

# Plant NLR diversity: the known unknowns of pan-NLRomes

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## Abstract

Plants and pathogens constantly adapt to each other. As a consequence, many members of the plant immune system, and especially the intracellular nucleotide-binding site leucine-rich repeat receptors, also known as NOD-like receptors (NLRs), are highly diversified, both among family members in the same genome, and between individuals in the same species. While this diversity has long been appreciated, its true extent has remained unknown. With pan-genome and pan-NLRome studies becoming more and more comprehensive, our knowledge of NLR sequence diversity is growing rapidly, and pan-NLRomes provide powerful platforms for assigning function to NLRs. These efforts are an important step toward the goal of comprehensively predicting from sequence alone whether an NLR provides disease resistance, and if so, to which pathogens.

# A brief history of plant NLRs

#### NLRs and disease resistance

Pathogens exert some of the strongest selection pressures on plants, and their prevalence is very dynamic in time and space. It is thus not surprising that the ecological and evolutionary forces resulting from plant–pathogen interactions play an important role in shaping both plant and pathogen genomes, including constant genetic innovation and preservation of long-standing variation on both sides.

Potential pathogens are ubiquitous, but only a few can typically colonize individuals from specific plant species. The first, most common reason for failure to colonize a potential host is that a pathogen lacks the genetic toolkit required for successful establishment and subsequent extraction of resources from this host. A second reason is that a pathogen is recognized by the host because it presents widely distributed, and often highly conserved molecules that mark it as a potential threat. These molecules have been termed

microbe- or pathogen-associated molecular patterns (MAMPs/PAMPs), and are detected by pattern recognition receptors (PRRs) that are localized in the plant membrane and that survey extracellular space [\(Jones and Dangl, 2006](#page-14-0)). Engagement of these receptors by PAMPs leads to a raft of cellular changes, collectively known as PAMP or pattern triggered immunity (PTI; [Jones and Dangl, 2006\)](#page-14-0).

To overcome this first layer of defense, a pathogen has to evade or suppress PTI, which it often does by delivering its own effector molecules into the plant cell. This in turn creates a second opportunity for the plant to detect the presence of the pathogen through intracellular receptors ([Chisholm et al., 2006\)](#page-13-0), and their effector-dependent activation leads to what is commonly called effector-triggered immunity (ETI). PTI and ETI, the two branches of the plant immune system, have often been looked at as largely independent from another, but there is increasing evidence of interdependence between PTI and ETI components, which

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enhances the robustness of the defense response ([Hatsugai](#page-14-0) [et al., 2017](#page-14-0); [Ngou et al., 2020;](#page-15-0) [Yuan et al., 2020\)](#page-17-0).

The central role of nucleotide binding (NB) site leucinerich repeat (NB-LRR, "nibbler") proteins in ETI was discovered in the mid-1990s, when several NB-LRR genes were isolated through map-based cloning or transposon tagging. The starting point was plant lines distinguished by presence or absence of disease resistances that followed Flor's genefor-gene paradigm [\(Flor, 1971](#page-13-0)), in which single pathogen genes trigger a strong defense response in hosts carrying the matching resistance genes [\(Mindrinos et al., 1994](#page-15-0); [Bent](#page-13-0) [et al., 1994](#page-13-0); [Whitham et al., 1994\)](#page-17-0). Such pathogen genes, which are typically found only in some races of a pathogen species, were originally called avirulence genes because their detection by the host makes the pathogen avirulent. Similarly, the host species is polymorphic for the presence of the corresponding resistance genes, resulting in variable gene-for-gene interactions in which a specific group of pathogen strains is recognized only by a subset of similarly dis-tinct host strains [\(Botella et al., 1998](#page-13-0); [Ellis et al., 1999](#page-13-0)).

Cloning of these resistance or R genes revealed several salient facts: (i) Many code for what we now call NLR (instead of NB-LRR) proteins, to emphasize their similarity to animal NOD-like receptor (NLR) immune proteins; (ii) the functional absence of a resistance gene in some cases translates into absence of the entire gene, but in other cases, into the presence of functionally diverged orthologs; and (iii) NLR genes are often organized in clusters that differ greatly between different host strains. Here, we will first describe what has been learned in the past 25 years about inter- and intraspecific NLR diversity. We will then review how much (or how little) progress has been made toward unraveling the known unknowns of intraspecific NLR diversity through pan-genome and pan-NLRome studies, and conclude by discussing challenges for the years ahead.

## NLR genes and NLR complexes are (almost) everywhere

NLR proteins are found in plants, animals, fungi, and protists, although the similarities in protein architecture are thought to result from convergent evolution ([Yue et al.,](#page-17-0) [2012](#page-17-0)). Generally, the basic NLR unit follows a modular tripartite structure. The N terminal domain of NLR proteins can be considered the business end, being the primary structural element of signal transduction [\(Bentham et al., 2017\)](#page-13-0). In plants, it is usually either a Toll/Interleukin-1 receptor/ Resistance protein (TIR) domain, a coiled-coil (CC) domain, or a RESISTANCE TO POWDERY MILDEW 8-like (RPW8 or CC-R) coiled-coil domain. The central and most conserved NLR domain is a signal-transducing ATPase (signal transduction ATPases with numerous domains superfamily, STAND) domain, which likely evolved from a common bacterial ancestor and then gave rise to two different subclades, NB-ARC, typical for plants, and NAIP, CIITA, HET-E, and TP1 (NACHT) domains, typical for animals and fungi [\(Urbach](#page-17-0) [and Ausubel, 2017](#page-17-0)). This domain is NB and serves as an ADP–ATP switch that regulates the ON/OFF state of the NLR. The C terminal domain, which is usually composed of repeated units, acts mostly as a ligand binding platform with autoinhibitory function. In plants and some animals, but not in fungi, it is often made up of leucine-rich repeats (LRRs; [Soanes and Talbot, 2010](#page-16-0); [Dyrka et al., 2014](#page-13-0)).

Plant NLRs recognize pathogen effectors by directly binding to them, or by recognizing effector-mediated modifications of another host component. In the latter scenario, the NLR acts as a guard and its host client is its guardee [\(Cesari,](#page-13-0) [2018\)](#page-13-0). Guardees are typically proteins that are targeted by effectors, with some having lost their original function in plant growth or immunity, and only acting as effector decoys ([Dangl and Jones, 2001;](#page-13-0) [van der Hoorn and Kamoun,](#page-14-0) [2008\)](#page-14-0).

Many of the inferences about NLRs came from the study of individual NLR domains and interactions between them and their guardees. The similarity in overall function between animal and plant NLRs was subsequently confirmed by structural analyses. In vertebrates, NLR proteins form wheel-shaped oligomers called inflammasomes, which are assembled upon pathogen recognition and activate a signaling cascade that leads to the formation of pores in the plasma membrane, resulting in localized cell death ([Broz and](#page-13-0) [Dixit, 2016\)](#page-13-0). In plants, the CC-NLR HOPZ-ACTIVATED RESISTANCE 1 (ZAR1) and the TIR-NLR Roq1 assemble into oligomeric complexes called resistosomes that are strikingly similar to inflammasomes ([Wang et al., 2019a,](#page-17-0) [2019b;](#page-17-0) [Martin](#page-15-0) [et al., 2020\)](#page-15-0). Elucidation of the ZAR1 structure also revealed that its CC domain is reminiscent of bacterial pore-forming toxins. Since ZAR1 localizes to the plasma membrane, it is tempting to speculate that the oligomerized CC domains directly insert into the membrane, creating pores and thereby potentially changing ion flow and triggering cell death [\(Wang et al., 2019a](#page-17-0)). In further support of a common cell death-inducing mechanism across kingdoms is the observation that the CC-R (RPW8) domain can be homology modeled on structures of the mammalian MLKL and the fungal HELo and HELL domains, all of which also form membrane pores and induce cell death [\(Seuring et al., 2012](#page-16-0); [Daskalov](#page-13-0) [et al., 2016;](#page-13-0) [Bentham et al., 2018\)](#page-13-0). However, it must be noted that while the idea that CC-NLRs, TIR-NLRs, and RPW8-NLRs can directly make holes in plasma membranes and thereby initiate further cell damage-associated signaling and/or cell death is attractive, it is unclear whether a onesize-fits-all model applies, since plant NLRs can apparently act in different cellular compartments, including the nucleus and the cytoplasm ( $Z$ hang et al., 2017a), with resistosomes potentially targeting diverse membranes ([Adachi et al.,](#page-12-0) [2019\)](#page-12-0).

If NLRs in plants, animals, and fungi are the product of convergent evolution, what is the advantage of having this particular multi-domain structure and mode of action? Immune receptors should ideally act as hair triggers, such that any threat is immediately met, but at the same time, they should also be robust to inadvertent activation, since

inappropriate immunity can have devastating consequences [\(Bomblies et al., 2007](#page-13-0); [van Wersch et al., 2016\)](#page-17-0). The multidomain structure allows for self-inhibition through intramolecular interactions, providing a primary safeguard against spurious activation. The formation of higher order complexes in turn may serve to amplify the triggering signal, but also help to prevent mis-regulation. Lastly, the modular NLR architecture may allow for facile reshuffling of individual domains, endowing them with versatility in recognition specificity, as well as allowing for different selection pressures to act on individual domains.

## NLR gene numbers vary greatly between plant species

After plants colonized land, plant genomes experienced a massive expansion of NLR genes, going from fewer than a dozen in green algae, where plant NLRs are thought to originate, to many hundreds in land plants, likely as a conse-quence of adaptation to new pathogen pressures [\(Shao](#page-16-0) [et al., 2019](#page-16-0); [Figure 1\)](#page-3-0). NLR numbers across different species are highly variable; among all coding genes, the percentage of NLR genes ranges over many orders of magnitude, from 0.003% in bladderwort (Utricularia gibba) to 2% in apple (Malus domestica; lia et al., 2015; [Baggs et al., 2020\)](#page-12-0). NLR genes appear to turn over rapidly, with frequent births and deaths [\(Michelmore and Meyers, 1998](#page-15-0)). Many of the NLR expansion and contraction events are lineage-specific; for example, TIR-NLRs, although present in some mosses and green algae, are largely absent from monocots ([Tarr and](#page-16-0) [Alexander, 2009](#page-16-0); [Gao et al., 2018](#page-14-0)), and in the Solanaceae, CC-NLRs are greatly expanded ([Seo et al., 2016](#page-16-0)), while Rosales and conifers have experienced independent expan-sions of RPW8-NLRs [\(Jia et al., 2015](#page-14-0); [Van Ghelder et al.,](#page-17-0) [2019;](#page-17-0) [Figure 1\)](#page-3-0). NLR numbers are generally low in the Cucurbitaceae, likely as a result of frequent gene losses and few subsequent duplication events ([Lin et al., 2013;](#page-15-0) [Figure 1\)](#page-3-0). Low NLR numbers may result from their functional dispensability; for example, Wolffia australiana, a duckweed with just over 15,000 genes that potentially represent a minimum set of genes necessary for survival in an aquatic environment, has only one canonical NLR [\(Michael](#page-15-0) [et al., 2020](#page-15-0); [Figure 1\)](#page-3-0). Bladderwort, another aquatic plant, has at most one, and perhaps no NLRs at all [\(Baggs et al.,](#page-12-0) [2020\)](#page-12-0). This raises the question of whether evolutionary innovations were required in these plants to compensate for the loss of NLRs. In support of such a scenario, another duckweed with a highly reduced NLR complement, Spirodela polyrhiza, appears to have more components for antimicrobial signaling than other plants ([An et al., 2019\)](#page-12-0).

## Diversity in NLR genomic organization: from single genes to clusters, pairs, and extras

A clustered genomic arrangement is often emphasized as an NLR characteristic. In Arabidopsis thaliana genomes for example, about half of NLR genes are found in clusters [\(Meyers et al., 2003;](#page-15-0) [Van de Weyer et al., 2019](#page-17-0)). NLR clusters often appear to be the products of tandem duplication events, sometimes followed by unequal crossing over, as well as intra-cluster rearrangements and gene conversion events ([Meyers et al., 1998](#page-15-0); Noël et al., 1999; [Kuang et al.,](#page-14-0) [2004](#page-14-0); [Figure 2\)](#page-4-0). Clusters, which are often, but not always, made up of phylogenetically related NLR genes, can range in size from tens of kilobyte, with RPP5 in A. thaliana, which contains five cluster members in the reference accession Col-0, being an example, to several megabytes, with RGC2 in lettuce being a record holder with  $\sim$ 3.5 Mb and consisting of 24 cluster members [\(Meyers et al., 1998](#page-15-0), [2003](#page-15-0)).

What are the evolutionary advantages and disadvantages of having clusters? Recombination during meiosis is reduced by structural differences, which are particularly high in NLR clusters, as we discuss in more detail below. In agreement, one of the first challenges during efforts to isolate disease resistance genes was often the suppressed recombination across many such loci (to name just a few examples, see [Ganal et al. \[1989\];](#page-14-0) Noël et al. [1999]; [Wei et al. \[1999\]](#page-17-0); and [Chin et al. \[2001\]](#page-13-0)). On the other hand, there is evidence of particularly high historical recombination rates around many NLR genes, as measured by linkage disequilibrium (LD) in natural populations ([Horton et al., 2012](#page-14-0); [Choi et al., 2013](#page-13-0)), and there is only weak evidence for NLR loci as a group to suppress recombination, even for loci that differ in arrange-ment between parents [\(Rowan et al., 2019\)](#page-16-0). These apparently contradictory observations likely reflect simply the high variation in recombination rates across NLR loci, with some acting as recombination coldspots, as expected, but others acting actually as recombination hotspots ([Choi](#page-13-0) [et al., 2016](#page-13-0)). This seems to be a function of the extent of structural variation between accessions, and many of the most structurally diverse regions of the A. thaliana genome indeed include NLR clusters with severely suppressed recombination ([Jiao and Schneeberger, 2020](#page-14-0)). As we also discuss below, excessive NLR variation can potentially reduce fitness because of intragenomic immune system conflict, and under reduced recombination, it is more difficult to select for advantageous alleles that are linked to disadvantageous alleles, known as the Hill–Robertson effect ([Hill and Robertson,](#page-14-0) [1966](#page-14-0)).

Perhaps then the major advantage of clustered gene arrangements comes from several linked, closely related genes providing a means for generating new functional diversity through unequal crossovers (also known as illegitimate recombination) as well as gene conversion involving genes that are not strictly orthologs [\(Kuang et al., 2004;](#page-14-0) [Wicker et al., 2007\)](#page-17-0). This is further facilitated by the repetitive structure of an important component of NLR exons, the LRR coding sequences.

Importantly, unequal crossover can support both expansion and contraction of sequences. Contraction is perhaps particularly relevant when considering events both at singlegene loci and clusters: At single-gene loci, any deletion will lead to truncation or loss of the gene. In contrast, in a cluster, illegitimate recombination between two genes can

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Figure 1 Plant phylogeny and NLR complement sizes. Adapted from [Gao et al. \(2018\)](#page-14-0), [Munch et al. \(2018\)](#page-15-0), and [Baggs et al. \(2020\)](#page-12-0).

simultaneously reduce gene numbers and lead to creation of a new full-length gene [\(Figure 2](#page-4-0)). Although new genes resulting from unequal crossovers will often be nonfunctional, they can serve as reservoirs for future evolution. In addition, they might combine the activities of the two original genes, or they could have a different activity all together [\(Smith and Hulbert, 2005](#page-16-0); [Figure 2](#page-4-0)). Furthermore, the more copies of a gene, the higher the chances are that beneficial mutations arise, both because multiple copies constitute a larger mutational target than a single copy, but also because duplicates can undergo relaxed selection due to their functional redundancy [\(Ohno, 1970;](#page-15-0) [Jiang and Assis, 2017\)](#page-14-0). Gene clusters thus provide a larger and more flexible genetic basis for evolving new resistance specificities through complete or partial domain swaps between closely related homologs.

A special type of clustered NLR genes comprises pairs of phylogenetically unrelated genes arranged in a head-to-head orientation, where one acts as an executor and the other as a sensor, and where the latter often carries a non-canonical integrated domain (ID) that can act as a bait for pathogen effectors [\(Cesari et al., 2014\)](#page-13-0). In A. thaliana, the sensor

TIR-NLR RRS1 features an integrated WRKY domain that is targeted by Ralstonia effector PopP2 and Pseudomonas syringae effector AvrRps4, triggering the formation of an active complex composed of RRS1 and its paired executor NLR RPS4 [\(Williams et al., 2014;](#page-17-0) [Sarris et al., 2015](#page-16-0)). In rice (Oryza sativa), RGA4/RGA5 and Pik-1/Pik-2 are two wellcharacterized head-to-head gene pairs, with RGA5 and Pik-1 encoding HMA domains that are targeted by multiple Magnaporthe oryzae effectors, activating an immune response through their paired NLR partners RGA4 and Pik-2, respectively [\(Cesari et al., 2013;](#page-13-0) [Maqbool et al., 2015\)](#page-15-0). Minor mutations in IDs can have massive effects on effector recognition; for example, in the aforementioned HMA domain in Pik-1, two adjacent amino-acid changes are sufficient to expand the response profile to different M. oryzae AVR-Pik ef-fector variants [\(De la Concepcion et al., 2019](#page-13-0)). Although there is a relatively wide variety of IDs, the most prevalent domains have kinase activity, bind DNA, or mediate protein–protein interactions [\(Kroj et al., 2016](#page-14-0); [Sarris et al.,](#page-16-0) [2016;](#page-16-0) [Van de Weyer et al., 2019\)](#page-17-0). Not only are some domains more prevalent than others, it also seems that

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Figure 2 Examples of possible changes in sequence and genomic organization in single-gene and clustered NLRs through evolution. Major changes can occur in an NLR locus, and often the reconstruction of the sequence of events will be impossible.

certain NLR subfamilies are particularly prone to attract IDs, many of which are found in paired arrangements ([Bailey](#page-12-0) [et al., 2018\)](#page-12-0). Not all paired NLRs carry IDs; however, for example the TIR-NLR SOC3 is found in a head-to-head orientation with either CHS3 or TN2, both of which are truncated TIR-NLRs. Both pairs monitor SAUL1, an E3 ligase, with SOC3-TN2 responding to overaccumulation of SAUL1, and SOC3-CHS3 to the absence of SAUL1 [\(Liang et al., 2019](#page-15-0)). Generally, these paired decoy-containing NLR genes may be evolutionarily advantageous; by being linked, they facilitate co-regulation and co-evolution, leading to effective pathogen recognition and downstream signal transduction.

## NLR haplotype diversity at individual loci: the known knowns

The starting point for the cloning of many NLR loci was naturally occurring resistant and susceptible plants from the same species. An obvious step was therefore to compare the genomic regions between resistant and susceptible individuals, including the underlying NLR genes. At single-gene loci, presence/absence variation (PAV) was found to be common, with entire genes being deleted in susceptible plants. RPS5 in A. thaliana is a classical example of P/A variation at single loci [\(Henk et al., 1999](#page-14-0); [Figure 3\)](#page-5-0). The fact that susceptible plants lack the gene completely, instead of simply having functionally divergent alleles, suggested early on that in the absence of the recognized pathogen, the NLR gene imparts a fitness cost, which could indeed be demonstrated experimentally ([Tian et al., 2003;](#page-16-0) [Karasov et al., 2014\)](#page-14-0). The two allelic variants in this and other cases, either P/A, or resistant/susceptible as observed for A. thaliana RPS2 ([Figure 3](#page-5-0)), are likely maintained in metapopulations through balancing selection [\(Caicedo et al., 1999](#page-13-0); [Stahl et al., 1999;](#page-16-0) [Tian et al., 2002](#page-16-0); [Mauricio et al., 2003;](#page-15-0) [Bakker et al., 2006;](#page-12-0) [Gos et al., 2012](#page-14-0); [Rose et al., 2012](#page-16-0); [Koenig et al., 2019\)](#page-14-0).

In contrast, RPP13 in A. thaliana and the L locus in flax (Linum usitatissimum) form extensive allelic series, where different functional alleles confer disease resistance to different races of the same pathogen [\(Figure 3\)](#page-5-0). At the L locus, 10 distinct specificities for recognition of particular races of the rust fungus Melampsora lini are encoded by 11 different alleles, in both cases, recognition differences seem to be mostly due to variation in the LRR domain [\(Ellis et al., 1999;](#page-13-0) [Bittner-Eddy et al., 2000\)](#page-13-0). To date, the A. thaliana CC-NLR RPP13 remains the most polymorphic single-gene NLR locus known, with 19 different haplotypes initially identified in 24 accessions. RPP13 directly binds to the Hyaloperonospora arabidopsidis (Hpa) effector ATR13, which is also highly polymorphic, making the RPP13/ATR13 system a paradigm for NLR-effector co-evolution ([Allen et al., 2004;](#page-12-0) [Hall et al.,](#page-14-0) [2009](#page-14-0)).

These two examples of allelic series, A. thaliana RPP13 and flax L, conform relatively well to the traditional notion of alleles being functionally related ([Muller, 1932](#page-15-0)). In other cases, NLR homologs from different accessions, but in the same genomic location, confer resistance to entirely different

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Figure 3 (A) Examples of allelic variation at individual NLR loci. P/A, presence/absence. R/S, resistant/susceptible. (B) RPP4/RPP5 NLR cluster across eight different A. thaliana accessions. Tall rectangles represent NLR genes, and short rectangles non-NLR genes. Colors indicate close sequence relatedness. There is a highly variable number of blue NLR genes in the cluster (including the functionally defined members RPP4, RPP5, and SNC1), while the adjacent RLM3 gene (magenta) shows P/A polymorphism. The RPP4/RPP5 cluster has also been invaded by an unrelated NLR gene (yellow). Finally, some of the non-NLR genes (pink, green) flanking the cluster are duplicated as well. (C) Average number of genes per NLR gene cluster across eight A. thaliana accessions. Most have on average fewer than three genes, while a few are both highly expanded and highly variable in numbers. Data for B and C from [Jiao and Schneeberger \(2020\)](#page-14-0).

pathogens. In A. thaliana, RPP8, HRT, and RCY1 all are at the same genomic location. RPP8 in the Ler accession endows plants with resistance to the Emco5 isolate of Hpa, HRT in the Dijon-17 accession to turnip crinkle virus, and RCY1 in the C24 accession to the yellow strain of cucumber mosaic virus [\(McDowell et al., 1998](#page-15-0); [Cooley et al., 2000](#page-13-0); [Takahashi et al., 2002\)](#page-16-0). Because of the complex nature and copy number variation (CNV) at the small RPP8/HRT/RCY1 cluster, it is unclear whether the different resistances are encoded by different alleles or different genes at the same cluster [\(MacQueen et al., 2019](#page-15-0)). In the latter case, the situation would be similar to what has been described for the viral resistance gene Rx1 and the cyst nematode resistance gene Gpa2 in potato (Solanum tuberosum). Both genes are not only located in the same cluster, but also in the same haplotype, and it is therefore clear that they are not allelic [\(van der Vossen et al., 2000\)](#page-17-0). An unresolved question is whether despite seemingly very different organismal function, such proteins have similar biochemical activity, e.g. if they act as guards, whether they converge on the same guardee that is targeted by effectors from different pathogens.

Finally, even though NLRs are modular proteins where each domain has an assigned function, the absence of some domains does not necessarily render NLR proteins nonfunctional, with some truncated NLRs being able to confer resistance to pathogens. In A. thaliana for example, the TIRonly protein RBAI can detect the bacterial HopBA1 effector [\(Nishimura et al., 2017](#page-15-0)), while the TIR-NB protein TN13 is required for basal resistance [\(Roth et al., 2017\)](#page-16-0). RLM3, which carries a TIR-NB-X configuration with C-terminal BRX motifs instead of LRRs, confers broad-spectrum resistance to necrotrophic pathogens ([Staal et al., 2008](#page-16-0)). In wheat, YrSP, a CC-NLR that is missing most of its LRR domain, retained the ability to confer resistance to yellow rust, but had different recognition specificities from its highly similar (99.8% identity), apparent full-length variant Yr5 ([Marchal et al.,](#page-15-0) [2018\)](#page-15-0). In some cases, truncated NLR may remain functional because they work together with other full-length proteins, including NLRs, to initiate an immune response ([Zhao et al.,](#page-17-0) [2015;](#page-17-0) [Zhang et al., 2017b](#page-17-0)).

#### Collateral damage of NLR diversity

Extensive sequence diversity found across NLR genes can backfire and lead to genetic incompatibilities, often resulting in hybrid necrosis, a phenomenon formally described as early as 1943 for crosses in wheat (Triticum aestivum; [Caldwell and Compton, 1943\)](#page-13-0). Hybrid necrosis cases have since been identified in numerous plants, including rice, lettuce (Lactuca sativa and Lactuca saligna), cotton (Gossypium barbadense and Gossypium hirsutum), tomato (Lycopersicon pimpinellifolium and Lycopersicon esculentum), Capsella spp., monkeyflower (Mimulus caespitosa, M. minor, and M. tilingii), and A. thaliana (Krüger et al., 2002; [Bomblies](#page-13-0) [et al., 2007;](#page-13-0) Alcázar et al., 2009; [Jeuken et al., 2009](#page-14-0); [Yamamoto et al., 2010](#page-17-0); [Chen et al., 2014;](#page-13-0) [Chae et al., 2014](#page-13-0); [Todesco et al., 2014](#page-16-0); [Sicard et al., 2015](#page-16-0); [Deng et al., 2019](#page-13-0); [BarragAn et al., 2019;](#page-12-0) [Sandstedt et al., 2020](#page-16-0)). The genetic architecture underlying these incompatibilities is usually simple, involving one or two mismatching loci that typically encode components of the plant immune system, very often NLR proteins. These incompatibilities lead to deleterious hyperimmunity in the hybrid, thereby limiting the combinations of beneficial alleles that can be assembled in a single genotype ([Chae et al., 2014](#page-13-0)). Conversely, mismatches among pyramided NLR genes can lead to the suppression of disease resistance. In wheat for example, different Pm3 alleles can suppress each other's ability to confer powdery mildew resistance [\(Stirnweis et al., 2014](#page-16-0)), and Pm3 can also suppress resistance provided by its rye (Secale cereale) ortholog Pm8 [\(Hurni et al., 2014](#page-14-0)).

## The age of NLR pan-genomes: discovering the known unknowns

#### Why pan-genomes?

From what we have discussed so far, several important points emerge. Foremost is that there is tremendous variation in NLR genes between and within species, and particularly at NLR clusters. In recent years, it has become abundantly clear that one must not speak of "the genome" of a species. In plants, one of the earliest indications that the concept of a single linear sequence as "the genome" was misleading came indeed from comparing haplotypes of indi-vidual NLR clusters [\(Botella et al., 1998;](#page-13-0) [McDowell et al.,](#page-15-0) [1998;](#page-15-0) Noël et al., 1999; [Kuang et al., 2004;](#page-14-0) [Srichumpa et al.,](#page-16-0) [2005\)](#page-16-0). Even from studies of just a few haplotypes, we already learned that there is extensive variation at all levels: in coding sequences, in copy numbers, and in genomic location. However, while we in principle know the molecular processes that can create the observed differences (point mutations, transposon insertions, deletions, duplications, other types of chromosomal rearrangements, gene conversion, illegitimate recombination events, etc.), we still largely do not know the true extent of NLR diversity within a species, both in terms of presence and absence of individual genes nor in terms of haplotype diversity, let alone knowing the underlying evolutionary forces generating and differentially maintaining this diversity.

A decisive factor in answering these questions will be how often, if ever, a sufficient number of haplotypes has survived in a population, so that we can begin to reconstruct the evolutionary history of all, or at least of most, NLR loci. The only way to find out is by assembling more and more genomes from the same species and comparing their NLR content. While this sounds straightforward, there are substantial hurdles in making sense of many genomes due to the exponential increase in all-against-all comparisons: with only 10 different genomes, there are already 45 comparisons, but with 100 genomes, this increases to 4,950 comparisons. In order to make such genome comparisons feasible, data structures built on the concept of pan-genomes must be considered. Below we discuss the challenges of pan-genomes and of pan-NLRomes, the NLR component of pan-genomes: How to produce pan-NLRomes (discovering the unknowns), how to estimate their completeness (estimating the known unknowns), and finally, and clearly the most substantial challenge, how to make sense of the pan-NLRomes (going from known unknowns to knowns).

#### A brief history of plant pan-genomes

A common definition of a pan-genome is the entire repertoire of DNA sequences and sequence variants in a given species. That closely related bacterial strains may differ by complete arrays of genes, such as pathogenicity islands, was already noticed in the 1980s [\(Groisman and Ochman, 1996](#page-14-0)), but one of the first formal mentions of the pan-genome concept was in a study where genomes of eight different Streptococcus agalactiae strains were compared, confirming that often genomes from many individuals are needed to capture the entire gene content of a species [\(Tettelin et al.,](#page-16-0) [2005](#page-16-0)). In eukaryotes, generating many high-quality whole genome assemblies was until recently difficult due to both technical and monetary constraints associated with sequencing large genomes, especially when compared with bacteria, where most initial pan-genomic studies were performed ([Mukherjee et al., 2019](#page-15-0)). This is changing quickly, but the mapping of short reads to a single reference genome has so far been the most common method to study variation within and across species. This approach can detect small indels, single-nucleotide polymorphisms (SNPs), or larger deletions in well-conserved regions, but it often fails to adequately capture highly structurally divergent regions and CNVs.

Strictly speaking, it is very rarely possible to know the complete pan-genome of a species. Therefore, as a first step, an initial pan-genome is constructed from a few individuals, and with such a sub-pan-genome in hand, one can estimate how much of the species-wide pan-genome is still missing. Although early pan-genome studies in plants relied mostly on assembling contigs from short reads (e.g. [Gordon et al.,](#page-14-0) [2017](#page-14-0); [Golicz et al., 2016](#page-14-0); [Hurgobin et al., 2018](#page-14-0); [Zhao et al.,](#page-17-0) [2018](#page-17-0)), they already led to the discovery of many genes missing from the reference, demonstrating the importance of sequencing more than one genome to capture inter- and intraspecific genetic diversity [\(Table 1\)](#page-7-0). For example, the first Brachypodium distachyon pan-genome from 54 lines doubled the number of genes known for this species ([Gordon](#page-14-0) [et al., 2017](#page-14-0)). Similarly, in a study of 66 accessions from different rice species, around  $\sim$  10,800 novel genes were identified that were missing from the Nipponbare reference [\(Zhao](#page-17-0) [et al., 2018\)](#page-17-0).

Although short reads only allow for highly fragmented assemblies, these attempts confirmed earlier observations made by assembling left-over reads that could not be mapped to the reference [\(Cao et al., 2011;](#page-13-0) [Gan et al., 2011;](#page-14-0) [Long et al., 2013](#page-15-0)), namely that NLR genes are overrepresented in the variable fraction of gene content across accessions [\(Golicz et al., 2016;](#page-14-0) [Montenegro et al., 2017](#page-15-0); [Hurgobin](#page-14-0) [et al., 2018](#page-14-0)). For example, while only 19% of the Brassica oleracea pan-genome was composed of genes missing from the reference, the number of NLR genes missing from the reference was almost 60% [\(Golicz et al., 2016](#page-14-0)). Similarly, 50% of the 307 NLR genes found across 53 Brassica napus acces-sions were absent in the Darmor-bzh reference ([Hurgobin](#page-14-0) [et al., 2018](#page-14-0)). These findings have been confirmed with more recent and more complete long-read assemblies of multiple accessions, for example in soybean ([Liu et al., 2020b\)](#page-15-0).

In these early studies, there were generally no attempts to estimate what fraction of gene content was not properly assembled, but given the repetitive nature of many NLR genes, it is almost certain that an unknown portion of the NLR complement was missed. This was partially solved by enriching first for regions of the genome carrying NLR-related DNA sequences by hybridization to baits designed based on

## <span id="page-7-0"></span>Table 1 Examples of plant pan-genome or pan-NRLome studies and key observations on NLRs or R genes

## Plant pan-genome or pan-NLRome studies



Plant pan-genome or pan-NLRome studies					
Organism (s)	N	Sequencing technology	Assembly strategy	Reference	Key observations on NLR or R genes
Glycine clade	29	Short and long reads. Hi-C and optical map	De novo assemblies and graph	Liu et al. (2020b)	- NLRs enriched in dispensable genome.
A. thaliana	64	RenSeq: short and long reads	Map to reference	Van de Weyer et al. (2019)	- NLR diversity saturation point reached after 40 accessions. - In 64 accessions, four times higher diversity in NLR architecture than in reference.
Solanum/N. ben- thamiana/C. annum	18	RenSeg: short and long reads	Map to reference	Seong et al. (2020)	- 128 of 314 NLR annotations improved. - Preferential expansion of the extended CC-NLR N terminal region.

Table 1 Continued

known NLR sequences, followed by short read sequencing, which reduces the complexity of the assembly task [\(Jupe](#page-14-0) [et al., 2013](#page-14-0); [Andolfo et al., 2014](#page-12-0); [Arora et al., 2019](#page-12-0)). This approach, called Resistance gene enrichment sequencing (RenSeq), was later improved upon by using long-read sequencing after enrichment ([Witek et al., 2016;](#page-17-0) [Van de](#page-17-0) [Weyer et al., 2019;](#page-17-0) [Seong et al., 2020\)](#page-16-0). Most recently, as the costs for long-read sequencing have dropped precipitously, more and more sets of highly contiguous de novo assemblies are being produced for a range of species, most notably rapeseed, soybean, tomato, rice, and A. thaliana [\(Alonge](#page-12-0) [et al., 2020;](#page-12-0) [Liu et al., 2020b;](#page-15-0) [Song et al., 2020](#page-16-0); [Zhou et al.,](#page-17-0) [2020;](#page-17-0) [Jiao and Schneeberger, 2020](#page-14-0); [Table 1\)](#page-7-0).

Genes in the pan-genome can be divided by their representation frequencies across accessions. Following the convention for bacterial pan-genomes, core genes are those genes that are present in all or almost all accessions (sometimes distinguished as core and soft core), shell genes are found at intermediate frequency, and cloud genes are rare, detected only in one or very few accessions [\(Figure 4](#page-9-0)). Shell and cloud genes together make up the dispensable genome [\(Tettelin et al., 2005;](#page-16-0) [Page et al., 2015\)](#page-15-0). The number of known genes in the dispensable genome is obviously highly dependent on the number of individuals analyzed, but even in comparable data sets their contribution to the pangenome seems to vary greatly. For example, in a recent comparison of nine B. napus high-quality long-read assemblies, only 56% of genes were assigned core status, while in a comparison of eight A. thaliana genome assemblies of similar quality, 80% of genes were identified as core genes, even though the definition of core genes allowed for absence in one of the studied genomes in B. napus, whereas the A. thaliana study considered only genes present in all genomes as core genes ([Jiao and Schneeberger, 2020;](#page-14-0) [Song et al.,](#page-16-0) [2020\)](#page-16-0). How much of this is due to differences in annotation pipelines and methods for identifying orthologs are currently unclear.

#### Key steps in constructing pan-NLRomes

We are interested in the NLR component of pan-genomes, or pan-NLRomes. The first step in constructing a panNLRome for a certain species is the same as for a pangenome: the informed selection of accessions to maximize genetic diversity with as few individuals as possible ([Figure 4](#page-9-0)). Next, one has to decide on the most appropriate approach, either whole-genome assemblies after long-read sequencing or focusing on the assembly of NLR sequences with RenSeq ([Witek et al., 2016](#page-17-0); [Figure 4\)](#page-9-0). The latter, while more cost effective, has several disadvantages: Anchoring assembled NLR contigs to the genome can be difficult, since this depends on the amount of sequences flanking the assembled NLR genes, and because baits are based on known NLR genes, not all NLR genes present in a target genome will be captured equally well. Nevertheless, particularly with very large genomes, this is still the method of choice, especially if the goal is the discovery of a specific resistance gene, rather than producing a complete inventory of all NLR genes in a species [\(Arora et al., 2019\)](#page-12-0).

De novo assemblies of entire genomes will ideally contain highly contiguous scaffolds that support the facile identification of syntenic sequences. Because RenSeq produces only contigs covering individual NLR genes, the ease with which contigs can be anchored on a reference genome, or linked to larger clusters, will depend on how much flanking sequences these contigs contain, which in turn is a function of the length of the original molecules captured in the enrichment step. As an example, in a recent A. thaliana study, genes that were not clearly orthologous to reference NLR genes could not always be easily anchored to the reference genome [\(Van de Weyer et al., 2019](#page-17-0)). The size of captured molecules could potentially be improved using technologies other than hybridization-based sequence capture. One of these is CRISPR/Cas9-based excision of target regions ([Gabrieli et al., 2017;](#page-13-0) [Tsai et al., 2017](#page-17-0)), but this comes with two disadvantages: First, large amounts of very highmolecular weight DNA must be available for size separation of excised molecules from the background. Second, universally conserved regions flanking NLR genes and clusters must be known. An exciting alternative is the capture of large fragments in microdroplets, followed by droplet sorting based on PCR-based detection of NLR-related sequences with fluorescent probes [\(Madsen et al., 2020](#page-15-0)), although the

<span id="page-9-0"></span>

Figure 4 Potential workflow of the construction, visualization, and analysis of a pan-NLRome. First, accessions to be sequenced are selected in a manner that maximizes diversity, in order to approach a saturated pan-NLRome as quickly as possible (1). Then either the whole genome or the NLR complement is sequenced and assembled (2). The assemblies from each genome are then aligned to each other and potentially combined into a genome graph (3). Core and accessory genes are identified and the saturation point of the NLRome is assessed, to determine whether the NLRome is open or closed (4). Finally, individual NLR groups can be studied, e.g. one can analyze the phylogeny of a certain cluster across accessions, by comparing individual alleles in those accessions that share the same genes, the structure of whole clusters, or the entire set of genes present in all clusters (5).

design of panels of oligonucleotide primers for comprehensive NLR detection will likely be challenging.

After assembling either whole-genome shotgun or RenSeq reads, NLR genes must be annotated in each individual assembly. Automatic annotation produces a non-negligible number of errors ([Meyers et al., 2003;](#page-15-0) [Van de Weyer et al.,](#page-17-0) [2019](#page-17-0)), even when using a specialized tool such as NLR-annotator ([Steuernagel et al., 2020\)](#page-16-0). As with other annotation efforts, RNA-seq data are often very helpful, and in this regard, manipulations such as various stresses that robustly induce the expression of many NLR genes are useful [\(Lai](#page-15-0) [and Eulgem, 2018](#page-15-0); [Steuernagel et al., 2020](#page-16-0)), although manual curation of gene models is still highly recommended. We would like to emphasize, however, that we consider genes or even gene fragments that are not expressed as an important component of the pan-NLRome, because such sequences can still have a role in evolution of the pan-NLRome, as discussed above.

#### The challenges of analyzing pan-NLRomes

A major unsolved issue in pan-genome and pan-NLRome analyses is how to best compare gene sets and arrangements in multiple genomes. A first step will often be to cluster genes from the different genomes into ortholog groups for further analysis ([Enright et al., 2002](#page-13-0); [Li et al.,](#page-15-0) [2003](#page-15-0); [Contreras-Moreira et al., 2017](#page-13-0); [Thanki et al., 2018\)](#page-16-0), which will already reveal NLR genes missing from the initial reference. As discussed above, NLR genes are particularly likely to be insufficiently captured by single reference genomes.

Conventional tools for phylogenetic and population genetic analyses can easily deal with single stand-alone NLR genes. They may show P/A polymorphisms, but if a gene is present, it can in most cases be treated as an orthologous member or true allele of this particular NLR locus. A caveat arises from truncated genes, which may be functional, but might not be correctly identified as an ortholog because orthology is a function of shared sequence.

Many NLR genes are found in clusters, and orthology relationships can be complex, e.g. it may be difficult to discern which, if any, of the multiple copies of an NLR gene in one genome is the true ortholog of that same NLR gene in another genome. This is in turn important if one wants to make inferences about shared functionality and understand the evolution of NLR clusters. In this regard, it is important to remember that there is value not only in discovering new NLR genes and new NLR cluster arrangements, but also in identifying accessions with similar or identical clusters. An excellent criterion for discovering conserved clusters in a principled manner was proposed in the context of comparing eight A. thaliana genomes [\(Jiao and Schneeberger, 2020\)](#page-14-0). The authors of this study developed a formal measure of pairwise synteny diversity, inspired by the measure of pairwise nucleotide diversity,  $\pi$ . Satisfyingly, many NLR clusters were found to have high synteny diversity across accessions, but some NLR clusters were also remarkably invariant, at least in the number of genes in the cluster [\(Figure 3](#page-5-0)). 2017<br>
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Unfortunately, there will be limits to pairwise comparisons, as discussed earlier, and ultimately one would want to move to a unified data structure for multiple genomes. An obvious solution is provided by genome graphs ([Herbig](#page-14-0) [et al., 2012](#page-14-0); [Nguyen et al., 2015;](#page-15-0) [Paten et al., 2017](#page-16-0); [Computational Pan-Genomics Consortium, 2018\)](#page-13-0). In addition, powerful tools are being developed for variant calling and gene annotation on such genome graphs [\(Paten et al.,](#page-16-0)

[et al., 2019](#page-16-0)), and in principle one could envision how these tools could be extended to support phylogenetic analyses of sequences present in graph genomes [\(Figure 4](#page-9-0)).

A major challenge in understanding the evolution of NLR clusters is to reconstruct the entire sequence of events such as duplications, deletions, invasions by non-NLR genes, and phylogenetically unrelated NLR genes [\(Figure 2\)](#page-4-0), which will be contingent on how many of the individual steps in the evolution of NLR clusters are still represented in a population. Methods for reconstructing the histories of tandem duplications and more complex gene clusters, including illegitimate recombination events, have been proposed [\(Elemento et al., 2002;](#page-13-0) [Vina](#page-17-0)ř [et al., 2009\)](#page-17-0), but they are not yet widely used, most likely because the available data have been too sparse in most cases.

These issues relate to another important question: How many individuals need to be analyzed to capture a substantial fraction of a species' pan-genome? This will greatly depend on the underlying diversity of the group of accessions considered, as nicely illustrated with a recent analysis of structural variants, which include the absence/presence polymorphisms typical for many NLR genes, in tomato. Nearsaturation for common structural variants was achieved much more quickly for domesticated tomato varieties, which have undergone a recent genetic bottleneck, than for wild tomato accessions ([Alonge et al., 2020\)](#page-12-0). If additional genomes lead to the addition of fewer and fewer new genes to the pan-genome, one can infer that saturation of pangenome content is achievable, and the pan-genome is said to be closed. If the number of new genes added by each additional genome analyzed does not diminish, and if it is therefore unclear whether saturation is achievable, a pangenome is said to be open [\(Figure 4\)](#page-9-0).

The openness of bacterial pan-genomes is primarily due to horizontal gene transfer (HGT) including uptake of environmental DNA by transformation [\(Vernikos et al., 2015;](#page-17-0) [Brockhurst et al., 2019](#page-13-0)). Although HGT exists in plants and should not be dismissed as a contributor to plant pan-genomes ([Bock, 2010\)](#page-13-0), a likely more important route for expanding pan-genomes is through homoploid hybridization, that is, hybridization between distant relatives without a change in chromosome number, which is very common across all major eukaryotic lineages including plants [\(Mallet](#page-15-0) [et al., 2016;](#page-15-0) [Payseur and Rieseberg, 2016;](#page-16-0) [Taylor and Larson,](#page-16-0) [2019\)](#page-16-0). Often, the genomes from the original two species contribute very unevenly to the extant genomes we observe today, and the sequences coming from the underrepresented genome are considered as introgressed segments. Of particular interest in this context is adaptive introgression, where alien sequences are favored under natural selection because they confer some fitness benefit ([Suarez-Gonzalez](#page-16-0) [et al., 2018\)](#page-16-0). It is not surprising that there are many instances of introgressions from wild material intro crops ([Janzen](#page-14-0) [et al., 2019](#page-14-0)), and introgression through wide crosses has a long history in breeding, including breeding for disease resistance [\(Stevens and Rick, 1988](#page-16-0); [Niu et al., 2014](#page-15-0); [Gaiero](#page-14-0) [et al., 2018\)](#page-14-0).

#### Pan-NLRomes: a step toward fewer unknowns

Two recent RenSeq studies focused specifically on the pan-NLRome, with the goal of capturing a substantial fraction of inter- and intraspecific NLR diversity [\(Table 1\)](#page-7-0). The first study compared 64 A. thaliana accessions ([Van de Weyer](#page-17-0) [et al., 2019](#page-17-0)), while the second study spanned 16 accessions from 5 different Solanum species, plus single accessions of Nicotiana benthamiana and Capsicum annuum, also from the Solanaceae family [\(Seong et al., 2020\)](#page-16-0). In both A. thaliana and Solanum, the average number of NLR genes discovered in the different accessions was similar to that in the reference genomes, with some accessions having more NLR genes than the reference, pointing to the completeness of the RenSeq efforts. This was similar in a comparison of four high-quality long read assemblies from domesticated rice accessions ([Wang et al., 2019c\)](#page-17-0). In A. thaliana, 26 new IDs in addition to the 10 known from the original reference genome were found, and the universe of different NLR domain architectures was expanded from 22 in the reference genome to 97 in the entire set of 64 accessions [\(Van de](#page-17-0) [Weyer et al., 2019](#page-17-0)), highlighting how incomplete our understanding of NLR diversity is with a single reference genome. This dataset also confirmed different modes of evolution for subsets of NLRs [\(Kuang et al., 2004](#page-14-0)). While the phylogeny of most groups of apparent orthologs behaved in a manner consistent with allelic series, some groups showed more complex patterns indicative of sequence exchanges and neofunctionalization. This was also recently observed in [Lee and](#page-15-0) [Chae \(2020\),](#page-15-0) where a distinction between conserved "high fidelity" and highly-variable "radiating" NLR genes was made. Notably, radiating NLRs included all three known cases in A. thaliana where genes at the same locus confer resistance to very different pathogens, such as the RPP8/HRT/RCY1 ex-ample discussed above [\(Van de Weyer et al., 2019\)](#page-17-0).

Both the A. thaliana and the Solanaceae pan-NLRome analyses showed that there are apparently core NLR genes that are present in all or almost all accessions, but that these account only for a minority of all NLR genes [\(Van de](#page-17-0) [Weyer et al., 2019](#page-17-0); [Seong et al., 2020\)](#page-16-0). Furthermore, although the length of sequencing reads in these studies was substantially below what would be possible in 2020 with current long-read sequencing technology, the assembled contigs often contained more than one NLR, and also often neighboring non-NLR genes, allowing at least for partial ordering of NLR genes in clusters and anchoring of a fraction of contigs to the corresponding place in the reference genome. A more conclusive analysis of variation in NLR cluster size was presented in a comparison of eight completely assembled A. thaliana genomes, which confirmed substantial CNV in most clusters, and also identified small clusters of up to three genes that appeared to be size invariant in this set ([Jiao and Schneeberger, 2020;](#page-14-0) [Figure 3\)](#page-5-0).

As alluded to earlier, a key question is how extensive NLR diversity within the target group is and how many genome

assemblies are needed to capture most of the variation. The increase in known diversity in the A. thaliana study with 64 accessions—more than quadrupling the number of NLR domain architectures known from the reference—may appear daunting, but it was also observed that a set of about 40 maximally diverse accessions would have been sufficient to discover the majority of NLR genes, as defined by phyloge-netic orthology ([Van de Weyer et al., 2019](#page-17-0); [Table 1\)](#page-7-0). This might perhaps appear surprising at first glance, but it likely reflects strong selection for NLR genes, which are therefore more likely to spread faster throughout the entire population, and are therefore more evenly distributed in the population. With pan-NLRome information, it should become possible to test such hypotheses, including whether longrange linkage among NLR genes due to population structure is lower than for other genes.

## Turning Pan-NLRome sequence knowledge into functional knowledge

The ultimate goal of pan-NLRome studies is to accelerate the discovery of NLR specificity in disease resistance. Such knowledge has practical value in breeding, but it is equally interesting to understand the diversification of NLR function in an evolutionary and ecological context. The first step is the generation of species-wide pan-NLRomes, which is clearly within reach. The second step, using the pan-NLRomes to better understand how NLR diversity is distributed and to discern the evolutionary history of NLRs in a species, is more challenging, but we are optimistic that this can be solved as well. A highly innovative approach to parsing NLR diversity has recently been introduced, with a focus on revealing potential functional sites in NLRs that will aid in the rational design of novel or broad-spectrum disease resistance ([Prigozhin and Krasileva, 2020](#page-16-0)).

What remains is to comprehensively assign function to all existing NLRs. The most common known molecular activities of NLRs are (i) detecting a specific pathogen, or a specific pathogen effector, and (ii) enabling the functioning of other, pathogen-detecting NLRs (helper function). An additional activity for which there is less direct evidence so far is NLR expression being part of broad-spectrum disease resistance after a pathogen attack.

Most NLRs with a known function in disease resistance were discovered through linkage mapping in experimental crosses, using resistant and susceptible parents, often from natural populations, and also from mutant screens [\(Kourelis](#page-14-0) [and van der Hoorn, 2018](#page-14-0)). Similar to other genes underlying traits that vary in natural or crop populations, the identification of disease resistance genes has benefited from advances with genome-wide association studies (GWASs). Disease resistance lends itself particularly well to GWAS because it is often controlled by loci of large effect ([Atwell et al., 2010](#page-12-0); [Liu et al., 2020a](#page-15-0)), although this advantage is partially reduced, at least in wild plant species, by genetic heterogeneity, where different loci may underlie resistance to the same pathogen race ([Nemri et al., 2010\)](#page-15-0). The identification of NLRs responsible for disease resistance by GWAS has been accelerated by focusing on the NLR complement of genomes with RenSeq and using alignment-free approaches for identifying the causal genes [\(Arora et al., 2019](#page-12-0)). Similarly, RenSeq without assembly is a powerful technique to search for evolutionary signals of selection in NLR genes, which can help to prioritize specific genes for functional follow-up studies [\(Stam et al., 2016,](#page-16-0) [2019\)](#page-16-0). In this regard, there is a virtuous circle: The better the pan-NLRome of a species is known, the better strategies can be designed for focused interrogation of NLR genes.

Despite the advances with genetic mapping in experimental crosses or in natural populations, these approaches do not easily scale. As an example, a review of the literature indicated that only 88 NLR genes in A. thaliana had some assigned function in the middle of the year 2020 [\(Kourelis](#page-14-0) [and Kamoun, 2020\)](#page-14-0), yet there are on the order of 500 orthologous groups of NLR genes that are represented in multiple accessions ([Van de Weyer et al., 2019\)](#page-17-0). Of course, a major unknown is how many NLR genes have a positive function in the sense that they can detect effectors that pose an actual threat to the NLR carrier, and how many no longer or not yet have such a function, but it seems likely that many more than 88 NLR genes in this species play a role in disease resistance in nature.

For comprehensive assignment of NLR function, clearly more direct functional tests need to be pursued. The simplest cases for direct functional tests are those where a function is already known for a particular NLR, and based on this knowledge a system can be devised for rapid investigation of sequence-related NLRs. An example is the RPP13 allelic series in A. thaliana, where transient expression was used to determine which RPP13 alleles recognized which allele of the matching effector ATR13 ([Allen et al., 2004,](#page-12-0) [2008](#page-12-0); [Krasileva et al., 2011\)](#page-14-0).

A more general approach introduces NLR genes into susceptible plant backgrounds followed by extensive testing with different races of the pathogen of interest ([Yang et al.,](#page-17-0) [2013;](#page-17-0) [Zhang et al., 2015](#page-17-0); [Wang et al., 2019c\)](#page-17-0). Direct functional testing of NLRs by expressing them in a foreign background may, however, not always be so straightforward. Many NLRs act as sensor NLRs and rely on helper NLRs for conferring disease resistance [\(Wu et al., 2017](#page-17-0); [Jubic et al.,](#page-14-0) [2019;](#page-14-0) [Feehan et al., 2020](#page-13-0)). In the case of paired NLRs, this limitation could be circumvented by always introducing both members of a pair, which are easily found due to their characteristic arrangement in the genome. More challenging are NLR networks that include unlinked NLRs, such as those found in the complex NRC immune network in the Solanaceae [\(Wu et al., 2017](#page-17-0)). While the sensor NLRs in this case are massively expanded, they converge on relatively few, conserved helper NLRs, although it cannot be excluded that some sensor NLRs require co-evolved helper NLRs. A clear opportunity that emerges from pan-NLR network studies is to identify combinations of interacting NLRs that do not currently exist in available breeding material, essentially

<span id="page-12-0"></span>generating a pan-NLR network, to extend both qualitative aspects (new resistances) or quantitative aspects (fine-tuning existing resistances).

In addition, focusing only on pathogens that cause disease on the focal plant species may not reveal the function of all of its NLRs, because some might provide resistance to microbes or other enemies that are not considered pathogens of that species ([Cevik et al., 2019](#page-13-0)). Even if one can assign function to NLRs in terms of recognizing a specific pathogen, the question remains which effectors they detect. Knowledge of both the NLR and matching effector in turn is required for predicting the spectrum of pathogen strains that is recognized by different plant accessions in the field. An alternative therefore is to identify the effector proteins from pathogens or the plant proteins with which NLRs interact, an approach taken in two large yeast-two-hybrid interactome studies [\(Mukhtar et al., 2011](#page-15-0); [Weßling et al.,](#page-17-0) [2014\)](#page-17-0). These studies identified interactors for a large fraction of tested NLR fragments; having defined such NLR partners, one should be able to easily test how the interactions are affected by sequence variation mined from pan-NLRome efforts. In principle, this approach could be scaled up to include all members of a pan-NLRome. As pan-genome efforts take off also for pathogens (Badet and Croll, 2020), pan-NLRomes could then be tested in this manner against pan-effectoromes and their host targets. Importantly, pan-NLRome resources will help to choose alleles that provide an agriculturally optimal response to a specific effector or pathogen race, with the best compromise between yield penalty in the absence of the pathogen, effective pathogen detection and minimal collateral damage from too strong an immune reaction ([Harris et al., 2013\)](#page-14-0). Finally, in order to derive predictive models for NLR function, functional information must be captured in a structured manner; RefPlantNLR is a recent initiative that aims to do just that [\(Kourelis and Kamoun, 2020\)](#page-14-0).

The ultimate goal is to arrive at a point where NLR function can be predicted from sequence alone. By combining knowledge of protein–protein interactions with other sources of information such as structures of NLRs and their interactors, as well as knowledge of host protein modification by pathogen effectors, we foresee that predictive models of NLR function will come into reach in the next decade. Finally, with global knowledge of NLRs and matching effectors in hand, mapping the co-occurrence of such pairs in hosts and pathogens in time and space should reveal the true extent of past and ongoing arms races between hosts and their pathogens.

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