

Short title
Vector system for *in vivo* ZAR1 resistosome study

A vector system for fast-forward studies of the HOPZ-ACTIVATED RESISTANCE1 (ZAR1) resistosome in the model plant *Nicotiana benthamiana*

Adeline Harant¹, Hsuan Pai¹, Toshiyuki Sakai^{1,2}, Sophien Kamoun^{1*}, Hiroaki Adachi^{1,3*}

¹The Sainsbury Laboratory, University of East Anglia, Norwich Research Park, Norwich NR4 7UH, UK.

²Laboratory of Crop Evolution, Graduate School of Agriculture, Kyoto University, Mozume, Muko, Kyoto 617-0001, Japan.

³Graduate School of Biological Sciences, Nara Institute of Science and Technology, Ikoma 630-0192, Japan.

*For correspondence: adachi.hiroaki@bs.naist.jp

Senior authors: Hiroaki Adachi and Sophien Kamoun

One sentence summary: An open source collection of 72 Golden Gate compatible plasmids accelerates discovery by enabling rapid functional and comparative studies of immune receptor complexes

The author responsible for distribution of materials integral to the findings presented in this article in accordance with the policy described in the Instructions for Authors (<https://academic.oup.com/plphys/pages/General-Instructions>) is Hiroaki Adachi (adachi.hiroaki@bs.naist.jp).

AUTHOR CONTRIBUTIONS

S.K. and H.A. designed the research, supervised the work and wrote the paper; A.H., H.P., T.S and H.A. performed the experiments and analyzed the data. All authors read and edited the manuscript.

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ABSTRACT

Nicotiana benthamiana has emerged as a complementary experimental system to *Arabidopsis thaliana*. It enables fast-forward *in vivo* analyses primarily through transient gene expression and is particularly popular in the study of plant immunity. Recently, our understanding of nucleotide-binding leucine-rich repeat (NLR) plant immune receptors has greatly advanced following the discovery of the *Arabidopsis* HOPZ-ACTIVATED RESISTANCE1 (ZAR1) resistosome. Here, we describe a vector system of 72 plasmids that enables functional studies of the ZAR1 resistosome in *N. benthamiana*. We showed that ZAR1 stands out among the coiled coil class of NLRs (CC-NLRs) for being highly conserved across distantly related dicot plant species and confirmed NbZAR1 as the *N. benthamiana* ortholog of *Arabidopsis* ZAR1. Effector-activated and autoactive NbZAR1 trigger the cell death response in *N. benthamiana* and this activity is dependent on a functional N-terminal α 1 helix. C-terminally tagged NbZAR1 remains functional in *N. benthamiana*, thus enabling cell biology and biochemical studies in this plant system. We conclude that

the NbZAR1 open source pZA plasmid collection forms an additional experimental system to Arabidopsis for *in planta* resistosome studies.

INTRODUCTION

Nicotiana benthamiana has developed into a popular model system in plant biology, particularly for the study of plant immunity. The appeal of *N. benthamiana* as an experimental system is primarily due to agroinfiltration, a rapid transient protein expression method that enables cell biology, biochemistry, protein–protein interaction analyses and other *in vivo* experiments (Goodin et al., 2008). In addition, *N. benthamiana* is a member of the asterid group of flowering plants and as such is relatively distant from the classic model plant *Arabidopsis thaliana*, which belongs to the rosid group. Therefore, *N. benthamiana* is a complementary experimental system to Arabidopsis that also allows for a broader perspective into the evolution of molecular mechanisms in dicot plants.

Plants use nucleotide-binding leucine-rich repeat (NLR) proteins to mount innate immunity against invading pathogens. NLRs are intracellular receptors that detect host cell-translocated pathogen effectors and activate a hypersensitive cell death response that is a hallmark of plant immunity. The recent elucidation of the structures of the inactive and active complexes formed by the Arabidopsis NLR protein ZAR1 (HOPZ-ACTIVATED RESISTANCE1) with its partners receptor-like cytoplasmic kinases (RLCKs) is a major breakthrough in understanding how these immune receptors are activated and function. Activated ZAR1 forms a wheel-like pentamer, termed a resistosome, that undergoes a conformational switch—the death switch—to expose a funnel-shaped structure formed by the N-terminal $\alpha 1$ helices (Wang et al., 2019a; 2019b). The funnel of the ZAR1 resistosome forms a pore on the plasma membrane and the resistosome itself acts as a Ca^{2+} channel (Bi et al., 2021). The N-terminal $\alpha 1$ helix matches a sequence motif known as MADA motif that is functionally conserved across ~20% of coiled-coil (CC)-type plant NLRs (Adachi et al., 2019). This indicates that the ZAR1 death switch mechanism may be widely conserved across plant species. Mutations in surface-exposed residues within the $\alpha 1$ helix/MADA motif impair cell death and disease resistance mediated by ZAR1 and other MADA-CC-NLRs (Wang et al., 2019b; Adachi et al., 2019; Hu et al., 2020).

Here, we describe a Golden Gate compatible vector system (pZA) that enables functional studies of the ZAR1 resistosome in *N. benthamiana*. Golden Gate enables modular cloning through assembly of DNA fragments in one-step Type IIS restriction enzyme reactions. Consistent with previous reports (Baudin et al. 2017; 2019; Schultink et al. 2019), we confirmed that Arabidopsis ZAR1 (AtZAR1) does not cause hypersensitive cell death in *N. benthamiana* when expressed on its own and that *N. benthamiana* carries an ortholog of ZAR1 (NbZAR1). We developed and validated the pZA plasmid collection and share it as an open source resource that should prove useful for functional and comparative studies of NLR resistosomes in plants.

RESULTS AND DISCUSSION

ZAR1 is conserved across distantly related dicot plant species

Baudin et al. (2017) previously reported two orthologs of Arabidopsis ZAR1 and named them NbZAR1 and NbZAR2. We confirmed their findings by performing phylogenetic analyses of NLR proteins using the NB-ARC (nucleotide-binding adaptor shared by APAF-1, certain *R* gene products and CED-4) domain from 5 dicot plant species (Arabidopsis, cassava (*Manihot esculenta*), sugar beet (*Beta vulgaris* ssp. *vulgaris* var. *altissima*), tomato (*Solanum lycopersicum*) and *N. benthamiana*) representing three major clades of dicot plants (rosids, asterids and Caryophyllales). The resulting phylogenetic tree included 829 NLRs, which expectedly clustered into the well-defined NLR classes (Fig. 1A). Arabidopsis ZAR1 (AtZAR1, AT3G50950) fell within a well-supported CC-NLR subclade that included NbZAR1 (NbS00000462g0011), NbZAR2 (NbS00013646g0007) and one gene each from the three other dicot plant species (Fig. 1A). As previously reported (Baudin et al. 2017), NbZAR2 is truncated in the reference *N. benthamiana* genome (Supplemental Fig. S1). The other four genes, NbZAR1, Manes.02G097400 (MeZAR1), Bv9_203570 geqf.t1 (BvZAR1) and Solyc02g084890 (SIZAR1), code for CC-NLR proteins of similar length to AtZAR1, suggesting them as ZAR1 orthologs in each dicot plant species (Supplemental Fig. S1).

Based on our experience, ZAR1 seemed atypical among NLRs for having well-defined orthologs across unrelated dicot species. To directly evaluate ZAR1 conservation relative to other CC-NLRs, we calculated the phylogenetic (patristic) distance between each of the 47 Arabidopsis CC-NLRs and their closest gene from the other plant species based on the phylogenetic tree (Fig. 1A). Interestingly, AtZAR1 stood out among the CC-NLRs as having the shortest phylogenetic distance to its four orthologs (Fig. 1B). Similarly, in a reverse analysis where we plotted the phylogenetic distance between each of the 153 *N. benthamiana* CC-NLRs to their closest gene from the other species, NbZAR1 again stood out as being exceptionally conserved across all five examined species relative to other *N. benthamiana* CC-NLRs (Supplemental Fig. S2). Multiple alignments of the ZAR1 orthologs from the examined species confirmed the relatively high conservation of ZAR1 with pair-wise sequence similarities ranging from 58% to 86% across the full NLR protein (Supplemental Fig. S1). Overall, these analyses confirm NbZAR1 as the *N. benthamiana* ortholog of AtZAR1 and reveal ZAR1 as possibly the most widely conserved CC-NLR in dicot plants.

Golden Gate compatible plasmids for *in vivo* ZAR1 resistosome studies

The Golden Gate cloning system enables rapid and high throughput assembly of multiple sequence modules, such as promotor, terminator or tags into a common vector (Patron et al., 2015). To facilitate *in vivo* resistosome studies using this cloning strategy, we first introduced synonymous mutations to *Bpil* restriction enzyme sites in the coding sequence of AtZAR1 and NbZAR1 to make them both Golden Gate compatible. Next, we generated a synthetic version of NbZAR1, termed NbZAR1^{syn}, for genetic complementation assays in NbZAR1 silencing plants. Shuffled synonymous codon sequences can avoid virus-induced and hairpin RNA-mediated gene silencing (Wu et al., 2016; Derevnina et al., 2021). We cloned the genes into Golden Gate compatible vectors (see Level 0 in Table 1). Each coding sequence was flanked by *BsaI* restriction enzyme sites and overhang sequences for optimal Golden Gate reactions (Supplemental Fig. S3). Therefore, the Level 0 vectors can be used to transfer genes into a variety of *Agrobacterium tumefaciens* binary vectors for *in planta* protein expression.

To activate ZAR1 in the *N. benthamiana* system, we cloned an RLCK XII protein *NbJIM2* (XOPJ4 IMMUNITY 2) and *Xanthomonas perforans* effector *XopJ4* with Golden Gate compatible sequences (Table 1). *NbJIM2* is an endogenous partner RLCK associating with *NbZAR1* and is required for *XopJ4* recognition in *N. benthamiana* (Schultink et al. 2019). As an alternative strategy for activating ZAR1, we generated autoactive ZAR1 by mutating the MHD motif, which is conserved across ZAR1 orthologs (Supplemental Fig. S1). Substitution from aspartic acid (D) to valine (V) in the MHD motif generally makes full-length NLRs autoactive (Tameling et al., 2006). MHD mutants of *AtZAR1*, *NbZAR1* and *NbZAR1^{syn}* (*AtZAR1^{D489V}*, *NbZAR1^{D481V}* and *NbZAR1^{syn-D481V}*) were cloned into Level 0 vectors (Table 1).

We also generated an inactive version of ZAR1 by substituting leucine 17 to glutamic acid mutation (L17E) in the N-terminal MADA motif, since mutations in the N-terminal MADA motif often impair NLR cell death activity (Supplemental Fig. S1; Wang et al., 2019b; Adachi et al., 2019). We generated a MADA mutant series in wild-type and MHD mutant backgrounds, namely MADA mutants (*AtZAR1^{L17E}*, *NbZAR1^{L17E}* and *NbZAR1^{syn-L17E}*) and MADA/MHD mutants (*AtZAR1^{L17E/D489V}*, *NbZAR1^{L17E/D481V}* and *NbZAR1^{syn-L17E/D481V}*) (Level 0 in Table 1).

We assembled all of the *ZAR1* genes into binary vector plasmids without epitope tags or with the C-terminal tags 4xMyc, 6xHA and eGFP (pZA17-48 and pZA57-68 in Table 1). We provide this collection of 72 plasmids (Table 1) as an open access resource for *in planta* ZAR1 resistosome studies.

Golden Gate-based genetic complementation assay evaluates effector-activated ZAR1 cell death in *N. benthamiana*

To validate constructs generated by the Golden Gate assembly, we first tested whether *XopJ4*-triggered cell death is dependent on *NbZAR1* in the *N. benthamiana* transient expression system. We used a hairpin-silencing construct (RNAi:*NbZAR1*), that mediates silencing of *NbZAR1* when transiently expressed in *N. benthamiana* leaves (Supplemental Fig. S4). Silencing of *NbZAR1* suppressed cell death triggered by *XopJ4*, compared to the RNAi:GUS silencing control (Fig. 2A and 2B). We further performed an agroinfiltration complementation assay using a silencing-resilient version of *NbZAR1* (*NbZAR1^{syn}*). Unlike the original *NbZAR1*, *NbZAR1^{syn}* complemented the cell death response triggered by *XopJ4* in *NbZAR1* silencing leaves (Fig. 2A and 2B). Consistent with this genetic complementation phenotype, *NbZAR1^{syn}* was expressed in the RNAi:*NbZAR1* background at a similar level to the RNAi:GUS control (Fig. 2C). These results indicate that our construct series enables functional studies of the ZAR1 resistosome in the *N. benthamiana* transient expression system.

L17E mutation in *NbZAR1* N-terminal $\alpha 1$ helix impairs *XopJ4*-triggered cell death in *N. benthamiana*

Given that mutations in surface-exposed residues within the N-terminal $\alpha 1$ helix/MADA motif impair cell death by Arabidopsis ZAR1 (Wang et al., 2019b), we hypothesized that the intact $\alpha 1$ helix is required for *NbZAR1*-mediated cell death in response to *XopJ4*. To test this hypothesis, we used a MADA mutant *NbZAR1^{syn-L17E}* in the complementation assay of RNAi:*NbZAR1*. Unlike *NbZAR1^{syn}*, *NbZAR1^{syn-L17E}* did not complement the *XopJ4*-triggered cell

death response in RNAi:NbZAR1 leaves (Fig. 2A and 2B). We confirmed that NbZAR1^{syn-L17E} protein was expressed in NbZAR1 silencing leaves (Fig. 2C). Thus, we conclude that similar to Arabidopsis ZAR1, the N-terminal α 1 helix/MADA motif in activated NbZAR1 mediates the cell death response in *N. benthamiana*.

Unlike Arabidopsis ZAR1, NbZAR1 triggers autoimmune cell death in *N. benthamiana*

Next, we investigated the effect of a MHD mutation on AtZAR1 and NbZAR1 autoimmunity in the *N. benthamiana* system (Fig. 3A). As recently reported by Baudin et al. (2019), AtZAR1 did not trigger macroscopic cell death in *N. benthamiana* leaves even when the full-length protein carries the D489V mutation in the MHD motif (Fig. 3B and 3C). In contrast, NbZAR1 caused macroscopic cell death when the MHD mutant NbZAR1^{D481V} was expressed in *N. benthamiana* leaves (Fig. 3D and 3E). NRC4 (NLR required for cell death 4) was used as a control MADA-CC-NLR, because its MHD mutant NRC4^{D478V} causes autoactive cell death in *N. benthamiana* leaves (Adachi et al., 2019). Our results indicate that unlike Arabidopsis ZAR1, full-length NbZAR1 can be made autoactive in *N. benthamiana* through mutation of the MHD motif. These results are consistent with a previous report that AtZAR1 does not complement NbZAR1 cell death activity when co-expressed with XopJ4/JIM2 (Schultink et al. 2019). We conclude that unlike AtZAR1, NbZAR1 is probably compatible with endogenous RLCK and can therefore display autoactivity when expressed on its own in *N. benthamiana* leaves.

The endogenous RLCK partner JIM2 is required for NbZAR1 autoimmune cell death

To test the hypothesis that the endogenous RLCK partner is required for NbZAR1 autoactivity (Schultink et al. 2019), we used RNAi:NbJIM2 (Supplemental Fig. S4) and coexpressed the hairpin silencing construct with autoactive NbZAR1^{D481V}. Compared to the RNAi:GUS silencing control, RNAi:NbJIM2 significantly reduced cell death by NbZAR1^{D481V} (Supplemental Fig. S5). Moreover, a synthetic version of NbJIM2 with a codon shuffled sequence (NbJIM2^{syn}) restored the NbZAR1^{D481V} autoimmune cell death response (Supplemental Fig. S5). Although a pathogen effector was not used to activate NbZAR1, this experiment shows that autoactive NbZAR1 is dependent on its partner RLCK NbJIM2. This indicates that NbZAR1^{D481V} may require an RLCK to form a signaling competent conformation like the AtZAR1 resistosome (Wang et al., 2019a; 2019b).

NbZAR1 autoimmune cell death is dependent on its N-terminal α 1 helix

Next, we tested whether NbZAR1 requires an intact N-terminal α 1 helix/MADA motif for triggering autoactive cell death. We expressed the double MADA/MHD mutant NbZAR1^{L17E/D481V} in *N. benthamiana* leaves using agroinfiltration and observed that the L17E mutation significantly reduced the cell death activity compared to NbZAR1^{D481V} (Fig. 3D and 3E). This result indicates that an intact N-terminal α 1 helix is necessary for NbZAR1^{D481V} autoactivity. Given that the α 1 helix forms a funnel-shaped structure that is a key structural component of the activated ZAR1 resistosome and its translocation into the plasma membrane (Wang et al., 2019b), we conclude that the NbZAR1^{D481V} experimental system recapitulates the resistosome model in *N. benthamiana*.

C-terminally tagged NbZAR1 is functional in *N. benthamiana*

N. benthamiana agroinfiltration is a powerful method for *in vivo* cell biology, biochemistry and protein–protein interaction analyses, but this experimental system requires that the target protein remains functional and intact after fusion to an epitope or fluorescent tag (Goodin et al., 2008; Bally et al., 2018). We therefore set out to determine the degree to which NbZAR1 can tolerate protein tags (Table 1). We focused on C-terminal tags, given that N-terminal tag fusion is known to interfere with the AtZAR1 cell death activity (Wang et al., 2019b). We tested NbZAR1^{D481V} constructs fused to three different tags, 4xMyc, 6xHA and eGFP by agroinfiltration. In all three cases, NbZAR1^{D481V} retained the capacity to trigger autoactive cell death although the response was weaker than the non–epitope-tagged NbZAR1 positive control (Fig. 4A to 4F). We also showed by western blot analyses that these C-terminally tagged NbZAR1 accumulated at detectable levels in *N. benthamiana* leaves and generally formed a single band indicative of minimal protein degradation (Fig. 4G to 4I). We conclude that C-terminal tagging of NbZAR1 doesn't dramatically interfere with the activity and integrity of this NLR protein.

Towards *in vivo* studies of the ZAR1 resistosome

We conclude that the open source pZA plasmid resource we developed can serve as valuable reagents for *in vivo* resistosome studies in *N. benthamiana* that complements the Arabidopsis system and provides a foundation for future studies. Effector-activated and autoactive NbZAR1 can be readily tested by agroinfiltration in *N. benthamiana* leaves and assayed for cell death activity. This indicates the effectiveness of our plasmid resource to monitor the activated state of ZAR1 in the *N. benthamiana* system. We also note one advantage of the ZAR1 MADA mutants, which are presumably activated but unable to trigger cell death. This mutation strategy enables protein purification or microscopic observation of effector-activated/autoactive ZAR1 using plant tissues without cell death response, that generally perturbs biochemical and cell biology studies. Since NbZAR1^{D481V} tolerates different C-terminal tags and each variant protein expresses well in *N. benthamiana*, the pZA plasmid resource can be applied for *in planta* cell biology, biochemical and protein-protein interactions studies of the ZAR1 resistosome.

The pZA toolkit complements existing resources in Arabidopsis. Of particular interest is the visualization of activated NLR oligomers *in vivo*. Recently, an ~900 kDa complex associated with activated AtZAR1 was detected in Blue Native polyacrylamide gel electrophoresis (BN-PAGE) analysis using Arabidopsis protoplasts (Hu et al., 2020). Bi et al. (2021) monitored subcellular dynamics of AtZAR1 resistosomes on the plasma membrane in Arabidopsis protoplasts under total internal reflection fluorescence microscopy. *N. benthamiana* agroinfiltration offers a rapid and effective alternative expression system to the Arabidopsis protoplast system used by Hu et al. (2020) and Bi et al. (2021) that can be explored in the future.

N. benthamiana is rapidly developing as an excellent experimental system to study oligomeric complexes by different types of NLRs. Jacob et al. (2021) detected oligomer formation of a CC_R-NLR (RPW8-type CC-NLR) AtNRG1.1 (N REQUIREMENT GENE 1.1) in BN-PAGE when they expressed an AtNRG1.1 autoactive MHD mutant in *N. benthamiana* leaves. Imaging analysis showed AtNRG1.1 MHD mutant forms punctate structures on the plasma membrane of *N. benthamiana* epidermal cells (Jacob et al., 2021). Martin et al. (2020)

solved a cryo-electron microscopy structure of activated TIR-NLR (Toll/interleukin-1 receptor-NLR) ROQ1 (RECOGNITION OF XOPQ 1) by purifying activated ROQ1 protein from *N. benthamiana* leaves. Therefore, the *N. benthamiana* system has potential as a protein expression platform for structural studies of both resting and activated states of NLRs *in vivo*.

Here we describe that ZAR1 stands out among CC-NLRs for being widely conserved across dicot plants. Recently, we expanded on these analyses and reconstructed the evolutionary history of ZAR1 across angiosperms (Adachi et al., 2021). Notably, ZAR1 orthologs were identified not only from dicot plant species but also from a monocot, taro (*Colocasia esculenta*). Thus, we conclude that the origin of ZAR1 traces back to early flowering plant lineages ~220 to 150 million years ago (Jurassic period). However, it still remains unknown how ZAR1 orthologs vary in their molecular and biochemical activities among flowering plant species. The pZA system enables evolutionary studies of the ZAR1 resistosome. Further comparative analyses between AtZAR1, NbZAR1 and other orthologs would help answer questions about the molecular evolution of the resistosome as a mechanism to trigger NLR-mediated immunity.

MATERIALS AND METHODS

Plant growth conditions

Wild type *Nicotiana benthamiana* was propagated in a glasshouse and, for most experiments, grown in a controlled growth chamber with temperatures of 22-25°C, humidity 45-65% and a 16/8-h light/dark cycle.

Bioinformatic and phylogenetic analyses

We used NLR-parser (Steuernagel et al., 2015) to identify NLR sequences from the protein databases of tomato (Sol Genomics Network, Tomato ITAG release 2.40), *N. benthamiana* (Sol Genomics Network, *N. benthamiana* Genome v0.4.4), *Arabidopsis thaliana* (<https://www.araport.org/>, Araport11), cassava (*Manihot esculenta*) (<https://phytozome-next.jgi.doe.gov/>, Phytozome v13, Manihot esculenta v7.1) and sugar beet (*Beta vulgaris* ssp. *vulgaris* var. *altissima*) (<http://bvseq.molgen.mpg.de/index.shtml>, RefBeet-1.2). The obtained NLR sequences from NLR-parser, were aligned using MAFFT v. 7 (Katoh and Standley, 2013), and the protein sequences that lacked the p-loop motif were discarded to make the NLR dataset (Supplemental File S1). The gaps in the alignments were deleted manually in MEGA7 (Kumar et al., 2016) and the NB-ARC domains were used for generating phylogenetic trees (Supplemental File S2). The neighbour-joining tree was made using MEGA7 with JTT model and bootstrap values based on 100 iterations (Supplemental File S3). Amino acid sequences of ZAR1 orthologs and the percent identity are listed in Supplemental File S4 and Supplemental File S5.

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 (Supplemental File S3) 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_plot).

Plasmid constructions

For Golden Gate cloning system, wild-type and MHD mutant clones of *AtZAR1* and *NbZAR1* were synthesized through GENEWIZ Standard Gene Synthesis with synonymous mutations to *BpiI* restriction enzyme sites. The synthetic fragment of *NbZAR1* was designed manually to introduce synonymous substitution by following *N. benthamiana* codon usage. The fragment was synthesized through GENEWIZ Standard Gene Synthesis and then subcloned into pICSL01005 (Level 0 acceptor for CDS no stop modules, Addgene no.47996) or pICH41308 (Level 0 acceptor for CDS modules, Addgene no.47998) together with the remaining *NbZAR1* fragment to generate full-length *NbZAR1^{syn}*. Primers used for amplifying the *NbZAR1* fragment are listed in Supplemental Table S1.

To generate MADA motif mutants of *AtZAR1* and *NbZAR1*, the leucine (L) 17 in the MADA motif was substituted to glutamic acid (E) by site-directed mutagenesis using Phusion High-Fidelity DNA Polymerase (Thermo Fisher Scientific). To make *AtZAR1* and *NbZAR1* clones for C-terminal tagging, the stop codons were removed by PCR using Phusion High-Fidelity DNA Polymerase (Thermo Fisher Scientific). All wild-type and mutant coding sequences of *AtZAR1* and *NbZAR1* were cloned into pCR8/GW/D-TOPO (Invitrogen) as level 0 modules (Weber et al., 2011). Primers used for the site-directed mutagenesis and removing the stop codon are listed in Supplemental Table S1.

The level 0 plasmids of *AtZAR1*, *NbZAR1* and *NbZAR1^{syn}* were used for Golden Gate assembly with or without C-terminal tag modules, pICSL50009 (6xHA, Addgene no. 50309), pICSL50010 (4xMyc, Addgene no. 50310) or pICSL50034 (eGFP, TSL SynBio) into the binary vector pICH86988 (Engler et al., 2014).

Golden Gate compatible *XopJ4* and *NbJIM2^{syn}* clones were synthesized through GENEWIZ Standard Gene Synthesis and were subcloned into the pICH86988 vector. pICSL50007 (3xFLAG, Addgene no. 50308) was used to make the C-terminal 3xFLAG fusion construct of *NbJIM2^{syn}*.

To prepare the RNAi construct, the silencing fragments were amplified from *NbZAR1* and *NbJIM2* cDNA by Phusion High-Fidelity DNA Polymerase (Thermo Fisher Scientific) using primers listed in Supplemental Table S1. The purified amplicons were cloned into the pRNAi-GG vector (Yan et al., 2012).

Transient gene-expression and cell death assays

Transient gene expression in *N. benthamiana* was performed by agroinfiltration according to methods described by Bos et al. (2006). Briefly, four-weeks old *N. benthamiana* plants were infiltrated with *A. tumefaciens* Gv3101 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).

To perform genetic complementation assays in RNAi:NbZAR1 and RNAi:NbJIM2 backgrounds, we initially infiltrated *A. tumefaciens* strains carrying RNAi constructs ($OD_{600} = 0.5$). At one day after the first agroinfiltration, we performed second agroinfiltration using mixtures of two different *A. tumefaciens* strains to complement NbZAR1/NbJIM2 expression and overexpress cell death triggering genes. In this second agroinfiltration, we adjusted the final concentration of each *A. tumefaciens* suspension to $OD_{600} = 0.3$.

RNA extraction and semi-quantitative RT-PCR

Plant total RNA was extracted using RNeasy Mini Kit (Qiagen). 500 ng RNA of each sample was subjected to reverse transcription using SuperScript® IV Reverse Transcriptase (Thermo Fisher Scientific). Semi-quantitative reverse transcription PCR (RT-PCR) was performed using DreamTaq (Thermo Fisher Scientific) with 22, 32 and 36 amplification cycles for *EF-1α*, *NbJIM2* and *NbZAR1*, respectively. Primers used for RT-PCR are listed in Supplemental Table S1.

Protein immunoblotting

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% (v/v) 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% (v/v) 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), c-Myc (9E10) HRP (Santa Cruz Biotech) or GFP (B-2) 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 Scientific).

Plasmid distribution

All pZA plasmids are available through Addgene through the Kamoun Lab page https://www.addgene.org/Sophien_Kamoun/.

Statistical analyses

HR indexes were visualized as a violin plot using the "ggplot2" package (Wickham, 2016). Tukey's HSD test was applied to HR indexes using the "TukeyHSD" function in R (version 3.6).

Accession Numbers

DNA sequence data used in this study can be found in reference genome or GenBank/EMBL databases under the following accession numbers: AtZAR1 (AT3G50950, NP_190664.1), NbZAR1 (NbS00000462g0011, AXY05280.1), NbZAR2 (NbS00013646g0007), SIZAR1 (Solyc02g084890, XP_004232433.1), MeZAR1 (Manes.02G097400, P_021604862.1), BvZAR1 (Bv9_203570_geqf.t1, XP_010688776.1), NbJIM2 (NbS00009407g0011, MH532571.1), XopJ4 (AF221058.1).

Supplemental Material

Supplemental Figure S1. Sequence alignment of full-length ZAR1 ortholog proteins.

Supplemental Figure S2. NbZAR1 is highly conserved across dicot species.

Supplemental Figure S3. Golden Gate Level 0 modules of AtZAR1 and NbZAR1 genes.

Supplemental Figure S4. *NbZAR1* and *NbJIM2* silencing in *N. benthamiana*.

Supplemental Figure S5. NbJIM2 is required for NbZAR1 autoactive cell death in *N. benthamiana*.

Supplemental Table S1. Primers Used for generating Golden Gate constructs in this study.

Supplemental File S1. Amino acid sequences of full-length NLRs in the NLR dataset.

Supplemental File S2. Amino acid sequences for NLR phylogenetic tree.

Supplemental File S3. NLR phylogenetic tree file.

Supplemental File S4. Amino acid sequences of ZAR1 orthologs.

Supplemental File S5. Percent identity matrix of ZAR1 orthologs by Clustal2.1.

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DECLARATION OF INTERESTS

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

472 **Table 1.** Golden Gate Plasmid resources prepared in this study.

Plasmids	Relevant features	Bacterial selection	Usage
pZA1	Golden Gate compatible (GG_) wild-type AtZAR1 with STOP codon	Spc ^r	Level 0
pZA2	GG_AtZAR1 ^{D489V} with STOP codon	Spc ^r	Level 0
pZA3	GG_AtZAR1 ^{L17E} with STOP codon	Spc ^r	Level 0
pZA4	GG_AtZAR1 ^{L17E/D489V} with STOP codon	Spc ^r	Level 0
pZA5	GG_wild-type NbZAR1 with STOP codon	Spc ^r	Level 0
pZA6	GG_NbZAR1 ^{D481V} with STOP codon	Spc ^r	Level 0
pZA7	GG_NbZAR1 ^{L17E} with STOP codon	Spc ^r	Level 0
pZA8	GG_NbZAR1 ^{L17E/D481V} with STOP codon	Spc ^r	Level 0
pZA9	GG_wild-type AtZAR1 without STOP codon	Spc ^r	Level 0
pZA10	GG_AtZAR1 ^{D489V} without STOP codon	Spc ^r	Level 0
pZA11	GG_AtZAR1 ^{L17E} without STOP codon	Spc ^r	Level 0
pZA12	GG_AtZAR1 ^{L17E/D489V} without STOP codon	Spc ^r	Level 0
pZA13	GG_wild-type NbZAR1 without STOP codon	Spc ^r	Level 0
pZA14	GG_NbZAR1 ^{D481V} without STOP codon	Spc ^r	Level 0
pZA15	GG_NbZAR1 ^{L17E} without STOP codon	Spc ^r	Level 0
pZA16	GG_NbZAR1 ^{L17E/D481V} without STOP codon	Spc ^r	Level 0
pZA17	GG_wild-type AtZAR1, Km ^r plant selection marker	Km ^r	Level 2
pZA18	GG_AtZAR1 ^{D489V} , Km ^r plant selection marker	Km ^r	Level 2
pZA19	GG_AtZAR1 ^{L17E} , Km ^r plant selection marker	Km ^r	Level 2
pZA20	GG_AtZAR1 ^{L17E/D489V} , Km ^r plant selection marker	Km ^r	Level 2
pZA21	GG_wild-type NbZAR1, Km ^r plant selection marker	Km ^r	Level 2
pZA22	GG_NbZAR1 ^{D481V} , Km ^r plant selection marker	Km ^r	Level 2
pZA23	GG_NbZAR1 ^{L17E} , Km ^r plant selection marker	Km ^r	Level 2
pZA24	GG_NbZAR1 ^{L17E/D481V} , Km ^r plant selection marker	Km ^r	Level 2

Spc^r: spectinomycin-resistant, Km^r: kanamycin-resistant.

475 **Table 1.** Continued.

Plasmids	Relevant features	Bacterial selection	Usage
pZA25	GG_wild-type AtZAR1-4xMyc, Km ^r plant selection marker	Km ^r	Level 2
pZA26	GG_AtZAR1 ^{D489V} -4xMyc, Km ^r plant selection marker	Km ^r	Level 2
pZA27	GG_AtZAR1 ^{L17E} -4xMyc, Km ^r plant selection marker	Km ^r	Level 2
pZA28	GG_AtZAR1 ^{L17E/D489V} -4xMyc, Km ^r plant selection marker	Km ^r	Level 2
pZA29	GG_wild-type NbZAR1-4xMyc, Km ^r plant selection marker	Km ^r	Level 2
pZA30	GG_NbZAR1 ^{D481V} -4xMyc, Km ^r plant selection marker	Km ^r	Level 2
pZA31	GG_NbZAR1 ^{L17E} -4xMyc, Km ^r plant selection marker	Km ^r	Level 2
pZA32	GG_NbZAR1 ^{L17E/D481V} -4xMyc, Km ^r plant selection marker	Km ^r	Level 2
pZA33	GG_wild-type AtZAR1-6xHA, Km ^r plant selection marker	Km ^r	Level 2
pZA34	GG_AtZAR1 ^{D489V} -6xHA, Km ^r plant selection marker	Km ^r	Level 2
pZA35	GG_AtZAR1 ^{L17E} -6xHA, Km ^r plant selection marker	Km ^r	Level 2
pZA36	GG_AtZAR1 ^{L17E/D489V} -6xHA, Km ^r plant selection marker	Km ^r	Level 2
pZA37	GG_wild-type NbZAR1-6xHA, Km ^r plant selection marker	Km ^r	Level 2
pZA38	GG_NbZAR1 ^{D481V} -6xHA, Km ^r plant selection marker	Km ^r	Level 2
pZA39	GG_NbZAR1 ^{L17E} -6xHA, Km ^r plant selection marker	Km ^r	Level 2
pZA40	GG_NbZAR1 ^{L17E/D481V} -6xHA, Km ^r plant selection marker	Km ^r	Level 2
pZA41	GG_wild-type AtZAR1-eGFP, Km ^r plant selection marker	Km ^r	Level 2
pZA42	GG_AtZAR1 ^{D489V} -eGFP, Km ^r plant selection marker	Km ^r	Level 2
pZA43	GG_AtZAR1 ^{L17E} -eGFP, Km ^r plant selection marker	Km ^r	Level 2
pZA44	GG_AtZAR1 ^{L17E/D489V} -eGFP, Km ^r plant selection marker	Km ^r	Level 2
pZA45	GG_wild-type NbZAR1-eGFP, Km ^r plant selection marker	Km ^r	Level 2
pZA46	GG_NbZAR1 ^{D481V} -eGFP, Km ^r plant selection marker	Km ^r	Level 2
pZA47	GG_NbZAR1 ^{L17E} -eGFP, Km ^r plant selection marker	Km ^r	Level 2
pZA48	GG_NbZAR1 ^{L17E/D481V} -eGFP, Km ^r plant selection marker	Km ^r	Level 2
pZA49	GG_synthesized NbZAR1 (NbZAR1 ^{syn}) with STOP codon	Spc ^r	Level 0
pZA50	GG_NbZAR1 ^{syn-D481V} with STOP codon	Spc ^r	Level 0

476 Spc^r: spectinomycin-resistant, Km^r: kanamycin-resistant.

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478 **Table 1.** Continued.

Plasmids	Relevant features	Bacterial selection	Usage
pZA51	GG_NbZAR1 ^{syn-L17E} with STOP codon	Spc ^r	Level 0
pZA52	GG_NbZAR1 ^{syn-L17E/D481V} with STOP codon	Spc ^r	Level 0
pZA53	GG_NbZAR1 ^{syn} without STOP codon	Spc ^r	Level 0
pZA54	GG_NbZAR1 ^{syn-D481V} without STOP codon	Spc ^r	Level 0
pZA55	GG_NbZAR1 ^{syn-L17E} without STOP codon	Spc ^r	Level 0
pZA56	GG_NbZAR1 ^{syn-L17E/D481V} without STOP codon	Spc ^r	Level 0
pZA57	GG_NbZAR1 ^{syn} -4xMyc, Km ^r plant selection marker	Km ^r	Level 2
pZA58	GG_NbZAR1 ^{syn-D481V} -4xMyc, Km ^r plant selection marker	Km ^r	Level 2
pZA59	GG_NbZAR1 ^{syn-L17E} -4xMyc, Km ^r plant selection marker	Km ^r	Level 2
pZA60	GG_NbZAR1 ^{syn-L17E/D481V} -4xMyc, Km ^r plant selection marker	Km ^r	Level 2
pZA61	GG_NbZAR1 ^{syn} -6xHA, Km ^r plant selection marker	Km ^r	Level 2
pZA62	GG_NbZAR1 ^{syn-D481V} -6xHA, Km ^r plant selection marker	Km ^r	Level 2
pZA63	GG_NbZAR1 ^{syn-L17E} -6xHA, Km ^r plant selection marker	Km ^r	Level 2
pZA64	GG_NbZAR1 ^{syn-L17E/D481V} -6xHA, Km ^r plant selection marker	Km ^r	Level 2
pZA65	GG_NbZAR1 ^{syn} -eGFP, Km ^r plant selection marker	Km ^r	Level 2
pZA66	GG_NbZAR1 ^{syn-D481V} -eGFP, Km ^r plant selection marker	Km ^r	Level 2
pZA67	GG_NbZAR1 ^{syn-L17E} -eGFP, Km ^r plant selection marker	Km ^r	Level 2
pZA68	GG_NbZAR1 ^{syn-L17E/D481V} -eGFP, Km ^r plant selection marker	Km ^r	Level 2
pZA69	Intron-containing hairpin RNA (ihpRNA)-NbZAR1	Km ^r	RNAi
pZA70	ihpRNA-NbJIM2	Km ^r	RNAi
pZA71	GG_synthesized NbJIM2 (NbJIM2 ^{syn})-3xFLAG, Km ^r plant selection marker	Km ^r	Level 2
pZA72	GG_XopJ4, Km ^r plant selection marker	Km ^r	Level 2

Spc^r: spectinomycin-resistant, Km^r: kanamycin-resistant.

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Figure legends

Figure 1. ZAR1 is a conserved CC-NLR across distantly related dicot plant species. (A) ZAR1 forms a small subclade that is composed of orthologous genes from distantly related dicot plant species. The phylogenetic tree was generated in MEGA7 by the neighbour-joining method using NB-ARC domain sequences of 829 NLRs identified from *N. benthamiana* (NbS-), tomato (Soly-), sugar beet (Bv-), Arabidopsis (AT-) and cassava (Manes-) (left panel). The ZAR1 subclade phylogenetic tree is shown on the right panel. Each branch is marked with different colors based on plant species. Red arrow heads indicate bootstrap support > 0.7 and are shown for the relevant nodes. The scale bars indicate the evolutionary distance in amino acid substitution per site. Amino acid sequences of the full-length ZAR1 orthologs can be found in Supplemental Fig. S1. (B) AtZAR1 is highly conserved among dicot CC-NLRs. The phylogenetic (patristic) distances 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 colors 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 Supplemental Fig. S2.

Figure 2. The NbZAR1 MADA motif mediates XopJ4-triggered cell death in *N. benthamiana*. (A) Cell death observed in *N. benthamiana* leaves after expression of ZAR1 and XopJ4. The leaves were infiltrated with Agrobacterium strains carrying ZAR1 and XopJ4 overexpression constructs at 1 day after initial agroinfiltration of RNAi constructs. *N. benthamiana* leaf panels expressing non-epitope-tagged wild-type and variant proteins of NbZAR1 were photographed at 5 days after the second agroinfiltration. (B) Violin plots showing cell death intensity scored as an HR index based on three independent experiments. Statistical differences among the samples were analyzed with Tukey's HSD test ($p < 0.01$). (C) *In planta* protein accumulation of wild-type NbZAR1 and the variant. For anti-HA immunoblot, total proteins were prepared from *N. benthamiana* leaves at 2 days after the second agroinfiltration. Empty vector control is described as EV. Equal loading was checked with Reversible Protein Stain Kit (Thermo Fisher Scientific).

Figure 3. NbZAR1 triggers autoimmune cell death in *N. benthamiana* leaves and requires an intact MADA motif for autoactivity. (A) Schematic representation of substitution sites in AtZAR1, NbZAR1 and NRC4. Mutated sites are shown with red asterisks on the sequence alignment. (B and D) Cell death observed in *N. benthamiana* leaves after expression of ZAR1 mutants. *N. benthamiana* leaf panels expressing non-epitope-tagged wild-type and variant proteins of AtZAR1 and NbZAR1 were photographed at 5 days after agroinfiltration. (C and E) Violin plots showing cell death intensity scored as an HR index based on three independent experiments. Statistical differences among the samples were analyzed with Tukey's HSD test ($p < 0.01$).

Figure 4. C-terminal tagged NbZAR1 triggers autoimmune cell death in *N. benthamiana* leaves. (A-C) Cell death observed in *N. benthamiana* leaves after expression of ZAR1 mutants. *N. benthamiana* leaf panels expressing wild-type and variant proteins of NbZAR1 with C-terminal 4xMyc, 6xHA or eGFP tag were photographed at 5 days after agroinfiltration. (D-F) Violin plots showing cell death intensity scored as an HR index based on three independent experiments. Statistical differences among the samples were analyzed with Tukey's HSD test ($p < 0.01$). (G-I) *In planta* protein accumulation of AtZAR1 and NbZAR1

variants. For anti-Myc, anti-HA and anti-GFP immunoblots of AtZAR1, NbZAR1 and the mutant proteins, total proteins were prepared from *N. benthamiana* leaves at 2 days after agroinfiltration. Empty vector control is described as EV. Equal loading was checked with Reversible Protein Stain Kit (Thermo Fisher).

Supplemental Figure S1. Sequence alignment of full-length ZAR1 ortholog proteins. Amino acid sequences of ZAR1 orthologs were aligned by ClustalW program. MADA and MHD motif sequences are marked with red and orange dot boxes, respectively. Red asterisks indicate substitution sites for MADA and MHD mutations.

Supplemental Figure S2. NbZAR1 is highly conserved across dicot species. The phylogenetic (patristic) distances 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 Fig. 1A. The closest patristic distances are plotted with different colors based on plant species.

Supplemental Figure S3. Golden Gate Level 0 modules of AtZAR1 and NbZAR1 genes. (A) *AtZAR1* and *NbZAR1* wild-type and the mutant genes were cloned in the pUC8-Spec vector (pZA1-16). The plasmid map was made with Benchling software. (B) *ZAR1* genes were cloned with the stop codon in pZA1-8 plasmids. (C) *ZAR1* genes were cloned without the stop codon for C-terminal tag fusion (pZA9-16). *BsaI* restriction enzyme sites for Golden Gate assembly are shown as red characters. 5' and 3' overhang sequences after *BsaI* digestion are described as 5'OV and 3'OV, respectively.

Supplemental Figure S4. NbZAR1 and NbJIM2 silencing in *N. benthamiana*. Leaf samples were collected 2 days after agroinfiltration expressing *RNAi:GUS*, *RNAi:NbZAR1* and *RNAi:NbJIM2*. Total RNA was extracted from three independent plant samples (#1-3). The expression of *NbZAR1* and *NbJIM2* were analyzed in semi-quantitative RT-PCR using specific primer sets. Elongation factor 1 α (EF-1 α) was used as an internal control.

Supplemental Figure S5. NbJIM2 is required for NbZAR1 autoactive cell death in *N. benthamiana*. (A) Cell death observed in *N. benthamiana* leaves after expression of *NbJIM2* and autoactive *NbZAR1*. The leaves were infiltrated with *Agrobacterium* strains carrying *NbJIM2*^{syn} and *NbZAR1*^{DV} overexpression constructs at 1 day after initial agroinfiltration of *RNAi* constructs. *N. benthamiana* leaves were photographed at 5 days after the second agroinfiltration. (B) Violin plots showing cell death intensity scored as an HR index based on three independent experiments. Statistical differences among the samples were analyzed with Tukey's HSD test ($p < 0.01$).

Supplementary files

Supplemental Table S1. Primers Used for generating Golden Gate constructs in this study.

Supplemental File S1. Amino acid sequences of full-length NLRs in the NLR dataset. The 829 NLR sequences used in this study.

Supplemental File S2. Amino acid sequences for NLR phylogenetic tree. NB-ARC domain sequences used for phylogenetic analysis are shown with the IDs, *N. benthamiana* (NbS-), tomato (Solyc-), sugar beet (Bv-), Arabidopsis (AT-) and cassava (Manes-).

Supplemental File S3. NLR phylogenetic tree file. The phylogenetic tree was saved in newick file format.

Supplemental File S4. Amino acid sequences of ZAR1 orthologs. The N-terminal 28 amino acids (MAMRNQLYILESASQFPWAAAFQKSEAE) of SIZAR1 (Solyc02g084890) were removed due to potential mis-annotation.

Supplemental File S5. Percent identity matrix of ZAR1 orthologs by Clustal2.1.

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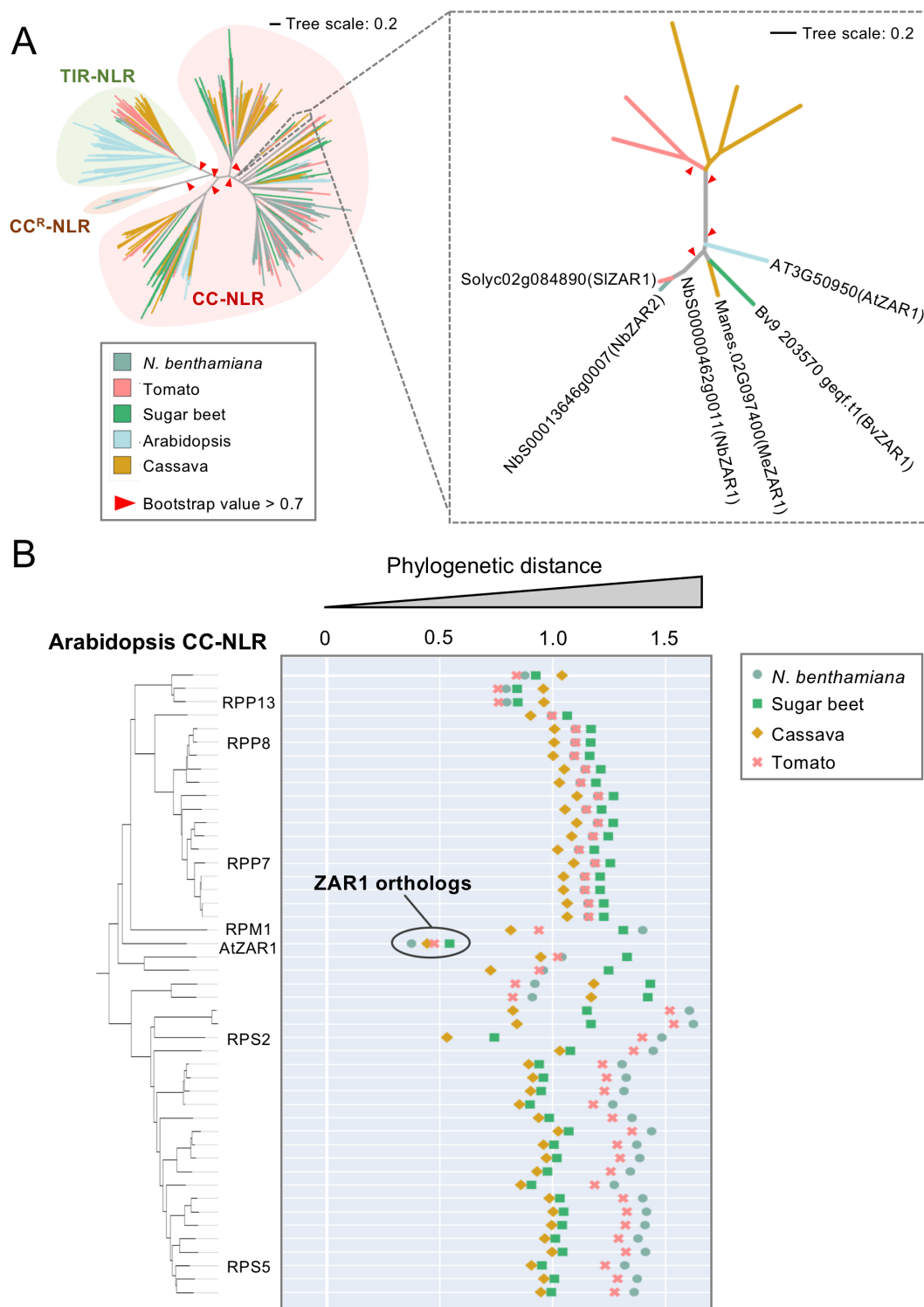


Figure 1. ZAR1 is a conserved CC-NLR across distantly related dicot plant species. (A) ZAR1 forms a small subclade that is composed of orthologous genes from distantly related dicot plant species. The phylogenetic tree was generated in MEGA7 by the neighbour-joining method using NB-ARC domain sequences of 829 NLRs identified from *N. benthamiana* (NbS-), tomato (Soly-), sugar beet (Bv-), Arabidopsis (AT-) and cassava (Manes-) (left panel). The ZAR1 subclade phylogenetic tree is shown on the right panel. Each branch is marked with different colours based on plant species. Red arrow heads indicate bootstrap support > 0.7 and is shown for the relevant nodes. The scale bars indicate the evolutionary distance in amino acid substitution per site. Amino acid sequences of the full-length ZAR1 orthologs can be found in Supplemental Fig. S1. (B) AtZAR1 is highly conserved among dicot CC-NLRs. 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 Supplemental Fig. S2.

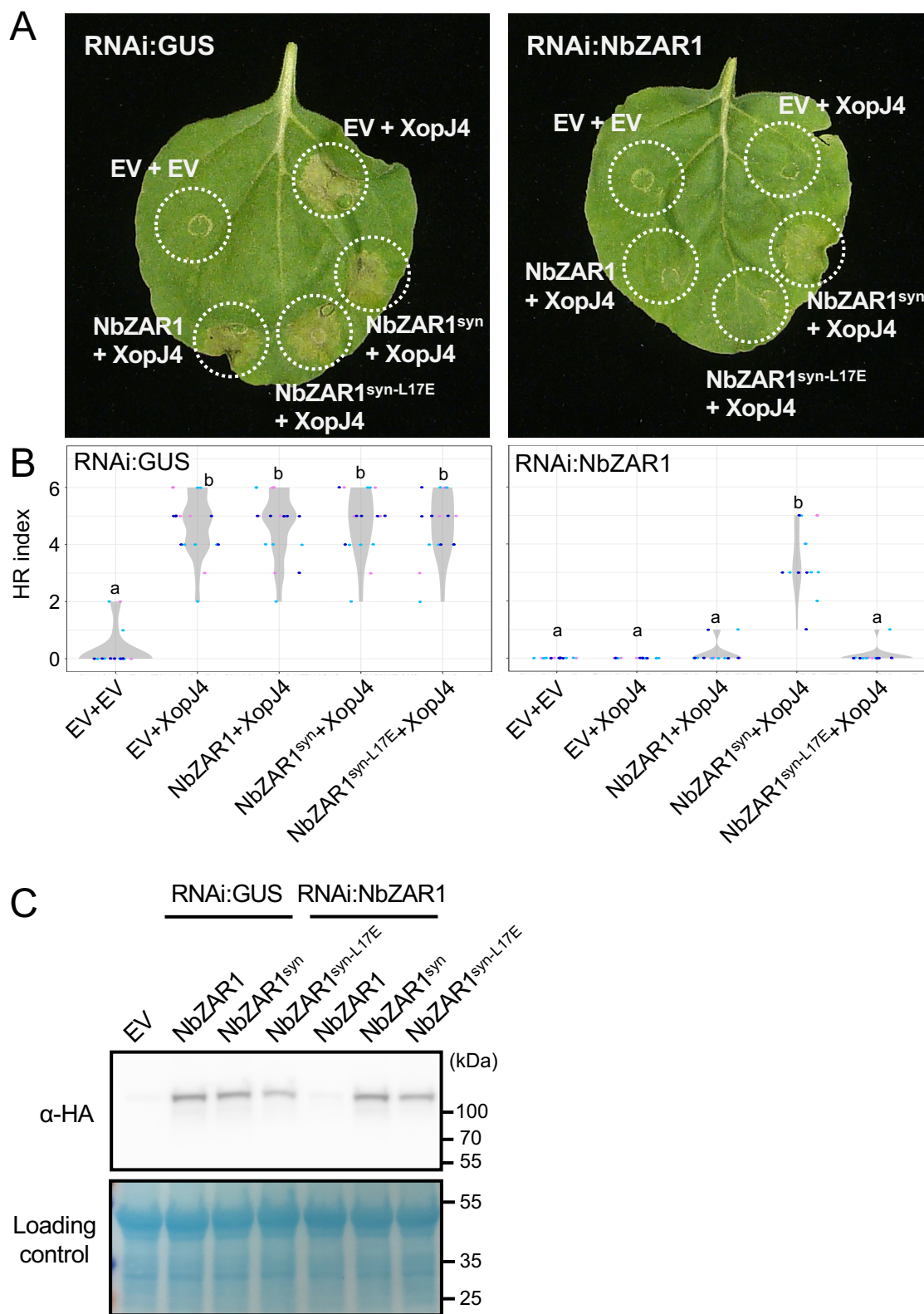


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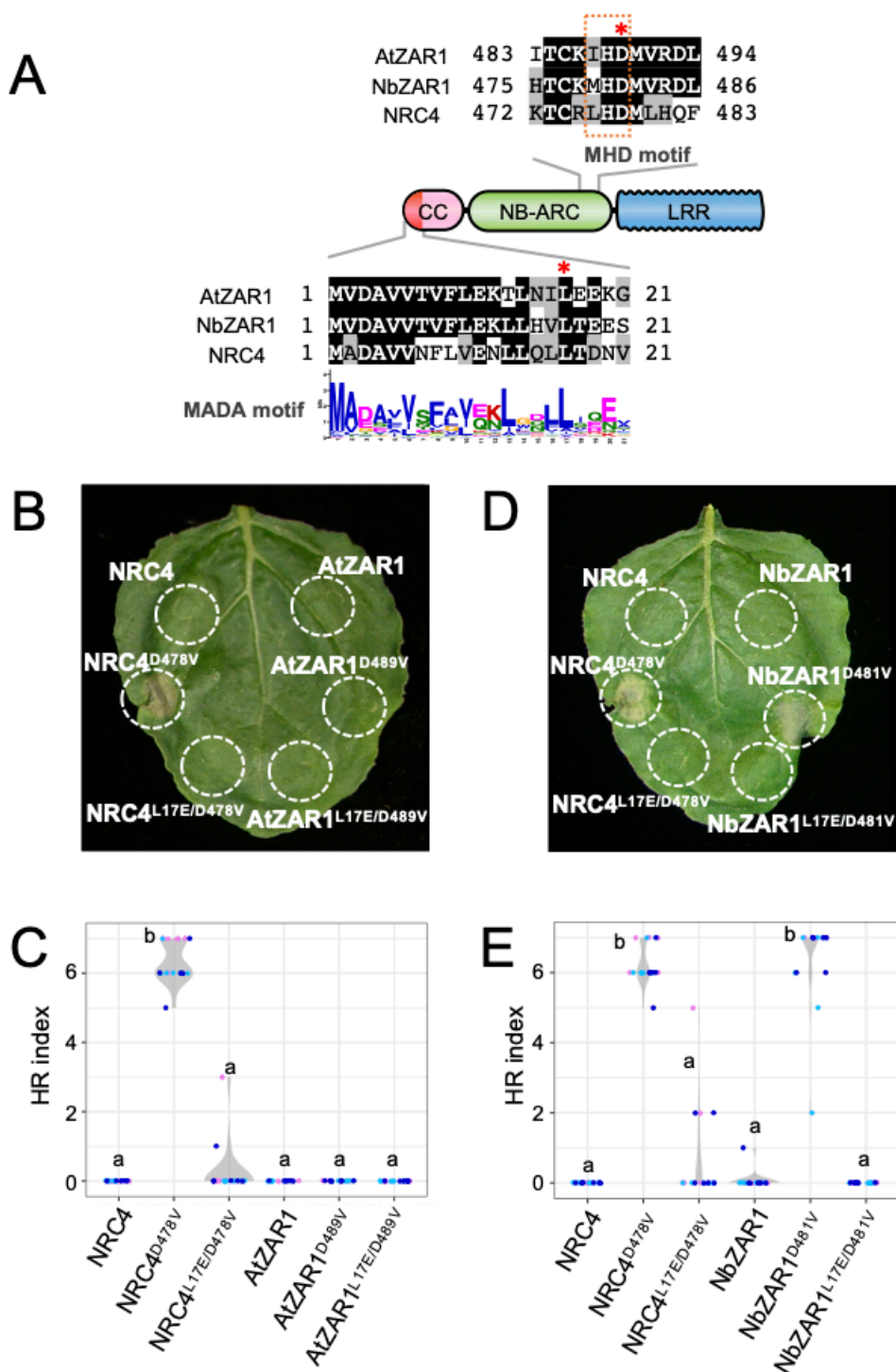


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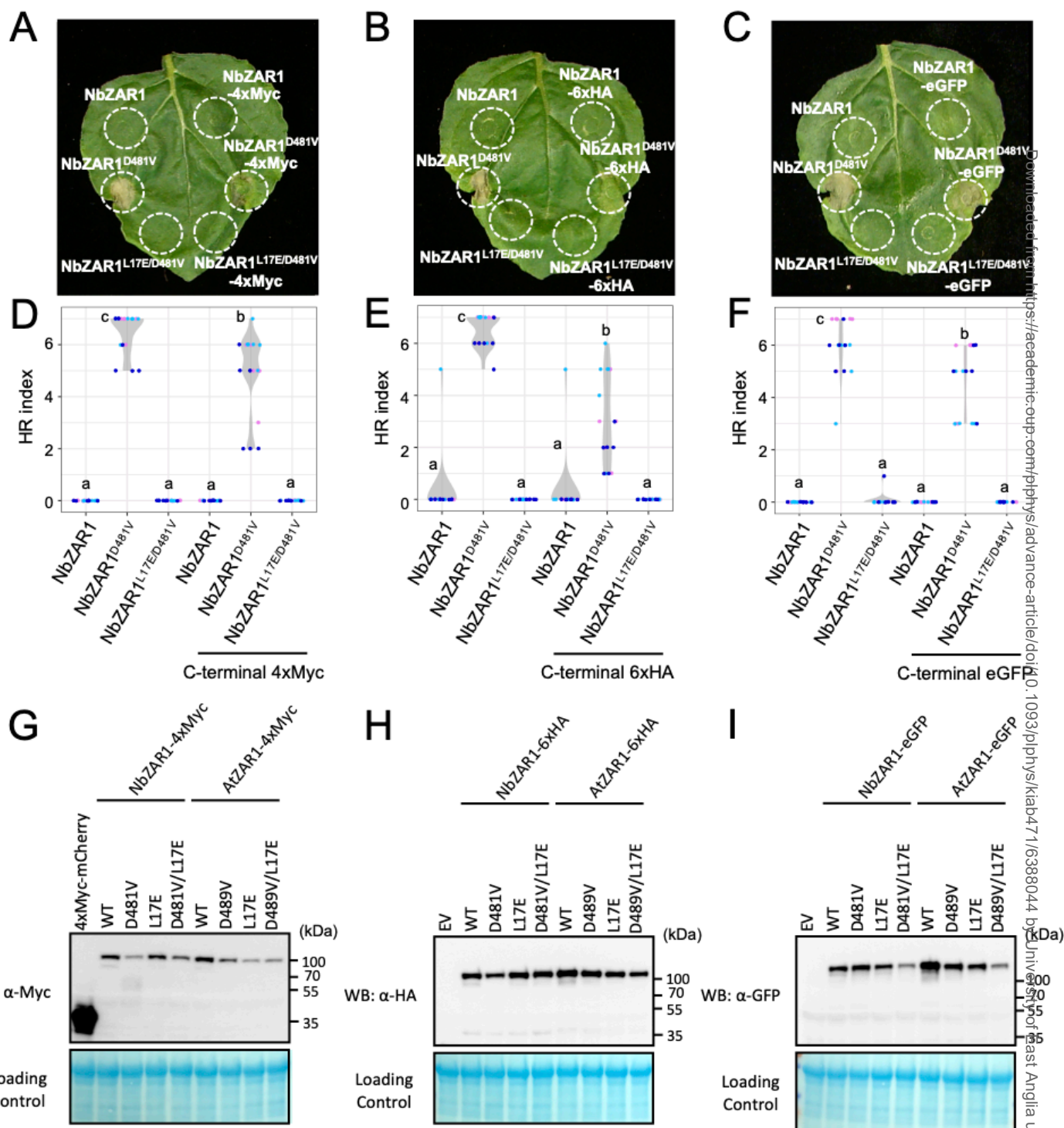


Figure 4. C-terminal tagged NbZAR1 triggers autoimmune cell death in *N. benthamiana* leaves. (A-C) Cell death observed in *N. benthamiana* leaves after expression of ZAR1 mutants. *N. benthamiana* leaf panels expressing wild-type and variant proteins of NbZAR1 with C-terminal 4xMyc, 6xHA or eGFP tag were photographed at 5 days after agroinfiltration. (D-F) 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 HSD test ($p < 0.01$). (G-I) *In planta* protein accumulation of AtZAR1 and NbZAR1 variants. For anti-Myc, anti-HA and anti-GFP immunoblots of AtZAR1, NbZAR1 and the mutant proteins, total proteins were prepared from *N. benthamiana* leaves at 2 days after agroinfiltration. Empty vector control is described as EV. Equal loading was checked with Reversible Protein Stain Kit (Thermo Fisher).

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