Signatures of Adaptation to Obligate Biotrophy in the *Hyaloperonospora arabidopsidis* Genome

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Many oomycete and fungal plant pathogens are obligate biotrophs, which extract nutrients only from living plant tissue and cannot grow apart from their hosts. Although these pathogens cause substantial crop losses, little is known about the molecular basis or evolution of obligate biotrophy. Here, we report the genome sequence of the oomycete *Hyaloperonospora arabidopsidis* (*Hpa*), an obligate biotroph and natural pathogen of *Arabidopsis thaliana*. In comparison with genomes of related, hemibiotrophic Phytophthora species, the *Hpa* genome exhibits dramatic reductions in genes encoding (i) RXLR effectors and other secreted pathogenicity proteins, (ii) enzymes for assimilation of inorganic nitrogen and sulfur, and (iii) proteins associated with zoospore formation and motility. These attributes comprise a genomic signature of evolution toward obligate biotrophy.

The oomycete *Hyaloperonospora arabidopsidis* (*Hpa*, formerly *Peronospora parasitica* or *Hyaloperonospora parasitica*) is a natural pathogen of *Arabidopsis thaliana* and a model for dissection of *A. thaliana* pathogen response networks (1, 2). *Hpa* belongs to a group of “downy mildew” pathogens, comprising more than 800 species that cause disease on hundreds of plant species (3). Downy mildew pathogens are related to other destructive oomycete plant pathogens (e.g., *Phytophthora* species) (4, 5). Oomycetes belong to the kingdom Stramenopila, which includes brown algae and diatoms. Although oomycetes and fungi share morphological and ecological similarities, they evolved independently to colonize plants.

*Hpa* hyphae grow between plant cells and establish feeding structures called haustoria, which have also evolved in fungal pathogens (2, 6). Downy mildews are obligately biotrophic and cannot be cultured apart from their hosts. In contrast, *Phytophthora* species are hemibiotrophic; an initial phase of biotrophic growth is followed by a necrotrophic phase. Molecular phylogenies show that downy mildews arose from a paraphyletic, *Phytophthora*-like, hemibiotrophic ancestor (4, 5, 7). Thus, insight into the genomic basis and evolution of obligate biotrophy can be obtained through comparison of the *Hpa* genome to the recently sequenced genomes of *Phytophthora* species (8, 9).

Genome analysis was performed on the *Hpa* Emo2 isolate (1) using DNA from asexual spores (10). Sanger shotgun sequencing at 9.5-fold coverage, combined with 97 sequenced bacterial artificial chromosome inserts, yielded an assembly of 77.8 Mb. Illumina sequencing at 46-fold coverage yielded 3.8 Mb of additional sequence, which was integrated into the Sanger assembly to form the 81.6-Mb final version (v8.3.2). Forty-two percent yielded 3.8 Mb of additional sequence, which was integrated into the Sanger assembly to form the 81.6-Mb final version (v8.3.2). Forty-two percent of 31,759 expressed sequence tag (EST) reads aligned to the assembly, indicating conserved single-copy eukaryotic genes (fig. S2). Moreover, 94% of 31,759 expressed sequence tag (EST) reads aligned to the assembly, indicating that the draft genome assembly encompasses a very high percentage of the *Hpa* gene space.

A total of 14,543 genes were computationally predicted in v8.3, of which 80% are supported by ESTs and/or Illumina cDNA tags. This predicted gene content is similar to *P. ramorum* (65 Mb, 15,743 genes) and lower than *P. sojae* (95 Mb, 19,027 genes) (9) or *P. infestans* (240 MB, 17).

## Table 1. Copy numbers of annotated *Hpa* genes for hydrolases, PAMPS, and effectors, compared with *Phytophthora* genomes.

<table>
<thead>
<tr>
<th>Gene product</th>
<th><em>H. arabidopsidis</em></th>
<th><em>P. sojae</em></th>
<th><em>P. ramorum</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Extracellular proteases</td>
<td>18</td>
<td>47</td>
<td>48</td>
</tr>
<tr>
<td>Glycosyl hydrolases</td>
<td>&gt;60</td>
<td>125</td>
<td>114</td>
</tr>
<tr>
<td>Endoglucanases (EGL12)</td>
<td>3</td>
<td>10</td>
<td>8</td>
</tr>
<tr>
<td>Polygalacturonases</td>
<td>3</td>
<td>25</td>
<td>16</td>
</tr>
<tr>
<td>Pectin methyl esterases</td>
<td>3</td>
<td>19</td>
<td>15</td>
</tr>
<tr>
<td>Cutinases</td>
<td>2</td>
<td>16</td>
<td>4</td>
</tr>
<tr>
<td>Chitinases</td>
<td>1</td>
<td>5</td>
<td>2</td>
</tr>
<tr>
<td>Elicitins</td>
<td>1</td>
<td>18</td>
<td>17</td>
</tr>
<tr>
<td>Elicitin-like</td>
<td>14</td>
<td>39</td>
<td>31</td>
</tr>
<tr>
<td>CBEL and CBEL-like</td>
<td>2</td>
<td>13</td>
<td>15</td>
</tr>
<tr>
<td>RXLR</td>
<td>134</td>
<td>396</td>
<td>374</td>
</tr>
<tr>
<td>NLP</td>
<td>10</td>
<td>29</td>
<td>40</td>
</tr>
<tr>
<td>Crinklers</td>
<td>20</td>
<td>40</td>
<td>8</td>
</tr>
<tr>
<td>PPAT12/24-like</td>
<td>8</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

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17,887 genes) (8). A total of 6882 predicted genes in Hpa had no identifiable ortholog in sequenced Phytophthora species or similarity to known proteins, and as such represent potentially lineage-specific genes. Some of these genes may play roles that are specific to biotrophy. For example, a novel family of secreted small, cysteine-rich proteins exists in Hpa (PPAT12/24-like) (Table 1) (11).

Pathogenicity genes were compared among Hpa and Phytophthora species, revealing that families encoding host-targeted, degradative enzymes (secreted proteinases, cell wall–degrading enzymes) are reduced in Hpa (Table 1). Two notable examples are the family 12 endoglucanases (EGL12) and pectin methyl esterases (Pect). Phylogenetic analyses delineated several EGL12 and Pect gene clades containing genes from P. sojae and P. ramorum but not Hpa (figs. S6 and S7). Because Hpa and P. sojae likely share a sister group relationship relative to P. ramorum (4, 5), it is probable that a number of EGL12 and Pect gene were lost from Hpa after divergence of the lineage leading to Hpa and P. sojae. Hydrolytic enzymes that target the host cell wall can release cell wall fragments that elicit host defenses. It is conceivable that in evolving a biotrophic lifestyle, Hpa has lost most of the secreted hydrolytic enzymes that were present in a hemibiotrophic ancestor.

Similarly, gene families encoding necrosis and ethylene-inducing (Nep1)-like proteins (NLPs) are significantly reduced in Hpa, compared with P. sojae and P. ramorum. NLPs in Phytophthora and Pythium can trigger plant cell death and defenses (12), which are implicated in the transition from biotrophy to necrotrophy. Only three of the 13 oomycete NLP clades contain genes from Hpa. However, one clade contains an expanded family that is specific to Hpa (Fig. 1A). All 10 HaNLP genes are supported by transcriptomic data. Of these, HaNLP3 is most closely related to the PsNLP and PiNLP1.1 proteins, but it did not induce necrosis in Nicotiana tabacum (Fig. 1B). These results suggest that downy mildew NLP genes may have evolved a different function than in Phytophthora. Copy number reduction was also evident for genes encoding known pathogen-associated molecular patterns (PAMPs) such as sterol-binding elicitors (13) and carbohydrate-binding CBEL (cellulose-binding, elicitor, lectin-like) genes (14) (Table 1). These examples further suggest that selection for “stealth” (avoidance of host defenses) was a major force during downy mildew evolution.

Phytophthora genomes encode hundreds of potential effector proteins (9, 15, 16) with RXLR cell entry motifs (16–18) that likely function to suppress host defenses (19, 20). The Hpa genome contained 134 high-confidence effector gene candidates (HaRXL genes), including the known effector genes AtR1 and AtR13 (21, 22), significantly fewer than in the Phytophthora genomes (9, 15). Single-nucleotide polymorphisms arising from heterozygosity in v8.3 occurred at a rate 5 times as high in RXLR effector candidates [1 per ~500 base pairs (bp)] than in other genes (1 per ~2500 bp). Only 36% of the high-confidence Hpa effectors had significant matches in any Phytophthora genome (sequence similarity >30%), consistent with strong divergent selection on

Fig. 1. Diversity, evolutionary history, and functional analysis of oomycete necrosis and NLPs. (A) Phylogeny of oomycete NLPs. A consensus tree from the Bayesian inference is shown. Thick lines indicate high support in minimum evolution (>90), maximum likelihood (>90), and Bayesian inference (>0.95). Hollow lines indicate branches highly supported in at least two analyses. Branches with high support in less than two analyses are represented by thin lines. (B) An Hpa NLP ortholog does not induce necrosis in plant leaves. NLP genes were transiently expressed in Nicotiana tabacum by agroinfiltration.
RXLR effector genes (15, 23). Moreover, Hpa effectors generally were not located in syntenic locations relative to Phytophthora genomes, except for three families of effectors, which have unusually high levels of sequence conservation (Fig. 2).

As obligate biotrophs, downy mildews may have lost some metabolic pathways. We identified several potential metabolic defects in Hpa compared with P. sojae and P. ramorum (fig. S9). For example, genes for nitrate and nitrite reductases, a nitrate transporter, and sulfite reductase were missing (fig. S10 and table S3), which is also a feature of the genomes of obligately parasitic powdery mildew fungi (24). Hpa also lacks genes required for synthesis of arachidonic acid and polyamine oxidases.

Flagellated zoospores are produced by many oomycetes (25). Contrasting, several downy mildew lineages germinate by extending infective germ tubes from nonmotile conidiospores, although evidence exists for a rare zoospore stage in some otherwise conidial downy mildews (26, 27). To conclusively determine whether spor motility has been lost from the Hpa lineage, we searched the Hpa genome for 90 flagellum-associated genes using Chlamydomonas sequences and their Phytophthora orthologs (28). No matches were detected in Hpa for any of these. Similarly, many Phytophthora mechanism-related genes are reduced in number or absent from Hpa, consistent with the lack of adherent cysts that normally develop from zoospores during infection.

Analysis of Hpa gene space revealed genomic signatures of major alterations in pathogenic strategy, metabolism, and development that occurred during the evolution of obligate biotrophy from a facultative, hemibiotrophic ancestor. Interestingly, some features of Hpa gene space (large numbers of secreted effectors, reduction in degradative enzymes, and loss of N and S assimilation) are mirrored in genomes of biotrophic fungi (24, 29, 30). These similarities indicate that convergent adaptations occurred during the independent evolution of biotrophy in fungal and oomycete lineages.

References and Notes
10. Materials and methods are available as supporting material on Science Online.
18. S. Kale et al., Cell 142, 284 (2010).
31. We thank E. Holub for providing the Emoy2 isolate, D. Greenhields and N. Bruce for technical assistance, A. Heck and M. Slijper for analysis of secreted Hpa proteins, R. Hubley for creating repeat modeller libraries, and participants in the 2007 Annotation Jamboree and in the 2008 and 2009 Oomycte Bioinformatics Training Workshops for sequence annotations. This research was supported by grants EF-0412213, IOS-0744875, IOS-0924861, and MCB-0639226 from the U.S. NSF and 2004-35600-15055 and 2007-35319-18100 from the U.S. Department of Agriculture National Institute of Food and Agriculture to B.M.T. and J.M.M.; Biotechnology and Biological Sciences Research Council (BBiSc) BB/E024153/1, BB/E024888/1, and Engineering and Physical Sciences Research Council (BBiSc) Systems Biology DTC student EPSF500025/S to J.B; Gatsby GATZ245 and BBiSc BB/F016190, BB/E024888/1, and BBiSc CASE studentship T12144 to J.D.G. Other support is detailed in the supporting online material. Genome browsers are maintained at the Virginia Bioinformatics Institute (vmd.vbi.vt.edu) and the Sainsbury Laboratories (gbrowse2.tsl.ac.uk/cgi-bin/gb/gbrowse/hpa_emo2_publication).

Supporting Online Material
www.sciencemag.org/cgi/content/full/330/6010/1549/DC1
Materials and Methods
Figs. S1 to S10
Tables S1 to S3
References
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The Major Genetic Determinants of HIV-1 Control Affect HLA Class I Peptide Presentation

The International HIV Controllers Study*†

Infectious and inflammatory diseases have repeatedly shown strong genetic associations within the major histocompatibility complex (MHC); however, the basis for these associations remains elusive. To define host genetic effects on the outcome of a chronic viral infection, we performed genome-wide association analysis in a multiethnic cohort of HIV-1 controllers and progressors, and we analyzed the effects of individual amino acids within the classical human leukocyte antigen (HLA) proteins. We identified >300 genome-wide significant single-nucleotide polymorphisms (SNPs) within the MHC and none elsewhere. Specific amino acids in the HLA-B peptide binding groove, as well as an independent HLA-C effect, explain the SNP associations and reconcile both protective and risk HLA alleles. These results implicate the nature of the HLA–viral peptide interaction as the major factor modulating durable control of HIV infection.

HIV infection is characterized by acute viremia, often in excess of 5 million viral particles per milliliter of plasma, followed by an average 100-fold or greater decline to a relatively stable plasma virus load set point (1). In the absence of antiretroviral therapy, the level of viremia is associated with the rate of CD4+ T cell decline and progression to AIDS. There is substantial interperson variability in the virus load set point, with most individuals having stable levels exceeding 10,000 RNA copies/ml. Yet a small number of people demonstrate sustained ability to control HIV replication without therapy. Such individuals, referred to as HIV controllers, typically maintain stable CD4+ cell counts, do not develop clinical disease, and are less likely to transmit HIV to others (2).

To determine the genetic basis for this rare phenomenon, we established a multinational consortium (www.hivcontrollers.org) to recruit HIV-1 controllers, who are defined by at least three measurements of plasma virus load (VL) < 2000 RNA copies/ml over at least a 12-month period in the absence of antiviral therapy. We performed a genome-wide association study (GWAS) in the HIV controllers (median VL, CD4 count, and disease duration of 241 copies/ml, 699 cells/mm³, and 10 years, respectively) and treatment-naïve chronically infected individuals with advanced disease (median VL and CD4 count of 61,698 copies/ml and 224 cells/mm³, respectively) enrolled in antiviral treatment studies led by the AIDS Clinical Trials Group. After quality control and imputation on the basis of HapMap...