LETTERS

Genome sequence and analysis of the Irish potato famine pathogen *Phytophthora infestans*

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Phytophthora infestans is the most destructive pathogen of potato and a model organism for the oomycetes, a distinct lineage of fungus-like eukaryotes that are related to organisms such as brown algae and diatoms. As the agent of the Irish potato famine in the mid-nineteenth century, *P. infestans* has had a tremendous effect on human history, resulting in famine and population displacement¹. To this day, it affects world agriculture by causing the most destructive disease of potato, the fourth largest food crop and a critical alternative to the major cereal crops for feeding the world's population¹. Current annual worldwide potato crop losses due to late blight are conservatively estimated at \$6.7 billion². Management of this devastating pathogen is challenged by its remarkable speed of adaptation to control strategies such as genetically resistant cultivars^{3,4}. Here we report the sequence of the *P. infestans* genome, which at ~240 megabases (Mb) is by far the largest and most complex genome sequenced so far in the chromalveolates. Its expansion results from a proliferation of repetitive DNA accounting for ~74% of the genome. Comparison with two other *Phytophthora* genomes showed rapid turnover and extensive expansion of specific families of secreted disease effector proteins, including many genes that are induced during infection or are predicted to have activities that alter host physiology. These fast-evolving effector genes are localized to highly dynamic and expanded regions of the *P. infestans* genome. This probably plays a crucial part in the rapid adaptability of the pathogen to host plants and underpins its evolutionary potential.

The size of the *P. infestans* genome is estimated by optical map and other methods at 240 Mb (Supplementary Information). It is several-fold larger than those of the related *Phytophthora* species *P. sojae*

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(95 Mb) and P. ramorum (65 Mb), which cause soybean root rot and sudden oak death, respectively5,6. We sequenced the genome of P. infestans strain T30-4 using a whole-genome shotgun approach, and generated a ninefold coverage assembly spanning 229 Mb (Table 1 and Supplementary Information). The unassembled fraction of the genome consists of high copy repeat sequences (Supplementary Information). The assembled genome sequence provides near complete coverage of genes, with 98.2% of P. infestans T30-4 complementary DNAs aligning (Supplementary Information). We identified 17,797 protein-coding genes by ab initio gene prediction, protein and expressed sequence tag (EST) homology, and direct genome-togenome comparative gene modelling with P. sojae and P. ramorum (Supplementary Information). Changes in gene content, number or length do not explain the marked difference in genome size (Table 1 and Supplementary Table 1). No evidence of whole-genome duplication or large-scale dispersed segmental duplication was detected. However, specific disease effector gene families are expanded in P. infestans (see later).

P. infestans, P. sojae and *P. ramorum* represent three major phylogenetic clades of *Phytophthora*⁶. Among the three genomes, we identified a core set of 8,492 orthologue clusters (including 9,583 *P. infestans* orthologues and close paralogues), of which 7,113 genes show 1:1:1 orthology relationships (Table 1, Supplementary Fig. 1 and Supplementary Table 2). The core proteome is enriched in genes involved in cellular processes including DNA replication, transcription and protein translation, whereas genes with functions involved in cellular defence mechanisms are underrepresented (Supplementary Fig. 2). Differences in gene family expansion, in particular dynamic repertoires of effector genes (see later), are probably responsible for different traits among *Phytophthora* species, such as altered host specificity.

Comparison of the three *Phytophthora* genomes reveals an unusual genome organization, comprised of blocks of conserved gene order in which gene density is relatively high and repeat content is relatively low, separated by regions in which gene order is not conserved, gene density is low and repeat content is high (Table 1 and Fig. 1). The conserved blocks represent ~90% of core orthologous groups in all three genomes, including ~70% (12,440) of all *P. infestans* protein-coding genes and ~78% of genes in both *P. sojae* (13,225) and *P. ramorum* (11,246). Within conserved blocks, genes are typically tightly spaced in all three genomes (Table 1 and Fig. 1), with median intergenic distances of 633 base pairs (bp) for *P. ramorum*, 804 bp for

Table 1	Genome	assembly	and	annotation	statistics
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P. sojae, and 603 bp for P. infestans. In regions between conserved blocks, intergenic distances are greater and increase with increasing genome size (median 1.5 kb for P. ramorum, 2.2 kb for P. sojae, and 3.7 kb for P. infestans). The differences in spacing between genes among the three genomes, within and outside regions of conserved gene order, are evident in Fig. 2a-f. The expansion of regions between conserved blocks results from increased density of repetitive elements (Supplementary Fig. 3), and overall differences in genome size among the three species are largely explained by proliferation of repeats in regions in which gene order is not conserved. This difference between conserved blocks and non-conserved regions is particularly apparent in the greatly expanded P. infestans genome (Fig. 2d, f). Further, it is evident that rapidly evolving secreted effector genes (see later) lie predominantly in the gene-sparse regions (Fig. 2g, h). This dual pattern of intergenic spacing and repeat content has been suggested for large, unsequenced genomes in the Poaceae such as maize⁷⁻⁹, but it is not seen in the genomes of other sequenced eukaryotes (Supplementary Fig. 4).

Recent proliferation of Gypsy elements in *P. infestans* underlies the genome expansion. Approximately one-third of the genome assembly corresponds to families of Gypsy elements (Supplementary Fig. 5). The two families with the highest relative expansion in *P. infestans* are Gypsy Pi-1 and a new Gypsy long terminal repeat (LTR) element we named 'Albatross', which together account for at least 29% of the genome (Supplementary Table 3). Albatross elements cover \sim 32 Mb and are enriched (>2-fold) in the regions in which gene order is not conserved (Supplementary Table 4 and Supplementary Fig. 6), contributing appreciably to relative expansion of gene-sparse regions (Supplementary Fig. 3). Gypsy Pi-1 elements cover \sim 22 Mb and, in contrast to Albatross elements, are relatively evenly distributed across the genome.

Overall, the *P. infestans* genome contains a strikingly rich and diverse population of transposons (Supplementary Table 3). We identified 273 full-length elements belonging to two large classes of autonomous rolling-circle type helitron DNA transposons (7.3-kb and 6.4-kb elements), in much larger numbers than described in any other genome (Supplementary Tables 3 and 5). Most helitron open reading frames (ORFs) are degenerate pseudogenes, but 13 are intact and presumed functional. Some apparently non-autonomous helitrons have intact termini so their transposition may be driven by gene products from the functional classes. In contrast, the *P. sojae* and *P. ramorum* genomes contain no intact helitron elements. The *P. infestans* genome carries

	P. infestans	P. sojae	P. ramorum
Genome			
Estimated genome size	240 Mb	95 Mb*	65 Mb*
Coverage (fold)	7.6	7.9*	5.6*
Number of scaffolds	4,921	1,810*	2,576*
N50 scaffold length	1,570 kb	463 kb*	308 kb*
Total scaffold length	228.5 Mb	86.0 Mb*	66.7 Mb*
Number of contigs	18,288	5,577*	7,588*
N50 contig length	44.5 kb	105.7 kb*	47.5 kb*
Total contig length	190 Mb	78 Mb*	54.4 Mb*
G+C content	51.0%	54.4%	53.9%
Repeat† (%)	74%	39%	28%
Collinear blocks‡	85 Mb	52 Mb	37 Mb
Repeat† (%) in collinear blocks‡	57%	28%	13%
Repeat† (%) outside collinear blocks‡	86%	60%	56%
Intergenic region spacing in collinear blocks: (25–75 percentiles)	224-3,070 bp	307-2,319 bp	270-1,551 bp
Intergenic region spacing outside collinear blocks‡ (25–75 percentiles)	664–19,144 bp	753-5,896 bp	566-4,351 bp
Genes			
Number of genes§	17,797	16,988	14,451
Phytophthora orthologues	11,893	12,427	12,136
Phytophthora core orthologues	9,583	9,550	9,664

* Statistics derived from supplementary materials accompanying ref. 5.

† Measured by RepeatMasker with *de novo* RepeatScout libraries (see Supplementary Information).

‡ Union of collinear blocks derived from pairwise genome comparisons (see Supplementary Information).

§ P. sojae and P. ramorum annotations were obtained from JGI (see Supplementary Information). Newly discovered mobile elements were removed, and specific gene families of interest were

reannotated (see Supplementary Information).

Core orthologous groups contain at least one orthologous gene from each of the three Phytophthora species.



Figure 1 Repeat-driven genome expansion in *Phytophthora infestans.* Conserved gene order across three homologous *Phytophthora* scaffolds. Genome expansion is evident in regions of conserved gene order, a

increased numbers of mobile elements across diverse families as compared to *P. sojae* and *P. ramorum*, with \sim 5 times as many LTR retrotransposons and \sim 10 times as many helitrons (Supplementary Fig. 7).

Consistent with a model of repeat-driven expansion of the *P. infestans* genome, the vast majority of repeat elements in the genome are highly similar to their consensus sequences, indicating a high rate of recent transposon activity (Supplementary Fig. 8). In addition, we have observed and experimentally confirmed examples of recently active elements (Supplementary Figs 9–11).

Phytophthora species, like many pathogens, secrete effector proteins that alter host physiology and facilitate colonization. The genome of *P. infestans* revealed large complex families of effector genes encoding secreted proteins that are implicated in pathogenesis¹⁰. These fall into two broad categories: apoplastic effectors that accumulate in the plant intercellular space (apoplast) and cytoplasmic effectors that are translocated directly into the plant cell by a specialized infection structure called the haustorium¹¹. Apoplastic effectors include secreted hydrolytic enzymes such as proteases, lipases and glycosylases that probably degrade plant tissue; enzyme inhibitors to protect against host defence enzymes; and necrotizing toxins such as the Nep1-like proteins (NLPs) and PcF-like small cysteine-rich proteins (SCRs) (Supplementary Table 6).

As in the other *Phytophthora* species⁵, candidate effector genes are numerous and typically expanded compared to non-pathogenic relatives (Supplementary Table 6). Most notable among these are the RXLR and Crinkler (CRN) cytoplasmic effectors, described later.

The archetypal oomycete cytoplasmic effectors are the secreted and host-translocated RXLR proteins¹². All oomycete avirulence genes (encoding products recognized by plant hosts and resulting in host immunity) discovered so far encode RXLR effectors, modular secreted proteins containing the amino-terminal motif Arg-X-Leu-Arg (in which X represents any amino acid) that defines a domain required for delivery inside plant cells¹¹, followed by diverse, rapidly evolving carboxy-terminal effector domains^{13,14}. Several of these C termini have been shown to exhibit virulence activities as host cell death suppressors^{15,16}. We exploited the known motifs and other conserved sequence features to predict 563 RXLR genes in the P. infestans genome (Supplementary Tables 6, 7 and Supplementary Information). RXLR genes are notably expanded in P. infestans, with $\sim 60\%$ more predicted than in *P. sojae* and *P. ramorum* (Supplementary Tables 6 and 7). We observed that 70 of these are rapidly diversifying (Supplementary Table 8). Approximately half of P. infestans RXLRs are lineage-specific, largely accounting for the expanded repertoire (Supplementary Figs 12 and 13). In contrast to the core proteome, RXLR genes show evidence of high rates of turnover with only 16 of the 563 genes with 1:1:1 orthology relationships (Supplementary Table 2) and many (88) putative RXLR

consequence of repeat expansion in intergenic regions. Genes are shown as turquoise boxes, repeats as black boxes. Collinear orthologous gene pairs are connected by pink (direct) or blue (inverted) bands.

pseudogenes (Supplementary Table 9). This high turnover in *Phytophthora* is probably driven by arms-race co-evolution with host plants^{5,13,14,17}.

RXLR effectors show extensive sequence diversity. Markov clustering (TribeMCL¹⁸) yields one large family (*P. infestans*: 85, *P. ramorum*: 75, *P. sojae*: 53) and 150 smaller families (Supplementary Fig. 14). The largest family shares a repetitive C-terminal domain structure (Supplementary Figs 15 and 16). Most families have distinct sequence homologies (Supplementary Fig. 14) and patterns of shared domains (Supplementary Fig. 17) with greater diversity than expected if all RXLR effectors were monophyletic.

In contrast to the core proteome, RXLR effector genes typically occupy a genomic environment that is gene sparse and repeat-rich (Fig. 2g and Supplementary Figs 18 and 19). The mobile elements contributing to the dynamic nature of these repetitive regions may enable recombination events resulting in the higher rates of gene gain and gene loss observed for these effectors.

CRN cytoplasmic effectors were originally identified from P. infestans transcripts encoding putative secreted peptides that elicit necrosis in planta, a characteristic of plant innate immunity¹⁹. Since their discovery, little had been learned about the CRN effector family. Analysis of the P. infestans genome sequence revealed an enormous family of 196 CRN genes of unexpected complexity and diversity (Supplementary Table 10), that is heavily expanded in P. infestans relative to P. sojae (100 CRNs) and P. ramorum (19 CRNs) (Supplementary Table 6). Like RXLRs, CRNs are modular proteins. CRNs are defined by a highly conserved N-terminal ~50-amino-acid LFLAK domain (Supplementary Fig. 20) and an adjacent diversified DWL domain (Fig. 3a, b). Most (60%) possess a predicted signal peptide. Those lacking predicted signal peptides are typically found in CRN families containing members with secretion signals (Supplementary Table 10). CRN C-terminal regions exhibit a wide variety of domain structures, with 36 conserved domains and a further eight unique C termini identified among the 315 Phytophthora CRN proteins (Supplementary Table 11). We observed evidence of recombination between different clades as a mechanism driving CRN diversity (Supplementary Figs 21-23).

We explored the ability of diverse CRNs to perturb host cellular processes. In assays for necrosis *in planta* (Supplementary Information), deletion mutants of the previously described CRN2 secreted protein¹⁹ defined a C-terminal 234 amino-acid region (positions 173–407, domain DXZ) that is sufficient to induce cell death when expressed inside plant cells (Supplementary Fig. 24). Assays with representative *P. infestans* CRN genes identified four other distinct C termini that also trigger cell death inside plant cells (Fig. 3c). These include the newly defined DC domain (*P. infestans*: 18 genes and 49 pseudogenes (ψ)) and the D2 (14 and 43 ψ) and DBF (2 and 1 ψ) domains, which have similarity to protein kinases (Supplementary Table 11). These results indicate that the CRN protein domains



Figure 2 | **The** *P. infestans* **genome shows an unusual distribution of intergenic region lengths.** The flanking distance between neighbouring genes provides a measurement of local gene density. *P. infestans* genes were sorted into two dimensional bins on the basis of the lengths of flanking intergenic distances to neighbouring genes at their 5' and 3' ends. **a**–**h**, The number of genes in each bin is shown as a colour-coded heat map on orthogonal projection. *P. infestans* whole-genome analysis (**a**) shows most genes with intergenic regions between 20-bp and 3-kb long, as well as sets of genes flanked by one or two intergenic region(s) between 5 kb and 36 kb.

expressed *in planta* are retained (lacking signal peptides and hence not secreted) by the plant cell and stimulate cell death by an intracellular mechanism, supporting the view that CRNs, like RXLRs, are cytoplasmic effectors. We propose that the conserved CRN N-terminal LFLAK domain may function similarly to the RXLR motif for delivery of CRN effectors into plant cells, and experiments to test this hypothesis are under way.

A further 255 CRN genes are fragmented or otherwise disrupted and presumably non-functional (Supplementary Table 10). CRN genes and pseudogenes are aggregated in large clusters at several genomic loci, typically clustered by domain type (Supplementary Fig. 25). One extraordinary example is scaffold 1.48 (\sim 1.2 Mb), containing 21 CRN genes and 31 CRN pseudogenes of the DXZ and D2 necrosis inducing domain-types (Fig. 3d). Many of the pseudogenes show only a few base changes, indicating recent conversion to pseudogenes. This high degree of expansion and pseudogene formation

Comparison with other *Phytophthora* genomes (\mathbf{b}, \mathbf{c}) indicates that this separation is observed in *P. infestans* but not the other two sequenced genomes. Genes in collinear blocks (\mathbf{d}) and the core orthologue clusters (\mathbf{e}) have primarily shorter intergenic distances, whereas genes outside of collinear blocks (\mathbf{f}) reside mostly in gene sparse regions. Genes belonging to the RXLR (\mathbf{g}) and Crinkler (CRN) (genes and pseudogenes) (\mathbf{h}) effector families have flanking intergenic distances among the longest. Genes found at the ends of scaffolds and hence lacking neighbouring genes were necessarily excluded.

suggests that, like RXLR effector genes, CRN genes have undergone relatively rapid birth and death evolution.

Both CRN and RXLR genes typically occur in repeat-rich, genesparse regions of the genome, where conserved gene order with *P. sojae* and *P. ramorum* is either absent or disrupted (Fig. 2g, h and Supplementary Fig. 19). Expansion of large RXLR and CRN effector gene families seems to have been driven by non-allelic homologous recombination and tandem gene duplication. Although the genome is heavily populated by mobile elements, no direct evidence of transposition of effector genes was observed. Instead, the repeat-rich regions of effector clusters probably facilitate non-allelic-homologousrecombination-based expansion. In one intriguing case, nearly identical tandem arrays of CRNs are present on scaffold 1.6 in a perfect head-to-tail arrangement that is similar to that observed for some helitrons (Supplementary Fig. 26). This region of the genome is heavily enriched for helitron elements, implicating helitron-based rolling



Figure 3 | Diverse Crinkler (CRN) families exhibit necrosis phenotypes in *planta.* a, CRN family phylogeny on the basis of the conserved N-terminal sequence, computed using PhyML with default parameters and 100 bootstrap replicates. CRN C-terminal domain structures are shown along the circumference. Branches are coloured according to organism: *P. infestans* in blue, *P. sojae* in yellow, and *P. ramorum* in red. Internal nodes with $\geq 80\%$ bootstrap support are marked with a black dot. **b**, Graphical representation of the CRN family domain architecture, exhibiting a conserved N-terminal region followed by diverse C-terminal domains. **c**, Phenotypes observed on *Nicotiana benthamiana* leaves upon *in planta* overexpression of CRN effectors. C-terminal effector domains of CRNs were

circle replication as a possible mechanism for establishing this CRN cluster.

To explore transcriptional responses to plant infection, we constructed a NimbleGen microarray based on the genome annotation. *P. infestans* gene expression during potato infection was monitored using samples from infected potato at 2–5 days post-inoculation (d.p.i.). In all, 494 genes were induced at least twofold during infection relative to mycelial growth. Days 2–4 of infection correlate with formation of infectious structures called haustoria. Mycelial necrotrophic growth on dead plant material occurs later at 5 d.p.i., and shows a similar expression profile to mycelial growth in plant extract media (Supplementary Fig. 27a and Supplementary Table 12). Seventy-nine RXLR genes exhibited this pattern of expression, including previously studied avirulence genes *Avr3a* (ref. 20), *Avr4* (ref. 21), and *Avr-blb1* (also known as *ipiO*) (ref. 22) (Supplementary Fig. 27b). Apoplastic tested for cell death phenotypes on *N. benthamiana* leaves by *Agrobacterium tumefaciens*-mediated transient expression of CRNs, *inf1* (positive control), *crn2* (positive control), and green fluorescent protein (GFP) (negative control). The domains DC, DBF, D2 and DXW-DXX-DXS, like the DXZ domain of *crn2*, were found to induce necrosis. Cell death phenotypes were visible at 4 days post infiltration. Photos were taken 7 days after infiltration. **d**, CRNs with necrosis domains D2 and DXZ along with pseudogene copies are found co-clustered across *P. infestans* scaffold 1.48 (~1.2 Mb). Genes and domain structures are illustrated according to the top and bottom strands of the genomic scaffold. Pseudogenes are indicated by Ψ ; non-CRN genes are shown as unfilled boxes.

effector genes, including protease inhibitors, cysteine-rich secreted proteins, and NPP1-family members, were among the most highly upregulated genes during infection of potato. Few CRNs were induced during infection; however most CRNs were very highly expressed, with ~50% of CRNs within the top 10% of gene expression intensities (Supplementary Fig. 28). Several genes encoding metabolic enzymes were upregulated *in planta* (Supplementary Table 12), suggesting considerable metabolic adaptation of the pathogen to the host environment²³. A related pattern of downregulation mirrors the induction of effectors, involving ~115 genes (Supplementary Table 12). Among those repressed were elicitin-like genes and pseudogenes, suggesting that reduced expression during infection or mutation to pseudogene could contribute to evading activation of host innate immunity²⁴.

P. infestans remains a critical threat to world food security, and the genome sequence is a key tool to understanding its pathogenic success.

The sequence of the P. infestans genome showed an extremely high repeat content (~74%) and unusual discontinuous distribution of gene density that correlate intriguingly with its biology. Gene-dense regions with conserved gene order across Phytophthora species are interrupted by repeat-rich expanded regions that are sparsely populated with genes, many of which are fast-evolving pathogenicity effectors such as the RXLR and CRN families. The localization of the effectors to dynamic regions of the genome probably both enables the rapid evolutionary changes and accounts for the considerable expansion in CRN and RXLR effector genes observed in P. infestans. This expansion provides a species-specific repertoire of effector genes, the dynamic nature of which probably provides an advantage in the arms race with host species. We postulate that these dynamic regions promote the evolutionary plasticity of effector genes, generating the enhanced genetic variation required to drive the rapid evasion of plant resistance that is a hallmark of the potato late blight pathogen.

METHODS SUMMARY

Genomic sequence and gene annotations. The updated *P. infestans* genome sequence and annotation can be accessed through GenBank accession number AATU01000000, and are available through the Broad Institute website at http:// www.broad.mit.edu/annotation/genome/phytophthora_infestans. All genome sequence reads have been deposited in the NCBI trace repository (http:// www.ncbi.nlm.nih.gov/Traces/home/). Paired reads of *P. infestans* cDNAs are available in dbEST with accessions in the range GR284383–GR301386. The NimbleGen microarray data are available in GEO under accession number GSE14480. Full methods description and associated references are provided as Supplementary Information.

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Supplementary Information is linked to the online version of the paper at www.nature.com/nature.

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