

BABA-induced pathogen resistance: a multi-omics analysis of the tomato response reveals a hyper-receptive status involving ethylene

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1 Title: BABA-induced pathogen resistance: a multi-omics analysis of the

2 tomato response reveals a hyper-receptive status involving ethylene

3 Running title: The multi-omics analysis of BABA-treated tomato

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29 **1. ABSTRACT**

30 Prior exposure to microbial-associated molecular patterns or specific chemical compounds can 31 promote plants into a primed state with stronger defence responses. β -aminobutyric acid (BABA) is 32 an endogenous stress metabolite that induces resistance protecting various plants towards diverse 33 stresses. In this study, by integrating BABA-induced changes in selected metabolites with transcriptome and proteome data, we generated a global map of the molecular processes operating 34 in BABA-induced resistance (BABA-IR) in tomato. BABA significantly restricts the growth of the 35 pathogens Oidium neolycopersici and Phytophthora parasitica but not Botrytis cinerea. A cluster 36 37 analysis of the upregulated processes showed that BABA acts mainly as a stress factor in tomato. The 38 main factor distinguishing BABA-IR from other stress conditions was the extensive induction of signaling and perception machinery playing a key role in effective resistance against pathogens. 39 40 Interestingly, the signalling processes and immune response activated during BABA-IR in tomato differed from those in Arabidopsis with substantial enrichment of genes associated with jasmonic 41 acid (JA) and ethylene (ET) signalling and no change in Asp levels. Our results revealed key 42 differences between the effect of BABA on tomato and other model plants studied until now. 43 44 Surprisingly, salicylic acid (SA) is not involved in BABA downstream signalization whereas ET and JA 45 play a crucial role.

46 2. INTRODUCTION

Throughout their lives, plants are constantly exposed to many stressful situations caused by 47 changing environmental conditions or attacks by various pests and pathogenic microorganisms. 48 Therefore, they have evolved structural barriers, microbicidal secondary metabolites, and inducible 49 defence mechanisms to repel potential attackers. Unfortunately, the basal immune responses of 50 51 plants are usually only sufficient to slow their colonisation by pathogens. As a result, a significant portion of the world's plant production is destroyed each year by fungi, oomycetes, bacteria, insects, 52 53 and nematodes. [1]. Although plants do not have the adaptive immune system of vertebrates, it has long been known that components of the innate immune system of plants can learn from the past 54 55 [2]. When exposed to microbe-associated molecular patterns (MAMPs) or specific chemical compounds, plants can enter a state of enhanced defence characterised by more rapid and robust 56 57 responses to stressful stimuli. Although the term "defence priming" was proposed in the 1930s, the molecular mechanisms underlying this phenomenon have only recently been partially elucidated, 58 particularly in the model plant Arabidopsis thaliana [3,4]. Defence priming causes increased 59 60 expression of genes related to stress and defence [5], including many transcription factors that regulate defence [6], and is now considered an essential component of several types of systemic 61

62 plant immunity, including acquired systemic acquired resistance (SAR), induced systemic resistance 63 (ISR) [3,7], wound-induced resistance [8], and resistance induced by chemical compounds. Unlike 64 strategies based on single resistance genes, defence priming activates multigenic defence 65 mechanisms, conferring relatively durable resistance [3]. One of the most effective priming agents is 66 the non-protein amino acid β -aminobutyric acid (BABA), which protects various plant species against 67 a wide range of stresses [9]. Importantly, this resistance is long-lasting and can be transferred to 68 vegetative progeny [10,11]. BABA was previously considered a xenobiotic, but it has recently been shown to accumulate in stress-exposed plants [12], suggesting that it is an endogenous stress 69 70 metabolite [13]. BABA induces resistance via the action of several hormones, including salicylic acid 71 (SA) [14,15], jasmonic acid (JA) [16], abscisic acid (ABA) [17], and ethylene (ET) [15]; the signalling 72 pathway that is activated appears to depend strongly on the particular plant-pathogen combination 73 [6]. Recently, it was discovered that the aspartyl-tRNA synthetase (AspRS) IBI1 in Arabidopsis 74 thaliana serves as an enantiomer-specific BABA receptor that interacts with the transcription factors 75 VOZ1 and VOZ2 [17,18]. In BABA -primed cells, this interaction represses the expression of ABA 76 genes, resulting in increased expression of PTI genes and callose-associated defence [17]. The 77 previous study of our laboratory showed that, as in potato, effective BABA-IR is also associated with 78 the formation of HR -like lesions in tomato [19]. However, while BABA-IR appears to activate SA signalling pathways in Arabidopsis and potato plants, our results suggest that ET signalling pathways 79 play a key role in BABA-IR in tomato plants [19]. 80

Here, we present a study in which a combination of nontargeted approaches was used to elucidate the molecular basis of BABA-IR in tomato (*Solanum lycopersicum* cv. *Marmande*), an important crop [20]. BABA was applied by spraying onto leaves, as this application strategy is easy to implement in practical agriculture. Global transcriptomic and proteomic analysis of tomato plants allowed us to identify the molecular processes and signalling pathways that occur in tomato at BABA-IR.

87 3. RESULTS

88

3.1 Growth of pathogens having different lifestyles after BABA treatment

To determine the protective effect of BABA treatment towards pathogens with different lifestyles (biotrophic *Oidium neolycopersici*, hemibiotrophic *Phytophthora parasitica*, and necrotrophic *Botrytis cinerea*) in *S. lycopersicum* cv. *Marmande* plants, we treated plants with 10 mM BABA. This dose was selected based on a previous study [21] showing maximal (95%) protection against *Phytophthora infestans* with no effect on the general growth of the *S. lycopersicum* cv. F1 hybrid cv Baby plants. Indeed, BABA was unable to trigger effective resistance, in our experimental

95 conditions, below concentration of 5mM, as shown previously [21]. More than two days after treatment we observed not reproducibly the appearance of HR-like microlesions on some BABA 96 97 spayed leaves (S1 Fig) not connected to plant age or leaf position, as described previously [21]. BABA treatment significantly reduced the sporulation of O. neolycopersici and the spreading of P. parasitica 98 99 but had no effect on B. cinerea disease (Fig 1A). These findings are in agreement with previous 100 studies showing that BABA-IR to biotrophic and hemibiotrophic pathogens in tomato and potato 101 plants but gave inconsistent results against the necrotrophic pathogen B. cinerea, possibly due to the 102 use of different methods to evaluate resistance [22-24].



103

104 Figure 1. Resistance, Transcriptomic, and Proteomic changes associated with BABA-IR, overlap 105 between BABA-induced genes in different plant species and overlap between genes induced by 106 BABA and selected elicitors. (A) Lesion surface area in 6-8 weeks-old tomato plants after spraying 107 with 10 mM followed by challenge with the phytopathogens Botrytis cinerea, Phythophtora 108 parasitica, and O. neolycopersici. (B) Total numbers of differentially expressed genes and proteins 109 during BABA-IR (calculated as the ratio of the expression in 10 mM BABA-sprayed and water-sprayed 110 leaves) 24 h after treatment. Cut-offs of a \geq 2-fold difference in expression and P \leq 0.01 (for genes) or 111 $P \le 0.05$ (for proteins) were applied. The number of genes or proteins in each category is shown. (C) 112 Degrees of overlap between orthologous groups identified in this study and previous transcriptomic

studies on BABA-treated potato (Bengtsson et al., 2014) and *Arabidopsis* plants (Zimmerli et al., 2008). (D) Overlap between genes induced in BABA-treated tomato Marmande plants and in tomato Rio Grande (RG)-PtoR resistant plants during MTI, ETI, or both (Pombo et al., 2014). (E) Overlap between genes induced in BABA-treated tomato Marmande plants and in tomato plants treated with the virulence factors coronatine (COR) from *Pst* DC3000 (Geng et al., 2014), flagellin from *P. syringae*

118 (Rosli et al., 2013), and elicitin INF1 secreted by *P. infestans* (Kawamura, 2009). Induced genes were

identified by applying cut-offs of a \geq 2-fold difference in expression and P <0.05.

120 3.2 Transcriptome and proteome modifications during BABA-IR in tomato

Three leaf biological replicates were sampled 24 h and 48 h after spraying with 10 mM BABA. 121 Six samples collected at 24 hrs were used for RNA sequencing and six samples collected at 48 hrs 122 were used for label-free LC-MS/ analysis of the proteome excluding microsomal fraction (thereafter 123 proteome). RNA sequencing generated 334,275,668, high-quality reads with an average of 124 55,712,611 reads per sample (S1 Table). The reads for each sample were mapped to the S. 125 lycopersicum reference genome sequence, with 74% of reads being mapped successfully. Only genes 126 with a median count above 20% in at least one treatment were considered in the subsequent 127 analysis. The cut-offs used in the comparison were P < 0.01, and ≥ 2 -fold expression change. Using 128 129 these criteria, we identified 24,562 genes from 34,725 annotated genes (ITAG 2.4), with 1,523 genes being differentially expressed (Fig 1B, S2 Table). In the proteome analysis, protein identification was 130 131 only performed for peptides of at least six amino acids with a statistically significant peptide score (q < 0.01, FDR 1 %; FDR based on decoy search against the reverse database). The cut-offs used for the 132 comparison were q < 0.05 and ≥ 2 -fold expression change. Using these criteria, we identified 1808 133 protein groups (S3 Table) and 319 differentially expressed proteins (Fig 1B, S2 Table). As with the 134 135 transcriptome, we found that far more proteins were upregulated (67%) than downregulated (33%) 136 after BABA treatment (Fig 1B, S2 Table), however the correlation between proteins and transcripts changes was only about 10% as discussed previously [25]. These results confirm that BABA treatment 137 causes extensive reprogramming of cellular processes in tomato plants, as previously observed in 138 139 Arabidopsis and potato [22,26]. However, a comparison of orthologous groups between our transcriptomic study and previous studies on Arabidopsis [26] and potato [22] plants exposed to 140 141 BABA revealed significantly greater overlap between the potato and tomato datasets than between Arabidopsis and tomato (Fig 1C, S4 Table). 142

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3.3

BABA exhibits common features with MAMPs-triggered immunity in tomato

Sets of genes whose transcript abundance was specifically increased at 6 hours post-infection (hpi) during MAMPs-triggered immunity (MTI) and effector-triggered immunity (ETI) were recently identified using RNA-Seq technology in tomato Rio Grande (RG)-PtoR resistant plants [27]. 147 Interestingly, almost 50% of the BABA-upregulated genes were also differentially regulated during 148 MTI, but there was no overlap between ETI-upregulated genes and BABA treatment (Fig 1D, S5 149 Table). Following this result, the response of tomato plants to BABA was far more similar to that induced by exposure to a well-characterized MAMP flagellin flgII-28 from the bacterium 150 Pseudomonas syringae [28] or elicitin INF1 secreted by Phytophthora infestans [29] than to the 151 response induced by the virulence factor coronatine (COR) secreted by Pst DC3000 [30] (Fig 1E, 55 152 153 Table). Specifically, almost 48% and 57% of BABA-upregulated genes were also upregulated following exposure to figli-28 analysed using RNA-Seq technology at 6 hpi [28] and INF1 analysed 154 155 using a GeneChip tomato genomic array 12 hpi [31], respectively. Almost 85 % of COR-upregulated 156 genes analysed using a TOM1 cDNA microarray 24 hpi of tomato var "Glamour" seedlings [32] were unaffected by BABA treatment (Fig 1E, S5 Table). The only genes upregulated by both BABA and COR 157 158 were associated with ET (ACC synthase and ACC oxidase) and JA signalling pathways.

159 3.4 BABA functions as a stress factor

A gene ontology (GO) term enrichment analysis was performed to identify critical processes upregulated by BABA treatment. The sets of terms obtained using the transcriptomic and proteomic data were similar (S2 Fig, S6 Table), with many common terms in the GO categories "Process" and "Function".

The protein-protein interaction network based on RNA-Seq data was generated by directly 164 165 mapping upregulated genes to proteins in the String database [33] (Fig 2A, S7 Table). The network is highly aggregated with clustered sub-networks comprising proteins associated with defence 166 responses (PR proteins), JA and ET signalling and synthesis, regulation of transcription related to 167 168 mitogen-activated protein kinase MPK3, and processes related to reactive oxygen species (ROS) production. This is also consistent with maps produced after the ReviGO [34] analysis showing 169 significant enrichment of stress-associated clusters (S2 Fig, S6 Table). The protein-protein interaction 170 171 network based on proteins exhibiting significant changes in abundance 48 hours after BABA treatment (Fig 2B, S7 Table) featured notable clusters relating to photosynthesis, secondary 172 173 metabolite biosynthesis, and translation. The photosynthesis cluster shows that BABA induces 174 complex changes in the regulation of photosynthesis-related energetic processes and carbohydrate 175 metabolism. The cluster related to secondary metabolites includes the enzymes prephenate 176 aminotransferase (PAT), arginase (ARG2), and lactate dehydrogenase. PAT plays a role in the 177 biosynthesis of aromatic amino acids, while lactate dehydrogenase plays an important role in 178 detoxifying D-lactate, a product of the glyoxalase pathway for detoxifying methylglyoxal, which 179 accumulates under stress conditions [35]. The arginase in tomato leaves was suggested to be 180 involved in ROS homeostasis when its expression was induced by JA signalling following wounding

and exposure to *Pseudomonas syringae* pv. *tomato* strain DC3000 [36]. The upregulation of the proteins in the cluster related to translation is probably related to translational switching from growth to defence [37] and supports the reported role of non-canonical functions of aminoacyl-tRNA synthetase in BABA responses [18].



Figure 2. Transcriptomic and proteomic comparisons of plants treated with BABA revealed the existence of sets of genes and proteins that are specifically induced during BABA-IR. Proteinprotein association networks were generated for significantly induced (A) genes and (B) proteins, applying cut-offs of a \geq 2-fold difference in expression and P \leq 0.01 (for genes) or P \leq 0.05 (for proteins). Interaction networks were constructed in STRING using a minimum required interactionscore of 0.7 [33] and visualized with Cytoscape [96].

To distinguish between stress-related genes upregulated following BABA treatment and 192 193 defence-related genes involved in BABA-IR, we compared the BABA-upregulated genes to the sets of 194 genes upregulated in previous transcriptomic studies on tomato plants subjected to temperature 195 [38] and salinity [39] stress as well as those upregulated in tomato plants infected by the fungus 196 Stemphylium lycopersici [40] and the FIRE (flagellin-induced repressed by effectors) genes, which 197 represent a pathogen-defined core set of immune-related genes [28]. We found that 50% of the BABA-upregulated genes overlapped only with the sets of genes upregulated by abiotic stress, 198 confirming the hypothesis that BABA acts on tomato plants primarily as a stress factor. However, 30 199 200 % of the BABA-upregulated genes were also FIRE genes or genes upregulated following S. lycopersici 201 infection, suggesting that these genes are associated with BABA-augmented defence expression against P. parasitica and O. neolycopersici. Moreover, about 20% of genes unique to the BABA 202 treatment were significantly enriched in KEGG pathways related to plant-pathogen interaction, 203 204 MAPK signalling, and phenylpropanoid synthesis, indicating that these genes are also involved in 205 BABA-IR (Fig 3B, S5 Table).



Intersection BABA/Abiotic Stress				
Category	1D	GO Term/Pathway	FDR	
Process	GO:0045449	regulation of transcription	2.9E-02	
	60:0010468	regulation of gene expression	2.9E-02	
	GO:0009889	regulation of biosynthetic process	2.98-02	
Function	GO:0030528	transcription regulator activity	1.4E-05	
	GO:0005509	calcium ion binding	1.48-03	
	GO:0016301	kinase activity	3.8E-03	
KEGG	SLY04626	Plant-pathogen interaction	9.0E-06	
	SLY04016	MAPK signaling pathway - plant	9.58-03	
	5LY04075	Plant hormone signal transduction	5.7E-03	
		Intersection BABA/FIRE		
Category	ID	GO Term/Pathway	FDR	
Process	GO:0006468	protein amino acid phosphorylation	3.0E-03	
	GO:0009607	response to biotic stimulus	3.0E-03	
Function	GO:0004674	protein serine/threonine kinase activity	7.8E-12	
	60:0019199	transmembrane receptor protein kinase activity	3.26-10	
	GO:0004872	receptor activity	2.5E-08	
KEGG	SLY04626	Plant-pathogen interaction	1.18-08	
	SLY00360	Phenylalanine metabolism	3.88-04	
	SLY00480	Glutathione metabolism	4.4E-04	
		BABA		
Category	ID	GO Term/Pathway	FDR	
Function	60:0016301	kingse activity	1.65-03	
KEGG	5LY04625	Plant-pathogen interaction	4.68-05	
	SLY04016	MAPK signaling pathway - plant	9.98-05	
	SLY00940	Phenylpropanoid biosynthesis	9.95-05	
	SLY00520	Amino sugar and nucleotide sugar metabolism	4.88-03	

Figure 3. Overlap between genes induced by BABA and genes induced in tomato plants subject to various abiotic and biotic stresses. (A) Overlap between genes induced in BABA-treated tomato Marmande plants and genes upregulated in tomato plants subject to abiotic heat stress [38], genes upregulated in abiotic salinity stress [39], genes upregulated following infection by the fungus

Stemphylium lycopersici (biotic stress; [40]), and FIRE (flagellin-induced repressed by effectors) genes, representing a pathogen-defined core set of immune-related genes [28]. Cut-offs of a \geq 2-fold difference in expression and q <0.05 were applied. (B) KEGG and Gene Ontology (GO) term analysis were performed for genes in each category. The most-enriched GO terms in the process and function categories are shown. (C) Protein-protein association network for genes significantly induced by both BABA treatment and abiotic stress. The interaction network was constructed in STRING using a minimum required interaction score of 0.7 [33] and visualized with Cytoscape [96].

225 3.5 The BABA stress response is orchestrated via ethylene and jasmonic acid signalling

The preceding analysis revealed a clear enrichment of genes associated with JA and ET 226 signalling pathways. Moreover, both ET and JA accumulated in tomato plants during the first few 227 hours after BABA treatment (Fig 4A, B). Accordingly, BABA induced upregulation of several isoforms 228 of ACC synthase (ACS) and ACC oxidases (ACOs) (Fig 4C, S8 Table), essential enzymes for ET 229 biosynthesis [41,42]. The high upregulation of the ACS2 and ACS6 isoforms is also consistent with 230 231 their reported activation during defence reactions in Arabidopsis [43] and with the observed upregulation of WRKY33, which activates ACS2 and ACS6 expression downstream of the MPK3/MPK6 232 cascade [44]. Our data also support the findings of an earlier study on Arabidopsis [45] showing that 233 the conversion of methylthioadenosine (MTA) to Met via the Yang (or Met salvage) cycle is generally 234 not controlled by ET signalling because BABA treatment had no detectable effect on the regulation of 235 the Yang cycle genes MTN, MTK, MTI, and ARD (Fig 4C, S8 Table). 236

237 Among the BABA-upregulated genes were the patatin-like proteins PLA1 and PLA3 (Fig 4D, S8 Table), which have been implicated in wound responses and resistance to necrotrophic pathogens 238 via JA signalling [46]. In addition, there was significant upregulation of genes encoding enzymes 239 involved in the synthesis of endogenous JA (TomloxD, AOS2 and OPR3). The upregulation of these 240 genes was accompanied by the accumulation of JA in BABA-treated leaves (Fig 4D). Also upregulated 241 242 were 6 of the 12 Jasmonate ZIM Domain (JAZ) genes, key regulators of JA signalling that govern host 243 and non-host pathogen-induced cell death in tomato. The most highly upregulated JAZ genes (SIJAZ1, SIJAZ2 and SIJAZ9-11) were also induced by treatment with COR [47]. 244

Surprisingly, unlike in previous studies on the effects of BABA in *Arabidopsis* [14] and potato plants [22], there was no significant enrichment of genes associated with the phytohormone SA. In keeping with this result, the levels of total SA declined slightly in leaves treated with BABA (Fig 4B). A recent study showed that ABA signalling is suppressed during BABA-IR in *Arabidopsis* plants [17]. In accordance with this result and our previous study [48], we observed no enrichment of genes associated with ABA signalling following BABA treatment in tomato and several genes involved in ABA catabolism were upregulated including *CYP707A* (*Solyc08g005610.2.1*) and UDPglucosyltransferase (*UGT*; *Solyc09q098080.2.1*) (Fig 4E, S8 Table).



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Figure 4. Involvement of signalling pathways in BABA-IR. (A) ET accumulation was measured at 24 254 255 hrs after BABA treatment (10 mM) of leaflets (n = 8) by gas chromatography. (B) Levels of JA, jasmonic acid-isoleucine (JA-IIe), methyl-jasmonate (MeJA) and SA were measured by LC-MS 24 h 256 after BABA treatment (10 mM) leaves (n = 6). The control tissue (CTRL) was a water-treated control 257 258 sample. Each bar represents the mean \pm SE. Asterisks denote mean values that differ significantly 259 from that for the control group based on Student's t-test at $P \leq 0.01$ (**). (C-E) The BABA-induced 260 induction of genes involved in the synthesis of ET (C) and JA (D) and the degradation of ABA (E) is 261 shown.

SAM – S-Adenosyl-L-methionine, MTA – 5'-Methylthioadenosin, MTR – 5'-Methylthioribose, MTR-P – 5' Methylthioribose-1-phosphate, MTRu-P – 5'-Methylthioribulose-1-phosphate, DHKMP – 1,2-Dihydroxy-3-keto 5-methylthiopentene, KMTB – 2-keto-4-methyl-thiobutyrate, SAM2 – S-Adenosyl-L-methionine synthase, ACS
 1-aminocyclopropane-1-carboxylic acid synthase, ACC – 1-aminocyclopropane-1-carboxylic acid, ACO – 1 aminocyclopropane-1-carboxylic acid oxidase, PLA – Phospholipase A, Lox – Lipoxygenase, (13S)-HPLA – 13-

hydroperoxylinolenic acid, AOS – Allene oxide synthase, 12,13-EOLA – 12,13-epoxyoctadecatrienoic acid, AOC –
 Allene oxide cyclase, OPDA – (9S,13S)-12-oxo-phytodienoic acid, OPR – 12-oxo-phytodienoic acid 10,11
 reductase, OPC – 3-oxo-2-(2´-pentenyl)-cyclopentane-1-octanoic acid, ACX1A – acyl-CoA oxidase, JA – Jasmonic
 acid, ABA – Abscisic acid, UGT – Abscisic acid glucosyl-transferase, ABA-GE – Abscisic acid glucosyl ester, 8´-OH
 ABA – 8´-hydroxy abscisic acid N.C. – transcript not changed, N.D. – transcript not detected.

272 3.6 BABA upregulates perception and signalling machinery related to abiotic stress

Protein kinases comprised 142 of the BABA-upregulated genes (S3 Fig, S9 Table), accounting 273 274 for 15% of the total set of expressed kinases in our analysis. Similarly, 44 protein kinases were 275 identified in the proteomic analysis, of which 14 were upregulated and only 3 were downregulated (S3 Fig). These results suggest that protein kinases play an important role in the BABA stress 276 response (S3 Fig, S9 Table). An enrichment analysis of the BABA-responsive families using the chi-277 278 squared test revealed overrepresentation of the large Receptor-Like Kinase/Pelle (RLK-Pelle) family, 279 which is crucial for plant-specific adaptation [49] (S3 Fig) at both the transcriptomic and proteomic levels. This finding agrees well with the upregulation of 27 Receptor Like Proteins (RLP), representing 280 23% of the total expressed RLPs in our analysis. It is also notable that all previously reported PRRs in 281 tomato plants were upregulated including those for flagellin (FLS2 and FLS3), the fungal elicitor EIX 282 (LeEix1 and LeEix2), Ave1 from V. dahlia (Ve1 and Ve2) [50], and Avr factors (Cf-2, Cf-4, Cf-5 and Cf-9) 283 [51]. This increased expression of perceptual proteins is consistent with the observed protein-protein 284 interaction between WRKY33 and the MPK3 kinase from the CMGC family (Fig 2B, S3 Fig). The 285 286 MPK3/MPK6 cascade causes the phosphorylation of WRKY33 and the closely related WRKY25, leading to ET production due to activation of the enzyme ACS as described above. Two other RLKs 287 interacting with RLPs involved in MAMPs perception, SOBIR1 [52] and TARK1 [53], were also 288 289 upregulated.

In parallel with the induction of pathogen perception machinery, we also observed massive 290 upregulation of genes encoding transcription factors (TFs): BABA treatment induced the upregulation 291 292 of 130 TF genes (S4 Fig). The largest numbers of induced genes were found in the ERF, WRKY, MYB 293 and NAC families (S4 Fig), which play essential roles in regulating stress responses in plants, mainly through ET and JA signalling pathways. The strong induction of ERFs following BABA treatment was 294 295 particularly notable: of the 137 identified ERF genes in tomato [54], 113 were expressed, 33 were upregulated, and 5 were downregulated. The BABA response has many features in common with the 296 297 responses to cold, salt, and mechanical stress observed in previous studies on tomato [38,55,56]. 298 However, the BABA-upregulated genes SIERF5 (ERF5), SIERF43 (RAV2), SIERF55 (TSRF1), SIERF60 299 (Pti5) or SIERF69 (ERF1) were previously linked to the activation of defence responses against diverse 300 pathogens (S9 Table). Unlike in previous studies on tomato, no WRKY genes were downregulated 301 after BABA treatment; their expression patterns resembled those induced by salt stress, the tomato

302 spotted wilt virus (TSWV), and the fungal elicitor *EIX* [57]. The largest class of plant MYB factors are 303 the 2R proteins, which regulate primary and secondary metabolism, hormone signal transduction, 304 development, and responses to biotic and abiotic stresses [58]. Of the 91 expressed 2RMYB genes, 305 11 were upregulated following BABA treatment and 6 were downregulated. Their expression profile 306 most closely resembled that seen in tomato plants treated with MeJA [59] or stressed by infection 307 with the bacterial pathogens *Pst* DC3000 or *P. putida* (S9 Table).

Finally, a comparison of the BABA-upregulated RLPs/WRKY, ERF, and MYB genes to recent RNA-Seq results for tomato plants under biotic [27] and abiotic stress [38] confirmed that the pattern of upregulation induced by BABA is significantly more similar to that for abiotic stress than that for biotic stress (S9 Table).

312 3.7 Amino acid metabolism in BABA-treated plants

Transcriptomic and metabolomic studies have consistently shown that amino acid (AA) 313 314 homeostasis plays a role in stress responses [60,61]. Moreover, BABA induces the stress-induced 315 morphogenic response (SIMR). In Arabidopsis, Asp levels were increased 3-fold by treatment with the 316 active (R)-BABA enantiomer but were unaffected by the (L) enantiomer, suggesting that BABA obstructs canonical AspRS activity [18]. Surprisingly, we observed no change in Asp levels in tomato 317 318 plants after BABA treatment. However, there was a significant increase in the levels of the enzymes Asparagine-tRNA synthetase (GInRS) and plastid Glutamyl-tRNA(GIn) amidotransferase (GATC) as 319 well as the Glu, GABA, Pro, Phe, and Tyr, together with a reduction in Ala levels (Fig 2B, 5A). The 320 321 glutamate family pathway is strongly activated under stress conditions, leading to the accumulation of GABA and proline. GABA is synthesized from Glu by a decarboxylation reaction in response to 322 323 abiotic stress, viral infection, and herbivore attack [60]; its increased concentration in BABA-treated tomato plants is almost certainly connected to the strong transcriptomic and proteomic upregulation 324 of its key biosynthetic enzyme glutamate decarboxylase (Solyc04g025530.2.1) (Fig 5C). Proline, which 325 326 plays a pixotal role in responses to abiotic stresses, osmotic, salinity, and low temperature stresses, is synthesized predominantly from glutamate by two successive reductions catalysed by P5C 327 328 synthetase (P5CS) and P5C reductase (P5CR), with P5CS being the rate-limiting enzyme for proline 329 synthesis [62]. However, although BABA treatment increased the abundance of P5CS, it did not 330 upregulate P5CS transcription. This suggests that the increase is due to a change in post-331 transcriptional regulation, as observed previously [63]. Conversely, there was significant 332 transcriptional upregulation of proline dehydrogenases (*ProDH*) (Solyc02g089630.2.1, 333 Solyc02g089620.2.1), which catalyse the catabolic conversion of Pro into the toxic intermediate P5C 334 (Fig 5C, S10 Table). Spraying Arabidopsis leaves with Pro or P5C causes the formation of HR-like 335 lesions resembling those induced by BABA [64]. We therefore suggest that the lesion formation observed in tomato after BABA foliar spraying [19,22] spraying can be attributed to the generation of
 very high local BABA concentrations on leaf surfaces (white deposits) and subsequent P5C
 accumulation.

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Figure 5. BABA-induced changes in amino acid levels and their metabolic pathways. (A) Amino acids whose abundance in BABA-treated Marmande tomato plants differs significantly from that in water-treated controls. Data are means from three replicates; the errors are standard errors of means. Statistically significant differences recorded for each amino acid as determined by the t-test are indicated with different numbers of asterisks (*P < 0.05, **P < 0.01). (B, C) Induction of genes belonging to the phenylpropanoid pathway (B) and glutamate metabolic pathway (C).

E4P – erythrose-4-phosphate, PEP – phosphoenolpyruvate, CRSM – chorismic acid, CM – chorismic acid
 mutase, PPA – prephenate, PDH – prephenate dehydrogenasa, PDT – prephenate dehydratase, PAL –
 Phenylalanine ammonia-lyase, CA – cinnamic acid, C4H – Cytochrome P450, PCA – p-coumaric acid, 4CL – 4 coumarate CoA ligase, THT – Tyramine N-(hydroxycinnamoyl) transferase, GM – Agmatine, ArgDC – Arginine

decarboxylase, 2OG – 2-ketoglutarate, GluDC – Glutamate decarboxylase, GABA – gamma-Aminobutyric acid,
 AS1 – Asparagine synthase, P5CS – Gamma-glutamyl phosphate reductase, AstD – N-succinylglutamate 5 semialdehyde dehydrogenase, P5C – 1-pyrroline-5-carboxylate, P5CR – Pyrroline-5-carboxylate reductase,
 ProDH – Proline dehydrogenase, N.C. – transcript not changed, N.D. – transcript not detected.

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357 BABA treatment also upregulated transcription of branched-chain amino acid 358 aminotransferase (Solyc03q043880.2.1) and cysteine desulfurase (Solyc11q005840.1.1) (Fig 5C). Both 359 of these enzymes are important in the degradation of branched AAs, whose complete oxidation in the mitochondria allows large amounts of ATP to be generated under stress conditions that impair 360 361 photosynthesis [60]. BABA treatment also caused upregulation of the glutamine-dependent asparagine synthetase AS1 (Solyc06g007180.2.1) and LHTs, which function as high-affinity 362 363 transporters of uncharged/acidic AAs in the mesophyll plasma membrane (Solyc11g066800.1.1, Solyc01q111980.2.1, Solyc10q055740.1.1). These changes indicate an effect on overall nitrogen 364 metabolism in the plant (Fig 5C, S10 Table). The upregulation of all these proteins was previously 365 366 observed during chlorosis caused by proteolytic activity and amino acid deamination during P. 367 syringae infections [65]. BABA also upregulated four glutamate receptor genes (SIGLR1.2, SIGLR2.1, SIGLR2.2 and SIGLR2.5) implicated in various processes including the response to aluminium [66] and 368 enhanced drought tolerance in plants [67]. 369

Two prephenate dehydrogenases (Solyc09g011870.1.1, Solyc06g050630.2.1) and one 370 prephenate dehydratase (Solyc06q074530.1.1) were strongly upregulated at the transcript level 371 following BABA treatment, and prephenate aminotransferase was upregulated at the protein level. 372 These changes are clearly connected to the BABA-induced upregulation of the phenylpropanoid 373 pathway and especially its lignin/lignan branch. Important upregulated enzymes of this branch 374 include 4-coumarate CoA ligase (4-CL) and several caffeoyl-CoA O-methyltransferase (COMT), 375 376 laccases (LAC), and peroxidases (PER), which catalyse the synthesis of several secondary metabolites (Fig 5B). BABA treatment also caused transcript-level upregulation of arginine decarboxylase 377 (Solyc10a054440.1.1), which converts arginine into agmatine, a precursor of polyamines including 378 putrescine, spermidine, and spermine (Fig 5B, S10 Table). Interestingly, however, we observed no 379 380 increase in polyamine levels, suggesting that agmatine serves some other metabolic purpose in 381 BABA-treated tomato plants [68]. It could possibly be converted into p-coumaroylagmatine; this 382 hypothesis is supported by the upregulation of 4-CL (SolycO6g035960.2.1) and PAL together with 383 several hydroxycinnamoyl-CoA:tyramine N-(hydroxycinnamoyl) transferases (THT1-3, THT1-4, THT7-1, THT7-8), which are also upregulated during incompatible interactions of Pst with tomato [69] (Fig 384 385 5B, S10 Table).

386 4. DISCUSSION

387 BABA has long been described as an agent inducing highly effective resistance against a wide 388 spectrum of biotic and abiotic stresses in multiple plant species [9,14]. Although much has been 389 learned about the perception and molecular action of BABA in Arabidopsis [17,18], many questions 390 remain regarding its mechanisms of action. Since recent studies have shown that during plant immune responses translation is tightly regulated and poorly correlated with transcription [25], as 391 observed here, we characterized BABA-induced changes in both the transcriptome and the 392 393 intracellular proteome. In our experimental conditions only high BABA concentration after foliar 394 spraying of the crop tomato cultivar Marmande (which would be a viable agricultural application 395 strategy) behaves as a strong stress inducer that does not only triggers highly effective resistance and 396 that deeply remodels transcriptome and subsequent proteome of tomato. Whether BABA primed 397 tomato defences at lower concentrations without any protective effect was not our aim since it 398 cannot be useful for crop protection. Transcriptomic and proteomic analyses showed a similar trend when most of differentially expressed genes and proteins were upregulated and have been linked to 399 general stress responses and defence. Besides, many proteins exhibiting decreased amounts can be 400 401 linked to the general decrease in the rate of photosynthesis and carbon metabolism under stress 402 conditions as previously described [61]. It is this stress-induced reduction in plant growth induced by BABA that is the major factor limiting its commercial exploitation [18]. 403

404 Analysis of the genes and proteins upregulated following BABA treatment revealed enrichment of cellular processes related to primary metabolism and responses to stimuli when BABA 405 406 increased the amount of enzymes involved in carbohydrate metabolism. Previous transcriptomic and genetic analyses have demonstrated the induction of genes involved in carbohydrate metabolism 407 408 upon challenge by pathogens or PAMPs, and the expression of these genes was shown to affect 409 downstream defence responses including ROS production and PR gene expression [61]. Induction of 410 these genes was also observed in genome-wide studies in Arabidopsis plants infected with the avirulent pathogen P. syringae pv. Tomato [70] and in rice leaf sheaths infected by Rhizoctonia solani 411 412 [71]. Thus, the genes upregulated in our study and publicly available datasets of potato [22] and Arabidopsis [26] (Fig 1C) can be seen as common defence response genes that contribute to BABA-IR. 413 414 Previous studies have also shown that BABA-IR is driven by different signalling molecules in 415 different plant species [15,16,72,73]. In Arabidopsis, BABA treatment induces accumulation of SA and A16 SA glucoside (SAG) and causes significant changes in the abundance of isochorismate synthase (ICS), 417 which is directly involved in SA biosynthesis which, in turn, is associated with the expression of acidic 418 PR proteins [15,26]. Although we paid a particular attention to the role of SA in BABA-induced 419 resistance, none of these responses were observed in tomato as in our recent study in which we 420 determined decreased level of SA after BABA treatment of tomato leaflets via petiole aspiration [48]. 421 In addition, unpublished results from our lab, clearly show that SA treatment on tomato leaves failed

422 to induce both a resistance to mildews and the classical defence marker genes outlined in the 423 literature and in this paper (mainly PR-proteins coding genes). It seems that SA is probably not so 424 important in tomato defence to biotic stresses since attempts to protect tomato with SA and SA 425 mimics always failed in available literature with the exception of a protective effect observed 426 towards tomato canker (Clavibacter michiganensis) with acibenzolar-S-methyl. Indeed, some plants, like rice or wintergreen, exhibit a high constitutive level of SA or SA methyl ester without being 427 428 protected from diseases [74] and we have previously demonstrated that SA sensitizes carnation to 429 disease by inhibiting N-benzoate-based phytoalexins biosynthesis [75]. Thus, what was demonstrated 430 in Arabidopsis and even in tobacco should not signify that all the plant kingdom, especially among 431 crops diversity, would follow the same defence regulation scheme. However, treatment of tomato with BABA did affect genes involved in the biosynthesis of JA and ET as well as basic isoforms of PR 432 433 proteins, which were also observed in an earlier study on BABA-treated potato and tomato plants [22,48]. This finding is in accordance with impaired BABA-IR towards *P. infestans* in the tomato *def* 434 435 mutant, which is defective in JA accumulation [73] and pivotal importance of ET in chitosan- and 436 Flg22-induced local and systemic defence responses of tomato plants previously proved in Never-ripe 437 (Nr) tomato mutants exhibiting insensitivity to ET in all vegetative tissues due to mutation in SIETR3 438 receptor [76,77]. All these findings clearly emphasize the need to study plant biology with a 439 necessary hindsight when comparisons to model plants fail to confirm established mechanisms.

A recent screening of Arabidopsis mutations affecting BABA-IR revealed defects associated 440 with the gene IBI1, which encodes aspartyl-tRNA synthetase (AspRS). The specific binding of R-BABA 441 to the L-Asp-binding domain of IBI1 primes the protein for non-canonical defence activity [18]. In 442 accordance with these findings, we found that BABA had no effect on the regulation of the IBI1 443 444 orthologue in tomato plants. However, we found no evidence of any change in aspartate levels driven by the accumulation of uncharged tRNA^{Asp} leading to inhibition of translation activity via 445 phosphorylation of the initiation factor eIF2 α [18]. Instead we observed elevated glutamate levels 446 and increased expression of the enzymes GlnRS and GATC, which are involved in tRNA^{Gln} synthesis 447 448 [78]. Following these findings, BABA treatment of tomato plants upregulated transcription of HSF24 449 (Solyc02q090820.2.1), a HSF-type homologue of TBF1. The heat shock factor(HSF)-like transcription 450 factor TBF1 was proven to play a crucial role in the growth-to-defence switch that activates multiple 451 defence mechanisms and inhibits primary growth and development upon pathogen challenge [79]. 452 Interestingly, one suggested activation mechanism for TBF1 is related to the GCN2-dependent 453 phosphorylation of eIF2 α , which is regulated via the accumulation of uncharged tRNAs. This is 454 consistent with the reported inhibitory activity of R-BABA towards the cellular AspRS activity of IBI1 in Arabidopsis [18]. Moreover, TBF1 controls distinct output genes in SAR and MTI, which could be 455 456 connected to our observation that BABA-IR in tomato is mediated by genes involved in MTI. Together, these observations convincingly explain the finding that BABA has dual activities in tomato, simultaneously activating defence mechanisms and downregulating protein synthesis and carbon metabolism. The latter activity is similar to the stress responses [61] associated with SIMR in *Arabidopsis*. It has been suggested that BABA primes defence responses simply via this stress imprinting process because L-glutamine treatment reduced both BABA-induced SIMR and BABA-IR [80]. In keeping with this hypothesis, 70% of the BABA-induced transcripts overlapped with the set of transcripts upregulated by abiotic salt [39].

Two important processes in pathogen recognition and the subsequent activation of plant 464 465 defence mechanisms are the secretion and spotting of diverse pattern-recognition receptors (PRRs) 466 to the plasma membrane and the activation of protein kinases involved in signal transduction cascades. Augmented perception of stress signals by plant cells seems to be essential in BABA-IR, as 467 468 demonstrated by the significant enrichment of GO terms and pathways related to receptor activity after BABA treatment. Moreover, extensive induction of signalling and perception machinery was 469 470 one of the main factors distinguishing BABA-treatment from the other stress conditions and put 471 forward in our data set. BABA induced a significant number of receptor and receptor-like kinases 472 involved in abiotic stress responses (L-type lectin receptor kinases) [81], MAMPs perception, Phytophthora resistance (LysM, Bti9, SOBIR1) [52,82], and responses to pathogen infection, 473 474 mechanical wounding, and oxidative stress (TPK1b) [83]. This is the first demonstration that BABA-IR in tomato is connected to a hyper-receptive status. In that way, BABA acts as a real priming agent, 475 476 preparing the plant to rapidly recognize pathogens and to set-up strong defences.

However, as stated above, BABA also induce a major plant stress. Whether these two aspects 477 478 can be disconnected is a pending question to only keep the hyper-receptive side unless this status could be a consequence of the major stress. Plant defence responses are also modulated by AA 479 480 homeostasis and treatment with high concentrations of AAs. For example, the Arabidopsis thaliana Iht1 mutant, which has reduced levels of Gln, Ala, and Pro, exhibits SA dependent resistance to a 481 482 wide range of diseases [84]. In addition, treatment of rice roots with the AAs Glu, Asn, Met, and Asp 483 induced systemic disease resistance against rice blast that was partially dependent on SA signalling 484 and did not cause any change in the content of free AAs in leaves [85]. In our experiments, treatment 485 with BABA increased levels of Pro and the expression of the Glu biosynthetic enzyme ProDH, as well 486 as the levels of free Glu in the leaves (Fig 5A, C). The ProDH is an enzyme that plays a crucial role in 487 plant metabolism. Recent studies have shown that ProDH activity is upregulated in response to 488 pathogen infection and contribute to HR and disease resistance, which apparently potentiates the 489 accumulation of ROS. In addition, ProDH may also regulate the balance between proline and P5C, 490 which has been shown to affect the accumulation of defence-related metabolites and the expression 491 of defence genes [86-88]. Decreases in Glu and Pro levels are also associated with microbial

community breakdown and disease incidence, supporting the idea that they play an important role 492 in the plant's defence response [89]. Free Glu may be recognized by glutamate receptor-like proteins 493 494 (GLRs), which have been implicated in enhanced resistance to Hyaloperonospora arabidopsidis, and Pseudomonas syringae in Arabidopsis [90]. The initial cellular events in BABA-IR in tomato may thus 495 involve GLRs as suggested previously in a study on AA-ISR to rice blast in leaves [85]. Interestingly, 496 497 despite significant differences, the upregulated gene clusters in BABA-treated tomato overlap 498 extensively with the sets of orthologous upregulated genes identified by microarray analysis of A. thaliana lht1 plants [84] and the genes upregulated in Glu-treated rice plants [85] (Fig 6A). Notably, 499 500 the orthologous genes common to all three sets exhibited functional enrichment in the plant-501 pathogen interaction pathway (KEGG) and in protein domains related to signal transduction (INTERPRO), suggesting that, in all three cases, induced resistance is driven by similar molecular 502 503 mechanisms based on sensitization to stress responsiveness (Fig 6B), which may be characteristic of 504 priming phenomena [3].



Figure 6. Overlap between genes induced by BABA and genes induced in selected plants with altered amino acid homeostasis. (A) Degrees of overlap between orthologous groups identified in this study and previous transcriptomic studies on A. thaliana lht1 plants [84], and Glu-treated rice plants [85]. (B) Functional enrichment of the plant-pathogen interaction pathway (KEGG) and protein domains related to signal transduction (INTERPRO) among the orthologous genes common to all three sets.

512 Collectively, here we demonstrate that a strong BABA-IR towards *Phytophthora parasitica* 513 and *Oidium neolycopersici* in tomato cv Marmande resemble in many aspects responses to general 514 stress. This resistance was largely explained by the activation of the ET and JA pathways resulting in 515 a strong defence set-up involving PR-proteins as well as phenyl propanoid pathway, lipid 516 peroxidation but in the same time revealed a complete remodelling of plant functions including 517 decrease in primary metabolism and in photosynthesis together with an enhanced ability to perceive 518 (P)(M)(D)AMPs and to set-up downstream signalling. In conclusion, much more attention should be

- 519 paid to comparative studies between plants of agronomic interest submitted to R-BABA treatment.
- 520 We clearly evidenced that multiomics as well as targeted approaches can bring original insight on 521 who to who, even though many black boxes still remain closed.
- 522 5. MATERIALS AND METHODS

523 5.1 Plant material and growing conditions

Tomato plants (*Solanum lycopersicum* cv. *Marmande*) were grown at 75% humidity and 14 hours of light (day 24°C, night 22°C). After 6 or 7 weeks of growth, whole tomato plants were sprayed with 10 mM DL-BABA or water. Leaflets were then removed from plants 24 or 48 hours after spraying for transcriptome and proteome analysis and processed immediately or stored at -80°C until use. Plants assigned to each treatment were randomly selected, labelled, and then returned to the growth chamber. Three biological replicates were selected for each treatment.

530 5.2 Botrytis cinerea and Phytophthora parasitica inoculation and measurement

531 Two days after spraying with 10 mM BABA or water, leaflets were removed from 7-8 weekold tomato plants and placed in clear Styrofoam boxes with moist absorbent paper to maintain high 532 533 relative humidity. The centre of each leaflet was inoculated with a mycelial plug (5 mm in diameter) from the growth margin of a 3-day-old culture of the BC21 strain of *B. cinerea*. Alternatively, leaflets 534 were pricked with a needle at a marked spot and 20 µl of a P. parasitica zoospore suspension (40 000 535 536 zoospores/ml) was applied to the wounded spot. Five replicates in styrofoam boxes with 3 leaflets each were used. After inoculation, the detached leaflets were incubated in a growth chamber under 537 conditions conducive to disease development (21°C, 14-h photoperiod, 114 µmol.s-1.m-2). 538 539 Symptoms were recorded after 3 days of incubation for P. parasitica or 4 days for B. cinerea. 540 Photographs were analysed using the ImageJ image analysis programme to quantify the surface area 541 of necrotic lesions (in mm²). Analysis of variance was used to evaluate whether differences between 542 controls and treatments were significant for each of the three independent experiments.

543 5.3 Oidium neolycopersici inoculation and measurement

544 Spraying with 10 mM BABA or water was done on whole 5-6 week old tomato plants. Two 545 days later, each plant was inoculated with approximately 10 ml of a spore suspension of *O*. 546 *neolycopersici* at a concentration of 10^4 sp/ml. The inoculated plants were then incubated in a growth 547 chamber under conditions conducive to disease development (21°C, RH > 80%, 14 hours of light). The 548 number of powdery mildew colonies was counted 14 days after inoculation on 2 leaves per plant 549 with 5 plants per test. Three independent tests were performed.

550 5.4 Identification and quantification of proteins by LC-MS/MS

551 Three biological replicates in the form of separate plants were subjected to analysis. For each 552 replicate, 4 g of leaflets from 3 different plants were harvested on ice and homogenised at 4°C in 20 553 ml of extraction buffer (50 mM Tris-Mes (pH=8.0), 20 mM EDTA, 500 mM sucrose, 10 mM DTT, 100 mM PMSF, cOmplete Mini Protease Inhibitor Cocktail tablets) using an Ultraturrax homogenizer (IKA, 554 DE) at 15 000 rpm. Samples were filtered through Miracloth and centrifuged at 20 000 x g and 4°C. 555 The supernatant was collected and centrifuged in a Beckman Optima (Beckman-Coulter) 556 557 ultracentrifuge at 35 000 rpm and 4°C with a Ti45 rotor. The supernatant was collected and 558 concentrated using Vivaspin[®] 3 kDa (GE Healthcare) sample concentrators. The concentrated 559 samples were dialyzed overnight to 10 mM ammonium acetate and finally concentrated to 0.5 ml 560 using 4 ml Amicon[®] Ultra 4 3 kDa (Merck Millipore Ltd.) sample concentrators. Each sample was then fractionated into 5 fractions by HPLC using an IEX PolyWAX LP mixed bed column (200 x 4.6 mm, 5 561 562 µm particles, PolyLC Inc., Columbia, USA) and a gradient of ammonium acetate. The collected 563 fractions were dried under vacuum and subjected to LC-MS /MS analysis (S3 Table). The dried protein fractions were processed using a philtre filter-aided sample preparation (FASP) method [83]. 564 565 LC-MS /MS analyses of the peptide mixtures were performed using the RSLCnano system connected to the Orbitrap Elite hybrid spectrometer (Thermo Fisher Scientific). For more details, see the 566 567 supplemental material (S1 Appendix).

568 5.5 RNA sequencing analysis

Total RNA was isolated using TRIzol reagent (Life Technologies, Grand Island, NY, USA) and 569 checked for integrity on a Bioanalyzer 2200 (RIN \geq 7.40). Libraries for sequencing were prepared 570 according to a standard protocol for the SOLiD 5500 system (Life Technology). Sequencing was 571 performed using the SOLID 5500W platform. Raw reads of 75 bp in length were mapped to the 572 Solanum lycopersicum build 2.40 reference using ITAG2.4 as the gene model in colour space with the 573 574 Maxmapper algorithm implemented in Lifescope software (Life Technologies, Ltd). RNA content was assessed using a whole-transcriptome workflow with the quality threshold set to 10, resulting in an 575 576 assignment probability of greater than 90. The raw sequencing data with corresponding metadata 577 are available in the NCBI Gene Expression Omnibus (GEO) repository under accession number GSE108421. Analytical comparison between BABA and the control treatments was performed using 578 the DESeq package [91]. 579

580 5.6 Orthology and Gene Ontology enrichment analysis

581 Orthologous gene clusters were compared and annotated using the OrthoVenn web platform [92], 582 and the results obtained were visualised using the eulerr R package [93]. Significantly differentially 583 expressed genes and proteins were analysed by Singular Enrichment Analysis (SEA) for GO term enrichment using agriGO [94] based on GO terms retrieved from the PLAZA 3.0 database [95]. Summarisation and visualization of SEA were done using REViGO [34] and Cytoscape [96] with DyNet [97]. Protein-protein association networks for significantly differentially expressed genes and proteins were generated using the STRING database with an interaction score of 0.9 [33] and visualized using Cytoscape [96].

589 5.7 HPLC analysis of amino acids

Twenty-four hours after treatment with BABA, tomato leaves were collected, frozen in liquid 590 591 nitrogen, and ground to a fine powder. A portion of 250 mg of this powder was then extracted with 1 592 ml of extraction buffer (0.1 M HCl and 4.6 µg/ml L-2-aminoadipic acid as an internal standard), mixed thoroughly, incubated on ice for 5 minutes, and centrifuged. A 500 μ l aliquot of the resulting 593 594 supernatant was then diluted with 100 µl methanol and loaded onto an SPE C18 column to adsorb interfering secondary metabolites, which had previously been wetted with 1 ml MeOH and 595 596 equilibrated with 20% MeOH in 0.1M HCI. The sample was loaded and the column was washed with 597 400 µl of 20% MeOH in 0.1 M HCl. Amino acids were recovered in both the flow-through and wash fractions and derivatized and analyzed as previously described [98]. See supplemental material (S1 598 599 Appendix) for further details.

600 5.8 Quantitative analysis of salicylic acid, jasmonic acid, and jasmonic acid-isoleucine

601 The tomato leaflest (100 mg) was frozen immediately after the harvest using liquid nitrogen, and the frozen materials ground under liquid nitrogen and extracted with 750 µl of MeOH-H₂O-HOAc 602 603 (90:9:1, v/v/v) containing 100 ng of o-anisic acid as an internal standard. The mixture was centrifuged 604 at 10,000 x g for 1 min, the supernatant was collected, and the pellet was repeatadly extracted. The pooled supernatants were dried under nitrogen, resuspended in 200 µl of 0.1% HOAc in H₂O-MeOH 605 606 (90:10, v/v), and a portion of the mixture $(2-5 \mu I)$ was subjected to LC-MS analyzes using a TOF mass 607 spectrometer (Agilent Technologies) as previously described [99]. Further details can be found in the 608 supplemental material (S1 Appendix).

609 5.9 Measurement of ethylene production

Single tomato leaflets 24 hours after treatment with BABA or water were placed in 20 ml test tube when the cut end of the petiole was in sterile water and sealed with a air-tight rubber syringe cap. ETwas accumulated for 4 hours before a 1-ml sample was withdrawn for analysis. ET production was measured using gas chromatography with a flame ionization detector quantified by using a gas chromatograph flame ionization detector (Agilent GC 6890, Agilent Technologies) as previously described [19].

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- 626 Contributions
- 627 Conceptualization: JL, MP, CR and ZZ; Data curation: MZ, CR, BI, MB, KLB, PN, VM, AS, PB, and DP;
- Data analysis: MZ, CR, DP, AS and BI; Funding acquisition: JL, MP and ZZ; Methodology: MZ, CR, JL,
- 629 MP and ZZ; Project administration: JL and MP; Resources: JL, MP and ZZ; Validation: MZ, CR, JL, MP
- and ZZ; Writing, review, and editing: MZ, JL, and MP; Supervision: JL and MP.
- 631 Data availability statement
- Raw sequencing data with appropriate metadata are available in the NCBI Gene Expression Omnibus
- 633 (GEO) repository under accession number GSE108421.
- 634 The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium
- via the PRIDE partner repository with the dataset identifier PXD038074
- 636 Competing interests
- 637 We declare that none of the authors have any competing interests.

638 7. SUPPLEMENTARY INFORMATION

- 639 S1 Appendix Supplementary methods
- 640 **S1 Figure.** HR-like microlesions on BABA spayed leaves.
- 641 **S2 Figure.** ReviGO analysis of enriched GO terms.
- 642 **S3 Figure.** BABA perception affects gene expression of certain protein kinase families.
- 643 **S4 Figure.** BABA perception affects gene expression of certain transcription factor families.
- 644 **S1 Table.** Workflow of sequence analysis.
- 645 **S2 Table.** Significant differentially expressed proteins and transcripts.

- 646 **S3 Table.** Identified protein groups.
- 647 S4 Table. Orthologous across our and previous transcriptomic studies of BABA treated potato and
 648 *Arabidopsis* plants.
- 649 **S5 Table.** Overlap between genes induced by BABA and different elicitors.
- S6 Table. GO analysis of up-regulated proteins and transcripts in tomato leaves after treatment with
 10 mM BABA.
- 652 S7 Table. Protein-protein association network for significantly induced genes and proteins
- 653 **S8 Table.** Involvement of signalling pathways in BABA-IR.
- 654 S9 Table. Transcripts of protein kinases, receptor like proteins and transcription factors found in our
- 655 study and their comparison with other studies.
- 656 **S10 Table.** Induction of genes in glutamate metabolic pathway and phenylpropanoid pathway.
- 657

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