#### 1 Transcriptomic profiling uncovers novel players in innate immunity in Arabidopsis

- 2 thaliana
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#### 36 Abbreviations

DAMPs: Damage-associated molecular patterns; DEGs: differentially expressed genes;
MAMPs: microbe-associated molecular patterns; NGS: next generation sequencing; PRR:
pattern-recognition receptor; PEPR: AtPep-receptor; PP2: phloem protein 2; PTI: patterntriggered immunity

#### 41 Abstract

42 In this research a high-throughput RNA sequencing based transcriptome analysis technique 43 (RNA-Seq) was used to evaluate differentially expressed genes (DEGs) in the wild type 44 Arabidopsis seedling in response to flg22, a well-known microbe-associated molecular pattern 45 (MAMP), and AtPep1, a well-known peptide representing an endogenous damage-associated 46 molecular patterns (DAMP). The results of our study revealed that 1895 (1634 up-regulated 47 and 261 down-regulated) and 2271 (1706 up-regulated and 565 down-regulated) significant 48 differentially expressed genes in response to flg22 and AtPep1 treatment, respectively. Among 49 significant DEGs, we observed that a number of hitherto overlooked genes have been found to 50 be induced upon treatment with either flg22 or with AtPep1, indicating their possible 51 involvement in innate immunity. Here, we characterized two of them, namely PP2-B13 and 52 ACLP1. pp2-b13 and aclp1 mutants showed an increased susceptibility to infection by the 53 virulent pathogen *Pseudomomas syringae pv tomato mutant hrcC*-, as evidenced by an 54 increased growth of the pathogen in planta. Further we present evidence that the *aclp1* mutant 55 was deficient in ethylene production upon flg22 treatment, while the pp2-b13 mutant, was 56 deficient in ROS production. The results from this research provide new information to a 57 better understanding of the immune system in Arabidopsis.

58 Keywords: Arabidopsis thaliana, AtPep1, FLS2 receptor, defense genes, gene expression,
59 immune signaling, next generation sequencing, pattern recognition receptor, pattern-triggered
60 immunity, plant immunology, RNAseq, transcriptome

#### 61 Introduction

As sessile organisms, plants are constantly under attack by a broad range of different microbes<sup>1-7</sup>. In a co-evolutionary arms race between plants and pathogens, plants initially sense the presence of microbes by perceiving microbe-associated molecular patterns (MAMPs) via membrane-resident pattern recognition receptors (PRRs) that are located on the cell surface; such MAMP perception generally leads to pattern-triggered immunity (PTI) 1,4,8,10.

68 The model plant Arabidosis thaliana can detect a variety of MAMPs, including fungal chitin 69 and bacterial elicitors such as flagellin and elongation factor-Tu (EF-Tu), or their respective peptide surrogates flg22 and elf18<sup>8-10</sup>. Flagellin and EF-Tu are perceived by FLS2 and EFR 70 71 receptors, respectively. Besides MAMPs, molecular patterns derived from the plant upon 72 pathogen attack can also trigger an immunity response. Examples of such damage-associated molecular patterns (DAMPs) are members of the family of AtPeps, recently discovered 73 74 endogenous and highly conserved peptides in A. thaliana. The different AtPeps (AtPep1-8) 75 originate from the conserved C-terminal portion of their respective precursors AtPROPEP1-8 <sup>11-14</sup>. The plant cell surface PRRs AtPEPR1 and AtPEPR2 have been identified as the AtPeps 76 receptors<sup>12,16,17</sup>. 77

DAMP/MAMP perception triggers a vast array of defense responses<sup>1,12</sup>. These include the 78 production of reactive oxygen species (ROS) in an oxidative burst<sup>18,19</sup>, the multi-level specific 79 reprogramming of expression profiles at transcriptional and also post-transcriptional levels<sup>20-</sup> 80  $^{23}$ , and downstream defense responses, including callose deposition<sup>24</sup>, MAP kinase activation, 81 82 synthesis of the defense hormone salicylic acid (SA), and seedling growth inhibition<sup>25</sup>. 83 MAMP treatment prior to the actual pathogen attack results in enhanced resistance to adapted 84 pathogens, and it has been observed that mutants impaired in MAMP recognition display enhanced susceptibility, not only to adapted but also to non-adapted pathogens<sup>10,19,23,26</sup>. This 85 86 indicates a contribution of pattern-triggered immunity (PTI) to both basal and non-host resistance, highlighting the importance of PTI in plant innate immunity<sup>27-30</sup>. 87

The proteobacterium *Pseudomonas syringae* is a bacterial leaf pathogen that causes destructive chlorosis and necrotic spots in different plant species, including monocots and dicots. *P. syringae* pathovars and races differ in host range among crop species and cultivars, respectively<sup>6,31</sup>. Many strains of *P. syringae* are pathogenic in the model plant *A. thaliana*, which makes *P. syringae* an ideal model to investigate plant–pathogen interactions<sup>31-33</sup>. The

ability of *P. syringae* to grow in plants and to multiply endophytically depends on the typethree secretion system (T3SS). T3SS enables the secretion into the cytoplasm of the plant cell
of effector proteins, which suppress or, in some cases, change plant defense responses<sup>34</sup>. *P. syringae* encodes 57 families of different effectors injected into the plant cell by the T3SS<sup>32</sup>.
Effectors inside plant cells are recognized by R proteins, which constitute the second level of
defense known as effector-triggered immunity (ETI)<sup>1,20,35</sup>.

99 PTI response is controlled by a complex, interconnected signaling network, including many 100 transcription factors (TFs); interference with this network can paralyze the adequate response upon pathogen infection<sup>36,37</sup>. A large fraction of genes in the plant genome respond 101 transcriptionally to pathogen attack<sup>21,38</sup>. In addition to specific reprogramming of 102 103 transcription, post-transcriptional regulation also plays a role in the plant immune response<sup>39</sup>. 104 The advent of advanced sequencing and proteomics technologies has led to the identification 105 of many novel players in defense signaling pathways and their characterization as important 106 components of innate immunity in Arabidopsis. However, for a fundamental understanding of 107 the plant's defense system and its response to pathogens, it is necessary to fill the remaining 108 gaps by further identifying genes and proteins involved in plant immunity<sup>1</sup>.

109 The highly conserved 22-amino-acid fragment (flg22) of bacterial flagellin that is recognized 110 by the FLS2 PRR can activate an array of immune responses in Arabidopsis<sup>1-4</sup>. In addition, resistance to *Pst* DC3000 is induced by pre-treatment with  $flg22^{1-4,9,23}$ . Previous studies 111 112 investigating flg22-induced transcriptional changes showed that among highly induced genes, 113 there were several with functions in the Arabidopsis immune pathway that had previously not been associated with immunity<sup>23,40,41</sup>. These studies profiled only a part of the Arabidopsis 114 115 gene space, and one can therefore speculate that a whole-genome transcriptome profiling of 116 elicitor-treated Arabidopsis plants would unveil additional new players in the immune 117 signaling system. Furthermore, given that both the MAMP flg22 and the DAMPs AtPeps 118 trigger immunity, analyzing their respective effects side by side in one coherently designed 119 experiment could increase the power to detect shared features and specific responses of the 120 respective immune response pathways.

Here, we performed whole-genome transcriptome profiling by RNA sequencing  $(RNA-seq)^{42}$ <sup>45</sup> of Arabidopsis seedlings treated with either flg22 or AtPep1 treatments. Filtering for genes induced in both treatments and those missing in previously published assays, we selected 85 candidate genes to be investigated for their role in plant immune response and systematically tested T-DNA insertion mutants of these genes for susceptibility towards *Pst*. For two loci,

#### 126 PHLOEM PROTEIN 2-B13 (PP2-B13) and ACTIN CROSS-LINKING PROTEIN 1 (ACLP1),

127 we identified mutant lines with altered pathogen response phenotypes and characterized these

- 128 genes as novel players in early PTI responses.
- 129 **Results**

#### 130 Whole-genome transcriptional profiling identifies two novel factors of PTI

131 To dissect transcriptional responses in response to flg22 and AtPep1, we extracted total RNA 132 from mock- and elicitor-treated one-week-old Arabidopsis plants and performed RNA-seq 133 transcriptome analysis on three biological replicates per treatment (Supplementary File S1 and 134 S2). Samples were collected 30 min after elicitor treatment. We used the R package DESeq2<sup>46</sup> 135 for differential gene expression analysis; all differentially expressed genes (DEGs) can be 136 found in Supplementary Files S3-S6. In response to flg22, we detected a total of 1,895 DEGs 137 compared to the control treatment (Fig. 1A), of which 1,634 genes were up- and 261 were 138 down-regulated in the flg22-treated seedlings (Supplementary Files S3 and S5). Treatment 139 with AtPep1 resulted in 2,271 DEGs, with 1,706 up-regulated and 565 down-regulated (Fig. 140 1A). When comparing the two treatments with each other, we detected only 511 DEGs, with 141 similar fractions of up- and down-regulated genes (265 and 246, respectively, in flg22 vs. 142 AtPep1) (Fig. 1A). Taken together, these results indicated that AtPep1 treatment causes 143 slightly more genes to be differentially regulated than flg22, and that the transcriptional 144 profiles are more similar between flg22- and AtPep1-treated samples than between either of 145 the treatments and the control. While a remarkable 70% of flg22-up-regulated genes were also 146 induced by AtPep1, 256 genes were exclusively up-regulated in response to flg22, while 328 147 were exclusively up-regulated in response to AtPep1 (Fig. 1; panel B). Of genes down-148 regulated upon flg22 treatment, only 23% were also down-regulated in response to AtPep1; 149 107 genes were exclusively down-regulated by flg22 treatment, vs. 411 genes by AtPep1 (Fig. 150 1; panel C). Detailed information on DEGs from different comparisons are presented in 151 Supplementary Files S1 to S16.

152 In response to flg22, the expression levels of *PP2-B13* and of *ACPL1* were 126-fold and 20-

153 fold induced, respectively (Fig. 1D). Similarly, AtPep1 treatment leads to 120-fold up-

regulation of *PP2-B13* and a 10-fold up-regulation of *ACLP1* (Fig. 1E).

Former studies showed that treatment of Arabidopsis seedlings with flg22 triggers robust PTIlike responses at the transcriptional level, activating ca. 1,000 genes that may have functions in PTI responses<sup>23,40,41</sup>. However, because these experiments were done using the ATH1 microarray, which does not cover all Arabidopsis protein-coding genes, we speculated that 159 there might be additional, so far unknown PTI-related genes affected by flg22 and other elicitors. Denoux et al., (2008)<sup>41</sup> performed a comprehensive microarray (Affymetrix ATH1) 160 161 transcript analysis in response to flg22 treatment. However, our RNA-seq analysis revealed 162 3,297 genes induced (without fold-change cutoff) in response to flg22 treatment that had not 163 been present on the ATH1 chip (Supplementary File S7). Comparing the upregulated DEGs 164 results in RNA-seq experiment analysis with fold change cutoff (adjusted p-value < 0.05 and 165 a minimum two-fold change) among the genes which are also present in ATH1 affymetrix 166 genechip showed that 1366 upregulated DEGs are present in both RNA-seq experiment and 167 ATH1 affymetirx genechip (Supplementary File S8). While our analysis showed that 268 168 genes with fold change cutoff are exclusively upregulated in RNA-seq analysis which were 169 not present in ATH1 affymetrix genechip and their expression only investigated in RNA-seq 170 analysis (Supplementary File S9). To identify yet unknown PTI factors, we first discarded all 171 genes from our list of DEGs that had been present on the ATH1 microarray chip and hence 172 would have been detected in the above-mentioned studies.

173 We then ranked the remaining DEGs by fold change induction (high induction of transcription 174 in response to both flg22 and AtPep1 treatments) and selected the 85 most strongly up-175 regulated genes as candidates (Supplementary File S10). Finally, we decided to focus on a 176 small set of genes that showed highest induction after flg22 treatment (Table 1). We checked 177 for availability of T-DNA insertion mutants for these genes and retrieved mutant lines for 178 AT1G56240, AT1G65385, AT4G23215, AT1G59865, AT1G24145, AT2G35658, 179 AT1G69900, AT2G27389, and AT1G30755. We confirmed homozygous T-DNA insertions 180 via PCR.

To test whether any of the candidate genes might play a role in immunity, we tested all homozygous T-DNA mutant lines for bacterial growth of the mutant pathogenic strain *P. syringae* pv. *tomato hrcC-* (*Pst hrcC-*), which is defective in T3SS. Comparing the bacterial growth titer in the mutant plants to that of wild-type Col-0 revealed that two of the lines, namely SALK\_144757.54.50 and SALK\_68692.47.55, showed significantly better bacterial growth (*P*-value = 0.0261 and 0.0089, respectively; Student's *T*-test) (Fig. 4), and that the underlying loci might play a role in defense signaling.

#### 188 Expression of the *PP2-B13* and *ACLP1* genes is induced following flg22 treatment

According to the Arabidopsis Information Resource<sup>47</sup> and the SIGnAL database (http://signal.salk.edu/), the predicted T-DNA insertion site in SALK\_144757.54.50 is located in the second of three exons of *PP2-B13*<sup>48</sup> (Fig. 3A); the T-DNA insertion in SALK\_68692.47.55 is located in the first of two exons of *ACLP1*. We confirmed that the T-

193 DNA insertion lines were null alleles for pp2-b13 and aclp1, respectively, via reverse-194 transcription polymerase chain reaction (RT-PCR) (Fig. 3B). PP2-B13 and ACLP1 transcripts 195 were not detectable in the respective T-DNA insertion lines; we therefore refer to these lines 196 as *pp2-b13* and *aclp1*, respectively. Visual inspection of plant growth did not reveal any 197 obvious phenotypic differences between any of the two insertion lines and wild-type Col-0 198 with regard to size and shape at the rosette stage. Two days post infection with Pst hrcC-, 199 neither *pp2-b13* nor *aclp1* showed symptoms different to this of with wild-type Arabidopsis 200 (Supplementary Fig. S4).

201 As can be seen in the volcano plot in Figure 1D, gene expression levels of PP2-B13 and 202 ACLP1 were strongly induced by flg22. To further monitor the gene expression of PP2-B13 203 and ACLP1 upon elicitor perception and to validate the RNA-seq results, we analyzed 204 expression levels by quantitative real-time PCR (qRT-PCR) in leaves of four-week-old 205 Arabidopsis plants at different time points. We confirmed that also at this later developmental 206 stage, expression of PP2-B13 and ACLP1 was strongly induced (100-fold for PP2-B13 and 207 12-fold for ACLP1) within 30 minutes after flg22 treatment (Fig. 2). Two and six hours after 208 elicitor treatment, expression levels of PP2-B13 had returned to pre-treatment levels, while 209 those of ACLP1 remained only slightly elevated (Fig. 2). This expression pattern suggests that 210 both genes might be involved in early defense response.

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## Increased susceptibility to *Pseudomonas syringae* pv. *tomato* mutant *hrc*C- in *pp2-b13*and *aclp1* mutant lines

Two days post inoculation of leaves with *Pst hrcC*-, the bacterial titer for wild type Arabidopsis reached 109,000 cfu/cm<sup>2</sup>, while for *pp2-b13* mutant lines it increased significantly to 325,000 cfu/cm<sup>2</sup> (p = 0.0261), albeit not as drastically as that of *sid2-2* mutants. The protein encoded by *PP2-B13* is a phloem protein containing the F-box domain Skp2. It also has a described function in carbohydrate binding<sup>48</sup>.

The protein encoded by *ACLP1* is of unknown function with the highest similarity to actin cross-linking proteins and includes a fascin domain. As it can be seen in Fig. 4, 48 hours post inoculation of leaves with *Pst hrcC*-, the bacterial titer for wild type Arabidopsis, reached 109000 cfu/cm<sup>2</sup> while for the *aclp1* mutant line, it increased significantly to 257000 cfu/cm<sup>2</sup> (p = 0.0089).

- In conclusion, these results suggest that PP2-B13, and ACLP1 play a role in defense signaling
- and that both genes are required for wild-type levels of resistance against *Pst hrcC*-.

### Differential ethylene production in *pp2-b13* and *aclp1* plants, as compared to the wild type Arabidopsis

To analyze the early defense responses upon elicitor treatment, we assessed ethylene (ET) production in response to flg22 treatment in the mutant lines *pp2-b13*, and *aclp1*. We observed that mutant line *aclp1* displayed a significantly reduced ET production in comparison to wild-type Arabidopsis upon treatment with 1  $\mu$ M flg22 (*p*-value = 0.0295; Fig. 5). This suggests that ACLP1 is involved in the enhancement of ET production in response to flg22 perception.

## Differential Reactive Oxygen Species Generation in *pp2-b13* and *aclp1* plants, as compared to the wild type Arabidopsis

236 One of the early responses triggered by MAMPs and DAMPs is the production of apoplastic 237 ROS by the Arabidopsis NADPH-oxidases RbohD and RbohF protein<sup>19</sup>. We observed that in 238 the treated leaf discs upon flg22 perception, pp2-b13 displayed a lower ROS production 239 compared to wild-type (Fig. 6A), indicating that PP2-B13 might play a role in early PTI by 240 enhancing the oxidative burst in response to the flg22 perception. In contrast, *aclp1*, although 241 exhibiting deficiency in ET production upon flg22 perception, showed robust enhancement of 242 ROS production at levels similar to that of wild-type (Fig. 6A). 243 When comparing maximum ROS production the two mutant lines and wild-type, we did not

observe statistically significant differences, in contrast to the fls2 mutant which displayed noROS production at all in response to flg22 (Fig. 6B).

# FLS2 receptor abundance in *pp2-b13* and *aclp1* mutants were similar to the wild type Arabidopsis

The *PP2-B13* and *ACLP1* genes were strongly induced upon elicitor treatment, as seen in the RNA-seq and qPCR data. Additionally, both mutant lines were deficient in early PTI responses (ET and ROS measurement). Hence it is conceivable that the products of the *PP2-B13* and *ACLP1* genes affect the abundance of FLS2 receptor. However, FLS2 analysis via immunoblots showed that both mutant lines had similar levels of FLS2 as the wild-type (Fig. 7), indicating that these genes do not play a role in regulating the abundance of the FLS2 receptor.

#### 255 **Discussion**

Plants are under constant exposure to microbial signals from potential pathogens, potential commensals, and mutualists. The plant cell immune sensors are able sense these signals and expand the defense against pathogens<sup>1,20,49-50</sup>. Host-pathogen interactions encompass a

complex set of events that are dependent on the nature of the interacting partners,
developmental stage, and environmental conditions<sup>1,50-51</sup>. These interactions are regulated
through diverse signaling pathways that ultimately result in altered gene expression<sup>1,23,42</sup>.
Membrane-resident pattern recognition receptors (PRRs) that are located on the cell surface
can sense and perceive microbe-derived signature components known as microbe-associated
molecular patterns (MAMPs) and also damage-associated molecular patterns (DAMPs),
leading to pattern-triggered immunity<sup>1-4</sup>.

- 266 In the current study, global gene expression profiling of wild type Arabidopsis seedlings 267 resulted in the identification of a large number of genes induced by flg22 and AtPep1 that had 268 not been detected by the ATH-1 array technology. Among them, we focused on two, namely 269 PP2-B13, and ACLP1. We observed noticeable up-regulation in wild type Arabidopsis for 270 both of these genes upon flg22 treatment (Fig. 2). Reverse-genetic studies of PP2-B13 and 271 ACLP1 genes showed that these genes are required to control infection by the bacterial 272 pathogens P. syringae py. tomato mutant hrcC- (Pst hrcC-; Fig. 4). Our results highlight the 273 general usefulness of transcriptomic approaches to identify new players in early defense 274 responses in innate immunity and reveal two new players, PP2-B13 and ACLP1, in this 275 pathway. It should be noted that extending the time points of the elicitor treatment in future 276 studies might help uncover additional players in innate immunity.
- PP2-B13<sup>48</sup> is an F-box protein with homology to PP2-B14<sup>52</sup>. The F-Box domain of PP2-B13
  is close to the N-terminus of the protein. PP2-B13 shows the highest similarity in amino acid
  sequence with AT1G56250, which formerly was reported as an F-box protein<sup>52</sup>. Zhang *et al.*,
  (2011)<sup>53</sup> showed that PP2-B13 and PP2-B14 were highly abundant in phloem upon aphid
  infection. These genes are located in a cluster of defense-related genes, which supports that
  hypothesis that they play a role in the defense signaling network.
- 283 Sequence alignment of PP2-B13 with homologues from other plant species revealed 284 conserved features (Fig. 8). A phylogenetic analysis supported high conservation of PP2-B13-285 like proteins across different plant species, suggesting similar function (Supplementary Fig. S7). In silico structural analysis using Raptor X<sup>54</sup> predicted two domains (Supplementary Fig. 286 287 S5): an N-terminal F-box domain (residues 4-46; Fig. 8) and a C-terminal PP2 domain 288 (residues 93-280; Fig. 8). PP2-domain proteins are one of the most abundant and enigmatic proteins in the phloem sap of higher plants<sup>55-56</sup>. It was reported that lectin domain proteins are 289 290 important in plant defense responses, and so far 10 membrane-bound lectin type PRRs, which are involved in plant defense signaling and symbiosis, have been identified<sup>56</sup>. Recently, 291 Eggermont et al.  $(2017)^{57}$  showed that lectins are linked to other protein domains which are 292

293 identified to have a role in stress signaling and defense.

294 Lectins are proteins containing at least one non-catalytic domain which enables them to 295 selectively recognize and bind to specific glycans that are either present in a free form or are 296 part of glycoproteins and glycolipids and help the plants to sense the presence of pathogens; as a defense response they use a broad variety of lectin domains to interact with pathogens<sup>58</sup>. 297 Additionally, Eggermont et al., (2017)<sup>57</sup> showed that among lectin proteins, the amino acids 298 299 responsible for carbohydrate binding are highly conserved. Furthermore, Jia et al (2015)<sup>58</sup>, showed that PP2-B11. (another member of the phloem lectin proteins<sup>48</sup>) is highly induced in 300 301 response to salt treatment at both transcript and protein levels. They showed that PP2-B11 302 plays a positive role in response to salt stress.

303 In order to predict PP2-B13 interaction partners, we submitted the PP2-B13 amino acid 304 sequence to the STRING database (version 11.0), which hypothetically determines proteinprotein interactions based on computational prediction methods<sup>59</sup>. This returned several major 305 306 players in innate immunity, specifically PBL1, RLP6 and RLP15, which are important 307 defense proteins, as potential interaction partners (Fig. 9)<sup>60-62</sup>. RLPs are regarded major players in immune system in Arabidopsis<sup>60-62</sup>. STRING also predicted interactions of PP2-308 309 B13 with major zinc transporter proteins (ZIPs), which have role in biotic and abiotic stress responses<sup>63</sup>. 310

In the region of the chromosome 1 where PP2-B13 is located, there are many genes which are activated upon biotic or abiotic stresses, such as AT1G56280. The protein product of this gene is named drought-induced protein 19 (Di19), because its expression increases due to progressive drought stress<sup>64</sup>. Importantly, we have found that the *WRR4* gene (*AT1G56510*) is downstream of the *PP2-B13* (Supplementary Fig. S2). WRR4 is one of the most important defense gene in *Arabidopsis thaliana*<sup>64-65</sup>.

ACLP1 is an actin cross-linking protein of 397 amino acids. Raptor X<sup>54</sup> predicted two Fascin 317 318 motifs in the N-terminal and C-terminal domains (residues 18-70 and 229-318, respectively; 319 10; Supplementary Fig. S6). The conserved domain database at NCBI Fig. 320 (https://www.ncbi.nlm.nih.gov/Structure/cdd/cdd.shtml) also identified two fascin domains in 321 ACLP1. Fascins are a structurally unique and evolutionarily highly conserved group of actin 322 cross-linking proteins. Fascins function in the organization of two major forms of actin-based structures: dynamic, cortical cell protrusions and cytoplasmic microfilament bundles<sup>67-69</sup>. For 323 324 ACLP1 the sequence Logo was created. As shown in the Figure 10, there are several

conserved regions in the ACLP1 and its homologues. Furthermore, a phylogenetic analysis
 supported high conservation of ACLP1-like proteins across different land plant species,
 suggesting similar function (Supplementary Fig. 9).

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MAMP perception changes actin arrangements and leads to cytoskeleton remodeling<sup>70-71</sup>. 329 Cytoskeleton rapidly responds to biotic stresses to supports cellular fundamental processes<sup>72-</sup> 330 <sup>74</sup>. Recently, Henty-Ridilla et al., (2014)<sup>75</sup> confirmed that Actin depolymerizing factor 4 331 (ADF4) has an important role in defense response through cytoskeleton remodeling. They 332 333 showed that the *adf4* mutant was unresponsive to a bacterial MAMP<sup>76</sup>. Using the STRING 334 database (version 11.0), we predicted many actin related proteins including ADF4, ACT2, 335 ACT12, PFN2, MRH2, ARK2 and ADF1 as putative interaction partner for ACLP1 (Fig. 9), 336 further corroborating a potential role for ACLP in defense-related actin reorganization. It is 337 noteworthy that downstream of the ACLP1, there is PP2-A5 gene (Supplementary Fig. S3). 338 The protein product of PP2-A5 gene is another member of the Phloem Protein 2 family. The 339 role of PP2-A5 in defense response against insect is already confirmed<sup>77</sup>.

#### 340 Conclusions and outlook

341 We observed a strong (>100fold) and very rapid but transient induction of PP2-B13 and 342 ACLP1 within 30 min of flg22 elicitor treatment (Fig. 2). Using a mutant approach, we 343 provide evidence that loss-of-function mutations in *PP2-B13* and *ACLP1* can affect the early 344 PTI responses including ET and ROS measurements (Fig. 5 and Fig. 6). We could show a 345 defect in activation of ET production for *aclp1* plants and also attenuated ROS generation in 346 pp2-b13 plants in response to flg22 treatment. ROS accumulation is regarded as an early PTI event occurring a few minutes after *Pst* inoculation<sup>1</sup>. These findings suggest that these genes 347 348 might have a function through interaction with PTI signaling pathways during bacterial 349 infection. However, we cannot yet determine at what point of the MAMP signaling cascade 350 the products of these genes function. Therefore, subsequent studies are needed to determine 351 the relationship of these genes in MAMP recognition and other signaling cascades in innate 352 immunity.

Furthermore, one important finding in this study is that our study reconfirm the importance of the chromosome 1 in innate immunity as there many resistant genes that their protein product have role in defense including ACLP1, Di19, PP2-A5, PP2-B13, WWR4 and VBF. Therefore, we suggest that in further studies, this region of the chromosome 1 should be evaluated in depth to identify more genes which have role in innate immunity.

358 In conclusion, based on what we have observed in different experiments, it can be concluded

that PP2-B13 and ACLP1 have a role in innate immunity. It is likely that the protein products

360 of these genes can have multiple functions in innate immunity in Arabidopsis. It has been

361 previously reported that the genes which have a function in innate immunity in Arabidopsis

362 can also have role in resistance against abiotic stress<sup>78</sup>. Hence, it will be interesting to see the

- 363 response of these mutant plants upon abiotic stresses such as salinity, cold, and drought.
- 364 Materials and Methods

#### 365 Plant material and growth conditions

366 All Arabidopsis genotypes were derived from the wild-type accession Columbia-0 (Col-0). 367 The plants were grown as one plant per pot at 10 h photoperiod light at 21°C and 14 h dark at 368 18°C, with 60% humidity for 4 to 5 weeks, or were grown on plates containing Murashige 369 and Skoog (MS) salts medium (Sigma, Aldrich), 1% sucrose, and 1% agar with a 16 h 370 photoperiod. Seeds of the sid2 mutant line were kindly provided by Jean-Pierre Métraux 371 (University of Fribourg). The *fls2* mutant line was previously published<sup>23</sup>. pp2-b13(AT1G56240; SALK\_144757.54.50), and aclp1 (AT1G69900; SALK\_68692.47.55) were 372 373 obtained from the Nottingham Arabidopsis Stock Centre (NASC).

#### 374 **Peptide treatments**

375 The peptides used as elicitors were flg22 (QRLSTGSRINSAKDDAAGLQIA), and AtPep1 376 (ATKVKAKQRGKEKVSSGRPGQHN). The peptides were ordered from EZBiolabs 377 (EZBiolab Inc., IN, USA), dissolved in a BSA solution (containing 1 mg/mL bovine serum 378 albumin and 0.1 M NaCl), and kept at -20°C. In order to prepare sterile seedlings, 379 Arabidopsis seeds were washed with 99% ethanol supplemented with 0.5% Triton for 1 min, 380 washed with 50% ethanol supplemented with 0.5% Triton for 1 min, then washed with 100% 381 ethanol for 2 min. Seeds were sown on MS salt medium supplemented with 1% sucrose and 382 0.8% Phytagel (Sigma-Aldrich) at pH 5.7. Subsequently, the plates were stratified for 2 d at 383 4°C and germinated at 21°C under continuous light (MLR-350; Sanyo chamber). One day 384 before treatment, the seedlings were moved from plates to ddH<sub>2</sub>O. One-week-old Arabidopsis 385 seedlings were treated with AtPep1 and flg22 (1  $\mu$ M) for 30 min. BSA solution was used for 386 the mock-treated control.

#### 387 **RNA isolation, Illumina sequencing and quality control**

388 Total RNA was isolated from one-week-old Arabidopsis seedlings using the RNeasy Plant 389 Mini Kit (Qiagen), according to the manufacturer's protocol. Three individual biological 390 replicates were used per condition. RNA purity, concentration, and integrity were determined 391 via spectrophotometric measurement on a NanoDrop 2000 (Thermo-Scientific). Libraries 392 were prepared using the RNA sample preparation kit (Illumina) according to the 393 manufacturer's instructions (Illumina). Libraries were sequenced on a HiSeq2000 instrument 394 (Illumina) as 100 bp single-end reads. Sequencing quality of the fastq files from the RNA-Seq 395 data was examined by FastQC software (version V0.10.1; 396 http://www.bioinformatics.babraham.ac.uk/projects/fastqc/). Adapter sequences were clipped 397 and low quality reads were either trimmed or removed.

## 398 Mapping reads to the reference genome and analysis of differentially expressed genes399 (DEGs)

400 RNA-seq reads were aligned against the A. thaliana cDNA reference genome (TAIR10; 401 (https://www.arabidopsis.org/). The reference genome index was constructed with Bowtie 402 v2.2.3 and reads were aligned to the Arabidopsis reference genome using TopHat v2.0.12 with default parameters<sup>46</sup>. The resulting alignments were visualized using Integrative 403 Genomics Viewer (IGV)<sup>79</sup>. To evaluate differentially expressed genes between elicitor-treated 404 and control samples, we used the DESeq2 R package<sup>80-81</sup>. Genes with an adjusted p-value <405 406 0.05 and a minimum two-fold change in expression were considered as differentially 407 expressed.

#### 408 Selection of candidate genes

409 Because we were interested in genes not yet classified as related to immune response, we 410 applied several filters: from the genes significantly up-regulated after 30 minutes of flg22 or 411 AtPep1 treatment, we discarded those which had previously been reported as differentially regulated and implicated in biotic and abiotic stress response<sup>23,40,41</sup>. We selected a subset of 85 412 413 genes (Supplementary File 10) based on the following criteria: 1) high induction of 414 transcription in response to both flg22 and AtPep1 treatments, 2) not present on Affymetrix 415 ATH-22k microarray chips, 3) no published function or at least not connected to defense, and 416 4) not a member of a large gene family (in order to avoid potential functional redundancy). 417 From this list, we eventually selected 20 genes as candidate genes for further analyses (Table 418 and ordered corresponding T-DNA insertion lines (http://signal.salk.edu/cgi-1) 419 bin/tdnaexpress) from NASC (www.arabidopsis.info).

#### 420 Determination of gene expression by quantitative real-time RT-PCR analysis

421 Discs of leaves of four-week-old Arabidopsis plants were cut out using a sterile cork borer 422 (d=7mm) and placed overnight in ddH<sub>2</sub>O in 5 cm Petri dish. Thereafter, the experiment 423 started (time zero) with the addition of 1  $\mu$ M flg22, dissolved in BSA solution (1 mg/mL 424 bovine serum albumin and 0.1 M NaCl). BSA solution without flg22 was used for the mock-425 treated control., In order to produce a time course in response to flg22 treatment, the 426 experiment was stopped after 30 min, 2 h and 6 h, Total RNA from leaves of four-week-old 427 Arabidopsis plants was extracted using the NucleoSpin RNA plant extraction kit (Macherey-428 Nagel) and treated with rDNase according to the manufacturer's extraction protocol. RNA 429 quality of all samples was assessed using NanoDrop 2000 (Thermo-Scientific). To synthesize 430 the cDNA, 10 ng of RNA was used with oligo (dT) primers and AMV reverse transcriptase 431 and reverse transcription was performed according to the manufacturer's instructions 432 (Promega). Using a GeneAmp 7500 Sequence Detection System (Applied Biosystems), 433 quantitative RT-PCR was performed in a 96-well format. The gene-specific primers used in 434 this study are listed in Supplementary Table S2. Expression of UBO10 (AT4G05320), which has been validated for gene expression profiling upon flg22 treatment<sup>82-84</sup>, was used as the 435 436 reference gene. Based on  $C_T$  values and normalization to UBO10 (AT4G05320) expression, the expression profile for each candidate gene was calculated using the qGene protocol<sup>83-84</sup>. 437

#### 438 Analysis of T-DNA insertion mutants

439 After grinding leaf material in liquid nitrogen, total DNA was extracted using EDM-Buffer 440 (200 mM Tris pH7.5; 250 mM NaCl, 25 mM EDTA; 0.5% SDS). Putative T-DNA insertion 441 mutants were genotyped by PCR. We designed gene specific primer pairs LP and RP based 442 on the predicted genomic sequence surrounding the T-DNA insertion (Supplementary Table 443 S2). The plants were considered homozygous mutants if there was a PCR product with T-444 DNA-specific border primers LP/ LBa1 but not with the LP/RP primers. (Table 1). We 445 obtained T-DNA insertion mutants of six single homozygous lines bearing a disruption in the 446 gene, including AT1G56240 (*PP2-B13*) and AT1G69900 (*ACLP1*) (Table 1).

#### 447 **RT-PCR experiment**

448 For total RNA extraction, samples of leaf tissue from 4-week-old Arabidopsis including wild 449 type plants (Col0), *pp2-b13*, and *aclp1* were harvested into liquid nitrogen and were grounded

- 450 with the sterile mortar and pestle. The NucleoSpin RNA Prep Kit (BioFACT<sup>TM</sup>, South Korea)
- 451 was used for RNA extraction according to the manufacturer's instructions and DNase-treated.

452 Reverse transcription was performed at 50°C for 45 minutes using total RNA, a reverse transcriptase (BioFACT<sup>TM</sup>, South Korea) and an oligo (dT)20 primer (BioFact, South Korea) 453 supplemented with 0.5ul RNase inhibitor (BioFACT<sup>TM</sup>, South Korea) and according to the 454 455 manufacturer's instructions. To ensure specificity and accuracy of each primer and to design 456 the highly specific primers for *PP2-B13* and *ACLP1* transcripts, the oligonucleotide primers were designed by AtRTPrimer program<sup>85</sup> which exclusively determine specific primers for 457 each individual transcript in Arabidopsis. The housekeeping gene ACTIN2 was used as a 458 459 positive control for each PCR. The primers for ACTIN 2 transcript were used as described 460 previously<sup>86</sup>. Primers that were used in these experiments are listed in Supplementary Table 461 S2.

#### 462 Bacterial growth assay

463 *Pseudomonas syringae* pathovar *tomato* mutant *hrcC*- (deficient in type three effector 464 secretion system)<sup>87,88</sup>; was grown in 20 ml liquid YEB medium supplemented with 50  $\mu$ g/ml 465 Rifampicin on a shaker at 28°C overnight. Infection assay and counting the bacterial titer was 466 done as described previously<sup>89</sup> with a bacterial suspension at OD<sub>600</sub> = 0.0002. Leaves of 4-5-467 week-old Arabidopsis plants were infiltrated using a syringe. The *sid2-2* mutant plants, which 468 are incapable of accumulating salicylic acid<sup>90</sup>, were used as a positive control. Mock-infected 469 plants were similarly treated with infiltration buffer.

#### 470 Measurement of ethylene production

Leaf material of Arabidopsis plants was cut into discs of 10 mm<sup>2</sup> using a sterile cork borer, at 471 472 the end of the light period. After mixing leaf strips from several plants, six leaf strips were 473 placed together in a 6 ml glass vial containing 0.5 ml of ddH<sub>2</sub>O. Vials with leaf strips were 474 incubated overnight in the dark in a short-day room (16 h dark / 8 h light). The following day 475 (approximately after 16 h), elicitor peptide was added to the desired final concentration (1 476 µM) and vials were closed with air-tight rubber septa and put in the short-day room. ET 477 accumulating in the free air space was measured by gas chromatography (GC-14A Shimadzu) 478 after 4 h of incubation with or without elicitor.

#### 479 **ROS measurement**

480 Using a sterile cork borer, leaf discs of approximately  $10 \text{ mm}^2$  were cut from several plants. 481 One leaf disc per well was left floating overnight in darkness in 96-well plates (LIA White, 482 Greiner Bio-One) on 100 µl ddH<sub>2</sub>O at 18°C. Horseradish peroxidase (1 µg/ml final

483 concentration), luminol (100  $\mu$ M final concentration) and elicitor peptide (1  $\mu$ M final 484 concentration) were added to the wells. Using a plate reader (MicroLumat LB96P, Berthold 485 Technologies) light emission of oxidized luminol in the presence of peroxidase was 486 determined over 30 min, starting from addition of the elicitor.

#### 487 Immunoblot analysis

488 150 mg of leaf material from 4-5-week-old Arabidopsis plants was shock-frozen and ground 489 in liquid nitrogen. 200  $\mu$ l Läemmli buffer containing 50 mM  $\beta$ -mercaptoethanol was added 490 and the ground homogenate was further mixed by vortexing. Proteins were denatured by 491 boiling for 10 min at 95 °C. Debris was pelleted by centrifugation for 5 min at 13,000 rpm. 492 Total proteins were separated by electrophoresis in 7% SDS-polyacrylamide gels and 493 electrophoretically transferred to a polyvinylidene fluoride membrane according to the 494 manufacturer's protocol (Bio-Rad). Transferred proteins were detected with Ponceau-S. The 495 abundance of FLS2 receptor was analyzed by immunoblot and immunodetection with anti-FLS2-specific antibodies as previously described<sup>91</sup>. 496

#### 497 Phylogenetic analysis and comparison consensus of the amino acid sequences

Protein sequences BB2-B13 and ACLP1 were used as queries using BLASTP<sup>92</sup> (Basic Local 498 499 Alignment Search Tool) search to identify the most similar proteins in A. thaliana and diverse 500 land plants. We applied a cutoff of 70% sequence identity on the top hit of the BLASTP 501 search for BB2-B13 and ACLP1 and their orthologous and prologues were identified. Protein 502 sequences with more than 70% sequence identity were downloaded from the NCBI database and multiple alignment analysis performed based on the ClustalW software<sup>93</sup>. Phylogenetic 503 504 analyses and graphical representation were carried out using MEGA software (Molecular Evolutionary Genetics Analysis) version  $6.0^{94}$ . A neighbor-joining phylogenetic tree was 505 506 constructed after 1,000 iterations of bootstrapping of the aligned sequences. All branches with 507 bootstrap values <60% were collapsed. To compare consensus of the amino acid sequences, 508 sequence logos were generated using WebLogo (http:// www.weblogo.berkeleky.edu/), using 509 the ClustalW alignment as input.

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#### 767 Author Contributions

T.B., M.S., and C.B. conceived and designed the work. T.B. conceptualized the research
work. M.S. and C.B. performed sequence analysis and data analysis. M.S. performed most of
the data mining and experiments. M.S. wrote the original draft of the manuscript. T.B. and
C.B. revised the manuscript. All authors read and approved the final manuscript.

#### 772 Disclosure of Potential Conflicts of Interest

773 No potential conflicts of interest were disclosed.

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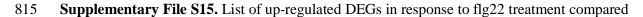
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783								
784	Supplementary Info	ormation						
785	Supplementary File	<b>S1.</b> List of all genes in response to flg22 treatment compared to the BSA						
786	treatment							
787	Supplementary File	e S2. List of all genes in response to AtPep1 treatment compared to the						
788	BSA treatment							
789	Supplementary File	e S3. List of top up-regulated DEGs in response to flg22 treatment						
790	compared to the BSA	treatment						
791	Supplementary File	e S4. List of top up-regulated DEGs in response to AtPep1 treatment						
792	compared to the BSA	treatment						
793	Supplementary File	e S5. List of top down-regulated DEGs in response to flg22 treatment						
794	compared to the BSA	treatment						
795	Supplementary File	e S6. List of top down-regulated DEGs in response to AtPep1 treatment						
796	compared to the BSA	treatment						
797	Supplementary File S7. List of the genes which their expression exclusively identified in the							
798	RNA-seq transcriptional profiling analysis in response to flg22 treatment and are not present							
799	in Affymetrix ATH1	GeneChip						
800	Supplementary File	<b>S8.</b> List of the up-regulated DEGs with fold change cutoff in response to						
801	flg22 treatment comp	bared to the BSA treatment that are present in both RNA-seq experiment						
802	analysis and ATH1 a	ffymetirx genechip						
803	Supplementary File	<b>S9.</b> List of the up-regulated DEGs with fold change cutoff in response to						
804	flg22 treatment com	pared to the BSA treatment that are exclusively present in RNA-seq						
805	experiment analysis							
806	Supplementary File	S10. List of 85 selected genes for subsequent study						
807	Supplementary File	e S11. List of DEGs exclusively up-regulated in response to flg22						
808	treatment compared t	o the BSA treatment						
809	Supplementary File	e S12. List of DEGs exclusively up-regulated in response to AtPep1						
810	treatment compared t	to the BSA treatment						
811	Supplementary File	e S13. List of DEGs exclusively down-regulated in response to flg22						
812	treatment compared t	to the BSA treatment						
813	Supplementary File	e S14. List of DEGs exclusively down-regulated in response to AtPep1						

814 treatment compared to the BSA treatment



816 to the AtPep1 treatment

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Fold Change (Log2)

- 817 Supplementary File S16. List of down-regulated DEGs in response to flg22 treatment
- 818 compared to the AtPep1 treatment
- A) Numbers of DEGs 265 246 fig22 VS Control AtPep1 VS Control flg22 VS AtPep1 B) C) flg22 flg22 AtPep1 AtPep1 (13%) (70.2%) (16.7%) (15.9%) (22.9%) (61.2%) D) E) P-Value (-log10) 0010)

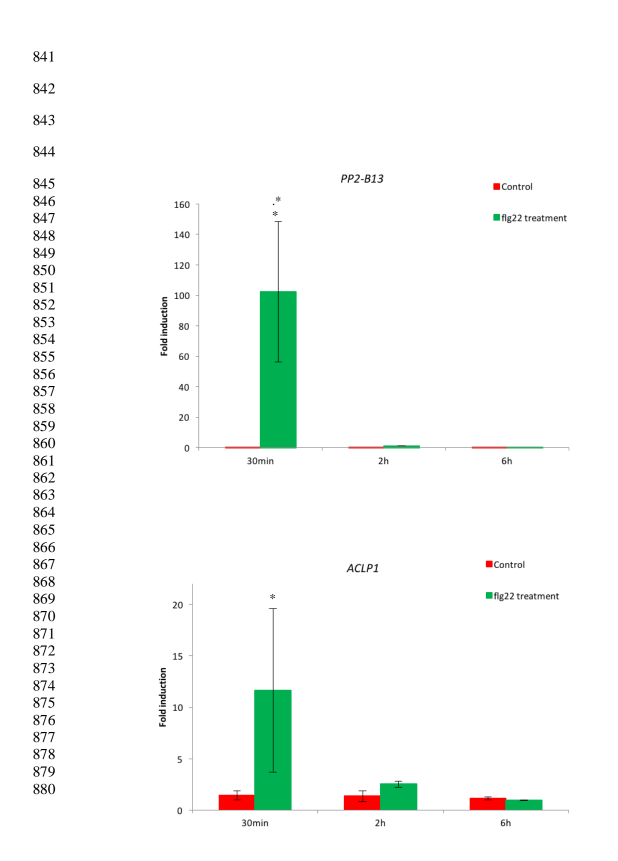
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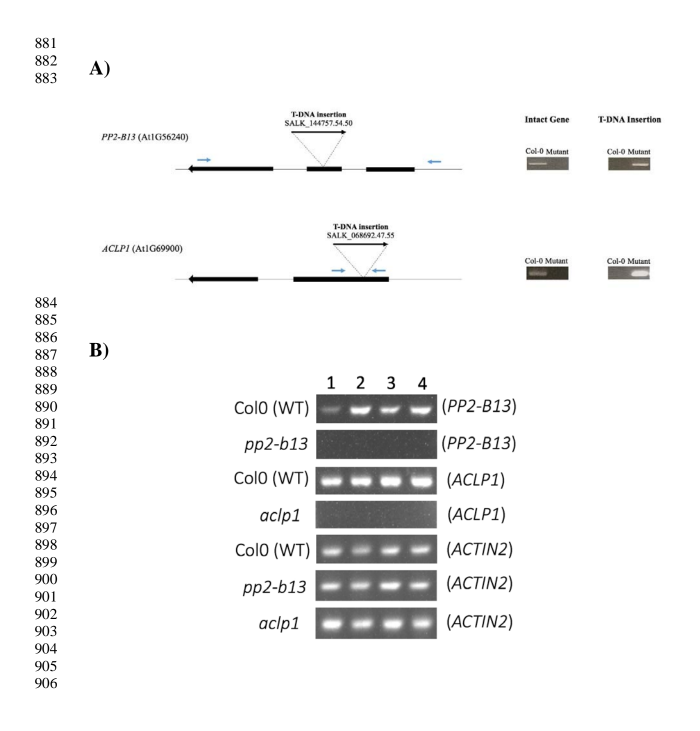
Fold Change (Log2)

**Table 1.** Detailed information on the top up-regulated genes based on RNA-seq analysis (fold change 30 minutes after flg22 and AtPep1 treatments compared to the control). Genes of interest are highlighted in bold.

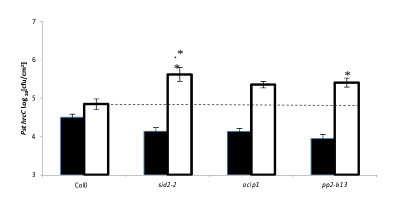
in in response to flg22 and AtPep1 treatments compared to the control investigated in this study. (**B**) Venn diagram of up-regulated DEGs between flg22 treatment and AtPep1 treatments. (**C**) Venn diagram of down-regulated DEGs between flg22 treatment and AtPep1 treatments. (**D**) Volcano plot of DEGs in response to flg22 treatment; (**E**) Volcano plot of DEGs in response to AtPep1 treatment. In (**D**-**E**), blue dots correspond to significantly up- and down-regulated DEGs, grey dots represent non-DEGs. *At1G56240 (PP2-B13)* and *At1G69900 (ACLP1)* are highlighted in red.

Accession Number	flg22 treatment		AtPep1 treatment		Putative function of the	Gene location	Consideration	T-DNA insertion	Final Genotyping
	Fold change	$\log_2$	Fold change	$\log_2$	— gene		for subsequent study	mutant/NASC Code	results confirmed by PCR
AT5G11140	503	8.97518	141	7.14031	Encodes an Arabidopsis phospholipase-like protein (PEARLI 4) family	Chr5:3545211-3546169	Not considered	Not available	
AT1G56240	126	6.97733	120	6.91095	Encodes a phloem protein 2- B13 (''PP2-B13''); function in: carbohydrate binding; F-box domain, cyclin-like, F-box domain, Skp2-like	Chr1:21056099- 21057577	"Consider"	detected/ SALK_144757.54.5 0	Homozygous line
AT2G32200	95	6.57146	36	5.16218	Encodes an unknown protein	Chr2:13676389- 13677306	Not considered		
AT1G05675	72	6.17964	63	5.97188	Encodes an UDP- Glycosyltransferase superfamily protein	Chr1:1701116-1702749	Not considered		
AT1G65385	65	6.01185	39	5.2804	Encodes an pseudogene, putative serpin	Chr1:24289566- 24291055	"Consider"	detected/ N570388, SALK_070388	Homozygous line
AT4G18195	60	5.9082	46	5.50896	Encodes the protein which is the member of a family of proteins related to PUP1, a purine transporter	Chr4:10069458- 10071115	Not considered		
AT5G36925	53	5.72585	55	5.78087	Encodes a protein with unknown protein	Chr5:14565476- 14566439	Not considered		
AT1G61470	33	5.03187	21	4.38478	Encodes a polynucleotidyl transferase protein which is, ribonuclease H-like superfamily protein	Chr1:22678092- 22679302	Not considered		
AT4G23215	30	4.9223	28	4.83246	Encodes a pseudogene of cysteine-rich receptor-like protein kinase family protein pseudogene	Chr4:12152900- 12153459	"Consider"	detected/ N605169, SALK_105169	Homozygous line
AT5G09876	29	4.83977	11	3.4823	Encodes an unknown protein	Chr5:3079887-3080435 Chr1:22032313- 22033297	Not considered		
AT1G59865	28	4.79972	39	5.27729	Encodes an unknown protein	Chr1:22032313- 22033297	"Consider"	detected/ N584779, SALK _084779	Not detected
AT2G35658	28	4.79818	22	4.46888	Encodes an unknown protein	Chr2:14990325- 14990935	"Consider"	detected/ N825604, SAIL_600_D01	Not detected
AT1G24145	26	4.71633	11	3.52029	Encodes an unknown protein, located in: endomembrane system	Chr1:8540838-8542053	"Consider"	detected/ N835081, SAIL_784_C07	Homozygous line
AT3G07195	24	4.58759	19	4.28252	Encodes a RPM1-interacting protein 4 (RIN4) family protein	Chr3:2288732-2290515	Not considered		
AT1G18300	22	4.48140	9	3.15319	Encodes a nudix hydrolase homolog 4 (NUDT4) protein	Chr1:6299669-6301139	Not considered		
AT2G24165	22	4.47869	16	4.03998	Encodes a pseudogene, similar to HcrVf3 protein	Chr2:10272672- 10273595	Not considered		
AT1G69900	20	4.34832	10	3.3191	Encodes an actin cross- linking protein; CONTAINS InterPro DOMAIN/s ("ACLP1")	Chr1:26326216- 26327965	"Consider"	detected/ N568692, SALK_068692 (AR)	Homozygous line
AT2G27389	20	4.288968	13	3.715253	Encodes an unknown protein	Chr2:11720294- 11721081	"Consider"	detected/ SALK_142825.23.9 5	Homozygous line
AT4G39580	18	4.16251	21	4.40726	Encodes a Galactose oxidase/kelch repeat superfamily protein	Chr4:18385684- 18386811	Not considered		
AT1G30755	14	3.83941	13	3.74068	Encodes an unknown protein	Chr1:10905731- 10909760	"Consider"	Not detected/ N666232, SALK_063010C	Not detected





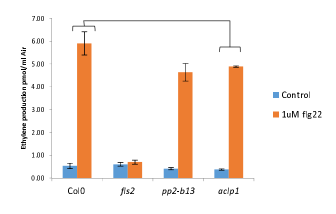
**Figure 3.** A) Schematic representation of homozygous T-DNA mutant lines *PP2-B13*, and *ACLP1*. Boxes indicate exons; thin lines indicate introns; bold arrows indicate T-DNA insertions; arrows indicate the direction of the gene. On the right side, the PCR results of the homozygous lines are shown, amplifying either the intact gene or the T-DNA. Small blue arrows indicate the primers position. **B**) RT-PCR results showing transcripts in Col-0 (WT), *pp2-b13* and *aclp1* mutant lines. The lower panel shows amplification of *ACTIN2* transcript as a control. Numbers 1 to 4 indicate individual plants for each genotype.



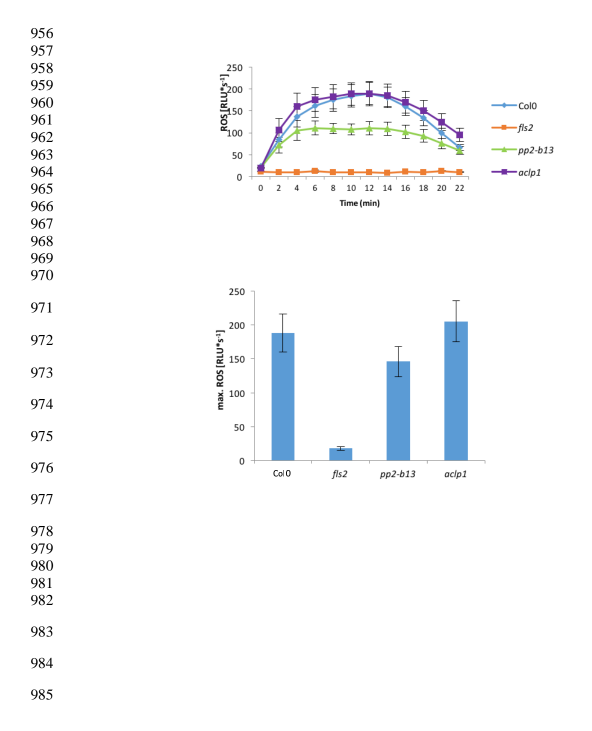
**Figure 4.** Bacterial susceptibility assay. Leaves of four- to six-week-old Arabidopsis plants (Col-0, *sid2-2*, *pp2-b13*, and *aclp1*) were pressure infiltrated with *Pseudomonas syringae* pv. *tomato* mutant *hrc*C- (OD<sub>600</sub>=0.0002, in infiltration buffer). *sid2-2* mutant plants, which are deficient in salicylic acid production, were used as a positive control. Black bars indicate bacterial colony from leaf discs of infected leaves just after infiltration (0 day); white bars represent colony-forming units

 $(cfu/cm^2)$  48 h post inoculation. Bars show the mean  $\pm$  s.e. (n=6). Similar results were observed in four independent experiments. Asterisks indicate a significant difference (\*p  $\leq 0.05$ , \*\*p $\leq 0.01$ ) from the wild type plants as determined by Student's *t-test*.

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**Figure 5.** Early PTI responses upon elicitor treatment. Ethylene accumulation after elicitor treatment. Leaf discs of four- to five-week-old plants of wild-type and mutant lines (pp2-b13, and aclp1) were treated with 1 µM of the flg22 elicitor peptide or without any peptide (control). In all cases, ethylene production was measured three and half hours after closing the tubes. Ethylene accumulation in pp2-b13 and aclp1 mutant lines was compared to the wild type Arabidopsis. *fls2* mutant line was used as a negative control. Columns represent mean ethylene concentration of six biological replicates. Error bars indicate standard deviation with n=6. Similar

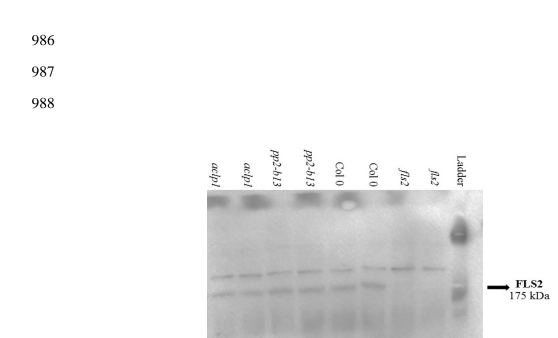


#### Figure 6.

(A) indicates ROS production in *pp2-b13* and *aclp1* mutant lines compared to wild-type Arabidopsis; (B) represents maximum ROS production in *pp2-b13* and *aclp1* mutant lines compared to wild-type Arabidopsis. *fls2* mutant line was used as a negative control.

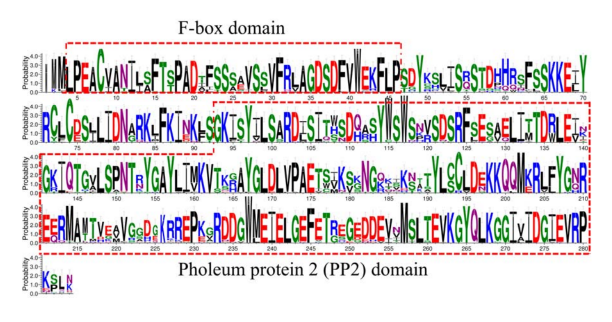
was repeated four times

with similar results. RLU= relative light units.



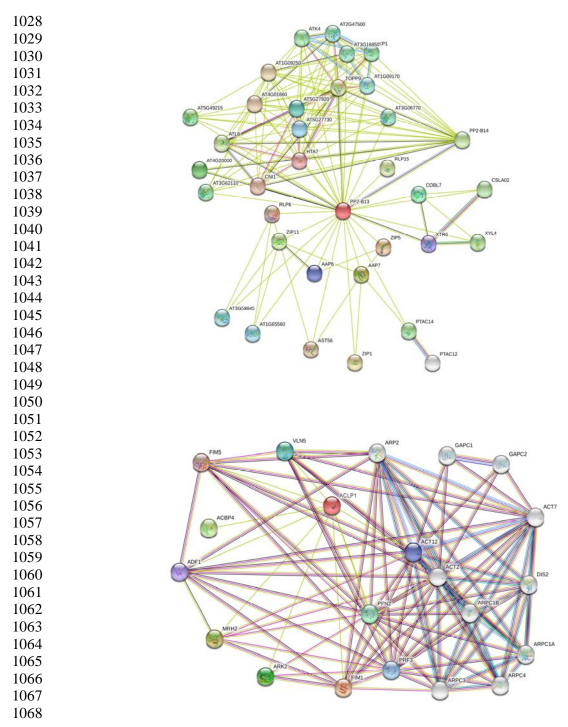
**Figure 7.** FLS2 protein levels. FLS2 protein levels of the mutant lines *pp2-b13* and *aclp1* as detected by immunoblot using a FLS2-specific antibody. *fls2* mutant plant is used as negative control. Ponceau S staining was used as loading control.

**Ponceau S** 



**Figure 8.** Different sequence conservation profiles in the PP2-B13 and its homologues in different plant species. Conservation plots were constructed using WEBLOGO. The y-axis represents the probability score. Y = 4 corresponds to 100% conservation. The predicted domains are highlighted in red boxes.

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**Figure 9.** The protein-protein interaction (PPI) network of the PP2-B13 and ACLP1 proteins in *Arabidopsis thaliana* based on STRING 11.0. analysis with a confidence threshold score of 0.4 (Szklarczyk et al., 2015).<sup>87</sup> Line colors indicate type of interaction used for the predicted associations: gene fusion (red), gene neighborhood (green), co-occurrence across genomes (blue), co-expression (black), experimental (purple), text mining (light green); association in curated databases (light blue). Line thickness represents the strength of data support. Proteins which have a known function in immune response are marked with dotted lines.

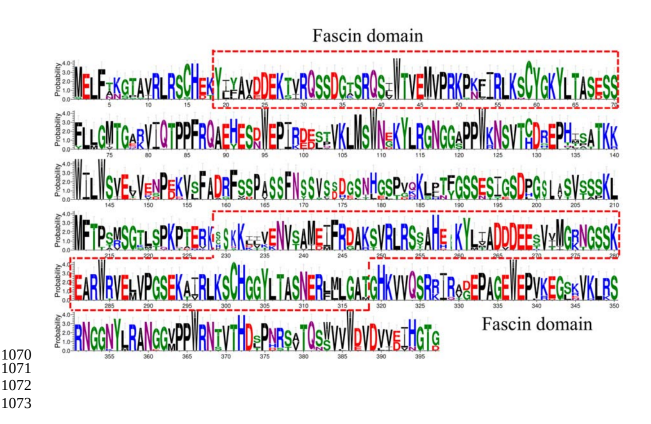


Figure 10. Different sequence conservation profiles in the ACLP1 and its homologues in different plant species. Conservation plots were constructed using WEBLOGO. The y-axis represents the probability score. Y = 4 corresponds to 100% conservation. The predicted domains are highlighted in red boxes.

### 1097

### 1098 Supplementary Information

#### 1099 Transcriptomic profiling uncovers novel players in innate immunity in Arabidopsis

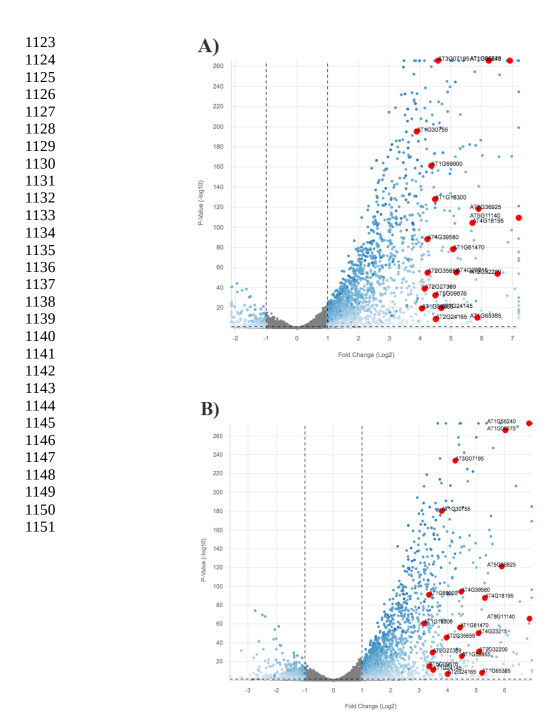
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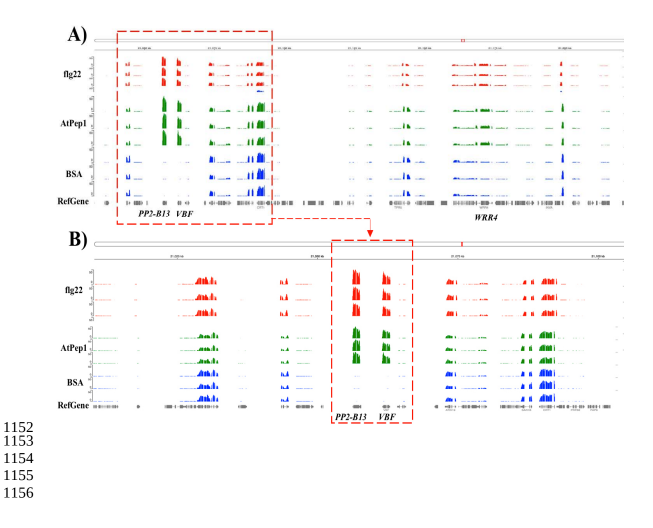
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**Supplementary Table S1.** Summary of Illumina sequencing data and mapped reads of *Arabidopsis thaliana* wild-type (Col-0) under BSA, flg22 and AtPep1 treatments.

	treatment								
		BSA			flg22			AtPep1	
	Repeat 1	Repeat 2	Repeat 3	Repeat 1	Repeat 2	Repeat 3	Repeat 1	Repeat 2	Repeat 3
Total clean read	1239300	12961695	14519871	17431550	9827458	14682108	12463709	10905959	12881404
Mapped reads of input	11934302	12646663	14157965	16833273	9581401	14350429	12162285	10620059	12538057
Read mapping rate of input (percent)	96.3%	97.6%	97.5%	96.6%	97.5%	97.7%	97.6%	97.4%	97.3%
Unmapped reads	210314	227871	273224	310845	170924	25657	22303	189382	222259
Unmapped reads (percent)	1.8%	1.8%	1.9%	1.8%	1.8%	1.8%	1.8%	1.8%	1.8%
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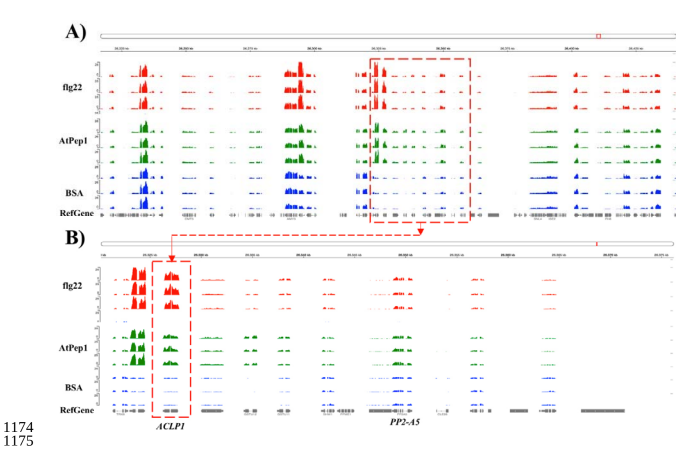


Volcano plot of gene expression in the seedling of Arabidopsis in response to flg22 treatment ( $\mathbf{A}$ ) and AtPep1 treatment ( $\mathbf{B}$ ). Blue dots correspond to significantly up- and down-regulated DEGs, while non-DEGs are in grey color. R



. Coverage depth graphs represent transcript abundance. (A) Overlaid depth graphs. (B) Zoomed in view of A. In the graph *PP2-B13*, *VBF* and *WRR4* genes are illustrated.





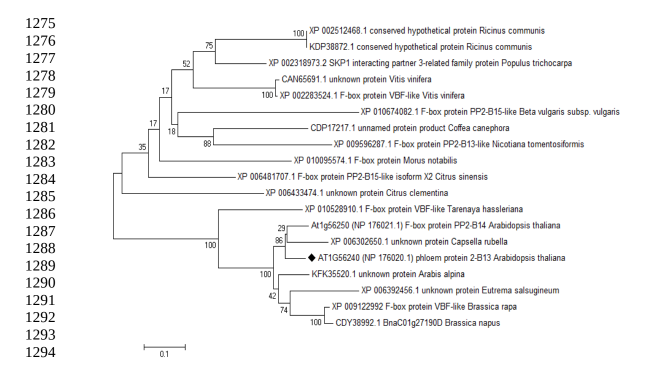
. (A) Overlaid depth graphs. (B) Zoomed in view of A. In the graph ACLP1 and PP2-A5 genes are illustrated.



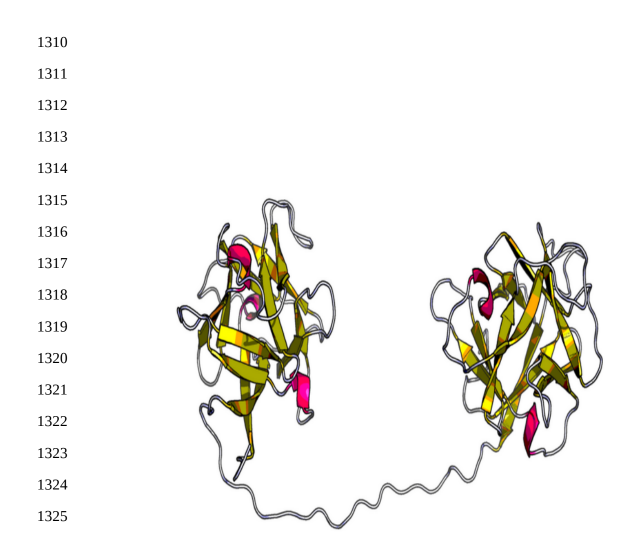
**Supplementary Figure S4.** Phenotype of five-week old Arabidopsis plants. Plant were grown under short-day conditions (ten hours light at 21°C and 14 hours dark at 18°C, with 60% humidity).

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1258	Supplementary Figure S5. Structure of PP2-B13 protein determined by Raptor X (Källberg et
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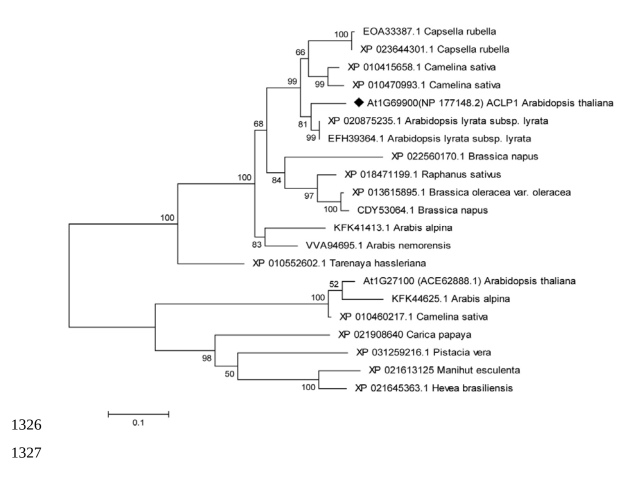
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**Supplementary Figure S7.** Phylogenetic analysis from the sequences of PP2-B13 protein in *Arabidopsis thaliana* and 15 representative land plants. The species indicated are *Ricinus communis, Populus trichocarpa, Vitis vinifera, Beta vulgaris, Coffea canephora, Nicotiana tomentosiformis, Morus notabilis, Citrus sinensis, Citrus clementine, Tarenaya hassleriana, Arabidopsis thaliana, Capsella rubella, Arabis alpina, Eutrema salsugineum, Brassica rapa* and *Brassica napus.* PP2-B13 protein in *Arabidopsis thaliana* was labelled. Sequences for comparisons were obtained from GenBank. The accession numbers and protein names (if available) are given. Analysis was done by maximum likelihood method implemented in MEGA6 (Molecular Evolutionary Genetics Analysis) version 6.0.



**Supplementary Figure S6.** Structure of ACLP1 protein determined by Raptor X (Källberg *et al.*, 2012).



**Supplementary Figure S8.** Phylogenetic analysis from the sequences of ACLP1 protein in *Arabidopsis thaliana* and 14 representative land plants. The species indicated are *Capsella rubella*, *Camelina sativa*, *Arabidopsis thaliana*, *Arabidopsis lyrata*, *Brassica napus*, *Raphanus sativus*, *Brassica oleracea*, *Arabis alpina*, *Arabis nemorensis*, *Tarenaya hassleriana*, *Carica papaya*, *Pistacia vera*, *Manihut esculenta* and *Hevea brasiliensis*. ACLP1 protein in *Arabidopsis thaliana* was labelled. Sequences for comparisons were obtained from GenBank. The accession numbers and protein names (if available) are given. Analysis was done by maximum likelihood method implemented in MEGA6 (Molecular Evolutionary Genetics Analysis) version 6.0.

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### **Supplementary Table S2.** List of the oligonucleotide primers which were used in this study. 1335

Name of primer	Purpose	DNA sequence (5'-to-3') 1336
SALK_144757.54.50_LP1	Genotyping	тсаатстссаассасссдтс 1337
SALK_144757.54.50_RP1	Genotyping	GGTCAGCAGAAATATGCCAATGATCACT
SALK_68692.47.55_LP1	Genotyping	1338 gagaccgacgagttaaaactag
SALK_68692.47.55_RP1	Genotyping	тааассааааттсатасдтстсаа 1339
SALK_LBa1	Genotyping	тдаттсасатадтададссатса 1340
SALK_RB	Genotyping	TCATGCGAAACGATCCAG 1341
SALK_LB2	Genotyping	gcgtggaccgcttgctgcaact 1342
SAIL-Pdap101_LB1	Genotyping	GCCTTTTCAGAAATGGATAAATAGCCTTG <b>C3143</b>
PP2-B13_RT_fw	RT-PCR	CCAGCCGATGCATTTTCGTCATC1344
PP2-B13_RT_rw	RT-PCR	тсттстсддттсссдтааатадс1345
ACLP1_RT_fw	RT-PCR	тсддатсстдддтсасттдтатса $1346$
ACLP1_RT_rw	RT-PCR	agcactccggtttggtaaatcatg <b>l</b> 347
ACTIN2_RT_fw	RT-PCR	agtgtctggatcggtggttc 1348
ACTIN2_RT_rw	RT-PCR	CCCCAGCTTTTTAAGCCTTT
<i>PP2-B13</i> (AT1G56240)_qRT_fw	gene expression	CGTGACACAGACTAAATAATAGATC
<i>PP2-B13</i> (AT1G56240)_qRT_rw	gene expression	CCTCTGAAATAGGGATCAAGATG
ACLP1 (AT1G69900)_qRT_fw	gene expression	GGAATATTTCCATCGCCGATAC
ACLP1 (AT1G69900)_qRT_rw	gene expression	GATCCTGGGTCACTTGTATCAG
<i>UBQ10</i> (AT4G05320)_qRT_fw	gene expression	GGCCTTGTATAATCCCTGATGAATAAG
<i>UBQ10</i> (AT4G05320)_qRT_rw	gene expression	AAAGAGATAACAGGAACGGAAACATAG