii* IBT 24891;
323 *Penicillium vulpinum* IBT 29486 [224]; *Penicillium chrysogenum* Wisconsin 54-1255 [225]; *Penicillium*
324 *digitatum* Pd1; *Penicillium digitatum* PHI26 [226]; *Penicillium expansum* d1; *Penicillium italicum*
325 PHI-1 [227]; *Penicillium griseofulvum* PG3 [228]; *Penicillium nordicum* DAOMC 185683 [229];
326 *Penicillium oxalicum* 114-2 [230]; *Penicillium subrubescens* FBCC1632 / CBS132785 [231];
327 *Penicillium thymicola* DAOMC 180753 v1.0 [232]; *Periconia macrospinosa* DSE2036 v1.0; *Cadophora*
328 sp. DSE1049 v1.0 [233]; *Phaeomoniella chlamydospora* UCRPC4; *Diplodia seriata* DS831; *Diaporthe*
329 *ampelina* UCDDA912 [234]; *Phanerochaete carnosa* HHB-10118-Sp v1.0 [235]; *Phanerochaete*
330 *chrysosporium* RP-78 v2.2 [236]; *Phialocephala scopiformis* 5WS22E1 v1.0 [237]; *Phialophora attae*

331 *CBS 131958* [238]; *Phlebia centrifuga* FBCC195 [239]; *Phlebia radiata* Fr. (*isolate 79*, FBCC0043)
332 [240]; *Phlebiopsis gigantea* v1.0 [241]; *Phyllosticta capitalensis* CBS 128856 v1.0; *Phyllosticta*
333 *citriasiiana* CBS 120486 v1.0; *Phyllosticta citribraziliensis* CBS 100098 v1.0; *Phyllosticta citricarpa*
334 CBS 127454 v1.0; *Phyllosticta citrichinaensis* CBS 130529 v1.0; *Phyllosticta paracitricarpa* CBS
335 141357 v1.0; *Phyllosticta* sp. CPC 27913 v1.0 [242]; *Pichia kudriavzevii* CBS573 [243]; *Pichia pastoris*
336 [244]; *Piriformospora indica* DSM 11827 from MPI [245]; *Pleurotus ostreatus* PC9 v1.0 [246];
337 *Pneumocystis jirovecii*; [247]; *Pochonia chlamydosporia* 170 [248]; *Podospora anserina* S mat+ [249];
338 *Polyporus brumalis* BRFM 1820 v1.0 [250]; *Postia placenta* MAD 698-R v1.0 [251]; *Postia placenta*
339 MAD-698-R-SB12 v1.0 [252]; *Pseudocercospora* (*Mycosphaerella*) *fijiensis* v2.0 [253];
340 *Pseudogymnoascus destructans* 20631-21 [254]; *Pseudozyma antarctica* T-34 [255]; *Pseudozyma*
341 *hubeiensis* SY62 [256]; *Psilocybe cubensis* v1.0; *Psilocybe serbica* v1.0 [257]; *Puccinia coronata*
342 *avenae* 12NC29; *Puccinia coronata avenae* 12SD80 [258]; *Puccinia graminis* f. sp. *tritici* 21-0
343 *haplotype A*; *Puccinia graminis* f. sp. *tritici* 21-0 *haplotype B*; *Puccinia graminis* f. sp. *tritici* Ug99
344 *haplotype A*; *Puccinia graminis* f. sp. *tritici* Ug99 *haplotype C* [259]; *Puccinia striiformis* f. sp. *tritici*
345 104 E137 A- [260]; *Puccinia striiformis* f. sp. *tritici* PST-130 [261]; *Puccinia striiformis* f. sp. *tritici*
346 PST-78 v1.0; *Puccinia triticina* 1-1 BBBD Race 1 [262]; *Pycnoporus cinnabarinus* BRFM 137 [263];
347 *Pycnoporus coccineus* BRFM 310 v1.0; *Pycnoporus puniceus* CIRM-BRFM 1868 v1.0; *Pycnoporus*
348 *sanguineus* BRFM 1264 v1.0; *Ramaria rubella* (*R. acris*) UT-36052-T v1.0 [264]; *Pyrenophora teres* f.
349 *teres* [265]; *Pyrenophora tritici-repentis* [266]; *Pyronema confluens* CBS100304 [267]; *Rhizoctonia*
350 *solani* AG-1 IB [268]; *Rhizophagus irregularis* A1 v1.0; *Rhizophagus irregularis* A4 v1.0; *Rhizophagus*
351 *irregularis* A5 v1.0; *Rhizophagus irregularis* B3 v1.0; *Rhizophagus irregularis* C2 v1.0; *Rhizophagus*
352 *irregularis* DAOM 197198 v2.0 [269]; *Rhizophagus irregularis* DAOM 181602 v1.0 [270]; *Rhizopogon*
353 *vesiculosus* Smith; *Rhizopogon vinicolor* AM-OR11-026 v1.0 [271]; *Rhizopus delemar* 99-880 from
354 Broad [272]; *Rhizopus microsporus* ATCC11559 v1.0; *Rhizopus microsporus* var. *microsporus*
355 ATCC52814 v1.0 [273]; *Rhizopus microsporus* var. *chinensis* CCTCC M201021 [274];
356 *Rhodosporidium toruloides* IFO0559_1; *Rhodosporidium toruloides* IFO0880 v2.0; *Rhodosporidium*
357 *toruloides* IFO1236_1 [275]; *Rhodosporidium toruloides* IFO0880 v4.0 [276]; *Rhodosporidium*
358 *toruloides* NP11 [277]; *Rhodotorula graminis* strain WP1 v1.1 [278]; *Rhodotorula* sp. JG-1b [279];
359 *Rickenella fibula* HBK330-10 v1.0 [280]; *Rickenella mellea* v1.0 (SZMC22713; [281]; *Rigidoporus*
360 *microporus* ED310 v1.0 [282]; *Rozella allomycis* CSF55 [283]; *Saccharomyces arboricola* H-6 [284];
361 *Saccharomyces cerevisiae* M3707 Dikaryon; *Saccharomyces cerevisiae* M3836 v1.0; *Saccharomyces*
362 *cerevisiae* M3837 v1.0; *Saccharomyces cerevisiae* M3838 v1.0; *Saccharomyces cerevisiae* M3839 v1.0;
363 *Arthroderma benhamiae* CBS 112371; *Trichophyton verrucosum* HKI 0517 [285]; *Saccharomyces*
364 *cerevisiae* S288C [286]; *Saksenaea vasiformis* B4078 [287]; *Scheffersomyces stipitis* NRRL Y-11545
365 v2.0 [288]; *Schizophyllum commune* H4-8 v3.0 [289]; *Schizopora paradoxa* KUC8140 v1.0 [290];
366 *Schizosaccharomyces cryophilus* OY26; *Schizosaccharomyces japonicus* yFS275;
367 *Schizosaccharomyces octosporus* yFS286 [291]; *Schizosaccharomyces pombe* [292]; *Sclerotinia*

368 *sclerotiorum* v1.0 [293]; *Serpula himantoides* (*S.lacrymans* var *shastensis*) MUCL38935 v1.0 [294];
369 *Serpula lacrymans* S7.3 v2.0; *Serpula lacrymans* S7.9 v2.0 [295]; *Smittium culicis* GSMNP; *Smittium*
370 *culicis* ID-206-W2; *Smittium mucronatum* ALG-7-W6; *Zancudomyces culisetae* COL-18-3 [296];
371 *Sodiomyces alkalinus* v1.0 [297]; *Sphaerospora brunnea* Sb_GMNB300 v2.0 [298]; *Spizellomyces*
372 *punctatus* DAOM BR117 [299]; *Sporisorium reilianum* SRZ2 [300]; *Stagonospora nodorum* SN15 v2.0
373 [301]; *Stemphylium lycopersici* CIDEFI-216 [302]; *Suillus brevipes* Sb2 v2.0 [303]; *Talaromyces*
374 *borbonicus* CBS 141340 [304]; *Talaromyces marneffei* ATCC 18224 [305]; *Taphrina deformans* [306];
375 *Thermomyces lanuginosus* SSBP [307]; *Tolypocladium inflatum* NRRL 8044 [308]; *Trametes pubescens*
376 FBCC735 [309]; *Trichoderma arundinaceum* IBT 40837; *Trichoderma brevicompactum* IBT40841
377 [310] ; *Trichoderma asperellum* CBS 433.97 v1.0; *Trichoderma citrinoviride* TUCIM 6016 v4.0;
378 *Trichoderma guizhouense* NJAU 4742; *Trichoderma harzianum* CBS 226.95 v1.0; *Trichoderma*
379 *longibrachiatum* ATCC 18648 v3.0 [311]; *Trichoderma asperellum* TR356 v1.0; *Trichoderma*
380 *harzianum* TR274 v1.0 [312]; *Trichoderma atrobrunneum* ITEM 908 [313]; *Trichoderma atroviride*
381 v2.0; *Trichoderma virens* Gv29-8 v2.0 [314]; *Trichoderma hamatum* GD12 [315]; *Trichoderma*
382 *parareesei* CBS 125925 [316]; *Trichoderma pleuroti* TPhu1 [317]; *Trichoderma reesei* QM6a [318];
383 *Trichoderma reesei* RUT C-30 v1.0 [319]; *Trichoderma reesei* v2.0 [320]; *Trichosporon asahii* var.
384 *asahii* CBS 2479 [321]; *Trichosporon asahii* var. *asahii* CBS 8904 [322]; *Trichosporon oleaginosus*
385 IBC0246 v1.0 [323]; *Tuber melanosporum* Mel28 v1.2 [324]; *Ustilaginoidea virens* [325]; *Ustilago*
386 *hordei* Uh4857_4 [326]; *Ustilago maydis* 521 v2.0 [327]; *Venturia inaequalis*; *Venturia pirina* [328];
387 *Verticillium alfalfae* VaMs.102; *Verticillium dahliae* VdLs.17 [329]; *Volvariella volvacea* V23 [330];
388 *Wallemia ichthyophaga* EXF-994 [331]; *Wallemia mellicola* v1.0 [332]; *Xylona heveae* TC161 v1.0
389 [333]; *Yarrowia lipolytica* CLIB89(W29) [334]; *Yarrowia lipolytica* FKP355 v1.0 [335]; *Yarrowia*
390 *lipolytica* PO1f v1.0; *Yarrowia lipolytica* YICW001 v1.0 [336]; *Yarrowia lipolytica* YB392 v1.0;
391 *Yarrowia lipolytica* YB419 v1.0; *Yarrowia lipolytica* YB420 v1.0; *Yarrowia lipolytica* YB566 v1.0;
392 *Yarrowia lipolytica* YB567 v1.0 [337]; *Zygosaccharomyces rouxii* CBS732 [338]; *Zymoseptoria*
393 *ardabilliae* STIR04_1.1.1; *Zymoseptoria pseudotritici* STIR04_2.2.1 [339]; *Zymoseptoria brevis*
394 Zb18110 [340]; *Melampsora allii-populina* 12AY07 v1.0; *Melampsora americana* R15-033-03 v1.0
395 (unpublished).

396

397 **2.3. Protein sequence analysis and search for potential subcellular targeting.**

398 The protein sequences were aligned using Clustal Omega [341]
399 (<https://www.ebi.ac.uk/Tools/msa/clustalo/>) and manually checked. The predictions for subcellular
400 localizations were done based on the assessment obtained from the TargetP [342]
401 (<http://www.cbs.dtu.dk/services/TargetP/>), SignalP [343]
402 (<http://www.cbs.dtu.dk/services/SignalP/abstract.php>), WolfPsor [344] (<https://wolfsort.hgc.jp/>) and
403 ESLpred [345] (<http://crdd.osdd.net/raghava/eslpred/>) software suites.

404

405 **2.4. Phylogenomic analyses and search for horizontal gene transfers.**

406 For each type of Msr, we built a multiple sequence alignment with all the identified sequences
407 using MAFFT version v7.429 [346]. Each alignment was trimmed to remove poorly aligned regions
408 using trimAl 1.2 [347] and manually inspected for the conservation of the catalytic residues. A
409 phylogenetic tree was constructed for each type of Msr using RAxML Master Pthread [348] version
410 8.2.12 (PROTGAMMAWAG model and 500 bootstraps). The phylogenetic tree was represented using
411 iTOL (<https://itol.embl.de/>) [349]. To investigate potential horizontal gene transfers, the non-canonical
412 Msr sequences along with the sequences whose phylogeny did not fit the species phylogeny were
413 retrieved. Those sequences were used as BLAST queries against the non-redundant protein sequence
414 (nr) database of NCBI
415 (https://blast.ncbi.nlm.nih.gov/Blast.cgi?PROGRAM=blastp&PAGE_TYPE=BlastSearch&LINK_LOC=blasthome) and the best 100 hits were retrieved. The fungal sequences for which protein accessions
416 outside the fungal kingdom were identified among the highest identity scores were in further used as
417 BLAST queries against nr excluding the taxon Fungi (taxid: 4751) and the 25 hits with the highest
418 identity scores were retrieved. In parallel, the same fungal genes were used as queries against the fungal
419 nr database and the 50 hits with the highest identity scores were retrieved, excluding those with 100%
420 identity with the query. The retrieved fungal hits were manually inspected to exclude incomplete
421 sequences. Non-canonical sequences (lacking canonical active site or of abnormal length) were also
422 removed to avoid potential inconsistencies and long branches in the phylogenetic trees. Finally, for each
423 type of Msr, a phylogenetic tree was constructed that included the candidate fungal gene for horizontal
424 gene transfer, plus 25 non-fungal and 25 fungal sequences with the highest identity scores. For each
425 candidate to HGT, the numbers of exons, the GC content of the coding sequence and the GC content at
426 the third positions of the codons were calculated using the gene structures and coding sequences
427 downloaded from the MycoCosm database. The GC content on the third positions of the codons were
428 calculated only for the coding sequences starting with ATG. All calculations were made using
429 Microsoft® Excel® version 2101.

431 **3. Results**432 **3.1. MsrA, MsrB and fRMSr are largely conserved across the fungal kingdom.**

433 Currently, around 136,000 species of fungi are known and classified into 9 phyla (**Figure 1**)
434 [37,350]. Together, the Ascomycota and the Basidiomycota form the subkingdom Dikarya, which
435 regroups more than 97 % of all described fungal species (~84,000 species and ~48,000 species,
436 respectively) [350]. Each phylum contains three monophyletic subphyla (Pucciniomycotina,
437 Ustilaginomycotina and Agaricomycotina for the Basidiomycota, and Taphrinomycotina,
438 Saccharomycotina and Pezizomycotina for the Ascomycota). The other seven phyla are described as
439 ‘early-diverging fungi’ [350] (**Fig. 1, Data S1**). Using the *S. cerevisiae* MsrA, MsrB and fRMSr protein
440 sequences as queries, we used the BLASTP and TBLASTN software suites to search for *msr* genes in
441 683 available genomes in the MycoCosm database [50]. The selected genomes were from 595 species
442 that spanned the kingdom Fungi (**Fig. 1, Data S1**), including 65% Ascomycota species, 25%
443 Basidiomycota species and 10% early-diverging fungi. We found that the very great majority of these
444 genomes contained one gene coding for a MsrA and one coding for a MsrB (**Fig. 1, Data S1**), indicating
445 that most fungi have a simple Msr system dedicated to protein oxidation repair, similarly to most other
446 known organisms [24,26]. Most interestingly, we found that a *fRmsr* gene was present in almost all the
447 analyzed genomes (**Fig. 1, Data S1**). The distribution of *fRmsr* across the fungal kingdom clearly
448 showed that the presence of fRMSr is not limited to bacteria and unicellular eukaryotes as previously
449 described [7–9]. Of note, the search for homologs of the bacterial molybdoenzymes able to reduce MetO
450 gave no significant hits (**data not shown**).

451

	Fungal groups	Genomes (Species)	<i>msrA</i>	<i>msrB</i>	<i>fRmsr</i>
BASIDIOMYCOTA	Pucciniomycotina	27 (16)	30	31	12
	Ustilaginomycotina	17 (17)	18	17	17
	Agaricomycetes	113 (102)	130	114	120
	Dacrymycetes	3 (3)	3	3	3
	Tremellomycetes	10 (8)	10	9	10
	Walleiomycetes	2 (2)	2	2	2
Dikarya	Pezizomycetes	12 (11)	12	12	12
	Orbiliomycetes	2 (2)	2	4	1
	Eurotiomycetes	139 (127)	136	139	137
	Dothideomycetes	114 (109)	114	115	114
	Lecanoromycetes	1 (1)	1	0	1
	Leotiomycetes	18 (16)	18	18	15
ASCOMYCOTA	Sordariomycetes	86 (70)	87	87	85
	Xylonomycetes	1 (1)	1	1	1
	Saccharomycotina	61 (46)	87	62	61
	Taphrinomycotina	9 (9)	8	9	3
GLOMEROMYCOTA	Glomeromycotina	9 (4)	10	10	0
MUCOROMYCOTA	Mortierellomycotina	2 (2)	4	2	2
	Mucoromycotina	21 (16)	35	21	21
ZOOPAGOMYCOTA	Entomophthoromycotina	2 (2)	3	3	1
	Kickxellomycotina	7 (6)	11	9	6
	Zoopagomycotina	3 (3)	4	1	1
BLASTOCLADIOMYCOTA	Blastocladiomycota	1 (1)	1	2	1
CHYTRIDIOMYCOTA	Chytridiomycetes	5 (5)	7	5	4
NEOCALLIMASTIGOMYCOTA	Monoblepharidomycetes	1 (1)	3	1	1
OPISTHOSPORIDIA	Neocallimastigomycetes	5 (5)	0	0	3
	Microsporidia	9 (9)	0	0	0
	Cryptomycota	2 (1)	2	0	0
	Total	683 (595)	739	677	634

452

453 **Figure 1.** Numbers of *msrA*, *msrB* and *fRmsr* genes identified in fungal genomes. The simplified
 454 phylogenetic tree was built according to [37,350]. The precise number of *msr* genes per genome is
 455 available in [Data S1](#).

456

457 We observed some variations in the numbers of *msr* genes in several genomes. We found 72
 458 genomes, corresponding to 57 species, that had more than one copy of at least one *msr* gene (**Table 1**,
 459 **Data S1-4**). We found 42 genomes having two copies of *msrA* and 17 having three or four copies (**Table**
 460 **1, Data S1-2**). The highest number of *msrA* copies was 4 in the genome of the Agaricomycetes
 461 *Crucibulum laeve* CBS 166.37 and *Dendrothele bispora* CBS 962.96 (**Table 1**). In the case of *msrB*,
 462 only 14 genomes had two gene copies (**Data S1, S3**), and the Kickxellomycotina *Smittium culicis*
 463 GSMNP was the only genome with three copies (**Table 1**). For *fRmsr*, only six genomes had two copies
 464 (**Data S1, S4**), and the Agaricomycetes *Ramaria rubella* (*R. acris*) UT-36052-T and *Coprinellus*
 465 *micaceus* FP101781 had 4 and 3 gene copies, respectively (**Table 1**). Overall, for the three types of Msr,
 466 the fungal species having more than one copy were sporadically spread across the fungal kingdom
 467 (**Table 1, Data S1-4**), indicating that gene enrichment was not a characteristic of a specific fungal group.

468 **Table 1.** Fungi with five or six Msr genes.

Genome	MsrA	MsrB	fRmsr	Lifestyle	Nb. Genes (Genome size in Mbp)	Ref.
Agaricomycetes						
<i>Crucibulum laeve</i> CBS 166.37 v1.0	4	1	1		14,218 (44)	[351]
<i>Dendrothele bispora</i> CBS 962.96 v1.0	4	1	1		33,645 (131)	[351]
<i>Tulasnella calospora</i> AL13/4D v1.0	3	1	1	Aerobic (saprotroph)	19,659 (62)	[352]
<i>Ramaria rubella</i> (<i>R. acris</i>) UT-36052-T v1.0	1	1	4		14,207 (62)	[353]
<i>Coprinellus micaceus</i> FP101781 v2.0	1	1	3		23,559 (77)	[351]
Dothideomycetes						
<i>Hortaea werneckii</i> EXF-2000 M0 v1.0	2	2	2	Aerobic	15,748 (50)	[167]
Sordariomycetes						
<i>Coniochaeta</i> sp. 2T2.1 v1.0	2	2	2	Aerobic (phytopathogen)	24,735 (74)	[121]
Saccharomycotina						
<i>Pichia kudriavzevii</i> CBS573	3	1	1		5,140 (11)	[354]
<i>Yarrowia lipolytica</i> (strain CLIB122)	3	1	1		6,447 (21)	[355]
<i>Yarrowia lipolytica</i> CLIB89(W29)	3	1	1		7,919 (21)	[356]
<i>Yarrowia lipolytica</i> FKP355 v1.0	3	1	1		6,858 (20)	[357]
<i>Yarrowia lipolytica</i> PO1f v1.0	3	1	1		6,798 (20)	[358]
<i>Yarrowia lipolytica</i> YB392 v1.0	3	1	1	Aerobic	6,750 (20)	[359]
<i>Yarrowia lipolytica</i> YB419 v1.0	3	1	1		6,751 (20)	[359]
<i>Yarrowia lipolytica</i> YB420 v1.0	3	1	1		6,772 (20)	[359]
<i>Yarrowia lipolytica</i> YB566 v1.0	3	1	1		6,764 (20)	[359]
<i>Yarrowia lipolytica</i> YB567 v1.0	3	1	1		6,776 (20)	[359]
<i>Yarrowia lipolytica</i> YICW001 v1.0	3	1	1		6,800 (20)	[358]
Mucoromycotina						
<i>Absidia repens</i> NRRL 1336 v1.0	3	1	1	Aerobic (saprotroph)	14,919 (47)	[360]
<i>Rhizopus microsporus</i> ATCC11559 v1.0	2	1	2		11,135 (24)	[273]
<i>Rhizopus microsporus</i> var. <i>chinensis</i> CCTCC M201021	2	1	2	Aerobic (phytopathogen)	17,676 (46)	[274]
Kickxellomycotina						
<i>Smittium mucronatum</i> ALG-7-W6	3	1	1		8,247 (102)	[361]
<i>Smittium culicis</i> GSMNP	2	3	1	Anaerobic (insect gut)	12,166 (77)	[361]
Monoblepharidomycetes						
<i>Gonapodya prolifera</i> v1.0	3	1	1	Aerobic (aquatic)	13,902 (49)	[362]

469 Also, we found only 74 genomes (58 species), in which one or more Msr types were absent
470 (**Table 2**). Among them, the nine Microsporidia species, two Neocallimastigomycetes species (out of
471 five species analyzed) and one Taphrinomycotina species, *Pneumocystis jirovecii* (out of eight species
472 analyzed) lacked all three Msr types. These species are either obligate intracellular parasites
473 (Microsporidia and *Pneumocystis jirovecii*) or live in the anaerobic conditions of the animal gut
474 (*Piromyces finnis* and *Piromyces* sp. E2) [37,363]. Interestingly, the three other Neocallimastigomycetes
475 species (*Anaeromyces robustus*, *Neocallimastix californiae* G1 and *Orpinomyces* sp.), also living in
476 anaerobic conditions [350], do not possess MsrA nor MsrB but have a gene coding for a fRMsR (**Table**
477 **2**). Our results show that, additionally to *Encephalitozoon cuniculi*, previously identified as the unique
478 eukaryote lacking both MsrA and MsrB [24], 14 other fungal species are devoid of any protein-bound
479 MetO reduction enzyme (**Table 2**). Furthermore, we found six species lacking *msrA* but having a *msrB*
480 and a *fRmsr* gene (**Table 2**). These are the agaricomycete *Moniliophthora perniciosa*, the
481 eurotiomycetes *Cladophialophora immunda*, *Penicillium coprophilum* and *Penicillium flavigenum*, the
482 sordariomycete *Magnaportheopsis poae* and the saccharomycotina *Saturnispora disporea*. To our
483 knowledge, they constitute the first species, over all kingdoms, described to have only a MsrB to reduce
484 and repair oxidized proteins, as none was found so far in genome surveys [24,26]. Of note, we did not
485 find fungal species having only a MsrB, as these six species also possessed a fRMsR (**Table 2**). We
486 found that the five species *Trichosporon asahii* (Tremellomycetes), *Clochesyomyces aquaticus*
487 (Dothideomycetes), *Cladonia grayi* (Lecanoromycetes) and *Piptocephalis cylindrospora*
488 (Zoopagomycotina) possessed a MsrA and a fRMsR but lacked a MsrB and that the Zoopagomycotina
489 *Syncephalis pseudoplumigaleata* had a MsrA only (**Table 2**). Finally, we found only 27 species having
490 both a MsrA and a MsrB but lacking a fRMsR (**Table 2**). These species were sporadically dispersed
491 among the fungal kingdom, but three groups stood out as remarkable: the Glomeromycotina, for which
492 none of the four species analyzed had a fRMsR, the Taphrinomycotina for which six species out of eight
493 analyzed (including the four *Schizosaccharomyces* species analyzed) were devoid of fRMsR, and the
494 Pucciniomycotina for which all the *Melampsora* and *Puccinia* species analyzed here lacked a fRMsR
495 (**Table 2**).

496 Altogether, these results showed that the great majority of fungi possess one gene coding for
497 each protein-repairing Msr type (i.e. MsrA and MsrB), as well as one gene coding for the free MetO
498 reductase fRMsR.

499

Table 2. Fungal genomes lacking one or more Msr types.

Genome	MsrA	MsrB	fRMsr	Lifestyle	Nb. Genes (Genome size in Mbp)	Ref.	
Microsporidia							
<i>Antonospora locustae</i> HM-2013	N	N	N		2,606 (6)	[364]	
<i>Encephalitozoon cuniculi</i> GB-M1	N	N	N		1,996 (2)	[365]	
<i>Encephalitozoon hellem</i> ATCC 50504	N	N	N		1,847 (2)	[366]	
<i>Encephalitozoon intestinalis</i> ATCC 50506	N	N	N	Intracellular parasite of metazoans	1,833 (2)	[367]	
<i>Encephalitozoon romaleae</i> SJ-2008	N	N	N		1,831 (2)	[366]	
<i>Enterocytozoon bieneusi</i> H348	N	N	N		3,632 (4)	[368]	
<i>Mitosporidium daphniae</i> UGP3	N	N	N		3,330 (6)	[369]	
<i>Nematocida parisii</i> ERTm1	N	N	N		2,661 (4)	[370]	
<i>Nosema ceranae</i> BRL01	N	N	N		2,060 (8)	[371]	
Neocallimastigomycetes							
<i>Piromyces finnis</i> v3.0	N	N	N		10,992 (56)	[372]	
<i>Piromyces</i> sp. E2 v1.0	N	N	N	Anaerobic (ruminant gut)	14,648 (71)	[372]	
<i>Anaeromyces robustus</i> v1.0	N	N	Y		12,832 (72)	[372]	
<i>Neocallimastix californiae</i> G1 v1.0	N	N	Y		20,219 (193)	[372]	
<i>Orpinomyces</i> sp.	N	N	Y		18,936 (101)	[373]	
Taphrinomycotina							
<i>Pneumocystis jirovecii</i>	N	N	N	Intracellular parasite of metazoans	3,520 (8)	[374]	
<i>Neolecta irregularis</i> DAH-1 v1.0	Y	Y	N		6,674 (15)	[375]	
<i>Schizosaccharomyces cryophilus</i> OY26	Y	Y	N		5,180 (12)	[376]	
<i>Schizosaccharomyces japonicus</i> yFS275	Y	Y	N	Aerobic	4,878 (12)	[376]	
<i>Schizosaccharomyces octosporus</i> yFS286	Y	Y	N		4,986 (12)	[376]	
<i>Schizosaccharomyces pombe</i>	Y	Y	N		5,134 (13)	[377]	
Agaricomycetes							
<i>Moniliophthora perniciosa</i> FA553	N	Y	Y	Aerobic (phytopathogen)	13,560 (18)	[378]	
Eurotiomycetes							
<i>Penicillium coprophilum</i> IBT 31321	N	Y	Y		8,999 (28)	[379]	
<i>Penicillium flavigenum</i> IBT 14082	N	Y	Y		10,994 (33)	[379]	
<i>Penicillium polonicum</i> IBT 4502	N	Y	Y	Aerobic	10,694 (32)	[379]	
<i>Aspergillus bombycis</i> NRRL 26010	Y	Y	N		12,265 (37)	[380]	
<i>Aspergillus zonatus</i> v1.0	Y	Y	N		9,886 (29)	[70]	
Sordariomycetes							
<i>Magnaporthe poae</i> ATCC 64411	N	Y	Y	Aerobic (phytopathogen)	12,335 (40)	[381]	
<i>Pochonia chlamydosporia</i> 170	Y	Y	N		14,204 (44)	[382]	
Saccharomycotina							
<i>Saccharomyces cerevisiae</i> Y-1447	N	Y	Y	Aerobic	4,881 (10)	[383]	
Tremellomycetes							
<i>Trichosporon asahii</i> var. <i>asahii</i> CBS 2479	Y	N	Y	Aerobic (human pathogen)	8,300 (24)	[384]	
<i>Trichosporon asahii</i> var. <i>asahii</i> CBS 8904	Y	N	Y		8,507 (25)	[385]	
Dothideomycetes							
<i>Clovesyomyces aquaticus</i> v1.0	Y	N	Y	Aerobic (saprotrophic)	15,810 (50)	[360]	
Lecanoromycetes							
<i>Cladonia grayi</i> Cgr/DA2myc/ss v2.0	Y	N	Y		11,389 (35)	[386]	
Zoopagomycotina							
<i>Piptocephalis cylindrospora</i> RSA 2659 single-cell v3.0	Y	N	Y	Aerobic (mycoparasite)	4,301 (11)	[387]	

<i>Syncephalis pseudoplumigaleata</i> Benny S71-1 single-cell v1.0	Y	N	N	6,123 (16)	[387]	
<i>Thamnocephalis sphaerospora</i> RSA 1356 single-cell v1.0	Y	Y	N	6,857 (18)	[387]	
Pucciniomycotina						
<i>Melampsora allii-populina</i> 12AY07 v1.0	Y	Y	N	23,089 (336)	-	
<i>Melampsora americana</i> R15-033-03 v1.0	Y	Y	N	15,984 (112)	-	
<i>Melampsora larici-populina</i> v2.0	Y	Y	N	19,550 (110)	[388]	
<i>Melampsora lini</i> CH5	Y	Y	N	16,335 (190)	[389]	
<i>Puccinia coronata avenae</i> 12NC29	Y	Y	N	28,270 (166)	[390]	
<i>Puccinia coronata avenae</i> 12SD80	Y	Y	N	26,323 (150)	[390]	
<i>Puccinia graminis</i> f. sp. <i>tritici</i> v2.0	Y	Y	N	15,979 (89)	[388]	
<i>Puccinia striiformis</i> f. sp. <i>tritici</i> 104 E137 A-	Y	Y	N	30,249 (157)	[391]	
<i>Puccinia striiformis</i> f. sp. <i>tritici</i> PST-130	Y	Y	N	18,021 (65)	[392]	
<i>Puccinia striiformis</i> f. sp. <i>tritici</i> PST-78 v1.0	Y	Y	N	20,482 (117)	[393]	
<i>Puccinia triticina</i> 1-1 BBBB Race 1	Y	Y	N	15,685 (135)	[393]	
Leotiomycetes						
<i>Blumeria graminis</i> f. sp. <i>hordei</i> DH14	Y	Y	N	7,118 (125)	[394]	
<i>Blumeria graminis</i> f. sp. <i>hordei</i> Race1	Y	Y	N	7,239 (116)	[394]	
<i>Blumeria graminis</i> f. sp. <i>tritici</i> 96224	Y	Y	N	6,525 (159)	[395]	
Glomeromycotina						
<i>Gigaspora rosea</i> v1.0	Y	Y	N	31,291 (568)	[396]	
<i>Rhizophagus cerebriforme</i> DAOM 227022 v1.0	Y	Y	N	21,549 (137)	[396]	
<i>Rhizophagus diaphanus</i> v1.0	Y	Y	N	23,252(129)	[396]	
<i>Rhizophagus irregularis</i> A1 v1.0	Y	Y	N	26,659 (126)	[397]	
<i>Rhizophagus irregularis</i> A4 v1.0	Y	Y	N	25,760 (138)	[397]	
<i>Rhizophagus irregularis</i> A5 v1.0	Y	Y	N	26,585 (131)	[397]	
<i>Rhizophagus irregularis</i> B3 v1.0	Y	Y	N	25,164 (125)	[397]	
<i>Rhizophagus irregularis</i> C2 v1.0	Y	Y	N	26,756 (123)	[397]	
<i>Rhizophagus irregularis</i> DAOM 197198 v2.0	Y	Y	N	26,183 (137)	[397]	
Mucoromycotina						
<i>Jimmerdemannia lactiflua</i> OSC166217	Y	Y	N	Aerobic (ectomycorrhizal)	12,651 (180)	[362]
<i>Lichtheimia corymbifera</i> JMRC:FSU:9682	Y	Y	N	Aerobic (saprotrophic/human pathogen)	13,404 (34)	[398]
Entomophthoromycotina						
<i>Basidiobolus meristosporus</i> CBS 931.73 v1.0	Y	Y	N	Aerobic (saprotrophic/animal pathogen)	16,111 (89)	[360]
Kickxellomycotina						
<i>Dimargaris cristalligena</i> RSA 468 single-cell v1.0	Y	Y	N	Aerobic (mycoparasite)	7,456 (31)	[387]
Chytridiomycetes						
<i>Caulochytrium protostelioides</i> ATCC 52028 v1.0	Y	Y	N	Aerobic (mycoparasite)	6,168 (22)	[387]

Y, presence of one or more genes coding for the Msr; N, absence of gene coding for the Msr.

501

502

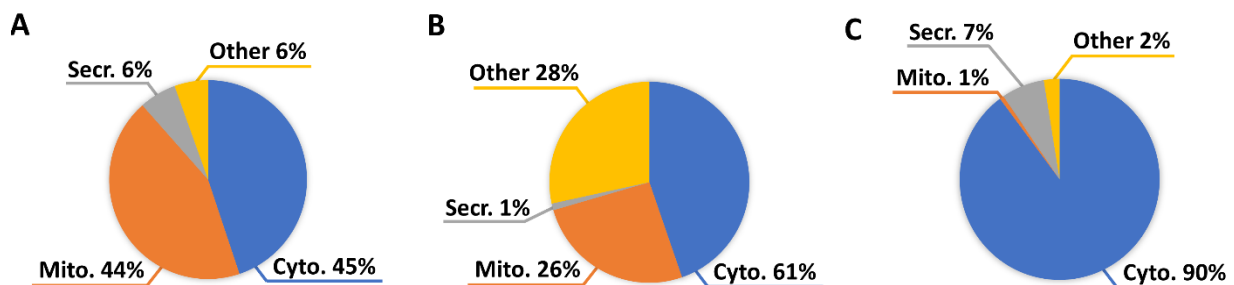
503 **3.2. Fungal Msrs globally retained canonical sequence features for activity.**

504 Sequence analysis and enzymatic characterizations of MsrAs from several organisms showed
505 that the catalytic residue is located within a G[C/U]FW motif, located in the N-terminal part of the
506 protein [6,16,17,399,400]. The great majority of known MsrAs possess a catalytic Cys, whereas a few
507 Sec-containing proteins are found in some insects, marine organisms and unicellular algae [25,400]. In
508 canonical MsrAs, the resolving Cys, involved in the regeneration of the catalytic Cys, is located in the
509 C-terminal part of the protein, although the number and positions of resolving Cys vary [6,17,22,23].
510 To determine whether the fungal MsrAs share these properties, we analyzed the 709 full length MsrAs
511 identified. These MsrAs ranged from 142 to 322 amino acids, and all of them were made of a single
512 MsrA domain (**Data S2A**). The length differences were mostly due to the presence of N-terminal
513 extensions of variable sizes, indicating the possible presence of signal peptides for protein distribution
514 in subcellular compartments. We used several targeting prediction programs (see section 2.3) to evaluate
515 the potential subcellular localization of the fungal MsrAs (**Data S2A**). Most MsrAs were predicted to
516 be localized into the cytoplasm (45 %) or in the mitochondria (44 %) (**Fig. 2A, Data S2A**). The other
517 sequences were predicted to be secreted (6%), to be localized in other compartments, or had no clearly
518 assigned localization (6%) (**Fig. 2A, Data S2A**). The alignment of the primary sequences revealed that
519 the great majority (~ 90 %) shared common features with the previously characterized *S. cerevisiae*
520 MsrA [6,401] (**Fig. 3A, Data S2B**). The catalytic Cys (position 25 in the *S. cerevisiae* MsrA) was located
521 in the conserved motif ²⁴GCFW²⁷. The Tyr⁶⁴, Glu⁷⁶, Asp¹¹¹ and Tyr¹¹⁶ residues involved in substrate
522 stabilization and catalysis [399], and the residues Gly⁴⁷, His¹⁰⁰, Gln¹⁰⁸, Gly¹¹³, His¹⁶³ and Tyr¹⁶⁶ were
523 also conserved. Finally, the Cys¹⁷⁶, previously identified as resolving Cys [6,401], was included in the
524 ¹⁷³GYXC¹⁷⁶ motif (**Fig. 3A, Data S2B**). Because of their predominance in all the fungal kingdom, we
525 defined the fungal MsrAs having these properties as '*canonical*' sequences. The fungal MsrAs that did
526 not match these sequence features were defined as '*non-canonical*' MsrAs (**Data S2A, C, D**).
527 Particularly, we observed that in ~ 5 % of the identified sequences, the Phe²⁶ residue in the ²⁴GCFW²⁷
528 motif containing the catalytic Cys, was substituted by a Tyr. We also observed the replacement of Asp¹¹¹
529 by an Asn residue in few sequences (~ 2 %) (**Data S2C**). Moreover, some variations were also observed
530 for the resolving Cys (**Data S2C**). Fourteen sequences (~ 2 %) lacked the conserved ¹⁷³GYXC¹⁷⁶ motif
531 but possessed two to four Cys in a Q[C/S/K]X₂KX[C/N][C/X]XI[R/L]CYG motif, similar to poplar
532 MsrAs [23]. Some other sequences possessed a Cys residue in the C-terminal region, but not in a GYXC
533 motif, and others lacked any potential resolving Cys (**Data S2C**). Finally, a special case could be made
534 for MsrAs from the early-diverging fungus *Gonapodya prolifera* (Monoblepharidomycetes). This
535 fungus has three non-canonical MsrAs, two of which had the catalytic Cys replaced by a Sec [402]. Each
536 had another Cys outside the conserved position of the resolving Cys in canonical fungal MsrAs. These
537 two MsrA sequences had high similarity with the Sec-MsrAs from the bacterium *Alkaliphilus*

538 *oremlandii* and the single-cell green alga *Chlamydomonas reinhardtii*, previously shown to use the Sec
539 residue for the regeneration of their activity [400,403] (**Data S2D**).

540

541



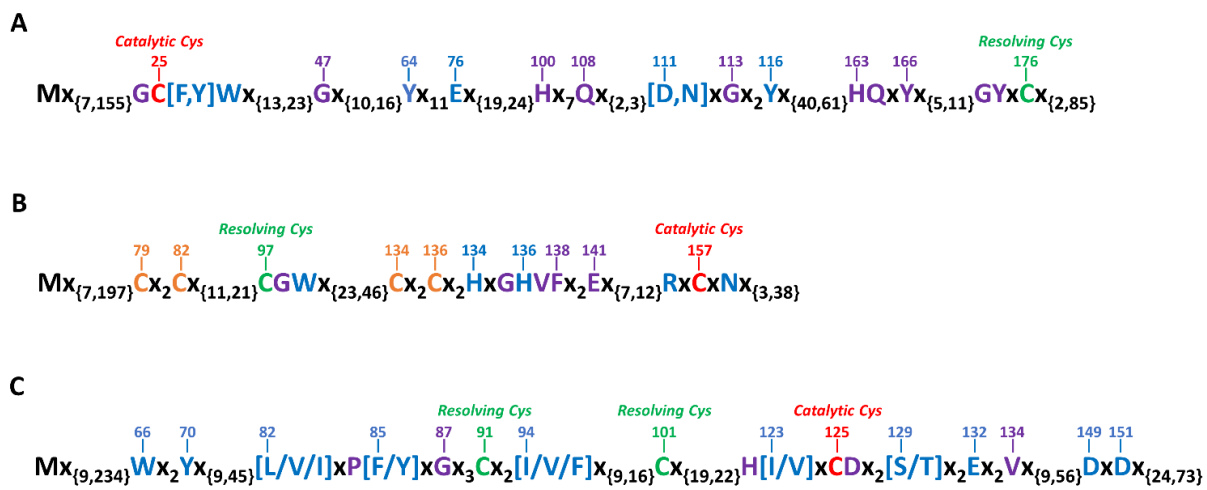
542
543 **Figure 2. Potential subcellular targeting of fungal Msrs.** The circle charts show the subcellular
544 predictions for fungal MsrA (A), MsrB (B) and fRMSr (C), the proportion of proteins predicted to be
545 localized in the cytosol ('Cyto.'), to be addressed to the mitochondria ('Mito.') or secreted ('Secr.'.). The
546 label 'Other' indicates the proportion of proteins predicted to be addressed to other compartments or for
547 which no consensus prediction was obtained.

548

549 In the case of MsrBs, previous biochemical characterizations demonstrated that the catalytic
550 Cys is located in a RXCXN motif in the C-terminal part of the protein [16,21,404–407]. Mammals
551 express a Sec-containing form in which the Asp is replaced by a Phe [404]. The resolving Cys is
552 generally located in a CGWP motif present in the N-terminal part of the protein [21,406]. In the case of
553 mammalian MsrBs, one or two resolving Cys, located in the N-terminal extremity of the protein, can be
554 involved in the regeneration process [404,408]. Of note, most MsrBs possess two CX₂C clusters
555 coordinating a structural Zn atom [409]. Here, we analyzed the 651 complete fungal MsrB protein
556 sequences. The fungal MsrBs consisted of a single domain and ranged in length from 95 to 289 amino
557 acids (**Data S3A**). Most of the variations in size were due to the presence of an N-terminal extension
558 potentially involved in subcellular targeting. The majority of fungal MsrBs were predicted to be
559 addressed to the cytosol (61 %) (**Fig. 2B, Data S3A**). The other proteins were either predicted to be
560 localized in the mitochondria (26 %), secreted (1 %), targeted to another compartment, or were not
561 clearly predicted to be addressed to a subcellular compartment (28 %) (**Fig. 2B, Data S3A**). Almost all
562 fungal MsrBs (> 99 %) possessed the features of the *S. cerevisiae* enzyme (**Fig. 3B, Data S3B**): i) the
563 two CX₂C motifs involved in the coordination of a Zn atom, ii) the resolving Cys⁹⁷ (according to *S.*
564 *cerevisiae* MsrB residue numbering) included in a ⁹⁷CGW⁹⁹ motif, iii) the conserved His¹³⁴ and His¹³⁶
565 implicated in substrate binding, together with Arg¹⁵⁵ and Asn¹⁵⁹ [16,405], and iv) the catalytic Cys¹⁵⁷
566 located in the ¹⁵⁵RXCXN¹⁵⁹ motif. The Gly⁹⁸, Gly¹³⁵, Val¹³⁷, Phe¹³⁸ and Glu¹⁴¹ residues were also
567 conserved in all these canonical fungal MsrBs (**Fig. 3B, Data S3B**). Only four sequences (< 1 %), from

568 the orbiliomycetes *Arthrobotrys oligospora* and *Monacrosporium haptotylum*, the taphrinomycotina
 569 *Protomyces lactucaeabilis* and the chytridiomycete *Blyttiomycetes helices*, presented remarkable
 570 differences in primary sequence features (**Data S3C**). These MsrBs lacked the resolving Cys at position
 571 97, which was substituted by a Ser or a Thr, like plant and human orthologs that use an unusual
 572 regeneration process [20,21,408]. Finally, another unusual feature was found in the MsrB from the
 573 taphrinomycotina *Protomyces lactucaeabilis*, with the location of the catalytic Cys in an HYCIN motif,
 574 instead of the typical RXCXN motif (**Data S3C**). Interestingly, searching in the NCBI nr database, we
 575 found 50 sequences, mainly from poorly characterized bacteria and archaea, that possessed this unusual
 576 motif (**Data S3D**). Considering His and Arg have similar physicochemical properties, we anticipate such
 577 non-canonical MsrBs might have conserved a catalytic activity.

578



579

580 **Figure 3. Protein sequence characteristics of canonical fungal Msrs.** In this representation of
 581 canonical MsrAs (**A**), MsrBs, (**B**) and fRMsrs (**C**), the catalytic Cys (in red), the resolving Cys (in green)
 582 and the residues previously shown to be involved in catalysis and/or substrate binding (in blue) are
 583 shown. The residues in purple are conserved in all canonical fungal Msrs. In **B**, the Cys residues labeled
 584 in orange correspond to Zn binding residues. The numberings are based on *S. cerevisiae* MsrA (**A**),
 585 MsrB (**B**) and fRMSr (**C**).

586

587 Very few fRMsrs have been characterized so far. However, sequence comparison studies and
 588 biochemical characterizations indicated that the catalytic Cys is located in the HIAC motif situated in
 589 the middle of the protein sequence and that the two resolving Cys are located ~30 and ~40 amino acids
 590 upstream in the N-terminal direction [7–9]. The 589 full length fRMSr sequences analyzed here had a
 591 single fRMSr domain. Their length varied from 77 to 394 amino acids, with variations in the size of the
 592 N-terminal extension (**Data S4A**). Most of the proteins (90 %) were predicted to be localized in the
 593 cytoplasm (**Fig. 2C, Data S4A**). A few proteins were predicted to be secreted (7 %), targeted to the

mitochondria (1 %), to other compartments or had no reliable prediction for subcellular targeting (2 %) (**Fig. 2C**, **Data S4A**). Similar to MsrAs and MsrBs, fRMsrs showed a strong conservation of the sequence features. Almost all (> 99 %) sequences possessed the catalytic Cys¹²⁵ included in a ¹²²H[I/V]XCD¹²⁶ motif and the resolving Cys in positions 91 and 101 (according to the *S. cerevisiae* fRMsrs residue numbering) (**Fig. 3C**, **Data S4B**). We also observed the strict conservation of Trp⁶⁶, Tyr⁷⁰, Glu¹³², Asp¹⁴⁹ and Asp¹⁵¹, previously shown to be involved in substrate binding and catalysis. Other important residues involved in substrate binding and catalysis [7] were also conserved or substituted by residues with similar properties in positions 82, 85, 94, 123, 129 and 132. The Pro⁸⁴, Gly⁸⁷, His¹²² and Val¹³⁴ were also strictly conserved in these canonical fungal fRMsrs (**Fig. 3C**, **Data S4B**). Only three fRMsrs sequences from the Agaricomycetes *Sclerotoderma citrinum*, *Dendrothele bispora* and *Pisolithus tinctorius* presented non-canonical characteristics. The first two lacked the potential resolving Cys¹⁰¹ and may be still able to reduce the free MetO, but in the latest, the catalytic Cys¹²⁵ was substituted by an Arg, likely precluding catalytic activity (**Data S4C**).

Altogether, these results uncover few proteins with non-canonical sequence features, but principally showed that for each Msr type, the residues involved in catalysis are globally conserved throughout the fungal kingdom.

610

611 3.3. The phylogenetic analysis of fungal Msrs revealed horizontal gene transfers from bacteria.

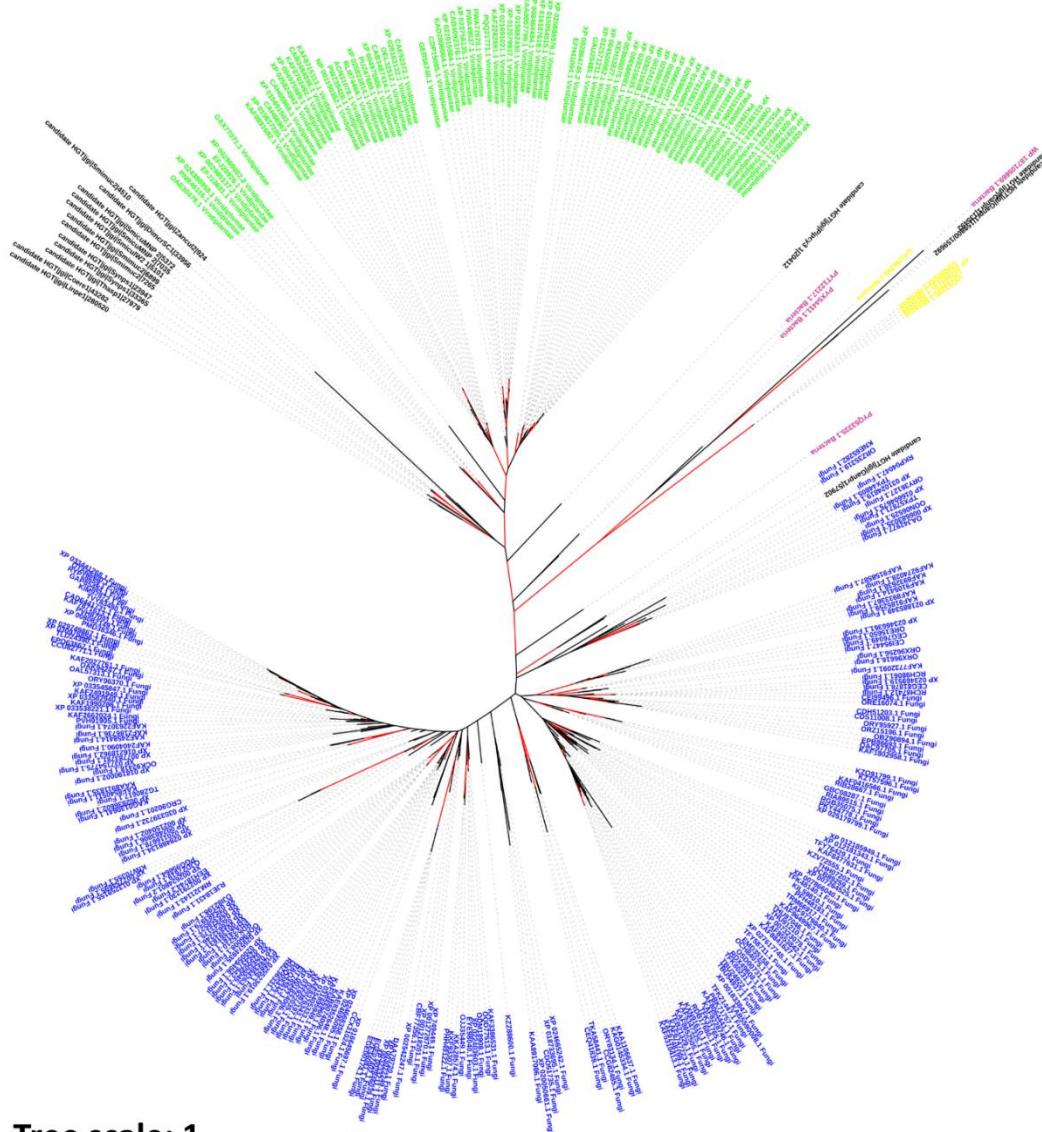
612 The phylogenetic relationship of fungal MsrAs globally matched the expected clustering for
613 early-diverging fungi, Ascomycota and Basidiomycota sequences (**Fig. S1-3**). However, a few
614 Ascomycota MsrA sequences clustered with Basidiomycota sequences (indicated by an asterisk on **Fig.**
615 **S1**). We also noticed the clustering of all the MsrB sequences from Pucciniomycotina (Basidiomycetes)
616 species with Ascomycota sequences (**Fig. S2**). Strikingly, the MsrB sequences from early-diverging
617 fungi did not group in a single cluster but were interspersed in clusters containing Basidiomycota or
618 Ascomycota sequences (**Fig. S2**). For fRMsrs, we observed three distinct clusters containing the protein
619 sequences from Basidiomycota, Ascomycota and early-diverging fungi, respectively (**Fig. S3**).
620 However, two sequences from early-diverging fungi were included in the cluster containing the
621 Ascomycota sequences (**Fig. S3**). Altogether, these results showed that the phylogeny of Msrs was
622 globally congruent with the phylogeny of the species, except for a few protein sequences.

623 Several discrepancies suggested that some Msrs could have arisen from horizontal gene transfer.
624 These discrepancies were: i) the positioning in phylogenetic clusters not reflecting the phylogeny of the
625 species from which they were isolated (**Fig. S1-3**), ii) the presence of non-canonical sequence features
626 (**Data S2-4**), and iii) the presence of *fRmsr* genes in the genome of three strictly anaerobic
627 neocallimastigomycetes, whereas other organisms from the same phylum had no *msr* genes (**Table 2**).

628 To evaluate the possibility of horizontal gene transfers, we selected all Msr sequences (67 MsrAs, 48
629 MsrBs and 8 fRMsrs) with one or more of these discrepancies and searched for their closest putative
630 homologs by BLASTP search in all organisms recorded in the NCBI nr database. We discarded all the
631 sequences for which the closest homologs were found in other fungal taxa. Indeed, because of the strong
632 conservation of the protein sequences, it would be difficult to ascertain the occurrence of fungus-to-
633 fungus horizontal gene transfers. We retrieved 17 MsrA, three MsrB and three fRMsrs protein sequences
634 for further analysis (**Fig. 4-6**). For each type of Msr, we used the selected protein sequences, together
635 with the respective 25 sequences with the highest identity score from fungi on the one hand and from
636 non-fungal organisms on the other hand for phylogenetic analyses (**Fig. 4-6**). The phylogenetic trees
637 highlighted two MsrA, three MsrB and three fRMsrs sequences, among the selected candidates,
638 clustering with bacterial and amoeba homologs (**Fig. 4-6, Table 3**). Moreover, we identified two
639 additional fungal fRMsrs, not included in our primary data set, which clustered with bacterial proteins
640 (**Fig. 6, Table 3**). Except for two MsrBs from Orbiliomycetes (Ascomycota), all these Msr sequences
641 were from early-diverging fungi (**Table 3**). Interestingly, in the cases of MsrA and MsrB, they were all
642 from organisms having another gene coding for a canonical enzyme in their genomes (**Data S1-3**).

643 For MsrAs, a potential horizontal gene transfer was observed for the two selenocysteine-
644 containing enzymes from *Gonapodya prolifera* (**Table 3**). Very interestingly, their closest homolog was
645 also a selenocysteine-containing MsrA, from the bacteria *Alkaliphilus oremlandii*. Bacteria from the
646 genus *Alkaliphilus* are found in sediments and ponds [410], and *Gonapodya prolifera* occurs on fruits
647 submerged into ponds [411]. The presence of both organisms in a same ecological niche could have
648 favored the horizontal gene transfer. The similarity between the two *Gonapodya prolifera* Sec-MsrA
649 suggested they arose from one horizontal gene transfer event followed by a gene duplication (**Fig. 4,**
650 **Fig. S4**).

651



652

653 **Figure 4. Phylogenetic analysis of MsrA candidates to horizontal gene transfer.** The fungal MsrAs
 654 tested for horizontal gene transfer (in black) are from the following fungal genomes (protein accessions
 655 in brackets): *Piptocephalis cylindrospora* RSA 2659 single-cell v3.0 (20412); *Syncephalis*
 656 *pseudoplumigaleata* Benny S71-1 single-cell v1.0 (33365; 23947); *Thamnocephalis sphaerospora* RSA
 657 1356 single-cell v1.0 (27979); *Coemansia reversa* NRRL 1564 v1.0 (43282); *Dimargaris cristalligena*
 658 RSA 468 single-cell v1.0 (33956); *Linderina pennispora* ATCC 12442 v1.0 (280520); *Smittium culicis*
 659 GSMNP (5372; 7035); *Smittium culicis* ID-206-W2 (5101; 8129); *Smittium mucronatum* ALG-7-W6
 660 (4510; 6899; 7265); *Zancudomyces culisetae* COL-18-3 (924) and *Gonapodya prolifera* v1.0 (135492;
 661 159800/159692; 57902). The MsrA sequences from fungi, plants, amoeba and bacteria are in blue,
 662 green, yellow and purple, respectively. The phylogenetic tree was built with RAxML v. 8.2 [348] and
 663 represented using iTOL (<https://itol.embl.de/>) [412]. The branches with bootstrap values over 70 are in
 664 red.

665

Table 3. Potential horizontal *msr* gene transfers from bacteria to fungi

Genome	Candidate gene				Mean of all genes from the scaffold				Organisms with the closest Msr homolog		
	Protein accession (Scaffold number)	Nb. of exons	GC content (%)	GC third base (%)	Nb. of exons	GC content (%)	GC third base (%)				
MsrA											
Monoblepharidomycetes											
<i>Gonapodya prolifera</i> v1.0	135492 ^a (36)	3	56.4	61.3	5.7 ± 4.8	56.5 ± 3.4	65.2 ± 8.5	<i>Alkaliphilus oremlandii</i> OhILAs	<i>Alkaliphilus oremlandii</i> OhILAs		
<i>Gonapodya prolifera</i> v1.0	159800/ 159692 ^a (105)	4	58.9	69.6	5.8 ± 3.8	57.3 ± 4.2	67.4 ± 8.2				
MsrB											
Orbiliomycetes											
<i>Arthrobotrys oligospora</i> ATCC 24927	9001 (00170)	1	55.6	56.5	3.0 ± 2.3	48.0 ± 2.4	48.0 ± 4.6	<i>Sphingomonas</i> bacterium	<i>Sphingomonas</i> bacterium		
<i>Monacrosporium haptotylum</i> CBS 200.50	10089 (00497)	1	55.8	58.0	4.3 ± 4.3	49.0 ± 2.7	52.9 ± 5.3	<i>Sphingomonas</i> bacterium	<i>Sphingomonas</i> bacterium		
Taphrinomycotina											
<i>Protomyces lactucaeabilis</i> 12-1054 v1.0	391028 (3)	1	54.9	52.1	1.6 ± 1.0	52.6 ± 1.9	54.4 ± 5.9	<i>Calothrix parasitica</i>	<i>Calothrix parasitica</i>		
fRMsR											
Neocallimastigomycetes											
<i>Anaeromyces robustus</i> v1.0	328892 ^b (243)	1	30.1	14.8	4.2 ± 4.5	26.3 ± 5.0	11.8 ± 3.3	<i>Pseudobutyrivibrio</i> sp. LB2011	<i>Pseudobutyrivibrio</i> sp. LB2011		
<i>Neocallimastix californiae</i> G1 v1.0	697339 (7)	1	27.5	9.8	4.6 ± 4.4	28.8 ± 4.4	16.7 ± 5.4	<i>Erysipelotrichaceae</i> bacterium	<i>Erysipelotrichaceae</i> bacterium		
<i>Orpinomyces</i> sp.	1191427 (d_7180000085738)	1	34.8	22.5	3.0 ± 2.9 ^d	27.2 ± 5.4 ^d	16.0 ± 7.8 ^d	<i>Pseudobutyrivibrio ruminis</i>	<i>Pseudobutyrivibrio ruminis</i>		
<i>Piromyces</i> sp.	AWI66787.1 ^{b,c}	1	31.3	9.9	n.d.	33.2 ± 5.7	23.3 ± 11.1	<i>Pseudobutyrivibrio ruminis</i>	<i>Pseudobutyrivibrio ruminis</i>		
Mucoromycotina											
<i>Mortierella</i> sp. GBA39	KAF9144191.1 ^c (77)	1	55.2	62.4	1.4 ± 0.7	57.1 ± 3.5	71.3 ± 6.0	<i>Paenibacillus rhizosphaerae</i>	<i>Paenibacillus rhizosphaerae</i>		

667 ^a For both *Gonapodya prolifera* selenocysteine-containing MsrAs, the MycoCosm accession numbers presented here refer to
668 only a part of the proteins. Their reconstructed complete gene structures are presented in Fig. S4.

669 ^b The horizontal gene transfers of *Anaeromyces robustus* and *Piromyces* sp. fRmsr genes have been shown recently [413].

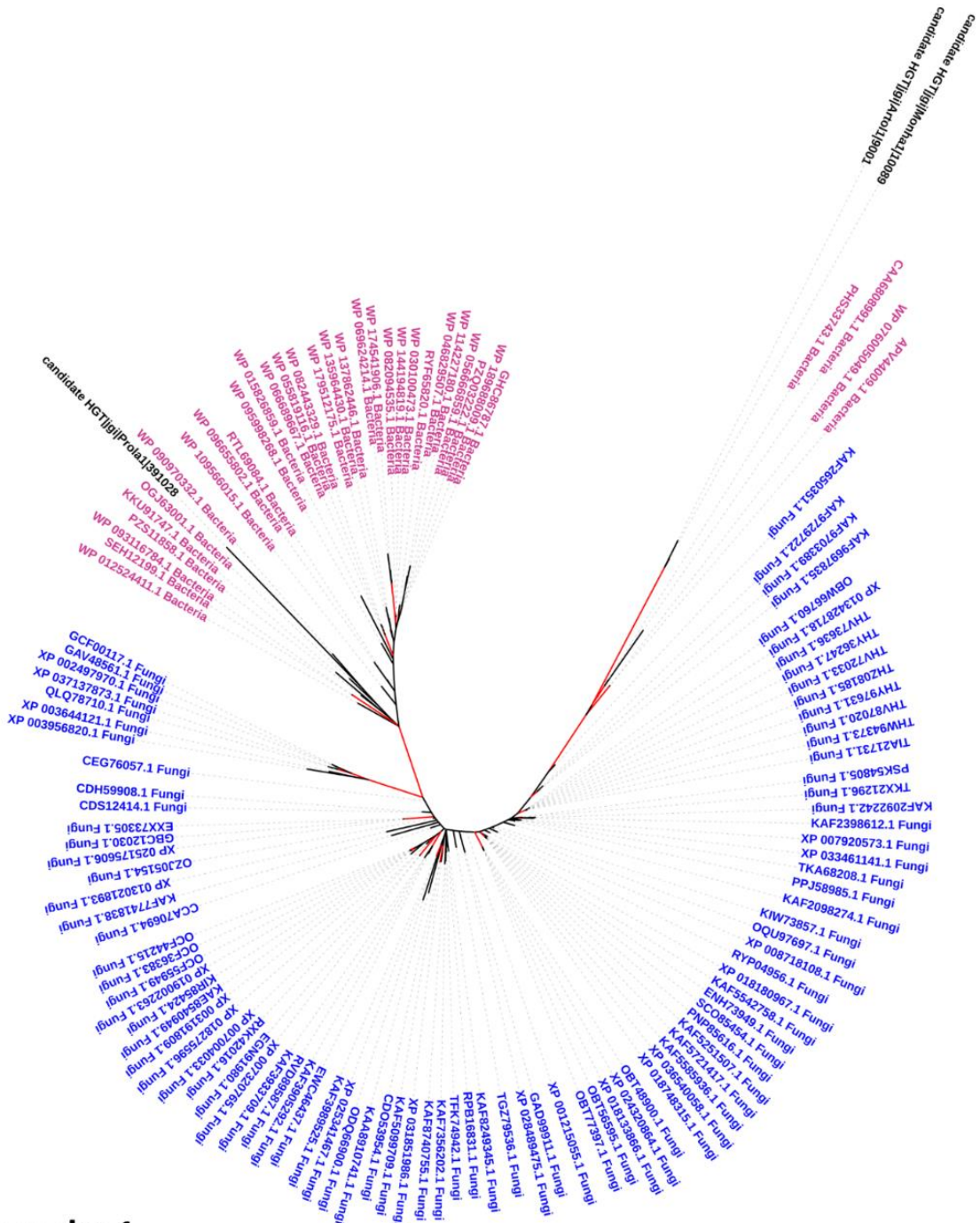
670 ^c The protein accessions are from NCBI (<https://www.ncbi.nlm.nih.gov/protein>).

671 ^d The positions of *Orpinomyces* sp. genes on the scaffolds are not publically available. The calculations were made on the
672 complete genome.

673 n.d., not determined.

674

675 In the case of MsrBs, the three selected sequences clustered with bacterial MsrBs (**Fig. 5**). The
676 two MsrBs from the Orbiliomycetes species *Arthrobotrys oligospora* and *Monacrosporium haptotylum*
677 had MsrBs from *Sphingomonas* bacteria as closest homologs. Both fungi trap nematodes in soil [197],
678 and sphingomonads have been isolated from many different land and water habitats [414], indicating
679 that the co-occurrence in the same habitat of a sphingomonad donor and a common ancestor of *A.*
680 *oligospora* and *M. haptotylum* is plausible, which could have allowed horizontal gene transfers. Another
681 potential horizontal gene transfer for MsrB was identified, from a donor cyanobacterium related to the
682 marine *Calothrix parasitica* to the plant pathogen fission yeast *Protomyces lactucaeabilis*
683 (Taphrinomycotina) (**Table 3**). In this case, the ecology of the extant candidate donor and acceptor do
684 not support the co-occurrence of both organisms in a same ecological niche.
685



Tree scale: 1 —

686

687 **Figure 5. Phylogenetic analysis of MsrB candidates to horizontal gene transfer.** The fungal MsrBs
 688 tested for horizontal gene transfer (in *black*) are from the following genomes (protein accessions in
 689 brackets): *Arthrobotrys oligospora* ATCC 24927 (9001); *Monacrosporium haptotylum* CBS 200.50
 690 (10089) and *Protomyces lactucaedebilis* 12-1054 v1.0 (391028). The MsrBs sequences from fungi and
 691 bacteria are in *blue* and *purple*, respectively. The phylogenetic tree was built with RAxML v. 8.2 [348]
 692 and represented using iTOL (<https://itol.embl.de/>) [412]. The branches with bootstrap values over 70
 693 are in *red*.

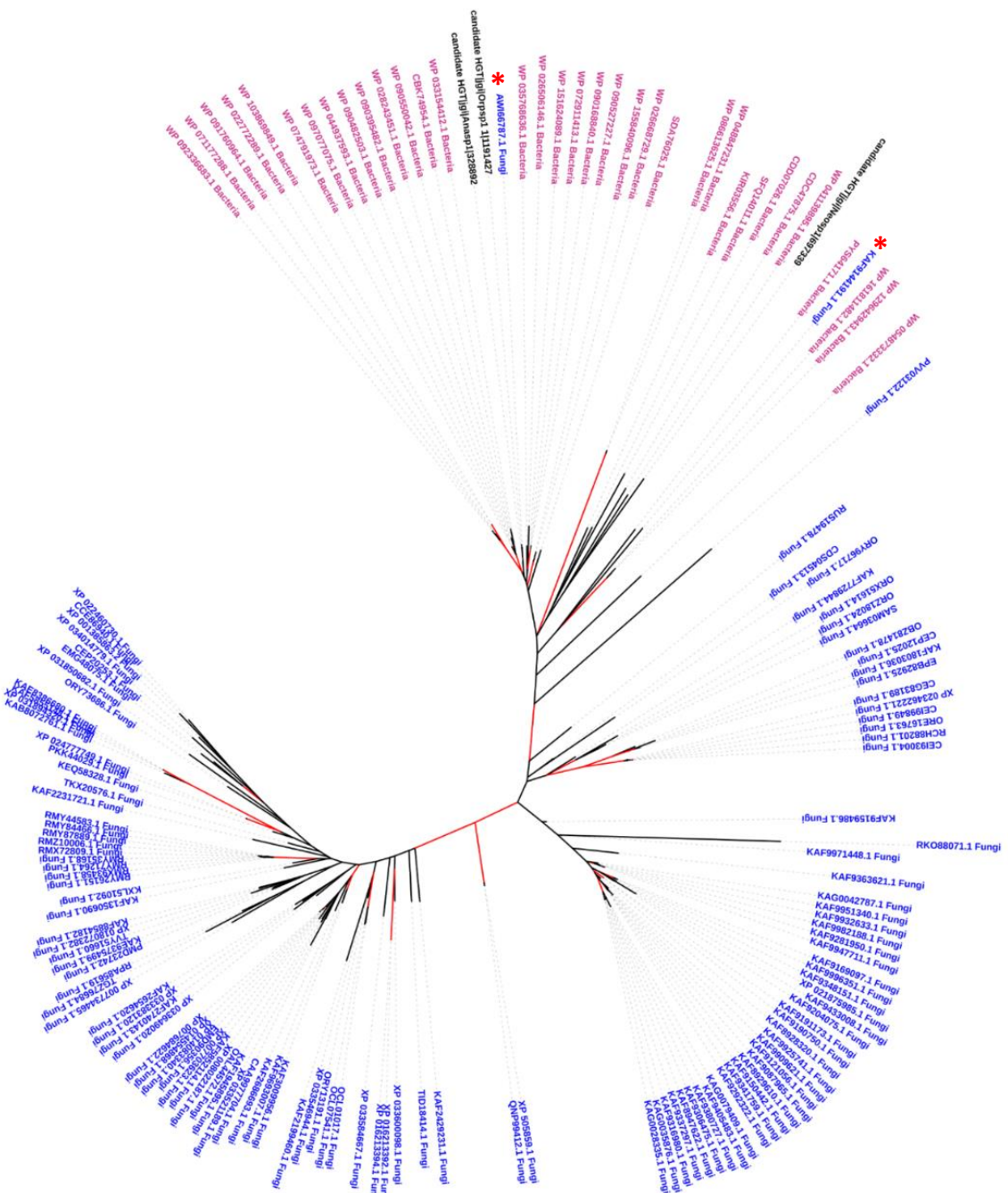
694

695

696 The three fRMsrs we selected as candidates to horizontal gene transfer were from the
697 neocallimastigomycetes species *Anaeromyces robustus*, *Neocallimastix californiae* and *Orpinomyces*
698 sp. (**Table 3**). In our phylogenetic analysis, they clustered with sequences of bacteria from the phylum
699 Firmicutes which, similarly to neocallimastigomycetes, live in the gut of ruminants (**Fig. 6**).
700 Interestingly, two other fungal fRMsrs sequences, one from *Piromyces* sp. and one from *Mortierella* sp.
701 GBA39, were also present in the same cluster. These two sequences were omitted from our genomic
702 search because of the absence of the corresponding genomes in the MycoCosm database. Of note, the
703 horizontal transfer of *fRmsr* genes from Firmicutes to neocallimastigomycetes has been shown recently
704 for *Anaeromyces robustus* and *Piromyces* sp. [413]. Altogether, these results strongly argue for the fact
705 that the presence of *fRmsr* in the genomes of these anaerobic fungi arose from horizontal gene transfers
706 from Firmicutes (**Table 3**).

707 In addition to the phylogenetic method, we performed a parametric analysis of the numbers of
708 exons, the overall GC contents of their coding sequence (CDS), and the GC contents of the third position
709 of each codon to identify potential bias as supporting arguments of horizontal gene transfer (**Table 3**).
710 The comparison of these parameters with those of the other genes included in the same genomic scaffold
711 were shown as potential indicators of horizontal gene transfers [415]. Excepted for the two *Gonapodya*
712 *prolifera msrA* genes, which had no distinguishable values from the other genes included in their
713 scaffolds, all other genes coding MsrBs or fRMsrs, had extreme values for at least two of the three
714 parameters considered (**Table 3**). First of all, all these eight genes were made of a single exon, like
715 bacterial genes. The global GC content of *Arthrobotrys oligospora* and *Monacrosporium haptotylum*
716 *msrB* genes coding sequences were the highest of all those included in their scaffold. Similarly, the
717 global GC content of *Protomyces lactucaeabilis msrB*, *Anaeromyces robustus fRmsr* and *Orpinomyces*
718 sp. *fRmsr* were among the highest (**Table 3**). In the cases of *fRmsrs* from *Neocallimastix californiae*,
719 *Piromyces* sp. and *Mortierella* sp., the GC contents of the third base of the codons were among the
720 lowest by comparison with the other genes present in the scaffold. Noticeably, we also observed a lower
721 GC content at the third position of the codons in the *fRmsr* gene from *Piromyces* sp., for which the
722 acquisition via HGT was previously shown [412].

723 Altogether, these analyses showed that the phylogeny of fungal Msrs globally matched the
724 phylogeny of the fungi, in accordance with vertical inheritance of the genes from bacterial ancestors
725 [8,9,24,26]. However, a few horizontal gene transfer events occurred in each Msr family, that
726 contributed to the distribution of Msr genes in extant species.



Tree scale: 1 —

727

Figure 6. Phylogenetic analysis of fRMSr candidates to horizontal gene transfer. The fungal fRMSrs tested for horizontal gene transfer (in *black*) are from the following genomes (protein accessions are in brackets): *Anaeromyces robustus* v1.0 (328892); *Neocallimastix californiae* G1 v1.0 (697339) and *Orpinomyces sp* (1191427). The fRMSrs sequences from fungi and bacteria are in *blue* and *purple*, respectively. Red stars indicate fRMSr genes possibly acquired by horizontal gene transfer, which were not identified in our genomics search but found as homologs of the selected fungal fRMSr candidates. The phylogenetic tree was built with RAxML v 8.2 [348] and represented using iTOL (<https://itol.embl.de/>) [412]. The branches with bootstrap values over 70 are in *red*.

736

737

738 **4. Discussion**

739 This global genomic search for *msr* genes in nearly 700 fungal genomes covering the fungal
740 kingdom showed that most fungi contain one gene coding for each thiol-oxidoreductase type, i.e., MsrA,
741 MsrB and fRMSr. The phylogenetic analyses and inspection of protein sequence features revealed that
742 Msrs from each type are globally strongly conserved across the fungal kingdom. This is consistent with
743 the prokaryotic origin of these genes. Yet, the identification of *fRmsr* genes in almost all genomes across
744 the fungal kingdom was surprising. Indeed, no *fRmsr* genes were identified so far from multicellular
745 eukaryotes [8]. Very likely, the most obvious reason was the smaller number of eukaryotic genomes
746 considered (i.e. 160), and the smaller number of fungal genomes available at the time of the previous
747 study [8,50]. Because the number of sequenced genomes has increased dramatically in the last years,
748 we took the opportunity of this study to reevaluate the presence of *fRmsr* genes in other multicellular
749 eukaryotes. We searched for *fRmsr* genes in the plant and animal genomes available in the nr database
750 of NCBI by BLAST search and found only a handful of multicellular eukaryotes apparently possessing
751 a gene coding for a fRMSr (**Table S1**). These organisms, a plant and a few insects, are not
752 phylogenetically related, indicating that the gene is not conserved in their lineage. This highlights the
753 prevalence of *fRmsr* genes in the fungal kingdom, as opposite to their extremely low occurrence in other
754 multicellular eukaryotes. What would be the advantage for the fungi to produce fRMSr whereas other
755 multicellular organisms do not? An obvious possibility would be to protect the intracellular free Met
756 during exposure to oxidative constraints as it was shown for *S. cerevisiae* [8]. Another possibility would
757 be to allow the reduction of the free MetO coming from the external environment or after the degradation
758 of proteins assimilated by the fungi and its use in protein synthesis and in sulfur metabolism. Combined
759 with MsrA, the presence of fRMSr could virtually allow to reduce the complete pool of MetO coming
760 from the external environment. This hypothesis arose from the observation of the presence of *fRmsr*
761 genes in four Neocallimastigomycetes species that lacked MsrA and MsrB. Living in the anaerobic
762 environment of ruminant gut, the fungi very likely do not suffer oxidative stress and the fRMSr could
763 reduce the MetO coming from the animal's food. Interestingly, the *fRmsr* genes in these
764 Neocallimastigomycetes have been probably acquired through horizontal gene transfer, reinforcing the
765 idea that a fRMSr would provide a selective advantage in an environment unlikely to generate oxidative
766 constraints.

767 We found few variations in the numbers of gene copies of each type of Msr, with only 74
768 genomes lacking one gene, and 72 genomes with an extra copy of at least one *msr* gene. In both cases,
769 it roughly corresponded to 10 percent of the analyzed genomes. Our study revealed the absence of both
770 MsrA and MsrB in 15 species of fungi, twelve of these being also devoid of fRMSr. The lack of all Msr
771 appears to be related to the fungal lifestyles, since it concerned the nine Microsporidia, and the
772 *Pneumocystis jirovecii* species, which live as intracellular parasites of metazoans. These 10 species have
773 small genomes with reduced numbers of genes, from 1,831 for *Encephalitozoon romaleae* SJ-2008 to

774 3,632 for *Enterocytozoon bieneusi* H348, as compared to the average ~12,200 genes for the fungal
775 genomes considered in this study. It has been proposed that the intracellular lifestyle allowed for genome
776 compaction and gene loss, making the fungi highly dependent on the infected cell for numerous
777 biochemical pathways [27]. Our results suggest the fungi could also rely on the host detoxification system
778 for the protection against oxidative constraints and limitation of protein oxidation. The absence of Msr
779 was also observed in Neocallimastigomycetes, which live in the anaerobic environment of ruminant gut,
780 where the protection of proteins from oxidative damage is likely not crucial. This hypothesis is
781 reinforced by the fact that the numbers of genes in Neocallimastigomycetes genomes is above the
782 average of the analyzed fungi (~15,500) and thus, the loss of *msr* genes was not due to a global genome
783 compaction, but potentially due to the lack of selection pressure. Besides these species, for which the
784 lack of Msr is consistent with the lifestyle, we also identified 12 other species for which *msrA* or *msrB*
785 genes were absent from the genomes. As they are very likely living in aerobic conditions and are
786 probably exposed to protein oxidation, the lack of MsrA or MsrB is surprising and we cannot exclude
787 that the missing genes are due to incomplete genome sequencing. On the other hand, most of the 37
788 analyzed genomes from Glomeromycotina, Pucciniomycotina (*Melampsora* and *Puccinia*) and
789 Taphrinomycotina (*Schizosaccharomyces*) species lacked the fRMs. These fungi may have lost the
790 capacity to reduce the free Met-R-O, as it has been shown for mammals [416], or the MsrB might have
791 significant activity on the free Met-R-O, as suggested for plants [417].

792 Regarding the presence of supernumerary copies of *msr* genes in few genomes, it seems not to
793 be related to the phylogeny, nor to the numbers of genes per genome, as the genomes of fungi having
794 two copies or more of one *msr* gene contain an average number of ~12,700 genes, similar to the average
795 number of genes in the considered genomes (~12,200). In most cases, the presence of several copies
796 was due to gene duplications, as indicated by the close phylogenetic relationships of paralogous proteins
797 and the conservation of canonical features. However, in few fungi, the presence of extra copies was due
798 to horizontal gene transfers, like for *msrsA* genes in *Gonapodya prolifera*, and *msrB* genes in
799 *Arthrobotrys oligospora*, *Monacrosporium haptotylum* and *Protomyces lactucaeabilis*. Together with
800 the horizontal transfers of *fRmsr* genes in *Neocallimastigomycetes*, most of these events were strongly
801 supported by the phylogeny and parametric values as well as by shared ecological niches for donor
802 bacteria and recipient fungi. The case of *Gonapodya prolifera msrA* was remarkably interesting as it
803 illustrated a prokaryote-to-eukaryote transfer of a selenoprotein gene. If such a transfer was observed
804 from bacteria to archaea [418], to our knowledge, the bacteria-to-eukaryote transfer of a selenoprotein
805 gene has never been described. As this fungus possesses an eukaryotic selenocysteine insertion
806 machinery [402], we hypothesize that the transferred prokaryotic *msrA* gene was compatible with the
807 eukaryotic machinery, or that it was modified after the transfer to allow proper Sec insertion. In all cases,
808 the presence of several Msr gene copies could allow a beneficial increase in gene dosage or,
809 alternatively, different spatial and temporal expression profiles during the life cycle of the fungi through

810 distinct transcriptional activities. It could also allow different subcellular targeting as shown for plant
811 Msrs [419]. Indeed, in 23 fungal genomes, out of the 59 having more than one *msrA* gene, the MsrAs
812 were predicted to be addressed to different subcellular compartments. Similarly, MsrBs from 7 genomes
813 out of the 15 with more than one gene were predicted to be addressed to different cellular compartments.
814 Conversely, all the multicopy fRMsrs were predicted to be cytoplasmic. Our results indicate that, as
815 experimentally demonstrated for yeast [39], most fungi might not have the ability to reduce the two
816 diastereomers of protein-bound MetO in each subcellular compartment, suggesting variations in protein
817 oxidation in the different cellular compartments. Another interesting aspect is the presence of potentially
818 secreted Msrs in few fungal species, mostly from the Ascomycota genera *Aspergillus* and *Penicillium*.
819 Although it remains to be demonstrated experimentally, the presence of Msrs in the extracellular
820 environment of the fungal cells could help to protect from oxidation the numerous secreted
821 carbohydrate-active enzymes ('CAZymes') and other enzymes used by saprotrophic and pathogen fungi
822 to degrade plant cell walls or insect chitins [231].

823 Finally, we observed that most MsrAs, MsrBs and fRMsrs had conserved canonical features,
824 but we also highlighted potentially interesting discrepancies. For the three types of Msrs, a few
825 sequences lacked the resolving Cys at conserved positions and might use alternative Cys residues for
826 the regeneration of their activity, whereas others possessed only the catalytic Cys and might be
827 regenerated by the direct reduction of the sulfenic acid formed after MetO reduction, as shown for Msrs
828 from other organisms [20,408]. The presence of Sec in two *Gonapodya prolifera* MsrAs might confer
829 them a catalytic advantage, similarly to other Sec-MsrAs [400]. We also observed that ~5% of fungal
830 MsrAs harbor a GCYW motif containing the catalytic Cys, where the Tyr residue replaces the Phe
831 residue of the canonical GCFW motif. To our knowledge, no MsrA with such motif has been described
832 to date. Because of the similarity between Phe and Tyr physicochemical properties, this substitution
833 should not prevent the catalysis, but it could induce a change in substrate specificity, as demonstrated
834 for the *E. coli* MsrA. Indeed, direct evolution assays showed that the substitution of the Phe by a Leu
835 conferred to the enzyme the capacity to efficiently reduce alkyl-aryl sulfoxides [420]. Similarly, the
836 atypical HYCIN motif of the *Protomyces lactucaeabilis* MsrB could modify the enzyme substrate
837 specificity. The substitution of residues around the catalytic Cys of Msrs could affect the specificity for
838 oxidized protein substrates or free MetO, but also potentially confer to the enzymes the ability to reduce
839 other molecules such as the oxidized thioether-containing metabolites involved in sulfur metabolism
840 (e.g S-adenosyl homocysteine, methylthio-ribose...) [421]. It would be interesting to evaluate the
841 catalytic properties of these atypical Msrs on various sulfoxide-containing substrates.

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843

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851

852 **Declaration of competing interest.**

853 None.

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