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1	Cortical waves mediate the cellular response to electric fields
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### 23 Abstract:

24 Electrotaxis, the directional migration of cells in a constant electric field, is important in 25 regeneration, development, and wound healing. Electrotaxis has a slower response and a smaller 26 dynamic range than guidance by other cues, suggesting that the mechanism of electrotaxis share 27 both similarities and differences with chemical-gradient-sensing pathways. We examined a 28 mechanism centered on the excitable system consisting of cortical waves of biochemical signals 29 coupled to cytoskeletal reorganization, which has been implicated in random cell motility. We 30 use electro-fused giant Dictyostelium discoideum cells to decouple waves from cell motion and 31 employ nanotopographic surfaces to limit wave dimensions and lifetimes. We demonstrate that 32 wave propagation in these cells is guided by electric fields. The wave area and lifetime gradually 33 increase in the first 10 minutes after an electric field is turned on, leading to more abundant and 34 wider protrusions in the cell region nearest the cathode. The wave directions display "U-turn" 35 behavior upon field reversal, and this switch occurs more quickly on nanotopography. Our 36 results suggest that electric fields guide cells by controlling waves of signal transduction and 37 cytoskeletal activity, which underlie cellular protrusions. Whereas surface receptor occupancy 38 triggers both rapid activation and slower polarization of signaling pathways, electric fields 39 appear to act primarily on polarization, explaining why cells respond to electric fields more 40 slowly than to other guidance cues.

41

### 42 Introduction

Electrotaxis, which refers to the directed migration of cells under the guidance of an electric field
(EF), is important in wound healing, development, and regeneration (Cortese et al., 2014; Lin et
al., 2008; Zhao et al., 2006). EFs have been shown to cause several key signaling molecules to be

46	distributed asymmetrically across cells (Sato et al., 2009; Zhao et al., 2002, 2006), setting up cell						
47	polarity. The one-order-of-magnitude range of EF strengths sensed by cells (Zhao et al., 2002) is						
48	considerably smaller than the four-orders-of-magnitude concentration sensitivity in chemotaxis						
49	(Harvath et al., 1991). Furthermore, whereas cells respond to chemical guidance cues on a time						
50	scale of seconds and develop polarity over several minutes, the response to EFs can take up to 10						
51	minutes or more after the EF is turned on (Wang et al., 2014; Zhao et al., 2006). These						
52	differences raise the possibility that the rapid gradient sensing mechanisms do not serve as						
53	primary mediators of EF sensing by cells. In this study, we examine whether, after turning on an						
54	EF, the gradual polarization of the excitable biochemical networks that organize actin						
55	polymerization comprises a slow-acting mediator of the cellular response to the EF.						
56							
57	Actin polymerization, coordinated with its associated signaling molecules, self-organizes into						
57 58	Actin polymerization, coordinated with its associated signaling molecules, self-organizes into microscale spatial regions that travel as waves across plasma membranes. These waves drive						
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58 59	microscale spatial regions that travel as waves across plasma membranes. These waves drive various cell behaviors, such as migration and division (Bhattacharya et al., 2019; Bretschneider						
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58 59 60 61	microscale spatial regions that travel as waves across plasma membranes. These waves drive various cell behaviors, such as migration and division (Bhattacharya et al., 2019; Bretschneider et al., 2009; Flemming et al., 2020; Gerhardt et al., 2014; Gerisch, 2010). The wave system can be described as a coupled signal transduction excitable network - cytoskeletal excitable network						
58 59 60 61 62	microscale spatial regions that travel as waves across plasma membranes. These waves drive various cell behaviors, such as migration and division (Bhattacharya et al., 2019; Bretschneider et al., 2009; Flemming et al., 2020; Gerhardt et al., 2014; Gerisch, 2010). The wave system can be described as a coupled signal transduction excitable network - cytoskeletal excitable network (STEN-CEN) (Devreotes et al., 2017; Miao et al., 2019). STEN-CEN has the characteristics of						
<ul> <li>58</li> <li>59</li> <li>60</li> <li>61</li> <li>62</li> <li>63</li> </ul>	microscale spatial regions that travel as waves across plasma membranes. These waves drive various cell behaviors, such as migration and division (Bhattacharya et al., 2019; Bretschneider et al., 2009; Flemming et al., 2020; Gerhardt et al., 2014; Gerisch, 2010). The wave system can be described as a coupled signal transduction excitable network - cytoskeletal excitable network (STEN-CEN) (Devreotes et al., 2017; Miao et al., 2019). STEN-CEN has the characteristics of an excitable system, including exhibiting an activation threshold for wave initiation and						
<ul> <li>58</li> <li>59</li> <li>60</li> <li>61</li> <li>62</li> <li>63</li> <li>64</li> </ul>	microscale spatial regions that travel as waves across plasma membranes. These waves drive various cell behaviors, such as migration and division (Bhattacharya et al., 2019; Bretschneider et al., 2009; Flemming et al., 2020; Gerhardt et al., 2014; Gerisch, 2010). The wave system can be described as a coupled signal transduction excitable network - cytoskeletal excitable network (STEN-CEN) (Devreotes et al., 2017; Miao et al., 2019). STEN-CEN has the characteristics of an excitable system, including exhibiting an activation threshold for wave initiation and experiencing refractory periods. It has been shown that the STEN-CEN wave properties dictate						

results (Bhattacharya et al., 2020; Bhattacharya & Iglesias, 2018). For simplicity, here we will
refer to STEN-CEN waves as cortical waves.

71	One challenge in investigating whether cortical waves can act as the mediators of EFs is that in				
72	many of the cell types that show a strong response to EFs, the wave area is comparable to the cell				
73	area. Furthermore, waves are generated at the leading edge of the cell during directed migration				
74	(Xu et al., 2003), so that wave dynamics are tightly coupled with cell dynamics. For instance,				
75	when a cell responds to an EF reversal, waves typically remain at the cell front as the cell turns.				
76	It is not known whether the waves drive cells to turn or the cell polarity keeps the previous				
77	leading edge more active so that this edge responds first.				
78					
79	To distinguish between wave response and cell motion, we produced electro-fused giant D.				
80	discoideum (Neumann et al., 1980) with diameters up to ten times larger than that of an				
81	individual cell. Multiple simultaneous waves can be generated across the surface contact area of				
82	a giant cell (Gerhardt et al., 2014). These waves also generate actin-filled macropinosomes on				
83	the dorsal membrane (Veltman et al., 2016). The giant cells provide an excellent opportunity to				
84	study cortical wave dynamics in multiple cell regions simultaneously.				
85					
86	We further use nanotopography to alter the waves' spatial structures and characteristic timescales.				
87	Upon contact with nanotopography, cells produce quasi-1D wave patches. The phenomenon of				
88	guided actin polymerization by nanotopography is known as esotaxis (Driscoll et al., 2014),				
89	which has been investigated in detail (Ketchum et al., 2018; Lee et al., 2020). There are several				
90	advantages of incorporating nanotopography in our study. First, these waves persist for a shorter				

91 time on nanotopography than on flat surfaces, enabling us to investigate whether wave systems 92 with different characteristic timescales respond to EFs differently. Second, waves on ridged 93 surfaces have shorter lifetimes than those on flat surfaces, and thus only propagate in local 94 regions of giant cells. Therefore, nanotopography allows us to distinguish between local and 95 global mediation of the EF response.

96

97 **Results** 

#### 98 Cortical waves and cell migration can be studied independently in giant cells

We imaged cells that simultaneously expressed both limE-RFP and  $PH_{Crac}$ -GFP. The former allows us to monitor filamentous actin (F-actin), which represents CEN activities. The latter enables us to monitor phosphatidylinositol-3,4,5-trisphosphate (PIP3), an indicator of STEN activities. In single, differentiated cells, usually only one wave is generated at the leading edge (Fig. 1a and Video 1), and the wave motion is coupled with cell motion. For instance, when the cell in Fig. 1a changed its direction of motion, the wave remained at the leading edge (72 s - 120 s).

106

In giant cells, multiple waves were initiated randomly and propagated radially across the basal
membranes (Fig. 1b and Video 2). CEN is driven by STEN, but has a substantially shorter
characteristic timescale. Thus, PIP3 waves displayed band-like shapes, whereas F-actin appeared
across the bands with higher levels at the rims of PIP3 waves (Miao et al., 2019). As shown in
Fig. 1b, colliding waves did not cross, but instead rotated by 90° (Fig. 1b, 150 s - 200 s). This
behavior is suggestive of a refractory period following excitation, which is a hallmark of an
excitable system. On nanoridges, the giant cells generated multiple, quasi-1D patches of F-actin

114	and PIP3 with shorter lifetimes than on flat surfaces (Fig. 1c and Video 3). Some waves formed				
115	and propagated for a short distance (Line 2 in Fig. 1c), whereas others formed and then quickly				
116	dissipated (Line 3 in Fig. 1c). The wave dissipation can be explained in terms of an excitable				
117	system with lateral inhibition, in which the dispersion of the inhibitor is faster than that of the				
118	activator. Thus, the waves eventually dissipate due to the spatial accumulation of the inhibitor.				
119	Prior studies have shown that in this situation, the excitable system threshold determines the				
120	wave duration (Bhattacharya et al., 2020; Ermentrout et al., 1984). As was the case on flat				
121	surfaces, 1D patches occurred throughout the basal surfaces on ridges, and thus were				
122	independent of cell motion.				
100					
123					
123 124	Waves were also generated on the dorsal planes. In contrast to basal waves, which propagated				
	Waves were also generated on the dorsal planes. In contrast to basal waves, which propagated across the surface contact (Video 2 and Video 3), dorsal waves were associated with membrane				
124					
124 125	across the surface contact (Video 2 and Video 3), dorsal waves were associated with membrane				
124 125 126	across the surface contact (Video 2 and Video 3), dorsal waves were associated with membrane deformations, and resembled macropinosomes (Video 4). Based on 3D lattice light-sheet images				
124 125 126 127	across the surface contact (Video 2 and Video 3), dorsal waves were associated with membrane deformations, and resembled macropinosomes (Video 4). Based on 3D lattice light-sheet images of a cell plated on nanoridges (Fig. 1d and Video 5), activation of PIP3 and F-actin was				
124 125 126 127 128	across the surface contact (Video 2 and Video 3), dorsal waves were associated with membrane deformations, and resembled macropinosomes (Video 4). Based on 3D lattice light-sheet images of a cell plated on nanoridges (Fig. 1d and Video 5), activation of PIP3 and F-actin was correlated in both basal waves and dorsal waves. However, the dorsal waves were primarily				
124 125 126 127 128 129	across the surface contact (Video 2 and Video 3), dorsal waves were associated with membrane deformations, and resembled macropinosomes (Video 4). Based on 3D lattice light-sheet images of a cell plated on nanoridges (Fig. 1d and Video 5), activation of PIP3 and F-actin was correlated in both basal waves and dorsal waves. However, the dorsal waves were primarily generated in cuplike structures, whereas the stripe-like basal waves spanned the entire basal				

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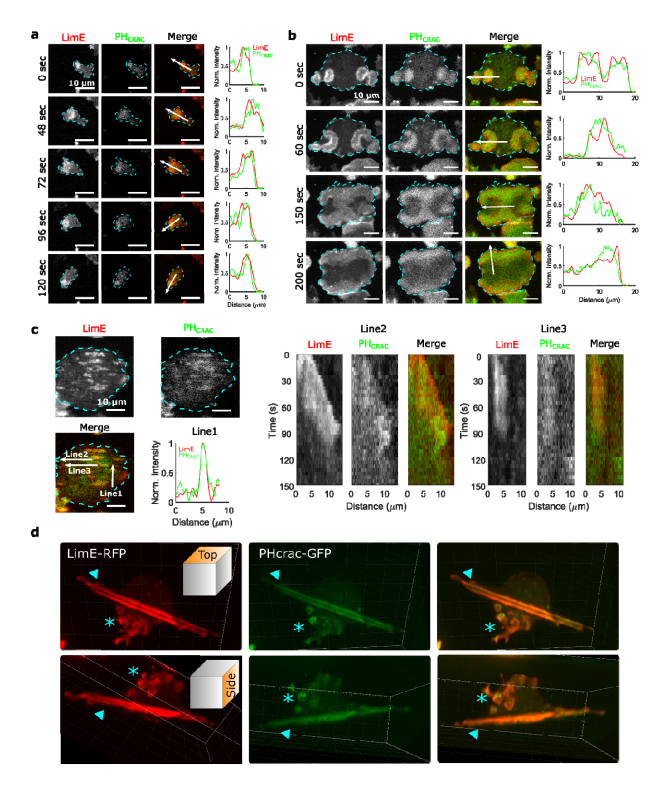


Fig. 1. STEN-CEN waves in single cells and giant cells. a. Snapshots of a differentiated, single *D. discoideum* cell expressing limE-RFP and PHcrac-GFP, with cell boundaries denoted with
blue dashed lines. The right column shows the normalized intensity of limE and PHcrac from the

137 arrows in the merge images. The scale bars are 10 um. b. Snapshots of an electrofused giant D. 138 *discoideum* cell on a flat surface, with scanning profiles in the right column. All scale bars are 10 139  $\mu$ m. c. A snapshot of an electrofused giant cell on the ridged surface. The left kymographs are 140 from the line 2 and line 3 specified in the merged image. Line 2 shows a wave propagating along 141 nanoridges, and line 3 shows a wave that existed briefly and then dissipated. d. 3D reconstruction (single time point) of a single D. discoideum cell plated on nanoridges, acquired 142 143 using a lattice light-sheet microscope. Here we show the top aspect view (top row) and the side 144 aspect view (bottom row). On the dorsal membrane of the cell, there are waves forming 145 microcytotic cups (triangle) on the curved membrane, and on the basal membrane, there are 146 streak-like waves (asterisk). The red channel represents limE-RFP, and the green channel 147 represents PHcrac-GFP. As both the side and top views show, the dorsal waves and basal waves 148 are independent structures, but both are composed of coordinated F-actin and PIP3.

149

#### 150 EFs increase the area, duration, and speed of waves on nanoridges.

151 We found that giant cells respond to a narrow range (15 V/cm to 20 V/cm) of EF amplitudes 152 (Video 7 and Video 8), and that higher voltage (35 V/cm) damaged cells. The 1D waves 153 generated on nanoridges related to esotaxis enabled us to quantify the effects of a 20 V/cm EF 154 (all EFs used here are of this magnitude, see Fig. 2 – figure supplement 1 and Methods for more 155 details about the electrotaxis experiments) on the areas, durations, and speeds of waves. Fig. 2a 156 shows snapshots of the dynamics of F-actin in a giant cell on parallel nanoridges with a 1.6 µm 157 spacing. In the absence of an EF (top row in Fig. 2a), individual actin polymerization events 158 were initiated in patches on the basal surfaces (Fig. 2a and Video 9).

160 In the first several minutes after turning on the EF, most patches propagated as a wave along a 161 single ridge (Fig. 2a, blue inset). After the EF was on for 10 min, some patches appeared to 162 undergo coordinated motion across several ridges (Fig. 2a, pink inset). We calculated the ratio of 163 F-actin occupancy to cell area, and found that an EF increased the overall level of actin 164 polymerization by a factor of two to three (Fig. 2b). Actin patches were larger in the presence of an EF and organized into larger groups located preferentially at the cell front (bottom row in Fig. 165 166 2a, 20 min, and 25 min), leading to wider protrusions at cell fronts that drove directed cell 167 migration (Fig. 2 – figure supplement 2). To determine whether the groups comprised a single, 168 large wave growing across multiple ridges or multiple, small patches nucleated in close 169 proximity, we measured the dynamics of the patches using optical flow (Lee et al., 2020), 170 focusing on the patch edges (Fig. 2c, left image. See Methods for more details). If both edges of 171 a patch were moving in the same direction, the structure was classified as a single, large wave. If 172 the edges were not coordinated, the patch was classified as multiple, individual structures. This 173 method enabled us to capture accurately waves that span across multiple ridges and are moving 174 coordinately.

175

Once the large actin structures were classified, their instantaneous dimensions were measured parallel and perpendicular to the ridges (Fig. 2c). Density scatter plots of both dimensions exhibit elliptical contours (Fig. 2d), suggesting that nanotopography constrains wave growth. With an EF parallel to the ridges, the waves broadened in both directions (Fig. 2d). The average increases in wave dimension parallel and perpendicular to the ridges were 20% and 13%, respectively, and the average increase in wave area was 44%. An increase in wave duration was also observed, with the minimum wave area correlated to the duration (Fig 2e, black circles). The wave area 183 depends exponentially on the maximum wave duration (Fig. 2e, solid black lines), allowing us to

184 extract a characteristic wave time scale via

185

186 
$$Area_{min} = C * e^{\frac{Duration}{T}} .$$
 (1)

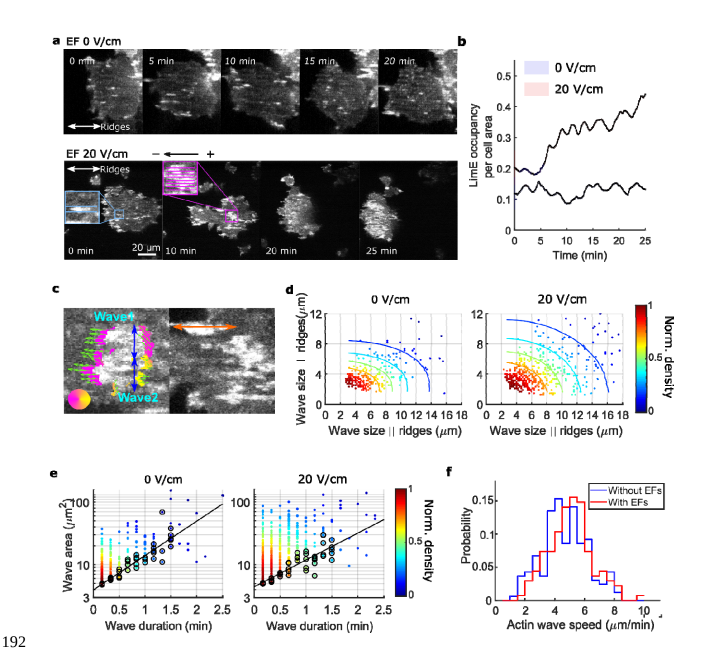
187

188 Here *C* is a constant, and *T* is the characteristic time scale, which is 48 s with no EF and 61 s in

189 the presence of a 20 V/cm EF. This difference is consistent with the EF drawing the system

- 190 closer to the excitability threshold. An average increase of 9% in wave propagation speed was
- 191 also observed in the presence of an EF (Fig. 2f).

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**Fig. 2. EFs alter F-actin wave properties. a.** limE images of a giant cell on nanoridges without an EF (top) and in a 20 V/cm EF turned on at 0 min (bottom). **b.** The temporal change of the percentage of the cell area occupied by limE without an EF (blue,  $N_{cell} = 5$ ) and in a 20 V/cm EF introduced at 0 min (red,  $N_{cell} = 4$ ). The shaded areas represent the mean plus or minus one standard deviation **c.** Division of groups of waves. The color represents the orientation of optical-flow vectors according to the color wheel. The green arrows are the optical-flow vectors,

- the length of which correspond to the magnitude of motion. The left image is an example of a
- 200 large structure composed of two independent substructures, where the vectors at the right edge
- 201 are not moving in the same direction. The wave scales in the directions perpendicular to (blue
- arrows) and parallel to the ridges (orange arrow) were measured on the preprocessed waves. **d.**
- 203 Density scatter plots of wave scales parallel to ridges vs. perpendicular to ridges. e. Density
- scatter plots of actin-wave dimension vs. actin-wave duration. For each wave duration, the 5
- 205 points with the smallest wave areas (black circles) were selected to fit the boundaries (solid black
- lines). **f.** Distributions of wave propagation speeds before (blue,  $N_{wave} = 125$ ) and after (red,  $N_{wave}$ )
- 207 = 163) applying an EF. The analyses in d-f were based on N = 4 independent experiments. The
- 208 two distributions are different (Two-sample t-test, P = 0.017).
- 209 Figure 2 source data 1. Related to Fig. 2b
- 210 Figure 2 source data 2. Related to Fig. 2d
- 211 Figure 2 source data 3. Related to Fig. 2e
- 212 Figure 2 source data 4. Related to Fig. 2f
- 213

### **EFs guide the direction of actin waves.**

215 Next, we consider the directional guidance of actin waves by EFs on nanoridges (Fig. 3a and

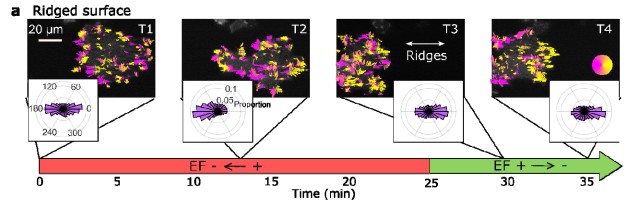
- 216 Video 9) and on flat surfaces (Fig. 3b and Video 10, see Methods for details). The EF was
- introduced at 0 min  $(T_1)$ , and in the first 2 min had little effect on the actin dynamics on any
- surface. On nanoridges, actin waves continued to propagate preferentially along the ridges (Fig.
- 219 3a,  $T_1$ ). On flat surfaces, the waves propagated radially in groups, as seen from the broad
- distribution at  $T_1$  in Fig. 3b. In the presence of an EF, the waves propagated preferentially

towards the cathode within ~15 min (Fig. 3a, 3b,  $T_2$ ). The perpendicular spread was significantly more limited on nanoridges (Fig. 3a,  $T_2$ ).

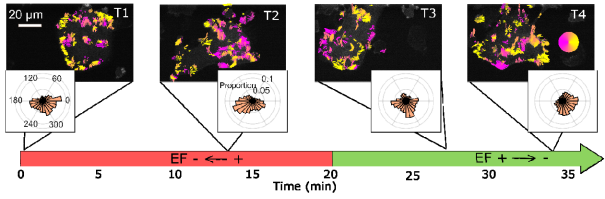
224	The direction of the EF was reversed after the cell had commenced steady directional migration,
225	which took ~20 to 25 min. Following the field reversal, waves on ridged surfaces reoriented
226	toward the new cathode within 5 min (Fig. 3a, $T_3$ ). On flat surfaces, the wave propagation
227	direction was perpendicular to both the previous and the new EF directions at ~7 min after the
228	field reversal (Fig. 3b, $T_3$ ). Preferential propagation towards the new cathode occurred after ~13
229	min (Fig. 3b, $T_4$ ). The difference in response time between nanoridges and flat surfaces may be
230	related to the fact that waves persist longer on flat surfaces than on nanoridges (Fig. 3c).
231	
232	Fig. 3d shows the continuous temporal changes of the orientation distributions. On nanoridges,
233	the preferred wave directions switched directly following EF reversal (left plot in Fig. 3d),
234	whereas on flat surfaces waves maintained a preferred direction that changed continuously in a
235	U-turn behavior (Right plot in Fig. 3d). In contrast, although single D. discoideum cells undergo
236	U-turns in response to EF reversal (Sato et al., 2007), giant cells did not (Video 10).
237	Wave turning may be related to differences in the patterns of wave expansion (Fig. 3e). On flat
238	surfaces, waves started from a small patch (Fig. 3e, S1) and eventually broke into band-shaped
239	waves (Fig. 3e, S4). During directed migration, the intermediate expansion of actin waves (Fig.
240	3e, S2, S3 in the top row) was biased by the EF, resulting in band-shaped waves propagating
241	preferentially towards the cathode (Fig. 3e, S4 in top row). After EF reversal, waves expanded in
242	all directions (Fig. 3e, S2, S3 in bottom row), such that optical-flow analysis captured turning
243	behavior more frequently.

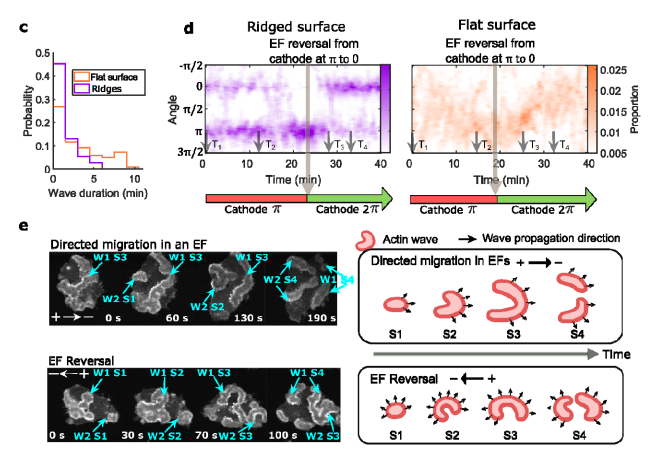
245	We also simultaneously imaged limE-RFP at the basal plane (near the surface contact) and the
246	dorsal plane (6 $\mu$ m higher). Dorsal waves (Video 11 and 12) are localized at cell fronts and
247	rearranged to the new fronts following EF reversal (Fig. 3 – figure supplement 1). Rather than
248	directly switching preferential direction, dorsal waves gradually turned toward the new cathode
249	(Fig. 3 – figure supplement 1), in a manner similar to that of basal waves on flat surfaces. Thus,
250	two different response times to EF reversal exist within the same cell, with a faster response for
251	basal waves guided by nanotopography and a slower response for the free dorsal waves. On flat
252	surfaces, the two responses are synchronized (Fig. 3 – figure supplement 2).

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**b** Flat surface





254 Fig. 3. EFs guide actin waves. a, b Optical-flow analysis of actin-wave dynamics in giant cells 255 on ridged and flat surfaces. The top row shows a time series of limE images for giant cells 256 overlaid with optical-flow vectors, the color of which is coded according to the color wheel. The 257 accompanying polar plots show the corresponding orientation displacements of optical-flow 258 vectors. For both a and b the EF was turned on at 0 min. The bottom time stamp indicates when 259 the EF was reversed from the cathode being on the right (red) to the cathode being on the left 260 (green). c. Distributions of wave duration from 3 independent days of experiments. The 261 distributions were weighted by wave area, because the number of long-lasting large waves on flat surfaces ( $N_{wave} = 359$ ) is smaller than the number of short-lived small patches on ridged 262 263 surfaces ( $N_{wave} = 658$ ). Correspondingly, the absolute waves counts do not match the pixel-based, 264 optical-flow analysis in a and b. Based on a two-sample t-test on the wave areas on flat surfaces vs. on ridges, the null hypothesis was rejected at the 5% significance level with  $p = 2 \times 10^{-15}$ . **d**. 265 266 Kymographs of orientation displacements of optical-flow vectors. The x-axes of the kymographs 267 represent time, and the y-axes represent orientation. The colors represent the proportions. The EF 268 was turned on at time  $T_1$ , and was reversed at the time denoted by the black arrow e. LimE 269 snapshots showing the patterns of actin-wave expansion during steady directed migration in a 270 constant EF (top) and after reversing the EF direction (bottom). The blue arrows point to specific 271 stages of wave expansion. W: Wave, S: Stage of wave expansion. The right panel is a cartoon 272 illustrating the patterns of actin-wave expansion during directed migration in EFs (top) and after 273 EFs were reversed (bottom).

- 274 Figure 3 source data 1. Related to Fig. 3c
- 275
- 276

#### 277 Subcellular spatial inhomogeneity of the response to EFs on nanoridges.

278 Although waves in migrating *D. discoideum* cells localize predominantly at the leading edge 279 (Weiner et al., 2007; Zhao et al., 2002), waves are observed across the basal layer in giant D. 280 *discoideum* cells. We analyzed the smaller, shorter-lived waves on nanoridges. Although the 281 wave locations were distributed essentially uniformly throughout cells in the absence of an EF, 282 more waves were generated at the cell fronts in the presence of an EF (Fig. 4a and Video 9). In 283 addition, the average area per wave was larger near the front of cells in an EF (Fig. 4b). We also 284 measured the wave properties in the single cells scattered throughout the field of view but did not 285 observe a corresponding gradient of wave properties among single cells closer to the cathode 286 versus the cells closer to the anode. This result indicates that the spatial inhomogeneity shown in 287 Figs. 4a, b was caused by the EF rather than by the absolute electrical potential relative to the ground (Fig. 4 – figure supplement 1). 288

289

We explored the response of this inhomogeneity to EF reversal by tracking each wave location relative to the cell centroid in the 12 min following EF reversal (Fig. 4c). New waves started to appear near the side of the cell facing the new cathode within 3 min (Fig. 4c, left region of  $P_2$ ), whereas the complete inhibition of wave generation near the old cell front took longer (Fig. 4c, right region of P<sub>5</sub>). This observation suggests that the initiation at a new cell front and the inhibition of waves at the old front are regulated by two distinct processes with different timescales.

297

Next, we looked at the time required to switch propagation direction in different subcellular
regions following EF reversal. The basal membrane was segmented into an "old front" region

- 300 (facing the original cathode) and a "new front" region (facing the new cathode), as illustrated in
- 301 Fig. 4d. The distributions of wave propagation directions show that waves in the new front
- 302 region switched their preferential direction at ~4 min. In contrast, waves in the old front region
- 303 changed their preferential direction on a time scale of ~ 7 min (Figs. 4 e, f). Our analysis further
- 304 shows that larger waves in the old fronts are less sensitive to EF reversal than those in the new
- 305 fronts (Fig. 4 figure supplement 2).

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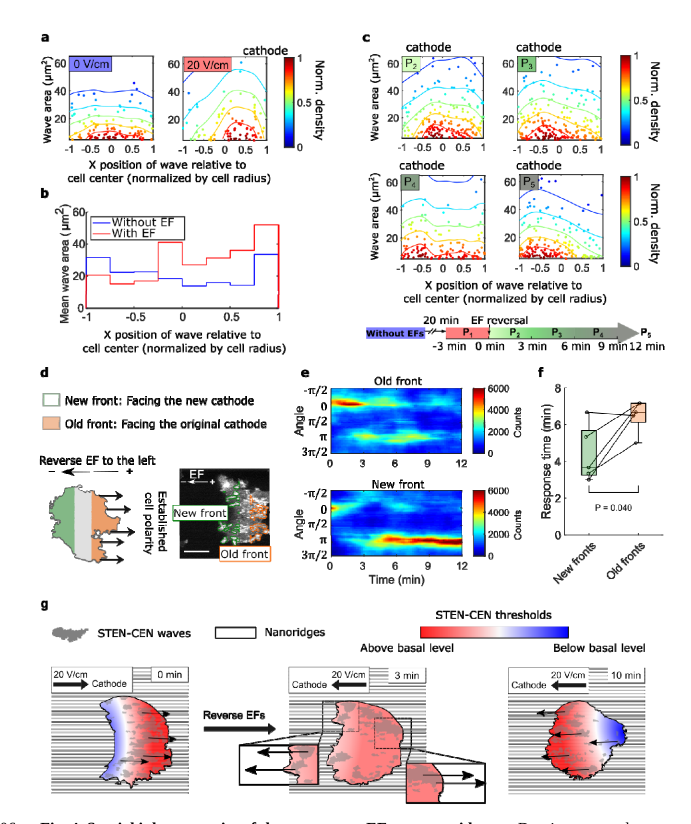


Fig. 4. Spatial inhomogeneity of the response to EFs on nanoridges. a. Density scatter plots
of the wave area vs. x position of the wave relative to the cell center. Nanoridges and EF are

308 orientated in the x-direction. The difference of x coordinates of cell center and wave location was 309 calculated, then the value was further normalized by the cell radius. Each point represents a wave, 310 and all the points were collected from 5 independent experiments. The left plot is for a period in 311 which there was no EF ( $N_{wave} = 296$ ), and the right plot is for a period in which there was a 20 312 V/cm EF, during which the cells exhibited steady directional migration ( $N_{wave} = 224$ ). For each 313 experiment without an EF, the EF was always turned on several minutes later. Thus, we defined 314 the direction in which cathode was located in the presence of an EF as the positive direction in 315 the absence of an EF. The color code corresponds to the density of points. b. Average wave area 316 in sub-cellular regions. The points in a were sectioned, based on their x position relative to the 317 cell center (normalized by cell radius) at a bin size of 0.25 (8 sections in total from -1 to 1), and 318 calculated the average wave area in each section. c. Changes in actin waves' spatial distribution 319 in response to EF reversal; data from 6 independent experiments. The color of each plot is coded 320 according to the timeline displayed at the bottom of the panel.  $P_2$ - $P_5$ : The EF was reversed, and 321 cells gradually developed polarization towards the new cathode. The number of waves in each 322 plot:  $N_{p2} = 272$ ,  $N_{p3} = 277$ ,  $N_{p4} = 193$ ,  $N_{p5} = 246$ . **d.** A schematic illustrating the old and new 323 fronts of giant cells when the EF was reversed. e. Time stacks of orientation distributions of 324 optical-flow vectors at an old front and a new front. The EF was reversed from the cathode being at the right (0) to the cathode being at the left ( $\pi$ ) at 0 min. **f.** Comparisons of response time 325 between new fronts (green) and old fronts (orange) from multiple experiments ( $N_{cell} = 5$ ). The P-326 327 value was calculated using a pairwise t-test at the 5% significance level. g. Cartoon illustrating 328 different time scales of local wave propagation and global rearrangement of STEN-CEN 329 thresholds, in response to EF reversal.

330 Figure 4 – source data 1. Related to Fig. 4a-c

## 331 Discussion

By employing giant cells, in which the cortical waves are disentangled from cell motion, we demonstrate that EFs modulate cortical wave dynamics directly, providing a mechanism for cell guidance by EFs (Fig. 1 c, b, d). Our use of nanoridges to generate quasi-1D waves that are small, short-lived, and unable to turn (Fig. 1c) enabled detailed quantification of wave properties, demonstrating that EFs directly affect the abundance, locations, and directions of cortical waves.

337

## 338 EFs guide cortical wave dynamics

339 Previous studies have suggested that the basal cortical waves in D. discoideum are insensitive to 340 external chemotactic gradients, whereas "pseudopods" at other regions in the same cells can be 341 guided (Lange et al., 2016). This conclusion is surprising because the biochemical events 342 traveling with the waves are the same as those occurring on pseudopods, and pseudopods with 343 the dorsal cups on the same cells do respond to chemoattractants. Also, similar cortical waves in 344 human mammary epithelial cells can be guided effectively by epidermal growth factors (Zhan et 345 al., 2020). Additional input from the greater contact of giant D. discoideum cells with the surface may outweigh the effect of applied chemical gradients on the basal waves. Other studies 346 347 have shown that single cells can integrate combinations of external chemical and mechanical 348 stimuli.

349

Our work shows that in giant cells, waves of both F-actin polymerization (Fig. 3) and its
upstream regulator PIP3 (Fig. 1 – figure supplement 1) are indeed guided by EFs. These biased
biochemical and biomechanical events lead to more protrusions at the cell front than at the cell
back, thus driving cell migration (Fig. 2 – figure supplement 2). The development of the biased

354	wave activities takes ~10 min following the introduction of an EF (Fig. 2a and Fig. 3), which is
355	much slower than the timescale of surface-receptor-regulated chemotaxis. The high resistance of
356	the cell membrane limits the effects of EFs on intracellular components, but EFs may act on the
357	charged lipids and molecular clusters. Thus, we suspect that the slow response results from the
358	electrophoresis of the charged membrane components involved in wave formation, which has a
359	characteristic time scale of 5 to 10 min (Allen et al., 2013; McLaughlin & Poo, 1981).
360	
361	We further explored the dynamics in response to EF reversal at the subcellular level using
362	nanotopography (Fig 4). We observed that the new waves are induced to propagate towards the
363	current cathode within 2 to 3 min (Fig. 4e and Fig.4 – figure supplement 2), suggesting that
364	waves themselves can adapt quickly to the changing electrical environments. Because we only
365	observed the fast adaptation on ridged surfaces, this phenomenon may be related to the shorter
366	wave lifetimes on nanoridges than on flat surfaces. A short lifetime allows waves to be nucleated
367	at a higher rate on the nanoridges, leading to a rapid directional response. During this process,
368	the EF may regulate the wave nucleation through locally changing specific charged lipids, ion
369	fluxes, or local pH gradients (Crevenna et al., 2013; Frantz et al., 2008; Köhler et al., 2012;
370	Martin et al., 2011; Zhou & Pang, 2018).
371	
372	EFs modulate the thresholds of the excitable wave system

373 Recent studies have shown that the cortical wave system can be described as a coupled signal
374 transduction and cytoskeletal excitable network. Based on both simulation and experimental
375 studies (Bhattacharya et al., 2020; Miao et al., 2017), it has been shown that the wave ranges,

durations, and speeds are determined by the local threshold of activation, which in turn are

377 regulated by the relative levels of activators and inhibitors (Miao et al., 2017, 2019).

378

379 Our quantification shows that guided waves become larger, faster, and more persistent in an EF 380 (Fig. 2), indicating that the excitable system is closer to its threshold for activation (Miao et al., 381 2019). This effect may arise from enhanced positive feedback, reduced negative feedback, or 382 both. We further find that wave nucleation is enhanced at the cell front and suppressed at the 383 back (Fig. 4a, b). This subcellular inhomogeneity is consistent with a biased excitable network 384 framework (Iglesias & Devreotes, 2012; Meinhardt, 1999; Tang et al., 2014; Xiong et al., 2010), 385 which was added to the STEN-CEN model to introduce an internal spatial gradient in the local 386 threshold of wave initiation, akin to cell polarity.

387

388 Local excitation and global inhibition (Xiong et al., 2010), LEGI, schemes have effectively 389 recreated the features of both fast directional sensing and stable polarity in response to chemical 390 signals, which can lead to robust biased excitable network. Both directional sensing and stable 391 polarity can lead to a robust biased excitable network. For chemical signals, the directional 392 response from PIP3 occurs within seconds, whereas the establishment of stable polarity usually 393 requires many minutes. However, based on our analysis, establishing both directional response 394 (Fig. 3) and polarity (Fig. 4) in response to EFs requires 5 to 10 min. It is worth noting that PIP3 395 waves also sense EFs on a time scale of minutes (Fig. 1 - figure supplement 1). Our observation 396 suggests that EFs act on the polarity establishment rather than directional sensing. This 397 hypothesis is supported by a recent study showing that G-protein-coupled receptors (GPCRs),

398 which are the regulator in the LEGI model for *D. discoideum* that allows for sensing

399 chemoattractant on timescales of seconds, are not essential for electrotaxis (Zhao et al., 2002).

400

#### 401 EFs act on waves, and waves determine cell behaviors

402 Our results raise the possibility that cortical wave dynamics are modulated directly by EFs and that the waves in turn mediate cellular response. Waves travel across cell membranes to 403 404 coordinate the trailing edge with the front edge, and the cytoskeletal components in cortical 405 waves are involved in developing the stable polarity. On the other hand, the duration and turning 406 capacity of STEN-CEN waves directly impact the speed and characteristics of the cellular 407 response to EFs (Fig. 3) on a longer timescale than that of surface-receptor-regulated chemotaxis. 408

409 Our results shed light on how EFs modulate protrusions. Previous studies have shown that 410 various protrusions that drive cell motion, such as filopodia, lamellipodia (Miao et al., 2019), and 411 macropinocytotic cups (Video 4), are always associated with expanding waves near cell 412 perimeter. Our previous work has shown that changing wave properties by perturbing STEN-413 CEN states leads to the transition of protrusion profiles, which indicates that wave properties 414 dictate the properties of the protrusions (Miao et al., 2019). Here we showed that EFs can alter 415 the waves differently on the two ends of the cell (Fig. 4a). As a result of these spatially 416 inhomogeneous wave properties, protrusions become more abundant and larger on one side of 417 the cell versus the other, which eventually leads to guidance of cell migration.

418

419 On flat surfaces, a slow U-turn is observed following EF reversal, whereas on nanoridges, faster 420 switching is observed. Thus, the response of migrating cells to a changing guidance cue can be

421 predicted from the characteristics of the waves driving the migration process. Indeed, the U-turn 422 behaviors of neutrophils and differentiated, single *D. discoideum* cells in response to EF reversal 423 (Hind et al., 2016; Sato et al., 2007; Srinivasan et al., 2003; Xu et al., 2003), which are usually 424 ascribed to stable cell polarity, may instead reflect the persistence and 2D turning behavior of 425 cortical waves in these environments (Fig. 3).

426

441

427 Nanoridges allow us to shed further light on the multiscale character of the system, because cells 428 include both short, 1D waves on the basal plane, and longer-lasting, 2D waves on the dorsal 429 plane. The different response times on the subcellular level due to different wave behaviors (Fig. 430 4 - figure supplement 2) provide strong evidence that cortical waves act as direct mediators of 431 EFs. Waves on different planes are similar in composition but are impacted differently by the EF. 432 We observed fast switching of wave directions in the basal plane near the ridged substrate and 433 slower turning of the waves in the dorsal plane within the same cell, indicating that the direction 434 of waves is controlled locally by external cues (Fig. 3 – figure supplement 1). 435 436 EFs provide a means to modulate cortical waves directly. On the other hand, biological 437 conditions that modulate wave characteristics may also speed up or suppress the cellular 438 response to directional cues. Longer-lasting waves offer persistence in the face of rapidly 439 changing gradients, whereas shorter waves yield faster adaptability to changing directional 440 signals. The durations of waves and their ability to turn together have a dominant effect on the

**Key Resources Table** 

response of cells to an EF.

Reagent type	Designation	Source or	Identifiers	Additional
(species) or		reference		information
resource				
Cell line (D.	Aca null	https://doi.org/1		The cell line
discoideum)		0.1016/S0092-		was a gift from
		8674(03)00081-		Carole A.
		3		Parent lab.
Cell line (D.	PHcrac-GFP	https://doi.org		The cell line
discoideum)	LimE-RFP	/10.1038/ncb		was a gift from
		3495		Peter N.
				Devreotes lab.
software,	Optical flow	https://doi.org		
algorithm	analysis (run by	/10.1091/mbc		
	MATLAB)	.E19-11-0614		

# 442

# 443 Materials and Methods

444 <u>Cell line</u>

445 In the study, we used LimE-RFP aca null Dictyostelium discoideum (D.d.) and PHcrac-

- 446 GFP/LimE-RFP D.d cell lines. LimE-RFP aca null was a gift from Carole A. Parent lab
- 447 (https://doi.org/10.1016/S0092-8674(03)00081-3), and PHcrac-GFP-LimE-RFP was a gift from

448 Peter N. Devreotes lab (https://doi.org/10.1038/ncb3495). We have conducted the mycoplasma

449 contamination testing for both cell lines and did not detected contamination.

450

451 <u>Cell culture</u>

452 Dictyostelium discoideum cell lines were grown axenically in the HL5 medium. Aggregation

453 adenylyl cyclase null (ACA-) mutants, which do not produce cAMP and do not have chemotaxis

454 signal relay (Kriebel et al., 2003), were used in electrotaxis experiments to avoid chemotaxis.

455 The cells used also express limE-RFP as a reference for filamentous actin structures. G418 was

456 used as the selection medium during cell culture. For the experiments in Fig. 1/Fig.1 -

457 supplement figure 1, we used Dictyostelium discoideum co-expressing PHcrac-GFP and LimE-

458 RFP, and we used G418 as the selective medium. Note that an enhancement in LimE

459 concentration is associated with protrusions, as seen in fig. 2 -figure supplement 2, and that the

460 protrusions are biased to the side facing the cathode. Because protrusions are driven by F-actin

461 polymerization, we believe this observation rules out the possibility that LimE binding/unbinding

462 to/from F-actin itself is sensitive to EFs.

463

### 464 <u>Electrofusion</u>

465 Cells were washed twice with 17 mM Sorensen buffer (15 mM KH<sub>2</sub>PO<sub>4</sub> and 2 mM Na<sub>2</sub>HPO<sub>4</sub>,

466 pH 6.0) and rolled for 30 min at a concentration of  $1.5 \times 10^7$  mL<sup>-1</sup>. Electrofusion was conducted

467 with a Gene Pulser Gen1 system. Three pulses of 1 kV at a 1 s interval were applied. After

468 electroporation, cells were relaxed for 5 min. Then cells were diluted to  $5 \times 10^5 \,\text{mL}^{-1}$  with normal

developing buffer (5 mM KH<sub>2</sub>PO<sub>4</sub>, 5 mM Na<sub>2</sub>HPO<sub>4</sub>, 2 mM MgCl<sub>2</sub> and 0.2 mM CaCl<sub>2</sub>, PH 6.5)

470 and seeded into a customized electrotactic chamber, with dimensions 20 mm  $\times$  5 mm  $\times$  0.25 mm.

471	The even will call line that we used did not	concrete many ways in the wasstative stage and
4/1	The aca null cell line that we used the not	generate many waves in the vegetative stage, and

472 electro-fusion with 1 kV pulses stressed the cells. Thus cells were starved for 2 h before

473 experiments to generate more actin waves.

474

475 <u>Nanotopography fabrication</u>

476 The nanotopographic pattern used in these cell studies was fabricated through a technique known

477 as multiphoton absorption polymerization (MAP), as described elsewhere (C. M. LaFratta et al.,

478 2006; C. N. LaFratta et al., 2004). An ultrafast, pulsed laser beam (Coherent Mira 900-F, 76

479 MHz) was passed through a high-numerical-aperture microscope objective onto a

480 photopolymerizable resin, sandwiched between glass coverslips. A LabVIEW (National

481 Instruments) program allowed for control of the stage position and the shutter state, determining

482 where polymerization occurred (and did not) in the resin and allowing patterning. Once

483 fabrication was completed, the patterned sample was developed in ethanol twice for 3 min each

484 to remove unreacted monomer. The polymerized structure was baked at 110 °C for at least 1 h.

485

486 To produce the necessary number of replicate patterns with the same dimensions, an adapted

487 version of replica molding was performed (Sun et al., 2018). A hard polydimethylsiloxane (*h*-

488 PDMS) film containing hexanes to increase the resolution of feature replication was spin-coated

489 onto the functionalized structure made from MAP. The film was allowed to sit on the structure

490 for 2 h at room temperature and was then baked at 60 °C for 1 h. Regular PDMS (Sylgard 184)

491 was prepared at a 10:1 ratio of elastomer base to the curing agent by degassing and mixing. The

492 PDMS was poured onto the *h*-PDMS film, and molding was completed by baking at 60 °C for an

493 additional 70 min. The final mold was peeled from the glass slide supporting the MAP-patterned494 structure.

495

496 The mold was used to produce replicas of the original pattern. A drop of the same acrylic resin 497 was placed on the patterned area of the PDMS mold, and then an acrylate-functionalized glass 498 coverslip was pressed firmly on top, spreading the sandwiched drop. Tape secured this system in 499 place. The resin was cured for a total of 5 min under a UV lamp (Blak-ray), producing a polymer 500 film. It should be noted that the PDMS mold is the negative relief pattern of the structure made 501 using MAP. Therefore, samples (or replicas) of the original pattern could be produced on a 502 relatively large scale with this method. The replicas were soaked in ethanol for at least 12 h 503 before use in the cell studies. We fabricated flat surface samples by using a PDMS mold with a 504 smooth surface. 505 506 Lattice light-sheet microscopy 507 The 3i lattice light-sheet microscope in the Johns Hopkins School of Medicine Microscope

508 Facility was used for 2-color, 3D imaging. Vegetative, single *Dictyostelium* cells were seeded on

a circular 5 mm coverslip patterned with nanoridges, which was immersed in a bath of standard

510 developing buffer throughout imaging.

511

512 <u>Electrotaxis experiments</u>

513 We 3D-printed electrotaxis chambers (Fig. 2 – figure supplement 1) with dimensions of 20 mm  $\times$ 

514 5 mm  $\times$  0.25 mm and composed of a clear resin using a Formlabs Form2 3D-printer. Agar

515 bridges were used to isolate cell media from electrodes to minimize electrochemical products

517 channel and the RFP/GFP channel were recorded using PerkinElmer spinning-disk microscope at

518 a frame rate of 0.1 frames/s (Yokogawa CSU-X1 spinning-disk scan head (5000 rpm)) with

- 519 Hamamatsu EMCCD camera and Volocity analysis software.
- 520

521 <u>Optical-flow analysis and model fitting of actin polymerization dynamics</u>

522 We applied the Lukas-Kanade optical-flow method to quantify the direction of the intensity flow

523 in fluorescence videos. This algorithm produced pixel-basis vector fields of intensity motion.

524 Before applying the optical-flow algorithm, each image was smoothed by a 2D Gaussian filter ( $\sigma$ 

525 = 3) to reduce noise. After the smoothing, we further removed the flow vectors created by noise

526 using optical-flow reliability as our criterion. The reliability is defined as the smallest

527 eigenvalues of the  $A^T w A$  matrix, where w is a Gaussian weight matrix and A is the intensity

528 gradient matrix. The size of the weight matrix for *D. discoideum* was set at  $19 \times 19$ , with

529 standard deviation  $\sigma = 2 (0.42 \,\mu\text{m})$ .

530

We built a bimodal von Mises model to compare the actin and cellular responses accurately. A
von Mises distribution is given by

533 
$$f_{VM}(\theta|\mu,\kappa) = \frac{e^{\kappa\cos(\theta-\mu)}}{2\pi I_0(\kappa)}.$$
 (2)

where the peak location is  $\mu$  and the concentration  $\kappa$ . The orientation distribution of optical-flow vectors at each time point is fit with two von Mises distributions

536 
$$f(\theta|\mu_1, \mu_2, \kappa, p_1, p_2) = p_1 f_{VM}(\theta|\mu_1, \kappa) + p_2 f_{VM}(\theta|\mu_1 + \pi, \kappa)$$
(3)

537 where *p* is the proportion of each component. We use the constraints

538 
$$\mu_1 - \mu_2 = \pi$$
 (4)

539 and

540

 $p_1 + p_2 = 1. (5)$ 

541 Maximum likelihood estimation (MLE) is applied to estimate model parameters based on the 542 orientation of all the optical-flow vectors every 12 frames (2 min). With this model, we can quantitatively study the temporal change of actin dynamics. The preferential direction is defined 543 544 as the  $\mu$  with the largest proportion p at each time point. 545 546 Quantification of actin wave properties 547 The segmentation of actin waves was conducted based on the combined information from 548 fluorescence intensities and optical flow. We first applied the kmeans (k = 3) cluster (Kanungo et 549 al., 2002) to pick up the bright regions in the limE-RFP videos, then only kept the moving 550 objects by applying the reliability mask from the optical-flow analysis.

551 To classify the large actin structures composed of substructures moving independently, we

considered the optical flow at the edges of the large structure (Fig. 2c). If the optical-flow vectors

553 were moving in the same direction at both edges, the large structure was classified as a single

554 wave. Otherwise, the large structure was divided into multiple smaller patches. In the latter case,

a pronounced boundary was detectable between two substructures, then we used the detected

556 boundary to divide the large structure into multiple substructures.

557

After classification, we measured properties such as wave speed, wave duration, and wave area to characterize STEN-CEN. Wave speed was measured by tracking the clusters of optical-flow vectors oriented in similar directions. The detailed algorithm can be found in a prior publication (Lee et al., 2020). To measure wave duration, we first tracked actin waves using a customized,

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	muni-oneci		Dascu on un	E OVEHADDINE AIEAS	DELWEEN HAINES.	A uniuuc

- 563 identification number was assigned to each wave, then wave duration, wave area (measured by
- the Matlab function regionprops) were recorded for each wave.
- 565
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567 The authors declare that they have no competing interests.

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569 References:
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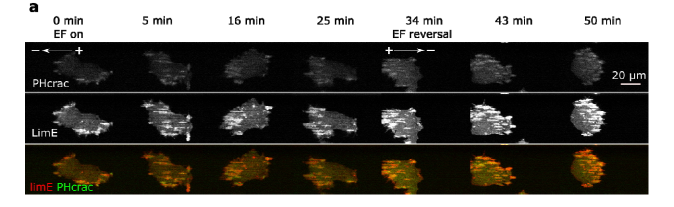
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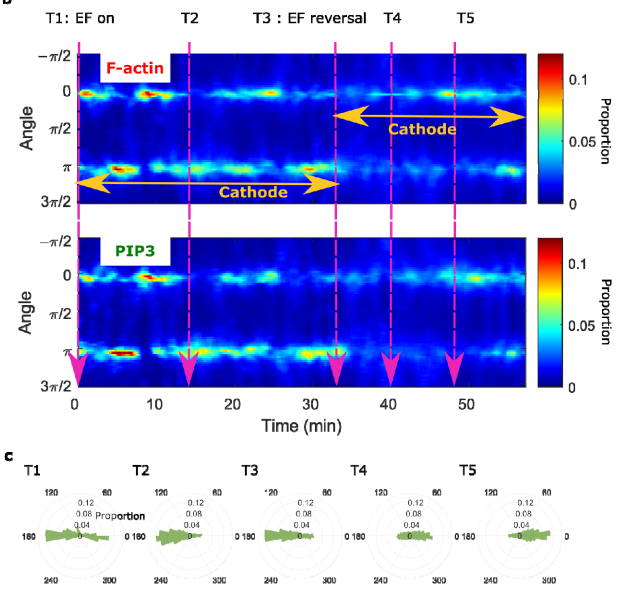
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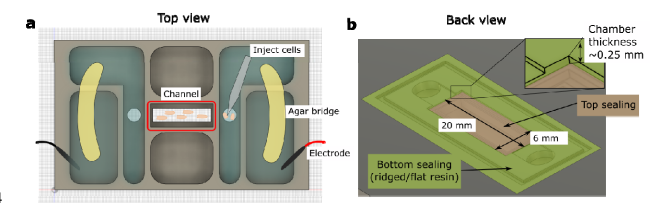




743 Fig. 1 – figure supplement 1. Colocalization of PIP3 and F-actin in an EF. a. Images of

744	PHcrac (top row) and LimE (middle row) in a 20 V/cm EF with the cathode located on the left.
745	The EF was turned on at 0 min and reversed at 34 min. From the combined images (bottom row),
746	it is clear that F-actin and PIP3 remained coordinated during electrotaxis. b. Kymograph of
747	angular distribution of F-actin (top) / PIP3 (bottom) wave motion. Optical-flow analysis was
748	applied to both the F-actin and PIP3 videos to measure wave motion. Then we stacked up the
749	angular distribution at each time point along the x-axis. The dynamics of F-actin/PIP3 in
750	response to EFs are similar. The color was coded according to the proportion of each bin in the
751	angular distribution. c. Polar plots of angular distributions from the time points specified in b.
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765 Fig. 2 – figure supplement 1. Schematic of the 3D-printed chamber used for electrotaxis

766 **experiments. a.** Top view of the experimental setup. **b.** Back view of the setup. Agar bridges

- 767 isolate the cell media from electrodes to avoid changes in pH and the generation of
- relectrochemical products. The cells were injected using pipettes into the center channel (red box
- highlighted in a). The height of the electrotaxis channel is ~0.25 mm. The channel was sealed
- with a large substrate coverslip and a smaller top coverslip.

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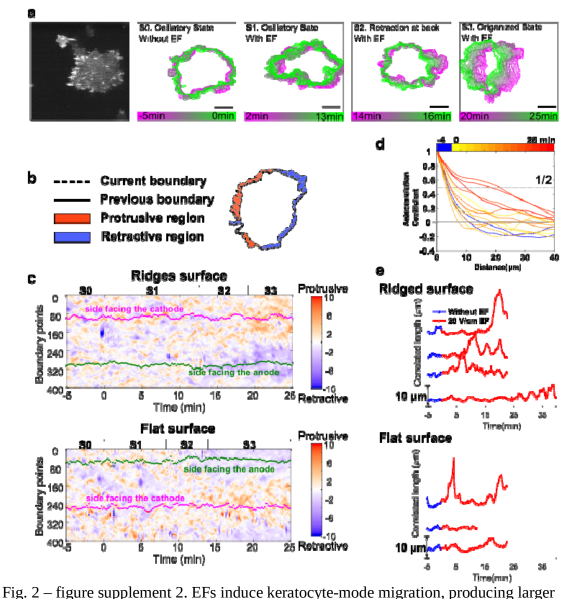
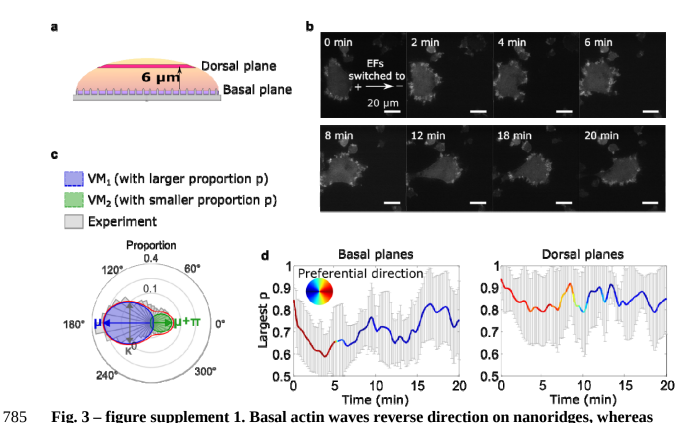
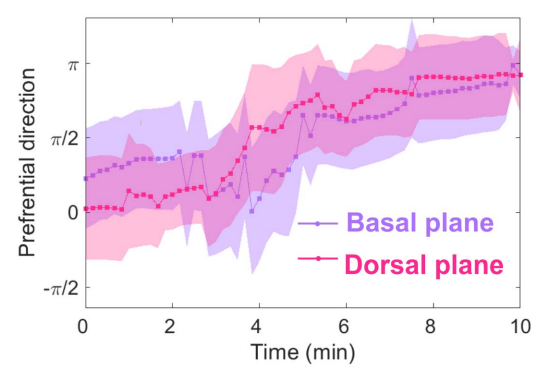


Fig. 2 – figure supplement 2. EFs induce keratocyte-mode migration, producing larger protrusions at cell fronts. **a**. Shape dynamics of a giant *D. discoideum* cell at 10 s intervals. Cell boundaries were outlined using an active contour algorithm. Boundaries are color-coded according to time. Four stages were categorized: (S0) Random motion in the absence of an EF. (S1) Random motion in the first 15 min in the presence of an EF. (S2) Transition state usually with a retraction at the back. (S3) Steady migration state in the presence of an EF. **b**. A schematic of local protrusion and retraction, where the solid line is the current frame, and the dotted line represents the cell boundary 1 min later. A protrusive region (yellow) is defined as

- one occupied in the new frame, not in the previous frame, and a retractive region (blue) is
- 780 defined as one occupied in the previous frame, not in the new frame. c. Kymograph of local
- boundary motion. The *x*-axis represents time, the *y*-axis indicates boundary points, and the color
- 782 of each pixel corresponds to the speed. **d**. Correlation curves of boundary motion at different
- time points. e. Correlation length (defined as the point at which curves in d reach a value of 1/2)
- vs. time. Each curve represents an independent experiment conducted on a different day.



786 dorsal waves turn. a. A schematic showing the two imaging planes used, with the morphology 787 of the substrate. **b.** LimE-RFP images recorded at the dorsal plane. Unlike the basal focal plane 788 images, which capture the complete basal wave dynamics, the dorsal plane images do not capture 789 the full dorsal wave motion. To avoid photobleaching and laser damage, we only imaged the 790 cross-section of the dorsal waves and tracked the cross-sections using optical-flow analysis. 791 c. A schematic introducing 4-parameter, bimodal von Mises (vM) fitting. The orientation 792 distribution (gray) was fitted to a bimodal vM distribution at each time point. The two vM 793 distributions were set to share the degree of concentration  $\kappa(\kappa_1 = \kappa_2)$ , peak locations  $\mu$  that were 794 apart ( $\mu_1 = \mu_2 + 180^\circ$ ). **d.** Temporal evolution of fitting parameters in response to set to be 795 EF reversal. Among two fitted vM distributions, the one with a larger proportion  $(vM_p)$  was 796 tracked. For both plots, the curves represent the mean p of  $vM_p$  with standard derivations from 797 multiple experiments ( ), and the colors represent the  $\mu$  of vM<sub>p</sub>.

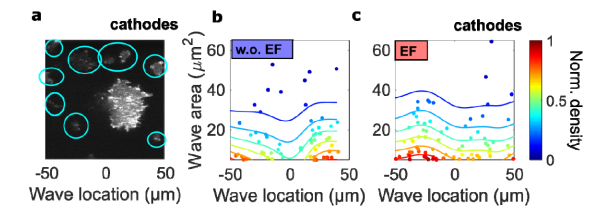


798 Fig. 3 – figure supplement 2. Basal and dorsal waves on flat surfaces both turn in response

799 to EF reversal. Temporal evolution of cosine function of the fitted preferential direction of actin

800 wave propagation for giant cells on flat surfaces. The analysis was based on N = 3 experiments.

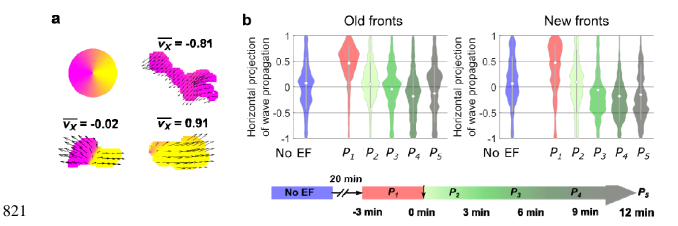
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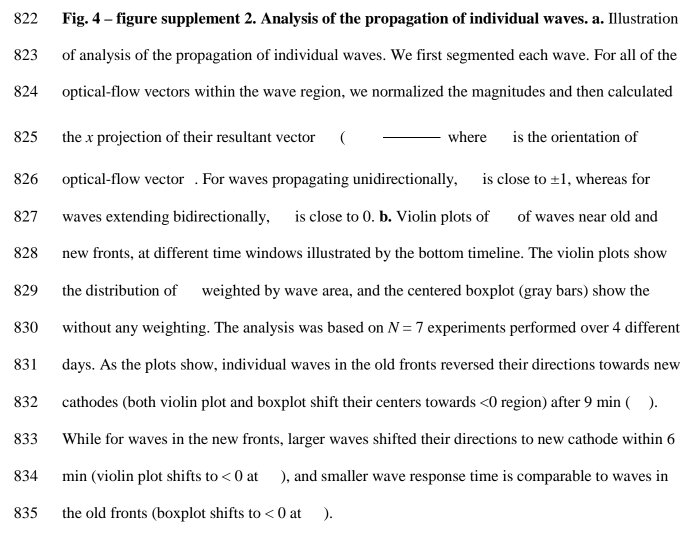


802 Fig. 4 – figure supplement 1. The spatial inhomogeneity of wave properties could be caused 803 either by the EF or by the external electrical potential gradient relative to the ground. To 804 explore these scenarios, we quantified the waves in single cells surrounding the giant cell in the 805 field of view, where the center of the field was defined as the origin. In contrast to the giant cells, 806 these single cells are scattered throughout the field of view but are not large enough for the 807 potential gradient to create significant intracellular polarization. Thus, if the spatial 808 inhomogeneity is caused by the external electrical potential gradient, we would observe a 809 gradient of wave properties from single cells located in the region between  $-50 \,\mu\text{m}$  and  $50 \,\mu\text{m}$ . 810 A limE image. Single cells are highlighted with blue circles. **b**, **c**. Density scatter plots of wave 811 location vs. wave area for single cells (highlighted by blue circles in a) in the absence (b) and the 812 presence (c) of EF. Unlike in giant cells (Fig. 4a, 4b), the wave areas in single cells are spatially 813 homogeneous. The ratio of mean wave areas in the regions nearer the cathode (location > 0) to 814 those in regions farther from the cathodes (location < 0) was calculated for both single cells and 815 giant cells. In giant cells, this ratio increases from 0.85 to 2.00 with an EF (Fig. 4a), whereas for 816 single cells, the ratio is almost unchanged (1.08 without an EF and 0.98 with an EF). This 817 analysis was based on the experiments from 4 different days. 818 Figure 4 – figure supplement 1- source data 1. Related to Fig. 4 – figure supplement 1.

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836 Figure 4 – figure supplement 2- source data 1. Related to Fig. 4 – figure supplement 2b.