Genomics of Compensatory Adaptation in Experimental Populations of Aspergillus nidulans

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ABSTRACT Knowledge of the number and nature of genetic changes responsible for adaptation is essential for understanding and predicting evolutionary trajectories. Here, we study the genomic basis of compensatory adaptation to the fitness cost of fungicide resistance in experimentally evolved strains of the filamentous fungus Aspergillus nidulans. The original selection experiment tracked the fitness recovery of lines founded by an ancestral strain that was resistant to fludioxonil, but paid a fitness cost in the absence of the fungicide. We obtained whole-genome sequence data for the ancestral A. nidulans strain and eight experimentally evolved strains. We find that fludioxonil resistance in the ancestor was likely conferred by a mutation in histidine kinase nikA, part of the two-component signal transduction system of the high-osmolarity glycerol (HOG) stress response pathway. To compensate for the pleiotropic negative effects of the resistance mutation, the subsequent fitness gains observed in the evolved lines were likely caused by secondary modification of HOG pathway activity. Candidate genes for the compensatory fitness increases were significantly overrepresented by stress response functions, and some were specifically associated with the HOG pathway itself. Parallel evolution at the gene level was rare among evolved lines. There was a positive relationship between the predicted number of adaptive steps, estimated from fitness data, and the number of genomic mutations, determined by whole-genome sequencing. However, the number of genomic mutations was, on average, 8.45 times greater than the number of adaptive steps inferred from fitness data. This research expands our understanding of the genetic basis of adaptation in multicellular eukaryotes and lays out a framework for future work on the genomics of compensatory adaptation in A. nidulans.

KEYWORDS fungicide resistance adaptive resistance trade-offs fludioxonil

A long-standing question in evolutionary biology concerns the number of genetic changes responsible for adaptation. A common view was that adaptation was underlain by many genes, each having small individual effects on fitness (Fisher 1930; Orr 1998). This interpretation initially gained wide acceptance, partly because there was little evidence with which it could be confronted (Orr 2005). A powerful approach for the study of the genetic basis of adaptation is through microbial selection experiments, especially when coupled with cost-effective whole-genome sequencing (Elena and Lenski 2003; Long et al. 2015). Not only can adaptive mutations be tracked in real time as they arise, but the number and nature of genetic changes underlying adaptive evolution can be determined directly.

A compelling result to emerge from this field of work is that much of the fitness gain associated with an adaptive walk (sequential substitution of beneficial mutations by selection) is attributable to a small number of genetic changes (e.g., Wichman et al. 1999; Riehle et al. 2001; Conrad et al. 2009; Schoustra et al. 2009; Anderson et al. 2010; Kvitek and Sherlock 2011). A recent review, for example, showed that the number of mutations fixed in microbial selection experiments that have reached a fitness plateau is typically between one and four (Kassen 2014). This result is consistent with the distribution of fitness effects among beneficial mutations being highly variable, such that in large populations, initial increases in fitness are associated with mutations that have large beneficial effects. Nonetheless, this result likely represents an
underestimate because many analyses infer the number of fixation events from fitness trajectories or genetic crosses rather than whole-genome sequencing. Moreover, most microbial experiments are conducted using very large population sizes that violate one of the key assumptions of models of adaptation: that beneficial mutations are rare events that, when they occur, sweep nearly instantaneously to fixation. Consequently, the number of genetic changes underlying adaptive walks may be substantially higher than previous estimates have inferred.

To gain a deeper understanding of the genetic basis of adaptation, we sequenced the genomes of experimentally evolved lines of the filamentous fungus Aspergillus nidulans. These lines were generated during a recently reported selection experiment (Schoustra et al. 2009) in which 118 selection lines were founded by a common ancestor (strain WG615) that carried a mutation that conferred resistance to the fungicide fludioxonil. As is often the case, a pleiotropic effect of the resistance mutation was a strong fitness cost in the absence of the fungicide (46% reduced growth rate; Schoustra et al. 2006). To allow for compensatory evolution, the populations were propagated in a fungicide-free environment for 800 generations with two different effective fitness costs. The recovery of fitness was tracked during the course of the selection experiment, and all lineages reached a fitness plateau by the end, suggesting that the bulk of fitness recovery had been accomplished. From data on the fitness trajectories, a novel maximum likelihood framework and genetic crosses were used to infer the number of beneficial, compensatory mutations. The main finding was that adaptive walks tended to be rather short, with an overall mean of only 2.20 steps (Schoustra et al. 2009).

The exact molecular mode of action of phenylpyruvole fungicides such as fludioxonil has yet to be elucidated, but they are known to act through the disruption of the HOG response pathway (Furukawa et al. 2007; Hagiwara et al. 2007, 2009, 2013). This pathway involves a two-component system and a mitogen-activated protein kinase (MAPK) signaling cascade, in which a histidine kinase serves as a sensor for multiple stressors and transmits the signal downstream via a response regulator to activate the MAPK. Previous work with Aspergillus has shown this stress response pathway to be activated by exposure to fludioxonil (Furukawa et al. 2007; Hagiwara et al. 2009) and that knocking out components of this pathway results in high levels of fludioxonil resistance (Hagiwara et al. 2007, 2013). Genes involved in the HOG MAPK pathway are therefore prime candidates for fludioxonil resistance determinants.

Here, we sequence the 30.5 Mb genomes of the ancestral A. nidulans strain and eight experimentally evolved strains (Schoustra et al. 2009) to investigate the evolutionary genomics of compensatory adaptation to the fitness cost of fungicide resistance. First, since the genetic basis of the fungicide resistance is not known, we identify candidate mutations for resistance in the ancestral strain. We then identify mutations within the experimentally evolved strains and analyze the characteristics of the mutated genes to determine how they may be related to compensatory, adaptive evolution. Finally, we use these data to determine the relationship between the number of genomic mutations and the predicted number of adaptive steps estimated from fitness data, and we also test hypotheses about how effective population size impacts the number and nature of mutations fixed during adaptation.

**MATERIALS AND METHODS**

**Strains**
The ancestral A. nidulans strain is fludioxonil-resistant WG615 (flA1, wA3, pyrA4, and veA1). When growing in the absence of fungicide, resistance confers a cost of ~46% relative to the sensitive strain from which WG615 was derived (Schoustra et al. 2006). All evolved strains were derived from WG615 through 800 generations of experimental evolution and adaptation to a permissive growth environment [complete medium (CM) without fungicide; Schoustra et al. 2009]. The wild-type sensitive strain was not evolved because, in previous experiments (Schoustra et al. 2005, 2006), it showed little evidence of adaptation to the permissive growth environment, suggesting that the sensitive strain already resides close to a fitness optimum. Every 80 generations, experimental lines were put through a bottleneck of either ~500 or 50,000 spores, corresponding to small (S) and large (L) effective population size treatments, respectively. For whole-genome sequencing, we chose four strains from each of the small and large population size treatments. Representative strains were chosen to cover the full range of observed fitness values and predicted step numbers (Supplemental Material, Table S1). The eight evolved strains characterized here (Table S1) were all put through a single-conidium transfer to ensure that a single haploid genotype was being sequenced. From the results of Schoustra et al. (2009), the number of predicted adaptive steps per strain ranged from one to three.

**DNA extraction and sequencing**

Strains were grown on CM agar plates and genomic DNA was extracted using the QIAGEN DNeasy Blood & Tissue Kit. Sequence data were generated using the Illumina Hi-Seq platform with paired-end reads (BC Cancer Agency). Reads generated for the evolved and ancestral strains were 75 and 100 bp in length, respectively.

**Reference-based mapping**
The A. nidulans reference genome was from strain FGSC A4 and is publicly available from NCBI (PRJNA13961). The assembly is divided into scaffolds (NT_106999 to NT_1070015) corresponding roughly to the 16 arms of the eight A. nidulans chromosomes. A modified version of the bioinformatics pipeline described in Dettman et al. (2012) was used for sequence data analyses. In brief, reads were trimmed using Popoolation (ver. 1.1, Koller et al. (2011)) with a phred quality threshold of 20 and a minimum retention length of 75% of original read length. Trimmed reads were mapped to the FGSC A4 reference genome using Novoalign (ver. 2.07).

**Coverage**
Coverage along scaffolds was calculated in 1 kb intervals and regions of high or low coverage were identified by bins with $>1.8 \times$ or $<0.20 \times$ the average coverage of the entire genome of that sample. Regions were confirmed by manual inspection of aligned reads and by results from other insertion/deletion (indel) calling programs (Pindel, ver. 0.2.4, Ye et al. 2009; Bresq, ver. 0.16, Barrick et al. 2014). Areas of high or missing coverage were reported if they were >500 bp in length, and regions that were within 200 bp of each other were collapsed into one.

**Variant calling**
Single nucleotide polymorphisms (SNPs) and indels relative to the reference were called using Samtools (ver. 0.1.19, Li et al. 2009; minimum coverage = 3 \times), VarScan (ver. 2.3.5, Koboldt et al. 2012; minimum coverage = 3 \times, P > 0.95), and Pindel (ver. 0.2.4). Results from the different methods were compared for cross-verification of mutation calls. A variant shared by WG615 and all evolved lines indicates that that variant was present in the ancestral WG615 strain at the start of the experiment. Variants that arose during the selection experiment were those that were absent in WG615 but present in an evolved line. For
variant calling, we filtered out the regions of low quality assembly in the reference genome, and correspondingly, regions of low mapping quality in the query genomes. Therefore, we may have underestimated the true number of sequence differences between the strains. Sequence data and positional alignments were inspected manually to confirm the called mutations in evolved lines. Significance of Pearson’s correlations were determined using the Student’s t-distribution.

**Annotation and functional analyses**

Genomic variation was annotated using SnpEff (ver. 3.3, Cingolani et al. 2012, minimum coverage = 3 x, minimum base quality = 20) to determine the types of mutation and what genes were affected. Amino acids were classified as hydrophilic uncharged (S, N, T, and Q), aliphatic uncharged (A, G, V, L, and I), nonpolar uncharged (C, M, and P), acidic (D and E), basic (K, R, and H), and aromatic (F, Y, and W). Amino acid changes within and between these classes were considered conservative and radical, respectively. Mutations that caused codon indels, premature stop codons, or frame-shifts were also considered radical. Intergenic mutations were investigated further if they occurred in a 5’-untranslated region or within 100 bp upstream of the transcriptional start site of a gene. Gene functions were inferred, in part, from automated Gene Ontology (GO) information. To summarize the functional information using broad, high-level GO terms, we used the GO Slim sets provided by the Aspergillus Genome Database (www.aspergillusgenome.org). For enrichment analyses, we aimed to identify the main functional classes, so the results were filtered to retain only the GO Slim terms that were represented by at least 5% of the queried genes. A GO Slim term was considered overrepresented if the observed gene number was > 1.2 times the expected gene number. Statistical significance of overrepresentation was calculated using one-tailed Chi-squared tests and the Benjamini–Hochberg false discovery rate (set to 0.15) correction for multiple comparisons. Given that “Cellular Component,” “Biological Process,” and “Molecular Function” GO Slim annotations were missing for 57, 56, and 54% of genes, respectively, we performed more comprehensive manual annotation of all affected genes using information from other sources, such as similarity and literature searches.

**Data availability**

Sequence data are available from the NCBI Short Read Archive under BioProject PRJNA356622.

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### Table 1: Types and numbers of derived mutations in evolved lines

| Mutation Type          | Small (S) Population Size | Large (L) Population Size |
|------------------------|---------------------------|---------------------------|
|                        | 9S | 16S | 25S | 42S | Total (%) | 8L | 37L | 45L | 59L | L Total (%) |
| Intergenic             | 14 | 6   | 5   | 6   | 31 (38.3) | 3  | 4   | 10  | 9   | 26 (41.3) |
| Intron                 | 1  | 1   | 0   | 1   | 3 (3.7)   | 0  | 1   | 0   | 0   | 1 (1.6)   |
| Coding, synonymous     | 5  | 1   | 4   | 7   | 17 (21.0) | 3  | 1   | 4   | 3   | 11 (17.5) |
| Coding, nonsynonymous  | 9  | 10  | 0   | 6   | 25 (30.9) | 4  | 5   | 7   | 6   | 22 (34.9) |
| Coding, other^         | 2  | 0   | 2   | 1   | 5 (6.2)   | 0  | 1   | 1   | 1   | 3 (4.8)   |
| All mutations          | 31 | 18  | 11  | 21  | 81         | 10 | 12  | 22  | 19  | 63        |

^Includes codon change with codon insertion, stop gained, and frame-shift mutations.

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and read length by only 4.3 and 1.1%, respectively, indicating that the overall sequence quality was relatively high (Table S1). All subsequent analyses were based on trimmed reads.

**Coverage distribution**

An average of 93.7% of reads from the ancestor WG615 and the eight evolved lines were successfully mapped to the FGSC A4 reference genome, resulting in an average coverage of 83.1 x (Table S1). Comparison of fold-coverage among and along scaffolds revealed that WG615 and all descendent, evolved lines shared numerous regions of missing coverage relative to ancestral strain FGSC A4 (Table S2). These absent regions could be up to 32.5 kb in length but still only comprised 1.7% of the reference genome (504.4 kb total). Also, some genome regions of poor alignment and/or low mapping quality were identified (Table S3). It is not clear if these areas represent genomic regions of poor reference assembly, repetitive regions, or regions of genome plasticity. Regardless, these low quality regions sum to only 82.6 kb, or 0.27% of the reference genome, and therefore have a negligible effect on our mutation detection ability. When filtering for changes that arose during the selection experiment, no evidence for large scale indels or aneuploidies were found.

**Variant calls for ancestral WG615 strain**

Variant calling revealed that the ancestral WG615 strain differed from FGSC A4 reference by 740 SNPs and 456 small indels, confirming that these two strains have moderate sequence divergence (0.004%). WG615 is described as having mutations in the wA (AN8209), pyroA (AN7725), and veA (AN1052) genes, so we examined these genes for quality control purposes. Mutations were confirmed for wA and pyroA, and WG615 and FGSC A4 shared the same veA mutant allele, as predicted.

**Candidate genes for fungicide resistance in ancestral WG615 strain**

To identify putative targets of fludioxonil resistance, we took a two-step strategy. We first filtered sequence differences between the fludioxonil-resistant WG615 strain and the fludioxonil-sensitive FGSC A4 strain to retain only those that caused protein sequence changes, under the assumption that these were the most likely to cause changes in gene function and phenotype. Filtering narrowed the field to 390 mutations, of which 237 had GO annotation information available. We screened for genes involved in the HOG MAPK pathway because they are prime candidates for fludioxonil resistance determinants. Just two of these candidate genes (AN2262 and AN4479) had described osmoregulation functions, and only the latter gene, AN4479, is known to respond to fungicides. Notably, AN4479 encodes nika, the histidine kinase sensor of the two-component system of the HOG MAPK pathway.

Second, we reviewed our previous genetic analyses (Schoustra et al. 2006) and found that the resistance mutation was located on the left arm of A. nidulans linkage group III, between the centromere and cys2
The HOG MAPK pathway (Hagiwara 2007) and is responsible for transmitting the phosphorelay signal down (Hagiwara et al. 2013), a fitness trade-off clearly observed in our experiment. Three lines of evidence support that nikA is the target of fludioxonil resistance in WG615. First, nikA is essential for fludioxonil-based growth inhibition (Hagiwara et al. 2007, 2013; Vargas-Perez et al. 2007) and is responsible for transmitting the phosphorelay signal down the HOG MAPK pathway (Hagiwara et al. 2009). Second, nikA mutants with increased resistance to fludioxonil are characterized by strong vegetative growth defects in the absence of the fungicide (Hagiwara et al. 2007; Vargas-Perez et al. 2007; De Souza et al. 2013), a fitness trade-off clearly observed in our fludioxonil-resistant strain (Schoustra et al. 2005, 2009). Third, the WG615 nikA mutation is a nonsynonymous substitution causing a radical amino acid change (K→N) at residue 854, which is within the highly conserved, signal transduction histidine kinase domain. Although formal genetic analyses are needed for full confirmation, the evidence points to the nikA mutation as the determinant of fludioxonil resistance.

**Derived mutations in compensatory evolution lines**

We identified 144 SNPs and indels that differed between the ancestor and the compensatory evolution lines after 800 generations of selection (Table 1, Table S5, and Table S6). An average of 16.75 SNPs and 1.25 small indels were detected in the ~30.5 Mb genome of each evolved line. Derived mutations were relatively evenly distributed throughout the genome (Figure 1), with only a few locations containing multiple mutations clustered together. Only one derived structural variant was called: a ~103 kb inversion was predicted in strain C24 (Table S6B). Substitution rates per chromosome did not differ drastically and ranged from 3.32 to 11.66 × 10⁻¹⁰ per nucleotide per generation (Table S7). The overall substitution rate was 7.58 × 10⁻¹⁰ per nucleotide per generation, similar to rates reported for other fungi such as yeast (Lynch et al. 2008; Nishant et al. 2010). Approximately 50% of the A. nidulans genome is coding, yet over 57% of the observed mutations (83/144) were located within coding regions (Table 1). This difference between observed and expected proportions in coding regions is not statistically significant (Chi-square, $P < 0.194$), but the overrepresentation of coding mutations is consistent with positive selection in our experiment.

**Genes mutated during compensatory adaptation**

The HOG MAPK pathway in A. nidulans senses and responds to multiple forms of potentially harmful stimuli, such as osmotic stress, oxidative stress, DNA damage, and fungicides (Furukawa et al. 2007; Miskei et al. 2009; Balazs et al. 2010; Jaimes-Arroyo et al. 2015). If the nikA resistance-conferring mutation disrupts the normal function of the HOG MAPK signaling pathway, there are likely to be pleiotropic effects on several other physiological processes and phenotypes (e.g., Vargas-Perez et al. 2007; Duran et al. 2010). Our observation of a substantial fitness cost associated with resistance is consistent with this idea. Recent studies of compensatory adaptation in yeast (Szamecz et al. 2014; Filteau et al. 2015) found that derived genetic changes were often specific to the original functional defect, and that compensatory evolution targeted genes that were functionally related to the disrupted gene. Thus, the obvious prediction was that the compensatory fitness gains observed here in A. nidulans would be caused by secondary modification of the HOG MAPK pathway.

Genome-wide analyses in A. nidulans are hampered by a lack of complete functional annotation for genes, particularly when compared to bacterial genomes or more intensively studied fungi like yeast. For example, the most recent A. nidulans genome snapshot indicates that only ~11% of ORFs are verified, while the rest remain uncharacterized. Given the high frequency of missing data, our power to detect statistically significant trends was quite low. However, such results are still
that were specifically associated with the HOG MAPK pathway itself (Table 2). The most obvious example was the atfA gene (AN2911), which encodes a transcription factor in the HOG MAPK pathway and regulates responses to a broad spectrum of environmental stressors (Hagiwara et al. 2009; Balazs et al. 2010; Lara-Rojas et al. 2011). The derived atfA mutation was a radical nonsynonymous change (Q→P) of a conserved amino acid within the functionally important basic-region leucine zipper (bZIP) domain. Another mutated gene with a direct role in the central stress response pathway is mpkC, which encodes a second MAPK that is activated by the same upstream MAPKK as hogA (sakA) MAPK. Overexpression of mpkC has been shown to suppress the high-osmolarity sensitivity observed in hogA-deletion strains (Furukawa et al. 2005), indicating that these two homologous genes have overlapping roles in a shared pathway. No compensatory mutations were observed in nikA itself.

Several mutated genes (Table 2) are known to be expressed in Aspergillus at increased levels in response to osmotic stress or stress-inducing chemicals such as menadione and camptothecin (Malavazi et al. 2006). Camptothecin is a cytotoxic quinoline alkaloid that contains a pyrrole moiety, which is a defining characteristic of phenylpyrrole fungicides such as fludioxonil. This shared chemical structure may explain why both compounds appear to activate similar stress response pathways.

**Signal transduction:** Genes associated with “signal transduction” (GO 7165) were overrepresented 2.3-fold among derived mutations (Chi-square, P < 0.029), many of which had direct roles in the phosphorylation-based transduction of stress response signals. For example, a radical nonsynonymous mutation (S→F) was observed in ppzA, the catalytic subunit of protein phosphatase Z. This serine/threonine phosphatase is known to modulate the response of fungi to oxidative and other stresses (Leiter et al. 2012). A radical nonsynonymous mutation (T→I) was also found in the histidine kinase phkB. Homologs of phkB are stress-activated sensor kinases of two-component systems, indicating a role in phosphorylation-based signal transduction, and a recent study with Aspergillus found that phkB expression increased greatly in response to fludioxonil treatment and osmotic shock in a manner dependent on the HOG pathway (Hagiwara et al. 2013).

**Plasma membrane:** Genes associated with the “plasma membrane” (GO 5886) were overrepresented 2.3-fold among derived mutations (Chi-square, P < 0.026). Remodeling of the fungal membrane and cell wall components is a common response to a wide range of stressors because it helps maintain cell wall integrity and allow for survival and growth (Ene et al. 2015). We found derived mutations in a number of genes (Table 2) with functions in the synthesis and breakdown of fungal cell wall components, such as glucans (agsB, bglM, and AN3504) and chitin (csnA and AN8765; de Groot et al. 2009). For example, a nonsynonymous mutation was observed in the main catalytic domain of agsB, which encodes an α-1,3 glucan synthase with roles in osmotic stress tolerance and cell wall integrity (Futagami et al. 2011). The mutated gene list also contained several membrane-bound transporters with specificities for a range of compounds. Transporters are known to mediate resistance to fungicides and other toxic compounds in many fungi, including Aspergillus (Del Sorbo et al. 2000).

**Intergenic mutations:** Intergenic mutations can alter 5'−untranslated regions (5'−UTRs) or promoter sequences, potentially resulting in changes in gene expression that affect fitness. We analyzed the 57 intergenic mutations (Table S6) and found five that were within 5'−UTRs,
### Table 2 Examples of mutated genes with functions relevant to stress response or membrane and cell wall

| Gene     | Mutation Type | Gene Product | Gene Function |
|----------|---------------|--------------|---------------|
| **Stress response** |               |              |               |
| AN2911   | Nonsynonymous | atfA, basic-region leucine zipper transcription factor activated by HOG MAPK | Stress response (osmotic, fungicide, oxidative) |
| AN3101   | Nonsynonymous | phkB, histidine kinase D5, stress-activated phosphotransfer protein | Phosphorelay sensor kinase activity, stress-activated (osmotic, oxidative, and fungicide) |
| AN3793   | Nonsynonymous | ppzA, serine/threonine-protein phosphatase, catalytic subunit of protein phosphatase Z (PPZ) | Resistance to oxidative stress |
| AN4668   | Synonymous    | mpkC, MAPK, highly similar to HogA | Response to osmotic stress, development |
| AN2204   | Nonsynonymous | Osmoadaptation protein with unknown function | Cellular response to osmotic stress |
| AN4789   | Nonsynonymous | uvsI, DNA polymerase | Response to UV-damage of DNA, postreplication DNA repair |
| AN8562   | Nonsynonymous | Ankyrin repeat protein | Stress response (camptothecin) |
| AN4206   | Nonsynonymous | DnaJ domain protein | Protein folding and chaperone binding, stress response (camptothecin) |
| AN1217   | Nonsynonymous | Putative LIM/homeobox transcription factor | Stress response (camptothecin) |
| AN6280   | Nonsynonymous | Uncharacterized | Stress response (camptothecin) |
| AN5217   | Codon change plus codon insertion | pilA, putative eisosome component | Protein localization, stress response (camptothecin) |
| AN6753   | Synonymous    | Putative NADH-dependent flavin oxidoreductase | Stress response (menadione) |
| AN1700   | Synonymous    | Putative 26S proteasomal regulatory subunit Rpn2 | Proteasome assembly, stress response (camptothecin) |
| **Membrane and cell wall** |               |              |               |
| AN3307   | Nonsynonymous | agsB, catalytic subunit of the major α-1,3 glucan synthase complex | Glucan synthesis, osmotic stress, and cell wall integrity |
| AN3504   | Nonsynonymous | Putative α-1,4-glucosidase | Starch and disaccharide degradation |
| AN7396   | Nonsynonymous | bgIM, putative β-glucosidase | Cellulose and polysaccharide degradation |
| AN8765   | Frame-shift   | Activator of chitin synthase | Response to osmotic stress, regulation of chitin synthase activity and cell wall formation |
| AN4686   | Synonymous    | csnA, chitosanase | Endo-chitosanase activity, predicted glycosyl phosphatidylinositol (GPI)-anchor |
| AN3247   | Frame-shift   | ABC multidrug transporter | Predicted ATP binding, ATPase activity, coupled to transmembrane movement of substances |
| AN2287   | Nonsynonymous | Putative transmembrane transporter | Predicted amino acid transport activity, GABA transport activity |
| AN0660   | Nonsynonymous | furA, putative nucleobase and allantoin transporter | Induced by allantoin and by uric acid |
| AN2699   | Nonsynonymous | Uncharacterized | Predicted transmembrane transport activity, integral component of membrane localization |
| AN3247   | Synonymous    | ABC multidrug transporter | Predicted ATP binding, ATPase activity, coupled to transmembrane movement of substances |
| AN5370   | Synonymous    | Putative MFS multidrug transporter | Predicted transmembrane transport activity, integral component of membrane localization |
| AN3113   | Synonymous    | ugtA, UDP-galactofuranose transporter, DMT family organic anion transporter, multidrug resistance efflux domain | Cell wall architecture, hyphal morphology, and drug sensitivity |
| AN10396  | Synonymous    | Putative farnesyl-diphosphate farnesyltransferase | Lipid biosynthetic process |

See Table S6 for details on specific mutations. HOG, high-osmolarity glycerol; MAPK, mitogen-activated protein kinase; UV, ultraviolet; NADH, nicotinamide adenine dinucleotide hydride; ATP, adenosine triphosphate; GABA, γ-aminobutyric acid; MFS, major facilitator superfamily; UDP, uridine diphosphate; DMT, drug/metabolite transporter.
and eight that were within potential promoter regions (defined as < 100 bp from the transcription start site of a gene). None of the potentially affected genes had known roles in the HOG MAPK pathway, osmotic stress tolerance, or phenylpyrrole resistance.

**Parallel evolution**

Parallel evolution is the repeated evolution of the same genotype or phenotype in independently evolving populations. At the level of fitness, all of the lines we studied evolved in parallel; that is, they all increased in fitness substantially when compared to their ancestor (Schoustra et al. 2009). Only one of the derived, mutated genes was hit in multiple evolved lines (uncharacterized gene AN2078, predicted RNA polymerase II transcription cofactor activity; strain 9S and 8L, Table 5S), providing little evidence for parallel evolution at the gene level. There may be multiple evolutionary pathways to increased fitness in the permissive environment, yet empirical evidence suggests that compensatory evolution would target pathways that were functionally related to the disrupted pathway that caused the original fitness deficit (Szamecz et al. 2014; Filetau et al. 2015).

Current understanding of the HOG pathway [KEGG (Kyoto Encyclopedia of Genes and Genomes), Furukawa et al. 2005; Hagiwara et al. 2009] indicates that it involves 22 of the 10,667 protein-coding genes in *A. nidulans*. We observed derived mutations in only two of these genes (mpkC and atfA), but the probability of 2/87 derived mutations occurring randomly in this gene subset is extremely small (~3.57 x 10^-6). Thus, compensatory adaptation was associated with the HOG pathway more commonly than by chance alone.

Previous analyses of multivariate phenotypic divergence among these *A. nidulans* lines suggested that fitness gains were achieved through distinct phenotypic routes (Schoustra et al. 2012), but the genetic causes were unknown. The lack of strong parallel evolution provided by our sequence data could indicate that the number of potential genes that allow for compensation is large, or it could be an artifact of our low sample size and lack of complete functional information. Future characterization of mutated genes with unknown function will determine if they interact with the HOG pathway, or if they are generally adaptive and completely independent from the pathway involved with fungicide resistance.

**Comparison of step number for fitness-based and sequence-based estimates**

A general result to emerge from genetic studies of adaptation in microbial experiments has been that just a few mutations are usually responsible for the bulk of adaptation achieved during an adaptive walk (Kassen 2014). Indeed, by analyzing fitness data with a maximum likelihood-based approach, we previously estimated that these evolved lines accumulated an average of 2.13 (SE = 0.30) fitness-increasing steps during their adaptive walks (Schoustra et al. 2009). The results from our genomic analyses reveal, at first glance, a strikingly different picture. An average of 18.0 (SE = 2.48) mutations were detected in the genome of each evolved line, or 8.45 times greater than the number of steps inferred from fitness trajectory data. The discrepancy between the number of fitness-based adaptive steps and the total number of sequence-based genomic mutations is likely due to two, nonexclusive reasons.

First, not all mutations observed in the evolved lines are adaptive. Some nonadaptive mutations (neutral, nearly neutral, or even slightly deleterious) may become fixed in the population by chance alone, or may hitch-hike to fixation alongside a beneficial mutation. Hitch-hiking is expected to be especially prevalent because reproduction in this experiment was strictly asexual, leading to complete linkage between beneficial mutations and any others that arose. This effect can be quite pronounced: in experimental evolution studies with yeast, for example, typically less than half of the derived genomic mutations have observed adaptive effects (Anderson et al. 2010; Kvitak and Sherlock 2011).

Second, not all adaptive mutations are detectable in fitness assays. Large fitness changes will be identified, but adaptive mutations that confer small fitness increases may be below the detection threshold of the phenotypic assays. Also, if multiple beneficial mutations arise in the same genome and fix together (Park and Krug 2007; Jerison and Desai 2015), these would be detected as a single fitness-increasing event. Moreover, fitness measures of mycelial growth rate do not assess all components of total fitness. Mutations that have adaptive effects on other fitness components (e.g., mycelial density, sporulation) would not be detected. In these cases, fitness-based step numbers would be underestimated.

Taken together, these explanations suggest that we should be more conservative in our estimate of the number of mutations substituted by focusing attention only on those mutations that are likely to be adaptive. Experimental determination of the individual fitness effects of all 144 derived *A. nidulans* mutations, alone and in combination, is a prodigious task and beyond the scope of the present paper, so we inferred potential adaptive effects based on the mutation location and context within genes. When restricted to mutations occurring within protein-coding DNA, evolved lines had an average of 10.38 (SE = 1.27) mutations, corresponding to 4.87 times the estimated step number. Similarly, when restricted to only those mutations that change the protein sequence, evolved lines had an average of 6.88 (SE = 1.04), or 3.23 times the estimated step number. When restricted to radical amino acid changes (mean = 6.00, SE = 0.96), the difference was only 2.82 times.

Reassuringly, there was a positive correlation between inferred step number and the various measures of genomic mutation number for the evolved lines (Figure 3 and Table 3). Given the low statistical power afforded by a sample size of only eight, it was not surprising that these positive relationships were all statistically nonsignificant. Deviation from linearity was mainly caused by the outlier strain 25S, which had
a high predicted number of steps but low number of genomic mutations. Examination of the data revealed that strain 25S had no detected nonsynonymous mutations, but was the only strain that had a predicted structural variant: an inversion that encompassed 13 genes, most of which are in a secondary metabolite gene cluster (AN7884 cluster; Andersen et al. 2013). This inversion could be contributing to fitness increases, but is not counted toward genomic mutation numbers. When outlier strain 25S was removed from the analyses, most mutation number measures were significantly correlated with step number (Table 3).

### Effect of small vs. large population size

The original selection experiment investigated the effects of population size on the properties of adaptive walks. Small (S) and large (L) effective population size treatments were imposed by bottlenecks of either 500 or 50,000 individuals, respectively, at every transfer. The effect of genetic drift was exaggerated in S populations, leading to lower mean fitness and a more rapid increase in the variance of fitness during the experiment (Schoustra et al. 2009). What effect does a 100-fold difference in population size have on the spectrum of genomic changes? While we have relatively little statistical power with which to detect effects, we do find some intriguing trends.

First, the variance in total mutation number for the S group is over twice as large as that for the L group (68.9 and 32.2, respectively), although the difference is not statistically significant (F-test, \( P > 0.27 \)). Similar patterns are found when each mutation class is analyzed separately. Second, the correlation between fitness-based and sequence-based step number estimates is significantly positive for the L group (\( R = 0.966, P < 0.034 \)) but not for the S group (\( R = 0.181, P > 0.81 \)), consistent with more pronounced effects of drift at smaller population sizes. Third, we might expect that, after controlling for differences in total mutation number per line, the S lines should have accrued more putatively neutral mutations than the L lines. To test this prediction, we calculated the proportion of coding SNPs that were synonymous (silent) and found that S lines did have a greater mean than L lines [0.497 (SE = 0.191) and 0.323 (SE = 0.056), respectively]. Similar results were obtained when we compared the proportion of all mutations that were putatively neutral [intergenic + intron + synonymous; S lines = 0.644 (SE = 0.077), L lines = 0.592 (SE = 0.032)]. While these trends are in the predicted direction (S > L), the differences were not significant (one-tailed t-test; \( P > 0.21 \) and \( P > 0.28 \), respectively), likely due to the low statistical power afforded by only four lines per treatment.

### Conclusions

Here, we examined the genomic basis of compensatory adaptation to the fitness cost of fungicide resistance in experimentally evolved strains of *A. nidulans*. Our work reveals three important insights into the genetics of adaptive evolution in this system.

First, fluoroxylin resistance in the ancestor was likely conferred by a mutation in the histidine kinase nikA, the sensor part of the two-component signal transduction phosphorelay system of the HOG MAPK stress response pathway.

Second, to compensate for the pleiotropic negative effects of the resistance mutation, the subsequent fitness gains observed in the evolved lines were likely caused by secondary modification of the HOG MAPK pathway activity. Candidate genes for the compensatory fitness increases were significantly overrepresented by stress response functions, and some mutations were within genes specifically associated with the HOG MAPK pathway itself. We have generated a candidate list of adaptive mutations, however, further detailed genetic and phenotypic analyses are required to confirm the function and fitness effects of each mutation.

Third, there was a positive relationship between the predicted number of adaptive steps, estimated from fitness data, and the number of genomic mutations, determined by whole-genome sequencing. It is clear, however, that fitness-based methods underestimate the number of genomic mutations because they miss mutations of small effect and cannot distinguish fixation events involving multiple mutations. On the other hand, sequencing-based methods likely overestimate the number of adaptive mutations, especially when recombination is absent, because many nonadaptive mutations rise to high frequency by hitchhiking alongside beneficial mutations.

Interestingly, the proportion of mutations predicted to be adaptive in our *Aspergillus* experiment is substantially lower than comparable experiments (Elena and Lenski 2003; Dettman et al. 2012; Kassen 2014; Long et al. 2015). If we assume, very approximately, that the fitness-based estimates of step number have correctly identified all adaptive mutations, then our sequencing results suggest that, on average, only 12% of the genomic mutations are adaptive. When considering only those mutations that alter the protein sequence, 31% of mutations are adaptive. Such low proportions of adaptive mutations may be due to the higher fraction of the *Aspergillus* genome that is noncoding, compared to bacterial genomes. Also, the spatially structured, multicellular growth form of filamentous fungi may reduce the efficiency of selection. For example, growing as an interconnected mycelial network may hamper competition among contending mutations, and competitive success may depend upon where within the mycelium the new variant arises. These reasons may explain why the proportion of total mutations we predict to be adaptive in *Aspergillus* are low compared to those reported for other, typically unicellular, organisms. In most bacterial experiments, for example, the vast majority of mutations tend to be adaptive (Dettman et al. 2012). The generality of our results cannot be determined because, to date, only a few studies have applied the evolve-and-resequencing approach to multicellular eukaryotes (e.g., *Drosophila*, Burke et al. 2010; Kang et al. 2016).

Our study with *Aspergillus* lays out a framework for future research on the genomics of compensatory adaptation. Now that we have

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**Table 3** Correlations between number of estimated steps and other variables

| Comparison | Eight Evolved Lines | Seven Evolved Lines* |
|------------|---------------------|----------------------|
|            | \( R \) | \( P \) | \( R \) | \( P \) |
| Step number vs. mycelial growth rate | 0.460 | 0.251 | 0.330 | 0.4701 |
| Step number vs. number of SNPs | 0.534 | 0.173 | 0.930 | **0.0024** |
| Step number vs. number of SNPs and indel combined | 0.562 | 0.147 | 0.884 | **0.0083** |
| Step number vs. number of SNPs and indels in protein-coding DNA | 0.555 | 0.153 | 0.970 | **0.0003** |
| Step number vs. number of protein-changing SNPs and indels | 0.181 | 0.667 | 0.689 | 0.0869 |
| Step number vs. number of radical amino acid changes | 0.251 | 0.548 | 0.689 | 0.0869 |

*Values in bold indicate \( P < 0.05 \). SNP, single nucleotide polymorphism.

*Testing for the effect of removing outlier (25S).
identified some clear trends (e.g., stress response and HOG pathway) based on analyses of only eight evolved lines, it will be interesting to see how these patterns hold when the sample size is expanded to dozens or even hundreds of evolved lines. Such increased sample sizes will mitigate our problems with lack of statistical power, and will also allow us to more thoroughly assess the degree of parallelism in genomic evolution during compensatory adaptation.

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LITERATURE CITED
Andersen, M. R., J. B. Nielsen, L. M. Petersen, M. Zachariassen et al., 2013 Accurate prediction of secondary metabolite gene clusters in filamentous fungi. Proc. Natl. Acad. Sci. USA 110: E99–E107.
Anderson, J. B., J. Funt, D. A. Thompson, S. Prabhu, A. Socha et al., 2010 Determinants of divergent adaptation and Dobzhansky–Muller interaction in experimental yeast populations. Curr. Biol. 20: 1383–1388.
Balazs, A., I. Pocsi, Z. Hamari, E. Leiter, T. Emri et al., 2010 AtfA bZIP-type transcription factor regulates oxidative and osmotic stress responses in Aspergillus nidulans. Mol. Genet. Genomics 283: 289–303.
Barrick, J. E., G. Colburn, D. E. Deatherage, C. C. Traverse, M. D. Strand et al., 2014 Identifying structural variation in haploid microbial genomes from short-read resequencing data using breseq. BMC Genomics 15: 1039.
Burke, M. K., J. P. Dunham, P. Shahrzadi, K. R. Thornton, M. R. Rose et al., 2010 Genome-wide analysis of a long-term evolution experiment with Drosophila. Nature 467: 587–590.
Cingolani, P., A. Platts, L. L. Wang, M. Coon, T. Nguyen et al., 2012 A program for annotating and predicting the effects of single nucleotide polymorphisms, SnpEff: SNPs in the genome of Drosophila melanogaster strain w(1118); iso-2; iso-3. Fly (Austin) 6: 80–92.
Conrad, T. M., A. R. Joyce, M. K. Applebee, C. L. Barrett, B. Xie et al., 2009 Whole-genome resequencing of Escherichia coli K-12 MG1655 undergoing short-term laboratory evolution in lactate minimal media reveals flexible selection of adaptive mutations. Genome Biol. 10: R118.
de Groot, P. W. J., B. W. Brandt, H. Horiiuchi, A. F. J. Ram, C. G. de Koster et al., 2009 Comprehensive genomic analysis of cell wall genes in Aspergillus nidulans. Fungal Genet. Biol. 46: S72–S81.
Del Sorbo, G., H. J. Schoonbeek, and M. A. De Waard, 2000 Fungal transporters involved in efflux of natural toxic compounds and fungicides. Fungal Genet. Biol. 30: 1–15.
De Souza, C. P., S. B. Hashmi, A. H. Osmani, P. Andrews, C. S. Ringelberg et al., 2013 Functional analysis of the Aspergillus nidulans kinome. PLoS One 8: e58008.
Dettman, J. R., N. Rodrigue, A. H. Melnyk, A. Wong, S. F. Bailey et al., 2012 Evolutionary insight from whole-genome sequencing of evolutionarily evolved microbes. Mol. Ecol. 21: 2058–2077.
Durán, R., J. W. Cary, and A. M. Calvo, 2010 Role of the osmotic stress regulatory pathway in morphogenesis and secondary metabolism in filamentous fungi. Toxins (Basel) 2: 367–381.
Elena, S. F., and R. E. Lenski, 2003 Evolution experiments with microorganisms: the dynamics and genetic bases of adaptation. Nat. Rev. Genet. 4: 457–469.
Ene, I. V., L. A. Walker, M. Schiavone, K. K. Lee, H. Martin-Yken et al., 2015 Cell wall remodeling enzymes modulate fungal cell wall elasticity and osmotic stress resistance. MBio. 6: e00986.
Filleul, M., V. Hamel, M.-C. Pouliot, I. Gagnon-Arsenault, A. K. Dube et al., 2015 Evolutionary rescue by compensatory mutations is constrained by genomic and environmental backgrounds. Mol. Syst. Biol. 11: 832.
Fisher, R. A., 1930 The Genetical Theory of Natural Selection. Oxford University Press, Oxford, UK.
Furukawa, K., Y. Hoshi, T. Maeda, T. Nakajima, and K. Abe, 2005 Aspergillus nidulans HOG pathway is activated only by two-component signalling pathway in response to osmotic stress. Mol. Microbiol. 56: 1246–1261.
Furukawa, K., A. Yoshimi, T. Furukawa, Y. Hoshi, D. Hagiwara et al., 2007 Novel reporter gene expression systems for monitoring activation of the Aspergillus nidulans HOG pathway. Biosci. Biotechnol. Biochem. 71: 1724–1730.
Futagami, T., S. Nakao, Y. Kido, T. Oka, Y. Kajiwara et al., 2011 Putative stress sensors WscA and WscB are involved in hypo-osmotic and acidic pH stress tolerance in Aspergillus nidulans. Eukaryot. Cell 10: 1504–1515.
Galagan, J. E., S. E. Calvo, C. Cuomo, L. J. Ma, J. R. Wortman et al., 2005 Sequencing of Aspergillus nidulans and comparative analysis with A. fumigatus and A. oryzae. Nature 438: 1105–1115.
Hagiwara, D., Y. Matsubayashi, J. Marui, K. Furukawa, T. Yamashino et al., 2007 Characterization of the NikA histidine kinase implicated in the phosphorelay signal transduction of Aspergillus fumigatus, with special reference to fungicide responses. Biosci. Biotechnol. Biochem. 71: 844–847.
Hagiwara, D., Y. Asano, J. Marui, A. Yoshimi, T. Mizuno et al., 2009 Transcriptional profiling for Aspergillus nidulans HogA MAPK signaling pathway in response to fluoroquinol and osmotic stress. Fungal Genet. Biol. 46: 868–878.
Hagiwara, D., A. Takahashi-Nakaguchi, T. Toyotome, A. Yoshimi, K. Abe et al., 2013 NikA/Tsc histidine kinase is involved in conidiation, hyphal morphology, and responses to osmotic stress and antifungal chemicals in Aspergillus fumigatus. PLoS One 8: e80881.
Jaimies-Arroyo, R., F. Lara-Rojas, O. Bayram, O. Valerius, G. H. Braus et al., 2015 The SrkA kinase is part of the SaA mitogen-activated protein kinase interactome and regulates stress responses and development in Aspergillus nidulans. Eukaryot. Cell 14: 495–510.
Jerison, E. R., and M. M. Desai, 2015 Genomic investigations of evolutionary dynamics and epistasis in microbial evolution experiments. Curr. Opin. Genet. Dev. 35: 33–39.
Kang, L. D. D. Aggarwal, E. Rashkovetsky, A. B. Korol, and P. Michalak, 2016 Rapid genomic changes in Drosophila melanogaster adapting to desiccation stress in an experimental evolution system. BMC Genomics 17: 233.
Kassen, R., 2014 Experimental Evolution and the Nature of Biodiversity. Roberts and Company, Greenwood Village, CO.
Koboldt, D. C., Q. Zhang, D. E. Larson, D. Shen, M. D. McLellan et al., 2012 VarScan 2: somatic mutation and copy number alteration discovery in cancer by exome sequencing. Genome Res. 22: 568–576.
Koříř, R., P. Orozco-terWengel, N. De Maio, R. V. Pandey, V. Nolte et al., 2011 PoPoolation: a toolbox for population genetic analysis of next generation sequencing data from pooled individuals. PLoS One 6: e15925.
Kvitv, D. J., and G. Sherlock, 2011 Reciprocal sign epistasis between frequently evolutionally adapted mutations causes a rugged fitness landscape. PLoS Genet. 7: e1002056.
Lara-Rojas, F., O. Sánchez, L. Kawasaki, and J. Aguirre, 2011 Aspergillus nidulans transcription factor AtfA interacts with the MAPK SakA to regulate general stress responses, development and spore functions. Mol. Microbiol. 80: 436–454.
Leiter, E., A. Gonzalez, E. Erdei, C. Casado, L. Kovacs et al., 2012 Protein phosphatase Z modulates oxidative stress response in fungi. Fungal Genet. Biol. 49: 708–716.
Li, H., B. Handsaker, A. Wysocker, T. Fennell, J. Ruan et al., 2009 The sequence alignment/map format and SAMtools. Bioinformatics 25: 2078–2079.
Long, A., G. Liti, A. Luptak, and O. Tenailleon, 2015 Elucidating the molecular architecture of adaptation via evolve and resequence experiments. Nat. Rev. Genet. 16: 567–582.
Lynch, M., W. Sung, K. Morris, N. Coffey, C. R. Landry et al., 2008 A genome-wide view of the spectrum of spontaneous mutations in yeast. Proc. Natl. Acad. Sci. USA 105: 9272–9277.
Malavazi, I., M. Savoldi, S. M. Zingaretti Di Mauro, C. F. Martins Menck, S. D. Harris et al., 2006 Transcriptome analysis of Aspergillus nidulans exposed to camptothecin-induced DNA damage. Eukaryot. Cell 5: 1688–1704.
Miskei, M., Z. Karanyi, and I. Pocsi, 2009  Annotation of stress-response proteins in the aspergilli. Fungal Genet. Biol. 46: S105–S120.

Nishant, K. T., W. Wei, E. Mancera, J. L. Argueso, A. Schlattl et al., 2010  The baker’s yeast diploid genome is remarkably stable in vegetative growth and meiosis. PLoS Genet. 6: e1001109.

Orr, H. A., 1998  The population genetics of adaptation: the distribution of factors fixed during adaptive evolution. Evolution 52: 935–949.

Orr, H. A., 2005  The genetic theory of adaptation: a brief history. Nat. Rev. Genet. 6: 119–127.

Park, S.-C., and J. Krug, 2007  Clonal interference in large populations. Proc. Natl. Acad. Sci. USA 104: 18135–18140.

Riehle, M. M., A. F. Bennett, and A. D. Long, 2001  Genetic architecture of thermal adaptation in Escherichia coli. Proc. Natl. Acad. Sci. USA 98: 525–530.

Schoustra, S. E., M. Slakhorst, A. J. M. Debets, and R. F. Hoekstra, 2005  Comparing artificial and natural selection in rate of adaptation to genetic stress in Aspergillus nidulans. J. Evol. Biol. 18: 771–778.

Schoustra, S. E., A. J. M. Debets, M. Slakhorst, and R. F. Hoekstra, 2006  Reducing the cost of resistance; experimental evolution in the filamentous fungus Aspergillus nidulans. J. Evol. Biol. 19: 1115–1127.

Schoustra, S. E., T. Bataillon, D. R. Gifford, and R. Kassen, 2009  The properties of adaptive walks in evolving populations of fungus. PLoS Biol. 7: e1000250.

Schoustra, S. E., D. Punzalan, R. Dali, H. D. Rundle, and R. Kassen, 2012  Multivariate phenotypic divergence due to the fixation of beneficial mutations in experimentally evolved lineages of a filamentous fungus. PLoS One 7: e50305.

Szamecz, B., G. Boross, D. Kalapis, K. Kovacs, G. Fekete et al., 2014  The genomic landscape of compensatory evolution. PLoS Biol. 12: e1001935.

Vargas-Perez, I., O. Sanchez, L. Kawasaki, D. Georgellis, and J. Aguirre, 2007  Response regulators SrrA and SskA are central components of a phosphorelay system involved in stress signal transduction and asexual sporulation in Aspergillus nidulans. Eukaryot. Cell 6: 1570–1583.

Wichman, H. A., M. R. Badgett, L. A. Scott, C. M. Boulianne, and J. J. Bull, 1999  Different trajectories of parallel evolution during viral adaptation. Science 285: 422–424.

Ye, K., M. H. Schulz, Q. Long, R. Apweiler, and Z. Ning, 2009  PinDLe: a pattern growth approach to detect break points of large deletions and medium sized insertions from paired-end short reads. Bioinformatics 25: 2863–2871.

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