1	Estradiol-inducible AvrRps4 expression reveals distinct properties of
2	TIR-NLR-mediated effector-triggered immunity
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13	
14	Highlight
15	Inducible expression of AvrRps4 activates RRS1/RPS4-mediated effector-
16	triggered immunity without the presence of pathogens, allowing us to
17	characterise downstream immune responses triggered by TIR-NLRs without
18	cell-surface receptor-mediated immunity.
19	
20	Abstract
21	Plant nucleotide-binding domain, leucine-rich repeat receptor (NLR) proteins
22	play important roles in recognition of pathogen-derived effectors. However, the
23	mechanism by which plant NLRs activate immunity is still largely unknown. The
24	paired Arabidopsis NLRs RRS1-R and RPS4, that confer recognition of
25	bacterial effectors AvrRps4 and PopP2, are well studied, but how the
26	RRS1/RPS4 complex activates early immediate downstream responses upon
27	effector detection is still poorly understood. To study RRS1/RPS4 responses
28	without the influence of cell-surface receptor immune pathways, we generated
29	an Arabidopsis line with inducible expression of effector AvrRps4. Induction

electrolyte leakage, which often correlates with plant cell death. Activation of
 RRS1 and RPS4 without pathogens cannot activate mitogen-associated

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does not lead to hypersensitive cell death response (HR) but can induce

protein kinase cascades, but still activates upregulation of defence genes, and
 therefore resistance against bacteria.

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36 Keywords

Plant innate immunity, NLR activation, protein complex, hypersensitive
 response, cell death, MAP kinase, defence gene expression, estradiol inducible expression system, Golden Gate modular cloning

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41 Introduction

To investigate plant immunity, researchers routinely conduct pathogen 42 inoculations on plants in a controlled environment. Upon pathogen attack, 43 44 plants activate innate immune responses via both membrane-associated and intracellular receptors, which makes it difficult to unravel the distinct contribution 45 46 of each component. Most plasma-membrane localized receptors perceive conserved pathogen-associated molecular patterns (PAMPs) or host-cell-47 48 derived damage-associated molecular patterns (DAMPs) and activate PAMPtriggered immunity (PTI) or DAMP-triggered immunity (DTI). Plant intracellular 49 50 immune receptors belong to a family of nucleotide-binding leucine-rich repeat (NB-LRR) proteins, also known as NLRs. NLRs recognize pathogen effectors 51 52 and activate effector-triggered immunity (ETI), which often leads to accumulation of reactive oxygen species (ROS) and a hypersensitive cell death 53 response (HR). Most plant NLRs carry either coiled-coil (CC) or Toll/interleukin-54 1 receptor (TIR) N-terminal domains. Both CC and TIR domains are believed 55 to function in signalling upon activation of NLRs, but the detailed mechanisms 56 are unknown. Many CC-NLRs localize at and function in association with the 57 plasma membrane, whereas TIR-NLRs can function in diverse locations, 58 including the nucleus. Regardless of the distinct localization patterns between 59 CC- and TIR-NLRs, their downstream outputs culminate in elevated resistance, 60 but have never been directly compared side-by-side. To study the specific 61 immune outputs generated by ETI, inducible expression tools have been 62 applied (McNellis et al., 1998; Tornero et al., 2002; Allen et al., 2004; Porter et 63 al., 2012). 64

In Arabidopsis, functionally paired NLRs RRS1-R and RPS4 confer resistance against a soil-borne bacterial pathogen *Ralstonia solanacearum* through the

recognition of an effector PopP2 secreted via Type III secretion system and a 67 hemibiotrophic ascomycetous fungal pathogen Colletotrichum higginsianum 68 (Narusaka et al., 2009). They can also confer resistance against bacteria 69 Pseudomonas syringae pv. tomato DC3000 carrying AvrRps4, an effector 70 protein from Pseudomonas syringae pv. pisi, causing bacterial blight in Pisum 71 sativum (pea) (Sohn et al., 2009; Narusaka et al., 2009). Previously, it was 72 reported that the 135th to 138th residues of AvrRps4, lysine-arginine-valine-73 tyrosine (KRVY), are required for the recognition of AvrRps4 by RRS1 and 74 75 RPS4 (Sohn et al., 2009). Crystal structure of the C-terminus of AvrRps4 revealed that the 187th residue glutamate (E187) is also required for HR and 76 immunity (Sohn et al., 2012). PopP2 recognition by RRS1 occurs by the 77 integrated WRKY domain at the C-terminal of a resistant allele of RRS1-R (from 78 the Ws-2 ecotype of Arabidopsis) but not the susceptible allele of RRS1-S (from 79 the Col-0 ecotype of Arabidopsis) (Sarris et al., 2015). Crystal structure 80 information of RRS1 and RPS4 on the TIR domains and the co-crystal 81 82 structures between RRS1-R WRKY domain and effector PopP2 have indicated some structural basis of how RRS1/RPS4 have been activated (Williams et al., 83 84 2014; Zhang et al., 2017). However, it is still unknown how the protein complex

assembles and functions.

Here we report tools for studying the immune complex of RRS1-R and RPS4 *in vivo*. We established a set of transgenic Arabidopsis lines to study RRS1/RPS4-mediated ETI in the absence of pathogens. Using these lines, we show that some but not all immune outputs induced by the conditionally expressed AvrRps4 resemble other reported effector-inducible lines.

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92 Materials and Methods

93 Plant material and growth conditions

Arabidopsis thaliana accessions Wassilewskija-2 (Ws-2) and Columbia-0 (Col0) were used as wild type in this study. The *eds1-2* mutant used has been
described previously (Falk *et al.*, 1999). Seeds were sown on compost and
plants were grown at 21°C with 10 hours under light and 14 hours in dark (10hL/14h-D), and at 70% humidity. Tabaco plants were grown at 22°C with 16hL/8h-D, and at 80% constant humidity. The light level is approximately 180-200
µmols with fluorescent tubes.

101

102 FastRed selection for transgenic Arabidopsis

103 Seeds harvested from the Agrobacteria-transformed Arabidopsis are 104 resuspended in 0.1% Agarose and exposed under fluorescence microscope 105 with DsRed (red fluorescent protein) filter. Seeds with bright red fluorescence 106 are selected as the positive transformants.

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108 GUS staining

109 Nicotiana benthamiana (N. b.) leaves were infiltrated with Agrobacteria carrying constructs with β-glucuronidase (GUS) reporter gene expressed under selected 110 Arabidopsis promoters (Table S1). Leaves were collected at 2 days post 111 infiltration (dpi), and vacuum-infiltrated with GUS staining buffer (0.1 M sodium 112 phosphate pH 7.0, 10 mM EDTA pH 7.0, 0.5 mM K₃Fe(CN)₆, 0.5 mM K₄Fe(CN)₆, 113 0.76 mM 5-Bromo-4-chloro-3-indolyl-β-D-glucuronide cyclohexylamine salt or 114 X-Gluc, and 0.04% Triton X-100). After vacuum-infiltration, the leaves were 115 incubated at 37°C overnight in the dark. The leaves were rinsed with 70% 116 ethanol until the whole leaf de-stains to a clear white. 117

118

119 Immunoblotting

N. b. leaves were infiltrated with Agrobacteria carrying our stacking constructs 120 (Table S2). At 2 dpi, same leaves were infiltrated with either DMSO or 50 µM 121 β-estradiol (E2) diluted in water. Samples were collected at 6 hpi of DMSO or 122 E2 treatment, and snap-frozen in liquid nitrogen. Proteins were extracted using 123 GTEN buffer (10% glycerol, 25 mM Tris pH 7.5, 1 mM EDTA, 150 mM NaCl) 124 with 10 mM DTT, 1% NP-40 and protease inhibitor cocktail (cOmplete™. 125 EDTA-free; Merck). For Arabidopsis seedlings, seedlings grown for 8 days after 126 germination were treated with DMSO or E2 with indicated time points and snap-127 frozen in liquid nitrogen. After centrifugation at 13,000 rpm for 15 minutes to 128 remove cell debris, protein concentration of each sample was measured using 129 the Bradford assay (Protein Assay Dye Reagent Concentrate; Bio-Rad). After 130 normalization, extracts were incubated with 3× SDS sample buffer at 95°C for 131 5 minutes. 6% SDS-PAGE gels were used to run the protein samples. After 132 transferring proteins from gels to PVDF membranes (Merck-Millipore) using 133 Trans-Blot Turbo System (Bio-Rad), membranes were immunoblotted with 134

HRP-conjugated Flag antibodies (Monoclonal ANTI-FLAG® M2-Peroxidase
HRP antibody produced in mouse, A5892; Merck-Millipore), HRP-conjugated
HA antibodies (12013819001; Merck-Roche) or Phospho-p44/42 MAPK
(Erk1/2) (Thr202/Tyr204) (D13.14.4E) XP® Rabbit monoclonal antibody (4370;
Cell Signalling Technology). Anti-Rabbit IgG (whole molecule)–Peroxidase
antibody produced in goat (A0545; Merck-Sigma-Aldrich) was used as
secondary antibody following the use of Phospho-p44/42 MAPK antibody.

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143 Bacterial growth assay

Pseudomonas syringae pv. tomato strain DC3000 carrying pVSP61 empty 144 vector was grown on selective King's B (KB) medium plates containing 15% 145 (w/v) Agar, 25 µg/ml rifampicin and 50 µg/ml kanamycin for 48 h at 28°C. 146 Bacteria were harvested from the plates, resuspended in infiltration buffer (10 147 mM MgCl₂) and the concentration was adjusted to an optical density of 0.001 148 at 600 nm (OD₆₀₀=0.001, representing approximately 5×105 colony forming 149 units [CFU] ml-1). Bacteria were infiltrated into abaxial surfaces of 5-week-old 150 Arabidopsis leaves with a 1-ml needleless syringe. For quantification, leaf 151 152 samples were harvested with a 6-mm-diameter cork borer (Z165220; Merck-Sigma-Aldrich), resulting in leaf discs with an area of 0.283 cm². Two leaf discs 153 per leaf were harvested as a single sample. For each condition, four samples 154 were collected immediately after infiltration as 'day 0' samples to ensure no 155 significant difference introduced by unequal infiltrations and six samples were 156 collected at 3 dpi as 'day 3' samples to compare the bacteria growth between 157 different genotypes, conditions and treatments. For 'day 0', samples were 158 ground in 200 µl of infiltration buffer and spotted (10 µl per spot) on selective 159 KB medium agar plates to grow for 48 h at 28°C. For 'day 3', samples were 160 around in 200 µl of infiltration buffer, serially diluted (5, 50, 500, 5000 and 50000 161 times) and spotted (6 µl per spot) on selective KB medium agar plates to grow 162 for 48 h at 28°C. The number of colonies (CFU per drop) was monitored and 163 bacterial growth was represented as in CFU cm-2 of leaf tissue. All results are 164 plotted using ggplot2 in R (Wickham, 2016), and detailed statistics summary 165 can be found in the supplemental materials. 166

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168 HR phenotyping in Arabidopsis

Pseudomonas fluorescens engineered with a type III secretion system (Pf0-1 169 'EtHAn' strains) expressing one of wild-type or mutant effectors, AvrRps4, 170 AvrRps4kRvy135-138AAAA, PopP2, PopP2c321A, AvrRpt2 or pVSP61 empty vector 171 were grown on selective KB plates for 24 h at 28°C (Thomas et al., 2009; Sohn 172 et al., 2014). Bacteria were harvested from the plates, resuspended in 173 infiltration buffer (10 mM MgCl₂) and the concentration was adjusted to OD₆₀₀= 174 0.2 (108 CFU ml-1). The abaxial surfaces of 5-week-old Arabidopsis leaves were 175 hand infiltrated with a 1-ml needleless syringe. Cell death was monitored 24 h 176 177 after infiltration.

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179 Electrolyte leakage assay

Either 50 µM E2 or DMSO were hand infiltrated in 5-week-old Arabidopsis 180 leaves with a 1-ml needleless syringe for electrolyte leakage assay. Leaf discs 181 were taken with a 2.4-mm-diameter cork borer from infiltrated leaves. Discs 182 were dried and washed in deionized water for 1 hour before being floated on 183 deionized water (15 discs per sample, three samples per biological replicate). 184 Electrolyte leakage was measured as water conductivity with a Pocket Water 185 186 Quality Meters (LAQUAtwin-EC-33; Horiba) at the indicated time points. All results are plotted using ggplot2 in R (Wickham, 2016), and detailed statistics 187 summary can be found in the supplemental materials. 188

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190 Trypan blue staining

Either 50 µM E2 or DMSO were hand infiltrated in 5-week-old Arabidopsis 191 leaves with a 1-ml needleless syringe for trypan blue staining. 6 leaves per 192 sample were collected 24 hours after infiltration. Leaves were boiled in trypan 193 blue solution (1.25 mg/ml trypan blue dissolved in 12.5% glycerol, 12.5% 194 phenol, 12.5% lactic acid and 50% ethanol) in a boiling water bath for 1 min 195 and de-stained by chloral hydrate solution (2.5 g/ml). De-stained leaves were 196 mounted, taken under on 197 and pictures were Leica fluorescent stereomicroscope M165FC. All images were taken with identical settings at 198 2.5x magnification. Scale bar=0.5mm. 199

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201 Reverse transcription-quantitative polymerase chain reaction (RT-qPCR) for 202 measuring relative gene expression

For gene expression analysis, RNA was isolated from 5-week-old Arabidopsis 203 leaves and used for subsequent RT-gPCR analysis. RNA was extracted with 204 Quick-RNA Plant Kit (R2024: Zymo Research) and treated with RNase-free 205 DNase (4716728001; Merck-Roche). Reverse transcription was carried out 206 using SuperScript IV Reverse Transcriptase (18090050; ThermoFisher 207 Scientific). gPCR was performed using a CFX96 TouchTM Real-Time PCR 208 Detection System. Primers for gPCR analysis of Isochorismate Synthase1 209 (ICS1), Pathogenesis-Related1 (PR1), AvrRps4 and Elongation Factor 1 Alpha 210 211 $(EF1\alpha)$ are listed in Table S4. Data were analyzed using the double delta Ct method (Livak and Schmittgen, 2001). All results are plotted using ggplot2 in R 212 (Wickham, 2016), and detailed statistics summary can be found in the 213 214 supplemental materials.

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216 Confocal laser scanning microscopy (CLSM) imaging

Transgenic plant materials were imaged with the Leica DM6000/TCS SP5 217 confocal microscopy (Leica Microsystems) for confirmation of expression of 218 inducible AvrRps4 fused with monomeric yellow-green fluorescent protein, 219 220 mNeonGreen or mNeon (Shaner et al., 2013). Roots from 3-week-old Arabidopsis seedlings were sprayed with 50 µM E2 and imaged at 1 day post 221 222 spray. Fluorescence of mNeon was excited at 500 nm and detected at between 520 and 540 nm. CLSM images of root cells from Arabidopsis seedlings are 223 recorded via the camera. The images were analyzed with the Leica application 224 225 Suite and Fiji software (Schindelin et al., 2012).

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227 Co-immunoprecipitation

Arabidopsis transgenic seedlings, and the background ecotype Col-0 grown for 228 7 days after germination (DAG) were treated with 0.1% DMSO or 50µM E2 for 229 3 hours. Proteins from seedlings were extracted using IP buffer (10% Glycerol, 230 50mM Tris-Cl pH 6.8, 50mM KCl, 1mM EDTA, 5Mm MgCl₂, 1% NP-40, 10mM 231 DTT, 1mM dATP). Crude extract of the seedlings was centrifuged, and 232 supernatants were incubated with Anti-HA-conjugated beads (EZviewTM Red 233 Anti-HA Affinity Gel; E6779; Sigma). A small portion of supernatants were taken 234 for input samples. At 2 hours after incubation of the extract with beads, beads 235 were washed three times with IP buffer containing 0.1% NP-40. Proteins bound 236

to beads were eluted by boiling the beads with SDS sample buffer.
Immunoblotting of the input and eluted samples were performed as described
above.

240

241 **Results**

242 RRS1 over-expression can compromise RPS1/RPS4 function

Overexpression of RPS4 leads to autoimmunity and dwarfism under standard 243 growth condition (see Materials and Methods) (Heidrich et al., 2013). This 244 245 autoimmunity is both temperature- and RRS1-dependent. In contrast, elevated expression of RRS1-R from ecotype Ws-2 in Col-0, an ecotype expressing a 246 dominant allele of RRS1-S, does not trigger auto-immunity (Huh et al., 2017). 247 Furthermore, high level RRS1-R expression does not confer recognition of 248 effector PopP2 (Fig 1). Overexpression of RRS1 in an RPS4 overexpression 249 line attenuates dwarfism and autoimmunity (Huh et al., 2017). We infiltrated 250 non-pathogenic strains of Pseudomonas (P.) fluorescens Pf0-1 engineered 251 with the type-III secretion system from P. syringae pv. tomato (Pst) DC3000 252 strain that are expressing effectors PopP2, mutant PopP2c321A, AvrRps4, 253 254 mutant AvrRps4kRvy-AAAA, AvrRpt2, and empty vector, respectively (Sohn et al., 2009; Thomas et al., 2009; Saucet et al., 2015). This enabled the assessment 255 256 of HR activated by individual effector with their corresponding NLR proteins without artefactual tissue damage from the carrier. Ws-2 ecotype containing 257 RRS1-R recognizes wild-type PopP2 (PopP2wT), whereas RRS1-S-containing 258 Col-0 ecotype show no HR with PopP2wT (Fig 1). Mutant PopP2 (PopP2c321), 259 260 mutant AvrRps4 (AvrRps4kRvy-AAAA) and empty vector served as nonrecognition negative controls which does not activate HR (Fig 1). AvrRpt2 is 261 known to be recognized by CC-NLR RPS2 (Axtell and Staskawicz, 2003; 262 Mackey et al., 2003), and therefore HR was observed in all tested lines. We 263 found that only simultaneously over-expressing RRS1-R and RPS4 can lead to 264 the gain-of-recognition of PopP2 in the susceptible ecotype Col-0 (Fig 1). No 265 HR was observed in rps4-2 rps4b-2 double mutant when infiltrated with Pf0-266 1:AvrRps4 (Fig 1). Thus, we propose that a balanced protein expression of 267 RRS1 and RPS4 is required for both suppressing autoimmunity and functional 268 recognition of the corresponding effectors. 269

270

A survey of leaf-expressed genes reveals promoters for moderate and

272 balanced expression levels of RRS1 and RPS4

Genome-wide expression profiling has revealed numerous genes altered by 273 PTI alone or PTI plus ETI at early time points of RRS1/RPS4-mediated immune 274 activation (Sohn et al., 2014). This analysis also enabled the discovery of genes 275 that are moderately and constitutively expressed without changing their 276 transcript abundance during immune activation. In plants, gene expression 277 patterns and levels are usually specified by their promoters. Based on the 278 279 endogenous expression relative transcript abundance in the 'stable gene set', we selected six promoters with 'moderate' expression (Table S1). We define 280 the 'moderate' expression based on two criteria: (1) the gene transcript 281 abundance with those promoters are at least 100 times more than the 282 endogenous transcript abundance of RRS1 and RPS4; (2) the gene transcript 283 abundance with those promoters is lower than that with the 35S promoter. The 284 selected genes encode proteins that are involved in essential biological 285 processes that we expect to be expressed in most mesophyll cells, including a 286 delta-tonoplast intrinsic protein (our name; At1, locus identifier AT3G16240, 287 288 protein symbol name TIP2-1), a ribosomal protein S16 (At2, AT4G34620, RPS16-1), a cysteine synthase isomer CysC1 (At3, AT3G61440, CYSC1), a 289 290 photosystem II subunit Q (At4, AT4G21280, PSBQ1), a xyloglucan endotransglucosylase/hydrolase 6 (At5, AT5G65730, XTH6), and a ubiquitin-291 like protein 5 (At6, AT5G42300, UBL5) (Table S1). 292

To test the strength of the selected Arabidopsis promoters (pAt1-pAt6) for 293 294 driving gene expression in planta, constructs were designed and generated to express β-glucuronidase (GUS) (pAt:GUS). Agrobacterium strains carrying 295 each pAt:GUS construct was infiltrated in Nicotiana (N.) benthamiana leaves 296 with the infiltration buffer as negative control and GUS expressed under the 297 CaMV 35S promoter (35S:GUS) as positive control. GUS expressed under 298 pAt4 shows similar level of activity to that with 35S, whereas GUS activities 299 detected from other pAt promoters are significantly weaker (Fig S1). 300

301

302 A T-DNA construct expresses RPS4, RRS1 and inducible AvrRps4

We designed a binary vector to reconstruct the effector ligand AvrRps4 and its receptors RRS1 and RPS4, using the Golden Gate Modular Cloning Toolbox

(Fig 2A) (Engler et al., 2014). We chose moderate and balanced promoters 305 pAt2 and pAt3 from our promoter survey experiment for expressing RRS1 and 306 RPS4, respectively. We have also cloned RRS1-R full-length coding 307 sequences (CDS) from Ws-2 and RPS4 full-length CDS from Col-0 for the 308 expression of RRS1-R and RPS4 proteins. We chose synthetic C-terminal-309 fusion epitope tags His6-TEV-FLAG₃ (HF) and HA6 for detecting RRS1 and 310 RPS4 protein expressions, respectively (Fig 2A, Table S2) (Gauss et al., 2005; 311 Soleimani et al., 2013). We have used an E2-inducible system for AvrRps4 312 313 expression (Zuo et al., 2000). We named this multi-gene stacking binary construct 'Super ETI', or SETI. We have also generated constructs inducing 314 mutant AvrRps4kRvy-AAAA or mutant AvrRps4E187A as negative controls, and 315 named them SETI KRVYmut and SETI E187A, respectively. SETI KRVYmut 316 and SETI E187A can induce the expression of mutant AvrRps4 alleles, but no 317 induction of immunity because these two mutant AvrRps4 alleles cannot be 318 recognized by RRS1 and RPS4 (Sohn et al., 2009, 2014). All restriction enzyme 319 sites for Bsal and Bpil in modules for promoters. CDSs for genes or epitope 320 tags and the terminators were synonymously eliminated (Fig 2A, Table S4). 321 322 More detailed information for the cloning can be found in supplemental materials. To verify the SETI construct, we used a transient expression system 323 in *N. benthamiana* by infiltrating Agrobacteria that deliver the SETI T-DNA. 324 Protein accumulation of RRS1-R-HF and RPS4-HA was detected (Fig S2). 325

326

The single-locus lines carrying the SETI T-DNA show inducible growth arrest

We generated transgenic Arabidopsis lines using the SETI, SETI KRVYmut 329 and SETI E187A construct expressing AvrRps4 (SETI WT), AvrRps4kRvy-AAAA, 330 and AvrRps4E187A, respectively. With the FastRed selection module, we have 331 selected approximately 20 positive SETI_WT T1 lines. The seedlings from the 332 T2 generation of 3 T1 lines were further tested for response to E2 treatment 333 (see Materials and Methods, Table S2). On E2-containing growth medium, 334 SETI WT transgenic lines display severe growth arrest (Fig S3). We selected 335 one of the lines (T1-#8 T2-#4; SETI WT) for subsequent experiments (Fig 2C, 336 Fig S3). We confirmed the protein expression of RRS1-R-HF and RPS4-HA 337 (Fig 2B). We also tested the expression of inducible AvrRps4-mNeon under 338

fluorescence microscope upon the treatment with E2. mNeonGreen signal was detected at 24 hours post spray on transgenic seedlings, consistent with the mRNA accumulation of *AvrRps4* at 4 hours post E2-infiltration in leaves (Fig 2D, Fig S2C).

343

344 RRS1-R and RPS4 form pre-activation complexes in Arabidopsis

The SETI lines enable detection of epitope-tagged RRS1-R and RPS4 (Fig 2B). 345 We investigated in vivo interaction of tagged RRS1-R and RPS4 by co-346 347 immunoprecipitation (co-IP) with SETI WT and SETI E187A seedling extracts with or without E2 induction. When RPS4-HA was immunoprecipitated using 348 HA beads, we found RRS1-R and RPS4 stay in association with each other 349 both before and 3 hours after the induction of AvrRps4 expression (Fig 3). 350 There were no significant differences of RRS1-R and RPS4 association upon 351 AvrRps4 induction. Induction of AvrRps4 E187A also had no effect on RRS1-R 352 and RPS4 association. While all previous studies in interactions of RRS1-R and 353 RPS4 was tested only using N. benthamiana transient expression system (Huh 354 et al., 2017), generation of SETI line enabled the detection of RRS1-R and 355 356 RPS4 interaction in its native system in Arabidopsis.

357

358 Some but not all defence responses are induced by E2 in SETI lines

The induced expression of multiple effectors, such as AvrRpt2, AvrRpm1 and 359 ATR13 can induce cell death or named macroscopic HR in Arabidopsis leaves 360 (McNellis et al., 1998; Tornero et al., 2002; Allen et al., 2004). We therefore 361 tested whether induced expression of AvrRps4 can trigger macroscopic HR in 362 Arabidopsis. We used SETI eds1 as control, in which SETI WT was crossed 363 with the mutant eds1. EDS1 is downstream genetic component of TIR-NLR-364 mediated ETI (Aarts et al., 1998; Falk et al., 1999). As seen in Fig 4A, no HR 365 can be observed after AvrRps4 expression is induced in the SETI leaves. 366 However, only the expression of AvrRps4 but not AvrRps4 KRVYmut leads to 367 electrolyte leakage (Fig 4C). We also observed slightly stronger trypan blue 368 stains in the SETI leaves treated with E2 compared to mock treatment; 369 suggesting that the expression of AvrRps4 causes microscopic or weak but not 370 macroscopic or strong HR in contrast to other known inducible effector lines 371 (Fig 4B). 372

Salicylic acid induction is another hallmark of ETI (Castel et al., 2019). Enzymes 373 such as Isochorismate Synthase 1 (ICS1), Enhanced Disease Susceptibility 5 374 (EDS5) and AvrPphB Susceptible 3 (PBS3) are involved in the biosynthesis of 375 salicylic acid and the expression of these genes is also highly induced during 376 ETI (Sohn et al., 2014). The expression of ICS1 after the AvrRps4 induction 377 was tested by guantitative real-time PCR. ICS1 was highly induced 4 hours 378 after the induction of AvrRps4 by E2 but not in the negative controls of 379 SETI KRVYmut or SETI eds1 (Fig 5A). In contrast, Pathogenesis-Related 380 381 protein 1 (PR1) was highly induced only 8 hours after the induction of AvrRps4 (Fig 5B). This shows that ETI triggered by RRS1/RPS4 is sufficient for the 382 induction of *ICS1* and the biosynthesis of salicylic acid, which subsequently 383 leads to expression of PR1. 384

Activation of mitogen-activated protein kinases (MAPKs) by PTI has been 385 reported under many cases and happens within a few minutes of the activation 386 of PTI. However, the activation of MAPKs by ETI is slower and lasts longer than 387 PTI-induced MAPK activation (Tsuda et al. 2013). We tested whether the 388 induced expression of AvrRps4 can lead to MAPK activation in SETI WT and 389 390 control lines Col-0, SETI KRVYmut, and SETI eds1. Treatment of flg22 for 10 minutes triggered phosphorylation of MAP kinases (Fig 5C). However, in 391 392 contrast to AvrRpt2-inducible transgenic Arabidopsis plants (Tsuda et al. 2013), induced expression of AvrRps4 does not activate MAPKs (Fig 5C). 393

We further tested if the induction of ETI would elevate resistance. We infiltrated the leaves with E2 or mock solution one day before we infiltrated plants with *Pst* DC3000 (see Materials and Methods). SETI_WT plants pre-treated with E2 are more resistant to the bacteria than those pre-treated with mock, while there was no significant difference between E2 and mock pre-treatment in Col-0 (Fig 6).

400

401 **Discussion**

To facilitate studying the functional complex of RRS1 and RPS4 *in vivo*, we generated an expression construct of E2-inducible AvrRps4 stacked with epitope-tagged RRS1 and RPS4. To achieve balanced expression levels higher than endogenous expression of *RRS1* and *RPS4*, we surveyed constitutively expressed gene promoters. Here, we report 6 new and tested

promoter modules that are compatible with the Golden Gate Modular Cloning 407 toolkit. We used two of the promoters to express RRS1 and RPS4, and we 408 avoided autoimmunity induced by excessive expression of RPS4 or non-409 recognition of PopP2 cause by excessive expression of RRS1-R. We were also 410 able to generate inducible AvrRps4 expression to activate RRS1/RPS4-411 mediated ETI under the control of E2 treatment. We thus were able to stack 412 genes for inducible expression of a pathogen effector and its NLR receptors in 413 one construct. In addition, with the epitope tags, we are able to monitor effector-414 415 dependent changes in the NLR proteins without interference from using a pathogen effector-delivery system. We could thus express any effectors or 416 pathogen ligands that will trigger immunity in plant cells with the E2-inducible 417 module, and their immune receptors using the same gene stacking strategy. 418

There are multiple advantages to enabling investigation of ETI without the 419 420 complication of co-activating PTI. Firstly, we could test the contribution of other genes to ETI activation by introducing mutants into the SETI background, either 421 using conventional crossing or using genome-editing such as CRISPR/Cas9. 422 These lines can also help investigating downstream signalling from plant NLRs. 423 424 Multiple forward genetic screens have been conducted, but few novel components have been found, and most mutations are either in the NLRs or 425 426 regulatory elements rather than signalling components (van Wersch et al., 2016). Another plausible explanation is that the signalling path downstream of 427 plants NLRs is very short, but this is debatable, because several significant 428 steps are required for immunity. EDS1, PAD4 and SAG101 are required for 429 TIR-NLR signalling (Falk et al., 1999; Gantner et al., 2019). NRC family proteins 430 in Solanaceae species required for many NLRs, and NRG1/ADR1s in 431 Arabidopsis required for TIR-NLRs and ADR1s for some CC-NLRs (Bonardi et 432 al., 2011; Dong et al., 2016; Wu et al., 2017, 2019; Castel et al., 2019). NRG1s 433 and ADR1s seem to function downstream of EDS1 and may function distinctly 434 with SAG101 and PAD4, respectively (Lapin et al., 2019). SETI lines carry 435 heterologously expressed RRS1-R/RPS4 and also endogenous RRS1-S/RPS4, 436 RRS1B/RPS4B, which together provide three redundant copies of NLR pairs 437 that can recognize AvrRps4. In theory, in an EMS-mutagenesis forward genetic 438 screen to identify suppressors of immunity induced by AvrRps4, there should 439 be a reduced background of mutations in the receptor(s), improving prospects 440

to reveal mutations in genes that are functionally important in NLR signallingand regulation.

With SETI, we are able to assess pure ETI response mediated by the TIR-NLRs, 443 RRS1 and RPS4. E2 induction provoked rapid transcriptional changes in 444 activation of defence genes and also ion leakage. AvrRps4-induced ETI 445 enhanced resistance against bacterial pathogens. However, neither MAPK 446 activation nor macroscopic HR, in contrast to other inducible ETI examples 447 (Tornero et al., 2002; Tsuda et al., 2013). This indicates that outputs of plant 448 449 NLRs might differ. Both TIR and CC domains alone are sufficient to activate plant immunity. However, whether they signal through similar or different 450 downstream components is still unknown. 451

In diverse multicellular eukaryotes, immune complexes are assembled into 452 oligomeric complexes to signal downstream. The mammalian inflammasome, 453 assembled in response to bacterial peptide recognition by NAIP proteins and 454 subsequent activation and binding of NLRC4 proteins, is a classic example 455 (Zhang et al., 2015). The plant CC NLR ZAR1 forms an effector-dependent 456 resistosome, which is a pentamer of ZAR1 assembled together with cofactors 457 458 PBL2 and RKS1 (Wang et al., 2019). The structure of TIR domains implies that activation might require the disassociation of the RRS1 and RPS4 TIR domains 459 460 and the oligomerization of RPS4 TIR domains (Williams et al., 2014). In SETI lines, RRS1 and RPS4 form a pre-activation complex in the absence of 461 pathogen effector. However, co-IP data cannot distinguish the ratio of which 462 RRS1 and RPS4 bind to each other. It will be interesting to check via various 463 non-denaturing methods if RRS1-R and RPS4 form a dimer or a higher order 464 oligomerization *in vivo*, or whether there is a conformational change in complex 465 upon effector recognition. Furthermore, with the SETI lines generated in this 466 study, we can ask what other co-factors are required for the activation of RRS1-467 R and RPS4 at native conditions. 468

The availability of SETI lines also will enable us to study how PTI and ETI interact with each other, especially in the context of RRS1- and RPS4-mediated immunity. Some models have been proposed in discussing on this topic (Tsuda *et al.*, 2009; Cui *et al.*, 2015). From the zig-zag model, PTI and ETI holds in different threshold on activating immunity (Jones and Dangl, 2006). With SETI line, we could specifically ask how physically PTI and ETI can influence each

other. A lot of evidence shows that the PTI receptors PRRs usually have very 475 specific post-translational modification events at early time points, there is also 476 some evidence showing ETI can activate somewhat overlapping but different 477 PTMs on immune-related proteins (Withers and Dong, 2017; Kadota et al., 478 2019). It will be interesting to know how the activation of RRS1/PRS4 leads to 479 the changes of PTMs and how those changes contribute to the robustness of 480 immunity. In addition, transcriptional changes are not the only process reported 481 as the early changes of ETI but also the changes in translations (Meteignier et 482 483 al., 2017; Yoo et al., 2019). Both work using inducible AvrRpm1 or AvrRpt2 reveal interesting observations on trade-off between defence and growth, and 484 the specific regulatory element in the genome (Meteignier et al., 2017; Yoo et 485 al., 2019). Both effectors are recognized by CC-type NLRs, so it will be 486 interesting to know what changes in translations will be induced by TIR-NLRs 487 488 using SETI line. One can also use proteomics tools to generate complex information using inducible SETI to fish for ETI-specific interaction networks. 489

490 Recently it has been shown plant NLRs can also form higher order protein complex, similar to inflammasome in mammalian immune system. However, it 491 is unknown if all plant NLRs form the same kind of complex or using the same 492 493 mechanism to activate defence. It was noted that NLRs have evolved to partner with other NLRs to function genetically, but if this model is also true 494 biochemically is still unknown (Adachi et al., 2019). Unlike ZAR1, RRS1 and 495 RPS4 requires each other to function, and they localized and function 496 exclusively in the nuclei but not the cell membrane, so it will be interesting to 497 compare them once the transmission electron cryomicroscopic (Cryo-EM) 498 structure of RRS1 and RPS4 complex is resolved. SETI line could be a very 499 good toolkit to make mutagenesis to verify the function based on the structural 500 information. 501

We have observed the activation of ETI alone in the absence of pathogens is sufficient to prime the resistance against bacterial pathogens in Arabidopsis (Fig 6). Previously, we have reported a group of upregulated genes at the early time point of activation of RRS1-R/RPS4 are related to salicylic acid pathway, so it will be interesting to know if the elevated or primed resistance against bacteria induced in SETI lines are due to the activation of salicylate pathway (Sohn *et al.*, 2014).

15

Another major question regarding the signalling pathways is that SAG101 and PAD4 seems to be redundant but functionally equivalent to EDS1 (Wagner *et al.*, 2013; Lapin *et al.*, 2019). They also have been shown to be genetically linked to helper NLRs NRG1s and/or ADR1s to function (Castel *et al.*, 2019; Wu *et al.*, 2019). Using SETI line, one can test their function more specifically in ETI in the absence of PTI and many other unwanted pathogen interferences.

515

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Figure 1



Fig. 1. Over-expression of RPS4 and RRS1-R reconstruct the recognition of PopP2 in Col-0. Arabidopsis transgenic lines overexpressing RPS4 (OE-*RPS4-HS*), RRS1-R (OE-*RRS1-R-HF*) or both generated by crossing (OE-*RPS4-HS*/OE-*RRS1-R-HF*) in the Col-0 background, with Col-0 and Ws-2 accession were tested for hypersensitive response (HR). 5-week old leaves were infiltrated with *Pseudomonas fluorescence* Pf0-1 strains carrying empty vector (EV), wild-type (WT) AvrRps4, mutant AvrRps4^{KRVY-AAAA}, WT PopP2, mutant PopP2^{C321A}, and WT AvrRpt2. Leaves were collected 1 day post infiltration (dpi) for imaging. Scale bar = 1cm. Yellow arrows indicate reconstructed PopP2 recognition of Col-0 background overexpressing RRS1-R and RPS4. Yellow dashed box highlights loss of AvrRps4 recognition in the double mutant *rps4-2 rps4b-2*. Infiltration of EV and AvrRpt2 serve as negative and positive controls of HR, respectively.



Fig 2. Single T-DNA expresses RRS1-R-HF, RPS4-HA and inducible wild-type AvrRps4 or AvrRps4 mutant variants

(A) Illustrative layout of the SUPER-ETI (SETI) construct. There are five individual expression units or Golden Gate Level 2 positional components listed, which are indicated position 1 to position 5. Position 1; expression unit of the FastRed selection marker (Shimada et al. 2010). Position 2, 5; chimeric transactivator XVE (LexA-VP16-ER) and the corresponding LexA inducible system to express AvrRps4 or its mutant variants under the control of β -estradiol (E2) treatment. Position 3, 4; full-length RRS1-R and RPS4 proteins with epitope tags His₆-Flag₃ and HA₆, respectively. All cloning details can be found in Methods and Materials. All individual units used for construct assembly can be found in the Supplemental Table 2 and 3.

(B) Protein accumulation of RRS1-R-HF (IB:Flag, black arrowhead) and RPS4-HA (IB:HA, white arrowhead) of SETI lines expressing AvrRps4 (SETI_WT) or mutant AvrRps4 KRVY-AAAA (SETI_KRVYmut). Seedlings were grown in liquid culture and induced with 50µM E2 for 2 hours at 7 days after germination (DAG). Ponceau staining of Rubisco large subunits were used as loading control.

(C) Seedling phenotype of SETI Arabidopsis transgenic line at 14 DAG in GM media containing Mock (0.1% DMSO) or 50μ M E2. Col-0 was sown as control for the effect of E2 on seedling growth. Scale bar = 0.5cm

(D) Confocal images of SETI_WT, SETI_KRVYmut, SETI_*eds1* root cells expressing AvrRps4-mNeon and AvrRps4^{KRVY-AAAA}–mNeon induced by 50 μ M E2 for 24h. mNeon channel shows nucleo-cytoplasmic localization of AvrRps4-mNeon and AvrRps4^{KRVY-AAAA}–mNeon. Bright field channel and merged image of mNeon and Bright field channel are shown together. Bars = 10 μ m.



Fig 3. RRS1-R and RPS4 interact in vivo.

Co-immunoprecipitation of RRS1-R-HF with RPS4-HA. Col-0, SETI_WT, and SETI_E187A seedlings at DAG7 were treated with 50µM E2 for 3hours. Crude extracts were centrifuged and RPS4-HA proteins were immunoprecipitated with Anti-HA-conjugated beads. Immunoprecipitation of RPS4-HA, and co-immunoprecipitation of RRS1-R-HF were determined by immunoblot analysis with HA (IB:HA) or Flag (IB:Flag). Ponceau staining indicates equal loading of the input samples. RRS1-R-HF (black arrowhead), and RPS4-HA (white arrowhead) are indicated.

Figure 4



Fig. 4. Induced expression of AvrRps4 in *Arabidopsis* cause microscopic but not macroscopic cell death.

(A) HR phenotype assay in Arabidopsis. 5-week old SETI_WT, SETI_KRVYmut and SETI_*eds1* leaves were infiltrated with Mock (1% DMSO) or 50µM E2. Images were taken at 1dpi. Numbers indicate the number of leaves displaying cell death from the total number of infiltrated leaves (18 for each genotype and treatment).

(B) Trypan blue staining. 5-week old SETI_WT, SETI_KRVYmut and SETI_eds1 were infiltrated with Mock (1% DMSO) or 50μ M E2. Leaves were stained with trypan blue solution at 1dpi. After destaining, leaves were imaged using stereoscopic microscope. Scale bar = 0.5mm

(C) Electrolyte leakage assay. 5-week old SETI_WT, SETI_KRVYmut and SETI_eds1 leaves were infiltrated with Mock (1% DMSO) or 50µM E2. Fifteen leaf discs were collected for each data point. Conductivity was measured at 1, 5, 20 and 24 hours post infiltration (hpi). Each data point represents one technical replicate and three technical replicates are included per treatment and genotype for one biological replicate. Black line represents the mean of the technical replicates. This experiment was repeated three times independently with similar results (Supplemental Figure 2). Significant differences relative to the mock treatment in each genotype was calculated with t-test and the P-values are indicated as ns (non-significant), P > 0.05; *, P < 0.05; **, P < 0.01; ***, P < 0.001.

Figure 5



Fig. 5. Induced expression of AvrRps4 in *Arabidopsis* leads to *ICS1* and *PR1* expression, but not MAPK activation.

(A and B) *ICS1* (A) and *PR1* (B) expression after induction with E2 for 2, 4, and 8h in SETI (left panel), SETI_KRVYmut (middle panel) and SETI_eds1 (right panel) leaf samples. 5-week old SETI and SETI_KRVYmut leaves were infiltrated with 50 μ M E2. Samples were collected at 0, 2, 4 and 8hpi for RNA extraction and subsequent qPCR. Expression level is presented as relative to *EF1a* expression. Each data point represents one technical replicate. Black line represents the mean of the technical replicates. This experiment was repeated three times independently with similar results. Significant differences relative to the untreated samples was calculated with t-test and the P-values are indicated as ns (non-significant), P > 0.05; *, P < 0.05; **, P < 0.01; ***, P < 0.001.

(C) Activation of MAP kinases in Col-0, SETI_WT, SETI_KRVYmut and SETI_eds1 seedlings by E2-induction of effector AvrRps4 or mutant AvrRps4^{KRVY-AAAA}. Seedlings grown in liquid culture at 7 dag were treated with 50µM E2 for indicated time points (0, 2, 4, 6, 8h) and collected for samples. Col-0, SETI_WT, SETI_KRVYmut and SETI_eds1 seedlings treated with 100nM flg22 for 10 minutes (10min*) were used as positive control. Proteins were extracted from these seedlings and phosphorylated MAP kinases were detected using p-p42/44 antibodies. Arrowheads indicate phosphorylated MAP kinases (black, pMPK6; grey, pMPK3; white, pMPK4/11). Ponceau staining were used as loading control.



Fig. 6. Effector-triggered immunity triggered by the expression of AvrRps4 leads to resistance against *Pseudomonas syringae pv. tomato* strain DC3000.

5-week old SETI_WT and Col-0 leaves were infiltrated with Mock (1% DMSO) or 50µM E2. At 1dpi, leaves were inoculated with *Pst* DC3000 (OD600=0.001). Bacteria in the leaves were then quantified as colony-forming units (CFU) at 0dpi and 3dpi. Each data point represents two leaves collected from one individual plant. Samples from four individual plants were collected for 0 dpi and samples from six individual plants were collected for 3 dpi. Black line represents the mean of the technical replicates. This experiment was repeated three times independently with similar results. Biological significance of the values were determined by one-way ANOVA followed by post hoc TukeyHSD analysis. Letters above the data points indicate significant differences (P<0.05).

Promoter Number	Promoter length (bp) ¹	Gene Symbol (UniProtKB)	AGI (TAIR)	Expression (TPM) ²
pAt1	1746	TIP2-1	AT3G16240	1600
pAt2	711	RPS16-1	AT4G34620	850
pAt3	550	CYSC1	AT3G61440	600
pAt4	348	PSBQ1	AT4G21280	557
pAt5	982	XTH6	AT5G65730	1000
pAt6	1659	UBL5	AT5G42300	280

Supplementary Table S1 Information of synthetic promoters used in this study.

1. The promoters are chosen from the first nucleotide next to the start codon ATG on the opposite direction of gene coding direction. The total length of each gene promoter is defined from the start codon of gene of interest up to either the 5' or 3' end of the immediate neighbouring gene.

2. TPM, tags per million. Transcripts of each gene under their endogenous promoters are indicated by the RNA sequencing profile data generated in Arabidopsis Ws-2 accession (Sohn *et al.*, 2013).

Position in Level2	Flank ¹	P+5U ²	CDS or genomic ³	cTag⁴	Ter⁵	Flank	Simplified Module
1	TGCC	AtOleosin ⁶	AtOleosin	RFP	AtOleosin	GCAA	FastRed
2	TTAC	AtActin2	XVE		AtuMas ⁷	CAGA	XVE
3	GCAA	AtSSR16 ⁸	AtRRS1-R	Hellfire ⁹	AtRRS1-R	ACTA	RRS1-R-HF
4	ACTA	AtCysC1 ¹⁰	AtRPS4	HA ₆ ¹¹	CaMV35S	TTAC	RPS4-HA
5	CAGA	LexA	PsAvrRps4 ¹²	BlmNeon ¹³	AtuOcs ¹⁴	TGTG	LexA:AvrRps4- mNeon
End Linker	TGTG		gaggatgcaca		GGGA	pELE-5	
Back Bone	TGCC		-				pAGM4723

Supplementary Table S2 Golden Gate stacking construct of R genes and effector.

1. Flank: the flank sequences indicate the overhang sequence generated by the restriction enzyme Bpil from the level 1 modules to the level 2 destination backbone, before the final ligation reaction.

2. P+5U: promoter and 5' untranslated region (UTR).

3. CDS or genomic: coding sequence or full-length genomic sequence that includes potential introns.

4. cTag: c-terminal in-frame coding sequence for epitope tag.

5. Ter: terminator.

6. AtOleosin: AT4G25140, a protein found in oil bodies, involved in seed lipid accumulation, that is specifically expressed in seed coat.

7: AtuMas: terminator of Mas1 agropine synthesis reductase from *Agrobacterium tumefaciens* (Engler *et al.*, 2014).

8. AtSSR16: SMALL SUBUNIT RIBOSOMAL PROTEIN 16, AT4G34620; was named as pAt2 in our 'moderate promoter' database for intermediate expressing control in transgenic Arabidopsis leaves.

9. HellFire: His₆-TEV-FLAG₃, a tandem epitope tag with 6× histidine, TEV protease cleavage site and 3× FLAG tag (Soleimani *et al.*, 2013); here we simplify it as HF.

10. AtCYSC1: CYSTEINE SYNTHASE C1, AT3G61440, was named as pAt3 in our 'moderate promoter' database for intermediate expressing control in transgenic Arabidopsis leaves.

11. HA₆: a tag with 6 tandem HA repeats (Gauss *et al.*, 2005).

12. PsAvrRps4: effector protein AvrRps4 from *Pseudomonas syringae* pv. *pisi*. Here this module can be placed with either wild-type or mutant AvrRps4 coding sequence.

13. BlmNeon: mNeonGreen protein, a bright monomeric green fluorescent protein derived from *Branchiostoma lanceolatum* (Shaner *et al.*, 2013).

14. AtuOcs: terminator of octopine synthase from Agrobacterium tumefaciens (Engler et al., 2014).

Supplementary Table S3

Supplementary Table S3 Golden Gate cloning modules used in this work.

Modules for Cloning	Modules for CloningTSL SynbioDescription of Inserts Name		Backbones	Overhangs	Overhangs for Ligation	
Level 1; Selection Cassettes	pICSL11015	See Table S2 FastRed module	pICH47732	TGCC	GCAA	
Level 1; Inducible Cassettes	pICSL11037	See Table S2 XVE module	pICH47742	GCAA	ACTA	
Level 0; Promoters + 5' Untranslated Regions (UTRs)	pICSL12028	AtSSR16 promoter, Col-0 allele	-	GGAG	AATG	
Level 0; Coding Sequence (CDS) Without A Stop Codon	pICSL80072	<i>AtRRS1-R</i> CDS from genomic DNA, Ws-2 allele, BbsI and Bpil sites are removed	-	AATG	TTCG	
Level 0; C-terminal Tag	pICSL50001	Hellfire tag, His ₆ -TEV-FLAG ₃	-	TTCG	GCTT	
Level 0; 3' UTRs and terminators	pICSL60019	AtRRS1-R terminator, Ws-2 allele	-	GCTT	CGCT	
Level 0; Promoters + 5' UTRs	pICSL12007	AtCysC1 promoter, Col-0 allele	-	GGAG	AATG	
Level 0; CDS Without A Stop Codon	pICSL80073	AtRPS4 CDS from genomic DNA, Col-0 allele, BbsI and Bpil sites are removed	-	AATG	TTCG	
Level 0; C-terminal Tag	pICSL50009	Human influenza hemagglutinin tag, HA ₆	-	TTCG	GCTT	
Level 0; 3' UTRs and terminators	pICH41414	CaMV 35S terminator	-	GCTT	CGCT	
Level 0; Promoters + 5' UTRs	pICSL12005	LexA inducible promoter		GGAG	AATG	
Level 0; CDS Without A Stop Codon	pICSL80070	AvrRps4 wild-type coding sequence (CDS) from <i>Pseudomonas syringae</i> , BbsI and Bpil sites are removed		AATG	TTCG	
Level 0; CDS Without A Stop Codon	pICSL80071	AvrRps4 CDS with KRVY135- 138AAAA substitutions, BbsI and Bpil sites are removed		AATG	TTCG	
Level 0; CDS Without A Stop Codon	pICSL80074	AvrRps4 CDS with E187A substitution, BbsI and Bpil sites are removed		AATG	TTCG	
Level 0; C-terminal Tag	pICSL50015	mNeonGreen fluorescent protein from <i>Branchiostoma</i> <i>lanceolatum</i> , Bbsl and Bpil sites are removed		TTCG	GCTT	
Level 1; Expression Cassettes	pICSL11162	See Table S2 RRS1-R-HF module	pICH47751	ACTA	TTAC	
Level 1; Expression Cassettes	pICSL11163	See Table S2 RPS4-HA module	pICH47761	TTAC	CAGA	
Level 1; Expression Cassettes	pICSL11164	See Table S2 LexA:AvrRps4- mNeon module	pICH47772	CAGA	TGTG	
Level 1; Expression Cassettes	pICSL11165	Similar to pICSL11164, but with KRVY135-138AAAA substitutions	pICH47772	CAGA	TGTG	
Level 1; Expression Cassettes	pICSL11166	Similar to pICSL11164, but with E187A substitution	pICH47772	CAGA	TGTG	

Supplementary Table S4

Primer Name	Directions	Nucleotide Sequence (5' to 3')						
	For real-time quantitative PCR							
AtEF1α_RT_Fw	forward	CAGGCTGATTGTGCTGTTCTTA						
AtEF1α_RT_Rv	reverse	GTTGTATCCGACCTTCTTCAGG						
AtICS1_RT_Fw	forward	CAATTGGCAGGGAGACTTACG						
AtICS1_RT_Rv	reverse	GAGCTGATCTGATCCCGACTG						
AtPR1_RT_Fw	forward	ATACACTCTGGTGGGCCTTACG						
AtPR1_RT_Rv	reverse	TACACCTCACTTTGGCACATCC						
AvrRps4_RT_Fw	forward	ATGACTCGAATTTCAACC						
AvrRps4_RT_Rv	reverse	GGTCCACCCAATAGGGATTTGGGTG						
		For cloning						
AvrRps4_dom_Fw	forward	GAGGGTCTCAAATGACTCGAATTTCAACCAGTTCAG						
AvrRps4_dom_Rv	reverse	GAGGGTCTCACGAACCTTGGTTGATTCTGCGGTCCTCG						
RPS4_dom_1_Fw	forward	agGAAGACAAAATGGAGACATCATCTATTTCCACTGTGGAgGAC						
RPS4_dom_1_Rv	reverse	agGAAGACAAGTCcTCATAGTCGTCGATAAAGAC						
RPS4_dom_2_Fw	forward	agGAAGACAAgGACAGAGGTCAACCTCTAGATG						
RPS4_dom_2_Rv	reverse	agGAAGACAAGTtTTCACCGCCTTCACAATTTCATTG						
RPS4_dom_3_Fw	forward	agGAAGACAAAaACAGCGTTGACCGGAATACCACCGG						
RPS4_dom_3_Rv	reverse	agGAAGACAAAtACACTGACAATATTAGGGCTGG						
RPS4_dom_4/5_Fw	forward	agGAAGACAAgtattccaagtgagttatgatgaattg						
RPS4_dom_4/5_Rv	reverse	agGAAGACAAcctccacttcagacaagtctagg						
RPS4_dom_6_Fw	forward	agGAAGACAAGAgGACGAAACGAGCTTAGACCGCGACCAC						
RPS4_dom_6_Rv	reverse	agGAAGACAATtTTCAGCGAACTACAGCCGTGTGCATCTAAGC						
RPS4_dom_7_Fw	forward	agGAAGACAAAAaACAGTTTCAAAGCCTTTGGCCCGTA						
RPS4_dom_7_Rv	reverse	agGAAGACAATaTCTTCATCTTTACTTTAAAGGTG						
RPS4_dom_8_Fw	forward	agGAAGACAAGAtAAGTCTTGGGTCGCATATACTTGTCC						
RPS4_dom_8_Rv	reverse	agGAAGACAAGAAtACATGGTCTAGCTCAATCTTATCTTT						
RPS4_dom_9_Fw	forward	agGAAGACAAaTTCATTGGATACACCAGTTG						
RPS4_dom_9_Rv	reverse	agGAAGACAAcgaaccGAAATTCTTAACCGTGTGCATGA						

Supplementary Table S4 Primers used in this study.

Supplementary Table S5

Supplementary Table S5 Statistical analysis results.

Figure 4C

Tukey multiple comparison of means **1hpi**

Condition 2	Diff ¹	Lower ²	Upper ²	p adjusted ³
SETI_eds1 E2	-0.33333330	-3.07588100	2.40921400	0.99814890
SETI_eds1 E2	-0.33333330	-3.07588100	2.40921400	0.99814890
SETI_eds1 E2	-0.66666670	-3.40921400	2.07588100	0.95879800
SETI_eds1 E2	0.00000000	-2.74254700	2.74254700	1.00000000
SETI_eds1 E2	-1.00000000	-3.74254700	1.74254700	0.81720430
SETI_eds1 Mock	0.00000000	-2.74254700	2.74254700	1.00000000
SETI_eds1 Mock	-0.33333330	-3.07588100	2.40921400	0.99814890
SETI_eds1 Mock	-0.33333330	-3.07588100	2.40921400	0.99814890
SETI_eds1 Mock	-0.66666670	-3.40921400	2.07588100	0.95879800
SETI_KRVYmut E2	-0.33333330	-3.07588100	2.40921400	0.99814890
SETI_KRVYmut E2	-0.33333330	-3.07588100	2.40921400	0.99814890
SETI_KRVYmut E2	-0.66666670	-3.40921400	2.07588100	0.95879800
SETI_KRVYmut Mock	-0.66666670	-3.40921400	2.07588100	0.95879800
SETI_KRVYmut Mock	-0.33333330	-3.07588100	2.40921400	0.99814890
SETI_WT E2	-1.00000000	-3.74254700	1.74254700	0.81720430
	Condition 2 SETI_eds1 E2 SETI_eds1 E2 SETI_eds1 E2 SETI_eds1 E2 SETI_eds1 Mock SETI_eds1 Mock SETI_eds1 Mock SETI_eds1 Mock SETI_eds1 Mock SETI_eds1 Mock SETI_KRVYmut E2 SETI_KRVYmut E2 SETI_KRVYmut E2 SETI_KRVYmut E2 SETI_KRVYmut Mock SETI_KRVYmut Mock SETI_KRVYmut Mock	Condition 2 Diff ¹ SETI_eds1 E2 -0.33333330 SETI_eds1 E2 -0.3333330 SETI_eds1 E2 -0.66666670 SETI_eds1 E2 -0.66666670 SETI_eds1 E2 -0.0000000 SETI_eds1 E2 -1.0000000 SETI_eds1 Mock 0.00000000 SETI_eds1 Mock -0.33333330 SETI_eds1 Mock -0.33333330 SETI_eds1 Mock -0.666666670 SETI_eds1 Mock -0.666666670 SETI_eds1 Mock -0.666666670 SETI_KRVYmut E2 -0.33333330 SETI_KRVYmut E2 -0.33333330 SETI_KRVYmut E2 -0.666666670 SETI_KRVYmut Mock -0.66666670 SETI_KRVYmut Mock -0.66666670 SETI_KRVYmut Mock -0.33333330	Condition 2 Diff ¹ Lower ² SETI_eds1 E2 -0.3333330 -3.07588100 SETI_eds1 E2 -0.3333330 -3.07588100 SETI_eds1 E2 -0.66666670 -3.40921400 SETI_eds1 E2 -0.66666670 -3.40921400 SETI_eds1 E2 0.00000000 -2.74254700 SETI_eds1 E2 -1.00000000 -3.74254700 SETI_eds1 Mock 0.00000000 -2.74254700 SETI_eds1 Mock -0.33333330 -3.07588100 SETI_eds1 Mock -0.33333330 -3.07588100 SETI_eds1 Mock -0.66666670 -3.40921400 SETI_eds1 Mock -0.666666670 -3.07588100 SETI_KRVYmut E2 -0.3333330 -3.07588100 SETI_KRVYmut E2 -0.66666670 -3.40921400 SETI_KRVYmut E2 -0.66666670 -3.40921400 SETI_KRVYmut Mock -0.66666670 -3.40921400 SETI_KRVYmut Mock -0.66666670 -3.40921400 SETI_KRVYmut Mock -0.33333330 -3.07588100 SETI_KRVYmut Mock -0.66666670 -3.40921400	Condition 2 Diff1 Lower2 Upper2 SETI_eds1 E2 -0.3333330 -3.07588100 2.40921400 SETI_eds1 E2 -0.3333330 -3.07588100 2.40921400 SETI_eds1 E2 -0.66666670 -3.40921400 2.07588100 SETI_eds1 E2 -0.66666670 -3.40921400 2.07588100 SETI_eds1 E2 0.0000000 -2.74254700 2.74254700 SETI_eds1 E2 -1.0000000 -3.74254700 2.74254700 SETI_eds1 Mock 0.00000000 -2.74254700 2.74254700 SETI_eds1 Mock -0.3333330 -3.07588100 2.40921400 SETI_eds1 Mock -0.33333330 -3.07588100 2.40921400 SETI_eds1 Mock -0.66666670 -3.40921400 2.07588100 SETI_kRVYmut E2 -0.3333330 -3.07588100 2.40921400 SETI_KRVYmut E2 -0.66666670 -3.40921400 2.07588100 SETI_KRVYmut E2 -0.66666670 -3.40921400 2.07588100 SETI_KRVYmut Mock -0.66666670 -3.40921400 2.07588100

<u>5hpi</u>

Condition 1	Condition 2	Diff	Lower	Upper	p adjusted
SETI_eds1 Mock	SETI_eds1 E2	-1.00000000	-8.05166000	6.05166000	0.99617570
SETI_KRVYmut E2	SETI_eds1 E2	-3.00000000	-10.05166000	4.05166000	0.71072430
SETI_KRVYmut Mock	SETI_eds1 E2	-3.33333330	-10.38499300	3.71832700	0.62047410
SETI_WT E2	SETI_eds1 E2	24.33333330	17.28167300	31.38499300	0.00000080
SETI_WT Mock	SETI_eds1 E2	2.33333330	-4.71832700	9.38499300	0.86763760
SETI_KRVYmut E2	SETI_eds1 Mock	-2.00000000	-9.05166000	5.05166000	0.92433520
SETI_KRVYmut Mock	SETI_eds1 Mock	-2.33333330	-9.38499300	4.71832700	0.86763760
SETI_WT E2	SETI_eds1 Mock	25.33333330	18.28167300	32.38499300	0.00000050
SETI_WT Mock	SETI_eds1 Mock	3.33333330	-3.71832700	10.38499300	0.62047410
SETI_KRVYmut Mock	SETI_KRVYmut E2	-0.33333330	-7.38499300	6.71832700	0.99998160
SETI_WT E2	SETI_KRVYmut E2	27.33333330	20.28167300	34.38499300	0.00000020
SETI_WT Mock	SETI_KRVYmut E2	5.33333330	-1.71832700	12.38499300	0.18678740
SETI_WT E2	SETI_KRVYmut Mock	27.66666670	20.61500700	34.71832700	0.00000020
SETI_WT Mock	SETI KRVYmut Mock	5.66666670	-1.38499300	12.71832700	0.14639040
SETI_WT Mock	SETI_WT E2	-22.00000000	-29.05166000	-14.94834000	0.00000250

<u>20hpi</u>

Condition 1	Condition 2	Diff	Lower	Upper	p adjusted
SETI_eds1 Mock	SETI_eds1 E2	-3.3333333	-12.656202	5.989535	0.8284027
SETI_KRVYmut E2	SETI_eds1 E2	-5.0000000	-14.322868	4.322868	0.4993132
SETI_KRVYmut Mock	SETI_eds1 E2	-5.3333333	-14.656202	3.989535	0.4350794
SETI_WT E2	SETI_eds1 E2	40.3333333	31.010465	49.656202	0.0000001
SETI_WT Mock	SETI_eds1 E2	-0.6666667	-9.989535	8.656202	0.9998581
SETI_KRVYmut E2	SETI_eds1 Mock	-1.6666667	-10.989535	7.656202	0.9889748
SETI_KRVYmut Mock	SETI_eds1 Mock	-2.0000000	-11.322868	7.322868	0.9755368
SETI_WT E2	SETI_eds1 Mock	43.6666667	34.343798	52.989535	0.0000000
SETI_WT Mock	SETI_eds1 Mock	2.6666667	-6.656202	11.989535	0.9218678
SETI_KRVYmut Mock	SETI_KRVYmut E2	-0.3333333	-9.656202	8.989535	0.9999954
SETI_WT E2	SETI_KRVYmut E2	45.3333333	36.010465	54.656202	0.0000000
SETI_WT Mock	SETI_KRVYmut E2	4.3333333	-4.989535	13.656202	0.6357179
SETI_WT E2	SETI_KRVYmut Mock	45.6666667	36.343798	54.989535	0.0000000
SETI_WT Mock	SETI_KRVYmut Mock	4.6666667	-4.656202	13.989535	0.5667788
SETL WT Mock	SETL WT E2	-41 000000	-50 322868	-31 677132	0.0000001

24hpi

2711pi					
Condition 1	Condition 2	Diff	Lower	Upper	p adjusted
SETI_eds1 Mock	SETI_eds1 E2	-3.666667	-13.298177	5.964843	0.7907177
SETI_KRVYmut E2	SETI_eds1 E2	-4.666667	-14.298177	4.964843	0.5976463
SETI_KRVYmut Mock	SETI_eds1 E2	-5.666667	-15.298177	3.964843	0.4072727
SETI_WT E2	SETI_eds1 E2	41.666667	32.035157	51.298177	0.0000001
SETI_WT Mock	SETI_eds1 E2	-1.333333	-10.964843	8.298177	0.9965824
SETI_KRVYmut E2	SETI_eds1 Mock	-1.00000000	-10.63151	8.63151	0.9991284
SETI_KRVYmut Mock	SETI_eds1 Mock	-2.00000000	-11.63151	7.63151	0.9787215
SETI_WT E2	SETI_eds1 Mock	45.333333	35.701823	54.964843	0.0000000
SETI_WT Mock	SETI_eds1 Mock	2.333333	-7.298177	11.964843	0.9593638
SETI_KRVYmut Mock	SETI KRVYmut E2	-1.00000000	-10.63151	8.63151	0.9991284
SETI_WT E2	SETI KRVYmut E2	46.333333	36.701823	55.964843	0.0000000
SETI WT Mock	SETI KRVYmut E2	3.333333	-6.298177	12.964843	0.8458559
SETI_WT E2	SETI_KRVYmut Mock	47.333333	37.701823	56.964843	0.0000000
SETI_WT Mock	SETI_KRVYmut Mock	4.333333	-5.298177	13.964843	0.664365
SETI_WT Mock	SETI_WT E2	-43.00000000	-52.63151	-33.36849	0.0000000

Figure 5A, B

Supplementary Table S5 (cont.)

PR1/EF1A in SETI_WT			• •		-	``
Group 1	Group 2	p ⁴	p adjusted	p.format⁵	p.signif ⁶	method
0h	Untreated	0.0173	0.035	0.0173	*	T-test
2h	Untreated	0.00493	0.028	0.0049	**	T-test
4h	Untreated	0.00258	0.026	0.0026	**	T-test
8h	Untreated	0.00352	0.028	0.0035	**	T-test

PR1/EF1A in SETI_KRVYmut

Group 1	Group 2	р	p adjusted	p.format	p.signif	method
0h	Untreated	0.248	0.25	0.24812	ns	T-test
2h	Untreated	0.062	0.19	0.06204	ns	T-test
4h	Untreated	0.0342	0.17	0.03423	*	T-test
8h	Untreated	0.0191	0.11	0.01912	*	T-test

ICS1/EF1A in SETI_WT

Group 1	Group 2	р	p adjusted	p.format	p.signif	method
0h	Untreated	0.075	0.11	0.07535	ns	T-test
2h	Untreated	0.0096	0.05	0.00969	**	T-test
4h	Untreated	0.00788	0.055	0.00788	**	T-test
8h	Untreated	0.00292	0.023	0.00292	**	T-test

ICS1/EF1A in SETI_KRVYmut

Group 1	Group 2	р	p adjusted	p.format	p.signif	method
0h	Untreated	0.00713	0.057	0.0071	**	T-test
2h	Untreated	0.182	0.36	0.1819	ns	T-test
4h	Untreated	0.0115	0.08	0.0115	*	T-test
8h	Untreated	0.0369	0.17	0.0369	*	T-test

Figure 6

0dpi

Tukey multiple comparison of means

Condition 1	Condition 2	Diff	Lower	Upper	p adjusted
Col-0 Mock	Col-0 E2	-0.009058683	-0.13119346	0.11307609	0.996022
SETI_WT E2	Col-0 E2	-0.031042838	-0.15317762	0.09109194	0.8730022
SETI_WT Mock	Col-0 E2	0.051462467	-0.07067231	0.17359724	0.6085929
SETI_WT E2	Col-0 Mock	-0.021984155	-0.14411893	0.10015062	0.9489591
SETI_WT Mock	Col-0 Mock	0.06052115	-0.06161363	0.18265593	0.4829018
SETI_WT Mock	SETI_WT E2	0.082505305	-0.03962947	0.20464008	0.2391907

T-test

Genotype	Group1	Group2	р	p.adj	p.format	p.signif
Col-0	Col-0 E2	Col-0 Mock	0.852	0.85	0.852	ns
SETI_WT	SETI_WT E2	SETI_WT Mock	0.0676	0.14	0.068	ns

3dpi

Tukey multiple comparison of means

Condition 1	Condition 2	Diff	Lower	Upper	p adjusted
Col-0 Mock	Col-0 E2	-0.08591854	-0.7827627	0.6109256	0.9854528
SETI_WT E2	Col-0 E2	-1.2506235	-1.9474676	-0.5537794	0.0003516
SETI_WT Mock	Col-0 E2	0.19040795	-0.5064362	0.8872521	0.8692549
SETI_WT E2	Col-0 Mock	-1.16470496	-1.8615491	-0.4678608	0.0007702
SETI_WT Mock	Col-0 Mock	0.27632649	-0.4205176	0.9731706	0.6877704
SETI_WT Mock	SETI_WT E2	1.44103146	0.7441873	2.1378756	0.0000637

T-test

1-1031						
Genotype	Group1	Group2	р	p.adj	p.format	p.signif
Col-0	Col-0 E2	Col-0 Mock	0.621	0.62	0.6213	ns
SETI_WT	SETI_WT E2	SETI_WT Mock	0.000905	0.0018	0.0009	***

- 1. diff: difference between means of the two groups
- 2. lower, upper: the lower and the upper end point of the confidence interval at 95% (default)
- 3. **p adjusted**: p-value after adjustment for the multiple comparisons.
- 4. p: p-value
- 5. p.format: formatted p value
- 6. p.signif: significance levels

Supplementary Figure S1



Supplementary Figure S1 GUS-staining activity of synthetic promoters in *N. benthamiana*.

Synthetic promoters At1-At6 were fused to β -glucuronidase (GUS) gene and infiltrated into *N. benthamiana* leaves. GUS expressed under the 35S promoter served as positive control. Mock treatment (infiltration with infiltration buffer) was used as negative control. Leaf samples were collected at 2 days post infiltration (dpi), and GUS staining was performed as in Materials and Methods.

Supplementary Figure S2



Supplementary Figure S2 Transient expression of Super-ETI (SETI) constructs in *N. benthamiana*.

(A) Schematc diagram of the SETI construct infiltration in *N. benthamiana*. Leaves were infiltrated with Agrobacterium containing SETI or SETI_E187A constructs. At 2dpi, leaves were re-infiltrated with 0.1% DMSO or 50µM E2 according to the diagram. Samples were taken 6h after infiltration with 0.1% DMSO or 50µM E2.

(B) Protein accumulation of RRS1-R-HF and RPS4-HA by transient expression in N. benthamiana. Crude extracts of leaf samples from (A) were immunoblotted with Flag antibody (IB:Flag) to detect RRS1-R-HF (black arrowhead) or HA antibody (IB:HA) to detect RPS4-HA(white arrowhead). Ponceau staining of Rubisco large subunits is the loading control.

(C) *AvrRps4* expression after induction with E2 for 4h in the SETI leaves. 5-week old SETI leaves were infiltrated with 50 μ M E2. Samples were collected at 0 and 4hpi for RNA extraction and subsequent qPCR. Expression level is presented as relative to *EF1a* expression. Each data point represents one technical replicate. Black line represents the mean of the technical replicates. This experiment was repeated three times independently with similar results.

Supplementary Figure S3



Col-0 T1-#12_T2-#14 Col-0 T1-#18_T2-#17



Supplementary Figure S3 SETI T2 lines grown under E2 treatment.

SETI-transformed

Arabidopsis transgenic seedlings were sown in GM either containing 50µM E2 or its solvent 0.1% DMSO. Images were taken at 14DAG. Further analysis of SETI lines were performed with the line T1-#8_T2-#4, indicated in red.