Jurassic NLR: conserved and dynamic evolutionary features of the atypically ancient immune receptor ZAR1

Hiroaki Adachi¹, Toshiyuki Sakai¹, Jiorgos Kourelis¹, Jose L. Gonzalez Hernandez², Abbas Maqbool¹ and Sophien Kamoun¹*

¹The Sainsbury Laboratory, University of East Anglia, Norwich Research Park, NR4 7UH, Norwich, UK
²Agronomy, Horticulture and Plant Sciences Department, South Dakota State University, Brookings, South Dakota, US

*For correspondence: sophien.kamoun@tsl.ac.uk

ABSTRACT

In plants, NLR immune receptors generally exhibit hallmarks of rapid evolution even at the intraspecific level. We reconstructed the evolutionary history of ZAR1, an atypically conserved NLR that traces its origin to early flowering plant lineages ~220 to 150 million years ago (Jurassic period). Ortholog sequence analyses revealed highly conserved features of ZAR1, including regions for pathogen effector recognition, intramolecular interactions and cell death activation. This uncovered a new conserved surface on the ZAR1 resistosome underside that is required for cell death induction. Throughout its evolution, ZAR1 acquired novel features, such as a C-terminal integration of a thioredoxin-like domain. ZAR1 duplicated into two paralog families, which underwent distinct evolutionary paths. We conclude that ZAR1 stands out among angiosperm NLRs for having experienced relatively limited gene duplication and expansion throughout its deep evolutionary history. Nonetheless, ZAR1 did also give rise to non-canonical NLR proteins with integrated domains and degenerated molecular features.

INTRODUCTION

Plants immune receptors, often encoded by disease resistance (R) genes, detect invading pathogens and activate innate immune responses that can limit infection (Jones and Dangl, 2006). A major class of immune receptors is formed by intracellular proteins of the nucleotide-binding leucine-rich repeat (NLR) family (Dodds and Rathjen, 2010; Jones et al., 2016; Kourelis and van der Hoorn, 2018). NLRs detect host-translocated pathogen effectors either by directly binding them or indirectly via host proteins known as guardees or decoys. NLRs are arguably the most diverse protein family in flowering plants (angiosperms) with many species having large (>100) and diverse repertoires of NLRs in their genomes (Shao et al., 2016; Baggs et al., 2017). They typically exhibit hallmarks of rapid evolution even at the intraspecific level (Van de Weyer et al., 2019; Lee and Chae, 2020; Prigozhin and Krasileva, 2020). Towards the end of the 20th century, Michelmore and Meyers (1998) proposed that NLRs evolve primarily through the birth-and-death process (Nei and Hughes, 1992). In this model, new NLRs emerge by recurrent cycles of gene duplication and loss—some genes are maintained in the genome acquiring new pathogen detection specificities, whereas others are deleted or become non-functional through the accumulation of deleterious mutations.
Such dynamic patterns of evolution enable the NLR immune system to keep up with fast-evolving effector repertoires of pathogenic microbes. However, as already noted over 20 years ago by Michelmore and Meyers (1998), a subset of NLR proteins are slow evolving and have remained fairly conserved throughout evolutionary time (Wu et al., 2017; Stam et al., 2019). These “high-fidelity” NLRs (per Lee and Chae, 2020) offer unique opportunities for comparative analyses, providing a molecular evolution framework to reconstruct key transitions and reveal functionally critical biochemical features (Delaux et al., 2019). Nonetheless, comprehensive evolutionary reconstructions of conserved NLR proteins remain limited despite the availability of a large number of plant genomes across the breadth of plant phylogeny. One of the reasons is that the great majority of NLRs lack clear-cut orthologs across divergent plant taxa. Here, we address this gap in knowledge by investigating the macroevolution of ZAR1 (HOPZ-ACTIVATED RESISTANCE1), an atypically ancient NLR, and asking fundamental questions about the conservation and diversification of this immune receptor throughout its deep evolutionary history.

NLRs occur across all kingdoms of life and generally function in non-self perception and innate immunity (Jones et al., 2016; Uehling et al., 2017). In the broadest biochemical definition, plant NLRs share a multidomain architecture typically consisting of a NB-ARC (nucleotide-binding domain shared with APAF-1, various R-proteins and CED-4) followed by a leucine-rich repeat (LRR) domain. Angiosperm NLRs form several major monophyletic groups with distinct N-terminal domain fusions (Shao et al., 2016; Kourelis and Kamoun, 2020). These include the subclades TIR-NLR with the Toll/interleukin-1 receptor (TIR) domain, CC-NLR with the Rx-type coiled-coil (CC) domain, CCζ-NLR with the RPW8-type CC (CCζ) domain (Tamborski and Krasileva, 2020) and the more recently defined CCζG10-NLR with a distinct type of CC (CCζG10) (Lee et al., 2020). Up to 10% of NLRs carry unconventional “integrated” domains in addition to the canonical tripartite domain architecture. Integrated domains are thought to generally function as decoys to bait pathogen effectors and enable pathogen detection (Cesari et al., 2014; Sarris et al., 2016; Wu et al., 2015; Kourelis and van der Hoorn, 2018). They include dozens of different modules indicating that novel domain acquisitions have repeatedly taken place throughout the evolution of plant NLRs (Sarris et al., 2016; Kroj et al., 2016). To date, over 400 NLRs from 31 genera in 11 orders of flowering plants have been experimentally validated as reported in the RefPlantNLR reference dataset (Kourelis and Kamoun, 2020). Several of these NLRs are coded by R genes that function against economically important pathogens and contribute to sustainable agriculture (Dangl et al., 2013).

In recent years, the research community has gained a better understanding of the structure/function relationships of plant NLRs and the immune receptor circuitry they form (Wu et al., 2018; Adachi et al., 2019a; Burdett et al., 2019; Jubic et al., 2019; Bayless and Nishimura, 2020; Feehan et al., 2020; Mermigka et al., 2020; Wang and Chai, 2020; Xiong et al., 2020; Zhou and Zhang, 2020). Some NLRs, such as ZAR1, form a single functional unit that carries both pathogen sensing and immune signalling activities in a single protein (termed ‘singleton NLR’ per Adachi et al., 2019a). Other NLRs function together in pairs or more complex networks, where connected NLRs have functionally specialized into sensor NLRs dedicated to pathogen detection or helper NLRs that are required for sensor NLRs to initiate immune signalling (Feehan et al., 2020). Paired and networked NLRs are thought to have evolved from multifunctional ancestral receptors through asymmetrical evolution (Adachi et al., 2019a, 2019b). As a result of their direct coevolution with pathogens, NLR sensors tend to
diversify faster than helpers and can be dramatically expanded in some plant taxa (Wu et al., 2017; Stam et al., 2019). For instance, sensor NLRs often exhibit non-canonical biochemical features, such as degenerated functional motifs and unconventional domain integrations (Adachi et al., 2019b; Seong et al., 2020).

The elucidation of plant NLR structures by cryo-electron microscopy has significantly advanced our understanding of the biochemical events associated with the activation of these immune receptors (Wang et al., 2019a; 2019b; Ma et al., 2020; Martin et al., 2020). The CC-NLR ZAR1, the TIR-NLRs RPP1 and Roq1 oligomerize upon activation into a wheel-like multimeric complex known as the resistosome. In the case of ZAR1, recognition of bacterial effectors occurs through its partner receptor-like cytoplasmic kinases (RLCKs), which tend to vary depending on the pathogen effector and host plant (Lewis et al., 2013; Wang et al., 2015; Seto et al., 2017; Schultink et al. 2019; Laflamme et al., 2020). Activation of ZAR1 induces conformational changes in the nucleotide binding domain resulting in ADP release, dATP/ATP binding and pentamerization of the ZAR1–RLCK complex into the resistosome. The ZAR1 resistosome exposes a funnel-shaped structure formed by the N-terminal α1 helices, which translocates into the plasma membrane and is thought to perturb membrane integrity to trigger cell death response (Wang et al., 2019b). The ZAR1 N-terminal α1 helix matches the MADA consensus sequence motif that is functionally conserved in ~20% of CC-NLRs including NLRs from dicot and monocot plant species (Adachi et al., 2019b). This suggests that the biochemical ‘death switch’ mechanism of the ZAR1 resistosome may apply to a significant fraction of CC-NLRs. Interestingly, unlike singleton and helper CC-NLRs, sensor CC-NLRs often carry degenerated MADA α1 helix motifs and/or N-terminal domain integrations, which would preclude their capacity to trigger cell death according to the ZAR1 model (Adachi et al., 2019b; Seong et al., 2020).

Comparative sequence analyses based on a robust evolutionary framework can yield insights into molecular mechanisms and help generate experimentally testable hypotheses. ZAR1 was previously reported to be conserved across multiple dicot plant species but whether it occurs in other angiosperms hasn’t been systematically studied (Baudin et al. 2017; Schultink et al. 2019; Harant et al. 2020). Here, we used a phylogenomic approach to investigate the molecular evolution of ZAR1 across flowering plants (angiosperms). We discovered 120 ZAR1 orthologs in 88 species, including monocot, magnoliid and eudicot species indicating that ZAR1 is an atypically conserved NLR that traces its origin to early angiosperm lineages ~220 to 150 million years ago (Jurassic period). We took advantage of this large collection of orthologs to identify highly conserved features of ZAR1, revealing regions for effector recognition, intramolecular interactions and cell death activation, along with a new conserved surface on the underside of the activated ZAR1 resistosome wheel. Throughout its evolution, ZAR1 also acquired novel features, including the C-terminal integration of a thioredoxin-like domain and duplication into two paralog families ZAR1-SUB and ZAR1-CIN. Members of the ZAR1-SUB paralog family have highly diversified in eudicots and often lack conserved ZAR1 features. We conclude that ZAR1 has experienced relatively limited gene duplication and expansion throughout its deep evolutionary history, but still did give rise to non-canonical NLR proteins with integrated domains and degenerated molecular features.
RESULTS

ZAR1 is the most widely conserved CC-NLR across angiosperms

To determine the distribution of ZAR1 across plant species, we applied a computational pipeline based on iterated BLAST searches of plant genome and protein databases (Figure 1A). These comprehensive searches were seeded with previously identified ZAR1 sequences from Arabidopsis, *N. benthamiana*, tomato, sugar beet and cassava (Baudin et al. 2017; Schultink et al. 2019; Harant et al. 2020). We also performed iterated phylogenetic analyses using the NB-ARC domain of the harvested ZAR1-like sequences, and obtained a well-supported clade that includes previously reported ZAR1 from the 5 eudicots (Arabidopsis, cassava, sugar beet, tomato and *N. benthamiana*) as well as new clade members from more distantly related plant species, notably *Colocasia esculenta* (taro, Alismatales), *Cinnamomum micranthum* (Syn. *C. kanehirae*), stout camphor, Magnoliidae) and *Aquilegia coerulea* (columbine, Ranunculales) (Supplementary table 1). In total, we identified 120 ZAR1 from 88 angiosperm species that tightly clustered in the ZAR1 phylogenetic clade (Figure 1B, Supplementary table 1). Among the 120 genes, 108 code for canonical CC-NLR proteins with 52.0 to 97.0% similarity to Arabidopsis ZAR1, whereas another 9 carry the three major domains of CC-NLR proteins but have a C-terminal integrated domain (ZAR1-ID, see below). The remaining 3 genes code for two truncated NLRs and a potentially mis-annotated coding sequence due to a gap in the genome sequence. In summary, we propose that the identified clade consists of ZAR1 orthologs from a diversity of angiosperm species. Our analyses of ZAR1-like sequences also revealed two well-supported sister clades of the ZAR1 ortholog clade (Figure 1B). We named these subclades ZAR1-SUB and ZAR1-CIN and we describe them in more details below.

We have recently proposed that ZAR1 is the most conserved CC-NLR between rosid and asterid plants (Harant et al. 2020). To further evaluate ZAR1 conservation relative to other CC-NLRs across angiosperms, we used a phylogenetic tree of 1475 NLRs from the monocot taro, the magnoliid stout camphor and 6 eudicot species (columbine, Arabidopsis, cassava, sugar beet, tomato, *N. benthamiana*) to calculate the phylogenetic (patristic) distance between each of the 49 Arabidopsis CC-NLRs and their closest neighbor from each of the other plant species. We found that ZAR1 stands out for having the shortest phylogenetic distance to its orthologs relative to other CC-NLRs in this diverse angiosperm species set (Figure 1—figure supplement 1). A similar analysis where we plotted the phylogenetic distance between each of the 159 *N. benthamiana* CC-NLRs to their closest neighbor from the other species also revealed ZAR1 as displaying the shortest patristic distance across all examined species (Figure 1—figure supplement 2). These analyses revealed that ZAR1 is possibly the most widely conserved CC-NLR in flowering plants (angiosperms).

Phylogenetic distribution of ZAR1 in angiosperms

Although ZAR1 is distributed across a wide range of angiosperms, we noted particular patterns in its phylogenetic distribution. Supplementary table 1 describes the gene identifiers and other features of ZAR1 orthologs sorted based on the phylogenetic clades reported by Smith and Brown (2018). 68 of the 88 plant species have a single-copy of ZAR1 whereas 20 species have two or more copies. ZAR1 is primarily a eudicot gene but we identified three
ZAR1 orthologs outside the eudicots, two in the monocot taro and another one in the magnoliid stout camphor. We failed to detect ZAR1 orthologs in 39 species among the 127 species we examined (Supplementary table 1). Except for taro, ZAR1 is missing in monocot species (17 examined), including in the well-studied *Hordeum vulgare* (barley), *Oryza sativa* (rice), *Triticum aestivum* (wheat) and *Zea mays* (maize). ZAR1 is also missing in all examined species of the eudicot Fabales, Cucurbitales, Apiales and Asterales. However, we found a ZAR1 ortholog in the early diverging eudicot columbine and ZAR1 is widespread in other eudicots, including in 63 rosid, 4 Caryophyllales and 18 asterid species.

**ZAR1 is an ancient Jurassic gene that predates the split between monocots, magnoliids and eudicots**

The overall conservation of the 120 ZAR1 orthologs enabled us to perform phylogenetic analyses using the full-length protein sequence and not just the NB-ARC domain as generally done with NLRs (Figure 2, Figure 2—figure supplement 1). These analyses yielded a robust ZAR1 phylogenetic tree with well-supported branches that generally mirrored established phylogenetic relationships between the examined plant species (Smith and Brown, 2018; Chaw et al., 2019). For example, the ZAR1 tree matched a previously published species tree of angiosperms based on 211 single-copy core ortholog genes (Chaw et al., 2019). We conclude that the origin of the ZAR1 gene predates the split between monocots, magnoliids and eudicots and its evolution traced species divergence ever since. We postulate that ZAR1 probably emerged in the Jurassic era ~220 to 150 million years ago (Mya) based on the species conservation of the orthologs.
Figure 2. ZAR1 gene is distributed across angiosperms. The phylogenetic tree was generated in MEGA7 by the neighbour-joining method using full length amino acid sequences of 120 ZAR1 orthologs identified in Figure 1. Each branch is marked with different colours based on the plant taxonomy. Red triangles indicate bootstrap support > 0.7. The scale bar indicates the evolutionary distance in amino acid substitution per site.

 divergence time estimate of Chaw et al. (2019) and consistent with the latest fossil evidence for the emergence of flowering plants (Fu et al., 2018).

ZAR1 is a genetic singleton in a locus that exhibits gene co-linearity across eudicot species

NLR genes are often clustered in loci that are thought to accelerate sequence diversification and evolution (Michelmore and Meyers, 1998; Lee and Chae, 2020). We examined the genetic context of ZAR1 genes using available genome assemblies of taro, stout camphor, columbine, Arabidopsis, cassava, sugar beet, tomato and N. benthamiana. The ZAR1 locus is generally devoid of other NLR genes as the closest NLR is found in the Arabidopsis genome 183 kb away from ZAR1 (Figure 2—figure supplement 2—supplementary table 1). We conclude that ZAR1 has probably remained a genetic singleton NLR gene throughout its evolutionary history in angiosperms.

Next, we examined the ZAR1 locus for gene co-linearity across the examined species. We noted a limited degree of gene co-linearity between Arabidopsis vs. cassava, cassava vs. tomato, and tomato vs. N. benthamiana (Figure 2—figure supplement 2). Flanking conserved
genes include the ATPase and protein kinase genes that are present at the ZAR1 locus in both rosid and asterid eudicots. In contrast, we didn’t observe conserved gene blocks at the ZAR1 locus of taro, stout camphor and columbine, indicating that this locus is divergent in these species. Overall, although limited, the observed gene co-linearity in eudicots is consistent with the conclusion that ZAR1 is a genetic singleton with an ancient origin.

**ZAR1 orthologs carry sequence motifs known to be required for Arabidopsis ZAR1 resistosome function**

The overall sequence conservation and deep evolutionary origin of ZAR1 orthologs combined with the detailed knowledge of ZAR1 structure and function provide a unique opportunity to explore the evolutionary dynamics of this ancient immune receptor in a manner that cannot be applied to more rapidly evolving NLRs. We used MEME (Multiple EM for Motif Elicitation) (Bailey and Elkan, 1994) to search for conserved sequence patterns among the 117 ZAR1 orthologs (ZAR1 and ZAR1-ID) that encode full-length CC-NLR proteins. This analysis revealed several conserved sequence motifs that span across the ZAR1 orthologs (range of protein lengths: 753-1132 amino acids) (Figure 3A, Figure 3—supplementary table 1). In Figure 3A, we described the major five sequence motifs or interfaces known to be required for Arabidopsis ZAR1 function that are conserved across ZAR1 orthologs.

Effectors recognition by ZAR1 occurs indirectly via binding to RLCKs through the LRR domain. Key residues in the Arabidopsis ZAR1-RLCK interfaces are highly conserved among ZAR1 orthologs and were identified by MEME as conserved sequence patterns (Figure 3A). Valine (V) 544, histidine (H) 597, tryptophan (W) 825 and phenylalanine (F) 839 in the Arabidopsis ZAR1 LRR domain were validated by mutagenesis as important residues for RLCK binding whereas isoleucine (I) 600 was not essential (Wang et al. 2019a; Hu et al. 2020). In the 117 ZAR1 orthologs, V544, H597, W825 and F839 are conserved in 97-100% of the proteins compared to only 63% for I600.

After effector recognition, Arabidopsis ZAR1 undergoes conformational changes from monomeric inactive form to oligomeric active state. This is mediated by ADP release from the NB-ARC domain and subsequent ATP binding, which triggers further structural remodelling in ZAR1 leading to the formation of the activated pentameric resistosome (Wang et al. 2019b). NB-ARC sequences that coordinate binding and hydrolysis of dATP, namely P-loop and MHD motifs, are highly conserved across ZAR1 orthologs (Figure 3A). Histidine (H) 488 and lysine (K) 195, located in the ADP/ATP binding pocket (Wang et al. 2019a; Wang et al. 2019b), are invariant in all 117 orthologs. In addition, three NB-ARC residues, W150, S152 and V154, known to form the NBD-NBD oligomerisation interface for resistosome formation (Wang et al. 2019b; Hu et al. 2020), are present in 82-97% of the ZAR1 orthologs and were also part of a MEME motif (Figure 3A).

The N-terminal CC domain of Arabidopsis ZAR1 mediates cell death signalling thorough the N-terminal α1 helix/MADA motif, that becomes exposed in activated ZAR1 resistosome to form a funnel like structure that perturbs the plasma membrane (Baudin et al., 2017; 2019; Wang et al. 2019b; Adachi et al., 2019b). We detected an N-terminal MEME motif that matches the α1 helix/MADA motif (Figure 3A). We also used the HMMER software (Eddy, 1998) to query the ZAR1 orthologs with a previously reported MADA motif-Hidden Markov
Figure 3. ZAR1 orthologs carry conserved sequence patterns required for Arabidopsis ZAR1 resistosome function. (A) Schematic representation of the Arabidopsis ZAR1 protein highlighting the position of conserved sequence patterns across ZAR1 orthologs. Consensus sequence patterns were identified by MEME using 117 ZAR1 ortholog sequences. Raw MEME motifs are listed in Figure 3—Supplementary table 1. Red asterisks indicate residues functionally validated in Arabidopsis ZAR1 for NBD-NBD and ZAR1-RLCK interfaces. (B) Conservation and variation of each amino acid among ZAR1 orthologs across angiosperms. Amino acid alignment of 117 ZAR1 orthologs was used for conservation score calculation via the ConSurf server (https://consurf.tau.ac.il). The conservation scores are mapped onto each amino acid position in Arabidopsis ZAR1 (NP_190664.1). (C, D) Distribution of the ConSurf conservation score on the Arabidopsis ZAR1 structure. The inactive ZAR1 monomer is illustrated in cartoon representation with different colours based on each canonical domain (C) and the conservation score (D). Major five variable surfaces (VS1 to VS5) on the inactive ZAR1 monomer structure are described in grey dot or black boxes in panel B or D, respectively.
Model (HMM) (Adachi et al., 2019b). This HMMER search detected a MADA-like sequence at the N-terminus of all 117 ZAR1 orthologs (Supplementary table 1).

Taken together, based on the conserved motifs depicted in Figure 3A, we propose that angiosperm ZAR1 orthologs share the main functional features of Arabidopsis ZAR1: 1) effector recognition via RLCK binding, 2) remodelling of intramolecular interactions via ADP/ATP switch, 3) oligomerisation via the NBD-NBD interface and 4) α1 helix/MADA motif-mediated activation of hypersensitive cell death.

**ZAR1 resistosome underside displays a novel conserved surface**

To identify additional conserved and variable features in ZAR1 orthologs, we used ConSurf (Ashkenazy et al., 2016) to calculate a conservation score for each amino acid and generate a diversity barcode for ZAR1 orthologs (Figure 3B). The overall pattern is that the 117 ZAR1 orthologs are fairly conserved. Nonetheless, the CC domain (except for the N-terminal MADA motif and a few conserved stretches), the junction between the NB-ARC and LRR domains and the very C-terminus were distinctly more variable than the rest of the protein (Figure 3B).

We also used the cryo-EM structures of Arabidopsis ZAR1 to determine how the ConSurf score map onto the 3D structures (Figure 3C, D and Figure 4). First, we found five major variable surfaces (VS1 to VS5) on the inactive ZAR1 monomer structure (Figure 3C, D), as depicted in the ZAR1 diversity barcode (Figure 3B). VS1 comprises α2/α4 helices and a loop between α3 and α4 helices of the CC domain. VS2 and VS3 corresponds to α1/α2 helices of NBD and a loop between α2 and α3 helices of HD1, respectively. VS4 comprises a loop between WHD and LRR and first three helices of the LRR domain. VS5 is mainly derived from the last three helices of the LRR domain and the loops between these helices (Figure 3B, D).

We also noted significant sequence variation at the glutamate rings (comprised of E11, E18, E130 and E134) inside the Arabidopsis ZAR1 resistosome (Figure 4—figure supplement 1). Mutations of glutamic acid (E) 11 and E18 impaired Arabidopsis ZAR1-mediated cell death without interfering with oligomerization and plasma membrane association (Wang et al. 2019b). Interestingly, activated Arabidopsis CC-NLRs, NRG1.1 and ADR1, were recently proposed to function as Ca\(^{2+}\) influx channels and require their N-terminal negatively charged residues (NRG1.1 E14 and ADR1 D11) for the Ca\(^{2+}\) influx and cell death (Jacob et al., 2021). The E130/E134 ring was previously discussed as potentially having Ca\(^{2+}\) transporter activity because of structural similarity to rings in the structures of the mitochondrial calcium uniporter from *Caenorhabditis elegans* and the calcium release-activated calcium channel ORAI from *Drosophila melanogaster* (Burdett et al., 2019). Whereas E11 is conserved in 94% of ZAR1 orthologs, only 3-18% retain E18, E130 and E134 in the same positions as Arabidopsis ZAR1. Therefore, it remains unclear whether the ZAR1 resistosome also functions as a Ca\(^{2+}\) influx channel given the lack of conservation of the N-terminal E/D-ring across ZAR1 orthologs and other CC-type NLRs.

Next, we examined highly conserved surfaces on inactive and active ZAR1 structures (Figure 4A, B). Consistent with the MEME analyses, we confirmed that highly conserved surfaces match to the RLCK binding interfaces (Figure 4A, B). We also confirmed that the N-terminal α1 helix/MADA motif is conserved on the resistosome surfaces, although the first four N-
Figure 4. ZAR1 orthologs across angiosperms display multiple conserved surfaces on the resistosome structure. Distribution of the ConSurf conservation score was visualized on the inactive monomer (A), active monomer (B) and resistosome (C) structures of Arabidopsis ZAR1. Each structure and cartoon representation are illustrated with different colours based on the conservation score shown in Figure 3. (D) Schematic representation of the conserved underside surface region among ZAR1 orthologs. The conserved underside regions are described with consensus sequence patterns identified by MEME. Red asterisks indicate residues exposed on resistosome surfaces. The raw MEME motif is listed in Figure 3—Supplementary table 1.
terminal amino acids are missing from the N terminus of the active ZAR1 cryo-EM structures (Figure 4B).

Remarkably, these analyses revealed a highly conserved ring that is exposed on the underside surface of the ZAR1 resistosome opposite to the funnel-shaped structure (Figure 4C). This conserved underside surface is mainly formed by residues located in α2 helix-loop, α3 helix-loop and α4 helix-loop of the NBD (Figure 4—figure supplement 2 and 3). Within the conserved patch, arginine (R) 214, K252 and K260, are positively charged residues that are exposed on the underside surface and are conserved in 94%, 96% and 99% of ZAR1 orthologs, respectively (Figure 4—figure supplement 4). The three residues form a positive electrostatic potential ring on the underside surface of the ZAR1 resistosomes (Figure 4—figure supplement 4). The conserved underside ring is composed of a 14 amino acid motif as revealed by MEME (Figure 4D). We propose that this underside sequence pattern has been maintained throughout the more than 150 million years of ZAR1 evolution and is likely to be functionally important.

**ZAR1 resistosome underside ring residues are required for cell death activity**

We hypothesized that the ZAR1 resistosome underside ring surface is functionally important given its high conservation across the orthologs. To test this, we performed site directed mutagenesis and investigated the contribution of the underside ring residues to the cell death activity of ZAR1. We introduced the mutations in the MHD motif mutant of *N. benthamiana* ZAR1 (NbZAR1**D481V**), which triggers autoactive cell death in the *N. benthamiana* expression system (Harant et al. 2020). We took advantage of ZAR1 structural information to make alanine substitutions in the 14 amino acid residues located on the underside surface of the ZAR1 resistosome (Figure 5A). This mutation strategy targeted residues located on NBD α2 helix-loop (referred to as Na2), α3 helix-loop (Na3) and α4 helix-loop (Na4) regions separately (NbZAR1**Na2**, NbZAR1**Na3**, NbZAR1**Na4**) or in combination (NbZAR1**14A**) (Figure 5A). We found that Na3, Na4 and 14A, but not Na2, mutations significantly reduced NbZAR1**D481V** cell death inducing activity (Figure 5B). Given that three of the 14 amino acid residues (R214, K252 and K260) form a clear positive electrostatic potential ring (Figure 4—figure supplement 4), we reasoned that a triple alanine substitution to the positively charged residues may also affect the cell death activity. We observed that the R214A/K252A/K260A mutation (RKA, NbZAR1**RKA**) abolished the cell death activity of NbZAR1**D481V** (Figure 5B). All of the NbZAR1**D481V** mutant proteins accumulated to similar levels as NbZAR1**D481V** when expressed in *N. benthamiana* leaves, indicating that the observed loss-of-function phenotypes were not due to protein destabilization (Figure 5C). These results indicate that the conserved α3 helix-loop, α4 helix-loop and the positively charged ring on the underside surface is required for cell death induction by the ZAR1 resistosome. Interestingly, the ZAR1 resistosome underside surface is not conserved in the resistosomes of the TIR-NLR proteins Roq1 and RPP1, which have a totally distinct conformation (Figure 6A-D). In Roq1 and RPP1 resistosomes, NBD is sandwiched between TIR and LRR domains unlike ZAR1 where NBD is exposed at the underside of resistosome (Figure 6A-D).

**Integration of a PLP3a thioredoxin-like domain at the C-termini of cassava and cotton ZAR1**
Figure 5. Mutations in the conserved underside surface impair cell death activity of autoimmune NbZAR1<sup>D481V</sup>. (A) Schematic representation of NbZAR1 and the mutated sites in the conserved underside surface of the resistosome. Mutated sites and substituted residues are shown as red characters in the ZAR1 sequence alignment. (B) Cell death observed in <i>N. benthamiana</i> after expression of NbZAR1 mutants. <i>N. benthamiana</i> leaf panels expressing NbZAR1<sup>D481V</sup> (ZAR1<sup>D481V</sup>), the MADA motif mutant (ZAR1<sup>L17E/D481V</sup>), and the conserved underside surface mutants (ZAR1<sup>Nα2/D481V</sup>, ZAR1<sup>Nα3/D481V</sup>, ZAR1<sup>Nα4/D481V</sup>, ZAR1<sup>RKA/D481V</sup>, ZAR1<sup>14A/D481V</sup>) were photographed at 5 days after agroinfiltration. Violin plots showing cell death intensity scored as an HR index based on three independent experiments. Statistical differences among the samples were analysed with Tukey’s honest significance difference (HSD) test (p<0.01). (C) In planta accumulation of the NbZAR1 variants. For anti-HA immunoblots of the NbZAR1 mutant proteins, total proteins were prepared from <i>N. benthamiana</i> leaves at 2 days after agroinfiltration. Empty vector control is described as EV. Equal loading was checked with Reversible Protein Stain Kit (Thermo Fisher).
Figure 6. Comparison of ZAR1, Roq1 and RPP1 resistosomes. (A) Schematic representation of ZAR1, Roq1 and RPP1 proteins highlighting domain architecture with different colour codes. Amino acid sequences of the NB-ARC domain were aligned by MAFFT version 7 program and are shown at a lower panel. P-loop and MHD motif are marked with blue boxes. A conserved underside sequence motif, which is required for cell death activity of ZAR1, is marked with a red box. (B-D) The structures of ZAR1, Roq1 and RPP1 resistosomes are depicted as surface representation in two different orientations. Individual domains are coloured according to the scheme in A. In ZAR1 resistosome, 10 amino acids which are conserved in the underside surface and are required for cell death activity of ZAR1 are depicted as red patch. Unlike ZAR1, NBD in Roq1 and RPP1 is sandwiched between TIR and LRR domains.
As noted earlier, 9 ZAR1 orthologs carry an integrated domain (ID) at their C-termini (Supplementary table 1). These ZAR1-ID include 2 predicted proteins (XP_021604862.1 and XP_021604864.1) from Manihot esculenta (cassava) and 7 predicted proteins (KAB1998109.1, PPD92094.1, KAB2051569.1, TYG89033.1, TYI49934.1, TYJ04029.1, KJB48375.1) from the cotton plant species Gossypium barbadense, Gossypium darwinii, Gossypium mustelinum and Gossypium raimondii (Supplementary table 1). The integrations follow an otherwise intact LRR domain and vary in length from 108 to 266 amino acids (Figure 7A). We confirmed that the ZAR1-ID gene models of cassava XP_021604862.1 and XP_021604864.1 are correct based on RNA-seq exon coverage in the NCBI database (database ID: LOC110609538). However, cassava ZAR1-ID XP_021604862.1 and XP_021604864.1 are isoforms encoded by transcripts from a single locus on chromosome LG2 (RefSeq sequence NC_035162.1) of the cassava RefSeq assembly (GCF_001659605.1) which also produces transcripts encoding isoforms lacking the C-terminal ID (XP_021604863.1, XP_021604865.1, XP_021604866.1, XP_021604867.1 and XP_021604868.1). Thus, cassava ZAR1-ID are probably splicing variants from a unique cassava ZAR1 gene locus (Figure 7—figure supplement 1).
To determine the phylogenetic relationship between ZAR1-ID and canonical ZAR1, we mapped the domain architectures of ZAR1 orthologs on the phylogenetic tree shown in Figure 2 (Figure 7—figure supplement 2). Cassava and cotton ZAR1-ID occur in different branches of the ZAR1 rosid clade indicating that they may have evolved as independent integrations although alternative evolutionary scenarios such as a common origin followed by subsequent deletion of the ID or lineage sorting remain possible (Figure 7—figure supplement 2).

We annotated all the C-terminal extensions as thioredoxin-like using InterProScan (Trx, IPR036249; IPR013766; cd02989). The integrated Trx domain sequences share sequence similarity to each other (Figure 7B). They are also similar to Arabidopsis AT3G50960 (phosphoducin-like PLP3a; 34.8-90% similarity to integrated Trx domains), which is located immediately downstream of ZAR1 in a tail-to-tail configuration in the Arabidopsis genome (Figure 2—figure supplement 2; Figure 7—figure supplement 3). We also noted additional genetic linkage between ZAR1 and Trx genes in other rosid species, namely field mustard, orange, cacao, grapevine and apple, and in the asterid species coffee (Figure 2—figure supplement 2—supplementary table 2). We conclude that ZAR1 is often genetically linked to a PLP3a-like Trx domain gene and that the integrated domain in ZAR1-ID has probably originated from a genetically linked sequence.

The ZAR1-SUB clade emerged early in eudicot evolution from a single ZAR1 duplication event

Phylogenetic analyses revealed ZAR1-SUB as a sister clade of the ZAR1 ortholog clade (Figure 1B, Figure 8). ZAR1-SUB clade comprises 129 genes from a total of 55 plant species (Supplementary table 2). 21 of the 55 plant species carry a single-copy of ZAR1-SUB whereas 34 species have two or more copies. Of the 129 genes, 122 code for canonical CC-NLR proteins (692-1038 amino acid length) with shared sequence similarities ranging from 36.5 to 99.9% (Figure 8).

Unlike ZAR1, ZAR1-SUB NLRs are restricted to eudicots (Figure 8—figure supplement 1, Supplementary table 2). Three out of 129 genes are from the early diverging eudicot clade Ranunculales species, namely columbine, Macleaya cordata (plume poppy) and Papaver somniferum (opium poppy) (Figure 8—figure supplement 1). The remaining ZAR1-SUB are spread across rosid and asterid species (Figure 8—figure supplement 1). We found that 11 species have ZAR1-SUB genes but lack a ZAR1 ortholog (Supplementary table 3). These 11 species include two of the early diverging eudicots plume poppy and opium poppy, and the Brassicales Carica papaya (papaya). Interestingly, papaya is the only Brassicales species carrying a ZAR1-SUB gene, whereas the 16 other Brassicales species have ZAR1 but lack ZAR1-SUB genes (Figure 8—figure supplement 1, Supplementary table 3). In total, we didn’t detect ZAR1-SUB genes in 44 species that have ZAR1 orthologs, and these 44 species include the monocot taro, the magnoliid stout camphor and 42 eudicots, such as Arabidopsis, sugar beet and N. benthamiana (Supplementary table 3).

In summary, given the taxonomic distribution of the ZAR1-SUB clade genes, we propose that ZAR1-SUB has emerged from a single duplication event of ZAR1 prior to the split between Ranunculales and other eudicot lineages about ~120-130 Mya based on the species divergence time estimate of Chaw et al. (2019).
Figure 8. ZAR1-SUB has emerged early in eudicots and diverged at MADA motif sequence. The phylogenetic tree was generated in MEGA7 by the neighbour-joining method using full length amino acid sequences of 120 ZAR1, 129 ZAR1-SUB and 11 ZAR1-CIN identified in Figure 1. Each branch is marked with different colours based on the plant taxonomy. Red triangles indicate bootstrap support > 0.7. The scale bar indicates the evolutionary distance in amino acid substitution per site. NLR domain architectures are illustrated outside of the leaf labels: MADA is red, CC is pink, NB-ARC is yellow, LRR is blue and other domain is orange. Black asterisks on domain schemes describe truncated NLRs or potentially mis-annotated NLR. The zoomed tree is described in Figure 8—figure supplement 3.

ZAR1-SUB paralogs have significantly diverged from ZAR1

We investigated the sequence patterns of ZAR1-SUB proteins and compared them to the sequence features of canonical ZAR1 proteins that we identified earlier (Figures 3, 4). MEME analyses revealed several conserved sequence motifs (Figure 9—supplementary table 1). Especially, the MEME motifs in the ZAR1-SUB NB-ARC domain were similar to ZAR1 ortholog motifs (Figure 9—supplementary table 2). These include P-loop and MHD motifs, which are broadly conserved in NB-ARC of 97% and 100% of the ZAR1-SUB NLRs, respectively (Figure 9A). MEME also revealed sequence motifs in the ZAR1-SUB LRR domain that partially overlaps in position with the conserved ZAR1-RLCK interfaces (Figure 9A, Figure 9—figure supplement 1). However, the ZAR1-SUB MEME motifs in the LRR domain were variable at the ZAR1-RLCK
Figure 9. Conserved sequence distributions in ZAR1-SUB and ZAR1-CIN. (A) Schematic representation of the ZAR1-SUB protein highlighting the position of the representative conserved sequence patterns across ZAR1-SUB. Representative consensus sequence patterns identified by MEME are described on the scheme. Raw MEME motifs are listed in Figure 9—supplementary tables 1 and 2. (B) Conservation and variation of each amino acid among ZAR1-SUB and ZAR1-CIN. Amino acid alignment of 129 ZAR1-SUB or 8 ZAR1-CIN was used for conservation score calculation via the ConSurf server (https://consurf.tau.ac.il). The conservation scores are mapped onto each amino acid position in queries XP_004243429.1 (ZAR1-SUB) and RWR85656.1 (ZAR1-CIN), respectively. (C) Schematic representation of the ZAR1-CIN protein highlighting the position of the representative conserved sequence patterns across 8 ZAR1-CIN. Raw MEME motifs are listed in Figure 9—supplementary tables 3 and 4.

Interface positions compared to ZAR1, and the motif sequences were markedly different between ZAR1-SUB and ZAR1 proteins (Figures 3A, 9A, Figure 9—supplementary table 2).

Remarkably, unlike ZAR1 orthologs, MEME did not predict conserved sequence pattern from a region corresponding to the MADA motif, indicating that these sequences have diverged across ZAR1-SUB proteins (Figure 9A). We confirmed the low frequency of MADA motifs in ZAR1-SUB proteins using HMMER searches with only ~30% (38 out of 129) of the tested proteins having a MADA-like sequence (Supplementary table 2, Figure 8). Moreover,
conserved sequence patterns were not predicted for the NBD-NBD interface and the conserved underside surface of the ZAR1 resistosome (Figure 9A, Figure 9—figure supplement 1). This indicates that the NB-ARC domain of ZAR1-SUB proteins is highly diversified in contrast to the relatively conserved equivalent region of ZAR1 proteins.

We generated a diversity barcode for ZAR1-SUB proteins using the ConSurf as we did earlier with ZAR1 orthologs (Figure 9B). This revealed that there are several conserved sequence blocks in each of the CC, NB-ARC and LRR domains, such as the regions corresponding to P-loop, MHD motif and the equivalent of the ZAR1-RLCK interfaces. Nonetheless, ZAR1-SUB proteins are overall more diverse than ZAR1 orthologs especially in the CC domain, including the N-terminal MADA motif, and the NBD/HD1 regions of the NB-ARC domain where the NBD-NBD interface is located.

Next, we mapped the ConSurf conservation scores onto a homology model of a representative ZAR1-SUB protein (XP_004243429.1 from tomato) built based on the Arabidopsis ZAR1 cryo-EM structures (Figure 9—figure supplement 2). As highlighted in Figure 9—figure supplement 2B and C, conserved residues, such as MHD motif region in the WHD, are located inside of the monomer and resistosome structures. Interestingly, although the prior MEME prediction analyses revealed conserved motifs in positions matching the ZAR1-RLCK interfaces in the LRR domain, the ZAR1-SUB structure homology models displayed variable surfaces in this region (Figures 9A, Figure 9—figure supplement 2A). This indicates that the variable residues within these sequence motifs are predicted to be on the outer surfaces of the LRR domain and may reflect interaction with different ligands.

Taken together, these results suggest that unlike ZAR1 orthologs, the ZAR1-SUB paralogs have divergent molecular patterns for regions known to be involved in effector recognition, resistosome formation and activation of hypersensitive cell death.

**Eleven tandemly duplicated ZAR1-CIN genes occur in a 500 kb cluster in the Cinnamomum micranthum (stout camphor) genome**

The ZAR1-CIN clade, identified by phylogenetic analyses as a sister clade to ZAR1 and ZAR1-SUB, consists of 11 genes from the magnoliid species stout camphor (Figure 1B, Figure 8, Supplementary table 4). 8 of the 11 ZAR1-CIN genes code for canonical CC-NLR proteins with 63.8 to 98.9% sequence similarities to each other, whereas the remaining 3 genes code for truncated NLR proteins. Interestingly, all ZAR1-CIN genes occur in a ~500 kb cluster on scaffold QPKB01000005.1 of the stout camphor genome assembly (GenBank assembly accession GCA_003546025.1) (Figure 8—figure supplement 2). This scaffold also contains the stout camphor ZAR1 ortholog (CmZAR1, RWR84015), which is located 48 Mb from the ZAR1-CIN cluster (Figure 8—figure supplement 2). Based on the observed phylogeny and gene clustering, we suggest that the ZAR1-CIN cluster emerged from segmental duplication and expansion of the ancestral ZAR1 gene after stout camphor split from the other examined ZAR1 containing species.

We examined the expression of the eleven CmZAR1 and ZAR1-CIN genes in seven tissues of C. micranthum based on the data of Chaw et al. (Chaw et al., 2019). The CmZAR1 gene is relatively highly expressed in seven different tissues of the stout camphor tree (Figure 8—
figure supplement 2). In contrast, only five of the eleven ZAR1-CIN genes displayed detectable expression levels. Of these, two ZAR1-CIN genes (RWR85656 and RWR85657) had different expression patterns across the tissues. Whereas RWR85657 had the highest expression level in flowers, RWR85656 displayed the highest expression levels in stem and old leaf tissues (Figure 8—figure supplement 2). The implications of these observations remain unclear but may reflect different degrees of tissue specialization of the ZAR1-CIN genes.

**Tandemly duplicated ZAR1-CIN display variable ligand binding interfaces on the LRR domain**

We performed MEME and ConSurf analyses of the 8 intact ZAR1-CIN proteins as described above for ZAR1 and ZAR1-SUB. The ConSurf barcode revealed that although ZAR1-CIN proteins are overall conserved, their WHD region and LRR domain include some clearly variable blocks (Figure 9B). MEME analyses of ZAR1-CIN sequences revealed that like ZAR1 orthologs, the MADA, P-loop and MHD motifs match highly conserved blocks of the ZAR1-CIN ConSurf barcode (Figure 9B, C, Figure 9—supplementary tables 3 and 4). Consistently, 87.5% (7 out of 8) of the ZAR1-CIN proteins were predicted to have a MADA-type N-terminal sequence based on MADA-HMM analyses (Supplementary table 4, Figure 8).

MEME picked up additional sequence motifs in ZAR1-CIN proteins that overlap in position with the NBD-NBD and ZAR1-RLCK interfaces (Figure 9C, Figure 9—figure supplement 3). However, the sequence consensus at the NBD-NBD and ZAR1-RLCK interfaces indicated these motifs are more variable among ZAR1-CIN proteins relative to ZAR1 orthologs, and the motif sequences were markedly different from the matching region in ZAR1 (Figures 3A, 9C).

We also mapped the ConSurf conservation scores onto a homology model of a representative ZAR1-CIN protein (RWR85656.1) built based on the Arabidopsis ZAR1 cryo-EM structures (Figure 9—figure supplement 2). This model revealed several conserved surfaces, such as on the α1 helix in the CC domain, the WHD of the NB-ARC domain and underside surface of the resistosome (Figure 9—figure supplement 2B, C, E). In contrast, the ZAR1-CIN structure homology models displayed highly varied surfaces especially in the LRR region matching the RLCK binding interfaces of ZAR1 (Figure 9—figure supplement 2A). This sequence diversification on the LRR surface suggests that the ZAR1-CIN paralogs may have different host partner proteins and/or effector recognition specificities compared to ZAR1.

**DISCUSSION**

This study of ZAR1 macroevolution originated from phylogenomic analyses we initiated during the UK COVID-19 lockdown of March 2020. We performed iterated comparative sequence similarity searches of plant genomes using the CC-NLR immune receptor ZAR1 as a query, and subsequent phylogenetic evaluation of the recovered ZAR1-like sequences. This revealed that ZAR1 is an ancient gene with 120 orthologs recovered from 88 species including monocot, magnoliid and eudicot plants. ZAR1 is an atypically conserved NLR in these species with the gene phylogeny tracing species phylogeny, and consistent with the view that ZAR1 originated early in angiosperms during the Jurassic geologic period ~220 to 150 Mya (Figure 10). The ortholog series enabled us to determine that resistosome sequences that are known to be functionally important and have remained highly conserved throughout the long
We propose that the ancestral ZAR1 gene has emerged ~220 to 150 million years ago (Mya) before monocot and eudicot lineages split. ZAR1 gene is widely conserved CC-NLR in angiosperms, but it is likely that ZAR1 has lost in a monocot lineage, Commelinales. A sister clade paralog ZAR1-SUB has emerged early in the eudicot lineages and may have lost in Caryophyllales. Another sister clade paralog ZAR1-CIN has duplicated from ZAR1 gene and expanded in the Magnoliidae *C. micranthum*. Trx domain integration to C terminus of ZAR1 has independently occurred in few rosid lineages. The evolutionary history of ZAR1. This also revealed a new conserved sequence ring on the underside of the resistosome, which has remained constrained in ZAR1 ortholog proteins and is required for ZAR1-mediated cell death induction (Figures 4 and 5). The only unexpected feature among ZAR1 orthologs is the acquisition of a C-terminal thioredoxin-like domain in cassava and cotton species (Figures 7 and 10). Our phylogenetic analyses also indicated that ZAR1 duplicated twice throughout its evolution (Figure 10). In the eudicots, ZAR1 spawned a large paralog family, ZAR1-SUB, which greatly diversified and often lost the typical sequence features of ZAR1. A second paralog, ZAR1-CIN, is restricted to a tandemly repeated 11-gene cluster in stout camphor. Overall, our findings map patterns of functional conservation, expansion and diversification onto the evolutionary history of ZAR1 and its paralogs (Figure 10). Phylogenomics analyses, such as this work, provide a unique evolutionary perspective on the function of a plant NLR immune receptor and generate experimentally testable hypotheses (Figure 10—figure supplement 1).

ZAR1 most likely emerged prior to the split between monocots, Magnoliids and eudicots, which corresponds to ~220 to 150 Mya based on the dating analyses of Chaw et al. (2019). The origin of the angiosperms remains hotly debated with uncertainties surrounding some of the fossil record coupled with molecular clock analyses that would benefit from additional genome sequences of undersampled taxa (Coiro et al., 2019). Recently, Fu et al. (2018) provided credence to an earlier emergence of angiosperms with the discovery of the fossil flower *Nanjinganthus dendrostyla*, which places the emergence of flowering plants at the
Early Jurassic. It is tempting to speculate that ZAR1 emerged among these early flowering plants during the period when dinosaurs dominated planet earth.

NLRs are notorious for their rapid and dynamic evolutionary patterns even at the intraspecific level. In sharp contrast, ZAR1 is an atypically core NLR gene conserved in a wide range of angiosperm species (Figures 3 and 4). Nevertheless, Arabidopsis ZAR1 can recognize diverse bacterial pathogen effectors, including five different effector families distributed among nearly half of a collection of ~500 *Pseudomonas syringae* strains (Laflamme et al., 2020) and an effector AvrAC from *Xanthomonas campestris* (Wang et al., 2015). How did ZAR1 remain conserved throughout its evolutionary history while managing to detect a diversity of effectors? The answer to the riddle lies in the fact that ZAR1 effector recognition occurs via its partner RLCKs. HopZ-ETI-deficient 1 (ZED1) and ZED1-related kinases (ZRKs) of the RLCK XII-2 subfamily rest in complex with inactive ZAR1 proteins and bait effectors by binding them directly or by recruiting other effector-binding RLCKs, such as the family VII PBS1-like protein 2 (PBL2) (Lewis et al., 2013; Wang et al., 2015). These ZAR1-associated RLCKs are highly diversified in Arabidopsis, with 8 of the 13 RLCK XII-2 members occurring in the expanded ZRK gene cluster (Lewis et al., 2013). In this ZRK cluster, RKS1/ZRK1 is required for recognition of *X. campestris* effector AvrAC (Wang et al., 2015) and ZRK3 and ZRK5/ZED1 are required for recognition of *P. syringae* effectors HopF2a and HopZ1a, respectively (Lewis et al., 2013; Seto et al., 2017). Therefore, as in the model discussed by Schultink et al. (2019), RLCKs have evolved as pathogen ‘sensors’ whereas ZAR1 acts as a conserved signal executor to activate immune response. Future phylogenomic analyses of the RLCK subfamilies coupled with functional analyses with ZAR1 across angiosperms will help test and sharpen this model.

Our MEME and ConSurf analyses are consistent with the model of ZAR1/RLCK evolution described above. ZAR1 is not just exceptionally conserved across angiosperms but it has also preserved sequence patterns that are key to resistosome-mediated immunity (Figures 3 and 4). In particular, within the LRR domain, ZAR1 orthologs display highly conserved surfaces for RLCK binding (Figure 4). We conclude that ZAR1 has been guarding host kinases throughout its evolution ever since the Jurassic period. These findings strikingly contrast with observations recently made by Prigozhin and Krasileva (2020) on highly variable Arabidopsis NLRs (hvNLRs), which tend to have diverse LRR sequences. For instance, the CC-NLR RPP13 displays variable LRR surfaces across 62 Arabidopsis accessions, presumably because these regions are effector recognition interfaces that are caught in arms race coevolution with the oomycete pathogen *Hyaloperonospora arabidopsidis* (Prigozhin and Krasileva, 2020). The emerging view is that the mode of pathogen detection (direct vs indirect recognition) drives an NLR evolutionary trajectory by accelerating sequence diversification at the effector binding site or by maintaining the binding interface with the partner garde/decoy proteins (Prigozhin and Krasileva, 2020).

ZAR1 orthologs display a patchy distribution across angiosperms (Figure 10, Supplementary table 1). Given the low number of non-eudicot species with ZAR1 it is challenging to develop a conclusive evolutionary model. Nonetheless, the most parsimonious explanation is that ZAR1 was lost in the monocot Commelinales lineage (Figure 10, Supplementary table 1). ZAR1 is also missing in some eudicot lineages, notably Fabales, Cucurbitales, Apiales and Asterales (Supplementary table 1). Cucurbitaceae (Cucurbitales) species are known to have reduced repertoires of NLR genes possibly due to low levels of gene duplications and frequent
deletions (Lin et al., 2013). ZAR1 may have been lost in this and other plant lineages as part of an overall shrinkage of their NLRomes or as a consequence of selection against autoimmune phenotypes triggered by NLR mis-regulation (Karasov et al., 2017; Adachi et al., 2019a). In the future, it would be interesting to investigate the repertoires of RLCK subfamilies VII and XII in species that lack ZAR1 orthologs.

We unexpectedly discovered that some ZAR1 orthologs from cassava and cotton species carry a C-terminal thioredoxin-like domain (ZAR1-ID in Figure 7). What is the function of these integrated domains? The occurrence of unconventional domains in NLRs is relatively frequent and ranges from 5 to 10% of all NLRs. In several cases, integrated domains have emerged from pathogen effector targets and became decoys that mediate detection of the effectors (Kourelis and van der Hoorn, 2018). Whether or not the integrated Trx domain of ZAR1-ID functions to bait effectors will need to be investigated. Since ZAR1-ID proteins still carry intact kinase binding interfaces (Supplementary table 1—source data 2), they may have evolved dual or multiple recognition specificities via RLCKs and the Trx domain. In addition, all ZAR1-ID proteins have an intact N-terminal MADA motif (Figure 7—figure supplement 2), suggesting that they probably can execute the hypersensitive cell death through their N-terminal CC domains even though they carry a C-terminal domain extension (Adachi et al., 2019b). However, we noted multiple splice variants of the ZAR1-ID gene of cassava, some of which lack the Trx integration (Figure 7—figure supplement 1). It is possible that both ZAR1 and ZAR1-ID isoforms are produced, potentially functioning together as a pair of sensor and helper NLRs.

Our sequence analyses of ZAR1-ID indicate that the integrated Trx domain originates from the PLP3 phosphoducin gene, which is immediately downstream of ZAR1 in the Arabidopsis genome and adjacent to ZAR1 in several other eudicot species (Figure 7—figure supplement 3). Whether or not PLP3 plays a role in ZAR1 function and the degree to which close genetic linkage facilitated domain fusion between these two genes are provocative questions for future studies.

ZAR1 spawned two classes of paralogs through two independent duplication events. The ZAR1-SUB paralog clade emerged early in the eudicot lineage—most likely tens of millions of years after the emergence of ZAR1—and has diversified into at least 129 genes in 55 species (Figure 10). ZAR1-SUB proteins are distinctly more diverse in sequence than ZAR1 orthologs and generally lack key sequence features of ZAR1, like the MADA motif and the NBD-NBD oligomerisation interface (Figures 8 and 9) (Adachi et al., 2019b; Wang et al. 2019b; Hu et al. 2020). This pattern is consistent with ‘use-it-or-lose-it’ evolutionary model, in which NLRs that specialize for pathogen detection lose some of the molecular features of their multifunctional ancestors (Adachi et al., 2019b). Therefore, we predict that many ZAR1-SUB proteins evolved into specialized sensor NLRs that require NLR helper mates for executing the hypersensitive response. It is possible that ZAR1-SUB helper mate is ZAR1 itself, and that these NLRs evolved into a phylogenetically linked network of sensors and helpers similar to the NRC network of asterid plants (Wu et al., 2017). However, 11 species have a ZAR1-SUB gene but lack a canonical ZAR1 (Supplementary table 3), indicating that these ZAR1-SUB NLRs may have evolved to depend on other classes of NLR helpers.
How would ZAR1-SUB sense pathogens? Given that the LRR domains of most ZAR1-SUB proteins markedly diverged from the RLCK binding interfaces of ZAR1, it is unlikely that ZAR1-SUB proteins bind RLCKs in a ZAR1-type manner (Figure 9—figure supplement 2). This leads us to draw the hypothesis that ZAR1-SUB proteins have diversified to recognize other ligands than RLCKs. In the future, functional investigations of ZAR1-SUB proteins could provide insights into how multifunctional NLRs, such as ZAR1, evolve into functionally specialized NLRs.

The ZAR1-CIN clade consists of 11 clustered paralogs that are unique to the magnoliid species stout camphor as revealed from the genome sequence of the Taiwanese small-flowered camphor tree (also known as Cinnamomum kanehirae, Chinese name niu zhang 牛樟) (Chaw et al., 2019). This cluster probably expanded from ZAR1, which is ~48 Mbp on the same genome sequence scaffold (Figure 8—figure supplement 2). The relatively rapid expansion pattern of ZAR1-CIN into a tandemly duplicated gene cluster is more in line with the classical model of NLR evolution compared to ZAR1 maintenance as a genetic singleton over tens of millions of years (Michelmore and Meyers, 1998). ZAR1-CIN proteins may have neofunctionalized after duplication, acquiring new recognition specificities as a consequence of coevolution with host partner proteins and/or pathogen effectors. Consistent with this view, ZAR1-CIN exhibit different patterns of gene expression across tissues (Figure 8—figure supplement 2). Moreover, ZAR1-CIN proteins display distinct surfaces at the ZAR1-RLCK binding interfaces and may bind to other ligands than RLCKs as we hypothesized above for ZAR1-SUB (Figure 9—figure supplement 2). ZAR1-CIN could be viewed as intraspecific highly variable NLRs (hvNLR) per the nomenclature of Prigozhin and Kasileva (2020).

Unlike ZAR1-SUB, ZAR1-CIN have retained the N-terminal MADA sequence (Figure 9, Figure 9—figure supplement 2). We propose that ZAR1-CIN are able to execute the hypersensitive cell death on their own similar to ZAR1. However, ZAR1-CIN display divergent sequence patterns at NBD-NBD oligomerisation interfaces compared to ZAR1 (Figure 9C, Figure 9—figure supplement 3). Therefore, ZAR1-CIN may form resistosome-type complexes that are independent of ZAR1. One intriguing hypothesis is that ZAR1-CIN may associate with each other to form heterocomplexes of varying complexity and functionality operating as an NLR receptor network. In any case, the clear-cut evolutionary trajectory from ZAR1 to the ZAR1-CIN paralog cluster provides a robust evolutionary framework to study functional transitions and diversifications in this CC-NLR lineage.

In summary, our phylogenomics analyses raise a number of intriguing questions about ZAR1 evolution. The primary conclusion we draw is that ZAR1 is an ancient CC-NLR that has been guarding RLCKs ever since the Jurassic Period. We propose that throughout at least 150 million years, ZAR1 has maintained its molecular features for sensing pathogens via RLCKs and activating hypersensitive cell death. We also identified an intriguingly conserved NB-ARC ring surface, located on the underside of the ZAR1 resistosome and functionally required for the cell death response (Figures 4 and 5). We propose that this underside surface plays an important function in the resistosome, possibly by associating with critical host factors. The equivalent region of the NB-ARC underside ring is not exposed onto the underside surface of the recently resolved TIR-NLRS RPP1 and Roq1 resistosome structures (Figure 6) (Ma et al., 2020; Martin et al., 2020). Future studies will indicate whether the ZAR1 underside surface may be a specific feature of CC-NLR resistosomes. Further comparative analyses, combining
molecular evolution and structural biology, of plant resistosomes and between resistosomes and the apoptosomes and inflammasome of animal NLR systems (Wang and Chai, 2020) will yield novel experimentally testable hypotheses for NLR research.

Materials and Methods

ZAR1 sequence retrieval

We performed BLAST (Altschul et al., 1990) using previously identified ZAR1 sequences as queries (Baudin et al. 2017; Schultink et al. 2019; Harant et al. 2020) to search ZAR1 like sequences in NCBI nr or nr/nt database (https://blast.ncbi.nlm.nih.gov/Blast.cgi) and Phytozome12.1 (https://phytozome.jgi.doe.gov/pz/portal.html#!search?show=BLAST). In the BLAST search, we used cut-offs, percent identity ≥ 40% and query coverage ≥ 95%. The BLAST pipeline was circulated by using the obtained sequences as new queries to search ZAR1 like genes over the angiosperm species. We also performed the BLAST pipeline against a plant NLR dataset annotated by NLR-parser (Steurnagel et al., 2015) from 38 plant reference genome databases (Supplementary table 5).

Phylogenetic analyses

For the phylogenetic analysis, we aligned NLR amino acid sequences using MAFFT v.7 (Katoh and Standley, 2013) and manually deleted the gaps in the alignments in MEGA7 (Kumar et al., 2016). Full-length or NB-ARC domain sequences of the aligned NLR datasets were used for generating phylogenetic trees. The neighbour-joining tree was made using MEGA7 with JTT model and bootstrap values based on 100 iterations. All datasets used for phylogenetic analyses are in source data files.

Patristic distance analyses

To calculate the phylogenetic (patristic) distance, we used Python script based on DendroPy (Sukumaran and Mark, 2010). We calculated patristic distances from each CC-NLR to the other CC-NLRs on the phylogenetic tree (Figure 1—source data 3) and extracted the distance between CC-NLRs of Arabidopsis or N. benthamiana to the closest NLR from the other plant species. The script used for the patristic distance calculation is available from GitHub (https://github.com/slt666666/ Phylogenetic_distance_plot2).

Gene co-linearity analyses

To investigate genetic co-linearity at ZAR1 loci, we extracted the 3 genes upstream and downstream of ZAR1 using GFF files derived from reference genome databases (Supplementary table 5). To identify conserved gene blocks, we used gene annotation from NCBI Protein database and confirmed protein domain information based on InterProScan (Jones et al, 2014).

Sequence conservation analyses
Full-length NLR sequences of the each subfamily ZAR1, ZAR1-SUB or ZAR1-CIN were subjected to motif searches using the MEME (Multiple EM for Motif Elicitation) (Bailey and Elkan, 1994) with parameters ‘zero or one occurrence per sequence, top twenty motifs’, to detect consensus motifs conserved in ≥90% of the input sequences. The output data are summarized in Figure 3—supplementary table 1, Figure 9—supplementary table 1 and Figure 9—supplementary table 3.

To predict the MADA motif from ZAR1, ZAR1-SUB and ZAR1-CIN datasets, we used the MADA-HMM previously developed (Adachi et al., 2019b), with the hmmsearch program (hmmsearch –max -o <outputfile> <hmmfile> <seqdb>) implemented in HMMER v2.3.2 (Eddy, 1998). We termed sequences over the HMMER cut-off score of 10.0 as the MADA motif and sequences having the score 0-10.0 as the MADA-like motif.

To analyze sequence conservation and variation in ZAR1, ZAR1-SUB and ZAR1-CIN proteins, aligned full-length NLR sequences in MAFFT v.7 were used for ConSurf (Ashkenazy et al., 2016). Arabidopsis ZAR1 (NP_190664.1), a tomato ZAR1-SUB (XP_004243429.1) or a Stout camphor ZAR1-CIN (RWR85656.1) was used as a query for each analysis of ZAR1, ZAR1-SUB or ZAR1-CIN, respectively. The output datasets are in Figure 3—source data 1, Figure 9—source data 1 and Figure 9—source data 2.

**Protein structure analyses**

The atomic coordinates of ZAR1 (protein data bank accession codes; 6J5T), Roq1 (protein data bank accession codes; 7JLV, 7JLU, and 7JLX) and RPP1 (protein data bank accession codes; 7CRC and 7CRB) were downloaded from protein data bank for illustration in ccp4mg. We used the cryo-EM structures of ZAR1 as templates to generate homology models of ZAR1-SUB and ZAR1-CIN. Amino acid sequences of a tomato ZAR1-SUB (XP_004243429.1) and a stout camphor ZAR1-CIN (RWR85656.1) were submitted to Protein Homology Recognition Engine V2.0 (Phyre2) for modelling (Kelley et al., 2015). The coordinates of ZAR1 structure (6J5T) were retrieved from the Protein Data Bank and assigned as modelling template by using Phyre2 Expert Mode. The resulting model of ZAR1-SUB and ZAR1-CIN, and the ZAR1 structures (6J5T) were illustrated with the ConSurf conservation scores in PyMol.

**Plant growth condition**

Wild-type *N. benthamiana* plants were grown in a controlled growth chamber with temperature 22-25°C, humidity 45-65% and 16/8 hr light/dark cycle.

**Plasmid constructions**

To generate the conserved underside ring mutants of NbZAR1, the full-length of NbZAR1D481V was amplified by Phusion High-Fidelity DNA Polymerase (Thermo Fisher) with two independent primer sets. Primers NbZAR1_Undermut_N_Fw (tgtgaagacaaAATGGTGGATGCGGTGGTC) and NbZAR1_Undermut_N_Rv (tgtgaagacaaCACCCTTTTGTCATTGAAAAC) were used for amplifying N-terminal fragment of NbZAR1 prior to the mutation site. Primers NbZAR1_Undermut_C_Fw (tgtgaagacaaTGATTGTTATGGATGATTTG) and NbZAR1_Undermut_C_STOP_Rv
(tgtgaagacaaAAGCTTAGTTCCTATGTTCTTC) or NbZAR1_Undermut_C_ns_Rv
(tgtgaagacaaCGAACCGTTCCTATGTTCTTCCTTC) were used for amplifying C-terminal fragment
of NbZAR1 with or without the stop codon. We used GENEWIZ Standard Gene Synthesis
service to synthesize mutated gene fragments (Figure 5—source data 1). Purified amplicons
and a mutated gene fragment were used in Golden Gate assembly with pICH41308 (Addgene
no. 47998) or pAGM1287 (Addgene no. 47996) to generate level 0 modules (Weber et al.,
2011). The level 0 plasmids were then used for Golden Gate assembly with or without C-
terminal tag module pICSL50009 (6xHA, Addgene no. 50309) into the binary vector
pICH86988 (Engler et al., 2014).

**Transient gene-expression and cell death assay**

Transient expression of NbZAR1 mutants in *N. benthamiana* were performed by
agroinfiltration according to methods described by Bos et al. (2006). Briefly, four-weeks old
* N. benthamiana * plants were infiltrated with *Agrobacterium tumefaciens* strains carrying the
binary expression plasmids. * A. tumefaciens * suspensions were prepared in infiltration buffer
(10 mM MES, 10 mM MgCl₂, and 150 μM acetosyringone, pH5.6) and were adjusted to OD₆₀₀
= 0.5. Macroscopic cell death phenotypes were scored according to the scale of Segretin et
al. (2014) modified to range from 0 (no visible necrosis) to 7 (fully confluent necrosis).

**Western blot analysis**

Protein samples were prepared from six discs (8 mm diameter) cut out of *N. benthamiana*
leaves at 2 days after agroinfiltration and were homogenised in extraction buffer [10%
glycerol, 25 mM Tris- HCl, pH 7.5, 1 mM EDTA, 150 mM NaCl, 2% (w/v) PVPP, 10 mM DTT, 1x
protease inhibitor cocktail (SIGMA), 0.5% IGEPAL (SIGMA)]. The supernatant obtained after
centrifugation at 12,000 xg for 10 min was used for SDS-PAGE. Immunoblotting was
performed with HA-probe (F-7) HRP (Santa Cruz Biotech) in a 1:5,000 dilution. Equal loading
was checked by taking images of the stained PVDF membranes with Pierce Reversible Protein
Stain Kit (#24585, Thermo Fisher).

**RNA-seq data analyses**

Public RNA-seq reads, which were previously obtained with Illumina HiSeq 2000 (Chaw et al.,
2019), were used to analyze expression profiles of CmZAR1 and ZAR1-CIN genes in the stout
camphor tree (Accession Numbers: SRR7416905, SRR7416906, SRR7416908, SRR7416909,
SRR7416910, SRR7416911, and SRR7416918). Reads were mapped to the stout camphor
genome assembly (GenBank assembly accession GCA_003546025.1) using the splice-aware
RNAseq tool in CLC Genomics Workbench vs 20.0.4 (https://digitalinsights.qiagen.com) and
transformed into a Transcripts Per Million (TPM) value according to Li et al. (2010). TPM
values were visualized by the heatmap. The heatmap was colored by eight ranges (0, 0–5,
5–20, 20–40, 40–60, 60–80, 80–100, 100<) of TPM values.

**ACKNOWLEDGEMENTS**

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AUTHOR CONTRIBUTIONS

H.A. and S.K. mainly wrote the paper; H.A., A.M. and S.K. designed the research and supervised the work; H.A., T.S., J.K., J.L.G.H. and A.M. performed research.

DECLARATION OF INTERESTS

S.K. receives funding from industry on NLR biology.

References


Figure 1—figure supplement 1. Arabidopsis ZAR1 is the most conserved CC-NLR across angiosperms. (A) Phylogenetic tree of NLR proteins from 8 plant species. The phylogenetic tree was generated in MEGA7 by the neighbour-joining method using NB-ARC domain sequences of 1475 NLRs identified from taro, stout camphor, columbine, Arabidopsis, cassava, sugar beet, tomato and N. benthamiana. The scale bars indicate the evolutionary distance in amino acid substitution per site. We used CC-NLR and CC\textsubscript{G10}-NLR superclades for calculating phylogenetic distances. (B) The phylogenetic (patristic) distance of two CC-NLR nodes between Arabidopsis and other plant species were calculated from the NB-ARC phylogenetic tree in A. The closest patristic distances are plotted with different colours based on plant species. Representative Arabidopsis NLRs are highlighted. The closest patristic distances of two CC-NLR nodes between N. benthamiana and other plant species can be found in Figure 1—figure supplement 2.
Figure 1—figure supplement 2. NbZAR1 is highly conserved across angiosperms. The phylogenetic (patristic) distance of two CC-NLR nodes between *N. benthamiana* and the closest NLR from the other plant species were calculated from the NB-ARC phylogenetic tree in Figure 1—figure supplement 1. The closest patristic distances are plotted with different colours based on plant species.
Figure 2—figure supplement 1. Sequence alignment of full-length ZAR1 ortholog proteins across angiosperms. Amino acid sequences of ZAR1 orthologs were aligned by MAFFT version 7 program. Conserved motif sequences highlighted in this study are marked with red boxes. Red asterisks indicate substitution sites for introducing gain or loss of ZAR1 protein function.
Figure 2—figure supplement 2. Schematic representation of the intragenomic relationship at ZAR1 loci across angiosperm genomes. We selected representative 8 plant species genome assemblies based on the phylogenetic tree in Figure 2 and used them for the synteny-based analysis of the ZAR1 loci. We highlight genes showing intragenomic linkages with different colours based on the gene annotations. Genes genetically linked to ZAR1 in eudicots are listed in Figure 2—figure supplement 2—Supplementary table 2.
Figure 4—figure supplement 1. E18, E130 and E134 on glutamate rings inside of the Arabidopsis ZAR1 resistosome are variable across the orthologs. The ConSurf conservation scores at E11 and E18 (A) or at E130 and E134 (B) are illustrated in cartoon representation of the Arabidopsis ZAR1 resistosome structure.
Figure 4—figure supplement 2. Residues in α2-loop, α3-loop and α4-loop of NBD make up conserved surface on underside of ZAR1 resistosome. NB-ARC domain of activated ZAR1 monomer is described as schematic representation. Residues that contribute to the conserved underside surface are shown as stick representation. The residues that are located on α3-loop and α4-loop and are shown as stick representation with carbon atoms coloured green. The residues that are located on α2 and on the loop between α2 and β2 are shown as stick representation with carbon atoms coloured yellow. A dATP molecule is present in nucleotide binding pocket and shown in stick representation with carbon atoms coloured orange.
Figure 4—figure supplement 3. Amino acid residues on the conserved underside surface of the ZAR1 resistosome. Conserved underside surface of the ZAR1 resistosome with five (top left) or single (bottom left) ZAR1 molecule(s). Both images are zoomed in from the underside view in Figure 4C. Conserved amino acids are labelled as numbers and listed in the right table.

<table>
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<tr>
<th>Number</th>
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<th>Location in αβ</th>
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<td>4</td>
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<td>14</td>
<td>Lys-260</td>
<td>9</td>
<td>loop</td>
<td></td>
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</table>
Figure 4—figure supplement 4. Three conserved residues form a positive electrostatic potential ring on undersurface of the ZAR1 resistosome. (A, B) Underside view of the ZAR1 resistosome with the ConSurf conservation score (A) or electrostatic potential (B). The color gradient from red to blue represents negative to positive electrostatic potentials. Black boxes indicate regions zoomed in panels C and D. (C, D) Conserved underside surface of the ZAR1 resistosome with five (C) or single (D) ZAR1 molecule(s). The underside surface is illustrated with the ConSurf conservation score (Left) and electrostatic potential (right). Regions marked by white or black dot lines are exposed to the underside surface from single ZAR1 protein. Yellow boxes indicate positive charged residues that are conserved on the underside surface across ZAR1 orthologs.
Figure 7—figure supplement 1. Cassava ZAR1 and ZAR1-ID are transcribed from a single locus on the genome. The gene locus of cassava ZAR1 (XP_021604863.1, XP_021604865.1, XP_021604866.1, XP_021604867.1 and XP_021604868.1) and ZAR1-ID (XP_021604862.1 and XP_021604864.1) is shown with RNA-seq exon coverage and is extracted from NCBI database (database ID: LOC110609538).
Figure 7—figure supplement 2. Trx domain integration occurred in two independent rosid ZAR1 subclades. The phylogenetic tree shown in Figure 2 was used to describe NLR domain architectures. Domain schemes are aligned to right side of the leaf labels: MADA is red, CC is pink, NB-ARC is yellow, LRR is blue and other domain is orange. Black asterisks on domain schemes describe truncated NLRs or potentially mis-annotated NLR. Each branch is marked with different colours based on the plant taxonomy. Red triangles indicate bootstrap support > 0.7. The scale bar indicates the evolutionary distance in amino acid substitution per site.
Figure 7—figure supplement 3. Integrated Trx domains show high sequence similarity to ZAR1-linked PLP3a gene in Arabidopsis. (A) Schematic representation of the intragenomic relationship at ZAR1 loci between Arabidopsis and cassava. We highlight sequence similarity of integrated Trx domain in Cassava ZAR1 (MeZAR1) to PLP3a gene genetically linked to Arabidopsis ZAR1 (AtZAR1). Details are explained in Figure 2—figure supplement 2. (B) Amino acid sequences of Arabidopsis PLP3a gene (AT3G50960) and integrated domains of an MeZAR1 (XP_021604862.1) and a cotton ZAR1 (GbZAR1; KAB1998109.1).
Figure 8—figure supplement 1. ZAR1-SUB gene is distributed across eudicots. The phylogenetic tree was generated in MEGA7 by the neighbour-joining method using full length amino acid sequences of 129 ZAR1-SUB orthologs identified in Figure 1. Each branch is marked with different colours based on the plant taxonomy. Red triangles indicate bootstrap support > 0.7. The scale bar indicates the evolutionary distance in amino acid substitution per site. Red asterisks on plant order term describe that NLRs from Malpighiales are distributed in three independent clades.
Figure 8—figure supplement 2. ZAR1-CIN gene cluster occurs in the Cinnamomum micranthum genome. (A) The subclades including ZAR1, ZAR1-SUB and ZAR1-CIN were zoomed in from the phylogenetic tree constructed in Figure 1—figure supplement 1. Red triangles indicate bootstrap support > 0.7. The scale bar indicates the evolutionary distance in amino acid substitution per site. Well supported subclades (I and II) in ZAR1-CIN are described with red or blue dot box. The gene IDs: taro (MQM-), stout camphor (RWR-), columbine (Aqcoe-), Arabidopsis (AT-), cassava (Manes-), sugar beet (Bv-), tomato (Solyc-) and N. benthamiana (NbS-). (B) Schematic representation of the ZAR1-CIN gene cluster on a C. micranthum (Stout camphor) scaffold. Stout camphor ZAR1 (CmZAR1) and ZAR1-CIN genes are highlighted in orange and yellow, respectively. (C) A heatmap showing Transcripts Per Million (TPM) values of the CmZAR1 and ZAR1-CIN genes. Public RNA-seq datasets from seven different tissue samples in C. micranthum were used for this heatmap analysis.
Figure 8—figure supplement 3. Phylogenetic analysis of ZAR1 and the sister clade members. The phylogenetic tree was generated in MEGA7 by the neighbour-joining method using full length amino acid sequences of 120 ZAR1, 129 ZAR1-SUB and 11 ZAR1-CIN identified in Figure 1. Each branch is marked with different colours based on the plant taxonomy. Red triangles indicate bootstrap support > 0.7. The scale bar indicates the evolutionary distance in amino acid substitution per site.
Figure 9—figure supplement 1. Sequence alignment of full-length ZAR1 and ZAR1-SUB proteins. (A) Schematic representation of the ZAR1-SUB protein highlighting the position of the representative conserved sequence patterns across ZAR1-SUB. (B) Amino acid sequences of ZAR1 orthologs and a representative ZAR1-SUB (XP_004243429.1 from tomato) were aligned by MAFFT version 7 program. ZAR1 motif sequences highlighted in this study are marked with red boxes. Positions of MEME motifs identified from ZAR1-SUB are marked in blue boxes. Raw MEME motifs are listed in Figure 9—supplementary tables 1 and 2.
Figure 9—figure supplement 2. ZAR1 and the sister subclade NLRs display different conserved surfaces on the resistosome structure. Distribution of the ConSurf conservation score was visualized on the inactive monomer (A), active monomer (B) and resistosome structures (C-E) of Arabidopsis ZAR1 or the structure homology models of ZAR1-SUB (XP_004243429.1) and ZAR1-CIN (RWR85656.1). Each structure and cartoon representation are illustrated with different colours based on the conservation score shown in Figures 3 and 9. Resistosome structures are shown from different angles, from side (C), from upper side (D) and from underside (E).
Figure 9—figure supplement 3. Sequence alignment of full-length ZAR1 and ZAR1-CIN proteins. (A) Schematic representation of the ZAR1-CIN protein highlighting the position of the representative conserved sequence patterns across ZAR1-SUB. (B) Amino acid sequences of ZAR1 orthologs and a representative ZAR1-CIN (RWR85656.1) were aligned by MAFFT version 7 program. ZAR1 motif sequences highlighted in this study are marked with red boxes. Positions of MEME motifs identified from ZAR1-CIN are marked in orange boxes. Raw MEME motifs are listed in Figure 9—supplementary tables 3 and 4.
Figure 10—figure supplement 1. Workflow from prediction to validation of novel protein function. The phylogenomics analysis in searching and classifying ortholog genes enables to predict motifs that have been conserved throughout the evolutionary history of plants. The comparison of the sequence conservation with 3D structures reveals highly conserved interfaces on the protein within its inactive and active states. This computational pipeline generates experimentally testable hypotheses to uncover novel protein functions.
Supplementary files

Supplementary table 1. List of ZAR1 in angiosperms.

Supplementary table 2. List of ZAR1-SUB.

Supplementary table 3. List of plant species with the number of ZAR1, ZAR1-SUB and ZAR1-CIN genes.

Supplementary table 4. List of ZAR1-CIN.

Supplementary table 5. Reference genome databases used for NLR annotation with NLR-parser.

Figure 2—figure supplement 2—Supplementary table 1. List of the closest NLR genes to ZAR1 locus.

Figure 2—figure supplement 2—Supplementary table 2. List of genes genetically linked to ZAR1 in eudicots.

Figure 3—supplementary table 1. List of MEME motifs predicted from ZAR1 in angiosperms.

Figure 9—supplementary table 1. List of MEME motifs predicted from ZAR1-SUB.

Figure 9—supplementary table 2. Comparison of MEME motifs between ZAR1-SUB and ZAR1. Black dot boxes indicate corresponding regions between ZAR1-SUB and ZAR1 based on MAFFT v7 alignment in Figure 9—figure supplement 1. Red boxes indicate motifs which are highlighted in Figures 3 and 4.

Figure 9—supplementary table 3. List of MEME motifs predicted from ZAR1-CIN.

Figure 9—supplementary table 4. Comparison of MEME motifs between ZAR1-CIN and ZAR1. Black dot boxes indicate corresponding regions between ZAR1-CIN and ZAR1 based on MAFFT v7 alignment in Figure 9—figure supplement 3. Red boxes indicate motifs which are highlighted in Figures 3 and 4.

Source data files

Figure 1—source data 1. Amino acid sequences of full-length NLRs used for phylogenetic analysis in Figure 1B. This file contains 1268 NLR amino acid sequences with the IDs, taro (MQM-), stout camphor (RWR-), columbine (Aqcoe-), Arabidopsis (AT-), sugar beet (Bv-) and tomato (Soly-).

Figure 1—source data 2. Amino acid sequences for NLR phylogenetic tree in Figure 1B. This file contains NB-ARC domain sequences used for phylogenetic analysis.
Figure 1—source data 3. NLR phylogenetic tree file in Figure 1B. The phylogenetic tree was saved in newick file format.

Figure 1—figure supplement 1—source data 1. Amino acid sequences of full-length NLRs used for phylogenetic analysis in Figure 1—figure supplement 1. This file contains 1475 NLR amino acid sequences with the IDs, taro (MQM-), stout camphor (RWR-), columbine (Aqcoe-), Arabidopsis (AT-), cassava (Manes-), sugar beet (Bv-), tomato (Solyc-) and N. benthamiana (NbS-).

Figure 1—figure supplement 1—source data 2. Amino acid sequences for NLR phylogenetic tree in Figure 1—figure supplement 1. This file contains NB-ARC domain sequences used for phylogenetic analysis.

Figure 1—figure supplement 1—source data 3. NLR phylogenetic tree file in Figure 1—figure supplement 1. The phylogenetic tree was saved in newick file format.

Figure 2—source data 1. NLR phylogenetic tree file in Figure 2. The phylogenetic tree was saved in newick file format.

Figure 3—source data 1. The ConSurf conservation score among ZAR1 proteins. The table contains conservation score on each position of amino acid sequences in Arabidopsis ZAR1.

Figure 5—source data 1. The conserved underside surface mutations in NbZAR1. The fasta file contains NbZAR1 fragment gene sequences carrying mutations. Blue characters indicate BpiI sites used for Golden Gate assembly.

Figure 8—source data 1. NLR phylogenetic tree file in Figure 6. The phylogenetic tree was saved in newick file format.

Figure 9—source data 1. The ConSurf conservation score among ZAR1-SUB proteins. The table contains conservation score on each position of amino acid sequences in a tomato ZAR1-SUB, XP_004243429.1.

Figure 9—source data 2. The ConSurf conservation score among ZAR1-CIN proteins. The table contains conservation score on each position of amino acid sequences in a cinnamomum ZAR1-CIN, RWR85656.1.

Supplementary table 1—source data 1. Amino acid sequences of 120 ZAR1 in angiosperms. This file contains 120 ZAR1 amino acid sequences identified from computational pipeline in Figure 1A.

Supplementary table 1—source data 2. Amino acid alignment file of 120 ZAR1 in angiosperms. Full-length amino acid sequences of ZAR1 orthologs were aligned by MAFFT version 7.
Supplementary table 1—source data 3. List of plant species carrying ZAR1 as single-copy gene.

Supplementary table 1—source data 4. List of plant species carrying 2 or more ZAR1 genes.

Supplementary table 2—source data 1. Amino acid sequences of 129 ZAR1-SUB. This file contains 129 ZAR1-SUB amino acid sequences identified from computational pipeline in Figure 1A.

Supplementary table 2—source data 2. Amino acid alignment file of 129 ZAR1-SUB. Full-length amino acid sequences of ZAR1-SUB were aligned by MAFFT version 7.

Supplementary table 3—source data 1. Amino acid sequences of 11 ZAR1-CIN. This file contains 11 ZAR1-CIN amino acid sequences identified from computational pipeline in Figure 1A.

Supplementary table 3—source data 2. Amino acid alignment file of 11 ZAR1-CIN. Full-length amino acid sequences of ZAR1-CIN were aligned by MAFFT version 7.