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Insect hapto-electrical stimulation of Venus flytrap triggers exocytosis in gland cells

- Authors: Sönke Scherzer¹, Lana Shabala², Benjamin Hedrich^{2†}, Jörg Fromm³, Hubert Bauer¹, 3 Eberhard Munz⁴, Peter Jakob⁵, Khaled Al-Rascheid⁶, Ines Kreuzer¹, Dirk Becker¹, Monika 4 Eiblmeier⁷, Heinz Rennenberg⁷, Sergey Shabala², Malcolm Bennett⁸, Erwin Neher^{9*} and Rainer 5 Hedrich^{1*} 6 Affiliations: 7 ¹ Institute for Molecular Plant Physiology and Biophysics, University Wuerzburg, D-97070 8 Wuerzburg, Germany; 9 ² School of Land and Food, University of Tasmania, Hobart, TAS, Australia; 10 ³ Universität Hamburg, Zentrum Holzwirtschaft, D-21031 Hamburg, Germany; 11 ⁴ Leibniz Institute of Plant Genetics and Crop Plant Research, D- 06466 Gatersleben, 12 Germany 13 ⁵ Experimental Physics 5, University of Würzburg, D-97070 Wuerzburg, Germany; 14
- ⁶Zoology Department, College of Science, King Saud University, Riyadh 11451, Saudi
 Arabia;
- ⁷ Chair of Tree Physiology, Institute of Forest Sciences, University of Freiburg, D-79110
 Freiburg, Germany;
- ⁸ Centre for Plant Integrative Biology, School of Biosciences, University of Nottingham,
 LE12 5RD, UK;
- ⁹ Department for Membrane Biophysics, Max Planck Institute for Biophysical Chemistry, D-37077 Goettingen, Germany

- ²⁴ *Correspondence to: R.H. (hedrich@botanik.uni-wuerzburg.de) or E.N. (eneher@gwdg.de).
- ²⁵ [†]Present address: Medical University of Graz, 8010 Graz, Austria
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26 Abstract

27 The Venus flytrap *Dionaea muscipula* captures insects and consumes their flesh (1, 2). Prey contacting touch-sensitive hairs trigger travelling electrical waves. These action potentials 28 (APs) cause rapid closure of the trap and activate secretory functions of glands, which cover its 29 inner surface (3, 4). Such prey-induced hapto-electric stimulation activates the touch hormone 30 jasmonate (JA) signaling pathway, which initiates secretion of an acidic hydrolase cocktail to 31 decompose the victim and acquire the animal nutrients (5-7). Although postulated since 32 Darwin's pioneering studies these secretory events have not been recorded so far. Using 33 advanced analytical and imaging techniques, such as vibrating ion selective electrodes, carbon 34 35 fiber amperometry and MRI, we monitored stimulus-coupled glandular secretion into the flytrap. Trigger hair bending or direct application of JA caused a quantal release of oxidizable 36 material from gland cells monitored as distinct amperometric spikes. Spikes reminiscent of 37 38 exocytotic events in secretory animal cells progressively increased in frequency, reaching steady state one day after stimulation. Our data indicate that trigger hair mechanical stimulation 39 40 evokes APs. Gland cells translate APs into touch-inducible JA signalling that promotes the formation of secretory vesicles. Early vesicles loaded with H⁺ and Cl⁻ fuse with the plasma 41 membrane, hyper-acidifying the 'green stomach'-like digestive organ, while subsequent ones 42 carry hydrolases and nutrient transporters together with a glutathione redox moiety, which is 43 likely to act as the major detected compound in amperometry. Hence, when glands perceive the 44 hapto-electrical stimulation, secretory vesicles are tailored to be released in a sequence, which 45 optimizes digestion of the captured animal. 46

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48 Significance statement:

The Venus flytrap has been in the focus of scientists since Darwin's time. Carnivorous plants,
with their specialized lifestyle, including insect capture, as well as digestion and absorption of

prey, developed unique tools to gain scarce nutrients. In this study we describe novel mechanistic insights into the cascade of events following the capture of insect prey. Action potentials evoked by the struggling prey are translated into touch-inducible hormone signals that promote the formation of secretory vesicles. A variety of digestive compounds are released sequentially into the flytrap's 'green stomach' and break down the captured animal. Amperometry provides insight into the kinetics and chemistry of the stimulus-coupled glandular secretion process.

58 **body**

59 Introduction:

Certain plants have turned the sword; they capture and consume animals, including potential 60 61 herbivores. Growing on mineral-deficient soils, the carnivorous Venus flytrap (Dionaea muscipula) lures (8), captures, and digests small arthropods, in order to feed on the nutrients 62 extracted from their flesh (1, 3, 9-12). Closure of the bilobed snap trap is initiated by mechanical 63 stimulation of trigger hairs located at the inner trap surface. Each trigger hair bending elicits 64 the firing of an action potential (AP). With the first AP, the trap stays open, but memorizes the 65 initial strike. If a second one fires within 20 s, it triggers rapid trap closure. In case an insect is 66 trapped and struggles to escape, two and more hapto-electric stimuli activate jasmonate 67 signaling and biosynthesis (3, 6, 7). From the fifth strike on, glands raise their expression levels 68 of hydrolase and nutrient transporter genes. When mechano-stimulation is replaced by 69 application of coronatine (COR), a mimic of the biologically active jasmonate hormone JA-Ile, 70 71 it can substitute for the mechano-electric stimulation of the flytrap (7). Hapto-electric signaling and touch hormone activation turn the closed trap into a 'green stomach', flooding the entrapped 72 prey with an acidic digestive fluid (3, 6, 13). Although prey capture and consumption of the 73 Venus flytrap has been known since Darwin's time (2), the molecular mechanisms of fluid 74 phase secretion underlying animal consumption remained unknown (14). In this study 75

amperometric carbon fibers were used for the first time in the plant field to monitor the
dynamics and kinetics of mechano-electric and JA stimulation of the secretory events,
providing insight into exocytosis-dependent liquor filling of the digestive organ.

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80 **Results:**

Upon hapto-electric trap activation the surface area of the multicellular gland cell complex 81 increases by 30% and an acidic protein moiety is released into the 'green stomach' formed by 82 the hermetically sealed lobes of the trap (3, 5-7, 13). In search of the membrane reservoir 83 responsible for the surface increase of stimulated glands, we exposed traps to the JA-Ile mimic 84 85 coronatine (COR). 48 h after stimulus onset membrane pits, observed in electron micrographs (EM) of glands suggested that secretory vesicle fusion had taken place predominantly at the 86 apical end of head cells (outermost cell layer; L1) (Fig. 1 and S3B). Head cells of non-87 88 stimulated glands only occasionally showed exocytotic vesicles (1.5 ± 0.4 per cell; Fig. 1A and C) but in the outermost layer of jasmonate-stimulated glands cells, we detected a pronounced 89 90 increase of pits associated with the more apical plasma membrane sections (16.5 ± 1.5 per cell, approximately 0.18 µm in diameter; Fig. 1B, D and E). These results indicate that the secretory 91 stimulation causes granule docking and membrane fusion. 92

93

94 <u>MIFE resolves early secretion of acidic vesicles</u>

In a previous study, we compared the transcriptomic profile of non-stimulated glands with that of glands stimulated either by insects or COR. Before stimulation, the transcription profile of resting glands is already dominated by secretory processes (7). *Dionaea* secretion is directly coupled to acidification; H^+ and chloride, Cl^- , are released into the digestive fluid of the tightly sealed trap (15). To test whether touch stimulation of the flytrap's trigger hairs is translated into ion fluxes across the gland plasma membrane, we used Ca^{2+} , Cl^- , and H^+ sensitive MIFE microelectrodes (3, 16), which measure fluxes by recording local concentration gradients. After

5 - 10 consecutive trigger-hair stimulations and a lag time of about 10 min, a rapid shift in the 102 net ion fluxes towards net Ca²⁺ uptake into the gland cells was observed (Fig. 2A). The mean 103 net Ca²⁺ flux after mechanical stimulation (5 APs) of the Venus flytrap was about 9.9 ± 1.8 104 nmol $m^{-2} s^{-1}$ (Fig. 2B; mean \pm SE, n = 6). Within the first hour following stimulation, the ion 105 fluxes were dominated by Ca^{2+} fluxes. Upon Ca^{2+} entry, the intracellular Ca^{2+} level rises (6), 106 and JA signaling is activated (3, 7). Either consecutive trigger hair stimulation alone or a direct 107 application of jasmonates or COR induces secretion. With jasmonates secretion in traps is 108 initiated before they close (6). Following application of JAs, however, Ca²⁺ sensitive MIFE 109 electrodes did not record net Ca²⁺ flux into glands (Fig. 2A, B; red symbols/bar). Hormone 110 stimulation however, triggered proton release, which appeared within 5 - 10 min following 111 stimulation onset (Fig. 2C, D). Net H⁺ efflux reached its peak between 1 and 2.5 h after stimulus 112 application and then gradually recovered (Fig. 2C). When comparing the time that glands 113 114 required to reach peak proton extrusion in response to mechanical or chemical stimulation, JAs were the fastest (Fig. 2C). Thus jasmonate-induced proton release was significantly faster than 115 that elicited by mechanical stimulation (insert in Fig. 2C), which reached peak currents of 54 \pm 116 7 nmol m⁻² s⁻¹ (mean \pm SE, n = 6). Also the lag time of H⁺ efflux resulting from the different 117 stimulations was longest in response to mechanical stimulation (Fig. 2D). This time dependence 118 fits the notion that the rise in gland JA is downstream of hapto-electrics and gland calcium 119 entry. 120

Regardless of whether stimulated or not, the resting membrane potential of glands remained in the range of -120 to -140 mV (12). This might indicate that trap acidification results from electroneutral exocytotic H⁺ release rather than the massive activation of plasma membrane proton pumps. This notion is supported by the COR induced increase in vacuolar AHA10-type proton pump transcripts (17), together with those of a ClC-type proton-chloride antiporter (18), two components required for hyper-acidification of secretory vesicles (see Fig. S1 and supplementary text S1). To test whether H⁺ fluxes are accompanied by Cl⁻ fluxes, we used 5

chloride-sensitive MIFE electrodes side-by-side with the pH microelectrodes. Confirming our 128 working model, we monitored pronounced Cl⁻ net efflux from glands in COR stimulated (Fig. 129 2E; blue symbols), but not in resting (Fig 2E; grey symbols) traps. COR induced chloride 130 currents appeared with a similar time dependence and amplitude as the proton fluxes (Fig. 2C, 131 E). Both fluxes were correlated with each other ($R^2 = 0.61$; P < 0.01), exhibiting a stoichiometry 132 between H⁺ and Cl⁻ close to 1:1 (Fig. 2F). The electrochemistry-based MIFE experiments 133 illustrated above can only be conducted in an aqueous environment. In such a wet scenario, we 134 monitored initial secretion-associated proton extrusion in response to COR about 9 min after 135 stimulation (Fig. 2D). To resolve the onset of gross gland fluid secretion in the initially dry 136 137 Dionaea trap, we followed the fluid production after COR stimulation by Infra Red Gas Analysis (IRGA) and magnetic resonance imaging (MRI). First fluid phase secretion-associated 138 trap water vapor emission was detected in IRGA recordings $151 \pm 13 \text{ min}$ (n = 3, mean \pm SD) 139 140 following trap stimulation with COR (Fig. S2A). After reaching peak humidity trap water emission slowly decreased and suddenly dropped after $445 \pm 84 \min (n = 3, \text{mean} \pm \text{SD})$ to the 141 142 basal level of evaporation prior to COR application and non-stimulated controls (Fig. S2A). This rapid drop in water emission reflects hermetical sealing of the trap lobes (6). Filling of the 143 closed trap with digestive fluid was visualized by MRI imaging (Fig. S2B and supplementary 144 video 1). 145

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147 Detection of digestive vesicles via Amperometry

With animal cells exocytotic events can be monitored non-invasively via amperometry, detecting redox currents when electrodes are placed near the membrane surface of secretory cells (19, 20). Given that amperometry detects oxidizable substances such as neurotransmitters, neuropeptides and hormones released from secretory vesicles, we adopted this electrochemical approach to probe for exocytotic events in active flytrap glands. Aiming to detect spikes associated with secretory cargo release from inner trap surface, we placed carbon fiber

microelectrodes in contact with the apical face of the glands upper head cells (Fig. S3A and B). 154 155 Under these experimental conditions no amperometric signals were detectable in nonstimulated glands (Fig. S3C). However, with glands stimulated by 5 - 20 trigger hair 156 displacements, signals similar to those measured with secretory animal cells could be monitored 157 (21, 22) (Fig. 3A), albeit with a much slower time course due to cell wall geometry. When 158 placing two electrodes next to each other both electrodes recorded characteristic increases in 159 160 amperometric current in close temporal relationship, as shown in Fig. 3A and B, excluding the possibility, that such discrete events were artifacts generated in one or the other electrode. In 161 these experiments we had to use strong pH-buffering in order to preserve the sensitivity of the 162 163 amperometric electrodes (see Methods).

The amperometrically detected chemical species is released to the apoplast at the point of 164 exocytosis. From that point source the released substance diffuses to the electroactive tip of the 165 166 carbon fiber where it is oxidized. It has been shown that placing electrodes more than several microns away from the cell surface results in a significant decrease in signal and spatio-167 168 temporal resolution (23, 24). Therefore, the best scenario for detecting exocytotic events without diffusional dilution is to actually touch the cell surface with the electrode. This 169 limitation by diffusion can be described by Fick's law. Thus we fitted the amperometrically 170 detected spikes with a 3D diffusion equation according to eq. 1 (see Methods). From this 171 calculation we gained a parameter t_c, which is a characteristic diffusion time, depending on the 172 distance between the point source of secretion and the carbon fiber tip, given a certain diffusion 173 coefficient D. Fitting sharp secretory events observed when electrodes were placed directly on 174 the Dionaea gland surface with a high spatio-temporal resolution, resulted in t_c-values of about 175 4-5 s. Plotting the relative signal abundance against the calculated t_c-values of detected 176 secretory events, a broadly homogenous distribution was obtained (Fig. S3D). In other words, 177 the amperometric approach we used, detects secretory events originating from various distances 178 to the tip of the carbon fiber, or else implies a range of diffusion coefficients. Interestingly, we 179

did not obtain any t_c-values ≤ 3.25 s in 63 analyzed spikes. Assuming a constant D-value in the 180 181 performed experiments and $t_c \ge 3.25$ s, we can calculate a lower bound for the geometrical distance between the point of secretion and the carbon fiber (eq.2, see methods) and D (diffusion 182 constant of the secreted substance in the medium) ((2) see Methods). In contrast to animal cells, 183 the plasma membrane of plant cells is covered with an extra layer of cellulose-based cell wall 184 and a lipid-based cuticle. Thus the minimal t_c-value obtained for *Dionaea* glands very likely 185 results from the cell wall-cuticle shell that keeps the fiber electrode at some distance (r) from 186 exocytotic vesicles fusing with the gland cell plasma membrane. From electron micrographs 187 similar to those shown in figure 1 we calculated a minimal distance between electrode and 188 189 secreted vesicle fusing with the head gland cell plasma membrane of ~0.5 μ m (Fig. 1B). Introducing this value in the equation (2) we are able to calculate the diffusion constant (D) of 190 the Dionaea secreted fluid in its diffusion medium (containing the cell wall and cuticle). The 191 calculated value of $D = 1.92 \times 10^{-10}$ cm²/s indicates a high diffusional resistance of the gland cell 192 wall. For comparison, the diffusion coefficient of dopamine in water was reported with 6.0x10⁻ 193 194 6 cm²/s (25). Also in the animal system diffusion in tissue or in solutions containing biological macromolecules is known to be hindered by the cellular matrix. Hafez et al. (26) have reported 195 that the diffusion coefficient of dopamine at the surface of an adrenal cell is one-tenth compared 196 to its diffusion in water. The small diffusion constant reported here for Dionaea also illustrates 197 the slow time characteristics of the detected amperometrical spikes with a half-life $(t_{1/2})$ time 198 constant of 87.82 ± 12.14 s (mean \pm SE; n = 92). Compared to the free aqueous diffusion of 199 catecholamine release in neuronal cells, $t_{1/2}$ of *Dionaea* plant secretory events is enlarged by a 200 factor of ~10,000 (24, 26). 201

To determine emergence and manifestation of gland cell exocytosis, we monitored the frequency of secretory events for up to 145 h. Traps were stimulated either mechanically by a series of 20 consecutive trigger hair bendings or by spraying COR onto the traps' inner surface. Within the first 4 - 5 h after stimulation onset, no significant signals could be monitored. First

exocytosis-type spiking was observed after about 6 h (Fig. 3D). Thereafter, exocytotic events 206 occurred more frequently, reaching about half-maximum spiking after 12 - 13 h. Maximal 207 spiking rates were detected after 24 h and remained high for another 2 days before slowly 208 declining at days 4 and 6 (Fig. 3D). Interestingly, COR stimulation and trigger hair bendings 209 resulted in a similar time dependence of spiking frequency. This indicates that jasmonate 210 induction of secretory vesicle formation, loading and membrane fusion, rather than touch 211 induction of jasmonate biosynthesis, represents the rate-limiting step during *Dionaea* gland cell 212 exocytosis. 213

We also found that the Ca^{2+} channel blocker gadolinium strongly reduced the volume of 214 secreted fluid (supplementary video 2). In order to further investigate the inhibitory effect of 215 Gd^{3+} on trap secretory fluid production, traps were sprayed with 10 mM Gd^{3+} (~2.5 µmol) 24 h 216 before mechanical stimulation. This Gd^{3+} challenge did, however, not affect the traps' naturally 217 218 fast closure in response to two trigger hair strikes. When traps were mechanically stimulated for secretion by 5 - 20 trigger hair displacements, gadolinium sprayed traps were found to be 219 220 strongly reduced in extruded fluid volume. Compared to control traps, which secreted 2.12 \pm 0.67 μ l/1000 glands within 48 h, Gd³⁺ pretreated traps released only 0.35 ± 0.25 μ l/1000) (Fig. 221 3C, black bars). At the same time, exocytotic events amperometrically determined with single 222 gland cells dropped from 14.3 \pm 4.17 events/h in controls to 2.1 \pm 2.45 events/h in the Gd³⁺ 223 exposed traps (Fig. 3C, red bars). The pronounced Gd³⁺ block of secretion seen by amperometry 224 and MRI suggests that JA and calcium signaling is required for hapto-electric and JA 225 stimulation of Dionaea gland cell secretion. 226

What kind of redox moiety *Dionaea's* secretory gland cells release? In order to gain and maintain functional integrity of cysteine-rich hydrolytic enzymes exuded into the digestive fluid (13, 27, 28), a defined redox status in the extracellular bioreactor is required. Glutathione (GSH) represents an important redox regulator of enzyme functions in plant cells (29, 30). Glutathione is synthesized via a well-known enzymatic pathway (see Fig. S4). Glutathione can be derived

from activated sulfate (APS, adenosine 5'- phosphosulfate) via a well-known enzymatic 232 233 pathway (see Fig. S4). Gene expression analysis based on RNAseq data (available at http://tbro.carnivorom.com, c.f. (7)) indicated that coronatine might induce genes involved in 234 GSH production and transport. These analyses were further confirmed by quantitative RT-PCR. 235 Among these genes the APS reductase (DmAPR3) is strongly upregulated 12 h after COR 236 stimulation (Fig. 4B). APS reductase represents the most important regulatory enzyme of the 237 pathway that determines the flux of sulfate into organic sulfur compounds in plants (31) (for 238 review see (32)). In addition, the availability of C-N skeletons for cysteine synthesis is 239 promoted in response to the JA mimic through enhanced serine O-acetyltransferase 240 241 (DmSERAT2) expression, and cysteine synthesis by itself, via elevated Oacetylserine(thiol)lyase (DmOASTL) expression (Fig. 4C, (33)). Moreover, the putative GSH 242 transporter DmOPT6 is transcriptionally induced after COR treatment as well. Interestingly, all 243 244 four transcripts are induced by both COR or prey capture in a similar fashion (Fig. 4B-D and http://tbro.carnivorom.com). Therefore, enhanced sulfate reduction and assimilation seems to 245 246 be required for both the synthesis of cysteine rich hydrolytic enzymes and additional synthesis of glutathione, which can be detected in the secreted fluid. 247

To test whether glutathione is released into the extracellular compartment, we sampled 248 digestive fluids from stimulated flytraps and analyzed the samples for the presence of anti-249 250 oxidants (34-36). Indeed, we could detect GSH in Dionaea's extracellular fluid (Fig. 4A). In contrast to glutathione however, ascorbate was not detectable by state-of-the-art methods (36, 251 37). While the GSH concentration in whole Dionaea traps was not significantly altered by COR 252 253 treatment, the stomach glutathione concentration was in the order of 10 µM 48 h after stimulation onset (Fig. 4A, red bars). In order to test the sensitivity of the used carbon fibers in 254 our amperometric analysis towards this ROS scavenger, we performed experiments with 255 defined GSH concentrations (Fig. S3E). In these experiments the reduced GSH was oxidized at 256

the positively charged carbon fiber resulting in a positive current. Interestingly, the amperometric current detected with a constant potential of +900 mV in solutions of defined GSH concentrations saturated with a half maximal concentration (K_m) of 10 μ M (Fig. S3E), which corresponds well with the actual GSH concentration in the secreted fluid. Thus it is likely that under our conditions secreted GSH is detected in the amperometric analysis. Nevertheless, we expect the amperometry to detect also additional electroactive substances besides GSH released in the secreted fluid of stimulated Venus flytraps.

264

265 **Discussion:**

The molecular machinery underlying secretory vesicle fusion with the plasma membrane in 266 267 animal cells is known in great detail (38-40). Upon chemical or electrical stimulation of secretory animal cells, exocytotic events can be detected within milliseconds (41-44). In these 268 fast responding cells, certain pools of preformed cargo-loaded vesicles are released immediately 269 after stimulus onset. Following hapto-electric calcium entry in Dionaea glands jasmonate-270 signalling triggers vesicle acidification and *de novo* synthesis of secretory proteins. The fact 271 that carbon fiber electrodes detect amperometric signals not earlier than about 6 h after 272 mechanical- and JA stimulation (Fig. 3D) may indicate that the oxidizable compound, most 273 likely the tripeptide glutathione, is contained only in those vesicles equipped with hydrolases. 274 In the acidic extracellular digestive fluid glutathione is very stable, providing for a proper redox 275 state for sustained hydrolase activity (45). 276

Dionaea's secretion events occur on a slow timescale. The apparent diffusion constant of released substances, as calculated from the waveform of the amperometric signal, was D =1.92x10⁻¹⁰ cm²/s, which indicates a high diffusional resistance of the gland cell wall. For comparison, the diffusion constant of catecholamines in aqueous solution is 1x10⁻⁶ to 8x10⁻⁷ cm²/s (19, 46). Thus diffusion in the cell wall of *Dionaea* glands is about four orders of magnitude slower than that of small molecules in aequeous solution. In contrast to fast synaptic signaling in the nervous system of animals, this slow diffusion as well as the slow time course of release reflects the biology of the insect-processing flytrap: Once *Dionaea* captures prey via its fast hapto-electric sensing system, exocytotic release and slow diffusion of a tailored hydrolase cocktail into the digestive fluid perfectly serves the long-term nutrient needs of the plant.

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289 Material and Methods:

290 Amperometric Recordings

In order to access the inner trap surface even in stimulated plants, unstimulated traps in the open 291 292 position were fixed in a chamber and mechanically locked to prevent trap closure upon stimulation. For inhibitor pre-treatments, plants were sprayed with 10 mM GdCl₃, or H₂O as 293 control. 24 h after pre-treatments traps were stimulated for secretion either mechanically (touch 294 of trigger hairs, 5 - 10 times within 1 min) or by hormone spraying (100 µM COR). At the given 295 time points after stimulation amperometric measurements were performed with open fixed traps 296 still attached to the plant. The chamber was filled with standard bath solution (1 mM KCl, 1 297 mM CaCl₂, 50 mM HEPES/NaOH pH 7) and placed on a microscope stage (Zeiss Axioscope 298 2 FS, Germany). A three-electrode configuration was employed where an Ag/AgCl electrode 299 served as the reference electrode grounding the bath solution. Two sensory carbon fiber 300 electrodes of 5 µm diameter (ALA Scientific Instruments, Westbury, NY) were used for 301 amperometric detection. Carbon fibers were gently placed on top of the gland head cells if not 302 stated otherwise. During amperometric recordings, electrodes were held at +900 mV with two 303 VA-10X amperometry amplifiers (ALA Scientific Instruments). Oxidative current was 304 acquired via VA-10X and digitized at 20 kHz through an ITC-18 digital to analog convertor 305 (InstruTECH, NY). Data were acquired using Patch master (HEKA Elektronik, Germany) and 306

analyzed with a custom-written fit running under Igor 6. Detected events were described byfollowing equations (47):

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$$f(x) = M / (t-t_0)^{1.5} * exp(-t_c / (t-t_0))$$
(1)

Here, t_0 is the time of signal onset (a free fitting parameter), and M depends on the amount of secreted substance as well as on the diffusion coefficient D. The parameter t_c depends on D and the distance, r, between the point source of secretion and the carbon fiber tip according to:

313 $t_c = r^2 / 4 D$ (2)

Further details on Materials and Methods can be found in 'Supplementary Materials andMethods'.

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317 Author Contributions:

E.N., and R.H. conceived the work; S.S. conducted initial feasibility studies; S.S., I.K., L.Sh.,

319 S.Sh., H.R., D.B., E.N., P.J., K.A.S. A.-R., and R.H. designed the experiments and analysed the

data; S.S., I.K., L.Sh., J.F., H.B., M.E., E.M., and B.H. performed the experiments; and S.S.,

321 I.K., D.B., E.N., M.B., and R.H. wrote the manuscript.

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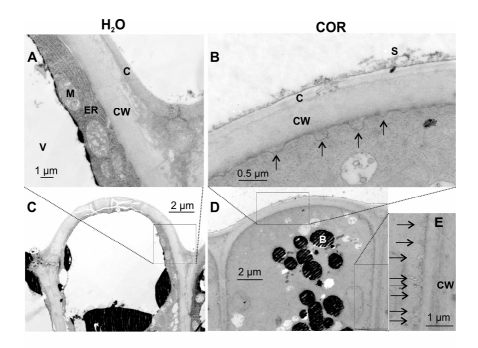
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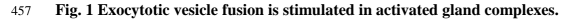
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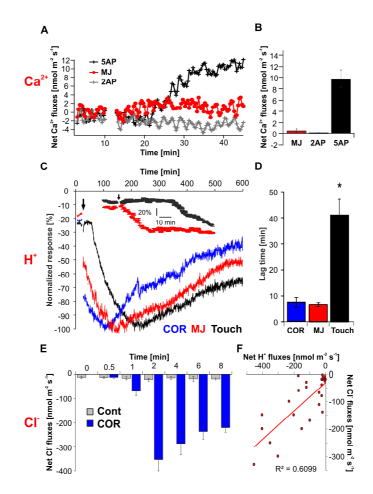
455 Figures:





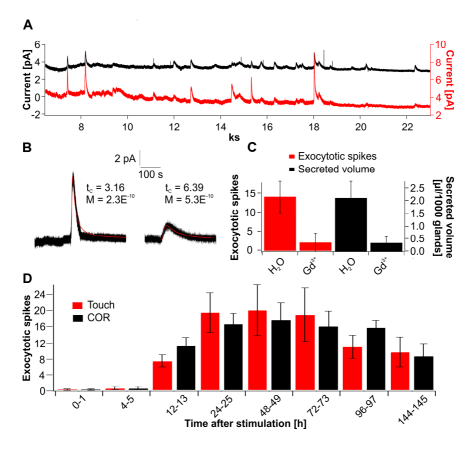


Electron micrographs of the outer layer of resting (**A** and **C**) and COR-stimulated (**B**, **D** and **E**) *Dionaea* gland complexes. **A**, **B** and **E** detailed view, **C** and **D** overview. Whereas resting glands do only exhibit few exocytotic events, a massive rise in exocytotic vesicle fusion with the plasma membrane (black arrows) could be detected 48 h after COR stimulation. B, dark stained body; C, cuticle; CW, cell wall; ER, endoplasmic reticulum; M, mitochondria; S, secreted fluid; V, vacuole.



466 Fig 2 Net ion fluxes measured from stimulated *Dionaea* glands via MIFE technique.

A) Net Ca^{2+} flux in response to mechanical (touched for either 2 or 5 times within 10 sec) and 467 chemical stimulation (1 mM methyl jasmonate, MJ). B) Peak Ca²⁺ flux response values for data 468 shown in panel A (mean \pm SE; n \geq 5). C) H⁺ flux kinetics in response to touch and Jasmonate 469 stimulation. Each flux was normalized to its maximum flux (100%) to illustrate the difference 470 in the peak time (mean \pm SE; n \geq 5). Insert: comparison of touch and MJ treatments at high 471 temporal resolution. **D**) Lag time in H^+ flux responses between treatments shown in panel C. 472 Jasmonate-induced proton release was significantly faster compared to mechanical induction 473 $(p \le 0.01; one way Anova)$. E) Net Cl⁻ fluxes measured in COR stimulated (blue bars) and non-474 stimulated (grey bars) glands at various time points after stimulation (mean \pm SE; $n \ge 4$). **F**) 475 476 Correlation of net H⁺ and Cl⁻ fluxes measured from COR stimulated glands at different time points illustrated in panel E. Each point represents a separate measurement. For all MIFE flux 477 data, the sign convention is 'influx positive'. 478





480 Fig 3 Amperometric detection of exocytotic events in *Dionaea glands*

A) Long-term spiking response of stimulated glands. Current spikes resulting from the 481 exocytosis of individual vesicles were detected with 2 electrodes simultaneously clamped to 482 +900 mV. B) Two examples of analyzed exocytotic current spikes are shown. Fitting these 483 events with equation (1) $f(x) = M / (t-t_0)^{1.5} \exp(-t_c / (t-t_0))$ (red line) reveals characteristics of 484 release quantified by M and t_c which reflect amount and distance of fusing vesicle to carbon 485 fiber. C) 10 mM Gadolinium was sprayed 24 h before mechanical stimulation of the Venus 486 flytraps. 24 h after stimulation the number of amperometrical detected events within 1 h (red) 487 and the secreted volume (black) was calculated. Compared to control traps, gadolinium inhibits 488 secretion as well as amperometrically detectable exocytotic spiking. Data are mean \pm SD (n \geq 489 25). D) Time course of exocytosis related spiking in response to touch (red) and COR (black). 490 For the given time points number of exocytotic events were calculated. Both stimuli lead to the 491 same long-term spiking response in flytrap glands. Data represent mean \pm SD (n \geq 54). 492

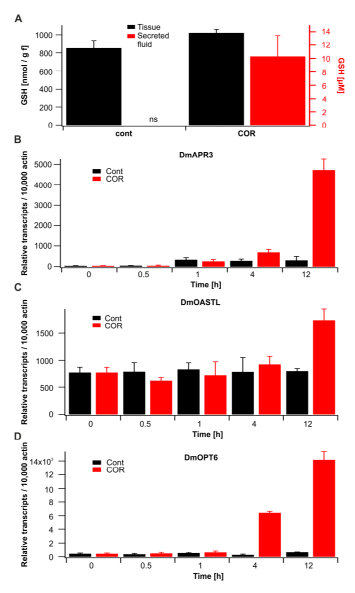


Fig 4 Synthesis of the ROS scavenger Glutathione is induced in stimulated Dionaea traps 494 A) GSH levels in traps (black) or secreted fluid (red bars) under non-stimulated conditions 495 (control) or 24 h after spray application of 100 µM COR. Please note that resting traps do not 496 secrete (ns) digestive fluid. Data represent mean \pm SD (n \geq 4). **B**)-**D**) Coronatine induces key 497 genes involved in GSH biosynthesis. Expression of DmAPR3, DmOASTL and DmOPT6 in 498 Dionaea gland complexes. Traps were sprayed with water (control, black) or 100 µM COR 499 (red) and gland complexes harvested at the time points indicated. Transcript numbers are given 500 relative to 10,000 molecules of DmACT1; (mean \pm SE, n = 6). 501