

1 **Root electrotropism in *Arabidopsis* does not depend on auxin distribution but**
2 **requires cytokinin biosynthesis**

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15 **AUTHOR CONTRIBUTIONS**

16 M.Sa., N.O. and G.S. conceived and designed the experiments. M.Sa. contributed to the
17 development of the V-box, performed and analysed most of the electrotropism assays. N.O.
18 contributed to the development of the V-box, performed and analysed some of the WT
19 electrotropism assays. D.T. contributed to the development of the V-slide and performed some
20 of the electrotropism assays. M.Sc. performed the root excision experiments and the
21 electrotropism assays for *cyp735a1*, and analysed the data. N.K. contributed to the
22 development of the V-slide and performed the *R2D2* and *WAVE131Y* imaging. S.J.K.
23 performed the heat-treatment experiments. M.Sa. and G.S. prepared the manuscript, with all
24 the authors contributing to the Methods section.

25

26 **ABSTRACT**

27 An efficient foraging strategy for plant roots relies on the ability to sense multiple physical and
28 chemical cues in soil and to reorient growth accordingly (tropism). Root tropisms range from
29 sensing gravity (gravitropism), light (phototropism), water (hydrotropism), touch
30 (thigmotropism) and more. Electrotropism, also known as galvanotropism, is the phenomenon
31 of aligning growth with external electric fields and currents. Although observed in a few species
32 since the end of the 19th century, the molecular and physical mechanism of root electrotropism
33 remains elusive, limiting the comparison to more defined sensing pathways in plants.

34 Here we provide a first quantitative and molecular characterisation of root electrotropism
35 in the model system *Arabidopsis thaliana*, showing that it does not depend on an asymmetric
36 distribution of the plant hormone auxin, but that instead it requires the biosynthesis of a second

37 hormone, cytokinin. We also show that the dose-response kinetics of the early steps of root
38 electrotropism follows a power law analogous to the one observed in some physiological
39 reactions in animals.

40 A future full molecular and quantitative characterisation of root electrotropism would
41 represent a step forward towards a better understanding of signal integration in plants, and an
42 independent outgroup for comparative analysis of electroreception in animals and fungi.

43

44 INTRODUCTION

45 Plant roots navigate the complex soil environment in search of water and nutrients, through
46 various sensing mechanisms reorienting their growth towards or away from signal sources
47 (tropism) (Muthert et al., 2020). For example, hydrotropism redirects growth towards high
48 moisture (Miyazawa and Takahashi, 2019), (negative) phototropism redirects away from light
49 sources (Kutschera and Briggs, 2012), and gravitropism induces growth downwards, following
50 the gravity vector (Su et al., 2017). The integration of overlapping and frequently contradicting
51 molecular and physical signals is as critical for plant roots as it is for other soil organisms.

52 Although many molecular aspects of a few root tropisms have been understood (Muthert
53 et al., 2020), several key sensing mechanisms remain elusive. One of these is the capacity of
54 plant roots to sense electric fields (electrotropism or galvanotropism) (Navez, 1927). The local
55 physical and chemical properties of soil determine the presence of mobile electrical charges
56 and the generation of spontaneous electric fields (Pozdnyakov and Pozdnyakova, 2002). For
57 example, electrostatic fields can appear from charge separation in minerals like clay (Ward,
58 1990), by electrokinetic conversion caused by a conducting fluid like water flowing through
59 rocks (Revil et al., 2003) or by local accumulation of mineral ions important for plant
60 metabolism such as ammonium and nitrate ions. Local electric fields in soil can have biological
61 origin as well, for example from negatively charged bacteria (Olitzki, 1932) or from ions and
62 charged molecules released by microorganisms (Chabert et al., 2015) or plant roots
63 (Takamura, 2006). All this suggests that transient electrostatic fields in soil encode unique
64 information regarding the localisation of water, micronutrients and organisms, and it is
65 plausible that a sensing mechanism to detect such signal provides a selective advantage.

66 In fact, the reception of electric fields (electroreception) has been observed in vertebrates
67 (Crampton, 2019) and invertebrates (Clarke et al., 2013), including common model systems
68 such as *C. elegans* (Sukul and Croll, 1978) and *Dyctostylium* (Shanley et al., 2006), as well
69 as in fungi (McGillivray and Gow, 1986). Interestingly, electric sensing structures have been
70 identified only in a few aquatic animal species (Peters et al., 2007) and more recently in
71 bumble-bees (Sutton et al., 2016).

72 First recorded in 1882 (Elfving, 1882) and rediscovered at the start of the 20th century
73 (Ewart and Bayliss, 1905), root electrotropism has been studied sporadically in maize (*Zea*

74 *mays*), peas (*Pisum sativum*) and bean (*Vigna mungo*) but with contradicting results
75 (Wolerton et al., 2000). Crucially, the anatomical and molecular details of sensing electric
76 fields are still largely unknown in roots. A quantitative description of root electrotropism's
77 kinetics is also missing, preventing a comparative analysis with animal electroreception and
78 electrotaxis.

79

80 **RESULTS**

81 *A novel root electrotropism assay*

82 To study root electrotropism in the plant model system *Arabidopsis thaliana*, we developed
83 a new setup to stimulate, image and quantitate electrotropism in its primary root. Briefly, roots
84 were grown vertically in a transparent chamber (V-box) containing a buffered liquid medium
85 and two immersed electrodes connected to a power supply to generate a uniform electric field
86 and an ionic current perpendicular to the growing roots (Fig. 1a and Methods).

87 To maintain constant temperature and minimise the pH gradient generated by electrolysis,
88 the liquid medium was continuously circulated in a closed loop between the V-box, a cooled
89 water bath and a 2-litre reservoir bottle (Supplemental Fig. S1 and Methods). In each V-box,
90 the primary roots of five seedlings were imaged every 10 minutes with a camera mounted in
91 front of the V-box (Supplemental Fig. S1 and Methods). We measured the actual field
92 generated in the medium by immersing voltmeter probes in the two neighbouring positions of
93 each seedling: as expected, the field measured in the medium was lower than the nominal
94 imposed in air between the two electrodes, but still relatively uniform across the five positions
95 (Supplemental Fig. S2a). At the same time, we measured the current passing through the
96 circuit (Supplemental Fig. S2b and Methods).

97 To quantify the root response to the applied electric field, we took an image of the roots
98 every 10 minutes and measured the angle between the root tip and the vertical gravity vector
99 (Fig. 1b).

100 To confirm that the circulation of the liquid medium was effective in damping any pH
101 gradient created by electrolysis and in eliminating any chemotropic effect, we imposed a field
102 of 1.5 V/cm in a V-box without plants for one hour while maintaining liquid circulation. We then
103 turned off the field and the liquid circulation, immediately positioned the plants in the V-box
104 and imaged the roots for the next 80 minutes: we did not observe any significant deviation in
105 root growth direction (t-test between 1.5 V/cm and 0 V/cm at 80 m, p=0.859), compared to
106 roots in a V-box that never experienced the electric field (Supplemental Fig. S2c), indicating
107 that no significant pH gradient was left in the medium.

108

109 *Root tip reorientation in external electric fields*

110 We measured the response of wild-type (WT) *Arabidopsis* primary roots to a continuous
111 electric field. The distribution of reorientations to a range of fields intensities between 0.5 V/cm
112 and 2.5 V/cm shows a quick tropism towards the negative electrode, or cathode (Fig. 1c).

113 We wondered how much of this effect was due to trivial electrostatic, *i.e.* the physical pull
114 towards the negative electrode due to a hypothetical net positive charge accumulated on the
115 root tip, rather than a more complex biological response involving molecular signalling. To
116 address this, we deactivated the roots by immersing them in a 50 °C bath for 10 min, until
117 growth and gravitropism were suppressed (0/9 roots growing and bending to gravity after
118 exposure to 50 °C, *vs.* 10/10 after exposure to 23 °C), transferred to the V-box and exposed
119 them to a 2.0 V/cm electric field: the root response shows that this simple treatment was
120 sufficient to completely inhibit electrotropism when compared to roots kept at a standard 23
121 °C temperature (t-test between 50 °C and 23 °C at 5 h, $p<0.001$), strongly suggesting that
122 electrostatic alone could not explain the root response and that this is in fact a biological
123 phenomenon (Supplemental Fig. S3).

124

125 *Response curves*

126 The progressively sharper root tip reorientation as the field intensity was increased (Fig.
127 1c) suggests that the sensing mechanism is not acting as a simple on/off switch but that it can
128 distinguish electric fields of different strengths. To quantitate this, we plotted the orientation
129 angle at 5 h (“response”) as a function of the electric field intensity (“stimulus”), and found a
130 best fit with a power function with exponent 0.45 (Fig. 2a, left), indicating that the resolution of
131 the sensor is higher at low-intensity stimuli (steep response curve) than at high-intensity stimuli
132 (shallow response curve). The analogous response curve as a function of the measured
133 current intensity is best fit with a power function with exponent 0.33 (Fig. 2a, right).

134 Since the root tip reorientations did not show any obvious overshoot (Fig. 1c), we looked
135 more closely at the angular velocity: the maximum average velocity in WT roots was reached
136 when the root tip was between 10 and 20 degrees orientation to the gravity vector and then
137 progressively decreased as the root tip approached its maximum reorientation (Supplemental
138 Fig. S4a). This indicates that the mechanism is able to sense and respond differently to the
139 changing relative orientation of the tip with the external electric field, and to progressively slow
140 down the root tip rotation as it approaches the target orientation. Moreover, the maximum
141 angular velocity appears to be roughly proportional to the electric field strength (Fig. 2b, left),
142 although this is much less evident as a function of the electric current (Fig. 2b, right).

143

144 *Root tips are not damaged*

145 Early reports noted that protracted exposure to external fields could cause physical
146 damage to plant root tips (Wawrecki and Zagórska-Marek, 2007). To control whether this was
147 the case in our experimental conditions, we developed a simple chambered slide (V-slide) to
148 be mounted on a standard confocal microscope stage (Supplemental Fig. S5a and Methods)
149 with electrodes on the chamber's sides and a circulating liquid medium for temperature control
150 similar to the one implemented in the V-box (Supplemental Fig. S5b). Seedlings from the
151 *Arabidopsis* transgenic line constitutively expressing the yellow fluorescent cell-membrane
152 marker WAVE 131Y (Geldner et al., 2009) were mounted on the V-slide and imaged at cellular
153 resolution while exposed to a 1.0 V/cm electric field (Methods). The field of view in our time-
154 lapse images comprises the meristem, the transition zone and the distal elongation zone, but
155 no cellular pattern perturbation was noticeable in any of these regions when compared to roots
156 not exposed to the field (Supplemental Fig. S6). Also, the time-lapse suggests that asymmetric
157 cell expansion in the elongation zone is causing the bending, as expected from other examples
158 of root tropism (Gilroy, 2008).

159 To further confirm that roots exposed to the electric field are not damaged and maintain
160 gravitropic response, we monitored root tips for 2 h after the electric field had been turned off,
161 observing a clear gravitropic behaviour (Supplemental Fig. S7).

162

163 *Regions of competence*

164 Since the root bending occurs in the elongation zone, we wanted to identify the region
165 responsible for sensing the electric field. We excised distal fragments of the root at 125 µm,
166 300 µm, 400 µm and 500 µm from the tip (Fig. 3a), and then exposed the cut root to 1.5 V/cm
167 (Methods).

168 In the first 2 h (Fig. 3b left), roots cut at 400 µm from the tip turned to angles
169 indistinguishable from uncut roots exposed to the same field (Wilcoxon between 400 µm cut
170 and uncut at 1.5 V/cm at 2 h, $p=0.211$), while roots cut at 500 µm from the tip did not turn and
171 at 2 h showed orientations indistinguishable from uncut roots not exposed to a field (t-test
172 between 500 µm cut at 1.5 V/cm and uncut at 0 V/cm at 2 h, $p=0.164$). These results indicate
173 that the 400 µm distal fragment is not necessary for the early (2 h) electrotropic response,
174 while the 500 µm distal fragment is; since the elongation zone involved in the bending spans
175 an extended region proximal to the 500 µm cut, we conclude that in *Arabidopsis* roots the
176 region between 400 µm and 500 µm from the tip is necessary for early sensing an external
177 electric field.

178 Between 2 h and 5 h of exposure (Fig. 3b right), while roots cut at 300µm on average
179 continue to turn like the uncut roots (t-test between 300 µm cut and uncut at 1.5 V/cm at 5 hrs,
180 $p=0.071$), roots cut at 400 µm quickly fail to sustain their response and at 5 h they show tip

181 orientations on average different than those of uncut roots exposed to the same field (t-test
182 between 400 μm cut and uncut at 1.5V/cm at 5 hrs, $p<0.001$). These results indicate that the
183 300 μm distal fragment is not necessary to maintain the electrotropic response up to 5hrs,
184 while the 400 μm distal fragment is; we conclude that in *Arabidopsis* roots the region between
185 300 μm and 400 μm from the tip is necessary for prolonged sensing of the imposed electric
186 field.

187 Interestingly, previous studies suggested that the movement of Ca^{++} ions accumulated in
188 the mucilage at the very tip of the root might be involved in the electrotropic sensing (Marcum
189 and Moore, 1990). Since any excision above 125 μm from the tip essentially removes all the
190 mucilage from the root, our results disprove this hypothesis for *Arabidopsis*.

191

192 *Auxin distribution is not altered by the electric field*

193 The fact that roots without tips could still respond to the electric field shows that even a
194 major disruption of the stereotypical auxin redistribution mechanism is not sufficient to inhibit
195 electrotropism.

196 On the other hand, an asymmetric accumulation of auxin is required for asymmetric cell
197 elongation and root bending in some tropisms, as suggested by the classic Chodlony-Went
198 model (Thimann and Went, 1937). To test whether this model applied to electrotropism, and
199 whether the external electric field is sufficient to induce an asymmetric distribution of auxin in
200 the root, we exposed roots expressing the auxin-sensitive fluorescent reporter *R2D2* (Liao et
201 al., 2015) to a field of 1.0 V/cm for 30 minutes, before quickly mounting them on a microscope
202 slide and imaging them with a confocal microscope (Methods). A ratiometric quantification of
203 *R2D2* signal (Kral et al., 2016) in each epidermal cell (Fig. 4a and Methods) showed that the
204 average auxin response measured in the epidermal cells on the side facing the negative
205 electrode and on the side facing the positive electrode were statistically indistinguishable (Fig.
206 4b; Methods), both in the distal (Wilcoxon test, $p=0.696$) and the proximal (Wilcoxon test,
207 $p=0.843$) region of the root, indicating that auxin is not asymmetrically distributed after
208 exposure to the electric field.

209 Moreover, both sides were also indistinguishable from the average between the two sides
210 in roots not exposed to the field (mock), both in the distal (Wilcoxon test between the negative
211 exposed side and mock, $p=0.775$; and between the positive exposed side and mock, $p=0.881$)
212 and in the proximal (Wilcoxon test between the negative exposed side and mock, $p=0.041$;
213 and between the positive exposed side and mock, $p=0.098$) regions.

214

215 *Auxin transport is not necessary for early response*

216 Previous work indicated that auxin transport inhibitors can inhibit electrotropism response
217 in maize roots (Ishikawa and Evans, 1990). To verify this in *Arabidopsis*, and further explore

218 the role of auxin in root electrotropism, we used N-1-naphthylphthalamic acid (NPA) to inhibit
219 polar auxin transport (Sabatini et al., 1999) while exposing the roots to the electric field. We
220 tested 0.1 μ M, 1.0 μ M and 10 μ M NPA and found that 10 μ M NPA was the lowest
221 concentration of NPA to inhibit gravitropism (Methods), a known auxin-dependent tropism.

222 We pre-treated the seedlings for 3 hours in liquid medium containing 10 μ M NPA
223 (Methods) and then transferred them to a V-box containing 10 μ M NPA in the medium and the
224 reservoir. In the first 1 hour of field exposure (Fig. 4c left) the NPA-treated roots reoriented at
225 angles indistinguishable from those of untreated ones (Wilcoxon test between NPA-treated
226 and untreated at 1 h, $p=0.350$). From 1 to 5 hours of exposure (Fig. 4c right), NPA-treated
227 roots respond less than the untreated (t-test between NPA-treated and untreated at 5 h,
228 $p<0.001$), but still significantly more than the untreated and not exposed to the field (t-test
229 between untreated at 1.5 V/cm and 0 V/cm at 5 h, $p=0.001$). These results indicate that auxin
230 polar transport is not necessary for an early electrotropic response, but might play a role in
231 maintaining a long-term orientation.

232 To further explore the role of auxin transport, we tested the mutants of PIN2, an auxin
233 cellular exporter (Chen et al., 1998), and mutants of AUX1, an auxin cellular importer
234 (Marchant et al., 1999). Since *pin2* and *aux1* mutants do not respond to gravity, to obtain a
235 sufficient number of roots growing vertically in preparation for the electrotropic assay, we
236 wrapped the sides and bottom of the nursery boxes with aluminium foil to induce negative root
237 phototropism towards the bottom of the box (Methods). The same setup was used to
238 germinate WT plants to be compared with these to mutants.

239 Roots of *pin2* mutants (Fig. 4d) showed a significant response (paired Wilcoxon between
240 the time-points 0 h and 2 h, $p<0.01$; and between the time-points 0 h and 5 h, $p<0.01$), but
241 weaker than WT (t-test between WT and *pin2* at 2 h, $p<0.01$; and between WT and *pin2* at 5
242 h, $p<0.01$) in the same conditions. Their angular velocity did not show an obvious decrease
243 before reaching the target orientation, as with WT roots (Supplemental Fig. S4b).

244 Analogously, roots of *aux1* mutants (Fig. 4d) showed a significant early response (paired
245 t-test between the time-points 0 h and 2 h, $p<0.01$), but weaker than WT (t-test between WT
246 and *pin2* at 2 h, $p<0.01$; t-test between WT and *pin2* at 5 h, $p<0.01$) in the same conditions.
247 Interestingly, *aux1* roots on average failed to maintain their orientation for longer period (paired
248 t-test between the time-points 0 h and 5 h, $p=0.051$). Moreover, their angular velocity on
249 averaged decreased while approaching the final orientation, as with WT roots (Supplemental
250 Fig. S4b).

251 Taken together, these results suggest that although auxin transport seems to play a role
252 in maintaining a sustained response to the electric field, it is not necessary in triggering early
253 electrotropism.

254

255 *Cytokinin biosynthesis is necessary for electrotropism*

256 Given our results suggesting a limited role for auxin during electrotropism, we wondered
257 which other plant hormones might be involved instead. Root hydrotropism, or the growth
258 towards high concentration of water, was also previously shown to be largely independent
259 from auxin distribution in *Arabidopsis* (Shkolnik et al., 2016), while it requires biosynthesis and
260 asymmetric distribution of cytokinin (Chang et al., 2019). Drawing an analogy with
261 hydrotropism, we considered the possibility that electrotropism might act through cytokinin as
262 well.

263 To test this hypothesis, we analysed root electrotropism in triple mutants of *AtIPTs*, a
264 family of adenosine phosphate-isopentenyltransferases required for the first step of isoprenoid
265 cytokinin biosynthesis (Miyawaki et al., 2006; Kamada-Nobusada and Sakakibara, 2009).
266 Within the tested mutants, although a high degree of redundancy is expected among the
267 *AtIPTs*, we found two distinct behaviours: the triple mutants *atipt1,3,5* and *atipt3,5,7* both
268 responded strongly to the electric field (Fig. 5a) (t-test between 1.5 V/cm and 0 V/cm at 5 h,
269 p<0.001 in both cases), while the triple mutants *atipt1,3,7* and *atipt1,5,7* responded well in the
270 first 2 h of exposure (Fig. 5b, left) (t-test between 1.5 V/cm and 0 V/cm at 2 h, p<0.001 for
271 both mutants) but showed a much weaker, although still significant, response at 5 h (Fig. 5b
272 right) (t-test between 1.5 V/cm and 0 V/cm at 5 h, p<0.01 for *atipt1,5,7* and p<0.001 for
273 *atipt1,3,7*; t-test between WT and mutant both at 1.5 V/cm at 5 h, p<0.001 for *atipt1,5,7* and
274 p<0.01 for *atipt1,3,7*). These results suggest that cytokinin biosynthesis is in part required for
275 long-term root electrotropism, and that for this phenotype *AtIPT1* and *AtIPT7* dominate their
276 family in a redundant way (the response is the weakest when both are mutated).

277 To confirm the role of cytokinin, we also tested the requirement for *CYP735A1*, a
278 cytochrome P450 monooxygenase enzyme acting downstream of *AtIPTs* and necessary for
279 the biosynthesis of the *trans*-zeatin (tZ) variation of cytokinin (Takei et al., 2004). Interestingly,
280 roots of *cyp735a1* mutants completely failed to respond to the electric field (Fig. 5c), with tip
281 orientations indistinguishable from that of WT roots not exposed to the field (t-test between
282 *cyp735a1* exposed to 1.5 V/cm and WT not exposed at 5 h, p=0.748). Crucially, this phenotype
283 could be rescued with the addition of the cytokinin *trans*-Zeatin (tZ) to the medium (Fig. 5c) (t-
284 test between *cyp735a1* +10nM tZ at 1.5 V/cm and 0 V/cm, p<0.001), while the same treatment
285 did not affect WT response (Fig. 5c).

286 Taken together, these results indicate that cytokinin is necessary for electrotropism in
287 *Arabidopsis* roots.

288 To further investigate a possible parallel between the molecular pathways involved in
289 electrotropism and hydrotropism, we tested mutants of *MIZU-KUSSEIN1* (*MIZ1*), which is
290 necessary for hydrotropism (Kobayashi et al., 2007) and its functional cytokinin asymmetric
291 distribution in roots (Chang et al., 2019). Translational fusion reporters have shown *MIZ1*

292 localisation in the lateral root cap and the cortex of meristem and elongation zone (Dietrich et
293 al., 2017): although the root tip is not necessary for hydrotropic response (Dietrich et al., 2017),
294 this requires MIZ1 in the transition zone (Dietrich et al., 2017).

295 When we exposed roots of *miz1* mutants to 1.5 V/cm (Fig. 6) they showed an unperturbed
296 electrotropic response in the first 2 h (Wilcoxon test between *miz1* and WT at 2 h, $p=0.859$)
297 and perhaps a weakened response at 5 h, although with only a weak statistical significance
298 (t-test between *miz1* and WT at 5 h, $p=0.033$), indicating that MIZ1 is not necessary for early
299 root electrotropism.

300 Taken together, these surprising results indicate that cytokinin plays an important role in
301 root electrotropism, but that the underlying molecular pathway differs early on from that of
302 hydrotropism.

303

304 **DISCUSSION**

305 The capability of plant roots to sense and combine numerous physical and chemical signals
306 in soil is quite extraordinary, notably in the absence of a centralised processing system. The
307 more we understand about the physical and molecular mechanism behind the various root
308 tropisms, the closer we will get to a complete understanding of signal integration in plants. In
309 this work we focused on the little-studied phenomenon of root electrotropism and present its
310 first quantitative characterisation in the flowering plant model system *Arabidopsis thaliana*.

311 A non-trivial observation from this work is that plant roots respond to weak (order of 1 V/cm
312 and 100 μ A) external electric fields and currents in a progressive way, with an increasingly
313 stronger tendency to align with the field as the field and current intensities increase. This is
314 interesting, because it reveals a sophisticated way to discriminate between highly charged
315 particles (e.g. ions, micro-organisms, other plant roots) and weakly charged ones. At the same
316 time, the absence of overshooting also points to a mechanism that perhaps can be modelled
317 along the lines of a proportional–integral–derivative control system (Chevalier et al., 2019).
318 Moreover, we show that the kinetics (response curve) of root electrotropism follows a power-
319 law with exponents similar to those that have been traditionally measured in physiological
320 responses to external stimuli in animal systems (Stevens, 1970): our results suggest that a
321 power-law response to external stimuli might be a universal feature across life kingdoms, and
322 that it does not require a nervous system. Moreover, it might reveal constraints on the genetic
323 architecture of the underlying sensory system (Adler et al., 2014).

324 A second, perhaps unexpected, result from this work is the identification of two regions in
325 *Arabidopsis* roots required for the detection of the field: the section between roughly 400 μ m
326 and 500 μ m from the tip is necessary for the early detection (within 2 h from exposure), while
327 the section between roughly 300 μ m and 400 μ m is necessary for a prolonged detection (up
328 to 5 h from exposure). Interestingly, these sections seem to correspond to the well-

329 characterised anatomical transition zone, between the meristem and the elongation zone in
330 roots, which has been previously involved in root sensing (Muthert et al., 2020). In future
331 works, it will be crucial to discern whether the electrotropism mechanism depends on a
332 relatively large region like the transition zone or if it can be narrowed to a more specific cell
333 population therein.

334 Finally, we present evidence against the assumption that auxin asymmetric distribution is
335 required for electrotropism in *Arabidopsis*, similarly to what found for hydrotropism (Shkolnik
336 et al., 2016) and phototropism (Kimura et al., 2018). Instead, we show that cytokinin is required
337 for a full electrotypic response, although its response-specific regulation seems to act through
338 a different pathway than the one established for hydrotropism.

339 Overall, our results show that root electrotropism requires a sensing mechanism likely
340 localised in the transition zone, a limited role for auxin but an important role for cytokinin. The
341 latter could be involved in a signalling or regulatory mechanism in the transition zone, in an
342 actuator mechanism (tissue bending) in the elongation zone, or both.

343 Cytokinins have been shown to regulate root meristem activity and size by controlling cell
344 proliferation (Beemster and Baskin, 2000) and the developmental progression from
345 proliferation to elongation in the transition zone (Iorio et al., 2007). It has been suggested that
346 root bending in hydrotropism is based on asymmetric distribution of cytokinin in the meristem
347 to induce asymmetric cell proliferation (Chang et al., 2019). This seems an unlikely
348 mechanism for electrotropism, because roots without meristem (a 400 μ m segment cut from
349 the tip) still respond to the electric field, suggesting that root bending in this case does not
350 depend much on cell proliferation. An alternative mode of action for cytokinins during
351 electrotropism could be based on its regulatory action in the transition zone, where an
352 asymmetric delay in elongation would result in root bending. Although we have shown that
353 MIZ1 is not required for root electrotropism, it is still possible that a MIZ1-independent
354 mechanism could generate an asymmetry in cytokinin distribution in the transition zone.

355 More recently, cytokinins have also been involved in stress response through regulation
356 of downstream factors and through crosstalk with other hormones (Li et al., 2021; Wu et al.,
357 2021). This signalling role could be relevant during elecrotpism, especially since the
358 transition zone has often been associated with signal integration in the root (Baluška et al.,
359 2010).

360 Future work on root electrotropism should focus on testing these hypotheses regarding
361 the role of cytokinin and on illuminating the still unknown molecular mechanism involved in
362 sensing an electric field or current.

363

364 **METHODS**

365 *Plant material*

366 Wild-type and mutant *Arabidopsis thaliana* plants were all from the Columbia (Col-0) ecotype; the following mutant alleles were used: *aux1-7* (NASC id 9583) for *aux1*; *eir1-1* (NASC id 8058) for *pin2*; SALK_093028C (NASC id N654306) for *cyp735a1*, *miz1-1* for *miz1* (courtesy of Prof. A. Kobayashi); *atipt* triple mutants as previously described with *atipt1-1*, *atipt3-2*, *atipt5-2*, *atipt7-1* (Miyawaki et al., 2006) (courtesy of Prof. O. Leyser).

371 The fluorescent line *WAVE131Y* is expressing *pUBQ10::WAVE131:YFP* (NASC id N781535); the fluorescent line *R2D2* is expressing *RPS5A-mDII-ntdTomato*, *RPS5A-DII-n3xVenus* (courtesy of Dr. Teva Vernoux).

374 Seeds were imbibed in water and kept in the dark for 2 days at 4 °C, in order to synchronise germination. All seeds were surface sterilised using 50% Haychlor bleach and 0.0005% Triton X-100 for 3 minutes and then rinsed 6 times with sterilised milliQ water. Seed germination protocols are described in the experiment-specific Methods sections below. Unless otherwise specified, all experiments were conducted with primary roots of seedlings 5-8 days post-germination, with roots approximately of the same length to be in the field of view of the V-box camera.

381

382 *Electrotropism assay (V-box)*

383 Seeds were sown individually inside PCR tubes filled with 1X MS gel medium: 0.44% Murashige and Skoog (MS) Basal medium (Sigma-Aldrich, M5519), 0.5% sucrose, 0.05% MES hydrate (Sigma-Aldrich M8250), 0.8% agar (Sigma-Aldrich 05040), pH adjusted to 5.7 with TRIS HCl (Fisher-Scientific 10205100). The PCR tubes had their end cut out to allow the root to grow through, and placed in a 3D-printed (Ultimaker 2+) holder inserted in a Magenta box (Sigma-Aldrich V8380). The Magenta box was filled with 150 ml of 1/500X MS liquid medium (0.00088% MS Basal medium, 0.5% sucrose, 0.05% MES hydrate, pH adjusted to 5.7 with TRIS HCl) to reach the end of the PCR tubes. These germination, or “nursery”, boxes were placed in a growth chamber at 22 °C, with a 16 h/8 h light/dark photoperiod and light intensity 120 $\mu\text{mol m}^{-2} \text{s}^{-1}$.

393 In preparation for the electrotropism assay, each PCR tube containing a single seedling was transferred to a modified 3D-printed holder in a Magenta box filled with 1/500X MS liquid medium. The modified module consisted of a main body with five holes for the PCR tubes containing the seedlings, and two side clips to position the electrodes consisting of platinum-iridium (Platinum:Iridium = 80:20; Alfa Aesar 41805.FF) foils (Fig. 1a), which were connected to an external power supply. In this paper, we refer to the Magenta box and its holder as the “V-box”. In addition to the five holes for the plants, four extra holes were designed: two at each end of the front side to pump the medium out of the V-box right on top of the two electrodes,

401 and two on the back to pump the medium in, using a tubing system and peristaltic pumps
402 (Verdeflex AU R2550030 RS1) to circulate the medium at a speed of 1 ml/sec to and from a
403 2 l reservoir bottle (Extended Data Fig. 1). This configuration was designed and tested to
404 ensure a slow and symmetric flow of medium in the box, eliminating any biased effect of the
405 flow on the roots, which was confirmed by the control experiments at 0 V/cm, performed with
406 the pump circulating the medium at the same speed as in the exposure experiments. Crucially,
407 between the V-box and the reservoir bottle, the tubings were immersed in a cooled water bath
408 (Grant Instruments, LTDGG) maintained at the constant temperature of 19 °C (Extended Data
409 Fig. 1), which was enough to maintaining the medium inside the V-box at a constant 22 °C, as
410 measured. All electrotropic experiments were performed at constant illumination.

411 The electric field was generated with a power supply attached to the platinum-iridium foil
412 electrodes that were immersed in the liquid medium in the V-box. Standard electric wires were
413 soldered on the top of two electrodes, always kept outside the liquid medium to avoid
414 contaminants from the solder. The voltage was set constant on the power supply, while the
415 current was measured independently with a multimeter in series.

416 In order to record the movement of the roots over time, a Raspberry Pi camera module V2
417 (913-2664) connected to a Raspberry Pi board module B+ (137-3331) was used. The
418 Raspberry Pi was programmed to take a picture every 10 minutes, using the command *crontab*
419 in the local Raspbian OS.

420 WT electrotropism assays (Fig. 1) were performed with the following sample sizes N and
421 number of replicates R: 0 V/cm, N=10, R=2; 0.5 V/cm, N=9, R=2; 1.0 V/cm, N=8, R=2; 1.5
422 V/cm, N=21, R=5; 2.0 V/cm, N=18, R=6; 2.5 V/cm, N=20, R=7; foil-wrapped and 1.5 V/cm
423 (control for *pin2* and *aux1*), N=15, R=3.

424 Mutants electrotropism assays (Fig. 4,5,6) were performed with the following sample sizes
425 N and number of replicates R: *aux1* at 1.5 V/cm, N=18, R=4; *aux1* at 0 V/cm, N=14, R=3; *pin2*
426 at 1.5 V/cm, , N=13, R=4; *pin2* at 0 V/cm, , N=13, R=3; *atipt3,5,7* at 1.5 V/cm, N=15, R=3;
427 *atipt3,5,7* at 0 V/cm, N=10, R=2; *atipt1,3,5* at 1.5 V/cm, N=15, R=3; *atipt1,3,5* at 0 V/cm, N=10,
428 R=2; *atipt1,5,7* at 1.5 V/cm, N=14, R=3; *atipt1,5,7* at 0 V/cm, N=10, R=2; *atipt1,3,7* at 1.5
429 V/cm, N=14, R=5; *atipt1,3,7* at 0 V/cm, N=10, R=2; *cyp735a1* at 1.5 V/cm, N=20, R=4; *miz1*
430 at 1.5 V/cm, N=9, R=2; *miz1* at 0 V/cm, N=10, R=2

431 The control for medium circulation efficiency (Fig. S2) was performed with the following
432 sample sizes N and number of replicates R: 1.5 V/cm, N=10, R=2; 0 V/cm, N=10, R=2.

433 The experiment showing gravitropism after 2 hours of electric field (Fig. S7) was performed
434 with a sample size N=10 and replicates R=2.

435

436 *High temperature treatment*

437 Wild-type Col-0 seeds were germinated and grown in the nursery boxes as described in
438 the Electrotropism Assay section. The boxes undergoing treatment were then immersed in a
439 water bath set at 50 °C, for 10 minutes. The PCR tubes containing the seedlings were then
440 transferred to a V-box, exposed to an electric field of 2.0 V/cm for 5 hours and imaged, as
441 described in the Electrotropism Assay section.

442 Electrotropism assays for treated roots were performed with the following sample sizes N
443 and number of replicates R: 23°C, N=18, R=6; 50°C, N=10, R=2.

444 For the gravitropism control, seedlings were treated with high temperature as described
445 above and then transferred on 1X MS agar plates, to complete the assay as described in the
446 Gravitropic Assay section.

447 Gravitropic assays for treated roots were performed with the following sample sizes N and
448 number of replicates R: 23°C, N=10, R=2; 50°C, N=9, R=1).

449

450 *Electrotropism on microscope (V-slide)*

451 Seeds of the transgenic reporter line *WAVE131Y* (see Plant Material section) were
452 germinated on 1X MS agar medium (0.44% MS Basal medium, 0.5% sucrose, 0.05% MES
453 hydrate, pH adjusted to 5.7 with TRIS HCl, 0.8% agar) in square plates kept vertical in a
454 growth chamber at 22 °C, with a 16 h/8 h light/dark photoperiod.

455 At 2 days post-germination the seedlings were mounted on the V-slide with 1/500X MS
456 liquid medium (0.00088% MS Basal medium, 0.5% sucrose, 0.05% MES hydrate, pH adjusted
457 to 5.7 with TRIS HCl). The V-slide was then connected to the medium perfusion system
458 circulating the same 1/500X MS medium, and the V-slide's electrodes were connected to the
459 power supply (Voltcraft PS-1302-D).

460 Imaging was performed on a Leica SP5 laser scanning confocal microscope with 20X air
461 objective, while the liquid medium was circulated and the electric field maintained constant.
462 Fluorophore was excited with the 514 nm Argon laser line and the emission collected with
463 PMT detector at 524-570 nm. Images were collected at intervals of 2 minutes.

464

465 *Root tip excisions*

466 Seeds were germinated in nursery boxes as described.

467 3 days post-germination seedlings still in their PCR tubes, as previously described, were
468 transferred to hard 1X MS agar medium (0.44% MS Basal medium, 0.5% sucrose, 0.05%
469 MES hydrate, pH adjusted to 5.7 with TRIS HCl, 5.0% agar) where the roots were manually
470 dissected using a dental needle (Sterican, 27G) under a dissecting microscope (Nikon
471 SMZ1000) at 180X magnification, following the published protocol (Kral et al., 2016).

472 After root tip excision, the PCR tubes containing the seedlings were immediately moved
473 into the V-box for the electrotropism assay.

474 Electrotropism assays on excised roots were performed with the following sample sizes N
475 and number of replicates R: 125 μ m, N=16, R=4; 300 μ m, N=7, R=2; 400 μ m, N=10, R=2; 500
476 μ m, N=10, R=2; uncut at 0 V/cm, N=10, R=2; uncut at 1.5 V/cm, N=21, R=5.

477

478 *R2D2 reporter*

479 Seeds of the transgenic reporter line *R2D2* (see Plant Material section) were germinated
480 in nursery boxes and at 3days post-germination exposed to 1.0 V/cm for 30min in V-boxes,
481 as described in the Electrotropism Assay section. After exposure, *R2D2* roots were quickly
482 mounted on standard microscope slides with sterile deionised water and imaged using Leica
483 SP5 laser scanning confocal microscope, with 63X water immersion objective.

484 *R2D2* expresses two versions of a protein that forms a complex with auxin: DII-
485 n3xVENUS, which is degraded within minutes upon binding with auxin; mDII-ntdTOMATO,
486 which contains a modified, non-degradable, version of DII. Both of the proteins are localised
487 in the nucleus. We followed the published protocol (Liao et al., 2015) to separately collect the
488 emission from mDII-ntdTOMATO (Extended Data Fig. 6a) and DII-n3xVENUS (Extended Data
489 Fig. 6b).

490 For each root, a mean background was defined as the average pixel intensity in the mDII
491 channel of a 80x80 pixels corner of the field of view not occupied by the root. The mean
492 background was then subtracted form all pixel intensities in both channels (mDII and DII). We
493 manually segmented with FIJI (Schindelin et al., 2012) the most visible cell nuclei in the
494 epidermis of each root, both in the distal (12 roots in mock conditions and 16 roots exposed
495 to the field) and in the proximal (11 roots in mock conditions and 13 roots exposed to the field)
496 regions of the root tip, and quantified the average pixel intensity (corrected after background
497 subtraction) for each segmented nucleus. Finally, we calculated the natural log of the ratio
498 between the average mDII (non-degradable, auxin-independent) and DII (degradable, auxin-
499 dependent) signals in each nucleus and mapped it on top of the root image (Extended Data
500 Fig. 6d).

501 For each root, we calculated the average and standard deviation of these ratios, for
502 epidermal nuclei facing the anode or the cathode (Fig. 4b).

503 Sample size as following: exposed to E field and imaged in distal region, N=16; exposed
504 to E field and imaged in the proximal region, N=13; not exposed to E field and imaged in the
505 distal region, N=24; not exposed to E field and imaged in the proximal region, N=22.

506

507 *NPA treatment*

508 To find the minimum concentration of N-1-naphthylphthalamic acid (NPA) that inhibits
509 gravitropism, seeds were germinated on 1X MS agar medium (0.44% MS Basal medium, 0.5%
510 sucrose, 0.05% MES hydrate, pH adjusted to 5.7 with TRIS HCl, 0.8% agar) and at 5-8 days
511 post-germination were transferred for 3 hrs into cell culture dishes containing 5 ml of 1/500X
512 MS liquid medium (0.00088% MS Basal medium, 0.5% sucrose, 0.05% MES hydrate, pH
513 adjusted to 5.7 with TRIS HCl) plus NPA at 0.1 μ M, 1.0 μ M and 10 μ M.

514 To quantify the effect of NPA on electrotropism, after being treated with a concentration of
515 10 μ M NPA for 3 hours as described above, the seedlings were transferred inside a V-box
516 and exposed to a 1.5 V/cm electric field as described in the Electrotropism Assay section.
517 Both the V-box and the reservoir bottle contained 10 μ M NPA throughout the experiment.

518 Sample sizes N and number of replicates R were the following: untreated, N=21, R=5; 10
519 μ M NPA treated, N=21, R=5.

520

521 *Cytokinin treatment*

522 Both the 1X MS agar medium (0.44% MS Basal medium, 0.5% sucrose, 0.05% MES
523 hydrate, pH adjusted to 5.7 with TRIS HCl, 0.8% agar) contained in the PCR tubes, and the
524 1/500X MS liquid medium (0.00088% MS Basal medium, 0.5% sucrose, 0.05% MES hydrate,
525 pH adjusted to 5.7 with TRIS HCl) in the nurseries, were supplemented with 10nM *trans*-zeatin
526 (Sigma-Aldrich Z0876) as previously suggested (Miyawaki et al., 2006). Still in their PCR
527 tubes, seedlings were transferred into V-boxes, filled with 1/500X MS liquid medium supplied
528 with 10nM *trans*-zeatin to conduct the electrotropism experiments.

529 Sample sizes N and number of replicates R were the following: WT + tZ, N=14, R=3;
530 *cyp735a1* +tZ at 1.5 V/cm, N=13, R=3; *cyp735a1* +tZ at 0 V/cm, N=14, R=3.

531

532 *Gravitropism assay*

533 In the high-temperature experiment, after the treatment the seedlings were transferred
534 onto 1X MS square agar plates (0.44% MS Basal medium, 0.5% sucrose, 0.05% MES hydrate,
535 pH adjusted to 5.7 with TRIS HCl, 0.8% agar).

536 In the NPA experiment, after the treatment the seedlings were transferred onto 1/500X MS
537 square agar plates (0.00088% MS Basal medium, 0.5% sucrose, 0.05% MES hydrate, pH
538 adjusted to 5.7 with TRIS HCl, 0.8% agar) containing the desired concentration of NPA.

539 In both cases, the plates were moved to a growth room (22 °C, 16 h/8 h light/dark
540 photoperiod), rotated by 90 degrees to position the roots in a horizontal orientation, and
541 monitored for root gravitropic response.

542

543 *Statistical analysis*

544 When comparing two samples of measurements, each distribution was first checked for
545 normality with the Shapiro-Wilk test with alpha-level = 0.05. Normal distributions were tested
546 with the two-tails Student's t-test without assuming equal variances (Welch two sample t-test);
547 if one of the two distributions was not normal, the non-parametric Mann-Whitney U (Wilcoxon)
548 test was used. Unless stated otherwise, all comparisons were performed assuming
549 independence (unpaired test). All statistical tests were performed with R.

550

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558

559 **COMPETING INTERESTS**

560 The authors declare no competing interests.

561

562 **REFERENCES**

563 **Adler M, Mayo A, Alon U** (2014) Logarithmic and Power Law Input-Output Relations in
564 Sensory Systems with Fold-Change Detection. *Plos Comput Biol* **10**: e1003781

565 **Baluška F, Mancuso S, Volkmann D, Barlow PW** (2010) Root apex transition zone: a
566 signalling-response nexus in the root. *Trends in Plant Science* **15**: 402–408

567 **Beemster GTS, Baskin TI** (2000) STUNTED PLANT 1 Mediates Effects of Cytokinin, But Not
568 of Auxin, on Cell Division and Expansion in the Root of *Arabidopsis*. *Plant Physiol* **124**: 1718–
569 1727

570 **Chabert N, Ali OA, Achouak W** (2015) All ecosystems potentially host electrogenic bacteria.
571 *Bioelectrochemistry* **106**: 88–96

572 **Chang J, Li X, Fu W, Wang J, Yong Y, Shi H, Ding Z, Kui H, Gou X, He K, et al** (2019)
573 Asymmetric distribution of cytokinins determines root hydrotropism in *Arabidopsis thaliana*.
574 *Cell Res* **29**: 984–993

575 **Chen R, Hilson P, Sedbrook J, Rosen E, Caspar T, Masson PH** (1998) The *Arabidopsis*
576 *thaliana* AGRAVITROPIC 1 gene encodes a component of the polar-auxin-transport efflux
577 carrier. *Proc National Acad Sci* **95**: 15112–15117

578 **Chevalier M, Gómez-Schiavon M, Ng AH, El-Samad H** (2019) Design and Analysis of a
579 Proportional-Integral-Derivative Controller with Biological Molecules. *Cell Syst* **9**: 338-
580 353.e10

581 **Clarke D, Whitney H, Sutton G, Robert D** (2013) Detection and learning of floral electric
582 fields by bumblebees. *Science* **340**: 66-69

583 **Crampton WGR** (2019) Electoreception, electrogenesis and electric signal evolution. *J Fish
584 Biol* **95**: 92-134

585 **Dietrich D, Pang L, Kobayashi A, Fozard JA, Boudolf V, Bhosale R, Antoni R, Nguyen
586 T, Hiratsuka S, Fujii N, et al** (2017) Root hydrotropism is controlled via a cortex-specific
587 growth mechanism. *Nature Plants* **3**: 17057

588 **Elfving F** (1882) Ueber eine Wirkung des galvanischen Stromes auf wachsende Wurzeln. *Bot
589 Zeit* 257-264

590 **Ewart AJ, Bayliss JS** (1905) On the nature of the galvanotropic irritability of roots. *Proc Royal
591 Soc Lond Ser B Contain Pap Biological Character* **77**: 63-66

592 **Geldner N, Déneraud-Tendon V, Hyman DL, Mayer U, Stierhof Y-D, Chory J** (2009)
593 Rapid, combinatorial analysis of membrane compartments in intact plants with a multicolor
594 marker set. *The Plant journal : for cell and molecular biology* **59**: 169-178

595 **Gilroy S** (2008) Plant tropisms. *Current Biology* **18**: R275-7

596 **Ioio RD, Linhares FS, Scacchi E, Casamitjana-Martinez E, Heidstra R, Costantino P,
597 Sabatini S** (2007) Cytokinins determine *Arabidopsis* root-meristem size by controlling cell
598 differentiation. *Current Biology* **17**: 678-682

599 **Ishikawa H, Evans ML** (1990) Electrotropism of maize roots. Role of the root cap and
600 relationship to gravitropism. *Plant physiology* **94**: 913-918

601 **Kamada-Nobusada T, Sakakibara H** (2009) Molecular basis for cytokinin biosynthesis.
602 *Phytochemistry* **70**: 444-449

603 **Kimura T, Haga K, Shimizu-Mitao Y, Takebayashi Y, Kasahara H, Hayashi K-I, Kakimoto
604 T, Sakai T** (2018) Asymmetric Auxin Distribution is Not Required to Establish Root
605 Phototropism in *Arabidopsis*. *Plant Cell Physiology* **59**: 828-840

606 **Kobayashi A, Takahashi A, Kakimoto Y, Miyazawa Y, Fujii N, Higashitani A, Takahashi
607 H** (2007) A gene essential for hydrotropism in roots. *Proc Natl Acad Sci U S A* **104**: 4724-
608 4729

609 **Kral N, Ougolnikova AH, Sena G** (2016) Externally imposed electric field enhances plant
610 root tip regeneration. *Regeneration* **3**: 156-167

611 **Kutschera U, Briggs WR** (2012) Root phototropism: from dogma to the mechanism of blue
612 light perception. *Planta* **235**: 443-452

613 **Li S-M, Zheng H-X, Zhang X-S, Sui N** (2021) Cytokinins as central regulators during plant
614 growth and stress response. *Plant Cell Rep* **40**: 271-282

615 **Liao C-Y, Smet W, Brunoud G, Yoshida S, Vernoux T, Weijers D** (2015) Reporters for
616 sensitive and quantitative measurement of auxin response. *Nature methods* **12**: 207–210

617 **Marchant A, Kargul J, May ST, Muller P, Delbarre A, Perrot-Rechenmann C, Bennett MJ**
618 (1999) AUX1 regulates root gravitropism in *Arabidopsis* by facilitating auxin uptake within
619 root apical tissues. *Embo J* **18**: 2066–2073

620 **Marcum H, Moore R** (1990) Influence of Electrical Fields and Asymmetric Application of
621 Mucilage on Curvature of Primary Roots of *Zea mays*. *American Journal of Botany* **77**: 446

622 **McGillivray AM, Gow NAR** (1986) Applied Electrical Fields Polarize the Growth of Mycelial
623 Fungi. *Microbiology* **132**: 2515–2525

624 **Miyawaki K, Tarkowski P, Matsumoto-Kitano M, Kato T, Sato S, Tarkowska D, Tabata S,**
625 **Sandberg G, Kakimoto T** (2006) Roles of *Arabidopsis* ATP/ADP isopentenyltransferases
626 and tRNA isopentenyltransferases in cytokinin biosynthesis. *Proc National Acad Sci* **103**:
627 16598–16603

628 **Miyazawa Y, Takahashi H** (2019) Molecular mechanisms mediating root hydrotropism: what
629 we have observed since the rediscovery of hydrotropism. *J Plant Res* **133**: 3–14

630 **Muthert LWF, Izzo LG, Zanten M van, Aronne G** (2020) Root Tropisms: Investigations on
631 Earth and in Space to Unravel Plant Growth Direction. *Front Plant Sci* **10**: 1807

632 **Navez AE** (1927) Galvanotropism of roots. *The Journal of general physiology* **10**: 551–558

633 **Olitzki L** (1932) Electric Charge of Bacterial Antigens. *J Immunol* **4**: 251–256

634 **Peters RC, Eeuwes LBM, Bretschneider F** (2007) On the electrodetection threshold of
635 aquatic vertebrates with ampullary or mucous gland electroreceptor organs. *Biol Rev* **82**:
636 361–373

637 **Pozdnyakov A, Pozdnyakova L** (2002) Electrical fields and soil properties. *Proceedings of*
638 *17th World Congress of Soil Science*. p N. 1558

639 **Revil A, Naudet V, Nouzaret J, Pessel M** (2003) Principles of electrography applied to self-
640 potential electrokinetic sources and hydrogeological applications. *Water Resources*
641 *Research* **39**: SBH 3: 1-15

642 **Sabatini S, Beis D, Wolkenfelt H, Murfett J, Guilfoyle T, Malamy J, Benfey P, Leyser O,**
643 **Bechtold N, Weisbeek P, et al** (1999) An Auxin-Dependent Distal Organizer of Pattern and
644 Polarity in the *Arabidopsis* Root. *Cell* **99**: 463–472

645 **Schindelin J, Arganda-Carreras I, Frise E, Kaynig V, Longair M, Pietzsch T, Preibisch S,**
646 **Rueden C, Saalfeld S, Schmid B, et al** (2012) Fiji: an open-source platform for biological-
647 image analysis. *Nature methods* **9**: 676–682

648 **Shanley LJ, Walczysko P, Bain M, MacEwan DJ, Zhao M** (2006) Influx of extracellular Ca²⁺
649 is necessary for electrotaxis in *Dictyostelium*. *J Cell Sci* **119**: 4741–4748

650 **Shkolnik D, Krieger G, Nuriel R, Fromm H** (2016) Hydrotropism: Root Bending Does Not
651 Require Auxin Redistribution. *Mol Plant* **9**: 757–759

652 **Stevens SS** (1970) Neural Events and the Psychophysical Law: Power functions like those
653 that govern subjective magnitude show themselves in neurelectric effects. *Science* **170**:
654 1043–1050

655 **Su S-H, Gibbs NM, Jancewicz AL, Masson PH** (2017) Molecular Mechanisms of Root
656 Gravitropism. *Current Biology* **27**: R964–R972

657 **Sukul NC, Croll NA** (1978) Influence of potential difference and current on the electrotaxis of
658 *Caenorhabditis elegans*. *J Nematol* **10**: 314–7

659 **Sutton GP, Clarke D, Morley EL, Robert D** (2016) Mechanosensory hairs in bumblebees
660 (*Bombus terrestris*) detect weak electric fields. *Proceedings of the National Academy of
661 Sciences* **113**: 7261–7265

662 **Takamura T** (2006) Electrochemical Potential around the Plant Root in Relation to Metabolism
663 and Growth Acceleration. *In* AG Volkov, ed, *Plant Electrophysiology, Theory and Methods*.
664 Springer, Berlin, pp 341–374

665 **Takei K, Yamaya T, Sakakibara H** (2004) Arabidopsis CYP735A1 and CYP735A2 Encode
666 Cytokinin Hydroxylases That Catalyze the Biosynthesis of trans-Zeatin. *J Biol Chem* **279**:
667 41866–41872

668 **Thimann KV, Went FW** (1937) Phytohormones. doi: 10.5962/bhl.title.5695

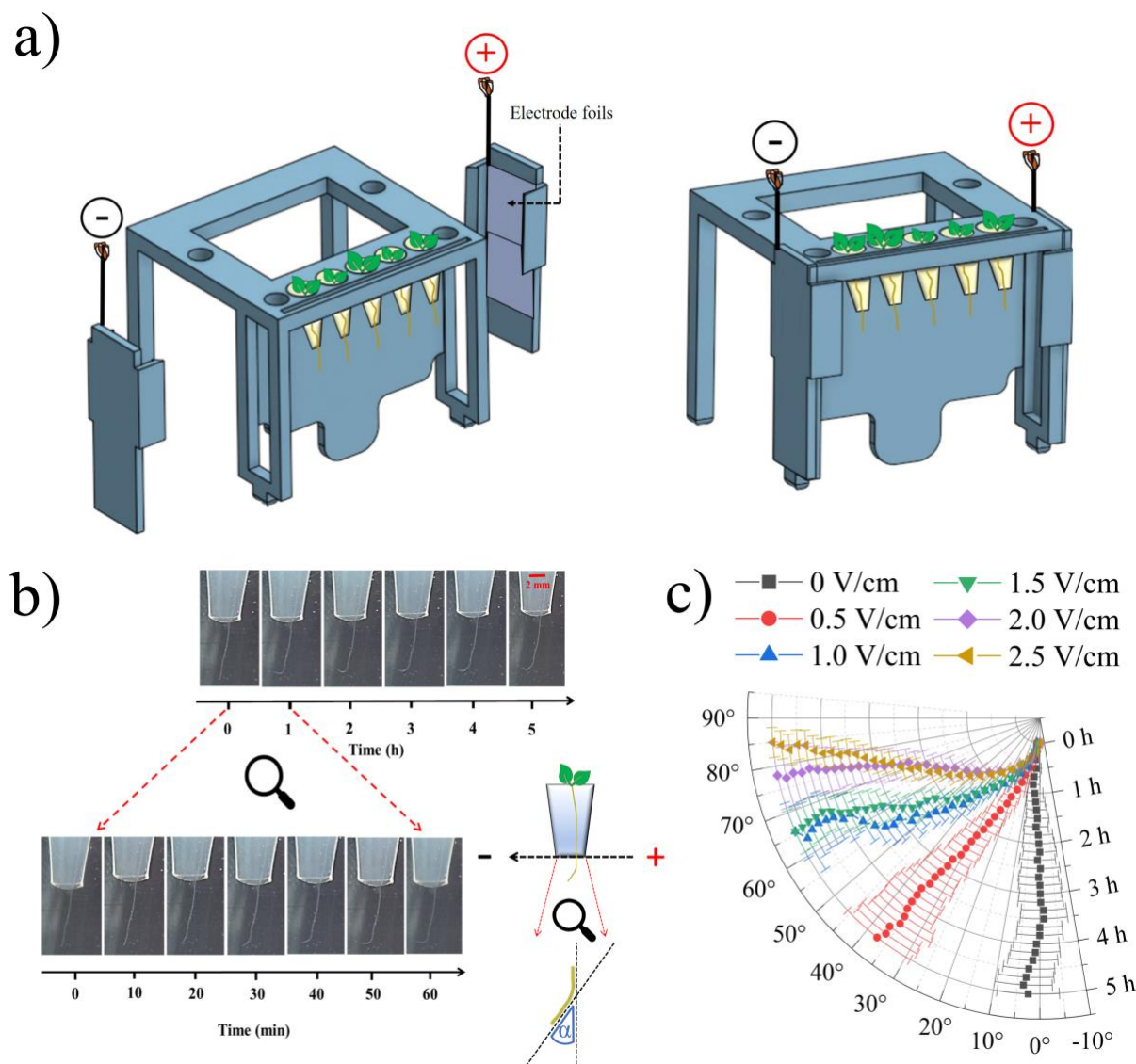
669 **Ward SH** (1990) Geotechnical and environmental geophysics. *Soc. of Exploration
670 Geophysicists*

671 **Wawrecki W, Zagórska-Marek B** (2007) Influence of a Weak DC Electric Field on Root
672 Meristem Architecture. *Ann Bot-london* **100**: 791–796

673 **Wolverton C, Mullen JL, Ishikawam H, Evans ML** (2000) Two distinct regions of response
674 drive differential growth in *Vigna* root electrotropism. *Plant Cell Environ* **23**: 1275–1280

675 **Wu Y, Liu H, Wang Q, Zhang G** (2021) Roles of cytokinins in root growth and abiotic stress
676 response of *Arabidopsis thaliana*. *Plant Growth Regul* 1–10

677



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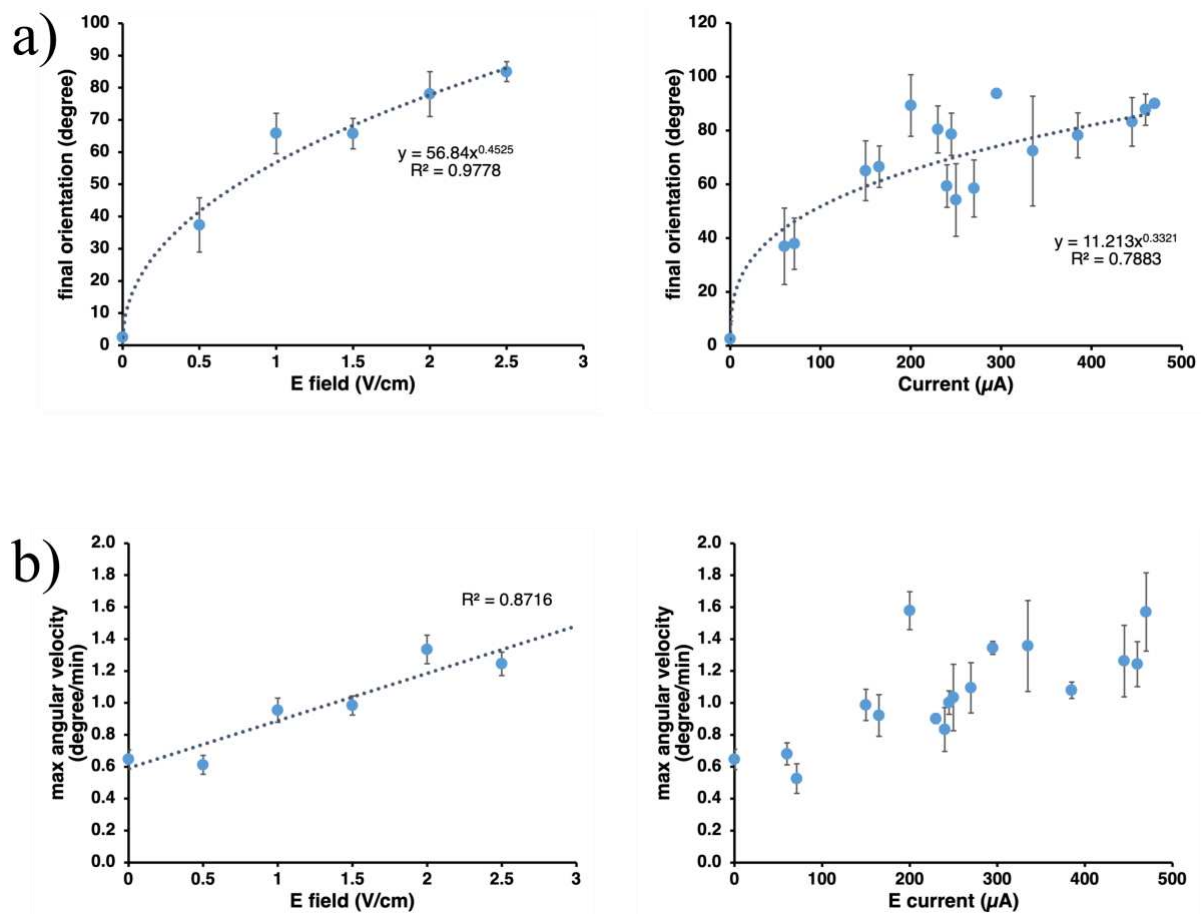
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681 **Figure 1 (a)** Schematic of the 3D-printed module used in the V-box. **(b)** Representative time-lapse series of a single root as imaged by the Raspberry camera from the V-box; inset, schematic of the angle measured. **(c)** Polar plot of the average root tip orientations relative to the gravity vector, with time on the radial axis and orientation angle on the circumferential axis; error bars, s.e.m.

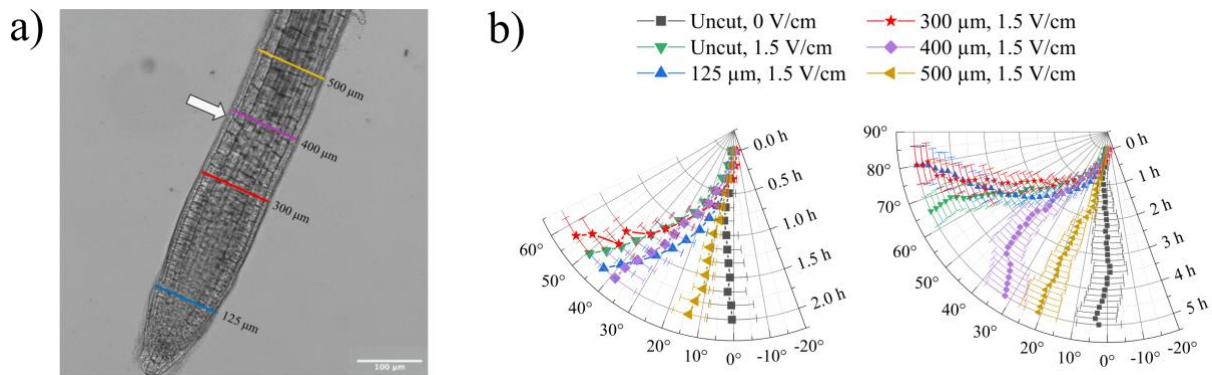
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689 **Figure 2. (a)** Root electrotropism response curves: average root tip orientations after 5 h of
690 exposure to a given electric field (left panel) or current (right panel); error bars, s.e.m.; R^2 ,
691 coefficient of determination. **(b)** Average maximum angular velocity reached by the root tip
692 after 5 h of exposure to a given electric field (left) or current (right); error bars, s.e.m.; R^2 ,
693 coefficient of determination.
694

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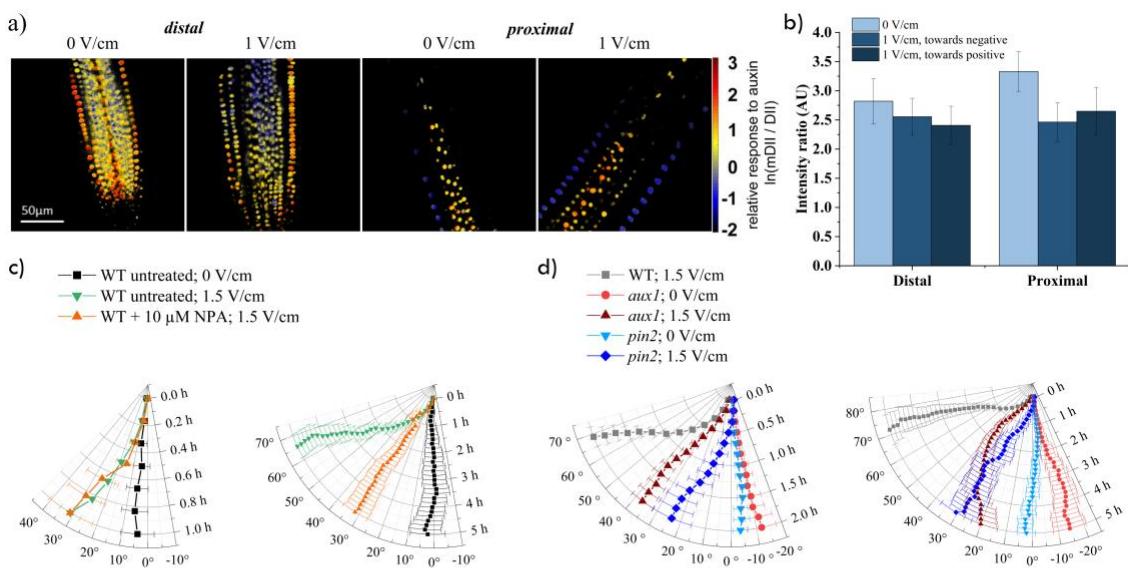
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698 **Figure 3.** (a) Points of excision overlapped to a representative microscope image of the
699 *Arabidopsis* primary root tip; arrow, transition point; scale bar, 100 μm. (b) Polar plot of the
700 average root tip orientations with respect to the gravity vector, with time on the radial axis and
701 orientation angle on the circumferential axis; the same data is presented for the first 2 h (left
702 panel) and for the full 5 h (right panel); error bars, s.e.m.

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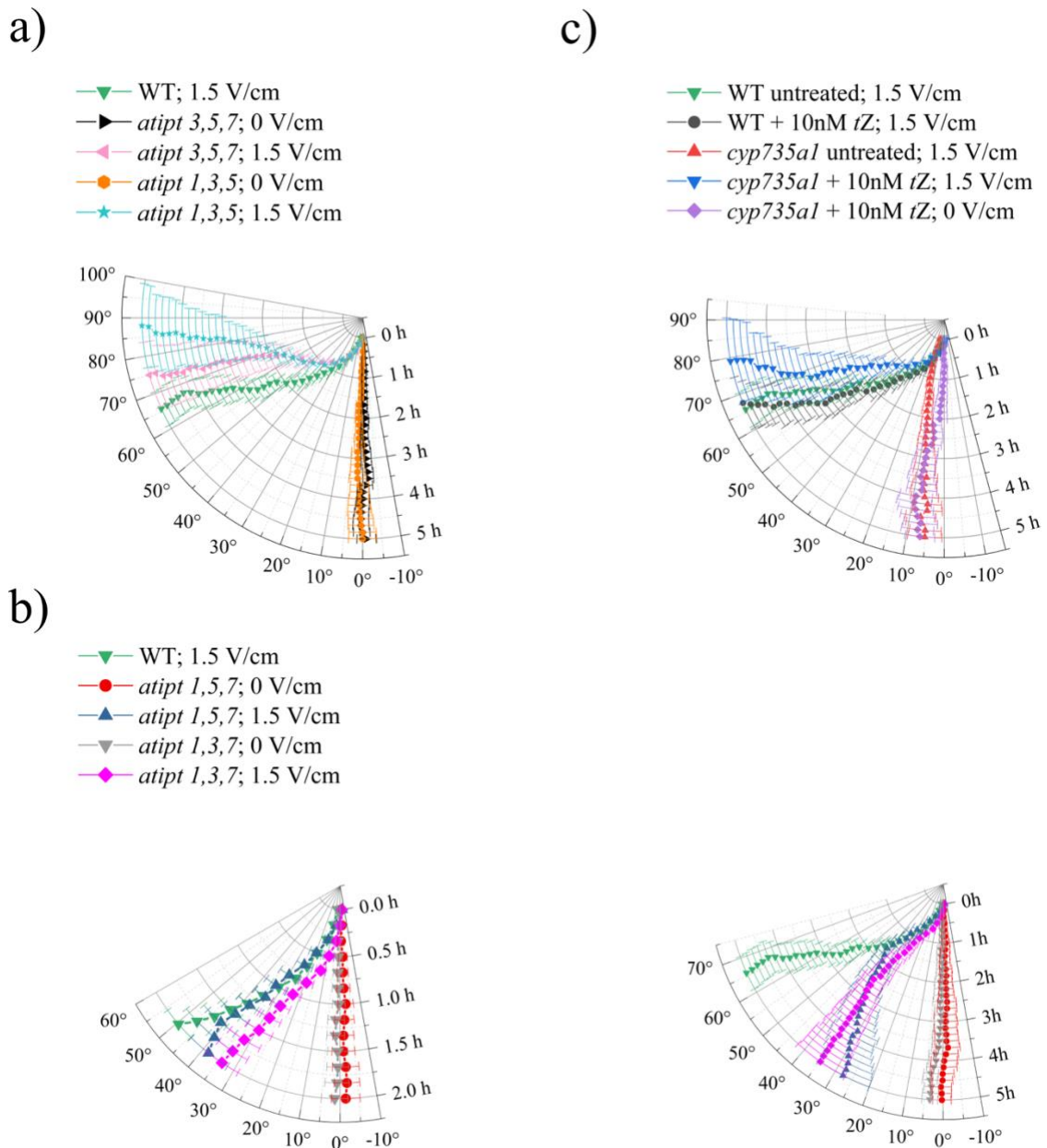
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706 **Figure 4. (a)** Relative response to auxin concentration in *Arabidopsis* primary root tips, indicated by the ratiometric fluorescent reporter R2D2 (see Methods); left panels: 707 representative tip (distal) regions at 0 V/cm and 1 V/cm; right panels: representative higher 708 (proximal) regions at 0 V/cm and 1 V/cm; scale bar, 50 μ m. **(b)** Quantification of the average 709 relative response to auxin (as shown in a) in the epidermis cell layer facing the positive or 710 negative electrode, in the distal and proximal regions; the data for 0 V/cm is an average 711 between the two sides; error bars, s.e.m. **(c-d)**, Polar plots of the average root tip orientation 712 relative to the gravity vector, with time on the radial axis and orientation angle on the 713 circumferential axis: NPA treatment **(c)**; aux1 and pin2 mutants **(d)**; error bars, s.e.m. 714

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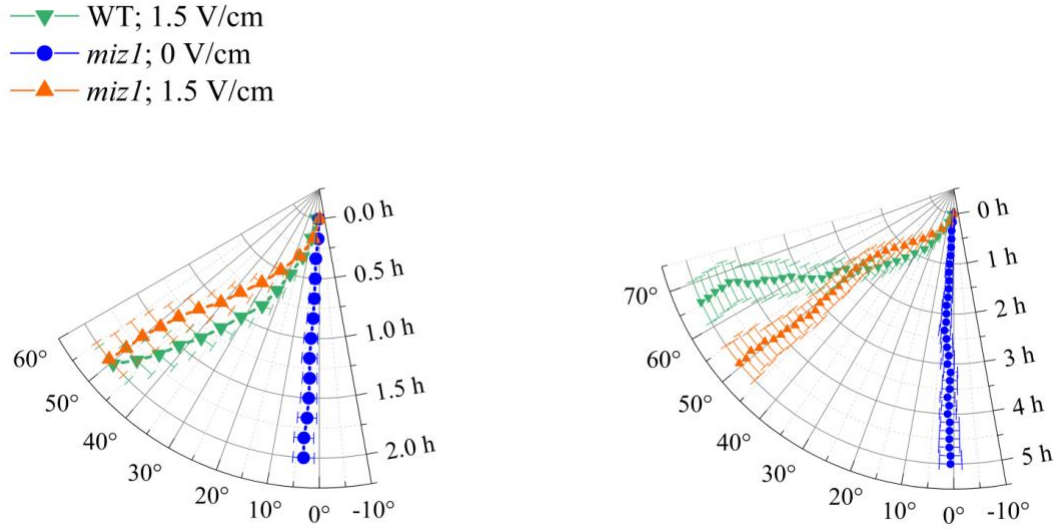


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718 **Figure 5.** Polar plots of the average root tip orientation relative to the gravity vector, with time
719 on the radial axis and orientation angle on the circumferential axis: **(a)** *atipt* triple mutants with
720 strong electro tropic response; **(b)** *atipt* triple mutants with weak electro tropic response; **(c)**
721 *cyp735a1* mutant; error bars, s.e.m.

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725 **Figure 6.** Polar plots of the average root tip orientation in *miz1* mutant relative to the gravity
726 vector, with time on the radial axis and orientation angle on the circumferential axis.