Plant cell surface pattern recognition receptors (PRRs) and intracellular immune receptors cooperate to provide robust resistance to microbial infection. Both receptor families have coevolved at an accelerated rate, but the evolution and diversification of PRRs is poorly understood. We have isolated potato surface receptor Pep-13 receptor unit (PERU) that senses Pep-13, a conserved immunogenic peptide pattern from plant pathogenic Phytophthora species. PERU, a leucine-rich repeat receptor kinase, is a bona fide PRR that binds Pep-13 and enhances immunity to Phytophthora infestans infection. Diversification in ligand binding specificities of PERU can be traced to sympatric wild tuber-bearing Solanum populations in the Central Andes. Our study reveals the evolution of cell surface immune receptor alleles in wild potato populations that recognize ligand variants not recognized by others.

Plant cell surface pattern recognition receptors (PRRs) and intracellular immune receptors cooperate to provide robust resistance to microbial infection (1–2). The synergistic activation of plant immunity by spatially separated plant immune receptors suggests their coevolution. A strong correlation in the number of genes encoding surface and intracellular immune receptors observed across the plant lineage supports the concept of mutual potentiation of immune responses initiated in different plant cell compartments (4, 5).

It is assumed that pathogen pressure in ecological niches drives plant immune receptor evolution (5, 6), but evidence for diversification of plant PRR sequences and functions among natural plant populations is lacking. We hypothesized that a PRR might recognize Pep-13, a conserved microbial immunogenic 13-amino acid fragment from a cell wall glycoprotein (GP42) with transglutaminase (TG) activity (7–9). TGs are produced by several plant-pathogenic oomycete Phytophthora species, including P. infestans, the causal agent of potato late blight disease and the Great Irish Famine (10–12). Pep-13 triggers a hypersensitive response and other immunity-associated responses in diverse plant species, including the solanaceous host plant, potato (10).

**Potato PRR senses oomycete-derived pattern Pep-13**

We screened a collection of wild Solanum species and cultivated potato genotypes for cell death induction when infiltrated with Pep-13 or its structural derivative, Pep-25 (7, 13). To identify the Pep-13-receptor by a map-based cloning approach, we crossed genotype Solanum tuberosum Group Phureja DM 1-3 516 R44 (DM) and genotype S. tuberosum RH89-039-16 (RH) (fig. 1A). DM is a Pep-13/Pep-25–sensitive genotype, which was previously used to establish the potato reference genome (14). RH is a Pep-13/Pep-25–insensitive genotype. We back-crossed the F1 generation 3240-4 to the RH parent and used the resulting F2 population (3648) for genetic mapping (15). Pattern sensitivity segregated in a 1:1 ratio, suggesting that a single, dominant gene encodes the corresponding receptor (fig. S1A). Pep-13/Pep-25 sensitivity was previously mapped to the top of chromosome 3 (13), and subsequent marker-assisted fine-mapping yielded a 55.2-kb fragment containing 7 open reading frames, three of which encode leucine-rich repeat receptor kinases (LRR-RKs a-c) (fig. S1B). LRR-RKs consist of an extracellular LRR-type domain, a transmembrane-spanning domain, and an intracellular serine/threonine protein kinase domain, which is absent in LRR receptor proteins (LRR-RRs). LRR-RR containing receptors are the predominant type of plant PRRs known to date and have evolved to recognize primarily proteinaceous microbial patterns or phytohormones (16–18).

To determine which LRR-RK candidate gene sequence confers Pep-13 sensitivity, we performed transient expression assays in the solanaceous model plant, Nicotiana benthamiana. Agrobacterium infection–mediated expression of LRR-RK b—but not LRR-RK a or c-encoded cDNA sequences—resulted in plant cell death after treatment with either Pep-13 or GP42 (Fig. 1B). We thus designated LRR-RK b Pep-13 receptor unit (PERU). PERU is a canonical plant LRR-RK that hosts an ectodomain composed of 27 LRRs linked by a transmembrane domain to an intracellular serine/threonine protein kinase domain (fig. S2). Stable expression of PERU cDNA in Pep-13–insensitive potato cultivar Atlantic resulted in Pep-13–inducible cell death, production of reactive oxygen species (ROS) and accumulation of the plant stress hormone, ethylene (fig. 1, C to E, and fig. S3). These responses were not observed in wild-type (WT) Atlantic or in control lines transformed with empty vector only. Inactivation of the PERU locus in Pep-13–sensitive DM by CRISPR-Cas9 mutagenesis provided direct proof for a causal role of PERU in Pep-13 pattern recognition. To abolish PERU gene expression, genotype DM was stably transformed with CRISPR-Cas9 and 4 sgRNA, CRISPR lines were genotyped, and deletion and frameshift mutations were found, resulting in loss of Pep-13–induced cell death, ROS burst, and ethylene production (Fig. 1, F to H). In sum, these results document a role for potato PERU in Pep-13 pattern recognition.

**A PERU-SERK3 complex mediates Pep-13/Pep-25–induced defenses and plant cell death**

LRR-type PRRs recognize their cognate ligands by binding to their LRR ectodomains (19). We investigated ligand-receptor binding in vitro and in planta. To test for physical interaction of PERU and Pep-25 in vitro, we incubated recombinant hexa-histidine (His6)–tagged PERU LRR ectodomain protein (PERULRR-His6) with biotinylated Pep-25 (Pep-25-bio) before treatment with the homo-bifunctional cross-linker ethylene glycol bis (succinimidyl succinate) (EGS) to stabilize the ligand-receptor complex (8). Pep-25-bio was as active as Pep-25 (fig. S4). Following PERU-His6 immunoprecipitation, bound Pep-25-bio was visualized by streptavidin/anti-streptavidin antisera (fig. S5). A large molar excess of free Pep-13 competitively abolished ligand-receptor complex formation, which suggests direct and specific ligand binding by PERU-His6. The affinity constant of the ligand/receptor interaction (Kd = 88.9 nM) is close to ligand concentrations required for immune activation in three P3SS:PERU-expressing Pep-13–insensitive Solanum hirtigii or in Pep-13–sensitive potato DM (EC50 = 9.8 nM or 44 nM respectively), indicating that the PERU ectodomain is sufficient for ligand binding (fig. S5). To analyze the Pep-25/bio/PERU interaction in planta, we treated leaves of N. benthamiana plants expressing green fluorescent protein (GFP)–tagged PERU (p3SS:PERU-GFP) with ligand prior to EGS treatment. Precipitation of PERU-GFP protein and subsequent detection of bound ligand corroborated

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1Plant Breeding, Wageningen University and Research, 6708 PB Wageningen, Netherlands. 2Department of Plant Biochemistry, Centre of Plant Molecular Biology (ZMBP), University of Tübingen, 72076 Tübingen, Germany. 3The Sainsbury Laboratory, University of East Anglia, Norwich, UK. 4Wageningen Food & Biobased Research, Wageningen University and Research, 6708 WG Wageningen, Netherlands. 5Department of Biochemistry, University of Johannesberg, Johannesberg 2006, South Africa.

*Corresponding authors, Email: vivianne.vleeshouwers@wur.nl (V.G.A.V.); nuernberger@uni-tuebingen.de (T.N.)

†Present address: State Key Laboratory of Plant Genomics, Institute of Microbiology, Chinese Academy of Sciences, Beijing 100101, China. ‡Present address: Department of Agronomy, Purdue University, West Lafayette, IN 47906, USA.
ligand-receptor binding observed in vitro (Fig. 2A). Again, excess of Pep-13 abolished ligand binding. We did not observe an inhibitory effect when a biologically inactive Pep-13 mutant peptide, Pep-13W231A (tryptophan residue 231 mutated to alanine, amino acid numbering corresponds to full-length GP42 sequence) (10), was used as a competitor (Fig. 2A). Notably, a W231A mutant of GP42 not only abolished its plant defense-eliciting activity but also reduced its TG activity by 98.5% (10). Altogether, these data demonstrate specific binding of Pep-13 to its high-affinity binding site, PERU.

The LRR-RK BAK1/SERK3 (BRASSINOSTEROID INSENSITIVE 1-ASSOCIATED KINASE1/SOMATIC EMBRYOGENESIS RECEPTOR KINASE 3) forms ligand-induced receptor/co-receptor complexes with plant LRR-RK-type PRRs (17, 18, 20). We found Pep-13 pattern-induced complex formation of PERU and SERK3A after transient coexpression of p35S::PERUΔ3′-GFP and p35S::SERK3AΔ3′-Myc in N. benthamiana plants (Fig. 2B). Virus-induced gene silencing of SERK3 (TRV::NhSERK3) in N. benthamiana resulted in a massive reduction in Pep-13–induced hypersensitive cell death and ROS production in TRV::NhSERK3 plants (Fig. 2C and fig. S6), suggesting that PERU recruits SERK3 in a pattern-dependent manner. Silencing of SOBIR1 (SUPPRESSOR OF BAK1-INTERACTING KINASE1) (TRV::NhSOBIR1) is exclusively required for the function of LRR-RP-type PRRs, did not

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affect Pep-13–induced cell death formation. By contrast, *Phytophthora infestans* elicitin INF1–induced cell death mediated by activation of LRR-RP-type ELR (ELICITIN RESPONSE) (21) is reduced in both TRV::NbSERK3 and TRV::NbSOBIR1/-like plants (Fig. 2C).

In solanaceous *N. benthamiana*, activation of plant immunity and cell death by LRR-RP-type PRRs requires lipase-like ENHANCED DISEASE SUSCEPTIBILITY 1 (EDS1) (22) and helper NUCLEOTIDE-BINDING LRR (hNLR) REQUIRED FOR HYPERSENSITIVE RESPONSE-ASSOCIATED CELL DEATH 2, 3, and 4 (NRC2, NRC3, NRC4) (23, 24). Because LRR-RK-type PRRs have not previously been implicated in activating plant cell death in any plant system, we tested whether these proteins are required for PERU signaling.

*N. benthamiana* plants transiently expressing p35S::PERUDM developed cell death symptoms upon infiltration of Pep-13 or GP42 (Fig. 1B) and produced ethylene upon treatment with Pep-25 (fig. S4). Transient expression of p35S::PERUDM::GFP in *N. benthamiana* mutants lacking EDS1 and related PHYTOALEXIN-DEFICIENT 4 (PAD4) (25, 26) or hNLRs NRC2/3/4 (23) had no reducing effect on elicitor–induced cell death and ethylene production (fig. S7). Altogether, we find substantial differences in the molecular mechanisms controlling plant immune responses upon activation of different classes of LRR-type PRRs in this plant.

PERU confers enhanced resistance to *P. infestans*

Potato varieties used in agricultural production are often susceptible to major plant pathogens,
experiments were performed 3 times with similar results and representative experiments are shown.

### Functional diversification of PERU in wild potato populations

Pep-13/25-induced plant defenses have been studied in parsley and potato cell suspensions, as well as in leaves of a cultivated potato clone, Désirée (7, 10). Alanine scanning mutagenesis of Pep-13 sequences revealed that mutant Pep-13W231A abolished elicitor activity, mutation of proline 234 (Pep-13P234A) reduced it, and replacement of the remaining amino acid residues (including tyrosine 241, Pep-13Y241A) did not significantly affect activities of the mutant peptides (7, 8, 10). We found the same pattern of ligand responses in the Pep-25–sensitive genotype DM, which was used for PERU identification (Figs. 1A, 3A, and data S1). To determine the frequency of biologically active PERU alleles in Solanum sect. Petota (30), we analyzed 476 genotypes corresponding to 98 species (97 wild, and 1 cultivated potato species) for cell death triggered by Pep-25 and its described mutant variants (data S1). 350 (74%) of these genotypes did not develop cell death in response to Pep-25, indicating that most wild Solanum genotypes in this collection lack an active PERU allele (data S1). Pep-25 and its mutants were tested for cell death induction on all 126 Pep-25–sensitive genotypes (Fig. 3A). Overall, we observed at least five different recognition specificities, including substantial qualitative and quantitative variations in their abilities to respond to Pep-25 and its mutants. Wild potato genotypes grouped in class 1 include DM and exhibit the same ligand response patterns as described for cultivated potato cultivar Désirée previously (7, 8, 10) (Fig. 3A). These accounted for 25% of all Pep-25–sensitive genotypes (Fig. 3A). Other genotypes showed responsiveness to Pep-13W231A but failed to respond to Pep-13P234A (class 2), others failed to respond to both Pep-13P234A and Pep-13W241A (class 3), and Pep-13/25 and Pep-25 mutant peptides yielded the same response pattern as observed in DM and LPH plants. (D) EC50 values were determined by quantification of elicitor-induced production of ethylene in DM or LPH plants, and in S. hirtii plants transiently transformed with either PERU or PERU.CGFP, nd, not determined. (E) Receptor/ligand binding assays show that PERU specifically bound both Pep-25-bio and Pep-25W231A-bio as ligands, and Pep-13W231A efficiently competes for ligand binding to PERU.CGFP. All experiments were performed 3 times with similar results and representative experiments are shown.

### Diversification of PERU ligand specificities in wild potato populations

Pep-13/25-induced plant defenses have been studied in parsley and potato cell suspensions, as well as in leaves of a cultivated potato clone, Désirée (7, 10). Alanine scanning mutagenesis of Pep-13 sequences revealed that mutant Pep-13W231A abolished elicitor activity, mutation of proline 234 (Pep-13P234A) reduced it, and replacement of the remaining amino acid residues (including tyrosine 241, Pep-13Y241A) did not significantly affect activities of the mutant peptides (7, 8, 10). We found the same pattern of ligand responses in the Pep-25–sensitive genotype DM, which was used for PERU identification (Figs. 1A, 3A, and data S1). To determine the frequency of biologically active PERU alleles in Solanum sect. Petota (30), we analyzed 476 genotypes corresponding to 98 species (97 wild, and 1 cultivated potato species) for cell death triggered by Pep-25 and its described mutant variants (data S1). 350 (74%) of these genotypes did not develop cell death in response to Pep-25, indicating that most wild Solanum genotypes in this collection lack an active PERU allele (data S1). Pep-25 and its mutants were tested for cell death induction on all 126 Pep-25–sensitive genotypes (Fig. 3A). Overall, we observed at least five different recognition specificities, including substantial qualitative and quantitative variations in their abilities to respond to Pep-25 and its mutants. Wild potato genotypes grouped in class 1 include DM and exhibit the same ligand response patterns as described for cultivated potato cultivar Désirée previously (7, 8, 10) (Fig. 3A). These accounted for 25% of all Pep-25–sensitive genotypes (Fig. 3A). Other genotypes showed responsiveness to Pep-13W231A but failed to respond to Pep-13P234A (class 2), others failed to respond to both Pep-13P234A and Pep-13W241A (class 3), and Pep-13/25 and Pep-25 mutant peptides yielded the same response pattern as observed in DM and LPH plants. (D) EC50 values were determined by quantification of elicitor-induced production of ethylene in DM or LPH plants, and in S. hirtii plants transiently transformed with either PERU or PERU.CGFP, nd, not determined. (E) Receptor/ligand binding assays show that PERU specifically bound both Pep-25-bio and Pep-25W231A-bio as ligands, and Pep-13W231A efficiently competes for ligand binding to PERU.CGFP. All experiments were performed 3 times with similar results and representative experiments are shown.

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and some genotypes failed to respond to all mutant peptides of Pep13 (class 4). Some Solanum genotypes, such as Solanum leptophyes (LPH) 680-5, exhibit sensitivities to all Pep-25 variants tested and were categorized as class 5 genotypes (Fig. 3A). Altogether, our findings reveal that wild Solanum species bear diverse PERU alleles that differ from PERU<sup>2DM</sup> and thus encode PRRs with distinct ligand specificities.

We assessed plant defense and cell death–inducing activities of Pep-25 WT and mutant peptides in potato genotypes expressing PERU<sup>2DM</sup> (class 1 genotype) or PERU<sup>4DM</sup> (class 5 genotype) (Fig. 3B) in greater detail. We found that PERU<sup>4DM</sup>-expressing plants recognized all Pep-25 variants, whereas PERU<sup>2DM</sup> did not mount cell death in response to Pep-25W231A. To corroborate these findings and to rule out that potato

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**Fig. 4. Phylogenetics and geographic distribution of PERU.** (A) Heatmap representation of presence/absence of PERU homologs across Solanum species. The PERU<sup>2DM</sup>, PERU<sup>4DM</sup>, and the Pep-13 responsive homologs grouped together in a single clade whereas the nonresponsive homologs were distributed throughout the phylogenetic tree. Bootstrap values are shown for the clade containing all tested homologs (0.87) and the clade containing the responsive PERU<sup>2DM</sup> homologs (0.98). The Solanum phylogeny was adapted from (48). (B) Kernel density distribution map of 266 genotypes insensitive to Pep-25 that are distributed from the Southern USA to Northern Chile; and (C) Kernel density distribution map of 98 sensitive genotypes that cluster in Peru and Bolivia. Red shades indicate high density, yellow shades indicate lower density, the blue dots represent individual geo-coordinates of accessions, and bar plots represent the number of genotypes along different latitudes of the continent. Available geographic coordinates of 364 genotypes (data S1) were used to elaborate the maps.
genotype-specific properties account for altered PERU ligand binding specificities, we transiently expressed PERU<sup>DM</sup> or PERU<sup>DPH</sup>-encoding sequences in Pep-13-insensitive <i>S. hjertingii</i>. Again, infiltration of Pep-25 WT and mutant peptides yielded the same response pattern as observed previously (Fig. 3B), with all Pep-25 mutants inducing cell death in <i>PERU<sup>W231</sup></i> plants only (Fig. 3C). Thus, differences in ligand specificities of PERU proteins from <i>PERU<sup>DM</sup></i> or <i>PERU<sup>DPH</sup></i>-expressing plants are features of the receptor proteins themselves rather than of co-receptors or other auxiliary factors.

We further analyzed the biological activities of Pep-25 and its mutants by quantifying elicitor-induced production of the stress hormone ethylene in <i>PERU<sup>DM</sup></i> or <i>PERU<sup>DPH</sup></i>-expressing potato plants and in <i>S. hjertingii</i> plants transiently transformed with either PERU allele (Fig. 3, D and E). Determination of elicitor concentrations required to induce half-maximal ethylene production required to induce half-maximal ethylene production (EC<sub>50</sub>) corroborated qualitative data from cell death assays (Fig. 3, B and C). Pep-25W231A proved as active as Pep-25 only when tested on <i>PERU<sup>W231</sup></i>-expressing plants. We found substantially reduced or no activity of this peptide in plants expressing <i>PERU<sup>DM</sup></i> (Fig. 3, D and E). In agreement with that, <i>PERU<sup>W231</sup></i> plants bound both Pep-25-bio and Pep-25W231A-bio in receptor-ligand binding assays (Fig. 3E). Likewise, Pep-13W231A efficiently blocked ligand binding to <i>PERU<sup>W231</sup></i> (Fig. 3E) but not to <i>PERU<sup>DM</sup></i> (Fig. 2A). Altogether, our data obtained from ligand binding assays and from plant defense activation studies confirm that <i>PERU<sup>DM</sup></i> and <i>PERU<sup>DPH</sup></i> encode related LRR-RK immune receptors that have diversified in ligand specificities. Our findings further suggest that functional diversification has occurred within this immune receptor family during evolution, resulting in PERU alleles that recognize Pep-13 variants not recognized by others.

**Evolutionary history of PERU**

To obtain information about the origin of PERU alleles, we studied their geographic distribution and genetic variation (Fig. 4). We developed a computational pipeline to extract PERU sequences from 6,630,292 predicted proteins from 124 Solanaceae genome assemblies and for comparison extracted sequences of FLS2 (FLAGELLIN-SENSING 2), a conserved LRR-RK that detects bacterial flagellin (fig. S8 and data S2) (3). Both PERU and FLS2 clustered in well-supported clades within the LRR-RK subgroup XII indicating a monophyletic origin within the Solanaceae (fig. S9) (5). All plant LRR-RK-type PRRs currently known fall into this clade, including <i>Arabidopsis</i> FLS2 and EF-TU RECEPTOR (EFR), or rice Xa21 (16, 31). This pipeline yielded 114 PERU clade sequences from 17 species and 180 FLS2 clade sequences from 33 species with the PERU clade sequences exhibiting markedly more diversity than the FLS2 clade (fig. S9 and data S3, S4, and S5). The same primers employed to isolate PERU (DM) facilitated the isolation of 26 responsive homologs and 25 nonresponsive homologs (data S6). The amplified and genome-extracted sequence datasets were then combined for phylogenomic analyses, which revealed that the PERU sequences encoding Pep-13-responsive PERU alleles fall into one clade whereas the nonresponsive homologs are scattered throughout the tree (Fig. 4A and fig. S10, A and B). The PERU<sup>DPH</sup> sequences are embedded within a tighter PERU<sup>DPH</sup> clade indicating that evolution of a new ligand specificity and, hence, functional diversification has occurred within the PERU receptor family of <i>Solanum</i> (Fig. 4A and fig. S10A). Our phylogenomics analyses of PERU sequences further suggest that potato PERU<sup>DM</sup> and the PEP-13 receptor in <i>Solanum</i> are distinct proteins, although they share similar ligand specificities (Fig. 3) (7, 10).

Metazoan and plant immune receptors have been targeted by positive, diversifying selection, which accelerates the divergence between homologous proteins (32, 33). To identify amino acids under diversifying selection in the proteins encoded by PERU alleles, we applied maximum likelihood models of codon substitution using the program codeML from PAML (34, 35). We found a total of 11 residues (S118, E172, L194, Q208, R245, E339, E691, L302, A414, Q469, R590) to be under positive selection according to the three models tested (table S1). We further used AlphaFold2 to predict the tertiary structure of the PERU<sup>DM</sup> ectodomain (fig. S11). All residues but one (S118) found to be under positive selection are located on the concave side of the LRR structure, consistent with observations made for binding of the bacterial flagellin epitope flg22 to the <i>Arabidopsis</i> LRR-RK FLS2 (36). As observed for other immune receptors, diversifying selection may have driven functional diversification of PERU receptors in wild potato populations.

We further observed that Pep-25-insensitive genotypes were found across the American continent ranging from the US to Chile and Argentina (Fig. 4B). By contrast, Pep-25 sensitivity clustered among species belonging to the section Tuberosa or Piurana—which thrive predominantly in the Andean region of Bolivia and Peru (Fig. 4C)—which suggests that the PERU receptor family arose in this region. Wild potatoes carrying PERU<sup>DPH</sup> alleles also cluster in this region, suggesting that functional diversification of PERU alleles in wild potato populations has occurred at its center of origin. PERU alleles from multiple potato cultivars used today for crop production all cluster with PERU<sup>DM</sup> (Fig. 4A and fig. S10A), suggesting that PERU has been maintained during domestication (43).

**Discussion**

In this study, we characterize potato PERU as a bona fide plant PRR conferring <i>P. infestans</i> recognition. PERU binds Pep-13/25 patterns that are conserved among species of the genus <i>Phytophthora</i>, hetero-dimerizes with BAK1 in a ligand-dependent manner, mediates activation of plant immunity, and increases resistance to a devastating potato disease.

Different ligand response specificities observed among wild <i>Solanum</i> accessions indicate that functional diversification within this PRR family has occurred at the site of origin of the predominant allele, PERU<sup>DM</sup>. The explicit use of wild potato populations instead of plant materials that have undergone substantial genetic rearrangements during crop breeding implies that natural forces have been major drivers of immune receptor diversification. The Pep-13 pattern is widespread and highly conserved among plant-associated oomycetes (10), a trait that has likely facilitated the evolution of plant PRRs that recognize it. Although residue W231 is invariant in known sequences of <i>Phytophthora</i> TGS, polymorphisms affecting the elicitor activity of WT Pep-13 might occur as pathogen pressure in defined ecological niches is assumed to shape immune receptor reservoirs in metazoa and plants (5). It is thus conceivable that functional diversification of PERU was driven by escape mutations within <i>Phytophthora</i> Pep-13 patterns that enable Pep-13-producing pathogen strains to elude recognition by the predominant allele, PERU<sup>DM</sup>. Microbial evasion strategies to avoid plant immune activation encompass alterations within immunogenic patterns, thus disabling their recognition by plant PRRs (37, 38). In turn, individual plant species have evolved to perceive polymorphic patterns or, alternatively, structurally unrelated immunogenic molecules (39–43). Likewise, phylogenetically distinct PRRs have evolved in different plant species to recognize structurally unrelated epitopes within individual microbial patterns (44, 45).

We report here the identification of a potato cell surface PRR from the Central Andes and its highlights PRR diversification in sympatric, natural potato populations.

**REFERENCES AND NOTES**

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Data and materials availability: Sequences of cloned homologs were deposited in the National Center for Biotechnology Information database, www.ncbi.nlm.nih.gov (accession nos. OQ999431 to OQ999482). All other data, sequences, datasets, and scripts for the phylogenetic analyses are available in the manuscript, the supplementary material or deposited at Zenodo (46). Most of the wild Solanum accessions were in possession of WUR Plant Breeding before Nagoya rules applied, part of the material was obtained later from the Center of Genetic Resources, The Netherlands for which we refer to Access and Benefit-Sharing (https://www.absfocalpoint.nl/en/absfocalpoint.html) at their website (https://cognegis.wur.nl/).

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SUPPLEMENTARY MATERIALS

science.org/doi/10.1126/science.adg5261
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Functional diversification of a wild potato immune receptor at its center of origin

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Editor’s summary
The oomycete Phytophthora infestans devastates potato crops, most famously during the Great Irish Famine of the mid-1800s. Torres Ascurra et al. examined wild potato variants from across the Americas and identified a pattern recognition receptor called PERU, which recognizes a P. infestans peptide. When PERU binds a protein fragment from P. infestans, the potato plant can mount an immune response. The authors established that different alleles of PERU from wild Andean potato relatives have different sensitivities to the P. infestans peptide. Their work provides mechanistic insight into P. infestans immunity, thus paving the way for improved crop resilience to a disease that has been challenging to control. —Madeleine Seale

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