

Paralog re-emergence: a novel, historically contingent mechanism in the evolution of antimicrobial resistance

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MBE Advance Access published April 26, 2014 Paralog Re-Emergence: A Novel, Historically Contingent Mechanism in the Evolution of Antimicrobial Resistance

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Abstract

Evolution of resistance to drugs and pesticides poses a serious threat to human health and agricultural production. CYP51 encodes the target site of azole fungicides, widely used clinically and in agriculture. Azole resistance can evolve due to point mutations or overexpression of CYP51, and previous studies have shown that fungicide-resistant alleles have arisen by de novo mutation. Paralogs CYP51A and CYP51B are found in filamentous ascomycetes, but CYP51A has been lost from multiple lineages. Here, we show that in the barley pathogen *Rhynchosporium commune*, re-emergence of CYP51A constitutes a novel mechanism for the evolution of resistance to azoles. Pyrosequencing analysis of historical barley leaf samples from a unique long-term experiment from 1892 to 2008 indicates that the majority of the *R. commune* population lacked CYP51A retains the same substrate as CYP51B, but with different transcriptional regulation. Phylogenetic analyses show that the origin of CYP51A far predates azole use, and newly sequenced *Rhynchosporium* genomes show CYP51A persisting in the *R. commune* lineage rather than being regained by horizontal gene transfer; therefore, CYP51A re-emergence provides an example of adaptation to novel compounds by selection from standing genetic variation.

Key words: standing variation, gene duplication, resistance, fungicides, triazoles, Rhynchosporium.

Introduction

Resistance to drugs and pesticides is a major challenge in medicine (McKenna 2013) and agriculture (Heckel 2012). Understanding the evolutionary mechanisms involved in the emergence and spread of resistance can aid resistance risk assessment and management (MacLean et al. 2010). One relevant evolutionary question is the extent to which populations adapt to novel environments, such as a new biocide, through de novo mutations or through selection from standing genetic variation, as this will affect the probability and speed of emergence of resistant alleles (Hermisson and Pennings 2005). Previous studies have indicated de novo origins for fungicide resistance mutations (Torriani et al. 2009; Camps et al. 2012). Selection from standing variation can be inferred from the population genetic signature of selection, or by tracing the history of the selected allele (Barrett and Schluter 2008). In this study, we have taken the second approach, following the evolutionary history of CYP51A in Rhynchosporium commune through comparative genomics and analysis of historical infected plant samples.

Duplicate genes have long been considered an important source of genetic adaptive potential (Ohno 1970; Zhang 2003; Dittmar and Liberles 2010). Extensive research has considered the origin and fate of duplicated genes, but focusing on the likelihood of emergence of copy number variants (Lynch and Conery 2000); loss or change of function pseudogenization, neofunctionalization, through or subfunctionalization (Conant and Wolfe 2008; Innan and Kondrashov 2010); and mechanisms of functional divergence before or after duplication (Hughes 1994; Näsvall et al. 2012); around the time of duplication under given conditions of selection and drift, rather than future changes if selective pressures alter. In this study, the use of historical samples enables changes in duplicate gene frequency over time to be measured directly, revealing responses to changes in selection.

The azoles are widely used as both clinical and agricultural fungicides (Kelly et al. 1993). Clinically, the importance of antifungal therapies has increased as HIV, transplant antirejection drugs, and some cancer treatments leave patients immunocompromised and vulnerable to fungal infections (Denning and Hope 2010). In agriculture, crop diseases result in yield losses of 10–15% of global food production (Oerke and Dehne 2004; Strange and Scott 2005), a continual obstacle to feeding a growing population. In 2005, azoles represented 25.3% of the \$8.9 billion global agricultural fungicide

© The Author 2014. Published by Oxford University Press on behalf of the Society for Molecular Biology and Evolution. This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http:// creativecommons.org/licenses/by-nc/3.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited. For commercial re-use, please contact journals.permissions@oup.com market (Morton and Staub 2008). The target site of the azoles is fungal CYP51. CYP51 is a cytochrome P450, sterol 14 α -demethylase, involved in the biosynthesis of ergosterol in fungi, cholesterol in mammals, and sitosterol in plants (Lepesheva and Waterman 2007).

Recently, some ascomycete fungi have been found to possess multiple CYP51 paralogs, with two paralogs, CYP51A and CYP51B, in Aspergillus fumigatus (Mellado et al. 2001), A. nidulans (Da Silva Ferreira et al. 2005), and Magnaporthe oryzae (Yan et al. 2011), and three in A. oryzae (Da Silva Ferreira et al. 2005) and Fusarium spp. (Deng et al. 2007). The reasons why some species possess multiple CYP51 paralogs, and others only one, are not yet known. Knocking out the CYP51A paralog from wild-type isolates of A. fumigatus (Mellado et al. 2005) and Fusarium graminearum (Fan et al. 2013) resulted in increased azole sensitivity, suggesting that the presence of additional CYP51 paralogs can confer reduced intrinsic azole sensitivity.

Acquired reductions in sensitivity to azoles may result from mutations or overexpression of CYP51. Various point mutations, with quantitative, interacting effects on azole sensitivity, have been reported in the single CYP51 paralogs of the clinical pathogen Candida albicans (Morio et al. 2010) and the wheat pathogen Mycosphaerella graminicola (Cools et al. 2010). Reduced azole sensitivity due to constitutive overexpression of CYP51 was first reported in citrus green mould, Penicillium digitatum (Hamamoto et al. 2000). A combination of promoter changes resulting in overexpression and point mutations in the coding sequence has been reported in the CYP51A paralog in clinical isolates of A. fumigatus (Mellado et al. 2007). Nontarget-site mechanisms, such as enhanced efflux (de Waard and van Nistelrooy 1980) or altered sterol metabolism (Kelly et al. 1995), may also reduce azole sensitivity. Reduced sensitivity to azoles has now been reported in the field or in field isolates of over 20 plant pathogens (Fungicide Resistance Action Committee 2013, www.frac. info, last accessed April 16, 2014).

Rhynchosporium commune Zaffarano, McDonald, and Linde is an ascomycete fungal pathogen causing barley leaf blotch or scald. Previously known as R. secalis (Oud.) Davis (Caldwell 1937), it has recently been shown that R. secalis s.l. comprises three distinct host-specialized lineages, with R. secalis s.s. now referring only to the rye-infecting species, R. agropyri infecting Agropyron spp. and R. commune infecting barley (Zaffarano et al. 2011). Rhynchosporium commune causes necrotic lesions on infected leaves, resulting in grain yield reductions of up to 30% if untreated (Mayfield and Clare 1991). Although some barley varieties provide partial resistance, control is still heavily reliant on fungicides. In 2010, winter barley crops in the United Kingdom received an average of 2.1 fungicide applications, with 98.5% of crop area treated, and 54% of treatments containing an azole (Garthwaite et al. 2011).

The first azole to be used on barley in the United Kingdom was introduced in the mid-1970s. By 1985, surveys of azole sensitivity in *R. commune* revealed a 5- to 10-fold decrease in sensitivity to the earlier azoles relative to baseline testing in 1975–1981 (Kendall et al. 1993). Since the introduction of

epoxiconazole in the late 1990s and prothioconazole in the mid-2000s, further sensitivity shifts have affected these compounds, but they retain some effectiveness in the field (Oxley et al. 2008). The mechanisms responsible for these azole sensitivity shifts in *R. commune* were not known.

The classical experiments at Rothamsted Research in Hertfordshire, UK, are field experiments studying the longterm effects of different agricultural practices and other factors affecting crop production and the environment. Broadbalk is the world's oldest continually running agricultural experiment, growing winter wheat since 1843, and the Hoosfield spring barley experiment started 9 years later. Throughout the history of the experiment, plant and soil samples have been retained in the Rothamsted Archive. This enables the use of new analytical techniques on historical samples, such as quantitative polymerase chain reaction (qPCR) to quantify pathogen levels (Bearchell et al. 2005), or pyrosequencing to quantify alleles within populations (Gruber et al. 2002).

Rhynchosporium commune isolates were characterized for sensitivity to four azoles, and a sensitivity shift was confirmed. Two CYP51 genes were sequenced from R. commune and shown to be paralogs CYP51A and CYP51B (sensu Mellado et al. 2001), along with a pseudogene, CYP51A-p. The azole sensitivity shift was not correlated with point mutations or constitutive overexpression of either paralog, but CYP51A was present only in less sensitive isolates. CYP51A was upregulated in response to tebuconazole and complements yeast CYP51; therefore, CYP51A, where present, encodes an additional sterol 14- α demethylase with inducible overexpression. Phylogenetic analysis of fungal CYP51s indicates that CYP51A and CYP51B originate from a gene duplication event basal to the filamentous ascomycetes, followed by multiple losses of CYP51A. Pyrosequencing analysis of samples from the Hoosfield long-term spring barley experiment revealed that CYP51A had declined in the R. commune population by the start of the 20th century, but re-emerged in the mid-1980s under selection by azoles.

Results

Ascomycete CYP51 Paralogs

CYP51 paralogs from fungal genomes on the Broad Institute server, along with R. commune CYP51 genes and pseudogene, are shown in supplementary figure S1, Supplementary Material online, with gene names according to Nelson (2009). The maximum-likelihood phylogenetic tree is summarized in figure 1 and shown in full in supplementary figure S1, Supplementary Material online. Basidiomycetes and hemiascomycete yeasts each possess a single CYP51 paralog, apart from an independent duplication in Coprinopsis cinerea. CYP51 gene duplication took place around the origin of the filamentous ascomycetes, forming clades CYP51A, CYP51B, and CYP51C. All species studied have retained a CYP51B paralog, but CYP51A has been lost from multiple lineages, and CYP51C is found only in Fusarium spp. Further duplications of CYP51A and CYP51B have taken place in some Aspergillus species. The A. flavus "CYP51C" described by Liu et al.



Fig. 1. Summary cladogram of maximum-likelihood phylogeny of coding nucleotide sequences of fungal and outgroup CYP51 genes, rooted with a CYP7 outgroup. The full phylogeny is shown in supplementary figure S1, Supplementary Material online.

(2012) appears from this phylogeny to be a second CYP51A. One R. commune CYP51 falls within the CYP51B clade, where its closest relatives are Botrytis cinerea and Sclerotinia sclerotiorum CYP51B. The other R. commune CYP51 falls within the CYP51A clade, where its closest relatives are sordariomycetes including M. oryzae as the sequenced leotiomycetes lack CYP51A. The R. commune CYP51 pseudogene is sister to R. commune CYP51A, and is therefore referred to as CYP51A-p.

Triazole Sensitivity in R. commune

Sensitivity of 57 R. commune isolates to the azole fungicides propiconazole, tebuconazole, epoxiconazole, and prothioconazole was measured by in vitro bioassay (supplementary table S2, Supplementary Material online). Triazoles demonstrate partial cross-resistance, so a principal component analysis (PCA) was carried out (fig. 2). 75.4% of variance was encompassed by PC1, representing overall positive crossresistance against the four triazoles, with all loadings positive. A further 17.2% of variance was encompassed by PC2, with positive loadings for propiconazole and tebuconazole but a negative loading for prothioconazole, showing incomplete cross-resistance between prothioconazole and the other azoles. The most sensitive group of isolates can be separated from the others by PC1 alone, indicating an initial sensitivity shift conferring positive cross-resistance against all four azoles tested. A further shift from intermediate to less-sensitive isolates, and further variation within the less-sensitive group,



Fig. 2. Biplot of principal component 2 against principal component 1 from PCA of triazole $\log_{10}[EC_{50}]$ data for 61 *Rhynchosporium commune* isolates. Percentage variation and loadings are listed in supplementary table S3, Supplementary Material online. Black crosses indicate isolates selected for sequence analysis. 1: RS 219; 2: K1124; 3: FI12-63; 4: R 9528.4; 5: R 9522.3; 6: QUB 30-10; 7: GKII 18-3-2; 8: GKII 18-2-3; 9: SAC 0003 1.4.8; 10: 788; 11: SAC 09/943/14; 12: QUB 12-3; 13: OSA 28-2-2; 14: RS 783. Bold numbers indicate isolates with a functional CYP51A.

involves variation in PC2 as well, so this shift shows only partial cross-resistance, with prothioconazole affected differently from the other compounds.

Isolates from each sensitivity group were selected for molecular analysis. Sensitivity profiles for these isolates are shown in figure 3. *CYP51A* and *CYP51B* were sequenced for these isolates (GenBank accession numbers KF753641–KF753674). No single nucleotide polymorphisms (SNPs) correlated with azole sensitivity, but functional *CYP51A* sequences were obtained from the intermediate and less-sensitive isolates, whereas a pseudogene, with a frameshift indel and premature stop codons, was sequenced from the sensitive isolates.

A polymerase chain reaction-restriction fragment length polymorphism (PCR-RFLP) assay was developed to distinguish CYP51A from CYP51A-p. This revealed that sensitive isolates possess CYP51A-p but not CYP51A, whereas intermediate and less sensitive isolates possess both CYP51A and CYP51A-p (supplementary fig. S2, Supplementary Material online). This difference in CYP51A copy number was confirmed by Southern blot (supplementary fig. S2, Supplementary Material online). CYP51A-p was sequenced from additional isolates (GenBank accession numbers KF753666–KF753674) and all contained premature stop codons.

Rhynchosporium commune CYP51A Expression

To test whether *R. commune* CYP51A still functions as a sterol 14α -demethylase, *R. commune* CYP51A and CYP51B were expressed in a *Saccharomyces cerevisiae* strain in which the native CYP51 is under the control of a tetracycline-repressible

promoter. *Rhynchosporium commune* CYP51A and CYP51B were each able to complement *S. cerevisiae* CYP51 (fig. 4). Yeast transformants were then tested for triazole sensitivity. There was little difference in fungicide sensitivity, to triazoles or to the non-DMI cycloheximide, between the CYP51A and CYP51B transformants (fig. 4b).

Expression analysis of *R. commune* CYP51A and CYP51B was carried out. No differences in constitutive expression of either paralog were correlated with azole sensitivity. However, following the addition of tebuconazole, CYP51B was only upregulated around 5- to 10-fold if at all, whereas CYP51A, where present, was upregulated around 100-fold. Therefore,



FIG. 3. Azole $\log_{10}[EC_{50}]$ values of *Rhynchosporium commune* isolates selected for target-site sequence analysis, and CYP51 amino acid substitutions. Black: propiconazole, gray: tebuconazole, dark gray: epoxiconazole, light gray: prothioconazole. Error bars indicate standard error between biological replicates. 1: RS 219; 2: K1124; 3: FI12-63; 4: R 9528.4; 5: R 9522.3; 6: QUB 30-10; 7: GKII 18-3-2; 8: GKII 18-2-3; 9: SAC 0003 1.4.8; 10: 788; 11: SAC 09/943/14; 12: QUB 12-3; 13: OSA 28-2-2; 14: RS 783. WT, wild type, identical to reference sequences from isolate 1130 (GenBank accession numbers KF753639 and KF753640); *no functional CYP51A sequenced.

MBE

Hoosfield Spring Barley Archive

Amplification of a 52-bp fragment of a reference gene (*R. commune* β -tubulin) was successful from samples from 33 different years, from 1892 to 2012. The CYP51A nested PCR amplified products from *R. secalis s.l.* and not from *R. orthosporum*. All *R. secalis s.l.* DNA present in these samples was confirmed as *R. commune* with 5% detection limit, except 2012, which was excluded from further testing.

A pyrosequencing assay was developed to distinguish R. commune CYP51A from CYP51A-p (fig. 6). For most of the 20th century, the majority of the *R*. *commune* population at the Hoosfield experimental site lacked CYP51A, but in 1985, levels of CYP51A rapidly increased, and subsequently the majority of the R. commune population possesses CYP51A. Logittransformed percentage data were analyzed with a Tukey multiple-comparison test. Levels of functional CYP51A in each year from 1985 onwards are significantly higher than in each year from 1892 to 1983. No year up to 1983 has significantly different levels of functional CYP51A from control isolates K1124 and FI12-63, which have only CYP51A-p, whereas no year from 1985 onwards has significantly different levels of CYP51A from isolates 788 and R9528.4, which have both CYP51A and CYP51A-p (supplementary table S4, Supplementary Material online). A further replicate DNA extraction from the homogenized plant material gave similar results (supplementary table S4, Supplementary Material online).

CYP51A in Rhynchosporium Species

The genomes of three *R. commune* isolates, and one isolate each of *R. agropyri*, *R. secalis s.s.*, and *R. orthosporum*, sister species to the *R. secalis s.l.* complex, have been sequenced. The genomes were searched for CYP51 paralogs (GenBank accession numbers KF753675–KF753687). All contain a single CYP51B. Two *R. commune* isolates contain CYP51A-*p* only, and one isolate contains CYP51A and CYP51A-*p*.



Fig. 4. (a) Growth of Saccharomyces cerevisiae strain YUG37::erg11 transformed with pYES3/CT (vector only) or pYES3-CT with Mycosphaerella graminicola CYP51 (Mg51), Rhynchosporium commune CYP51A (Rc51A, two transformants) or R. commune CYP51B (Rc51B, two transformants) on SD + GAL + RAF agar, with and without 3 μ g ml⁻¹ doxycycline (DOX), with six inoculum concentrations of 1.25 × 10⁶, 2.5 × 10⁵, 5 × 10⁴, 1 × 10⁴, 2 × 10³, and 4 × 10² cells μ l⁻¹. DOX represses expression of the native S. cerevisiae CYP51 in strain YUG37::erg11. (b) Triazole and cycloheximide log₁₀[EC₅₀] values of yeast transformants S. cerevisiae strain YUG37::erg11 transformed with R. commune CYP51A (black) or R. commune CYP51B (gray). A: propiconazole, B: tebuconazole, C: epoxiconazole, D: prothioconazole, E: cycloheximide. Error bars indicate standard error of biological replicates.



Fig. 5. Expression of (*a*) CYP51A and (b) CYP51B in *Rhynchosporium commune*, relative to isolate R 9528.4 constitutive expression, with β -tubulin as endogenous control. Light gray: constitutive expression, without fungicide; dark gray: induced expression, following the addition of $1 \mu \text{g ml}^{-1}$ tebuconazole. Error bars indicate standard error of biological replicates. *CYP51A not present in these isolates: A low level of amplicon results from nonspecific amplification of CYP51A-*p*.



FIG. 6. Percentage of functional CYP51A in *Rhynchosporium commune* isolates, and populations sampled from the Hoosfield experiment at Rothamsted Research, UK. Isolate 1: K1124; 2: FI12-63; 3: 788; 4: R 9528.4. Isolates K1124 and FI12-63 contain CYP51A-*p* only; isolates 788 and R 9528.4 contain both CYP51A-*p* and functional CYP51A. Error bars indicate standard error of five technical replicates.

Rhynchosporium agropyri and *R. secalis s.s.* contain a CYP51A pseudogene, but with different stop codons from *R. commune* CYP51A-p, whereas *R. orthosporum* contained only a functional CYP51A. Therefore, a functional CYP51A was present in the most recent common ancestor of *R. orthosporum* and *R. commune*, followed by duplication of CYP51A and pseudogenization of one copy in the *R. commune* lineage, and loss of the functional copy from some isolates (fig. 7).

Discussion

A 10-fold shift in azole sensitivity in *R. commune* is correlated with the presence of the CYP51A paralog. All isolates possess CYP51B and a CYP51A-p pseudogene, but the functional CYP51A is absent from azole sensitive isolates and present in less-sensitive isolates. Further selection for reduced azole sensitivity among isolates with CYP51A has resulted in a further 10-fold difference in sensitivity to some azoles, which is not correlated with mutations or overexpression of either



Fig. 7. Maximum-likelihood phylogeny of coding nucleotide sequences of CYP51 paralogs sequenced from *Rhynchosporium* species, with *Magnaporthe oryzae* CYP51A, *Botrytis cinerea* CYP51B, and *Fusarium graminearum* CYP51C. Rooted with *Candida albicans* CYP51 outgroup. Node labels indicate bootstrap support.

CYP51 paralog, implying that an additional, nontarget-site mechanism is responsible for this second shift.

Analysis of *R. commune* populations in barley samples from Hertfordshire, UK, shows an increase in *CYP51A* frequency in the population from 1985, with *CYP51A* present in the majority of the population thereafter. Similarly, the oldest studied isolates lack *CYP51A*, whereas most modern UK isolates have *CYP51A*. This correlates with a shift in triazole sensitivity reported in UK field populations between baseline monitoring from 1975 to 1981 and further surveys from 1985 onwards (Kendall et al. 1993). This demonstrates the value of the Rothamsted Classical Experiment archives to analyze past evolutionary changes involving mutations which have only now been identified.

The mechanism by which CYP51A reduces azole sensitivity appears to be inducible target-site overexpression. Rhynchosporium commune CYP51A is able to complement S. cerevisiae CYP51, so it is a functional sterol 14α demethylase, and in R. commune, CYP51A is upregulated following exposure to an azole, so CYP51A may be able to maintain CYP51 function when CYP51B is inhibited by azoles. Previously reported cases of reduced azole sensitivity due to target-site overexpression have all involved constitutive overexpression of an existing paralog due to promoter inserts or gene duplication. Acquired reductions in azole sensitivity due to CYP51 duplication have been reported due to chromosome duplication in C. glabrata (Marichal et al. 1997); isochromosome formation, with duplication of chromosome arm 5L containing CYP51, two efflux pump encoding genes and one ABC transporter transcription factor, in C. albicans (Selmecki et al. 2006); and disomy of chromosome 1, containing CYP51 and ABC transporter encoding gene AFR1, in Cryptococcus neoformans (Sionov et al. 2010). However, in all of these cases, this involved duplication of an existing gene, producing a second identical copy rather than a functionally divergent paralog. CYP51A demonstrates functional divergence from CYP51B through different patterns of transcriptional regulation, including greater upregulation in response to azoles. Neofunctionalization of fungal genes often involves changes in transcriptional regulation rather than changes in substrate (Skamnioti et al. 2008), and often altered expression evolves primarily in one copy whereas the other retains the original pattern (Gu et al. 2005). Intrinsic sequence differences between CYP51A and CYP51B contribute to reduced intrinsic azole sensitivity in A. fumigatus (Martel et al. 2010) and F. graminearum (Fan et al. 2013), but R. commune CYP51A conferred similar azole sensitivity to CYP51B when expressed in yeast, suggesting that upregulation of CYP51A, rather than reduced azole binding due to intrinsic sequence differences, is the main mechanism of reduced azole sensitivity in R. commune isolates possessing CYP51A.

Although CYP51 paralogs have been extensively studied in a few species, such as A. fumigatus where CYP51A mutations confer azole resistance, in others species, knowledge of CYP51 paralog presence is based only on a single isolate selected for genome sequencing. Therefore, genome resequencing of multiple isolates may reveal paralog presence polymorphisms in other species, and possible multiple CYP51 paralogs should be considered in future when investigating azole sensitivity shifts. Further investigation is also needed concerning the reasons for the emergence of CYP51A, and its retention in some fungal lineages, in the absence of selection by azoles. CYP51A expression is induced in CYP51B deletion mutants (Fan et al. 2013), as well as following CYP51 inhibition by azoles, suggesting that CYP51A is upregulated in response to disruption of ergosterol biosynthesis. Therefore, presence of this inducible CYP51, in addition to the constitutively expressed CYP51B found in all filamentous ascomycetes, would allow a fungus to respond to fluctuating ergosterol requirements, but it is not yet clear which aspects of fungal biology and lifestyle make this necessary in some species and not others.

Phylogenetic analysis places R. commune CYP51A in a monophyletic clade with other filamentous ascomycete CYP51As, which diverged from CYP51B around the origin of the filamentous ascomycetes, 320-520 Ma (Lücking et al. 2009). This indicates a single origin of CYP51A followed by multiple losses, rather than an independent recent origin in R. commune. However, the origin of CYP51A-p, and partial loss of CYP51A, in the R. commune lineage took place far more recently. Rhynchosporium orthosporum possesses a functional CYP51A and not a pseudogene, whereas the sequenced isolates of R. agropyri and R. secalis s.s. possess only a pseudogene. The R. orthosporum CYP51A forms a sister group to R. commune CYP51A and CYP51A-p. Therefore, CYP51A duplication took place in the R. commune lineage after it diverged from R. orthosporum, 14.5–35 ka BP (Zaffarano et al. 2008), but before the divergence of R. agropyri, R. secalis s.s., and R. commune from their most recent common ancestor, 1.3–3.6 ka BP (Zaffarano et al. 2008). This was followed in the R. commune lineage by pseudogenization of one copy, and loss of the functional copy in some of the population. Further study of R. agropyri and R. secalis s.s. would be needed to ascertain whether CYP51A pseudogenization occurred independently in these lineages or once in the R. secalis s.l. common ancestor, and whether any isolates of these species possess a functional CYP51A. The gene phylogeny confirms that CYP51A was retained within the R. commune lineage, and not regained by horizontal gene transfer, in contrast to the acquisition of antibiotic resistance genes in clinical bacteria by horizontal gene transfer from soil bacteria previously exposed to naturally occurring microbial antibiotics (Forsberg et al. 2012).

Having declined in the R. commune population, CYP51A later re-emerged under selection by azoles in the late 20th century. Most research into the origin and fate of duplicate genes considers possible outcomes in terms of loss or fixation in a species, and it has been demonstrated that most intraspecific gene copy number variations result from very recent duplication events, as older duplicates have already been lost or fixed, and most established, diverged duplicates are maintained by purifying selection (Schrider and Hahn 2010). Comparative genomics has revealed cases of fixed paralog losses following a change in selection in some taxa, such as repeated gene losses from the olfactory receptor gene family in microsmic mammals (Rouquier et al. 2000), or extensive gene losses in obligate parasites (Spanu et al. 2010). However, the Rothamsted archive samples, covering a time frame including a known change in selection, provide the temporal resolution to detect changes in paralog frequency over time, enabling us to detect the previous near-loss of a paralog followed by its re-emergence due to a further change in selection.

Furthermore, as phylogenetic analyses have shown that the functional CYP51A in *R. commune* originated long before azole use, its re-emergence provides an example of an adaptation to a new environment that has been selected from standing genetic variation rather than arising from a de novo mutation (Barrett and Schluter 2008), whereas previous studies of other fungicide resistance alleles have pointed to de novo mutational origins (Torriani et al. 2009; Camps et al.

2012). Although ancient origins have previously been demonstrated for resistance genes to antibiotics based on natural microbial products (D'Costa et al. 2011), the triazoles are synthetic in origin with no know natural analogs and therefore their use imposes a previously unencountered selective pressure. Standing variation may allow resistance to evolve more quickly, or may result in increased probability of fixation where the fitness benefit is smaller and a de novo mutation may have been lost through genetic drift (Hermisson and Pennings 2005). Furthermore, as this standing variation comprises a trait that has been lost from multiple ascomycete lineages, the availability of CYP51A re-emergence as an adaptive pathway is historically contingent upon CYP51A still having been present at low levels in the population at the point when selection pressures altered due to anthropogenic fungicide use.

Materials and Methods

Bioinformatic and Phylogenetic Analysis

CYP51 sequences were obtained from fungal genomes on the Broad Institute server (http://www.broadinstitute.org/sci ence/data, last accessed April 16, 2014), along with Mycobacterium tuberculosis, Trypanosoma cruzi, Homo sapiens, Arabidopsis thaliana CYP51G1, and Mus musculus CYP51, with Mus musculus CYP7A1 as an outgroup (Aoyama et al. 1996). A text search was carried out for annotated CYP51 genes, followed by BLAST searches against each genome with S. cerevisiae CYP51, and with CYP51A and CYP51C from the nearest available relative for species where these paralogs were not found. Start sites, introns, and stop sites were checked manually, and introns removed as they were too variable to align unambiguously. Predicted amino acid sequences were aligned using M-Coffee (Notredame et al. 2000), and the corresponding coding nucleotide alignment generated using PAL2NAL (Suyama et al. 2006). Model selection was carried out in jModelTest 0.1.1 (Posada 2008), selecting by AICc. Maximum-likelihood phylogenetic analysis was carried out in PhyML (Guindon and Gascuel 2003), implemented through the TOPALi v2.5 platform (Milne et al. 2009), using a TVM + I + G model, with 100 bootstrap runs.

Fungicide Sensitivity Testing

Isolates used are listed in supplementary table S1, Supplementary Material online. Isolates had been stored as spore suspensions on silica gel at -80 °C. Isolates were grown on Czapek dox agar with 0.5% mycological peptone, with an inoculum density of 1.25×10^5 spores per 90 mm Petri dish, at 18 °C for 10 days. Sensitivity testing was carried out as described by Pijls et al. (1994), using Sabouraud medium, *R. commune* spores at 1.25×10^5 spores ml⁻¹, and the following technical-grade fungicides: epoxiconazole, 50 µg ml⁻¹ with 2.5-fold dilutions; prothioconazole, propiconazole, tebuconazole, 100 µg ml⁻¹ with 3-fold dilutions, repeated with 300 µg ml⁻¹ for less sensitive isolates. Plates were incubated at 18 °C for 7 days, then fungal growth was measured by optical absorbance at 620 nm using an Optima Fluostar plate reader (BMG Labtech, Germany), in well-scanning mode with a 4×4 matrix of scanning points within a 3-mm diameter. The accompanying software was used to fit a dose–response curve (4-parameter fit) and calculate EC₅₀ values. Cross-resistance was assessed by PCA using the correlation matrix of log₁₀ transformed EC₅₀s for the four fungicides, in GenStat 14th Edition (VSN International, Hertfordshire, UK).

CYP51 Sequencing

Initial fragments were amplified using degenerate primers CYP51 3F and CYP51A CIGEK rv for CYP51A, and DMIRES F1 and 14DM R2 for CYP51B (supplementary table S3, Supplementary Material online). The remainder of each gene (GenBank accession numbers KF753639-KF753640) was found by genome-walking using the GenomeWalker Universal Kit (Clontech, CA). Libraries were prepared from isolate 1130 (Rohel et al. 1998) genomic DNA, with restriction enzymes Dral, Pvull, EcoRV, and Stull. Three reactions in the 5'-direction and one in the 3'-direction were needed to obtain the complete coding sequence of CYP51A, and two reactions in the 5'-direction and one in the 3'-direction for CYP51B. Genome-specific primers (GSP1 and GSP2) used in successive reactions are listed in supplementary table S3, Supplementary Material online. PCR products were purified using the Wizard SV Gel and PCR Clean-Up System, ligated into pGEM-T Easy plasmid vector using T4 DNA ligase and cloned in JM109 competent cells (all Promega). Plasmid DNA was extracted and purified using the GeneElute (Sigma) or QIASpin (Qiagen) mini-prep kits, then sequenced by Eurofins MWG (Germany) using sequencing primers M13uni and M13rev (Messing 1983) and the sequences assembled in Vector NTI 10 (Invitrogen, CA).

CYP51B, CYP51A, and CYP51A-p were sequenced from isolates with a range of triazole sensitivities (fig. 2): K1124, FI12-63, RS-219, QUB 30-10, R 9528.4, 9522.3, 788, SAC 1-4-8 (0003), GKII 18-2-3, GKII 18-3-2, SAC 09/943/14, QUB 12-3, OSA 28-2-2, and RS 783. Isolates were grown in Sabouraud liquid medium for 10 days. Fungal material was harvested by filtration and freeze-dried. DNA was extracted as in Motteram et al. (2009).

CYP51A and CYP51B were amplified using PCR, carried out with Phusion High-Fidelity DNA Polymerase (Finnzymes Oy, Espoo, Finland), with CYP51B and CYP51A primers Forwards 1 and Reverse 1 (supplementary table S3, Supplementary Material online). Products were purified and sequenced by Eurofins MWG (Germany), with the nested and internal primers listed in supplementary table S3, Supplementary Material online. CYP51A-p was amplified using Easy-A High-Fidelity PCR Cloning Enzyme (Stratagene) with primers listed in supplementary table S3, Supplementary Material online. Products were cloned and sequenced as described for genome walking, with additional sequencing reactions using internal primers (supplementary table S3, Supplementary Material online).

PCR-RFLP Assay

Two PCR reactions were carried out per isolate, one to amplify CYP51A and one to amplify CYP51A-p, using Phusion

High-Fidelity DNA Polymerase (Finnzymes Oy) with primers listed in supplementary table S3, Supplementary Material online. Restriction digests were carried out with enzymes *Pstl* and *HindIII* (Promega), with 3-h incubation time.

CYP51A copy number was confirmed by Southern blot as described by Motteram et al. (2009). Ten micrograms of genomic DNA from isolates 788, K1124, and FI12-63 was digested with high concentration *Pstl* and *EcoRV* (Promega). The probe was amplified from pGEM-T Easy plasmids containing CYP51A from isolate 788 cDNA, using primers Southern F and Reverse 10D (supplementary table S3, Supplementary Material online) to amplify a 734-bp fragment of exon 3.

Yeast Complementation

Rhynchosporium commune CYP51A and CYP51B were amplified from a 1/10 dilution of isolate 788 cDNA using Easy-A High-Fidelity PCR Cloning Enzyme (Stratagene), with primers CYP51A pYES F2 and pYES R and CYP51B pYES F and pYES R (supplementary table S3, Supplementary Material online). PCR product was purified using the Wizard SV Gel and PCR Clean-Up System (Promega), and a sample of the CYP51A product digested with Pstl and HindIII to confirm it was CYP51A and not CYP51A-p. Kpnl and a Sacl restriction sites were incorporated by the forward and reverse primers, respectively. An internal Kpnl site was removed from CYP51B using the QuikChange II Site-Directed Mutagenesis Kit (Agilent Technologies, CA, USA) with primer pYES mut (supplementary table S3, Supplementary Material online). Cloning in yeast expression vector pYES2/CT (Invitrogen), transformation into S. cerevisiae YUG37:erg11 strain, and complementation analysis were carried out as described by Cools et al. (2010), with pYES-Mg51wt as a positive control. Fungicide sensitivity testing was carried out as in Cools et al. (2010), but with maximum fungicide concentrations of $0.005 \,\mu \text{g ml}^{-1}$ for epoxiconazole and propiconazole, $0.05 \,\mu g \,m l^{-1}$ for prothioconazole and tebuconazole and $5 \,\mu\text{g}\,\text{m}\text{l}^{-1}$ for cycloheximide, all with 2-fold serial dilutions.

CYP51 Expression

Rhynchosporium commune isolates QUB 30-10, 788, K1124, QUB 12-3, FI12-63, R 9528.4, GKII 18-2-3, and RS 783 783 were grown liquid culture, in 100-ml Sabouraud liquid medium at 2.5×10^4 spores ml⁻¹ at 20 °C, 150 rpm, with two biological replicates of treatment and control culture for each isolate. After 7 days, tebuconazole solution in acetone to a final concentration of $1 \,\mu g \,m l^{-1}$ was added. An equal volume of acetone without tebuconazole was added to the control cultures. After 10 days, cultures were harvested by vacuum-filtration, then freeze-dried. RNA extraction was carried out with TRIzol Reagent (Invitrogen), according to manufacturer's instructions, followed by overnight precipitation with 4 M lithium chloride and treatment with the TURBO DNA-free Kit (Applied Biosystems, CA). cDNA was synthesized with the Superscript III first-strand synthesis system (Invitrogen). End-point PCR reactions were carried out using Red Hot Taq (ABgene, Epsom, UK) with intron-spanning primer pair

CYP51B Expression F and CYP51B Expression R (supplementary table S3, Supplementary Material online) to check for genomic DNA contamination.

Quantitative PCR was carried out using SYBR Green JumpStart Taq ReadyMix (Sigma), with CYP51B, CYP51A, and β -tubulin primers Expression F and Expression R (supplementary table S3, Supplementary Material online). Relative quantification was calculated by the $2^{-\Delta\Delta C_T}$ method (Livak and Schmittgen 2001), with β -tubulin as endogenous control and isolate R9528.4 without fungicide as calibrator sample.

Hoosfield Long-Term Experiment and Pyrosequencing Assay

Spring barley has been grown on the Hoosfield plot at Rothamsted Research, in Harpenden, UK, every year since 1852, apart from 1912, 1933, 1943, and 1967 when it was left fallow to control weeds. Dried leaf, grain, and soil samples have been retained each year. Straw samples used were from Plot 4A, receiving 48 kg N ha⁻¹ in the form of ammonium sulfate. From 1968, this plot was divided into four subplots receiving different nitrogen doses in the form of calcium ammonium nitrate, and samples used were from the subplot receiving 48 kg N ha⁻¹.

One gram of material was taken from the archive sample, ground using a sterilized pestle and mortar, then extraction buffer was added and DNA extractions carried out as described by Motteram et al. (2009), with two replicate extractions from each homogenized sample. Rhynchosporium secalis s.l. DNA presence and quality were tested by endpoint PCR amplification of a 520-bp fragment of reference gene β -tubulin using primers Rs F3 and Rs R3 (supplementary table S3, Supplementary Material online), and up to 3 years per decade were selected for analysis. Rhynchosporium secalis s.l. species present were identified based on a C/T SNP at nucleotide 391 of β -tubulin (Zaffarano et al. 2011), with a nested PCR using primers RhyUF1 and RhyUR1 then PyroRhF1Bio and PyroRHR1, followed by a pyrosequencing assay using the PyroMark system (Qiagen) and the PSQ96 instrument (Biotage, Uppsala, Sweden), programmed to analyze the sequence TC/TGCAGATCCAAGGTAGAACTTACA CT, using the nucleotide dispensation order ATCTGCAGA, with primer PyroRhS1 (supplementary table S3, Supplementary Material online).

To distinguish CYP51A from CYP51A-p, a pyrosequencing assay was developed to detect a 4-bp indel at nucleotide 418. A nested PCR was carried out with primers Pyro F3 and Pyro R2 then primers Del F1 and Del R1 (supplementary table S3, Supplementary Material online) and the products pyrosequenced using the PyroMark system (Qiagen) and the PSQ96 instrument (Biotage), to analyze the sequence CA[TTTC]CTTCCAGCCCAAGATAAACAGTCAT, using the nucleotide dispensation order GCATCGTCA, with primer Del S1 (supplementary table S3, Supplementary Material online).

Rhynchosporium Spp. Genome Analysis

The genomes of R. *commune* isolates UK7, AU2 (Lehnackers and Knogge 1990), and 13-13 (A. Avrova, The James Hutton

Institute); R. secalis s.s. isolate 02CH4-6a.1 (Zaffarano et al. 2008); R. agropyri isolate 04CH Rac A.6.1; and R. orthosporum isolate 02CH4 Bär A.1.1.3 (B. McDonald, ETH Zurich) have been sequenced and are currently prepared for publication. A BLAST search was carried out against R. commune CYP51A, CYP51A-p, and CYP51B. Introns were predicted and coding sequences translated. Each CYP51 was cloned and resequenced by Sanger sequencing, as described for CYP51A-p sequencing with primers listed in supplementary table S3, Supplementary Material online, to confirm the genome assembly. Predicted amino acid sequences, along with B. cinerea CYP51B and M. oryzae CYP51A (the closest relatives of R. commune in supplementary fig. S1, Supplementary Material online), F. graminearum CYP51C, and C. albicans CYP51 as an outgroup were aligned with M-Coffee, and aligned coding nucleotides were with PAL2NAL. Substitution models were predicted with iModeltest, and phylogenetic reconstruction carried out in PhyML with the GTR + G model and 100 bootstrap runs.

Supplementary Material

Supplementary tables S1–S4 and figures S1 and S2 are available at *Molecular Biology and Evolution* online (http://www.mbe.oxfordjournals.org/).

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References

- Aoyama Y, Noshiro M, Gotoh O, Imaoka S, Funae Y, Kurosawa N, Horiuchi T, Yoshida Y. 1996. Sterol 14-demethylase P450 (P450 14DM*) is one of the most ancient and conserved P450 species. *J Biochem.* 119:926–933.
- Barrett RDH, Schluter D. 2008. Adaptation from standing genetic variation. *Trends Ecol Evol.* 23:38–44.
- Bearchell SJ, Fraaije BA, Shaw MW, Fitt BDL. 2005. Wheat archive links long-term fungal pathogen population dynamics to air pollution. *Proc Natl Acad Sci U S A*. 102:5438–5442.
- Caldwell R. 1937. *Rhynchosporium* scald of barley, rye and other grasses. *J Agric Res.* 55:175–198.

- Camps SMT, Rijs AJMM, Klaassen CHW, Meis JF, O'Gorman CM, Dyer PS, Melchers WJG, Verweij PE. 2012. Molecular epidemiology of Aspergillus fumigatus isolates harboring the TR34/L98H azole resistance mechanism. J Clin Microbiol. 50:2674–2680.
- Conant GC, Wolfe KH. 2008. Turning a hobby into a job: how duplicated genes find new functions. *Nat Rev Genet.* 9:938–950.
- Cools HJ, Parker JE, Kelly DE, Lucas JA, Fraaije BA, Kelly SL. 2010. Heterologous expression of mutated eburicol 14 alpha-demethylase (CYP51) proteins of *Mycosphaerella graminicola* to assess effects on azole fungicide sensitivity and intrinsic protein function. *Appl Environ Microbiol.* 76:2866–2872.
- Da Silva Ferreira ME, Colombo AL, Paulsen I, Ren Q, Wortman J, Huang J, Goldman MHS, Goldman GH. 2005. The ergosterol biosynthesis pathway, transporter genes, and azole resistance in *Aspergillus furnigatus*. *Med Mycol.* 43:S313–S319.
- D'Costa VM, King CE, Kalan L, Morar M, Sung WWL, Schwarz C, Froese D, Zazula G, Calmels F, Debruyne R, et al. 2011. Antibiotic resistance is ancient. *Nature* 477:457–461.
- de Waard MA, van Nistelrooy JGM. 1980. An energy-dependent efflux mechanism for fenarimol in a wild-type strain and fenarimol-resistant mutants of Aspergillus nidulans. Pestic Biochem Phys. 13:255–266.
- Deng JX, Carbone I, Dean RA. 2007. The evolutionary history of cytochrome P450 genes in four filamentous Ascomycetes. *BMC Evol Biol.* 7:22.
- Denning DW, Hope WW. 2010. Therapy for fungal diseases: opportunities and priorities. *Trends Microbiol.* 18:195–204.
- Dittmar K, Liberles DA. 2010. Evolution after gene duplication. New Jersey: John Wiley & Sons.
- Fan J, Urban M, Parker JE, Brewer HC, Kelly SL, Hammond-Kosack KE, Fraaije BA, Liu X, Cools HJ. 2013. Characterization of the sterol 14alpha-demethylases of *Fusarium graminearum* identifies a novel genus-specific CYP51 function. *New Phytol.* 198:821–835.
- Forsberg KJ, Reyes A, Wang B, Selleck EM, Sommer MOA, Dantas G. 2012. The shared antibiotic resistome of soil bacteria and human pathogens. *Science* 337:1107–1111.
- Garthwaite DG, Barker I, Parrish G, Smith L, Chippindale C, Pietravalle S. 2011. Pesticide Usage Survey report 235: arable crops in the United Kingdom 2010. York (United Kingdom): Food and Environment Research Agency.
- Gruber J, Colligan P, Wolford J. 2002. Estimation of single nucleotide polymorphism allele frequency in DNA pools by using Pyrosequencing. *Hum Genet.* 110:395–401.
- Gu X, Zhang Z, Huang W. 2005. Rapid evolution of expression and regulatory divergences after yeast gene duplication. *Proc Natl Acad Sci U S A*. 102:707–712.
- Guindon S, Gascuel O. 2003. A simple, fast, and accurate algorithm to estimate large phylogenies by maximum likelihood. *Syst Biol.* 52: 696–704.
- Hamamoto H, Hasegawa K, Nakaune R, Lee YJ, Makizumi Y, Akutsu K, Hibi T. 2000. Tandem repeat of a transcriptional enhancer upstream of the sterol 14 alpha-demethylase gene (CYP51) in *Penicillium digitatum*. Appl Environ Microbiol. 66:3421–3426.
- Heckel DG. 2012. Insecticide resistance after silent spring. Science 337: 1612–1614.
- Hermisson J, Pennings PS. 2005. Soft sweeps: molecular population genetics of adaptation from standing genetic variation. *Genetics* 169: 2335–2352.
- Hughes AL. 1994. The evolution of functionally novel proteins after gene duplication. *Proc R Soc Lond B Biol Sci.* 256:119–124.
- Innan H, Kondrashov F. 2010. The evolution of gene duplications: classifying and distinguishing between models. *Nat Rev Genet.* 11: 97–108.
- Kelly S, Arnoldi A, Kelly DE. 1993. Molecular genetic analysis of azole antifungal mode of action. *Biochem Soc Trans*. 21:1034–1051.
- Kelly SL, Lamb DC, Corran AJ, Baldwin BC, Kelly DE. 1995. Mode of action and resistance to azole antifungals associated with the formation of 14[alpha]-methylergosta-8,24(28)-dien-3[beta],6[alpha]diol. Biochem Biophys Res Commun. 207:910–915.

- Kendall SJ, Hollomon DW, Cooke LR, Jones DR. 1993. Changes in sensitivity to DMI fungicides in *Rhynchosporium secalis*. Crop Prot. 12: 357–362.
- Lehnackers H, Knogge W. 1990. Cytological studies on the infection of barley cultivars with known resistance genotypes by *Rhynchosporium commune. Can J Bot.* 68:1953–1961.
- Lepesheva GI, Waterman MR. 2007. Sterol 14 alpha-demethylase cytochrome P450 (CYP51), a P450 in all biological kingdoms. *Biochim Biophys Acta*. 1770:467–477.
- Liu W, Sun Y, Chen W, Liu W, Wan Z, Bu D, Li R. 2012. T778G mutation in cyp51C gene confers voriconazole-resistance in Aspergillus flavus causing aspergillosis. Antimicrob Agents Chemother. 56:2598–2603.
- Livak KJ, Schmittgen TD. 2001. Analysis of relative gene expression data using real-time quantitative PCR and the 2(-Delta Delta C(T)) method. *Methods* 25:402–408.
- Lücking R, Huhndorf S, Pfister DH, Plata ER, Lumbsch HT. 2009. Fungi evolved right on track. *Mycologia* 101:810–822.
- Lynch M, Conery JS. 2000. The evolutionary fate and consequences of duplicate genes. *Science* 290:1151–1155.
- MacLean RC, Hall AR, Perron GG, Buckling A. 2010. The population genetics of antibiotic resistance: integrating molecular mechanisms and treatment contexts. *Nat Rev Genet.* 11:405–414.
- Marichal P, VandenBossche H, Odds FC, Nobels G, Warnock DW, Timmerman V, VanBroeckhoven C, Fay S, MoseLarsen P. 1997. Molecular-biological characterization of an azole-resistant *Candida* glabrata isolate. Antimicrob Agents Chemother. 41:2229–2237.
- Martel CM, Parker JE, Warrilow AGS, Rolley NJ, Kelly SL, Kelly DE. 2010. Complementation of *Saccharomyces cerevisiae* ERG11/CYP51 (sterol 14α-demethylase) doxyxycline regulated mutant and screening the azole-sensitivity of *Aspergillus fumigatus* isoenzymes CYP51A and B. *Antimicrob Agents Chemother*. 54:4920–4923.
- Mayfield AH, Clare BG. 1991. Effects of *Rhynchosporium secalis* at specific growth stages on barley grown in a controlled environment. *Plant Pathol.* 40:561–567.
- McKenna M. 2013. Antibiotic resistance: the last resort. *Nature* 499: 394–396.
- Mellado E, Diaz-Guerra TM, Cuenca-Estrella M, Rodriguez-Tudela JL. 2001. Identification of two different 14-alpha sterol demethylase-related genes (*cyp51A* and *cyp51B*) in *Aspergillus fumigatus* and other aspergillus species. J Clin Microbiol. 39:2431–2438.
- Mellado E, Garcia-Effron G, Alcazar-Fuoli L, Melchers WJG, Verweij PE, Cuenca-Estrella A, Rodriguez-Tudela JL. 2007. A new Aspergillus fumigatus resistance mechanism conferring in vitro cross-resistance to azole antifungals involves a combination of *cyp51A* alterations. *Antimicrob Agents Chemother*. 51:1897–1904.
- Mellado E, Garcia-Effron G, Buitrago MJ, Alcazar-Fuoli L, Cuenca-Estrella A, Rodriguez-Tudela JL. 2005. Targeted gene disruption of the 14alpha sterol demethylase (*cyp51A*) in *Aspergillus fumigatus* and its role in azole drug susceptibility. *Antimicrob Agents Chemother*. 49: 2536–2538.
- Messing J. 1983. New M13 vectors for cloning. In: Ray Wu, Grossman L, Moldave K, editors. Methods in enzymology. New York: Academic Press. p. 20–78.
- Milne I, Lindner D, Bayer M, Husmeier D, McGuire GI, Marshall DF, Wright F. 2009. TOPALi v2: a rich graphical interface for evolutionary analyses of multiple alignments on HPC clusters and multi-core desktops. *Bioinformatics* 25:126–127.
- Morio F, Loge C, Besse B, Hennequin C, Le Pape P. 2010. Screening for amino acid substitutions in the *Candida albicans* Erg11 protein of azole-susceptible and azole-resistant clinical isolates: new substitutions and a review of the literature. *Diagn Microbiol Infect Dis.* 66: 373–384.
- Morton V, Staub T. 2008. A short history of fungicides. *APSnet* Features [Internet]. St. Paul (MN): American Phytopathological Society; [cited 2014 Apr 24]. Available from: http://www.apsnet.org/.
- Motteram J, Küfner I, Deller S, Brunner F, Hammond-Kosack KE, Nürnberger T, Rudd JJ. 2009. Molecular characterization and functional analysis of MgNLP, the sole NPP1 domain-containing protein,

from the fungal wheat leaf pathogen *Mycosphaerella graminicola*. *Mol Plant Microbe Interact.* 22:790–799.

MBE

- Näsvall J, Sun L, Roth JR, Andersson DI. 2012. Real-time evolution of new genes by innovation, amplification, and divergence. *Science* 338: 384–387.
- Nelson DR. 2009. The cytochrome P450 homepage. Hum Genomics. 4: 59-65.
- Notredame C, Higgins DG, Heringa J. 2000. T-Coffee: a novel method for fast and accurate multiple sequence alignment. *J Mol Biol.* 302: 205–217.
- Oerke EC, Dehne HW. 2004. Safeguarding production—losses in major crops and the role of crop protection. *Crop Prot.* 23:275–285.
- Ohno S. 1970. Evolution by gene duplication. Berlin (Germany): Springer-Verlag.
- Oxley S, Burnett FJ, Hunter EA, Fraaije B, Cooke LR, Mercer PC, Gilchrist A. 2008. Understanding fungicide mixtures to control *Rhynchosporium* in barley and minimise resistance shifts. London: HGCA.
- Pijls CFN, Shaw MW, Parker A. 1994. A rapid test to evaluate in vitro sensitivity of *Septoria tritici* to flutriafol, using a microtitre plate reader. *Plant Pathol.* 43:726–732.
- Posada D. 2008. jModelTest: phylogenetic model averaging. *Mol Biol Evol*. 25:1253-1256.
- Rohel EA, Payne AC, Hall L, Barker H, Butters JA, Holloman DW. 1998. Isolation and characterisation of *alpha-tubulin* genes from *Septoria tritici* and *Rhynchosporium secalis*, and comparative analysis of fungal *alpha-tubulin* sequences. *Cell Motil Cytoskeleton*. 41:247–253.
- Rouquier S, Blancher A, Giorgi D. 2000. The olfactory receptor gene repertoire in primates and mouse: evidence for reduction of the functional fraction in primates. *Proc Natl Acad Sci U S A*. 97: 2870–2874.
- Schrider DR, Hahn MW. 2010. Gene copy-number polymorphism in nature. Proc R Soc Lond B Biol Sci. 277:3213–3221.
- Selmecki A, Forche A, Berman J. 2006. Aneuploidy and isochromosome formation in drug-resistant Candida albicans. Science 313:367–370.
- Sionov E, Lee H, Chang YC, Kwon-Chung KJ. 2010. *Cryptococcus neoformans* overcomes stress of azole drugs by formation of disomy in specific multiple chromosomes. *PLoS Pathog.* 6:13.
- Skamnioti P, Furlong RF, Gurr SJ. 2008. Evolutionary history of the ancient cutinase family in five filamentous Ascomycetes reveals differential gene duplications and losses and in *Magnaporthe grisea* shows evidence of sub- and neo-functionalization. *New Phytol.* 180: 711–721.
- Spanu PD, Abbott JC, Amselem J, Burgis TA, Soanes DM, Stüber K, Loren van Themaat EV, Brown JKM, Butcher SA, Gurr SJ, et al. 2010. Genome expansion and gene loss in powdery mildew fungi reveal tradeoffs in extreme parasitism. *Science* 330:1543–1546.
- Strange RN, Scott PR. 2005. Plant disease: a threat to global food security. Annu Rev Phytopathol. 43:83–116.
- Suyama M, Torrents D, Bork P. 2006. PAL2NAL: robust conversion of protein sequence alignments into the corresponding codon alignments. *Nucleic Acids Res.* 34:W609–W612.
- Torriani SFF, Brunner PC, McDonald BA, Sierotzki H. 2009. Qol resistance emerged independently at least 4 times in European populations of *Mycosphaerella graminicola*. *Pest Manag Sci*. 65:155–162.
- Yan X, Ma WB, Li Y, Wang H, Que Y-W, Ma ZH, Talbot NJ, Wang ZY. 2011. A sterol 14[alpha]-demethylase is required for conidiation, virulence and for mediating sensitivity to sterol demethylation inhibitors by the rice blast fungus *Magnaporthe oryzae*. Fungal Genet Biol. 48:144–153.
- Zaffarano PL, McDonald BA, Linde CC. 2008. Rapid speciation following recent host shifts in the plant pathogenic fungus *Rhynchosporium*. *Evolution* 62:1418–1436.
- Zaffarano PL, McDonald BA, Linde CC. 2011. Two new species of *Rhynchosporium. Mycologia* 103:195–202.
- Zhang JZ. 2003. Evolution by gene duplication: an update. *Trends Ecol Evol.* 18:292–298.