- Reshaping of the *Arabidopsis thaliana* proteome landscape and
 co-regulation of proteins in development and immunity
- 3 4 Mona Bassal, Petra Majovsky, Domenika Thieme, Tobias Herr, Mohammad Abukhalaf, 5 Mohamed Ayash, MHD Rami Al Shweiki, Carsten Proksch, Ali Hmedat, Jörg Ziegler, 6 7 Steffan Neumann and Wolfgang Hoehenwarter 8 Leibniz Institute of Plant Biochemistry, Weinberg 3, Halle/Saale D-06120, Germany 9 10 Keywords: Arabidopsis, deep proteomics, protein co-expression. proteome, 11 senescence, ribosome, seed proteome, photosynthesis, plant immunity, jasmonate 12 13 Address correspondence to 14 Wolfgang Hoehenwarter 15 16 Proteome Biology of Plant Interactions Research Group Leibniz Institute of Plant Biochemistry 17 Email: wolfgang.hoehenwarter@ipb-halle.de 18 19 20 21 22
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24 Abstract

Proteome remodeling is a fundamental adaptive response and proteins in complex and 25 functionally related proteins are often co-expressed. Using a deep sampling strategy we 26 define Arabidopsis thaliana tissue core proteomes at around 10,000 proteins per tissue 27 28 and absolutely quantify (copy numbers per cell) nearly 16,000 proteins throughout the plant lifecycle. A proteome wide survey of global post translational modification revealed 29 amino acid exchanges pointing to potential conservation of translational infidelity in 30 eukaryotes. Correlation analysis of protein abundance uncovered potentially new tissue 31 and age specific roles of entire signaling modules regulating transcription in 32 photosynthesis, seed development and senescence and abscission. Among others, the 33 34 data suggest a potential function of RD26 and other NAC transcription factors in seed development related to desiccation tolerance as well as a possible function of Cysteine-35 rich Receptor-like Kinases (CRKs) as ROS sensors in senescence. All of the 36 components of ribosome biogenesis factor (RBF) complexes were co-expressed tissue 37 38 and age specifically indicating functional promiscuity in the assembly of these little described protein complexes in Arabidopsis. Treatment of seedlings with flg22 for 16 39 40 hours allowed us to characterize proteome architecture in basal immunity in detail. The results were complemented with parallel reaction monitoring (PRM) targeted 41 proteomics, phytohormone, amino acid and transcript measurements. We obtained 42 strong evidence of suppression of jasmonate (JA) and JA-Ile levels by deconjugation 43 and hydroxylation via IAA-ALA RESISTANT3 (IAR3) and JASMONATE-INDUCED 44 OXYGENASE 2 (JOX2) under the control of JASMONATE INSENSITIVE 1 (MYC2). 45 This previously unknown regulatory switch is another part of the puzzle of the as yet 46 understudied role of JA in pattern triggered immunity. The extensive coverage of the 47 Arabidopsis proteome in various biological scenarios presents a rich resource to plant 48 biologists that we make available to the community. 49

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53 Introduction

Proteome biology is receiving increasing interest in the last years but still proves difficult 54 on a genome wide scale in plants. The proteome is the most fundamental active 55 determinant of an organism's phenotype and its landscape is large, complex and 56 dynamic, entailing changes in protein abundance, interaction, post translational 57 58 modification (PTM) and sub-cellular localization. 59 Steady state protein abundance at a certain time point is to a considerable part determined by the abundance of its transcript and the latter's translation rate. Synthesis 60 is however only half of the equation governing protein abundance, indeed Arabidopsis 61 has more than 600 F-box proteins as components of diverse E3 ubiquitin ligase 62 complexes that direct protein degradation. Newer evidence has shown that post 63

transcriptional and translational mechanisms (Ponnala et al., 2014; Merchante et al.,
2017) and phenomena such as cell-to cell mobile mRNAs (Thieme et al., 2015) and

66 proteins (Han et al., 2014; Guan et al., 2017) play equally important roles in determining

the proteome's temporal and spatial plasticity foremost in steady state shifts. Therefore,

68 despite the continued practice of quantifying the abundance of proteins' cognate

transcripts to estimate and quantify functional protein coding gene expression, direct,

⁷⁰ large scale measurement of protein abundance and PTM should be the explicit end

71 point of functional genomics.

72 The problem with this is that proteomics has long been beset by a lack of sensitivity,

r3 especially in plants. Indeed, only a handful of true deep proteomics studies in this

kingdom can be found. Walley and co-workers constructed a protein co-expression

network based on measurement of 17,862 proteins and 6,227 phosphorylated proteins

in maize (Walley et al., 2016). The tissue and development specific wheat proteome has

been mapped with measurement of 15,779 proteins (Duncan et al., 2017) as well as the

tomato fruit where 7,738 proteins were measured in one or more developmental stages

79 (Szymanski et al., 2017). Song and co-workers reported optimized FASP sample

80 preparation in conjunction with 2D-LC-MS/MS allowed measurement of 11,690 proteins

from a single Arabidopsis leaf sample (Song et al., 2018) and Baerenfaller and co-

workers reported measurement of nearly 15,000 *Arabidopsis* proteins more than ten
years ago (Baerenfaller et al., 2008).

Here we go beyond this classic study and elucidate *Arabidopsis thaliana* tissue specific proteome remodeling in development and immunity in unprecedented resolution and detail. We describe a procedure that allows deep sampling of up to 9,000 proteins per plant sample and does not require the expertise necessary for 2D-LC-MS. We quantified nearly 16,000 proteins in absolute terms (copy number per cell) and conducted a global analysis of protein PTM.

90 A long standing caveat in proteomics is proteomics "dark matter" (Skinner and Kelleher,

2015) meaning an abundance of high quality MS2 spectra that do not result in PSM.

Many of these MS2 spectra are derived from peptides bearing PTM, however classic

search algorithms necessitate PTM predefinition, limiting potential PTM identification to

⁹⁴ a handful. Recent years have seen the advent of "open search" algorithms (Chick et al.,

2015; Kong et al., 2017; Bagwan et al., 2018; Chi et al., 2018) that allow unrestricted

96 precursor mass shifts in PSM, and so have the potential to identify the vast array of

biologically occurring modifications on a proteome wide scale. Here we performed such

a survey using the MSFragger (Kong et al., 2017) software suite in *Arabidopsis*.

99 Large scale protein co-expression analysis is a potentially powerful strategy to

100 determine functional relationships between proteins because it circumvents the

101 limitations inherent to transcriptomics measurements described above. We applied

102 clustering algorithms to our data set that provides extensive coverage of the

103 Arabidopsis proteome to uncover tissue and developmentally specific protein

104 expression patterns. The approach was effective in pinpointing co-regulation of all

105 components of protein complexes and developmentally timed signaling modules. More

importantly, it then allowed inference of previously unknown functions of proteins,

107 protein families and entire signaling modules based on the same expression patterns.

108 This included processes such as tissue and age specific ribosome biogenesis,

109 photosynthesis, ABA signaling and NAC transcription factors in seed development and

110 establishment of dormancy and senescence and abscission.

The various sequential steps of ribosome biogenesis including the involved RPs and 111 112 RBFs are well described in yeast and human (Henras et al., 2008; Henras et al., 2015). Mutations in numerous RPs and RBFs are known to cause severe developmental 113 defects, so called ribosomopathies in humans. This also holds true in Arabidopsis 114 where a number of mutations in RBFs affect gametophyte development and 115 116 embryogenesis (Byrne, 2009; Weis et al., 2015). Furthermore Arabidopsis ribosomes are extensively heterogenic, each individual RP being encoded by two to seven 117 118 paralogs (Weis et al., 2015). This heterogeneity of ribosome species is dependent on developmental stage, tissue and environmental stimuli, suggesting that the specific 119 ribosome constituency may play a regulatory role in these processes. 120

121 Cotyledons of young seedlings are characterized by light induced chloroplast

biogenesis. Photomorphogenesis includes tetrapyrrole and chlorophyll biosynthesis to

establish photosystems along with carotenoid synthesis as an accessory pigment and

124 ROS scavenger. Once photoautotrophic, protein biosynthesis is ramped up, hallmarked

by the increased abundance of ribosomal proteins and proteins involved in translation

and folding. Concomitantly, nuclear encoded proteins synthesized in the cytosol are

imported. Finally mature chloroplasts with their established structures proliferate by

division to accommodate cell expansion and division in the growing leaves.

Senescence is a coordinated process with several stages and developmental check
points (Bleecker and Patterson, 1997; Rogers and Munne-Bosch, 2016). Early

150 points (Dieecker and Fatterson, 1997, Rogers and Munne-Dosch, 2010). Lany

senescence syndrome involves reprogramming of gene expression (Breeze et al.,

132 2011) and redox and ROS signaling. This is followed by ordered dismantling of the

photosynthetic apparatus leading to ROS production and involving ROS control and

nutrient remobilization to other plant parts in the case of leaves and to the developing

ovary in the case of floral petals. The final result is dell death lastly followed by

abscission in some organs such as floral petals.

137 Several post-transcriptional/translational mechanisms controlling protein abundance

have just recently been uncovered in plant immunity (Meteignier et al., 2017; Xu et al.,

139 2017; Tabassum et al., 2019). Therefore we investigated reshaping of the proteome in

the steady state shift from growth to pattern triggered immunity (PTI). We measured

changes in protein abundance of more than 2,000 proteins in all avenues of PTI as well 141 as photosynthesis and primary metabolism, giving a comprehensive picture of altered 142 proteome architecture in basal immunity. The focus was set on hormone signaling and 143 the deep proteomics measurements were complemented by targeted parallel reaction 144 monitoring (PRM) proteomics, qPCR and phytohormone and amino acid 145 measurements. Salicylic acid (SA) and Jasmonate (JA) are the phytohormones 146 quintessentially associated with plant defense, the former constituting the backbone of 147 148 resistance to biotrophic pathogens, the latter to necrotrophic pathogens as well as wounding (Pieterse et al., 2012). There mutual antagonism to prioritize one defense 149 strategy over the other as needed is a long standing paradigm in the field of plant 150 immunity. However research in the last decade has made it clear, that JA also plays a 151 152 role in resistance to biotrophs and pattern triggered immunity (PTI) which is however not yet fully understood (Nickstadt et al., 2004; Hillmer et al., 2017; Mine et al., 2017). The 153 integrative –omics strategy applied here brought to light a previously unknown 154 regulatory switch wherein a MYC2 dependent negative feedback loop controls JA-Ile 155 156 and JA levels via deconjugation and hydroxylation by IAR3 and JOX2 respectively. It reconciles the well-known dampening of JA levels and suppression of JA signaling 157 158 downstream of JA synthesis by SA, adding another feature to the picture of JA activity in PTI. 159

Finally we make both the raw and meta data produced in this study available to the general public (submission to Proteome Exchange and TAIR pending). The sampling depth of the proteome in development and immunity should prove to be a valuable resource to plant biologists.

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166 Results

167 Deep proteomics method

168 We conducted discovery proteomics measurements of several Arabidopsis thaliana Col-

169 0 tissues throughout the lifetime of the plant. These were roots, leaves, cauline leaves,

stem, flowers and siliques/seeds as well as whole plant seedlings as early as seven
days up to 93 days of age when the plant was in late senescence (Supplemental figure
1 and Supplemental methods and data tables 1 and 2). In addition we measured
proteomes of PTI elicited plants treated with the peptide flg22. Primarily we were
interested in reshaping of proteome architecture in these different biological scenarios
to capture protein co-regulation and predict tissue and developmentally specific protein
function.

To this end it was essential to develop a method that allows deep sampling of the plant proteome in a reasonable time frame. It was clear that this would have to begin with comprehensive extraction of tissue proteins and further entail multi-step fractionation of the complex extract. We settled on and optimized 4% SDS protein extraction and Gel-LC MS combining SDS-PAGE protein separation and reverse phase (RP) LC peptide separation on-line with ESI MS (see Supplementary methods and data and

183 Supplemental figure 2 for full method and optimization details).

Plant tissue is more recalcitrant to proteomics analysis than other samples. The plant
cell wall requires harsh disruption techniques. Plant tissue contains an abundance of
secondary metabolites, oils and waxes and more significantly, in the case of green
tissue, pigments as part of the light harvesting complexes, that all interfere with LC-MS.
Green tissue also contains the most abundant protein on earth, Ribulose bisphosphate
carboxylase (RuBisCo) leading to suppression of less prominent ion signals in the mass
analyzer and severely hampering detection of less abundant proteins.

SDS-PAGE alleviated all of these issues (Supplemental figure 3 A): i) It allowed the depletion of high amounts of SDS used for protein extraction. ii) Pigments and small molecules conglomerated in a green low molecular weight band below the protein front effectively partitioning them from the proteins. iii) RuBisCo large and small subunits (RBCL and RBCS) migrated to two prominent bands facilitating their separation from the rest of the proteome.

Five gel slices were in gel digested with trypsin and individually injected into the LC-MS
neatly fractionating the proteome according to the molecular weight of its constituents
(Supplemental figure 3 B). The most abundant plant proteins including RBCL and RBCS

were separated in individual fractions in both leaves and roots, diminishing the 200 suppressive effects of over abundant proteins on peptide and protein identification in 201 202 single shot LC-MS measurement of the entire proteome (Supplemental figure 3 C and D). This allowed the identification of between 5977 and 9524 protein groups per sample, 203 each identified with at least 1 unique peptide at protein and peptide FDR thresholds of 204 205 1%. These protein groups will henceforth be referred to as proteins (Supplemental table 1 and Supplemental figure 3 E). Measurements of the individual proteomes showed 206 intra sample variability that was considerably lower that the observed inter sample 207 variability (Supplemental figure 3 F and G). 208

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210 The Arabidopsis proteome

211 In the entire study we identified and quantified 15926 Arabidopsis thaliana proteins in 212 total, 15845 encoded in the nuclear genome (Supplemental table 2). For further details on the parameters of the entire data set see Supplemental methods and data Table 3. 213 Nuclear protein identifications were evenly distributed over all five chromosomes. We 214 complemented the proteomics data with transcriptomics data from an extensive study of 215 Arabidopsis development using microarrays that measured gene expression on 22157 216 nuclear loci including 1058 non-protein coding genes (Schmid et al., 2005). Our 217 218 proteomics data presents mass spectrometric evidence for cognate protein expression of 60% (and locally more) of the Arabidopsis nuclear genome and close to 70% or more 219 220 of the transcriptome, including 1886 proteins for which no transcripts were identified or 221 which were not on the microarray (Figure 1 A and B).

222 We then asked what precluded more extensive coverage of the Arabidopsis proteome, 223 i.e. the missing proteome. We first looked at a potential impact of transcript abundance on the detection of cognate proteins and found that missing (not detected) proteins are 224 evenly distributed over the range of transcript abundance with the exception of the most 225 226 abundant transcripts for nearly all of which a cognate protein was detected (Figure 1 C). 227 This indicates that transcript abundance is not the primary factor impeding protein detection. Next we investigated if gene length in base pairs as a proxy for protein length 228 229 may have an impact and found that indeed the distribution of missing proteins was



shifted markedly towards smaller genes from the distribution of detected proteins

232 Figure 1 Deep coverage of the Arabidopsis thaliana proteome. A. Mapping protein expression to the 5 Arabidopsis nuclear 233 chromosomes (scale in Megabases (Mb), centromers are indicated). Tracks from outside to inside: Number of protein coding 234 genes per 500 Kb; scale 0 to 167 (maximum) [dark green]. Number of non-protein coding genes per 500 Kb; scale 0 to 167 [light 235 green]. Number of transcripts per 500 Kb; scale 0 to 167 [dark blue]; Log₁₀ transcript abundance normalized to maximum 236 transcript abundance; scale 0 to 1 [light blue]. Number of proteins per 500 Kb; scale 0 to 167 [dark red]. Log₁₀ protein 237 abundance normalized to maximum protein abundance; scale 0 to 1 [light red]. Fraction of protein coding genes for which 238 cognate proteins were detected; scale 0 to 1 [magenta]. Fraction of transcripts for which cognate proteins were detected; scale 239 0 to 1 [light purple]. B. Scaled VENN diagram showing the coverage of protein coding genes by cognate transcripts and proteins. 240 C. Relationship between transcript abundance and detection of its cognate protein. Green: all protein coding transcripts, red: 241 protein coding transcripts with a detected cognate protein, blue: protein coding transcripts without a detected cognate protein. 242 Frequency polygon drawn with 20 bins D. Relationship between protein coding gene size in bp and detection of its cognate 243 protein. Green: all protein coding genes, red: protein coding genes with a detected cognate protein, blue: protein coding genes 244 without a detected cognate protein. Frequency polygon drawn with bin size 190 bp. E. Size in bp of genes assigned to the gene 245 ontology terms regulation of transcription, DNA templated (GO:0006355), extracellular region (GO:0005576), mitochondrion 246 (GO:0005739) and nucleus (GO:0005634). Solid lines indicate all members of the GO term, dashed lines indicate members of the 247 GO term not identified in the proteomics measurements, i.e. the missing proteome. Gene size of all proteins not identified in the 248 proteomics measurements, the entire missing proteome is also indicated. Frequency polygon drawn with 350 bins and bin size 249 190 bp. F. Cumulative increase in identified proteins, i.e. proteome coverage, as different tissues and flg22 treatment are 250 sampled. G Copy numbers per cell of all identified proteins and leaf proteins as determined by proteomics ruler method 251 (Wisniewski et al., 2014). Quartiles are indicated by dashed lines. Significantly enriched GO annotations by DAVID are shown for 252 the individual quartiles. H Global survey of PTM in the Arabidopsis thaliana proteome. The relative abundance PTMs comprising 253 0.1% or more of the total PSMs in at least 1 tissue type or biological scenario (flg22 treatment) are shown.

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255 (Figure 1 D). This suggests that protein size is a factor limiting protein detection. This effect could be observed for several gene ontology categories that were 256 overrepresented in the missing proteome (Supplemental figure 4). Then we examined 257 how tissue specific protein expression contributed to the cumulative expression of the 258 259 entire proteome (Figure 1 F). While every tissue contributed to cumulative protein expression curve with a steep slope, it can also be seen that the curve converges on 260 261 saturation as sample numbers increase. This was also the case for flg22 treated tissues sampled last, where one could expect expression of a host of immunity specific proteins 262 not detected in normally developing tissue. This suggests that further sampling of 263 Arabidopsis would not greatly increase coverage of the entire proteome. 264 We quantified all 15,927 measured proteins in terms of protein copy number per cell in 265 the entire sample set and in leaf tissue (LF) by way of the proteomics ruler approach 266 (Wisniewski et al., 2014). It equates the total histone MS signal (total number of histone 267

PSMs in our case) to the cellular DNA mass allowing the conversion of PSMs to mass

units and calculation of the total cellular protein mass. Individual to total protein PSM

ratios can then be used to calculate individual protein copy numbers per cell.

The total cellular protein mass was calculated as 256.5 pg, in agreement with amounts 271 from cell lines and tissues (Wisniewski et al., 2014). Protein copy numbers spanned a 272 273 range of more than 5 orders of magnitude from just below 100 to 2.29e+07 copies per 274 cell (Figure 1 G, Supplemental table 3). The most abundant protein was RBCL with 4.73e+07 cellular copies in leaves. The total copy number of all detected small subunits 275 276 was 5.4e+07 so the ratio of large to small subunits was 0.876, close to a 1:1 ratio. Gene ontology analysis of the proteins in the four quartiles showed that nuclear proteins and 277 regulators of transcription were very specifically among the least abundant proteins 278 (1750 of 3970 1st quartile proteins were annotated as nucleus). Membrane spanning 279 proteins and kinases were generally also lower abundant, 1108 of 3970 2nd quartile 280 proteins being annotated as integral membrane. 281

282 Protein post translational modification (PTM) is an essential modulator of protein

function. Therefore we performed a proteome wide survey of global PTM in Arabidopsis

thaliana with our deep proteomics data set using the "open-search" algorithm MS-

Fragger (Kong et al., 2017). This led to the identification of more than 3.5 million PSMs

from more than 11.6 million total acquired MS2 spectra. The most abundant PTMs

comprising more than 0.1% of total PSMs in at least one tissue type or biological

scenario are shown in figure 1 H (for full data see Supplemental table 4). Next to

289 predominantly experimentally induced PTM (protein oxidation and

carbamidomethylation of cysteine residues to reduce disulfide bonds), serine or

threonine phosphorylation and N-terminal acetylation were abundant naturally occurring

292 modifications affecting approximately 1.25% of total protein abundance.

293 Transpeptidation reaction, a non-translational mechanisms for the formation of peptide

bonds, derived addition of amino acids was also detected. Furthermore, a number of

amino acid substitutions in protein primary structure were common.

296 Architecture of Tissue and Developmental Proteomes

297 We utilized our extensive MS data collection as a resource to investigate the individual

tissue proteomes during the *Arabidopsis* life cycle and co-regulation of protein

abundance between them (Supplemental table 5). First we grouped all of the data

tissue specifically also merging rosette and cauline leaves and qualitatively compared

the tissue proteomes which each comprised around 10,000 proteins (Supplemental
table 6). Around 6,500 proteins (by far the largest set), were ubiquitous to all tissues
whereas 500 to 600 proteins were unique to each tissue with exception of leaves which
showed nearly 1,000 unique proteins, perhaps as a consequence of the larger number
of aggregated leaf samples (Figure 2 A). Also around 1,000 proteins were absent in
roots but present in all other tissues, reflecting the former tissues below ground nature.

The large volume of the data made PCA an attractive method to reduce its 307 dimensionality and explore the relationships of the sampled proteomes. The first two 308 principal components accounted for nearly 50% of the total variance indicating that the 309 310 linear projection in this two dimensional subspace reflects the predominant data structure (Figure 2 B). The individual tissue proteomes were clearly separated with the 311 312 exception of flowers and the inflorescence stem at the same point in development (66 and 73 days). More interestingly, a developmental component was also visible in all 313 314 sampled tissues, reflecting proteome remodeling during ageing (increasing sample age bottom right to top left; arrow). Hierarchical cluster analysis (HCL) corroborated the PCA 315 316 results (Supplemental figure 5).

To investigate the dynamics of proteome structure and protein co-expression in tissue development in detail and thereby extrapolate protein function in previously undisclosed contexts, a noise robust soft partitioning technique that does not assign a feature exclusively to a single group, called fuzzy c-means clustering was applied (citation). The procedure produced 7 out of 16 clusters (clusters numbered 3, 4, 7, 8, 10, 11, 13) with biologically meaningful changes in protein abundance with a permutation based FDR of less than 1%.

324 Root Proteome

The largest set of 577 proteins assigned to a cluster were root specific and not abundant or present in any other tissues at any of the sampled developmental stages (Cluster 3, Figure 2 C, Supplemental table 7). Gene ontology analysis of these proteins indicated that many of them were involved in processes related to the extracellular region, metal- binding and oxidation and reduction. We described the expression of these proteins in the context of phosphate metabolism previously (Hoehenwarter et al.,

- 2016). We compared our results to an exhaustive study of the root proteome by Lan et
- al (Lan et al., 2012) and found that around 70% of the total root proteins we detected
- 333 were also present there. This intersection of 7577 proteins between our two respective



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335 *Figure 2 Tissue and development specific dynamic architecture of the* Arabidopsis

- 336 proteome. A. Qualitative comparison of Arabidopsis tissue proteomes. All
- 337 measurements of a tissue regardless of plant age were aggregated, leaves and cauline
- 338 leaves as well. B. Principal component analysis (PCA) of the deep proteomics
- 339 measurements of sampled tissues. Numbers indicate sample, i.e. plant age in days.

Tissues are color coded. Purple indicates late flowering stages, i.e. siliques and leaves 340 341 in senescence. C. FCM cluster 3 showing proteins exclusively abundant in roots. Right panel: Qualitative comparison of all root proteins identified in this study and all root 342 343 proteins identified in Lan et al. FCM core clusters were extracted wherein all clustered proteins have membership values α exceeding a threshold of 0.5 meaning cluster 344 members all have similarity to the cluster centroid greater than 0.5 (Futschik and 345 Carlisle, 2005). D. FCM cluster 7 showing proteins increasing in abundance in seed 346 development and exclusively present in seeds or siliques. Right panel: Protein 347 abundance of kinases and transcription factors (PQI given is raw #PSMs) in the 348 respective samples. E. FCM cluster 4. Right panel: Physical protein interaction networks 349 produced with STRING database based on homology to yeast and human studies. Solid 350 spheres in the UTP-B, t-UTP and PeBoW networks are proteins designated as DWD40 351 352 by DAVID gene ontology analysis and part of the respective STRING input dataset (WD40/YVTN repeat-like-containing domain list Supplemental Table 9). Unfilled 353 spheres are not part of the WD40 input set but are also members of cluster 4. Thin 354 black edges indicate bona fide physical interactions identified in humans (Wan et al., 355 356 2015). Green edges indicate bona fide physical interactions identified in Arabidopsis (Ishida et al., 2016). Red underlines indicate deletion of the gene has a developmental 357 phenotype. Yeast homologs of Arabidopsis gene names are given in italics. F. FCM 358 359 cluster 13.

- 360 studies allows us to define the core set of proteins most abundant in the root, the root
- 361 core proteome.
- 362 Seed Proteome

363 Cluster 7 (Figure 2 D) contains proteins whose abundance increased specifically in

- seed. As expected, many of these were the more abundant seed storage proteins such
- as oleosins, albumins, cruciferins and enzymes of sugar and fatty acid metabolism.
- However, many of the core transcription factor, phosphatase and chromatin modifier
- 367 modules that regulate seed development were also detected and quantified
- 368 (Supplementary table 8 and Figure 2 D). The PP2C POL directs *WUS/WOX5* gene
- 369 expression and is essential for meristem establishment and stem cell maintenance in
- the early embryo together with PLL1 (Song et al., 2008). Similarly, the AIL transcription
- factor BBM and HDG1 act antagonistically to balance stem cell proliferation and
- differentiation (Horstman et al., 2015) and AN3 establishes cotyledon identity upstream
- of PLETHORA1 (Kanei et al., 2012). The ABA signaling proteins which are known to
- play a major role in seed development were underrepresented in cluster 7 mainly
- because they either did not accumulate exclusively in seed or not in both of the
- measured stages (developing green and ripe brown siliques) but there abundance could

be easily reconstituted from the data. LEC1, the master inducer of seed development 377 was highly abundant in the earlier stage together with AREB3 and EEL, two ABA 378 379 responsive bZIP transcription factors that govern early seed maturation (Agarwal et al., 2011). The ratio of the bZIP transcription factor ABI5 to AFP1, a repressor of ABA 380 signaling, both detected exclusively in mature to post mature seeds was greater than 1 381 indicative of high ABA levels and induction and possible maintenance of seed dormancy 382 in the brown siliques sample. Concomitantly the DOG1 protein and the RDO5 PP2C 383 that are essential for seed dormancy (dormancy is completely abolished in the dog1 384 knockout mutant also in the presence of ABA (Nee et al., 2017)) were abundant in the 385 post mature seed proteome. A number of other PP2C proteins also accumulated to high 386 levels specifically in siliques, particularly in brown siliques. The same was true for 387 several members of a clade of NAC transcription factors also hitherto not known to play 388 a role in seed development indicating potential functions for them in seed development 389 and establishment of dormancy. 390

391 Ribosomal Proteins in Development

The abundance of the 315 proteins in cluster 4 strongly increased or was exclusively measured in young roots, young stem and early flowers/floral buds. There abundance decreased in the latter two tissues as development progressed (Figure 2 E). More than half of these proteins were localized in the nucleus and a substantial number of them pertained to nucleolar processes and ribosome biogenesis with WD40 proteins, containing the WD40 repeat molecular interaction domain, being highly significantly enriched (Supplemental table 9).

One of these was LEUNIG (LUG), a transcriptional co-repressor and master regulator of flower development that directly modulates antagonistic A and C class gene expression in the inner and outer whorls (Grigorova et al., 2011). TAIR10 annotated six WD40 proteins as DWD components of CUL4 RING E3 ubiquitin ligases (Supplemental table 9) but closer inspection revealed that AT3G56990 (EDA7) and AT4G04940 actually lacked the canonical DWD motif (Supplemental material and data), while the others belonged to the A clade of Arabidopsis DWD proteins (Lee et al., 2008). Among them

406 was DWA2, a negative regulator of ABA signaling that targets ABI5 to degradation (Lee407 et al., 2010).

We investigated potential interactions among the 18 WD40 proteins in cluster 4 using 408 the STRING database set to produce essentially only true positive binary physical 409 410 interactions. Five of the six components of the UTP-B complex, a sub-complex of the 411 SSU-processome/90S pre-ribososme, were found to interact based on experimental evidence from yeast and human (Gavin et al., 2002; Gavin et al., 2006; Krogan et al., 412 2006; Wan et al., 2015), showing tissue and development specific expression of this 413 entire protein complex (Figure 2 E, Supplemental figure 6 and Supplemental table 9 for 414 415 raw abundance values). The complex interacted with EDA7, the homolog of enp2 (Soltanieh et al., 2014), a putative ribosome biogenesis factor (RBF) which is not known 416 417 in Arabidopsis. NuGWD1 a sugar inducible WD40 protein was reported to interact with SLOW WALKER 1 (SWA1) which is a component of the t-UTP sub-complex of the SSU 418 419 processome and which was also shown to interact with several UTP-B members by co-IP MS in Arabidopsis (Ishida et al., 2016). SWA1 is not a WD40 protein but the pattern 420 421 of its abundance was very similar to the protein expression pattern of the cluster 4 proteins (Supplemental Table 9 for raw abundance values). Another WD40 protein 422 423 containing complex, all of whose components were measured and showed the same 424 expression pattern, elevated in roots, young shoots and early flowers/floral buds was the PeBoW complex which is essential for pre-rRNA processing (Cho et al., 2013; Ahn 425 et al., 2016). Additionally, extensive physical interaction between a large number of 426 427 ribosomal (RPs) and other ribosome associated proteins with this expression pattern became evident when interactions between all 315 cluster 4 proteins was assayed 428 (Supplemental figure 6). However, the cluster 4 expression pattern of the RBF protein 429 430 complexes was significantly more conserved than expression patterns of ribosomal proteins (Supplemental Figure 7). 431

Molecular chaperones were also highly significantly enriched among the cluster 4
proteins. Nine of these 21 proteins putatively interact physically based on x-ray
crystallography and tandem affinity purification studies in yeast and human (Dekker et
al., 2011; Hauri et al., 2016). All of these TCP-1/cpn60 proteins are homologs of the

436 yeast and human cytosolic chaperonin CCT proteins suggesting they constitute the
437 hardly described CCT complex in *Arabidopsis* and that it's abundance is tissue and
438 developmentally specific (Figure 2 E and Supplemental table 9 for raw abundance
439 values).

440 Vesicle Trafficking and Transport

441 A set of 153 proteins were abundant in roots, increased during ageing and peaking in senescence in leaves and increased in the young inflorescence stem and flowers 442 (Cluster 13, Figure 2 F Supplemental table 10). These proteins were largely related to 443 vesicle trafficking, Golgi apparatus and membrane transport. They contained numerous 444 445 exocyst complex members, SNAREs and cytoskeletal proteins such as actin. Vesicle 446 trafficking and cytoskeletal remodeling and organization are central processes in tip growing cells which are well studied in root hairs (Rounds and Bezanilla, 2013), but are 447 also prevalent in fast growing tissues such as the emerging inflorescence stem and 448 flowers (Chen et al., 2009). On the other hand, autophagy, a process which also 449 involves the formation of lytic compartments and vesicle trafficking for the degradation 450 451 of cytoplasmic material is known to play an important role in senescence in leaves and other tissues (Wojciechowska et al., 2018). 452

453 The Proteome in Establishment and Maintenance of Photosynthesis

454 Clusters 11 and 8 (Supplemental tables 11 and 12 respectively) were almost exclusively comprised of proteins annotated as chloroplast localized (297 of 326 nuclear plus 18 455 plastid encoded and 223 of 264 nuclear plus 20 plastid encoded, respectively) and 456 essential to photosynthesis by DAVID GO. Proteins were absent in roots and mature 457 458 brown siliques and predominant in green tissues, primarily leaves and cauline leaves. The abundance of cluster 11 proteins peaked in cotyledons of 7 and 10 day old 459 seedlings and young cauline leaves and declined in these tissues in the course of 460 ageing, whereas cluster 8 proteins were consistently abundant throughout the plant 461 lifecycle, in rosette leaves with a slight maximum at 40 days and declining in 462 senescence (Figure 3 A and B). 463



464

465 Figure 3 Proteome remodeling in establishment and maintenance of photosynthesis as well as in degradation of photosynthesis 466 apparatus and senescence in leaves. A. FCM cluster 11. Right panel: Physical / functional protein interaction networks generated 467 with the STRING database using all proteins assigned to the GOTERM category Chloroplast (Supplemental Table 11) as input set. 468 Various biochemical pathways in pigment and photosynthesis related protein synthesis and chloroplast biogenesis are color 469 coded and indicated. B. FCM cluster 8. Right panel: Physical / functional protein interaction networks generated with the STRING 470 database using all proteins assigned to the GOTERM categories Stroma, Thylakoid, Carbon metabolism (C Metabolism) and 471 Photosynthesis as input sets (Supple, emtal Table 12). Light independent reactions are highlighted in green (Calvin-Benson cycle 472 dark, photorespiration light), light dependent reactions in orange (photosystem II PSII) and blue (photosystem I PSI). C. FCM Cluster 10. Right panel Cumulative abundance (PQI given is raw #PSMs) of core abscission signaling module proteins in leaves, 473 474 cauline leaves, flowers and siliques.

475 The proteins in cluster 11 primarily function in chloroplast biogenesis and the

establishment of the functional photosynthetic machinery. A number of central players in

these molecular events were identified including HEMA1 and GUN5 (as well as GUN4), 477 (Liu et al., 2017) that also represent hubs in STRING generated networks of cluster 478 479 member proteins interacting in tetrapyrrole and chlorophyll synthesis, respectively (Figure 3 A, complete annotated network Supplemental figure 8). Other cluster proteins 480 were interaction partners in carotenoid synthesis. Around 30 cluster proteins were 481 classified as ribosomal proteins or related to translation by DAVID GO which was 482 reflected by a highly connected interaction network composed of chloroplast specific 483 ribosomal proteins, ribosome associated proteins or RBFs, the gene knockouts of many 484 of which have embryo lethal phenotypes. CPN60 and CPN10 class molecular 485 chaperones formed another cluster of interactors. Protein import from the cytosol is 486 essential to chloroplast biogenesis and constituents of the Toc/Tic complex including 487 488 TOC33, which is known to be most strongly expressed in young seedlings were cluster members (Waters and Langdale, 2009). The two proteins forming the inner division ring 489 mediating chloroplast division, FtsZ1 and FtsZ2, were also cluster members (Waters 490 and Langdale, 2009). DAVID GO mapped the cluster proteins to several plastid 491 492 structures thereby giving a possible inkling of their localization within the chloroplast.

Cluster 8 mostly contained proteins prevalent in mature chloroplast directly related to 493 494 photosynthesis. Cluster members constituted extensive parts of both the light dependent and independent reactions. Regarding the former, most components of the 495 oxygen evolving photosystem II (PSII) and photosystem I (PSI), providing the final 496 strong reducing potential, were identified, many of them being close interactors (Figure 497 498 3 B). Components of the electron transferring Cytb₆f complex and the thylakoid ATP synthase CF₁ were also present. Regarding carbon fixation, nearly all components of 499 the Calvin-Benson cycle and many of the proteins involved in photorespiration were 500 mapped as functional interaction partners (Figure 3 B). Furthermore, functional 501 interaction mapping could resolve organelle specificity of the individual reactions in 502 photorespiration. Interaction networks produced from GO category stroma and carbon 503 metabolism and from GO category thylakoid and photosynthesis protein sets were 504 specific and showed extensive overlap attesting to the quality of protein extraction from 505 the chloroplast compartments (full annotated networks Supplemental figure 9 to 12). 506

507 The Proteome in Senescence

508 Cluster 10 contained 241 proteins whose abundance increased substantially during the 509 course of plant life and peaked in the latest developmental stage, i.e. during leaf and 510 flower senescence and fruit ripening, predominantly in rosette leaves but also in cauline 511 leaves, stem and flowers (Supplemental table 13). They were not abundant in young 512 tissues (Figure 3 C).

513 Senescence is a controlled developmental process that entails disassembly of the

514 photosynthetic apparatus in leaves for the purpose of nutrient remobilization and

resource allocation to fruit ripening in flower petals (Bleecker and Patterson, 1997).

516 Conversely, numerous proteins involved in chlorophyll and carotenoid degradation were

found including Pheophorbide a oxidase that is the key enzyme in the formation of

518 primary fluorescent chlorophyll catabolites (pFCCs). It has been shown that

nonfluorescent dioxobilin-type chlorophyll catabolites (NDCCs) represent the major end-

520 products of chlorophyll catabolism as opposed to NCCs and that the Cytochrome P450

521 monooxygenase CYP89A9 is responsible for their accumulation (Christ et al., 2013).

522 This protein was also a cluster member as well as 13 other CYPs. Twelve of the 14 total

523 CYPs belonged to the CYP71 clan suggesting a broader role for these proteins going

524 beyond CYP89A9.

525 During senescence large amounts of ROS are produced which must be controlled so as

not to lead to tissue damage and premature cell death (Rogers and Munne-Bosch,

2016). Many cluster proteins were involved in oxidative-reductive processes and ROS

scavenging including the centrally important cytosolic ascorbate peroxidase APX6.

529 More interestingly, a significant group of cluster proteins were kinases. Six of these

530 (CRK7, CRK8, CRK10; CRK14, CRK21, CRK41) were Cysteine-rich Receptor-like

531 Kinases (CRKs) that have two extracellular DUF26 domains that each contain 4

conserved cysteine residues. Next to the programmed loss of redox control leading to

ROS accumulation and ultimately cell death at the later stages of senescence, ROS

may also play an important role in signaling, mediating genetic reprogramming during

senescence (Breeze et al., 2011), the mechanisms of which however are largely

unexplored. The CRK cysteine thiol groups will likely be sensitive to the redox state,

potentially implicating these proteins as ROS sensors and ROS signaling initiators in
 senescence.

Abscission of floral organs after fertilization is another developmental process that 539 occurs late in the Arabidopsis life cycle. Most of the proteins of the canonical abscission 540 signaling module (Meng et al., 2016) including the receptor like protein kinases HAESA 541 (HAE), HAESA-like 2 (HSL2), and EVERSHED (EVR/SOBIR1), the co-receptor 542 SOMATIC EMBRYOGENESIS RECEPTOR-LIKE KINASE 4 (SERK4) and MKK4 and 543 MPK3, the mitogen associated protein kinase cascade downstream of the HAE receptor 544 complex were cluster members. The proteins accumulated in flowers and later siliques 545 546 as development progressed (Figure 3 C right panel). Interestingly, their abundance also increased in leaves and cauline leaves although leaf abscission is not developmentally 547 548 timed suggesting an unknown function of this abscission signaling module.

549 Proteome Remodeling in Steady State PTI

550 In addition to tissue proteomes and their altercation in the course of development, we were interested in assaying more rapid changes in proteome architecture such as 551 determined by steady state shifts from ordinary growth to immunity. Seven and ten day 552 old seedlings grown in liquid culture were treated with a final concentration of 1 µM flg22 553 in the medium for 16 hours. Flg22 is the 22 amino acid N-terminal epitope of flagellin 554 555 and elicits pattern triggered immunity (PTI) downstream of the receptor like kinase (LRR-RLK) FLAGELLIN-SENSITIVE 2 (FLS2). Deep proteomics measurement of the 556 untreated and treated samples identified 8344 proteins in all (Supplemental table 14). 557 HCL showed that the abundance of 1774 proteins increased whereas the abundance of 558 559 915 decreased in both samples as a result of flg22 exposure (Supplemental table 15). 560 These proteins were categorized by mapping them to a self-constructed model of PTI

using the MapMan software (Supplemental data for details, Supplemental table 16,
 Supplemental figure 13). The result was a comprehensive picture of proteome
 remodeling in PTI showing extensive changes in protein abundance in most of the
 major perception, signaling and response pathways. Dynamics of the flg22 receptor
 complex, including decreased abundance of FLS2 as a result of internalization and
 degradation (Robatzek et al., 2006) and a host of other RLKs were quantified. Vesicle

trafficking and transport proteins including exocyst and SNARE complex members 567 increased in abundance along with proteins involved in the early respiratory burst. 568 569 primarily RESPIRATORY BURST OXIDASE HOMOLOGUE D (RBOHD). All of the 570 components of both MAPK signaling cascades central to plant immunity, MKK4/5-MPK3/6/11 and MKK2-MPK4 (Bigeard et al., 2015) were measured and also showed a 571 572 slight increase in their abundance. The same goes for the calcium-dependent protein kinases (CDPKs) integral to PTI signaling, CPKs 4, 5 and 6 (Boudsocq et al., 2010), as 573 well as a host of other CDPKs, calmodulin (CAM) and CAM binding proteins and 574 calcineurin (Supplemental table 17). Interestingly several proteins showing changes in 575 their abundance upon PAMP stimulus were mapped to other avenues of plant immunity 576 such as effector triggered immunity (ETI) and programmed cell death (PCD) and 577

578 systemic acquired resistance (SAR).

We investigated proteome plasticity in more detail beginning with the category hormone 579 580 signaling and branching out from it to produce a proteomics model of phytohormone activity in PTI (Figure 4 Supplemental table 18). The deep proteomics results were 581 582 complemented by measurements of the plant hormones themselves, amino acids and secondary metabolites after 16 hours of flg22 treatment. Furthermore, we undertook a 583 584 retention time scheduled, parallel reaction monitoring (PRM) based targeted proteomics study that allowed accurate quantification and inference of statistical significance of fold 585 changes of 52 model proteins in three sample pools of 10 day old Arabidopsis seedlings 586 again following 16 hours of flg22 treatment (Figure 5, Supplemental Table 19). The 587 588 PRM based estimates of protein abundance fold changes were in full agreement with the deep proteomics quantification highlighting the latter's accuracy and power of the 589 deep proteomics strategy in general. 590

The abundance of 43 proteins playing roles in photosynthesis decreased slightly after
16 hours of exposure to flg22 (Supplemental table 20). Fifteen of these were assigned
to the photosynthesis light reaction and 6 of these in turn to photosystem II by the
MapMan software (MapMan bins PS.lightreaction and PS.lightreaction.photosystem



595 II.LHC-II respectively, Supplemental table 20). Both bin assignments were statistically

596

597 Figure 4 Proteomics model of phytohormone activity in PTI. The figure shows the proteins quantified in the jasmonate (blue), 598 tryptophan and IAA (green), secondary defense compounds/indolic glucosinolate (red) biosynthesis pathways as well as IAA and 599 JA signaling pathways (purple). Note all or nearly all components of these biochemical pathways were measured and quantified 600 in the deep proteomics study. Proteins pertaining to ET and SA synthesis as well as reversible hormone conjugation (GH3s) and 601 modification are also shown. Protein with names in black were measured; bar charts next to protein names indicate relative 602 changes in abundance after flg22 treatment; values are the sum of z-score transformed raw #PSMs in flg22 treated samples. 603 Proteins in grey were not detected. Red nucleotide sequence indicates an ARF1 binding site in the cognate genes promotor 604 region, blue a MYC2 binding site. Solid arrows indicate direct functional interaction or immediately neighboring steps in 605 biochemical pathways. Dashed arrows indicate more distal interactions. Phytohormone, tryptophan and indole glucosinolate 606 abundance 16 hours after flg22 treatment are also shown.

- significant (Benjamini Hochberg corrected p-values < 0.01) indicating these categories
- were enriched among the 43 proteins. Other proteins were assigned to the carbon
- reactions, more specifically the Calvin-Benson cycle (Supplemental table 20). The
- relatively small decrease in abundance of a set of these proteins was corroborated by
- small yet significant fold changes (maximum decrease -0.6 log₂ fold change) in the PRM
- study (Supplemental figure 14 and Supplemental table 19). These results suggest that
- 613 photosynthesis is inhibited upon PAMP perception and conversely in PTI.



614

Figure 5 Targeted PRM based quantification of model proteins. Framework is in analogy to figure 4. Bars represent log2 fold changes of protein abundance after 16 hours of flg22 exposure (1 μ M concentration in medium) estimated by area under the curve label free protein quantification index (PQI) of the 6 most intense product ions from MS2 spectra of targeted proteotypic peptides. Bars represent median PQI of all quantified proteotypic peptides for a given protein in 9 measurements (3 biological replicates each measured 3 times). Standard error is indicated. Star indicates significance α =0.05 if fold change of at least one of the quantified peptides was significant.

- The abundance of 93 proteins playing roles in plant primary metabolism, especially
- carbohydrate metabolism, increased upon exposure to flg22. 22 were categorized as
- 623 pertaining to glycolysis (MapMan bin glycolysis), 19 to the TCA cycle (MapMan bins
- TCA / org. transformation.TCA and TCA / org. transformation.other organic acid
- transformations) and 6 to major CHO metabolism (mapMan bin major CHO
- 626 metabolism.degradation.starch), among others by the MapMan software. Increases in
- abundance were more pronounced than the downregulation of photosynthesis related
- proteins and log₂ fold changes up to 1.6 were quantified and significant in the PRM
- study (Supplemental figure 14 and Supplemental table 19).
- 630 Jasmonate and Salicylic Acid Cross Talk
- 631 Sixteen hours of flg22 exposure led to an increase in the abundance of both salicylic
- acid (SA) synthesis pathway proteins, primarily ISOCHORISMATE SYNTHESIS 1

- 633 (ICS1) but also upstream transcription factors TEOSINTE
- 634 BRANCHED1/CYCLOIDEA/PCF 8 and 22 (TCP8, TCP22) (Wang et al., 2015) and
- 635 NTM1-Like 9 (NTL9) (Zheng et al., 2015) and free SA levels (nearly 3-fold).

Jasmonate (JA) and jasmonate-isoleucine (JA-IIe), its bioactive conjugate, levels were

- both low, although the former increased around 2-fold, the latter showed no significant
- change and absolute levels of both were in the range of a few ng/g FW. Concurrently,
- (+)-12-oxo-phytodienoic acid (OPDA) levels increased from 1.6 to 7 μ g/g FW so were
- 640 high and elevated more than 4-fold following flg22 treatment. OPDA is a primary JA
- 641 precursor synthesized from allene oxides by ALLENE OXIDE CYCLASE 1 to 4 (AOC1
- to 4). ACC, the ethylene precursor and a proxy for the phytohormone's abundance
- 643 increased more than 5-fold.

Controversially, the deep proteomics measurements showed elevated protein amounts 644 of all components of the JA biosynthesis pathway, corroborated as significant in the 645 PRM study. Moreover the core JA receptor complex / signaling proteins CORONATINE 646 INSENSITIVE 1 (COI1), TOPLESS RELATED PROTEINS 1 to 3 (TPR1 to 3) and S 647 PHASE KINASE-ASSOCIATED PROTEIN 1 (SKP1) and CULLIN 1 (CUL1) proteins, 648 members of the E3 ubiquitin ligase SCF complex, were all significantly increased in 649 abundance, albeit slightly. The abundance of a large number of GRETCHEN HAGEN 650 651 (GH) enzymes, the amidohydrolase IAA-ALA RESISTANT3 (IAR3) and JASMONATE-652 INDUCED OXYGENASE 2 (JOX2) also increased following exposure to the PAMP, the latter two significantly around 5-fold. These enzymes are all involved in conjugating or 653 deconjugating phytohormones, specifically JA and auxin (IAA) to amino acids or small 654 molecules or hydroxylating them (JOX2) thereby modulating their sub-cellular location 655 656 and/or rendering them active or inactive.

The proteomics results imply, that flg22 induced PTI prioritizes SA over JA signaling. On the other hand, the induction of both JA synthesis and signaling pathways on the protein level and the highly elevated abundance of both IAR3 and JOX2, which deconjugate JA-IIe to JA and hydroxylate the latter to 12-OH-JA, together with the low JA and JA-IIe levels themselves, suggests that phytohormone conjugation may play a pivotal role in this context and a role for JA in PTI.

663 Auxin/IAA homeostasis

The detection of numerous GH proteins known to conjugate auxin/IAA (GH3.2,

665 GH3.5/WES1 and GH3.17/VAS2) and others prompted us to investigate the role of this

666 phytohormone in PTI. The levels of GH3.14, GH3.15 and GH3.17 increased upon flg22

exposure, the first significantly 1.2 fold. GH3.15 (AT5G13370) function in conjugating

the IAA precursor IBA has just recently been elucidated (Sherp et al., 2018). GH3.14

(AT5G13360), the neighboring gene, does not shown any significant sequence

670 homology indicating a potentially different uncharacterized function in PTI.

Auxin/IAA is synthesized by tryptophan dependent and independent pathways. Both

672 deep and targeted proteomics results showed significant substantial increase in the

abundance of all proteins in the tryptophan biosynthesis pathway and tryptophan levels

were also elevated almost 2-fold upon 16 hours of PAMP treatment. Tryptophan

channels into a host of defense related secondary metabolite synthesis pathways

676 particularly indole glucosinolates (IGs), the protein abundances of which all increased

highly (2-fold or more) and significantly. IG levels themselves, indol-3-

4- ylmethylglucosinolate (I3M), 1-methoxy-indol-3-ylmethylglucosinolate (1MOI3M) and 4-

679 methoxy-indol-3-ylmethylglucosinolate (4MOI3M), did not change significantly.

680 Presumably they were hydrolyzed by the mirosinase PENETRATION 2 (PEN2), whose

abundance, along with CYP81F2, which showed an 8.2 fold change in abundance,

increased 2.3 fold in response to flg22, to play a role in callose deposition (Clay et al.,

683 2009).

684 Several proteins potentially involved in auxin/IAA synthesis pathways were identified.

The abundance of ALDEHYDE OXIDASE 1 (AAO1) increased nearly 2-fold and that of

the NITRILASES 1 and 4, 1.2 and 23 fold, respectively. Auxin/IAA levels however

decreased slightly (0.75 fold) but significantly (p-value 0.051, two sample t-test equal

variance, n=5, α =0.1). AAO1 has recently been shown to play a role in converting the

I3M downstream hydrolysis product indole-3-carbaldehyde (ICHO) to indole-3-

carboxylic acid (ICOOH) in the abiotic stress response (Muller et al., 2019).

691 Pseudomonas syringae pv tomato DC3000 (Pst DC3000) infection induced a strong

transcriptional response of the nitrilases NIT2, NIT3 and NIT4 which was corroborated

by increased protein abundance in the case of NIT2. It was postulated that NIT2 is
involved in IAA signaling in R gene mediated resistance and defense against biotrophic
pathogens (Choi du et al., 2016). The more than 20-fold increase in the abundance of
NIT4 that we measured suggests an unknown function of this protein in basal resistance
to biotrophic pathogens.

While auxin/IAA levels did not change substantially even 16 hours after PTI induction, 698 the abundance of polar auxin transporters was markedly affected. Particularly, the 699 abundance of PIN-FORMED 3 (PIN3), a polar efflux carrier with well documented 700 functions in lateral root growth and tropism decreased significantly 2-fold. A similar 701 decrease was apparent for PIN-FORMED 7 (PIN7) in the deep proteomics results which 702 is also known to affect lateral root growth. Both PIN 3 and 7 have important roles in 703 704 establishing local auxin concentration maxima (Jang et al., 2018). The abundance of AUXIN RESISTANT 4 (AXR4) an ER protein that is responsible specifically for the 705 706 distribution and localization of the influx carrier AUXIN RESISTANT 1 (AUX1) was also elevated slightly. PIN3, PIN7 and AUX1 are the primary components of the polar auxin 707 708 transport system so our results indicate active, cell-to-cell auxin transport is perturbed in steady sate PTI. 709

710 Chronology and Model of Phytohormones in PTI

711 It has been documented that JA, IAA and SA act in chronological order in the establishment of SAR and that JA plays an early role (Truman et al., 2010a). Therefore, 712 we measured hormone levels 1, 3 and 16 hours and OPR3 (as a marker for JA 713 synthesis), JOX2 and COI1 and TPR1 (as markers for JA signaling) transcript levels 3 714 715 and 16 hours after flg22 exposure (Supplemental tables 21 and 22 and Figure 6). SA accumulated after 1h and levels remained elevated until 16h after exposure. JA and JA-716 717 Ile levels remained basal throughout the entire time course. OPDA levels however 718 increased substantially already 1 hour after PAMP treatment and remained elevated over time and OPR3 transcript abundance increased significantly 5.7 fold 3 hours after 719 720 PAMP exposure and decreased to basal levels at the 16 hour time point. Together this 721 suggests that the JA biosynthesis pathway is induced early in PTI. JOX2 transcripts



were elevated substantially and significantly 3 hours and remained high up to 16 hours

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Figure 6 Absolute quantification of phytohormones without and 1, 3 and 16 hours after flg22 exposure (1μM concentration in medium) in Col-0 Wt and myc234 mutant backgrounds using GC-MS. Bars represent mean of three biological replicates;
 standard error is given. qPCR based transcript abundance profiling of selected genes without and 3 and 6 hours after flg22
 exposure (1μM concentration in medium). Bars represent mean of 6 measurements (3 biological replicates each measured twice), standard error is given.

- after exposure to flg22 again reconciling initiation of JA synthesis with low JA levels by
- way of hydroxylation already at the early stages of biotrophic pathogen defense. The
- abundance of COI1 and TPR1 transcripts did not change markedly over time
- suggesting little or no JA signaling in the absence of JA-IIe itself.
- 733 Finally, as we uncovered several potentially new aspects of JA synthesis and
- homeostasis in PTI, we measured the abundance of the phytohormones at the same

735 time points following flg22 treatment in the myc234 background. JASMONATE INSENSITIVE 1 (MYC2) is one of the most important transcription factors in JA 736 737 signaling downstream of COI1 with a host of diverse regulatory functions and the triple knockout exhibits essentially no functional redundancy. SA hyperaccumulated in the 738 triple KO under standard growth conditions as described previously (Nickstadt et al., 739 2004). Its abundance decreased markedly 0.54- and 0.78-fold 1 and 3 hours, but 740 increased 16 hours after flg22 perception, suggesting that JA signaling plays a role in 741 SA accumulation in PTI and that it is an early one. 742

Strikingly, JA and JA-Ile levels both greatly increased (maximum JA-Ile 12.1-fold 1 hour, 743 744 JA 13.5-fold 3 hours after flg22 exposure) throughout the time course. OPDA also increased in the mutant, its profile mirroring that of JA in the wild type, reaching a 745 746 maximum increase of 4-fold three hours after flg22 perception. This shows induction of JA biosynthesis pathway and synthesis of JA intermediates independently of MYC2. 747 748 The highly elevated JA levels and JOX2 transcript abundance 3 hours post PAMP treatment prompted us to investigate the IAR3 and JOX2 promotor regions for MYC2 749 750 binding sites and indeed two and three were identified as the top ranking hits repsectively. This supports a hypothesis wherein JOX2 expression is under the control 751 752 of MYC2 and wherein JA is synthesized but continuously depleted by JOX2 by way of hydroxylation in biotrophic pathogen defense. Our combined measurements represent 753 754 strong evidence of a MYC2 dependent negative feedback loop controlling JA-Ile and JA levels in PTI via deconjugation and hydroxylation by IAR3 and JOX2 (Figure 7). 755

The other phytohormones showed similar levels over time in both the wild type and the 756 mutant. The amount of free IAA did not change dramatically, increasing slightly at 757 758 earlier time points and decreasing as stated above at the 16 hour time point presumably 759 because steady state PTI is achieved. ACC/ET abundance increased already 1 hour after exposure of the seedlings to the PAMP and remained elevated. ABA decreased 3 760 761 hours after flg22 treatment and remained low throughout the time course, presumably because of its known antagonism with SA (Cao et al., 2011). Additionally we measured 762 763 changes in the levels of the 20 proteinogenic amino acids and some others and found

alanine, glycine, tryptophane and taurine to increase during the course of flg22



766 *Figure 7 A MYC2 dependent negative feed-back loop controls JA-Ile and JA levels in equilibrium with JA synthesis by way of* **767** *deconjugation and hydroxylation via IAR3 and JOX2 in immunity to biotrophic pathogens.*

resposure (Supplemental figure 15). The abundance of leucine, isoleucine, lysine,

proline and ornithine decreased over the time of exposure.

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772 Discussion

773 Deep Proteomics Study of the Arabidopsis thaliana Proteome

In this report we describe the proteome biology of the model plant Arabidopsis thaliana 774 comprehensively. We elucidated the proteome landscapes of the plant and tissues 775 776 throughout its lifecycle as well as in the immune response to PAMP in unprecedented resolution and detail. Importantly we placed an emphasis on remodeling of proteome 777 architecture between these diverse biological scenarios to extrapolate genome wide 778 protein co-regulation and function. This approach has just recently been documented in 779 humans (Kustatscher et al., 2019) and draws its power from deep coverage of the 780 proteome as we have done here, quantifying close to 16,000 Arabidopsis proteins. In-781 782 depth sampling allowed quantitative measurement of all of the components of entire biochemical pathways such as JA synthesis in PTI, protein complex components such 783 784 as RBF complexes and protein co-expression of genes in local neighborhoods such as 785 cysteine rich kinases (CRKs), facilitating confident inference of protein abundance co-786 regulation and functional and temporal connections for instance in seed development

and dormancy, the establishment, maintenance and deconstruction of the
photosynthetic machinery and in senescence and abscission. With this we alleviated an
inherent weakness of most previous plant proteomics studies that were underpowered,
quantifying only a few thousand proteins if at all. To do this we developed and optimized
a strategy that in our hands allowed quantification of more than 9,000 proteins from a
sample in 3.5 days, a duration that is acceptable for medium scale proteomics studies.

Despite our unprecedented sampling depth, we did not delve much deeper into the 793 entire Arabidopsis proteome than Baerenfaller and co-workers (or as a matter of fact the 794 other deep proteomics studies in plants). In their classic study published more than ten 795 796 years ago individual sampling depth was modest however they utilized more than 1,300 low resolution LC-MS measurements to quantify nearly 15,000 proteins. Both studies 797 798 show a similar impact of transcript abundance on detection of cognate proteins (Figure 2C Baerenfaller et al, Figure 1C here) however we show protein size has a more 799 800 pronounced effect presumably because small proteins will generate less tryptic peptides amenable to MS detection. Also transcription factors and nuclear proteins whose 801 802 expression may be highly transient and biological context specific were conspicuously 803 overrepresented in the missing proteome (Supplemental figure 4). Capturing these 804 proteins could require exhaustive sampling of different organs at all developmental 805 stages (imagine par example the diversity of the 12 stages of organ development in the floral bud (Bowman et al., 1989; Smyth et al., 1990)) and environmental interactions and 806 even then may escape detection. Indeed, the more than 8,000 proteins identified in the 807 808 context of flg22 treatment did not substantially increase the amount of total proteins 809 previously identified in the plant tissues (Figure 1F) Also proteomics ruler method allowed us to quantify proteins with less than 100 copies per cell, which is in general 810 agreement with LOQs of modern large scale proteomics studies in humans (Bekker-811 Jensen et al., 2017). This suggests that even more comprehensive sampling of the 812 813 Arabidopsis thaliana proteome may be confounding with current technologies.

Next to the prevalence of known artificial and biologically occurring modifications, amino acid substitutions were found to be common. Such exchanges are generally caused by ribosome infidelity, have a number of biological implications and have been reported

previously in human cell lines (Chick et al., 2015), human tissue (Bagwan et al., 2018) 817 and in *E.coli* and yeast in detail (Mordret et al., 2019). The authors of the latter study 818 819 define exchanges as near cognate errors (NeCE) when two of the three bases between 820 error bearing origin and destination codon match and as non-cognate error (NoCE) when there is no such match between possible error bearing origin and destination 821 codons. Interestingly, 5 of the 8 substitutions we found to be abundant in Arabidopsis 822 (Asn->Met, Ala->Asp, Pro->Glu, Asp->Asn, Glu->Gln) were also abundant in yeast but 823 not *E.coli*, and three (Ala->Asp, Asp->Asn, Glu->Gln) were NeCEs suggesting 824 conserved patterns of translational error in eukaryotes, a possible method of generating 825 random phenotype variants in genetically identical organisms (Mordret et al., 2019). 826 This phenomenon may also highlight the limitations of searching protein databases 827 828 derived from reference genomes with proteomics data.

829 Root and Seed Proteomes

The root is a highly specialized, plastic tissue that confers structural stability to the 830 above ground portion of the plant and is responsible for the uptake of nutrients and 831 water from the soil. Thus it exudes a number of metabolites, peptides and proteins that 832 allow it to interact with the rhizobiome. The function of the root specific proteins in 833 cluster 3 are well documented. Interestingly the transcription factors and proteins that 834 pertain to hormone signaling in the developing root (Paez Valencia et al., 2016) were 835 836 not overly prominent among the set of cluster proteins. Comparative evaluation of different deep proteomics studies as we have done here can be useful in defining the 837 core set of proteins ubiquitously expressed in a cell type, tissue or organism. 838

839 The proteins in cluster 7 are very specific to the seed proteome. They were exclusively or significantly highly abundant in siliques only and not ubiquitously abundant in other 840 841 tissues. These include many of the proteins involved in ABA dependent regulation of 842 seed development. The amount of ABI5, a positive regulator of ABA signaling is post translationally controlled by AFP1, presumably promoting its degradation by the 843 proteasome (Lynch et al., 2017). The high ABI5 to AFP1 ratio measured in post mature 844 seeds indicates high ABA levels and ABI5 activity also in the induction and maintenance 845 of dormancy. ABA and DOG1 pathways converge on PP2C phosphatases such as 846

AHG1 to suppress germination (Nee et al., 2017). DOG1 and the PP2Cs RDO5 and 847 AHG1 were some of the most abundant measured proteins suggesting a possibly 848 849 dominant role in seed dormancy by this pathway over ABA signaling. In addition two 850 other PP2Cs (AT3G15260 and AT4G31860 of the F and I clades respectively) previously not known to be involved in seed development also accumulated to very high 851 levels implicating them in the same processes. The abundance of RD26/ANAC72 852 (AT4G27410) and its two closest homologs ANAC19 (AT1G52890) and ANAC03 853 (AT3G15500) as well as two other NAC transcription factors, ANAC02 (AT3G15510) 854 and ANAC14 (AT1G33060) increased specifically and significantly in seeds particularly 855 in post mature seeds (brown siliques). RD26 and homologs have been shown to be 856 ABA responsive (Fujita et al., 2004) and to be expressed ubiquitously in Arabidopsis 857 858 vegetative tissues in response to drought and salt stress (Tran et al., 2004; Nakashima et al., 2012) and coronatine (Zheng et al., 2012). They are also known to play a role in 859 jasmonate (Bu et al., 2008) and brassinosteroid signaling (Ye et al., 2017) as well as 860 leaf senescence (Takasaki et al., 2015). Given their role in promoting drought tolerance, 861 862 their seed specific accumulation in the absence of stress may indicate a previously unknown role in the onset and maintenance of desiccation tolerance. No seed specific 863 864 or reproductive phenotype has been reported for the respective single NAC gene knockout mutants possibly reflecting functional redundancy and to our knowledge no 865 866 multiple knockout mutants have been produced.

867 Ribosomal Proteins in Development

In our study we identified a large number of RPs and RBFs that all show the same 868 tissue and developmentally specific pattern of protein abundance, increased in young 869 870 (seedling) roots, young stem and early flowers/floral buds. The deep proteome 871 coverage of our approach allowed us to identify entire RBF protein complexes, all of whose members had the same expression pattern. The RBF complexes were part of 872 873 the SSU-processome or the pre-60S pre-ribosomal particle and mutation of many of their constituent RBFs lead to aberrant gametophyte development. This suggests that 874 875 particularly these complexes regulate gametophyte development in early flowers in the context of ribosome biogenesis and translation, underscored by the their much more 876

conserved, significant expression pattern as opposed to ribosomal proteins in general. It 877 has been reported that the transcripts of several of these genes are especially abundant 878 879 in developing tissue such as young roots, stem and flowers (Missbach et al., 2013). Furthermore, the extensive physical interaction between RPs and RBFs with the protein 880 abundance pattern found here may indicate these particular isoforms assemble 881 882 ribosomes specific to young roots, young stem and early flowers/floral buds with possible function specific to these tissue states. Closer inspection of the protein 883 interactions may also bring to light new RBFs and RP/RBF complex conformations that 884 have not yet been described. Indeed, although the complexes detected here are well 885 described in yeast, much of what is known in *Arabidopsis* is by inference, so this study 886 presents evidence of their tissue specific expression on the protein level, particularly 887 888 pertaining to the CCT complex.

ABI5 is a positive regulator of ABA signaling that has been shown to repress the 889 890 flowering transition (Wang et al., 2013) next to its well documented function in the induction and maintenance of seed dormancy (Finkelstein et al., 2008). So far a handful 891 892 of proteins have been reported to target ABI5 to degradation and thus negatively impact the ABA response, including AFP1 (Lopez-Molina et al., 2003) and DWA2 (Lee et al., 893 894 2010). Both had distinct abundance patterns in this study. AFP1 was most abundant in ripe brown siliques (see above). DWA2 abundance was elevated in young roots, young 895 stem and early flowers/floral buds wherein it presumably mediates ABI5 degradation to 896 induce flowering. These results suggest different developmental and tissue specific 897 898 mechanisms post-translationally control ABI5 levels and ABA activity.

899 Vesicle Trafficking

Interestingly many of the membrane trafficking cluster 13 proteins were abundant in
both fast growing tissues, primarily roots but also stem and flowers and senescent
leaves. Vesicle trafficking is preeminent in the targeted deposition of new cell wall
material in the clear zone at the apex of tip growing cells as is the formation of an apical
actin structure. Autophagy, which also involves transport of cytoplasmic components in
membrane vesicles is important in senescence, both in counteracting premature cell
death and degradation of cellular structures such as the photosynthetic apparatus in

nutrient remobilization. Our results at least suggest, that some of the molecular playersin these distinct processes in different organs may be the same.

909 Photosynthesis

910 The tissue and developmental abundance profile of cluster 11 proteins implicates them in chloroplast biogenesis and indeed, large sets of interacting proteins as well as known 911 912 individual proteins central to its molecular processes were the major constituents of this 913 cluster. These processes are known to be most prevalent in the young plant. The proteins of the photosynthetic apparatus are abundant in green tissues in nearly all 914 developmental stages beginning in the cotyledons of the young seedling prior to 915 916 photoautotrophy. This can be seen in the abundance profile of the proteins in cluster 8 917 which decreases in leaves in the later senescent stages, concomitantly with an increase of proteins that facilitate disassembly of the apparatus and pigment degradation in 918 919 cluster 10.

920 Senescence and Abscission

921 As such it is no surprise that a large number of antioxidant proteins and proteins involved in redox regulation and ROS scavenging increased in abundance as ageing 922 923 progressed, peaking in senescence in leaves, cauline leaves and flowers/siligues. Chlorophyll and carotenoid catabolism are central aspects of leaf senescence and 924 925 CYP89A9 is important in this respect catalyzing the oxidative deformylation of FCCs to dioxibilin-type FDCCs and ultimately the accumulation of NDCCs (Christ et al., 2013). 926 927 CYP89A2 has been shown to be co-expressed with CYP89A9 (Obayashi et al., 2009) and we have observed this here for the cognate proteins. In addition to these two, 10 928 929 other CYP71 clan members had the same protein abundance pattern of cluster 10. This 930 makes it likely, that a larger number of CYPs play undescribed roles in chlorophyll catabolism, as was also speculated by Christ and co-workers. 931 We detected 6 CRKs showing the characteristic abundance pattern associated with 932 senescence. This RLK gene family consists of 44 members all in close proximity on 933

chromosome 4 with a host of different physiological functions (Wrzaczek et al., 2010;

Burdiak et al., 2015). Their extracellular regions have two conserved DUF domains

each with 4 cysteine residues as potential targets for thiol redox regulation. Members of 936 the gene family have been shown to be expressed on the transcript and protein level in 937 938 response to pathogen challenge, flg22 and SA, underpinning a potential role in immunity (Wrzaczek et al., 2010; Yadeta et al., 2017). Transcripts of five of the six 939 CRKs we identified were also significantly upregulated following ozone treatment 940 (Wrzaczek et al., 2010). Collectively these processes as well as senescence all involve 941 ROS, although the spatio-temporal mechanisms of ROS production, interaction and 942 signaling in the senescence syndrome are not well understood. A probable ROS 943 dependent role of CRK5 in cell death and senescence is documented (Burdiak et al., 944 2015). Our results further support an as of yet little known role of the CRKs in 945 senescence. Considering CRKs as ROS sensors and signaling molecules leads to the 946 947 question of chloroplast derived ROS production affecting the extra-cellular redox state akin to the oxidative burst upon pathogen perception. 948

949 Arabidopsis floral organ abscission as the culmination of petal senescence and nutrient remobilization for fruit ripening and ultimately release is a well-studied model of general 950 951 abscission processes (Patharkar and Walker, 2018). Here we detected most of the 952 components of the abscission signaling cascade increasing in abundance in flowers as 953 they age and set seeds in siliques following fertilization. Recently it has been shown that the same mechanisms also act in Arabidopsis cauline leaf abscission in response to 954 drought (Patharkar and Walker, 2016) and indeed it is common that plants shed their 955 leaves in response to various environmental stimuli. However, leaf abscission is not 956 957 known to be on a developmental clock and *Arabidopsis* does not abscise rosette leaves 958 (Stenvik et al., 2006), making it intriguing that we also found the abundance of the proteins to increase in both rosette and cauline leaves in the course of ageing and 959 960 senescence.

ABA plays a major role in fruit ripening (Jia et al., 2011) and the leaf senescence
syndrome (Song et al., 2016), yet its role in abscission is debated. ABA promotes
ethylene biosynthesis during the later stages of fruit ripening in tomato (Zhang et al.,
2009) and ethylene is known to be a regulator of abscission in *Arabidopsis* (Patterson
and Bleecker, 2004). The hormone is also a key factor in the response to drought

966 (Huang et al., 2008). HAE, HLS2 and IDA expression in Arabidopsis seedlings is

⁹⁶⁷ induced by ABA and to a lesser extent ethylene (eFP browser (Goda et al., 2008)). It is

possible, that the elevated ABA and ethylene levels in the course of senescence led to

the accumulation of the abscission signaling proteins in rosette and cauline leaves.

Therefore one may speculate on possible other yet unknown functions of this signaling

971 module outside of organ abscission.

972 Photosynthesis and Primary Metabolism in PTI

It is known that pathogen infection leads to inhibition of photosynthesis and it has been

shown that this is an active response of the plant to the invading pathogen.

Downregulation of a host of genes related to the light dependent reaction and

976 particularly photosystem II and parameters of photosynthetic activity has recently been

reported to be dependent on constitutive MPK3/MPK6 activation in ETI (Su et al., 2018).

The resulting increase in chloroplast localized ROS is hypothesized to support the

programmed cell death in the hypersensitive response (HR). Measurement of the

photosynthetic parameters upon exposure to 100 nM flg22 for up to 24 hours or

981 infiltration with a Pseudomonas syringae pv tomato DC3000 (Pst DC3000) strain that

982 lacks a type III secretion system to deliver effectors in this study indicated that

downregulation of photosynthesis does not occur in PTI, which feature only transient

984 MPK3/MPK6 activation. An older proteomics study however discloses downregulation of

some photosynthetic proteins and non-photochemical quenching (NPQ) 2 hours after

flg22 treatment (100 nM to 10 μ M) and decrease of electron flux through PSII upon 7

days of exposure, suggesting inhibition of photosynthesis may occur in some PTI

scenarios (Gohre et al., 2012). This is in line with our findings where plants were

exposed to 1 µM flg22 for sixteen hours. Conceivably higher flg22 doses may lead to

prolonged MPK3/MPK6 activation and photosynthetic inhibition also in PTI. As an

afterthought, the abundance of both MPK3 and MPK6 increased somewhat in the

discovery proteomics results (Supplemental table 14) upon PAMP exposure although

we did not measure phosphorylation levels.

Jasmonate and Salicylic Acid Cross Talk

The induction of PTI downstream of FLS2 binding flg22 is a commonly accepted model 995 of bacteria induced basal immunity in plants. As the majority of bacteria adopt biotrophic 996 997 lifestyles, it can also be considered a model of plant resistance to biotrophs although not 998 exclusively, as some bacteria, such as *Erwinia carotovora* are necrotophs. Also bacteria will be recognized by more than one pattern recognition receptor (PRR) such as EF-TU 999 RECEPTOR (EFR) and others in addition to FLS2, inducing multiple partially 1000 overlapping responses, so the flg22/FLS2 model may be overly specific and somewhat 1001 artificial. 1002

1003 The importance of SA and JA in resistance to biotrophs and necrotrophs respectively as 1004 well as their generally antagonistic modes of action are well documented (Pieterse et al., 2012). It has also been shown that both hormones interact and play roles in flg22 1005 1006 induced PTI (Denoux et al., 2008; Yi et al., 2014; Hillmer et al., 2017; Mine et al., 2017). Our results suggests flg22 induced PTI at the steady state of 16 hours after continuous 1007 1008 flg22 exposure prioritizes SA mediated defenses over JA mediated defenses because of dampening of JA and JA-Ile accumulation in culture grown seedlings. SA has been 1009 1010 shown to accumulate in and be essential for flg22 triggered defenses (Tsuda et al., 2008). The suppressive effect of SA on JA levels has been extensively explored and 1011 1012 shown to be NPR1 dependent in *Pst* DC3000 infected *Arabidopsis* plants (Spoel et al., 1013 2003) and integral to the trade-off between defense against biotrophic and necrotrophic pathogens (Spoel et al., 2007). However most of this interplay is downstream of JA 1014 synthesis at the level of inhibition of JA responsive gene transcription by transcriptional 1015 1016 regulators primarily WRKY70 and TGAs and ORA59 (Li et al., 2004; Leon-Reyes et al., 1017 2010; Shim et al., 2013; Van der Does et al., 2013; Zander et al., 2014). Here we propose a model wherein a MYC2 dependent negative feed-back loop in equilibrium 1018 1019 with JA synthesis controls JA-Ile and JA levels via clearance of the phytohormones by deconjugation and hydroxylation by IAR3 and JOX2 respectively. Four major arguments 1020 1021 for this model can be extrapolated from our measurements.

10221. JA and JA-Ile levels were very low and did not increase significantly following1023flg22 exposure in the wild type. However, the abundance of all of the proteins in1024the JA synthesis pathway and of the OPR3 transcript was significantly increased

1025after induction of PTI (16 and 3 hours respectively). It has been reported, that JA1026biosynthesis and signaling gene expression was upregulated in *Arabidopsis* at1027earlier time points (1h and 3h) following flg22 exposure (Denoux et al., 2008).1028The OPR3 transcript levels were again basal 16 hours post flg22 perception1029whereas protein abundance was still elevated, highlighting the importance of1030measuring protein levels directly.

- 1031 2. Absolute levels of the JA intermediate OPDA were very high (μ g/g FW) and 1032 increased more than 4-fold following treatment with the PAMP in both the wild 1033 type and the *myc234* background.
- 3. JA and JA-Ile levels were strongly elevated in the *myc234* mutant as opposed to
 the wild type. This indicates suppression of the phytohormone and its bioactive
 conjugate is dependent on MYC2 in PTI. Moreover, this result together with point
 3. above leads us to doubt that arrest of the JA synthesis pathway at the point of
 OPDA synthesis is the reason for the basal JA and JA-Ile levels. Also OPDA
 REDUCTASE 3 (OPR3) which reduces OPDA was the pathway component with
 the greatest fold change in abundance.
- 4. The amidohydrolase IAR3 and the 2OG oxygenase JOX2 were among the
 proteins with the most significantly increased fold change in abundance following
 16 hours of flg22 exposure. The same goes for the JOX2 transcript after three
 hours. MYC2 binding sites were identified as the top scoring motifs in *in silico*analysis of both the JOX2 and IAR3 promotor regions. These proteins
 deconjugate JA-Ile and hydroxylate free JA respectively, thereby inactivating the
 former as well as depleting both of them.

1048 These results lead us to hypothesize that deconjugation and subsequent modification of 1049 JA-Ile play an important role in control of the JA pathway in defense against biotrophs 1050 as has been shown against the necrotrophic fungus Botrytis cinerea (Caarls et al., 1051 2017). JA/JA-Ile clearance is presumably in some type of equilibrium with JA synthesis because it has been reported that the JA pathway is party to PTI induction and SA 1052 accumulation via EDS5 (Mine et al., 2017). The involvement of JA signaling in SA 1053 1054 accumulation is underscored in our data by the diminished SA levels in the myc234 1055 background at the earlier time points (1h and 3h after flg22 exposure). Thus our model

reconciles significant induction of the JA synthesis pathway proteins and known SA JA
antagonism at the level of transcriptional regulation with the low observed JA and JA-Ile
levels we measured and which were reported previously (Spoel et al., 2003).

1059 Auxin/IAA homeostasis

Auxins are among the most studied phytohormones and next to their preeminent role in 1060 1061 growth and development have established functions in plant immunity. Free IAA and 1062 IAA signaling both increase plant susceptibility to biotrophic pathogens (Kunkel and Harper, 2018) whereas IAA acts synergistically with JA in resistance to necrotrophic 1063 pathogens (Qi et al., 2012). Flg22 induces transcription of a microRNA miR393 that 1064 1065 targets the auxin receptor TRANSPORT INHIBITOR RESPONSE 1 (TIR1), thereby 1066 dampening auxin signaling (Navarro et al., 2006). Free IAA levels were not reported to 1067 change dramatically in response to flg22 (Navarro et al., 2006) nor to PStDC3000 1068 infection or SA (Qi NewPhytol 2012) and we also observed a slight but significant decrease after 16 hours of exposure, presumably by downregulation of the IAM 1069 pathway indicated by a decrease in the abundance of AMIDASE1 (AMI1) when steady 1070 1071 state PTI is reached. IAA is synthesized primarily from tryptophan by way of several biosynthesis pathways and tryptophan also channels into defense compound synthesis, 1072 especially indolic glucosinolates and camalexin. Thus branching points in 1073 tryptophan/IAA biosynthesis pathways, particularly indole-3-acetaldoxime (IAOx) 1074 1075 (Sugawara et al., 2009) may represent an important nexus in the growth defense trade 1076 off. Here we measured substantial, significant increase in protein abundance in the entire tryptophan as well as in defense compound biosynthesis pathways in response to 1077 flg22. The abundance of enzymes (AAO1, NIT1 and NIT4) in some alternative auxin 1078 1079 synthesis pathways with known secondary roles in defense also increased.

It is known that JA induces these pathways upon *Alternaria brassicicola* infection also
leading to increase in free IAA levels in the response to necrotrophic pathogens (Qi et
al., 2012). In our study JA and JA-IIe levels remained basal. However, we measured a
significant increase in the abundance of ethylene biosynthesis proteins ACC OXIDASE
2 (ACO2) and 1-AMINO-CYCLOPROPANE-1-CARBOXYLATE SYNTHASE 7 (ACS7)
and a significant, more than 5-fold increase of ET 16 hours after PAMP exposure. Clay

and co-workers reported expression of tryptophan and IG biosynthesis pathway
transcripts to be mediated by ET via myb transcription factors. These pathways play
important roles in defense against both biotrophic and necrotrophic pathogens and our
results suggest that they are induced independently by different phytohormones, ET
and JA respectively. These two hormones are generally synergistic posing new
questions about the role of JA in the flg22 response.

Auxins are unique among phytohormones in that they are transported directionally from 1092 1093 cell to cell by a polar transport system with many components (Friml, 2003). Indeed, the distribution and local IAA concentration have a profound impact on cellular processes 1094 1095 more so than absolute IAA levels or aspects of synthesis and catabolism (Teale et al., 1096 2006) and auxin transport mutants have been reported to be defective in mounting SAR 1097 (Truman et al., 2010b). Concurrently, we did not observe major changes in free IAA levels, however the abundance of two of the most well studied auxin efflux carriers, 1098 1099 PIN3 and PIN7, decreased and that of a protein directing the localization of the influx carrier AUX1 increased after flg22 exposure. This suggests that IAA transport and local 1100 IAA gradients also play an important role in the defense against biotrophic in addition to 1101 1102 necrotrophic pathogens (Qi et al., 2012).

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1105 Methods Summary

For detailed description of methods and optimization see Supplemental Methods and 1106 Data. In brief, proteins were extracted from Arabidopsis thaliana suspension cultures or 1107 soil grown plants and tissues with 4% SDS, separated into 5 bands by SDS-PAGE, in 1108 gel digested and measured on an Orbitrap Velos Pro mass spectrometer with a 1109 conventional DDA scan strategy using a 50 cm C18 liquid chromatography column and 1110 1111 an extended gradient of 9 hours. MS data was searched using the Mascot and Andromeda search engines. Search results were concatenated and peptides and 1112 proteins identified with global FDR thresholds of 0.01 in the Scaffold software. Proteins 1113 were quantified by way of PSM counting. For parallel reaction monitoring (PRM) 1114

proteins were extracted with 4% SDS, digested with an optimized FASP protocol and 1115 peptides measured on a QExactive Plus mass spectrometer. Up to three proteotypic 1116 peptide m/z per protein were targeted, peptides and proteins were identified using the 1117 Mascot search engine and data analysis and AUC quantification of 6 fragment ions was 1118 done using the Skyline software. Statistical significance of protein fold changes was 1119 1120 inferred at a threshold of 0.05 if one of the quantified peptides tested significant. qPCR was performed with the EvaGreen Kit from Bio&Sell according to manufacturer's 1121 1122 instructions, phytohormone and amino acid measurements according to (Ziegler et al., 2014). Bioinformatics analysis including gene ontology analysis and pathway 1123 visualization and mapping was done with DAVID, MapMan, STRING; PTM analysis with 1124 MSFragger. Multivariate data analysis was done in case of fuzzy c-means clustering 1125 1126 with the m-fuzz R package, hierarchical clustering and PCA with Perseus. Coexpression of proteins from neighboring genes was analyzed using an in-house 1127 1128 program programmed in R.

- 1129
- 1130
- 1131 Supplemental Figure Legends

Supplemental figure 1. Sampling of different *Arabidopsis thaliana* tissues and seedlings
at various stages in the plants life (in days). Samples correspond to Supplemental
methods and data tables 1 and 2.

Supplemental figure 2. Experiments to optimize GeLC-MS approach for deep 1135 proteomics analysis of plant tissues A. Total amount of extracted proteins and 1136 concentration of proteins in solution for various tissue amounts and tissue to buffer 1137 ratios using the SDS based protein extraction described in this work. For 250 mg / 750 1138 µl and 250 mg / 1000 µl mean values and standard deviations (error bars) of four 1139 1140 experiments are shown, for 500 mg / 500 μ l, 500 mg / 750 μ l and 250 mg / 500 μ l of eight and for 500 mg / 1000 µl of 18 experiments are shown. For phenol extraction (500 1141 mg / 1000 µl) mean values and standard deviations of six experiments are shown. B. 1142 Number of proteins identified (Mascot, no significance filters) in bands 1 through 5 and 1143

in total (cumulative sum of all bands) employing a 180 minute LC gradient when loading 1144 different amounts of phenol extracted proteins onto the SDS-PAGE as compared to 40 1145 µg of SDS extracted proteins. Mean values and standard errors of 2 experiments are 1146 shown for Phenol 40 µg C. Number of proteins identified (Mascot, no significance filters) 1147 in bands 1 through 5 and in total (cumulative sum of all bands) employing a 180 minute 1148 1149 LC gradient when separating 80 µg of SDS extracted proteins over different distances in SDS-PAGE. Mean values and standard errors of 2 experiments are shown for 2.5 cm 1150 1151 SDS-PAGE. D. Number of proteins identified (Mascot, no significance filters) in bands 1 through 5 and in total (cumulative sum of all bands) employing a 180 minute LC 1152 gradient when separating different amounts of SDS extracted proteins for 2.5 cm on 1153 SDS-PAGE. Mean values and standard errors of 3 experiments are shown for 40 µg 1154 1155 and 80 μ g of SDS extracted proteins. Only band 1 was measured for 10 μ g, 20 μ g and 240 µg of SDS extracted proteins. E. Number of proteins identified (Mascot, no 1156 1157 significance filters) in bands 1 through 5 and in total (cumulative sum of all bands) employing a 180 minute LC gradient when separating 80 µg of SDS extracted proteins 1158 1159 for 2.5 cm on 12% and 16% SDS-PAGE.

Supplemental figure 3. GeLC-MS approach for deep proteomics analysis of plant 1160 1161 tissues A. SDS extracted plant protein extract is separated on a 12% SDS-PAGE for a 1162 total distance of exactly 2.5 cm. High staining intensity bands are excised and analyzed separately leading to 5 protein fractions. Contaminants such as secondary metabolites, 1163 chromophoric compounds and leaf pigments (colored green) precede the proteins and 1164 1165 can be readily eliminated. B. Proteins are fractionated in the gel bands according to their size. Box plots show molecular weight (Mw) of all proteins identified with at least 1166 one unique peptide and a peptide FDR threshold (q-value) < 1% with the Mascot 1167 1168 software linked to Proteome Discoverer from 8 green tissue samples (LF7, LF10, LF40, LF66, LF73, JLF66, JLF73, JLF90; see Supplementary Methods and Data for 1169 1170 explanation) in the respective bands. Boxes represent the inter-guartile range (IQR) between the first and third quartile. Whiskers extend up to 1.5*IQR plus the third quartile 1171 and down to first quartile minus 1.5*IQR. C. Highly abundant leaf proteins involved in 1172 photosynthesis are fractionated. Mean number of peptide spectral matches (#PSMs) 1173 1174 identified as above for the 8 green tissues samples as above are plotted for each of the

indicated proteins as a percentile of the total PSMs of all proteins per band. Error bars 1175 indicate standard errors (SE). D. As C. for two root samples R7 and R10 (see 1176 Supplementary Methods and Data for explanation). E. Between 6,000 and 9,000 1177 proteins and 35,000 and 65,000 unique peptides were identified with a peptide and 1178 protein q-value < 1% with the Mascot and MaxQuant software integrated in the Scaffold 1179 1180 software per tissue sample. Box plots show protein and peptide numbers from all 23 samples; otherwise as above. F. Protein identification and quantification is repeatable. 1181 1182 Principal component analysis of two repeated GeLC-MS analyses of flowering stage 4 sample (FF4) and FF3 and JLF3 samples. The two principal components capture more 1183 than 96% of the total variance indicating the two dimensional subspace almost perfectly 1184 represents the relationship of the higher dimensional samples. G. Protein identification 1185 1186 and quantification is repeatable. Pair wise scatter plots of the four samples in F.

- 1187 Pearson correlation coefficients are given.
- 1188 Supplemental figure 4. DAVID Missing and Covered GO terms. Gene ontology
- classification of all proteins identified in the study, the "covered" Arabidopsis proteins
- and of all remaining protein coding genes for which no cognate peptide was identified,
- the "missing" proteome. The percentage of the respective proteome is given for each
- 1192 GO bin as well as the Benjamini corrected p-value for enrichment.
- 1193 Supplemental figure 5. Hierarchical cluster analysis of the deep proteomics
- measurements of sampled tissues. Row tree was generated using Spearman
- 1195 correlation, column tree using Pearson correlation. Values were z-score transformed so
- color gradient goes from -3 (dark blue) to 0 (mean value; black) to 4 (bright red).
- 1197 Supplemental figure 6. All cluster 4 proteins STRING physical interactions. Physical
- 1198 protein interaction networks produced with STRING database of all cluster 4 proteins.
- 1199 Thick blue edges highlight the UTP-B, t-UTP complexes, thick yellow edges the PeBoW
- 1200 complex. Red underlines indicate gene deletion has a developmental phenotype.
- 1201 Ribosomal core complexes are color coded.
- 1202 Supplemental figure 7. Significance of RBF complex protein co-expression. A 95%
- 1203 confidence interval was calculated for the mean expression value of RBF complex
- 1204 proteins in cluster 4 and all ribosomal proteins and ribosome constituents as

background. To assess the significance of RBF protein tissue and developmental 1205 specific expression according to the cluster 4 expression pattern, the number of times 1206 the expression value for samples R7, R10, SF66 and FF66 were outside the interval 1207 were counted for each protein and divided by the number of expression values of all 1208 samples that were outside of the interval. A ratio of 1 indicates only the 4 samples 1209 1210 mentioned above were significantly removed from the mean, indicating 95% confidence in protein expression specific to these tissues and developmental stages, as found for 1211 1212 many of the RBFs (red bars). Ribosomal proteins in blue had much lower ratios indicating expression values in more diverse tissues and developmental stages were 1213 significantly removed from the mean and therefore alternate and non-specific 1214 expression patterns. 1215

1216 Supplemental figure 8. Cluster 11 proteins Chloroplast. Complete physical / functional

1217 protein interaction network generated with the STRING database using all proteins

assigned to the GOTERM category chloroplast (Supplemental table 11) as input set.

1219 Proteins are labelled.

1220 Supplemental figure 9. Cluster 8 proteins Stroma. Complete physical / functional protein

interaction network generated with the STRING database using all proteins assigned to

the GOTERM category Stroma (Supplemental table 12) as input set. Proteins are

1223 labelled.

1224 Supplemental figure 10. Cluster 8 proteins Thylakoid. Complete physical / functional

1225 protein interaction network generated with the STRING database using all proteins

assigned to the GOTERM category Thylakoid (Supplemental table 12) as input set.

- 1227 Proteins are labelled.
- 1228 Supplemental figure 11. Cluster 8 proteins Carbon fixation. Complete physical /
- 1229 functional protein interaction network generated with the STRING database using all
- 1230 proteins assigned to the GOTERM category Carbon metabolism (C Metabolism)
- 1231 (Supplemental table 12) as input set. Proteins are labelled.

1232 Supplemental figure 12. Cluster 8 proteins Stroma. Complete physical / functional

1233 protein interaction network generated with the STRING database using all proteins

assigned to the GOTERM category Photosynthesis (Supplemental table 12) as inputset. Proteins are labelled.

Supplemental figure 13. Flg22 MapMan. The MapMan software was used to 1236 characterize and map proteins with changes in their abundance following 16 hours of 1237 1238 flg22 treatment to various avenues of plant immunity. The PTI pathway was drawn by 1239 ourselves and populated with a mapping concatenating the the MapMan Affymetrix mapping depicted on the pathway and related to PTI with corresponding GO terms from 1240 the TAIR GO slim ontology. Protein abundance is shown as the sum of z-score 1241 transformed spectral counts acquired for the respective proteins in measurements of 1242 1243 flg22 treated 7 and 10 day old liquid culture grown seedlings. Red indicates an increase in abundance blue a decrease (black is no change). Max and minimum values in the 1244 1245 color bar are 1.8 and – 1.8 respectively.

Supplemental figure 14. Targeted PRM based quantification of proteins involved in 1246 photosynthesis and primary metabolism. Bars represent log2 fold changes of protein 1247 abundance after 16 hours of flg22 exposure (1µM concentration in medium) estimated 1248 by area under the curve label free protein quantification index (PQI) of the 6 most 1249 intense product ions from MS2 spectra of targeted proteotypic peptides. Bars represent 1250 median PQI of all quantified proteotypic peptides for a given protein in 9 measurements 1251 1252 (3 biological replicates each measured 3 times). Standard error is indicated. Star 1253 indicates significance α =0.05 if fold change of at least one of the quantified peptides was significant. 1254

Supplemental figure 15. Absolute quantification of amino acids without and 1, 3 and 16
hours after flg22 exposure (1µM concentration in medium) in Col-0 Wt and *myc234*mutant backgrounds using GC-MS. Bars represent mean of three biological replicates;
standard error is given.

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1626 Supplemental figure 1



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1636 Supplemental figure 2



1647 Supplemental figure 3



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1650 Supplemental figure 4

Percent of Proteome 50.0 40.0 30.0 20.0 10.0 0.0 10.0 20.0 30.0 1.10E-42 Oxidation-reduction process 1.00E-03 Protein phosphorylation 4.90E-73 Cytoplasm 5.50E-275 Cytosol 1.20E-178 Chloroplast 5.20E-101 Chloroplast stroma 8.80E-90 Chloroplast envelope 1.40E-52 Membrane 1.40E-93 Plasma membrane 2.90E-11 Integral component of membrane 1.30E-90 Golgi apparatus 2.00E-53 Endoplasmic reticulum Plasmodesma 4.00E-77 Vacuole 2.60E-66 5.30E-46 ATP binding RNA binding 1.90E-11 Protein binding 2.50E-03 Serine/threonine kinase activity 3.70E-03 1.90E-127 Regulation of transcription 2.10E-109 Transcription, DNA-templated 3.50E-64 Nucleus 1.30E-12 Mitochondrion 5.30E-12 Extracellular region 2.90E-215 Transcription factor activity 3.20E-106 DNA binding 3.30E-45 Zinc ion binding Missing Covered Biological Process (BP) Cellular Compartment (CC) Molecular Function (MF) 1651 1652 1653 1654 1655 1656 1657 1658 1659 1660 1661

1662 Supplemental figure 5







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1665 Supplemental figure 6



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1672 Supplemental figure 7



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1710 Supplemental figure 10



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1747 Supplemental figure 14



1758 Supplemental figure 15



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