¹ The dynamics of root cap sloughing in

² Arabidopsis is regulated by peptide signalling

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29 ABSTRACT

30

The root cap protects the stem cell niche of Angiosperm roots from damage. In Arabidopsis 31 32 lateral root cap (LRC) cell files covering the meristematic zone are regularly abolished 33 through programmed cell death, while the outermost layer of the root cap covering the tip is 34 regularly sloughed. Efficient coordination with stem cells producing new layers is needed to 35 maintain a constant size of the cap. We present a signalling pair, the peptide IDA-LIKE1 36 (IDL1) and its receptor HAESA-LIKE2 (HSL2), mediating such communication. Live imaging over several days characterized the sloughing process from initial fractures in LRC 37 38 cell files to full separation of a layer. Enhanced expression of IDL1 in the separating root cap 39 layers resulted in increased frequency of sloughing, balanced with generation of new layers in a HSL2-dependent manner. Mutations in either IDL1 or HSL2 slowed down cell division, 40 maturation and separation, and suggested involvement of programmed cell death in the 41 42 sloughing event. Transcriptome analyses linked IDL1-HSL2 signalling to the transcription 43 factors BEARSKIN1/2.

45	The root cap provides protection for the root apical meristem (RAM) and senses
46	environmental conditions ¹ . The RAM encompasses stem cells, surrounding the Quiescent
47	Centre (Fig. 1a), both for the tissues of the root proper, and for the root cap. Although both
48	forms of stem cells continuously produce new cells, the root cap has a defined size, due to
49	homeostatic balance between generation and loss of root cap cells ² . Many species shed living,
50	single cells, so called border cells, but in Brassicaceae, including Arabidopsis thaliana, the
51	outermost layer of the root tip is released intact from the underlying cell layer ³ . The
52	Arabidopsis root cap distal to (below) the QC, consists of a core of 5-6 tiers of gravity-sensing
53	columella cells (COL) surrounded by layers of lateral root cap (LRC) cells (Fig. 1a). The LRC
54	extends to cover the meristematic zone (MZ), and there the outermost layer regularly
55	undergoes programmed cell death (PCD) whereby it is replaced by a new LRC layer
56	developing underneath the old one ⁴ . Less is known regarding the genetic and molecular
57	mechanisms governing the detachment of the outer layers of the distal root cap, how the
58	frequency of sloughing is regulated and how the homeostatic balance with stem cells is
59	secured ² .

Root cap sloughing requires breakdown of cell walls and is in that respect similar to other cell separation processes such as flower or fruit abscission ⁵, however, sloughing is a recurrent event in the same organ. Hence, the challenge is to compensate for each lost root cap layer by generation of a new, and to detach the oldest layer when a new layer has been formed. In the distal root cap, this requires communication between old separating layers and the stem cells of COL and LRC, as well as a sequential coordination of generation, maturation and separation of root cap cells.

Mutations in four genes encoding closely related NAC (<u>NAM, ATAF, CUC</u>)
transcription factors (TFs) – FEZ, SOMBRERO (SMB), BEARSKIN1 (BRN1) and BRN2 –
have provided insights into the development and differentiation of the root cap ^{4,6-8}. FEZ is

implicated in the formative division of the common stem cells for the LRC and the epidermal
layer (EPI). SMB regulates differentiation and maturation⁶⁻⁸, while the double mutant *brn1 brn2* is impaired in cell wall degradation and accumulates additional undetached root cap
layers⁸.

Downstream of these TFs, genes have been identified that encode enzymes involved 74 cell wall remodelling and synthesis⁸. Similar genes have been discovered in other cell 75 separation processes in flowering plants ^{9,10}. Floral organ abscission and emergence of the 76 77 lateral root though the overlying tissues in *Arabidopsis*, are regulated by the small peptide 78 INFLORESCENCE DEFICIENT IN ABSCISSION (IDA) that signals through two closely 79 related leucine-rich repeat receptor-like kinases (LRR-RLKs), HAESA (HAE) and HAESA-LIKE2 (HSL2)¹¹⁻¹⁵. *IDA* belongs to a family of *IDA-LIKE* (*IDL*) genes, encoding 80 81 preproproteins with a common C-terminal motif (Supplementary Fig. 1), hypothesized to have 82 a function similar to IDA. Here, we address the function in root cap sloughing of IDL1, expressed in the root tip, and identify HSL2 as its receptor. Rather than specifically 83 84 controlling the cell separation step of the sloughing process, this signalling pair regulates the 85 dynamics of root cap detachment and generation of new root cap layers. 86

87 **RESULTS**

88 Identification of HSL2 as the receptor of IDL1

89 *IDL1* is prominently expressed in COL cells at the centre of the two adjacent root cap layers

- 90 of the primary root that will undergo separation (Fig. 1a-b). IDL1 can rescue the abscission
- 91 deficiency of the *ida* mutant 16 , indicating that IDL1 peptide can interact with the receptors of
- 92 IDA. Using promoter- β -glucuronidase (GUS) reporter lines for *HAE* and *HSL2* we found that
- only *pHSL2:GUS* had an expression pattern in the primary root tip, with the strongest
- expression in the youngest LRC cell files (Supplementary Fig. 1). This was confirmed when

95	using a construct with the HSL2 promoter coupled to a nuclear targeted florescent protein
96	(pHSL2::Venus-H2B) (Fig. 1c). The N-terminal signal sequence of the IDL1 prepropeptide
97	(Supplementary Fig. 1) indicates that the mature peptide is exported and can move in the
98	apoplastic space to interact with the extracellular domain of the preferred receptor ^{17,18} . Thus,
99	the expression patterns were compatible with a hypothesized function of IDL1 and HSL2 as a
100	ligand-receptor pair involved in communication between cells in the root cap.
101	To investigate whether IDL1 could activate and bind the receptor, we took advantage
102	of methods previously used for IDA and HSL2 12 . Addition of a synthetic IDA peptide of 12
103	amino acids, with hydroxylation of the central proline residue (mIDA), to Nicotiana
104	benthamiana leaf pieces expressing the HSL2, results in an immediate receptor activation
105	measurable as oxidative burst. The mIDA peptide (Fig. 1d) activates HSL2 down to a half-
106	maximal efficient concentration (EC50) of 1 nM ¹² . The corresponding hydroxylated dodeca
107	IDL1 peptide (mIDL1, Fig. 1d), tested in the same assay, functioned also in a dose-dependent
108	manner with a similar efficiency (EC50 ~3 nM, Fig. 1e).
109	Competition assays with acridinium-labelled and unlabelled IDA-derived peptides
110	have demonstrated specific and reversible binding of the ectodomain of HSL2 expressed in <i>N</i> .
111	benthamiana leaves ¹² . The same experimental set-up showed that unlabelled mIDL1
112	competed for binding of the HSL2 ectodomain with an efficiency similar to the unlabelled
113	IDA peptide, with a half-maximal inhibitory concentration (IC50) of ~30 nM (Fig. 1f).
114	Root cap expression, together with binding and activation of HSL2 by IDL1,
115	suggested that this peptide and receptor might serve as a root cap signalling module.
116	
117	IDL1 expression increases the frequency of sloughing in a HSL2-dependent manner
118	Since a number of LRR-RLKs with high similarity in their ectodomains are expressed in the
119	root ¹⁹ , ectopic ligand expression or exogenously applied synthetic peptides might lead to

120	interaction with non-native receptors ²⁰ . With the aim of testing whether IDL1-HSL2 was
121	specifically involved in root cap sloughing, we therefore used the promoter of <i>IDL1</i> itself to
122	generate Enhanced IDL1 (EnhIDL1) plants with estradiol (ES)-inducible enhanced expression
123	of both <i>IDL1</i> and <i>GUS</i> (Fig. 2a-b) ²¹ . EnhIDL1 plants were crossed to an <i>hsl2</i> knockout
124	mutant ¹⁶ to test the importance of HSL2 for IDL1 function. Cell-specific and efficient ES-
125	induction was confirmed by strongly enhanced GUS expression in EnhIDL1 roots both in
126	wild type Col-0 and hsl2 mutant background as compared to the pIDL1:GUS expression
127	pattern and by qRT-PCR (Fig. 1b, 2b and Supplemental Fig. 1). These lines did not deviate in
128	root length (Supplementary Fig. 1).
129	Cell elongation, cell division and differentiation take place in the root tip during the
130	first days after germination. To ensure that homeostatic balance between gain and loss of root
131	cap layers had been established, our experiments were performed on roots with full-sized
132	RAM and root cap (Supplementary Fig. 2). Seven days after germination (DAG) roots of Col-
133	0, EnhIDL1 and EnhIDL1 <i>hsl2</i> seedlings grown vertically on agar with 10 μ M ES, covered by
134	a cover slip, were tracked, using a laser scanning microscope with a vertical stage ²² . Images
135	taken in two experiments every 15 minutes for 64 and 70 hours (h) revealed a stepwise
136	sloughing process (Fig. 2c-e). The first signs of a forthcoming sloughing event were
137	characteristic fractures or gaps in LRC cell files in a region just above the Quiescent Centre
138	(QC) and about 5h later at the level of the youngest starch-filled COL tiers (Fig. 2c, red and
139	yellow arrows). These gaps initiated a gradual loosening of the outermost LRC layer towards
140	the tip. Finally, the COL cells at the centre of the cap separated from the layer underneath
141	followed by detachment of the whole layer as it slid to the side and was lost (Fig. 2c, white
142	arrow and Supplementary Video 1). On average, 0.6 root cap layers were shed per root per
143	day in Col-0 (Fig. 2f), with an average of ~18h for a completed sloughing process, and

thereafter another ~18h before a new event of root cap layer separation would be initiated (Fig.
2g and Supplementary Table 1).

146	The sloughing frequency in EnhIDL1 was 50 % higher than for Col-0, i.e. 0.9 events
147	per root per day (Fig. 2f). This increase seemed to be caused by a shorter interval between the
148	completion of one event and the initiation of the next (Fig. 2g and Supplementary Table 1),
149	and overlapping events with fractures in the second outermost layer, before the previous event
150	was fully completed (Fig. 2d, green arrow, 2h, and Supplementary Video 2).
151	In hsl2 background the increased sloughing frequency of EnhIDL1 was rescued and
152	did not deviate from that of Col-0 (Fig. 2f). However, the sloughing process developed most
153	often differently in that the outermost layer was split open close to the tip at the border
154	between the COL and LRC cells (Figure 2e and h). The split cap layer, sometimes leaving the
155	central COL region behind, seemed to be still attached to the LRC sheet surrounding the
156	meristematic region (Fig. 2e, white arrows, and Supplementary Video 3).
157	Live imaging was supplemented with data for roots grown on vertical agar plates. In
158	this set-up, shed layers of the distal cap, recognizable by detached LRC files, did not slide
159	away immediately, but accumulated in front of the growing roots, most likely due to less
160	friction when the roots were not covered by a cover slip. Caps were, however, easily lost
161	when removed from the agar, and sloughed caps were therefore counted directly on the plates
162	at 6, 8 and 10 DAG, with and without 10 μ M ES (Fig. 3a-b). Col-0, EnhIDL1 hsl2, and
163	EnhIDL1 on non-inducing medium sloughed about 0.6 caps per day (Fig. 3a-b). On ES
164	medium, on the other hand, EnhIDL1, accumulated up to eight sloughed root cap layers
165	(Supplementary Fig. 3) and differed significantly with an average loss of one root cap layer
166	per day (Fig. 3a). Thus, these results were in full accordance with the live imaging results.
167	

169 Higher frequency of IDL1-induced sloughing is compensated by more stem cell divisions

170 We questioned whether a higher frequency of sloughing reduced the number of attached

171 layers, however, all three lines maintained stably ~6 attached COL tiers (including the stem

cell layer) (Supplementary Fig. 3), and did not differ significantly from each other (Student's

t-test P \ge 0.1), irrespective of ES in the growth medium (Fig. 3c).

174 More frequent shedding, but at the same time constant cell number and size of the root cap, imply an increased frequency of new initiations of COL tiers and LRC cell files to 175 176 compensate for the lost caps. Recent divisions of COL stem cells, positioned just below the 177 QC (Fig. 1a), can be recognized by the smaller cell size in the new 1st COL tier and their weaker cell wall staining with propidium iodide (PI) compared to older COL cells ⁶. The COL 178 179 stem cells are surrounded by the shared stem cells for LRC and epidermal cell files, and in 180 medial optical sections of the root tip, they are seen as four COL stem cells with one EPI/LRC 181 stem cell on each side (Fig. 1a). New epidermal cells are generated by anticlinal division, while periclinal divisions represent initiation of a new LRC cell file. On inductive medium, 182 183 cell wall patterns indicating recent cell divisions were found for COL in 80% and for LRC in 184 60% of EnhIDL1 PI-stained root tips (Fig. 3d-e). This was in contrast to Col-0 and even more 185 so for EnhIDL1 *hsl2*, where < 15 % of the root tips showed signs of recent stem cell divisions (Fig. 3d-e). 186

187 These findings imply that increased expression level of *IDL1* not only leads to
188 increased sloughing, but also an increased frequency of initiation of new root cap layers.

189

190 Mutations in *IDL1* and *HSL2* slow down the sloughing process

191 To further explore the function of the IDL1-HSL2 signalling in the root cap, we investigated

the sloughing behaviour of $hsl2^{16}$ and an *idl1* CRISPR/Cas9 mutant line (Supplementary Fig.

4) compared to Col-0. While the number of attached layers remained close to 6 for both Col-0

and the mutants, a subtle, but significant difference was observed regarding the number of 194 sloughed cap layers at 10 DAG with on average 3.0 caps for Col-0 and 2.4 for the mutants 195 196 (Fig. 4a). Inspection of the youngest LRC cell files furthermore suggested that the frequencies of generation of new LRC cell layers had slowed down in both mutants. After the initial 197 198 periclinal division of a EPI/LRC stem cell, the new LRC cell file is extended by anticlinal 199 divisions. The new file will be neighbouring an older and longer LRC cell file on the lower side and an epidermal cell file on the upper side until the next new LRC layer is initiated (Fig. 200 201 4b-d). In the wild type, one cell of the two youngest cell files neighboured the epidermal cell 202 file, in contrast to 2-3 cells in both the *hsl2* and *idl1* mutants (Fig. 4b-c). This is an expected 203 result if the periclinal divisions initiating a new LRC file take place less frequently relative to 204 the anticlinal divisions of already initiated LRC cell files (Fig. 4d).

205

206 Mutation in *HSL2* affects genes in both the LRC and the COL cell layers

To gain a better understanding of the function of the signalling initiated by activation of HSL2, 207 208 RNA sequencing was performed on RNA isolated from the RAM of root tips of Col-0 and 209 hsl2. In line with the subtle phenotype of hsl2, few genes were grossly affected by the 210 mutation. However, using a P adjusted (Padj) cut-off value of 0.1, 156 genes were identified with an expression level < 70 % of the Col-0 level (Supplementary Table 2). In an attempt to 211 identify where the genes affected by HSL2 signalling were expressed, we took advantage of a 212 data set for tissue-specific gene expression in the Arabidopsis RAM²³. This dataset provides 213 214 information on differential gene expression in the different root tissue, QC, LRC, COL, EPI, 215 cortex (COR) and endodermis (END), based on cell sorting using marker genes for these tissues ²³. Eighty-one of the *hsl2* down-regulated genes were identified as differentially 216 217 expressed according to this data set. We defined a gene as characteristic for a specific tissue if it had the highest expression level for this tissue compared all the other tissues. Evaluation 218

according to a binomial test (p=0.1, see Methods for details) found significant enrichment of
genes expressed in LRC and COL (Table 1). Genes encoding extracellular proteins were well
represented among the LRC expressed genes, while many genes most highly expressed in
COL encoded proteins involved in catalytic and hydrolytic activity including starch and cell
wall degradation.

224 Activation of RLK receptors is transduced via phosphorylation cascades and 225 activation of transcription factors, which at present are unknown for HSL2 in the root tip. 226 However, we found an overlap between the *hsl2* down-regulated genes and genes identified in a recent transcriptome analysis of NAC TFs⁸. Of sixty genes reported to be positively 227 228 regulated by overexpression of SMB and downregulated in the triple mutant *smb brn1 brn2*, twenty-five genes, a highly significant number (Fischer's exact test $P < 10^{-10}$), was also down 229 230 regulated in *hsl2* mutant root tips (Supplementary Table 2). There was an overrepresentation 231 of genes encoding extracellular proteins and enzymes, hereunder enzymes involved in cell wall remodelling and starch metabolism (Supplementary Table 2)^{8,24,25}. 232 233 The three NAC TFs were not significantly down regulated in hsl2 according to our 234 RNAseq analyses, however, to investigate whether IDL1 signalling could affect expression 235 patterns, lines with translational fusion constructs for BRN1 and BRN2 were crossed into the EnhIDL1 line and the *idl1* mutant. Both *BRN1* and *BRN2* are expressed in the outer LRC cell 236 files⁸. In EnhIDL1, the expression pattern of *pBRN1:BRN1-GFP* was very similar to Col-0 237 238 grown with ES in the medium, while *pBRN2:BRN2-GFP* consistently showed an additional 239 layer of expressing cells at the tip (Supplementary Fig. 5). In the *idl1* mutant, on the other 240 hand, *pBRN2:BRN2-GFP* expression was similar to Col-0, but fewer cells expressed 241 *pBRN1:BRN1-GFP* in the LRC cells (Supplementary Fig. 5).

The *ROOT CAP POLYGALACTURONASE* (*RCPG*), a direct target of BRN1⁸, was of the most strongly downregulated genes both in the triple mutant *smb brn1 brn2* and *hsl2*

244	(Supplementary Table 2). In loosened Col-0 root cap layers the construct pRCPG:nYG, with
245	nuclear targeted YFP driven by the <i>RCPG</i> promoter ⁸ , was expressed in more cells and with
246	higher intensity than in hsl2 and idl1 background (Fig. 5a). Moreover, pRCPG:nYG was
247	expressed in undetached LRC cells in the mutants (Fig. 5a), suggesting delayed cell
248	separation. pRCPG:nYG expression was also seen in undetached root cap layers in the
249	EnhIDL1 line, in this case suggesting premature expression (Supplementary Fig. 5).
250	This influence on RCGP expression suggests that IDL1-HSL2, like the BRN TFs, are
251	involved in the final step of the sloughing process – the actual cell separation.
252	
253	IDL1-HSL2 regulates BFN1 expression at PCD Site II
254	We noted that in contrast to hsl2 and idl1, most Col-0 cells expressing RCPG had initiated the
255	sloughing process and their nuclei stained with PI (Fig. 5a and Supplementary Fig. 5), which
256	is indicative of plasma-membrane permeabilization. Such PI entrance has been used as a
257	hallmark of initiation of developmental programmed cell death (dPCD) associated with the

recurrent turnover of the LRC covering the MZ⁴. Interestingly, in the *hsl2* transcriptome five

out of eight genes strongly associated with dPCD were down-regulated; the BIFUNCTIONAL

260 NUCLEASE1 (BFN1), METACAPASE9 (MC9), SAPOSIN-LIKE ASPARTYL PROTEASE3

261 (*PASPA3*), *EXITUS* (*EXI*) and *RIBONUCLEASE3* (*RNS3*) ^{4,26} (Supplementary Table 2). To

investigate the potential involvement of IDL1-HSL2 in dPCD in the root cap, a transcriptional

fusion construct with the *BFN1* promoter and a nuclear-targeted GFP $(BFN1_{pro}:nGFP)^{27}$ was

crossed into EnhIDL1 and the mutants. In Col-0, two known sites of dPCD, at the upper end

of the MZ (Site I) and close to the QC (Site II), were identified as expected (Fig. 5b).

266 Interestingly, in addition to the strong expression of pHSL2::Venus-H2B in young LRC cell

267 files, HSL2-driven nuclear-targeted Venus was found in LRC cell files surrounding the MZ

and stretching past Site II (Fig. 5c). A third PCD site was found in loosened LRC cell files

consistent with the high number of nuclei there staining with PI (Fig. 5a-b), and suggestive of
PCD in the sloughed cells. The pattern of BFN1_{pro}:nGFP expression in EnhIDL1 at these sites
was similar to Col-0. In contrast, *idl1* and *hsl2* root tips had significantly fewer cells
expressing the BFN1_{pro}:nGFP construct, in particular at Site II (Fig. 5b and Supplementary
Fig. 6). Interestingly, Site II coincides with the position of the fractures initiating the
detachment of the distal root cap (Fig. 5b), which suggestively may provide a molecular link
between the IDL1-HSL2 signalling pair and initiation of the sloughing process.

276

277 DISCUSSION

278 Dynamics of root cap sloughing and mechanisms securing homeostatic balance between loss 279 and gain of layers of the distal root cap have been difficult to study because of the duration of the process, the easily lost, small cap layers, and few available mutants². Continuous live 280 imaging over three days, analyses of transcriptome data and cellular expression patterns have 281 282 enabled us to detail the different phases of root cap sloughing and the involvement of IDA-283 HSL2 in this process (Fig. 6). Under our conditions, 0.6 distal cap layers were shed per root 284 per day in Col-0. Enhanced expression of *IDL1* increased this frequency by shortening the 285 interval from one event to the next, or by detaching two layers at the same time. We have 286 identified IDL1 as a peptide ligand that in nanomolar concentrations binds and activates the HSL2 receptor. Both IDL1-increased sloughing and the balancing increase in stem cell 287 288 divisions were dependent on HSL2, i.e. IDA-HSL2 signalling controls the dynamics of sloughing at the root tip. 289

We had expected cell separation phenotypes of *idl1* and *hsl2* similar to the total
deficiency in floral organ abscission seen in the *ida* mutant and the double mutant *hae hsl2*¹⁶.
However, homeostasis appears maintained in the two mutants, albeit at a much lower
frequency. Cell division, maturation and separation were slowed down, but not abolished. It is

294 likely that other factors, including other peptides and receptors, are involved in the separation 295 step. A few mutants have been identified that release single cells instead of entire root cap layers, like nin-like protein 7 (nlp7), quasimodo1-1 (qua1-1) and qua2-1^{25,28}. In contrast, the 296 brn1 brn2 double mutant, and active quiescent center1-1 (aqc1-1), are delayed in separation 297 of root cap layers ^{8,29}. The latter mutant is defect in the gene encoding tyrosylprotein 298 299 sulftransferase (TPST), an enzyme responsible for sulfation of a number of peptide ligands. 300 The disorganized root cap and stunted growth in this mutant is likely to be a consequence of 301 several dysfunctional peptides lacking sulfation.

302 Live imaging of roots and marker-gene expression in the EnhIDL1 line and the *idl1* 303 and *hsl2* mutants suggest that IDL1-HSL2 signalling is controlling initiation of the sloughing 304 process. Intriguingly, the region where the first formative fractures or gaps in the LRC cell 305 files are found, seems to coincide with dPCD Site II, and a speculative suggestion is that these 306 fractures are created by PCD. dPCD starting from Site I is involved in the recurrent removal of the LRC surrounding the MZ⁴. Differential control of dPCD Sites I and II would make it 307 308 possible for the plant to operate with different frequencies of renewal of the LRC surrounding 309 the MZ and of sloughed layers at the tip.

310 IDL1-HSL2 control of the position of initiation may also explain the split caps still connected to the LRC surrounding the MZ observed for EnhIDL1 in hsl2 background. With 311 fewer fractures, MZ cell division and COL cell elongation is likely to exert mechanical stress 312 313 that tears apart cells at other, weaker sites. Similarly, in *ida* and *hae* mutants, separation of 314 cells in the cortex and epidermis overlying developing lateral root primordia (LRPs) is impaired, but the LRPs eventually emerge forcibly through these tissues ¹⁰. These two 315 316 examples suggest that regulation of growth of the LRP and the primary root is not tightly 317 coupled to the respective cell separation events.

BRN1, BRN2 and SMB are implicated in root cap maturation and root cap detachment 318 ^{7,8}. pHSL2::Venus-H2B was found to have overlapping expression pattern with the BRN TFs 319 in the LRC, and a subset of genes regulated by these TFs was also found downstream in the 320 321 HSL2 signalling pathway. Differences in spatial expression patterns compared to Col-0 of 322 transcriptional and translational reporter construct crossed into our lines, indicated faster 323 maturation in EnhIDL1 and slower maturation in the mutants. Together this suggests HSL2 to play a role in maturation and cell layer separation in addition to the role in initiation. $BFNI_{pro}$ -324 325 controlled maker-gene expression and PI-stained nuclei in LRC cells of root caps undergoing 326 sloughing, indicate that root cap sloughing involves both cell separation and cell death. 327 RNAseq data for *hsl2* combined with publicly available information on differential 328 gene expression in root tissues were used in an attempt to identify the cell types that primarily 329 respond to HSL2 activation. This analysis identified overrepresentation of genes preferably 330 expressed in LRC and COL cells. The major effect of receptor activation will occur in the cells where the receptor is expressed, indicating that HSL2 is present and responsive for 331 332 activation both in LRC and COL cells. Additionally, this analysis suggests that activation of 333 HSL2 gives different outcomes in different tissues.

334 IDL1-HSL2 communication facilitates dynamic regulation of the homeostatic balance between stem cell division and sloughing activity. Since the root cap is guiding root growth 335 by sensing and responding to external cues, like gravity, water potential or the presence of 336 obstacles in the soil ^{1,2}, it is highly likely that the frequency of root cap shedding is triggered 337 338 by environmental factors, possibly by influencing the expression level of IDL1 and/or HSL2. 339 At present, it is not known whether or how external condition might feed into the circle of 340 generation and loss of root cap layers. Factors involved in transcriptional regulation of *IDL1* 341 and HSL2, peptide processing and export, and interacting partners have not been identified. Additionally, several parallel pathways partake in regulation of stem cell activity, the NAC 342

343	transcription factors FEZ and SMB, the RETINOBLASTOMA-RELATED protein, the
344	WOX5 transcription factor, and the AUXIN RESPONSE FACTORS ARF10 and ARF16 30,32 ,
345	but their relationship to the IDL1-HSL2 signalling pathway have not yet been unravelled.
346	It will be exciting to take advantage of root tip tracking by live imaging and to explore
347	these relations in the future.
348	
349	
350	Methods
351	Accession numbers of the genes studied in this work: <i>IDL1 – At3g25655, HSL2 –</i>
352	At5g65710; HAE – At4g28490; BRN1 – At1g33280, BRN2 – At4g10350, BFN1 – AT1G11190
353	and RCPG–At1g65570.
354	Constructs and plant lines. The <i>pIDL1:GUS</i> , <i>pHAE:GUS</i> , <i>pHSL2:GUS</i> , <i>BFN1</i> _{pro} :nGFP,
354 355	Constructs and plant lines. The <i>pIDL1:GUS</i> , <i>pHAE:GUS</i> , <i>pHSL2:GUS</i> , <i>BFN1</i> _{pro} : <i>nGFP</i> , <i>pRCPG:nYG</i> , <i>pBRN1:BRN1-GFP</i> and <i>pBRN2:BRN2-GFP</i> lines have been described
354 355 356	Constructs and plant lines. The <i>pIDL1:GUS, pHAE:GUS, pHSL2:GUS, BFN1_{pro}:nGFP,</i> <i>pRCPG:nYG, pBRN1:BRN1-GFP</i> and <i>pBRN2:BRN2-GFP</i> lines have been described previously ^{8,16,27} . A 2300 bp long fragment upstream of the start codon of <i>HSL2</i> ³¹ was cloned
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pMDC221 line (checked on 50 mg/l Kanamycin (Km)). EnhIDL1 plants homozygous for both
constructs were identified by selection on Hyg and Km and checked non-segregating GUS
expression. Of two lines with comparable sloughing behaviour, GUS expression level and
pattern, one was chosen for the experiments shown here. This line was furthermore used for
all crosses to mutants and marker-lines.

371 Two idl1 mutants were made with CRISPR/Cas9 technology using an IDL1-targeting vector construct based on the pDe-Cas9 systems provided by Holger Puchta³³. A 20 base pair 372 373 gRNA target site, unique for IDL1 including BslI recognition motif, was selected for the 374 mutagenesis. Vector construction and generation of mutants were performed as described previously ³⁴. Through selections with BASTA and CAPS analyses, mutants containing 2 bps 375 376 deletion (*idl1-cr1*) and 4 bps deletion (*idl1-cr2*) in the open reading frame of *IDL1* were 377 isolated. Both mutations result in frame shifts generating premature stop codons in the central 378 variable region of IDL1 prepropeptide and the C-terminal peptide motif is thereby deleted (Supplementary Fig. 4). Phenotypic differences between the alleles were not observed 379 380 (Supplementary Fig. 4). Data for *idl1-cr1* (named *idl1* here) is presented. 381 Primers for genotyping and generation of constructs are listed in Supplementary Table

382 3.

Growth conditions and registration of number of root cap layers. Sterilized seeds were 383 stratified at 4 °C for 72 h. Seedlings were grown at 22 °C, 8 h dark and 16 h light (100 µE/m2 384 385 light intensity) on square vertical plates with sucrose-free 0.5 x MS medium adjusted to pH 386 6.0, with 10μM 17-β-estradiol for induction of enhanced expression of *IDL1* in EnhIDL1 lines ²¹. Sloughed root cap layers were counted and imaged directly on the agar plates (using a 387 388 Zeiss Axioplan2 imaging microscope) as they easily fall off when roots are moved. Attached 389 caps were imaged and counted after immersion in Lugol solution (Sigma 6265) for 1 minute for detection of the COL cell tiers with amyloplasts (Supplementary Fig. 3). No significant 390

difference in length was observed between genotypes or with ES in the medium

392 (Supplementary Fig. 1).

393	Oxidative burst measurements and receptor-ligand binding assay. N. benthamiana leaves
394	transiently expressing HSL2 constructs were used for oxidative burst measurements and
395	peptide binding assays as previously described ¹² . EC50 and IC50 curves are representative
396	for N=3 independent experiments, each with three technical replicates.
397	Microscopy. H istochemical GUS assays were performed as previously described ¹⁶ . Confocal
398	laser scanning microscopy (CLSM) was carried out with Olympus FV1000 and ZEISS
399	LSM880 confocal microscopes at the Oslo NorMIC imaging node. Roots were stained with
400	10 μM propidium iodide (PI).
401	For live imaging seeds were surface sterilized by chlorine gas, sown on 1% agar AM-
402	medium (0.5 x MS, pH 5.8, 1% agar), stratified for 3 days at 4°C and cultivated in a growth
403	incubator at 22°C in a 16/8 h day/night cycle with 120–140 μ mol/m ² /s light.
404	Eight day-old plants were put on blocks (rectangular parallelepipeds) of 1.5% agar AM-
405	medium (0.5 x MS, pH 5.8) supplemented with 10 μ M 17- β -estradiol. Roots and shoots were
406	put on the blocks' basis and sides, respectively ²² . These samples were put into the chambered
407	cover-glass (Thermo Scientic Nunc, catalogue no: 15536), which were then mounted onto a
408	laser scanning microscope with a vertical stage (roots down, shoots up). Photographs of root
409	tips were taken every 15 minutes, in two experiments over 64 and 70 h, with in total 9 roots
410	for Col-0 and 10 roots for EnhIDL1 and EnhIDL1 hsl2.
411	RNA extraction and RNAseq analysis. Total RNA was extracted in three replicates from 0.5
412	mm segments of primary root tips of 7-day-old Col-0 and hsl2 seedlings using
413	Trizol/chloroform extraction follow by purification using the RNeasy Plant Mini Kit

414 (Qiagen). The quality of the RNA was ascertained using the Agilent 2100 Bioanalyzer.

- 415 RNAseq was performed at the Functional Genomics Center Zurich, Switzerland. Deseq2 with
- 416 standard parameters was used to analyse expression levels
- 417 (https://bioconductor.org/packages/3.7/bioc/vignettes/DESeq2inst/doc/ DESeq2.html).
- 418 **Statistical methods** *Student's t-test* (2-tailed) was used to analyse whether root cap
- 419 phenotype and marker-gene expression in different lines deviated significantly from Col-0 cf.
- 420 Fig. 2, 3 and 4, and Supplementary Fig. 1, and 6. If not otherwise stated in the figure legends
- 421 ≥ 10 roots were used for each genotype, treatment and age. Two to five biological independent
- 422 repeats were used. In all graphs, standard deviations are shown as bars.
- 423 Binomial test was applied to test for enrichment of maximal expression level for a specific
- tissue across all other tissues. The probability to find k maximal expressions out of N genes in
- 425 total is given by

$$P(X \ge k) = \sum_{i=k}^{N} {N \choose i} p^{i} (1-p)^{N-i}$$

- 426 whereby p is the per gene probability to be maximal and () is the binomial coefficient.
- 427 Fisher's exact test was used to calculated the likelihood of overlap of regulated genes by
- 428 *SMBox /smb brn1 brn2* (NAC TFs) and *hsl2*.

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525 Material requests

- 526 Constructs or seeds, as well as the raw data underlying the RNAseq data in Supplementary
- 527 Table 2 are available from R.B.A. upon request, e-mail: reidunn.aalen@ibv.uio.no.

528

529 Contributions

- 530 C.-L.S., T.I., S.S., U.H., M.A.B, M.W., M.K.A. and V.O. generated *Arabidopsis* lines; J.F.
- and D.v.W. designed, D.v.W. and I.K. performed, and D.v.W., I.K. and R.B.A. analysed live
- imaging experiments; G.F., M.A., M.A.B designed and M.W. performed IDL1-HSL2
- 533 interaction studies; M.C. and R.B.A. designed and A.K., M.C. and R.B.A. analysed RNAseq
- data; the rest of the experiments were designed by C.-L.S. and R.B.A., performed by C.-L.S.
- together with U.H., M.W., V.O. and analyzed by R.B.A., C.-L.S., M.W. and U.H.; C.-L.S.
- drafted the manuscript; R.B.A. wrote the paper with input from other authors.

537 **Competing interests**

538 The authors declare no competing interests.

539¹²

540 Figure legends:

541 Fig. 1 | IDL1 peptide interacts with and activates HSL2. a, The tissues of the Arabidopsis

542 root tip. LRC – lateral root cap. **b**, pIDL1:GUS after overnight incubation with X-gluc. **c**,

- 543 Confocal medial optical section of 7 DAG PI-stained root tip expressing pHSL2::Venus-H2B.
- 544 The QC is indicated by an ellipse. d, Alignment of mIDA and mIDL1 peptides. * -
- 545 hydroxylated proline. e, mIDL1 activates HSL2 transiently expressed in *N. benthamiana* in a
- 546 dose-dependent manner. Activation resulted in oxidative burst which was monitored as
- relative light units (RLU) using a luminol-based assay ⁸. EC50 half-maximal efficient
- 548 concentration. **f**, Unlabelled mIDL1 compete efficiently for binding of HSL2. The mIDA
- 549 peptide with an N-terminal Valine residue labelled with acridinium (acri-V-IDA) bound to *N*.
- 550 benthamiana-expressed HSL2, was treated with increasing concentrations of unlabelled
- 551 mIDL1 or V-IDA. 100% binding assay without competitors; negative control (background
- signal) leaf tissue not expressing HSL2 incubated with 10 nM acri-V-IDA peptide and
- 553 increasing concentrations of unlabelled V-IDA. IC50 half-maximal inhibitory concentration.

554

555 Fig. 2 | Live imaging details the sloughing process in Col-0, the effect of enhanced

556 expression of IDL1 and dependency on HSL2. a, The two-component constructs for

inducible enhanced expression of IDL1 and GUS. The IDL1 promoter controls expression of

- 558 XVE, which in the presence of estradiol (ES) binds the OlexA-TATA box. b, ES-induced
- 559 GUS expression in Col-0 and *hsl2* mutant background after 1 hour (h) incubation with X-Gluc.
- 560 Controls were grown without ES. c, Representative stages of the standard ~ 18h sloughing
- 561 process of Col-0, captured by live imaging, i.e. initial fracture in LRC cell files, first above,
- after several h just below the QC (red and yellow arrows), and finally separation of a whole
- 563 cap layer (white arrow). **d**, Overlapping sloughing events in EnhIDL1 root tips, i.e. a new

564	initiating gap (green arrow) occurring before completion of the ongoing event. e, Split root
565	caps of EnhIDL1 <i>hsl2</i> , i.e. gaps at the LRC-COL border (blue arrow) followed by detachment
566	of a split cap (white arrows) still attached to the LRC covering the meristematic zone. f ,
567	Number of initiations and detachments per root per 24h. Bars indicate standard deviation. * -
568	significantly different from Col-0 (Student's t-test P< 0.05). Number of roots - Col-0 9,
569	EnhIDL1 and EnhIDL1 $hsl2^{10}$. g, Average time from initiation to detachment and from
570	detachment to initiation of a new event. N – number of events per genotype. Standard
571	deviations are given as bars. * - significantly different from Col-0 (Student's t-test P< 0.04). h,
572	Total number of sloughed caps and their phenotypes (cf. c-e) documented by live imaging
573	over 64 to 70 hours.

575 Fig. 3 | Higher frequency of sloughing is compensated with more stem cell divisions. a,

576 Number of detached root cap layers 6, 8 and 10 days after germination (DAG). * -

significantly different from other lines (grey), and without ES (blue) (Student's t-test P<

578 0.005). N=10-12 roots per genotype per treatment and per time point. Standard deviations are

given as bars. **b**, Root tips after 10 days on medium with 10 μ M ES, and control without ES. **c**,

580 Number of attached COL tiers at 8 and 10 DAG. d, Percentage of root caps with new 1st tier

581 COL cells, and new LRC cell generated after division of the respective stem cells. * -

significantly different from Col-0 (Student's t-test P < 0.01). e, Medial optical sections of 7

583 DAG PI-stained stem cell niches of roots grown with ES, with schematic drawing below.

584 Dots mark novel cell walls in EnhIDL1.

585

Fig. 4 | *idl1* and *hsl2* mutants influence sloughing frequency and LRC division patterns.

a, Number of sloughed and attached layers. * - significantly different from Col-0 (Student's t-

test $P_{idl1} < 10^{-3}$, $P_{hsl2} < 10^{-6}$, N = 10). **b**, Stem cell niche of the indicated genotypes. Asterisks indicate LRC/Epidermis stem cells (pink), and LRC cells directly neighbouring the epidermal cell file (light and dark blue). **c**, Schematic presentation of cellular patterning shown in b. Arrows show the directions of cell divisions. **d**, Number of cells in the first and second LRC cell file neighbouring the epidermal cell file. * - significantly different from Col-0 (Student's t-test P < 10-5). $N_{Col-0}=22$, $N_{hsl2}=17$ and $N_{idl1}=18$. Standard deviations are given as bars.

594



604

Fig. 6 | IDL1-HSL2 signalling affects the dynamics of root cap sloughing. IDL1-activated
HSL2 regulates the dynamics of sloughing events, through regulation of PCD and cell wall
remodelling genes, and triggering of COL and LRC stem cell divisions.

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				611
Tissue	Marker ¹	# genes ² with highest expression ³	P-values	FDR adjusted ^p ‡ ² values
LRC	LRC	20	9.0 E-04	3.1E-02
COL	pet111	29	1.16E-06	8.14E-06
QC	AGL42	8	0.908	0.999614
Epidermis (hair cells)	COBL9	12	0.494	0.999
Epidermis (non-hair cells)	gl2	1	0.999	0.999
Cortex	CORTEX	10	0.737	0.999616
Endodermis	scr5	1	0.999	0.999
				617

¹Makers used for sorting cells of a particular tissue prior to RNA isolation (Nawy et al. 2005)

²Genes down regulated in the *hsl2* mutant relative to Col-0, cf. Supplementary Table 2.

620 ³Based on expression data in radial tissue types (Nawy et al. 2005). Genes were assigned

to the tissue with the highest expression level across all tissues.