

37 **Abstract**

38 Ca^{2+} is an important intracellular messenger and regulator in both physiological and
39 pathophysiological mechanisms in the hearing organ. Investigation of cellular Ca^{2+}
40 homeostasis in the cochlea of hearing mammals is hampered by the special anatomy and high
41 vulnerability of the organ. A quick, straightforward and reliable Ca^{2+} imaging method with
42 high spatial and temporal resolution in the mature organ of Corti is missing. Cell cultures or
43 isolated cells do not preserve the special microenvironment and intercellular communication,
44 while cochlear explants are excised from only a restricted portion of the organ of Corti and
45 usually from neonatal pre-hearing murines. The hemicochlea, prepared from hearing mice
46 allows tonotopic experimental approach on the radial perspective in the basal, middle and
47 apical turns of the organ. We used the preparation recently for functional imaging in
48 supporting cells of the organ of Corti after bulk loading of the Ca^{2+} indicator. However, bulk
49 loading takes long time, is variable and non-selective, and causes the accumulation of the
50 indicator in the extracellular space. In this study we show the improved labeling of supporting
51 cells of the organ of Corti by targeted single-cell electroporation in mature mouse
52 hemicochlea. Single-cell electroporation proved to be a reliable way of reducing the duration
53 and variability of loading and allowed subcellular Ca^{2+} imaging by increasing the signal-to-
54 noise ratio, while cell viability was retained during the experiments. We demonstrated the
55 applicability of the method by measuring the effect of purinergic, TRPA1, TRPV1 and ACh
56 receptor stimulation on intracellular Ca^{2+} concentration at the cellular and subcellular level. In
57 agreement with previous results, ATP evoked reversible and repeatable Ca^{2+} transients in
58 Deiters', Hensen's and Claudius' cells. TRPA1 and TRPV1 stimulation by AITC and
59 capsaicin, respectively, failed to induce any Ca^{2+} response in the supporting cells, except in a
60 single Hensen's cell in which AITC evoked transients with smaller amplitude. AITC also
61 caused the displacement of the tissue. Carbachol, agonist of ACh receptors induced Ca^{2+}
62 transients in about a third of Deiters' and fifth of Hensen's cells. Here we have presented a
63 fast and cell-specific indicator loading method allowing subcellular level functional Ca^{2+}
64 imaging in supporting cells of the organ of Corti in the mature hemicochlea preparation, thus
65 providing a straightforward tool for deciphering the poorly understood regulation of Ca^{2+}
66 homeostasis in these cells.

67

68 **1. Introduction**

69 The mammalian organ of Corti has a uniquely spiraled structure covered with bony walls in
70 the adulthood. The special anatomy, high vulnerability and the calcification of the temporal
71 bone makes the organ hardly attainable and hampers its investigation significantly. Therefore,
72 most of the experimental studies in the organ of Corti are implemented in preparations made
73 from younger animals, e.g. the explant from 3-5 days old (P3-5) mice or rats (Lahne and Gale,
74 2008; Landegger et al., 2017). At this age the organ of Corti is immature yet and the rodents
75 are deaf, although the mechanotransducer channels are expressed and working in hair cells
76 from P0-P2 (Fettiplace and Kim, 2014; Lelli et al., 2009; Michalski et al., 2009). The mouse
77 and rat organ of Corti and hearing are considered to be mature both anatomically and
78 functionally at >P15 (Ehret, 1976; Rybak et al., 1992). Furthermore, hemicochlea is a
79 preparation available at mature stages providing the accessibility to the organ of Corti in three
80 different turns of the cochlea, and hence, the opportunity to investigate the cellular and
81 molecular mechanisms of tonotopy. The preparation, preserving the delicate cytoarchitecture
82 of the organ of Corti was originally developed for morphological, kinematic, and
83 mechanoelectric investigations (Edge et al., 1998; He et al., 2004; Hu et al., 1999; Keiler and
84 Richter, 2001; Richter et al., 1998). Our group was the first using it recently for real
85 functional Ca^{2+} imaging measurements in supporting cells of the organ of Corti bulk loaded
86 with acetoxymethyl ester conjugated (AM) Ca^{2+} indicator (Horváth et al., 2016). AM-dyes
87 load all types of cells and support the imaging of synchronized activity of cell groups and
88 their intercellular communication. Bulk loading of the tissue is simple, however, takes longer
89 time, is variable and non-selective, and causes the accumulation of the indicator in the
90 extracellular space. In this study, we aimed at developing a novel method which requires
91 shorter loading time, increases the selectivity and decreases the variability of labeling, and
92 results in lower extracellular dye spillover and light scattering from adjacent structures, thus
93 improves spatial resolution and reliability.

94 Ca^{2+} is a major intracellular second messenger (Berridge, 2016; Horváth et al., 2016;
95 Mammano et al., 2007) and Ca^{2+} indicators are the most reliable and pervading sensors of
96 intracellular messengers in functional imaging studies. Beside the cell permeable AM forms,
97 small-molecule Ca^{2+} sensors are available as membrane impermeable salts. By their targeted
98 loading into individual cells the background noise can be decreased significantly. Salt
99 indicators can be loaded into the cell by a patch pipette via diffusion in whole-cell
100 configuration (Beurg et al., 2009; Denk et al., 1995; Lagostena et al., 2001; Lagostena and
101 Mammano, 2001; Lorincz et al., 2016; Zelles et al., 2006) or single-cell electroporation.
102 Single-cell electroporation is faster, and prevents the wash-out of intracellular compounds
103 (Nevian and Helmchen, 2007), thus does not change the physiology of the cell and does not
104 modify the experimental results (Ishikawa et al., 2002; Vyleta and Jonas, 2014). Genetically
105 encoded Ca^{2+} indicators are wide-spread (Horikawa, 2015) and have the advantage of being
106 relatively selective for the cells expressing the target gene, however they are not available for
107 every cell type and their use is not always feasible.

108 Glia-like supporting cells of the organ of Corti are less investigated than the receptor hair
109 cells. Their structural, physical supporting roles are complemented with functional ones. They
110 are important in the development, macro- and micromechanics and in sensing the harmful
111 stimuli and initiating protective mechanisms in the inner ear, and also serve as a regenerative

112 pool for the lost hair cells (Monzack and Cunningham, 2013). Unfortunately, the majority of
113 information on supporting cells is from studies on neonatal and young pre-hearing animals.

114 In this study, we set up and validated a simple, rapid and reliable method of Ca^{2+} indicator
115 loading into individual supporting cells of the organ of Corti prepared from hearing mice. We
116 demonstrated that the single-cell electroporation in the hemicochlea is selective to the target
117 cell and causes little dye spill-over in the extracellular space. Using this technique we were
118 able to investigate the P2, TRPA1, TRPV1 and acetylcholine receptor (AChR) agonist-evoked
119 cellular and subcellular dynamics of intracellular Ca^{2+} concentration in Deiters', Hensen's and
120 Claudius' cells (DCs, HCs, CCs). These experiments also verified the technique. Furthermore,
121 the functional role of AChRs in HCs and the lack of functional role of TRPA1 and TRPV1
122 channels in Ca^{2+} signaling in the three supporting cell types have not been described before.

123

124 **2. Materials and Methods**

125

126 *2.1 Tissue Preparation*

127 All animal care and experimental procedures were in accordance with the National Institute of
128 Health Guide for the Care and Use of Laboratory Animals. Procedures were approved by the
129 Animal Use Committee of Semmelweis University, Budapest. Acutely dissected cochleae of
130 BALB/c mice from postnatal day 15 (P15) to P21 were used. Hemicochlea preparation was
131 carried out according to the Dallos' group method (Edge et al., 1998; Horváth et al., 2016).
132 Briefly, mice were anesthetized superficially by isoflurane then decapitated. The head was
133 divided in the medial plane and the cochleae were removed and placed in ice-cold modified
134 perilymph-like solution (composition in mM: NaCl 22.5; KCl 3.5; CaCl_2 1; MgCl_2 1;
135 HEPES:Na 10; Na-gluconate 120; glucose 5.55; pH 7.4; 320 mOsm/l), which was
136 continuously oxygenated. The integrity of the preparations was assessed by the gross
137 anatomy, location and shape of the supporting cells, hair cells, and the basal-, tectorial- and
138 Reissner's membranes. The perilymph-like solution with reduced $[\text{Cl}^-]$ minimizes swelling
139 and deformation of the cochlear tissue and preserves the morphological and functional
140 integrity of the preparation beyond 2hrs (Emadi, 2003; Teudt and Richter, 2007). We reduced
141 the Cl^- influx by iso-osmotic replacement of 120 mM NaCl for Na-gluconate, another
142 chemical efficiently used against cellular swelling in brain slice preparations (Rungta et al.,
143 2015). The medial surface of the cochlea was glued (Loctite 404, Hartford, CT) onto a plastic
144 plate with the diameter of 7 mm. Then the cochlea was placed into the cutting chamber of a
145 vibratome (Vibratome Series 3000, Technical Products International Inc., St. Louis, Mo,
146 USA) bathed in ice cold experimental solution and cut into two halves through the middle of
147 the modiolus with a microtome blade moving with a 30 mm/min speed and 1 mm amplitude
148 of vibration (Feather Microtome Blade R35, CellPath Ltd, Newtown, UK) under visual
149 control through a stereomicroscope (Olympus SZ2-ST, Olympus Corporation, Philippines).
150 Only the half that was glued to the plastic plate was used for imaging.

151

152 *2.2 Targeted single-cell electroporation dye-loading*

153 The method of Nevian and Helmchen in acute brain slices was adopted (Nevian and
154 Helmchen, 2007). The experiments were performed at room temperature (22-24 °C). The
155 hemicochleae were placed into an imaging chamber filled with the oxygenated perilymph-like
156 solution on the microscope stage. The perfusion speed was 3.5 ml/min in the chamber. The
157 cells were chosen in oblique illumination under a LUMPlanFl 40x/0.80w water immersion
158 objective (Olympus, Japan) with 3.3 mm working distance. Borosilicate pipettes (5–7 M Ω)

159 were filled with the Ca²⁺ indicators Oregon Green 488 BAPTA-1 hexapotassium salt (OGB-
160 1) or fura-2/K⁺ (ThermoFisher Scientific) dissolved in distilled water at a final concentration
161 of 1 mM. The pipettes were mounted onto an electrode holder attached to a micromanipulator
162 (Burleigh PCS-5000, Thorlabs, Munich, Germany). Each chosen cell was approached and
163 gently touched by the pipette under visual control; a single square wave current impulse of 10
164 ms duration and amplitude of 10 μ A were sufficient to load the cells with the Ca²⁺ indicator.
165 The pulses were generated by pCLAMP10 software-guided stimulator system (Biostim STE-
166 7c, Supertech Ltd, Pecs, Hungary; MultiClamp 700B Amplifier and Digidata 3200x,
167 Molecular Devices, Budapest, Hungary).

168 *2.3 Calcium imaging*

169 The OGB-1 dye-filled cells were illuminated by 494 ± 5 nm excitation light (Polychrome II
170 monochromator, TILL Photonics, Germany) and the emitted light was monitored after
171 passage through a band-pass filter (535 ± 25 nm). Fura-2/K⁺ loaded cells were alternately
172 illuminated by 340 ± 5 nm and 380 ± 5 nm excitation light and the emitted light was detected
173 behind a 510 ± 20 nm band-pass filter. Fluorescent images were obtained with an Olympus
174 BX50WI fluorescence microscope (Olympus, Japan) equipped with a Photometrics Quantix
175 cooled CCD camera (Photometrics, USA). The system was controlled with the Imaging
176 Workbench 6.0 software (INDEC BioSystems, USA). The image frame rate was 1 or 0.5 Hz
177 during the ATP-evoked responses and 0.1 or 0.05 Hz otherwise (OGB-1 or fura-2/K⁺,
178 respectively) to reduce phototoxicity and photobleaching. Fura-2/AM was simply used to
179 contrast the difference between single cell and bulk loading (Fig. 1A). Method of fura-2/AM
180 loading have been described previously (Horváth et al., 2016). Briefly, the hemicochlea was
181 incubated with 10 μ M fura-2/AM in the presence of pluronic F-127 (0.05 %, w/v) for 30 min,
182 then deesterified in standard experimental solution for 15 min before recording. The whole
183 experiment was performed within 1.5-2 h after decapitation. Cells with not preserved
184 morphology were excluded from further analysis.

185

186 *2.4 Drug Delivery*

187 ATP, allyl isothiocyanate (AITC), capsaicin and carbachol (Sigma-Aldrich, USA) were added
188 to the perfusion for 30 seconds. The perfusion reached the chamber in 27-30 sec and the
189 responses started in 60-80 sec. The buffer volume in the perfusion chamber was about 1.9 ml.
190 ATP, as a standard stimulus on supporting cells (Horváth et al., 2016), was always
191 administered at the beginning and at the end of experiments to confirm the cellular
192 responsiveness and the preparation viability. Before the first ATP application, an at least 3-
193 minute long baseline period was registered in each experiment. At least 10 minutes had to
194 elapse between two ATP stimulus, and if the solution was changed to Ca²⁺ free one
195 (composition in mM: NaCl 22.5; KCl 3.5; MgCl₂ 2; Hepes 10; Na-gluconate 120; glucose
196 5.55; EGTA 1; pH 7.4; 320 mOsm/l) the time lag before the 2nd ATP application was 15
197 minutes, similarly to our previous experiments (Horváth et al., 2016).

198

199 *2.5 Data Analysis*

200 Data analysis was performed off-line. Region of interest was drawn around the soma of the
201 stained cell and the phalangeal process in case of Deiters' cell imaging. Cell image intensities
202 were background-corrected using a nearby area devoid of loaded cells. Using OGB-1, the
203 relative fluorescent changes were calculated as follows:

$$204 \quad \frac{\Delta F}{F_0} = \frac{F_t - F_0}{F_0}$$

205 where F_0 is the fluorescent intensity of the baseline, and F_t is the fluorescent intensity at time
206 t .

207 In case of fura-2/ K^+ , the ratio of emitted fluorescence intensities (F_{340}/F_{380}) were calculated.
208 The response amplitudes were defined as the maximal change in intensity. Area under curves
209 and averages of the responses (Fig. 3) were calculated in Igor Pro 6.37.

210 Signal-to-noise ratio (S/N) in fura-2/AM and fura-2/ K^+ loaded cells were calculated from
211 ATP response curves of 12-12 randomly selected cells as follows:

$$212 \quad \frac{S}{N} = \frac{\Delta R}{\delta_R}$$

213 where ΔR is the amplitude of the ATP induced transients and δ_R is the standard deviation of
214 the baseline ratio prior to the ATP administration (at least 200 sec).

215 Data are presented as mean \pm standard error of the mean (SEM). The number of experiments
216 (n) indicates the number of cells. Testing of significance ($p < 0.05$) was performed based on the
217 distribution of the data. In case of normal distribution (tested by Shapiro-Wilk test) ANOVA,
218 in other cases Kruskal-Wallis test were used, both followed by Bonferroni post-hoc tests.

219 Levels of significance were as follows: * $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$.

220

221 **3. Results**

222 *3.1 Targeted single-cell electroporation is suitable to load Ca^{2+} indicators into cells in the* 223 *hemicochlea prepared from hearing mice*

224 The organ of Corti matures during the second postnatal week of life in mice (Ehret, 1976)
225 therefore we used P15-P21 hemicochlea preparations (Fig. 1) to investigate mature hearing
226 (Edge et al., 1998). The preparation allowed us to image all three turns of the cochlea (Fig.
227 1E) and the organs of Corti were well preserved in all turns (Fig. 1A and B show an apical
228 and a middle turn organ, respectively). The anatomical structures (e.g. membranes, stria
229 vascularis, spiral limbus) and cells were clearly visible, identifiable and exposed for
230 electroporation.

231 We optimized the electroporation described by Nevian and Helmchen (Nevian and Helmchen,
232 2007) for supporting cells in the hemicochlea preparation. Electroporation was fast and
233 efficient (10 minutes overall from the positioning of the preparation in the tissue chamber on
234 the microscope to the removal of the loading pipette, including filling up the pipette with the
235 dye) compared to bulk loading (Fig. 1A; 30 min loading plus 15 min deesterification;
236 (Horváth et al., 2016)) promoting the health of the tissue. We approached the cell, first using
237 the manipulator under mechanical then piezoelectric control. After approaching the cells with
238 the pipette filled with dye, we placed the tip gently on the cell membrane, and applied a 10 ms
239 long, 10 μA square pulse to deliver the charged molecules into the somas (Fig. 2A, D, E).

240 Forming a seal around the pipette tip by gently pushing the membrane is crucial to the
241 selective dye injection without any spillover into the extracellular space (Fig. 1B). A single 10
242 ms long pulse at lower current amplitudes (2-5 μA) resulted in insufficient loading of OGB-1.
243 Single pulses with larger currents (50-100 μA) loaded the cells with the sufficient amount of
244 dye, but a large proportion of the cells were damaged and lost their fluorescent intensities
245 quickly. A single 10 μA pulse could load the cells with sufficient amount of dye reliably. The
246 cells kept their morphology and did not loose their fluorescence till the end of the
247 experiments. Even in the case of a second loading pulse the cells survived and were
248 responsive to stimuli. Mistargeting the pipette caused instant cellular damage and dye leakage
249 (Supplementary Fig. 1A). The direction and speed of the loading pipette during removal was

250 critical. A faster removal could cause the rupture of the cell membrane with consequent dye
251 loss. Slow, fine movement preserved the cell integrity. Vertical pipette elevation gave
252 typically the best outcome, however a diagonal pipette removal was more advantageous for
253 deeper cells.

254 We have not observed any punctate dye accumulation in the cytoplasm which is a sign of dye
255 loading into the cytoplasmic organelles. However, in accord with the literature (Lagostena et
256 al., 2001; Lagostena and Mammano, 2001) we occasionally found higher fluorescence
257 intensity over the nucleus of the Hensen's and Claudius' cells (see Fig. 2, 3).

258 OGB-1 was tested in variable concentrations (100, 300, 500 μ M and 1 mM). In the lower
259 concentration range (100-500 μ M) multiple pulses were necessary to load the cells elevating
260 the chance of cell damage. To keep the membrane integrity we increased the dye
261 concentration to 1 mM at which concentration a single pulse was sufficient. The pulse and the
262 dye concentration parameters we applied for OGB-1 were appropriate for Fura2/ K^+ and OGB-
263 6F (OGB-6F data are not shown).

264 The diffusional equilibration of the dye took approximately 5 seconds. A rapid loss of the
265 intracellular fluorescence after loading indicated the damage of cell membrane
266 (Supplementary Fig. 1A). We discarded these hemicochleae. The success rate of the targeted
267 electroporation was ~60 % and most of the loaded cells survived. The single cell loading
268 procedure ensured the lower loading variability of supporting cells, the unambiguity of
269 fluorescent light sources (Fig. 1B), and the decrease in dye spill over into the extracellular
270 space (Fig. 1A and C) resulting in a significantly improved S/N and cell border contrast
271 compared to the bulk-loading method (compare Fig. 1C and D). These improvements together
272 enabled us to perform subcellular imaging in the phalangeal processes of Deiters' cells in
273 addition to their somas (Fig. 1B). The Deiters' and the Hensen's cells were easily loaded (Fig.
274 1B, 2, 3A), as they are large, even in the basal turn of the cochlea where they are shorter than
275 in the apical and middle turns (Keiler and Richter, 2001). Targeting of the laterally positioned
276 Claudius' cells was more difficult because of their smaller size (Fig. 3A). Loading of the
277 pillar cells was mostly unsuccessful, as their somas were too flexible to target them.
278 Interestingly, their apical or basal part did not load through the stalk (Supplementary Fig. 1B).
279 We could successfully load the inner and outer hair cells using the same parameters we
280 implemented for supporting cells (Fig. 2, D, E). The inner hair cell loading was more
281 challenging because of their close contacts with the inner border and inner phalangeal cells
282 occasionally resulting in the accidental electroporation of these supporting cells (Fig. 2E).

283 In order to validate the method and demonstrate its applicability in real functional imaging of
284 receptor-mediated Ca^{2+} signaling, we tested the effect of P2, TRPA1, TRPV1 and ACh
285 receptor stimulation. P2 purinergic Ca^{2+} signaling in supporting cells of the mature organ of
286 Corti is well substantiated (Dulon et al., 1993; Horváth et al., 2016; Housley et al., 2009,
287 1999; Lagostena et al., 2001; Lagostena and Mammano, 2001; Matsunobu and Schacht,
288 2000), while the functional role of TRP and ACh receptors in different supporting cells is
289 largely unexplored.

290
291 *3.2 ATP evoked reversible and repeatable Ca^{2+} transients in Deiters' cell soma and process,*
292 *Hensen's and Claudius cells*

293 Perfusion of ATP (100 μ M, 30 sec), acting on both P2X and P2Y receptors (Horváth et al.,
294 2016), evoked reversible and repeatable Ca^{2+} transients in all three supporting cell types
295 (Deiters', Hensen's and Claudius' cells) and the phalangeal processes of Deiters' cells (DCp)
296 loaded by electroporation (Fig. 3). High S/N attained by targeted single-cell electroporation
297 was indispensable to image subcellular compartments. ATP responses in cells loaded with
298 electroporation (fura-2/ K^+) had better S/N than ATP responses in bulk loaded cells (fura-
299 2/AM; Fig. 1F). DCp (25 apical, 4 middle, 3 basal turn responses) showed the largest ATP-
300 evoked Ca^{2+} transient expressed in relative amplitude (dF/F_0 ; Fig. 3B, C) and response
301 integral (area under the curve, AUC, $sec \cdot dF/F_0$; Fig. 3B, D). The amplitudes and AUCs of
302 ATP-evoked Ca^{2+} transients were not significantly different from each other in Deiters' (24
303 apical, 4 middle, 3 basal turn responses), Hensen's (10 apical, 12 middle, 2 basal turn
304 responses) and Claudius' cell (6 apical, 5 middle, 3 basal turn responses) somas (p-values of
305 the amplitudes: CC-DC: 1; CC-HC: 1; DC-HC: 0.4511; DCp-DC: 0.0018; DCp-HC: $1.47 \cdot 10^{-6}$;
306 $DCp-CC: 2.85 \cdot 10^{-4}$; p-values of the AUCs: CC-DC: 0.0919; CC-HC: 1; DC-HC: 0.1412;
307 $DCp-DC: 0.0163$; $DCp-HC: 2.46 \cdot 10^{-6}$; $DCp-CC: 1.01 \cdot 10^{-5}$; Bonferroni post-hoc test; Fig.
308 3B, C). The shape of Hensen's cells transients was two-peaked in several cases modifying the
309 average response trace. Ca^{2+} transients in Claudius' cells had the fastest decay (Fig. 3B).

310 Omission of Ca^{2+} from the perfusion buffer decreased the ATP-evoked Ca^{2+} transients in all
311 three supporting cell types, and the Deiters' cell process, although the inhibition was
312 statistically not significant in the Hensen's and Claudius' cells. Readministration of Ca^{2+}
313 resulted in the recovery of the ATP response (Fig. 4) indicating the viability of the cells in the
314 hemicochlea during the whole experiment. Cells not responding to the third ATP stimulus
315 were removed from the analysis.

316 Inner hair cells could also be stimulated by ATP (Fig. 2C).

317

318 *3.3 Stimulation of TRPA1 and TRPV1 channels did not induce Ca^{2+} signaling (except AITC in*
319 *a single Hensen's cell), but TRPA1 activation resulted in the slight movement of the tissue*

320 Anatomical studies (Ishibashi et al., 2008; Velez-Ortega, 2014; Zheng et al., 2003) indicated
321 the presence of TRPA1 and TRPV1 non-selective cation channel receptors on supporting cells
322 of the organ of Corti. In this study, the possible functional role of TRPA1 channels in Ca^{2+}
323 signaling in Deiters', Hensen's and Claudius' cells was tested by the perfusion (30 sec) of its
324 agonist, AITC (Sághy et al., 2015). Before and after AITC the cells were challenged with
325 ATP (100 μ M) to demonstrate the viability and responsiveness of the cells during the whole
326 experiment (Fig. 5B, C). Cells not responding to any of these stimulations were excluded
327 from the analysis.

328 AITC, tested in 200 μ M, 400 μ M and 2 mM concentrations did not evoke any Ca^{2+} transients,
329 but caused a faint fluctuation of the baseline in a dose-dependent manner (Fig. 5A). Because
330 the cells in the images moved out from and into the focal plane after AITC application, we
331 electroperated the supporting cells with the double excitation Ca^{2+} indicator, fura-2/ K^+ . The
332 ratio of fluorescence at 340 and 380 nm (F_{340}/F_{380}) is independent of the focal position and
333 geometrical factors (Grynkiewicz et al., 1985) thus it is free of the movement artifacts present
334 on the 340 and 380 nm excitation traces induced by the 400 μ M and 2 mM AITC perfusion
335 (Fig 5B and 5B inset). By using fura-2/ K^+ in the ratiometric mode, we found no Ca^{2+} response
336 for TRPA1 stimulation by AITC either in Deiters' or Claudius' cells. However, the agonist

337 evoked transients with smaller amplitude in one Hensen's cell (P15) out of 7 (~14 % response
338 rate; Fig. 5C). The transients of this cell showed a ~40 sec slower onset. Subcellular imaging
339 in Deiters' cells was also feasible with fura-2/K⁺ (Fig. 5C). The amplitude of the second ATP
340 stimuli were similar to the first ones except in Claudius' cells which showed a declined in the
341 second ATP response after AITC application (p= 0.008498).

342 Capsaicin (330 and 990 nM), the agonist of TRPV1 channels (Sághy et al., 2015) did not
343 induce any Ca²⁺ response in the supporting cells (Fig. 6). The experimental arrangement (Fig.
344 6A) was similar to the one testing TRPA1 function. ATP (100 μM) was used to confirm cell
345 viability. Capsaicin administration, unlike AITC, was not followed by any movement in the
346 preparation. The ATP responses recovered after capsaicin, either in Claudius' cells (p=
347 0.2413).

348

349 *3.4 Activation of ACh receptors by carbachol induced Ca²⁺ response in Deiters' and* 350 *Hensen's cells*

351 In order to further demonstrate the applicability of targeted electroporation in hemicochlea
352 preparation we applied carbachol, the agonist of ACh receptors. Deiters' and Hensen's cells
353 receive efferent innervation, including cholinergic input (Bruce et al., 2000; Burgess et al.,
354 1997; Fechner et al., 2001; Nadol and Burgess, 1994; Raphael and Altschuler, 2003) and
355 evidence supports the presence of the highly Ca²⁺ permeable functional α9 subunit-containing
356 nicotinic ACh receptors (nAChRs) in Deiters' cells isolated from adult guinea-pigs
357 (Matsunobu et al., 2001). Functional role of ACh receptors on Hensen's cells has not been
358 investigated so far.

359 Carbachol was perfused in 100 μM concentration (30 sec). Both compartments of the Deiters'
360 cells were activated by carbachol in 33 % of the experiments (Fig. 7A). The amplitudes of
361 these responses were similar to the ATP-induced ones (Fig. 7A, C), but their duration looked
362 shorter, reaching statistically significant difference in the process (ATP: 41.34±5.94 sec,
363 carbachol: 17.87±3.43 sec, p-value= 0.01667).

364 One Hensen's cell (in the middle turn of the cochlea) out of 5 was activated by carbachol at
365 100 μM (Fig. 7C). The response was small, but clearly visible both in its amplitude and AUC.
366 It had only one peak in contrast to a typical ATP induced response in Hensen's cells (Fig. 3).

367 Viability of the cells was confirmed by ATP application again. Cells not responding to ATP
368 were excluded from the study.

369

370 **4. Discussion**

371

372 *4.1 Advantages of the mature hemicochlea preparation and drawbacks of bulk loadings in* 373 *Ca²⁺ imaging*

374 Although the hemicochlea (Edge et al., 1998; Richter et al., 1998) lacks the normal
375 hydrodynamic properties and amplification of the cochlea, the preparation provides several
376 advantages for investigations in the hearing organ: it i) sustains the delicate cytoarchitecture
377 of the organ of Corti, ii) allows tonotopic experimental approach on the radial perspective of
378 the organ in the basal, middle and apical turns, and iii) provides all of these in a preparation
379 from hearing mice (>P15; (Ehret, 1976)). Cell cultures of certain cochlear cell types or

380 acutely isolated cells (Ashmore and Ohmori, 1990; Dulon et al., 1993) do not preserve the
381 special microenvironment and intercellular communication in the organ of Corti. Cochlear
382 explants lack some of these disadvantages, but in their case a restricted portion of the organ of
383 Corti is excised from its environment (Chan and Rouse, 2016; Moser and Beutner, 2000). The
384 explants are usually prepared from neonatal pre-hearing murines (Landegger et al., 2017;
385 Piazza et al., 2007), similarly to the cochlear slices (Lin et al., 2003; Morton-Jones et al.,
386 2008; Ruel et al., 2008). Dissected temporal bone preparation from the guinea-pig provides
387 access only to the apical coil (Fridberger et al., 1998; Mammano et al., 1999). Thus in many
388 characteristics the hemicochlea preparation is superior for physiological investigations in the
389 mature cochlea, identification of the pathomechanisms leading to sensorineural hearing losses
390 (SNHLs) in the adults or deciphering potential drug targets for SNHLs (Lendvai et al., 2011)
391 and testing candidate therapeutic compounds acting on these targets. The preparation was first
392 used by our group for real functional imaging of intracellular Ca^{2+} signaling, which is
393 implicated in the aforementioned phenomena (Horváth et al., 2016). In that study, the
394 indicator dye was bulk loaded in its AM form, as in the majority of Ca^{2+} imaging studies on
395 cells in the cochlea (Chan and Rouse, 2016; Dulon et al., 1993; Matsunobu and Schacht,
396 2000; Piazza et al., 2007). Bulk loading is convenient, but the dye remains in the extracellular
397 space resulting in significant background staining and low S/N. AM dyes can be taken up by
398 every cell, contaminating the responses of the cell of interest by fluorescence from adjacent
399 responding cells (Fridberger et al., 1998). Furthermore, loading and deesterification take
400 longer time compromising the survival of the preparation. Here, we show the novel method
401 and validation of targeted single-cell electroporation of identified supporting cells in the
402 hemicochlea preparation of the adult mouse cochlea. The improved technique is rapid,
403 reliable and has a significantly better S/N, which enables functional imaging of single cells in
404 the hemicochlea preparation with higher spatial resolution.

405
406 *4.2 Single-cell electroporation – rapid and specific Ca^{2+} indicator loading of supporting cells*
407 *with low S/N and retained viability*

408 Single-cell electroporation allows dye loading of selected cells. It has been successfully used
409 in brain slices to load neurons and measure Ca^{2+} signals even in fine structures as dendritic
410 spines (Nevian and Helmchen, 2007). Previously Lin and coworkers (Lin et al., 2003) have
411 reported the targeted electroporation of a spiral ganglion cell, an outer hair cell and an
412 epithelial cell in the Reissner's membrane, but their actual experiment was performed on
413 cochlear slices from P0-P7 rats and the technique has never been used in follow-up studies.
414 Our success rate of Ca^{2+} indicator loading by electroporation into identified supporting cells
415 in the hemicochlea was similarly high as in the brain slices and the successfully loaded cells
416 nearly all survived. The quick approach of the selected cell and the lack of pressure on the
417 pipette minimized the spillover of the indicator from the pipette. The negligible amount of
418 extracellular fluorescent dye and the specific cell loading enabled subcellular functional
419 imaging of the soma and the process of Deiters' cells, i.e. the stalk and the phalangeal process
420 of the Deiters' cells were not obscured by the fluorescence of outer hair cells. Ca^{2+} imaging in
421 Deiters' cells at the subcellular level has only been performed before in isolated cells (Dulon
422 et al., 1993) or with simultaneous whole-cell patch-clamp recording (Lagostena and
423 Mammano, 2001), which is a laborious technique and washes out the intracellular
424 biomolecules involved in signaling (Ishikawa et al., 2002; Vyleta and Jonas, 2014).
425 Electroporation is suitable for loading more cells in a preparation. We have also managed to

426 do that in the hemicochlea preparation (Fig. 2D). However, electroporation and bulk loading
427 are not mutually exclusive. The latter one is favorable if loading of high number of cells is
428 required, e.g. for investigating Ca^{2+} waves travelling through a larger population of supporting
429 cells in the cochlea. On the other hand, if spatial resolution and a radial view of the adult
430 organ of Corti is important for a given cochlear study, targeted single-cell electroporation in
431 the hemicochlea preparation is a simple, rapid and reliable choice.

432 The electroporation worked well for the Deiters', Hensen's and Claudius' cells. In contrast,
433 the pillar cells could not be loaded homogenously, because the dye did not diffuse through the
434 stalk part of the cell. We have not experienced any problem of dye diffusion through the stalk
435 of the Deiters' cells. Dye compartmentalization in this cell type only appeared when the glass
436 pipette was mistargeted, pushed deep inside the cell and reached the microtubule bundle
437 directly. This happened rarely with an experienced experimenter and became easily
438 recognizable by the visible bundles (Supplementary Fig 1.). Inner and outer hair cells could
439 also be loaded successfully.

440

441 *4.3 ATP evoked Ca^{2+} transients in the soma of Deiters', Hensen's and Claudius' cells and the* 442 *phalangeal process of the Deiters' cells - validation of (sub)cellular imaging*

443 Viability of the loaded cells and applicability of the method for functional imaging of
444 intracellular Ca^{2+} signaling were tested by measuring the ATP-evoked responses. ATP is a
445 ubiquitous transmitter in the hearing organ and its role in purinergic receptor-mediated Ca^{2+}
446 signaling is well substantiated (Housley et al., 2009; Lee and Marcus, 2008; Mammano et al.,
447 2007). Previously, we have also demonstrated its effect in Deiters', Hensen's and pillar cells
448 in the hemicochlea preparation after bulk loading with fura-2/AM (Horváth et al., 2016). ATP
449 induced reversible and repeatable Ca^{2+} transients in all three electroporation loaded
450 supporting cell types with higher S/N compared to bulk loading. In Deiters' cells, which have
451 two well defined compartments the selective loading and the low background fluorescence
452 allowed us to perform subcellular imaging, thus we could measure ATP- and carbachol-
453 evoked Ca^{2+} transients in the soma and the plate of the phalangeal process. The ATP
454 responses had somewhat different characteristics in different supporting cells. The Hensen'
455 cells frequently had two-peak Ca^{2+} responses while the Claudius' cells showed the fastest
456 recovery after stimulation.

457 The processes of Deiters' cells had the largest Ca^{2+} transients, expressed in $\Delta F/F_0$, probably
458 because of the largest density of ATP receptors on their surface. However, the lower baseline
459 fluorescence (F_0), or the tiny volume of the process with larger surface-to-volume ratios may
460 further contribute to the difference by promoting the Ca^{2+} accumulation compared to the
461 somas with smaller surface-to-volume ratios (Helmchen et al., 1997). Quantification of basal
462 Ca^{2+} concentration and its changes in absolute concentration values requires dual wavelength
463 indicators or dual indicators loading and calibration (Yasuda et al., 2004). Nevertheless, our
464 hemicochlea electroporation method provides a reliable tool to investigate the supporting cell
465 Ca^{2+} signaling at the single cell and subcellular level in more details.

466 Functional expression of both ionotropic P2X and metabotropic P2Y receptors of ATP have
467 been shown on supporting cells in the organ of Corti in neonatal rodents and hearing mice
468 (P15-21) (Horváth et al., 2016; Housley et al., 2009; Lee and Marcus, 2008). Partial inhibition
469 of the ATP transients by omission of Ca^{2+} from the perfusion buffer, a blunt way of separating
470 the extracellular Ca^{2+} -dependent P2X- and intracellular store-dependent P2Y receptor

471 responses reproduced the results in the literature and further validated the method.
472 Furthermore, this arrangement of the experiment, when Ca^{2+} transients are evoked in the
473 absence then in the presence of Ca^{2+} in the same cell, demonstrated the way how
474 pharmacological interventions can be tested by internal control and provide a lower variability
475 of the effects. The development of the 3rd stimulus in the absence of the pharmacological
476 inhibitor or modulator can confirm the viability of the cell and the effect of the tested drug.

477
478 *4.4 TRPA1 stimulation did not induce Ca^{2+} response in Deiters' and Claudius' cells but*
479 *raised the possibility of TRPA1 role in Hensen's cell Ca^{2+} homeostasis*

480 TRP channels have mostly been studied by anatomical methods and their presence has been
481 shown in the inner ear. We tested the effect of the TRPA1 agonist AITC and the TRPV1
482 agonist capsaicin on Ca^{2+} regulation in the supporting cells of the mouse organ of Corti.
483 TRPA1 channels have also been shown in the supporting cells, mostly in Hensen's cells
484 (David P. Corey et al., 2004; Stepanyan et al., 2011; Velez-Ortega, 2014), but also in Deiters',
485 Claudius' and pillar cells (Velez-Ortega, 2014). In newborn rodent cochlear explant the
486 TRPA1 antibodies seems to be nonspecific or appears in the endoplasmic reticulum in
487 Hensen's and Claudius' cells (David P Corey et al., 2004). However, indirect immunolabeling
488 (against TRPA1 promoter connected reporter gene) confirmed TRPA1 presence in the
489 neonatal cochlear explants (Velez-Ortega, 2014). Contrarily, Takumida et al. (Takumida et
490 al., 2009) reported immunoreactivity to TRPA1 channels exclusively in nerve fibers of the
491 spiral ganglion cells and in nerves innervating the outer or inner hair cells in the mouse inner
492 ear. We could not detect Ca^{2+} response at any AITC concentrations in the investigated
493 supporting cells, except reduced-amplitude and late-onset transients in a single Hensen's cell.

494
495 *4.5 TRPA1 stimulation displaced the organ of Corti*

496 On the other hand, we detected a dose-dependent movement 'artefact' in the images after
497 AITC application. This probably represents a displacement of the whole organ of Corti and
498 could be caused by AITC-evoked contraction of cells in the cochlear epithelium. Outer hair
499 cells may be involved in this contraction (David P. Corey et al., 2004). However, Velez-Ortega
500 (Velez-Ortega, 2014) suggested the contraction of pillar and Deiters' cells as the origin of
501 TRPA1 stimulation-evoked tissue movement in P0-P7 wild type mice. The contraction was
502 not induced in *Trpa1*^{-/-} mice. Our study is in contrast to the idea of TRPA1-evoked
503 contraction of mature Deiters' cells or, alternatively, it is not exerted by intracellular Ca^{2+}
504 increase. Use of TRPA1 KO mice could contribute to decipher the role of TRPA1 channels.

505
506 *4.6 TRPV1 stimulation did not evoke any Ca^{2+} response in the supporting cells*

507 The presence of TRPV1 channels has also been shown in the cochlear epithelium. Their
508 expression was dependent on rodent species and age. In mouse cochlea the TRPV1 RNA level
509 first increased then declined in the E18-P8 period, similarly to TRPA1 (Asai et al., 2009). On
510 the contrary, Scheffer et al. (Scheffer et al., 2015) did not detect RNA for TRPV1 in hair cells
511 and surrounding cells in E16-P7 mice. Immunohistochemistry was used in adult guinea-pigs
512 and rats to show the presence of TRPV1 in some supporting cells, particularly in Hensen's
513 and outer and inner pillar cells (Takumida et al., 2005; Zheng et al., 2003). The lack of
514 capsaicin response in our experiments may indicate the absence of TRPV1 channels in

515 Deiters', Hensen's or Claudius' cells in the P15-21 mouse cochlea. Indeed they have not been
516 directly demonstrated on these cell types yet. Alternatively, they are functionally not involved
517 in intracellular Ca^{2+} regulation in these cells. We did not observe any movement in response
518 to capsaicin in the preparation either, suggesting that TRPV1 is not involved in contraction of
519 cells in the organ of Corti in hearing mice.

520 The decrease in the amplitudes of ATP transients after AITC applications in Claudius' cells
521 may be the consequence of a functional cross-inhibition between co-expressed TRPA1 and
522 the purinergic P2X receptors in that cells (Stanchev et al., 2009). Note that in the absence of
523 these insults the ATP response recovered (Fig. 3).

524

525 *4.7 ACh receptor activation evoked Ca^{2+} transients in some Deiters' and Hensen's cells*

526 Cholinergic efferent innervation of the motile outer hair cells has a well-known role in setting
527 cochlear amplification (Dallos et al., 1997; Kujawa et al., 1994). Deiters' and Hensen's cells
528 also receive efferent innervation (Bruce et al., 2000; Burgess et al., 1997; Fechner et al., 2001;
529 Nadol and Burgess, 1994; Raphael and Altschuler, 2003). Matsunobu and his coworkers have
530 shown acetylcholine-evoked Ca^{2+} increase in isolated Deiters' cells from guinea-pigs and
531 suggested the involvement of $\alpha 9$ -subunit containing nAChRs (Matsunobu et al., 2001). The
532 presence of $\alpha 10$ -subunit of nAChRs was not ruled out either in adult rat Deiters' cells
533 (Elgoyhen et al., 2001). Both homomeric $\alpha 9$ and heteromeric $\alpha 9\alpha 10$ nAChRs are highly
534 permeable for Ca^{2+} what can be detected by Ca^{2+} imaging methods (Fucile et al., 2006;
535 Matsunobu et al., 2001). There are no similar receptor expression or functional data on
536 Hensen's cells in the literature, thus we investigated the effect of carbachol, a partial agonist
537 on both native and $\alpha 9$ -subunit containing nAChRs (Verbitsky et al., 2000), also on Hensen's
538 cells. The proportion of Deiters' cells (33 %) responding for carbachol was very similar to the
539 one Matsonubu et al. (Matsunobu et al., 2001) reported in isolated guinea-pig Deiters' cells
540 for acetylcholine (42-44 %). The response rate of Hensens' cells was only 20 % and the
541 amplitude of the Ca^{2+} transient was smaller than that of the ATP-evoked one, differing from
542 Deiters' cells in which carbachol and ATP transients were comparable in amplitude. In
543 addition to confirming the cholinergic responsiveness of Deiters' cells in an *in situ*
544 preparation, we also raised the possibility of cholinergic regulation in Hensens' cells, the
545 other innervated supporting cell type in the organ of Corti.

546

547 **5. Conclusions**

548 Here we presented the method of Ca^{2+} indicator loading of supporting cells in the organ of
549 Corti in the mature mouse hemicochlea preparation using targeted single-cell electroporation.
550 Ca^{2+} is an important intracellular messenger and regulator and the method is a reliable and
551 straightforward tool for elucidating its role in these cells. Indicator loading is always a crucial
552 step in functional imaging. Our method provides the advantages of being i.) performed in the
553 adult hearing cochlea, ii.) rapid, thus extends the experimental time window, iii.) selective,
554 therefore lowers S/N and allows subcellular imaging, iv.) free from washing out the
555 intracellular biomolecules involved in signaling and metabolism and v.) suitable for tonotopic
556 investigations on the radial perspective in the basal, middle and apical turns of the cochlea.

557 Confirming the effect of ATP in Deiters', Hensen's and Claudius' cells and supporting the
558 functional role of AChRs in Deiters' and Hensen's cells in an *in situ* preparation also served
559 as a validation of the method. Showing the lack of involvement of TRPA1 and TRPV1
560 channels in Ca²⁺ regulation in Deiters' and Claudius' cells and in Deiters', Hensen's and
561 Claudius' cells, respectively, and raising the possibility of the functional role of ACh and
562 TRPA1 channels in Hensen's cell Ca²⁺ homeostasis demonstrated the applicability of the
563 method in the exploration of new Ca²⁺ signaling pathways in supporting cells of the mature
564 cochlea.

565

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572

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785 **Figure 1.** Targeted single-cell electroporation provides rapid and reliable loading of
786 fluorescent Ca^{2+} indicators in hemicochlea preparation of hearing mice. Hemicochlea
787 preparation (E) is suitable to investigate the mature organ of Corti in different regions of the
788 frequency map (A, B: hearing organ from the apical and the middle turn, respectively). The
789 cells of the organ are recognizable by their location and morphology. In case of bulk loading
790 of an AM dye (fura-2/AM) different cells take up different amount of dye and the
791 extracellular matrix also bind some (A), whilst single-cell electroporation of the dye (OGB-1)
792 specifically and reliably loaded up the targeted cell (B). This resulted in a significant
793 improvement in the S/N and the cell border contrast making visible the stalk and the
794 phalangeal process of the Deiters' cell, which is covered by the fluorescence of OHCs in case
795 of bulk loading. Surface plot of the fluorescent images of the electroporated Deiters' cells
796 demonstrates the low intensity of the background (D) versus the cell, while the bulk loading
797 surface plot (Fiji analysis platform; (Schindelin et al., 2012)) shows the intensity of multiple
798 cells and the noisier background (C). S/N of ATP responses (response amplitude / standard
799 deviation of the baseline) in 12-12 randomly selected cells loaded by fura-2/AM (bulk
800 loading, AM) or fura-2/ K^+ (electroporation, EP; F). (OHCs, outer hair cells; DCs, Deiters'
801 cells; DCp, phalangeal process of the Deiters' cell; OPC, outer pillar cell; IPC, inner pillar
802 cell; IHC, inner hair cell; HC, Hensen's cell; BM, basal membrane; TM, tectorial membrane)

803

804 **Figure 2.** Procedure of single-cell electroporation. The cells were selected at oblique
805 illumination and approached by glass pipette filled with fluorescent dye (OGB-1, 1 mM). A
806 single square wave current impulse (10 ms, 10 μA) was enough to load the cells within
807 seconds. (A) Course of dye loading by electroporation. Hensen's cell from the middle turn of
808 the mouse cochlea. (B) Schematic drawing of pipette and preparation arrangement. The same
809 pipette orientation was used in all three cochlear turns. (C) ATP evoked Ca^{2+} transient in an
810 inner hair cell. (D) Loading of inner and outer hair cells in the basal turn of a preparation. Left
811 image, moment of electroporation; middle image, oblique illuminated image and location of
812 the pipette; right image, loaded cells 10 min after electroporation of the outer hair cell. (E)
813 Loading of an inner boarder and an outer hair cell in the apical turn (same sequence of
814 images). (OHCs, outer hair cells; DCs, Deiters' cells; PCs, pillar cells; IBC, Inner border cell;
815 IHC, inner hair cell; HC, Hensen's cell; BM, basal membrane; TM, tectorial membrane)

816

817 **Figure 3.** ATP evoked repeatable Ca^{2+} transients in different supporting cells (Deiters',
818 Hensen's and Claudius') loaded with OGB-1 by electroporation. Perfusion of ATP (100 μM ,
819 30 sec), agonist of purinergic receptors, caused the elevation of fluorescent intensity in all
820 types of supporting cells electroporated by the Ca^{2+} sensitive dye OGB-1. The good S/N
821 enabled subcellular imaging in the Deiters' cells. The ATP responses were reversible and
822 repeatable (A) in all three cell types. Averages of 10 random responses (dF/F_0 , average \pm
823 SEM; B) showed different shapes of responses. The phalangeal process subcompartment
824 showed the largest Ca^{2+} response in dF/F_0 , the transient amplitudes in the somas of the three
825 supporting cell types did not differ significantly (B, C). Responses of Hensen's cells were
826 often 2 peak-shaped (A, B). Claudius' cells had a rapidly increasing and decaying response
827 (A, B). Area under the response curve values showed similar relations to each other as the
828 amplitudes (D). Example experiments of Deiters' and Claudius' cells were from the apical

829 and of Hensen's cells from the middle cochlear turn. Number of cells responded to ATP are
830 given in the respective bars. * p<0.05; ** p<0.01; *** p<0.001.

831

832 **Figure 4.** The model system is suitable for testing the effect of pharmacological
833 manipulations on evoked Ca^{2+} transients - omission of Ca^{2+} inhibited the ATP response. The
834 viability of cells loaded by single-cell electroporation in the hemicochlea preparation allows
835 functional imaging of triple stimulation, thus making the model feasible for comparing the
836 effect of pharmacological manipulations (2nd stimulus) to an internal control (1st stimulus).
837 The recovery of the response for the 3rd stimulation again in the absence of the
838 pharmacological intervention support the validity of the result and reassure preparation
839 viability. Cells without a response for the 3rd stimulus were excluded from the analysis. (A)
840 subcellular imaging of the effect of Ca^{2+} withdrawal on ATP stimulation in the process and
841 soma of an apical Deiters' cell. (B) The absence of extracellular Ca^{2+} decreased the ATP
842 response in all three cell types loaded with OGB-1, including the process of the Deiters' cells.
843 Number of experiments are given in the respective bars. * p<0.05; ** p<0.01.

844

845 **Figure 5.** TRPA1 agonist AITC did not induce $[\text{Ca}^{2+}]_i$ changes in the supporting cells of the
846 organ of Corti (except in a single Hensen's cell), but evoked a faint movement of the tissue.
847 (A) Cells loaded with OGB-1 and stimulated by different concentrations of AITC (200 μM ,
848 400 μM and 2 mM; 30 sec perfusion) moved out from the focal plane of imaging, but did not
849 respond with measurable $[\text{Ca}^{2+}]_i$ changes. See dose-dependent fluctuations of fluorescence on
850 individual and average curves of Deiters' cell soma. (B) Ratiometric imaging after successful
851 loading of fura-2/ K^+ by targeted electroporation. Ratiometric measurement eliminated the
852 moving artefact visible on the individual 340 nm and 380 nm excitation curves (*inset*) of a
853 representative experiment. ATP (100 μM) responses at the beginning and at the end of the
854 experiments prove that the lack of AITC (400 μM and 2 mM) effect is not because of lost
855 cellular viability. Cells not responding for ATP were excluded from further analysis. (C)
856 AITC (400 μM and 2 mM) did not induce Ca^{2+} transients in the investigated supporting cell
857 types, except the reduced amplitude transients in a single Hensen's cell (P15) out of the seven.
858 Bars represent change in F_{340}/F_{380} ratio relative to the baseline ($\Delta F_{340}/F_{380}$). Number of
859 responding/imaged cells are in parentheses. The amplitude of the Ca^{2+} increase is calculated
860 from the responding cells only. ** p<0.01.

861

862

863 **Figure 6.** TRPV1 agonist capsaicin did not induce $[\text{Ca}^{2+}]_i$ changes in any of the supporting
864 cell types investigated. (A) Representative trace shows the experimental arrangement.
865 Capsaicin was perfused (30 sec) in the concentration of 330 and 990 nM. ATP (100 μM),
866 administered at the beginning and the end of the experiments to prove the viability and
867 responsiveness of the cells. Not responding cells were excluded. (B) Bar graphs show the
868 averages of evoked transients. Capsaicin induced Ca^{2+} response in neither of the cell types
869 (Deiters', Hensen's and Claudius') loaded with OGB-1. Number of responding/imaged cells
870 are in parentheses.

871

872 **Figure 7.** Carbachol, a cholinergic receptor agonist, induced $[Ca^{2+}]_i$ transients in Deiters' and
873 Hensen's cells. (A) Deiters' cells loaded with OGB-1 were activated by the perfusion of
874 carbachol (100 μ M, 30 sec perfusion) in 33 % of the experiments. (B) A representative
875 carbachol-evoked response in a middle-turn Deiters' cell. The amplitude of the transient was
876 comparable to the ATP induced responses, its duration was shorter. (C) We detected intensity
877 elevation in one out of five Hensen's cells (20 %) after the carbachol stimulus. The amplitude
878 of the Ca^{2+} increase is calculated from the responding cells only. Number of
879 responding/imaged cells are in parentheses.

880

881 **Supplementary Fig. 1.** Signs of mistargeted electroporation and the block of dye diffusion in
882 pillar cells. (A) If the electroporation pipette, filled with OGB-1 was pushed too hard and sank
883 deep inside the cell membrane of the Deiters' cells the indicator dye tended to
884 compartmentalize into the nucleus and the microtubule bundles running along the cell while
885 the intracellular fluorescence was leaking out. (B) OGB-1 was unable to diffuse to the other
886 end of the pillar cells from the electroporation site. Both inner and outer pillar cells have long
887 microtubule bundle in the soma ended in actin mesh in the head and foot parts of the cell. The
888 indicator seems to have high affinity to these intracellular elements. (nuc., nucleus; m.b.,
889 microtubule bundle; IPC and OPC, inner and outer pillar cell; OHC and IHC, outer and inner
890 hair cell; DC, Deiters' cell; BM, basal membrane)