Organ-on-chip model shows that ATP release through connexin hemichannels drives spontaneous Ca\textsuperscript{2+} signaling in non-sensory cells of the greater epithelial ridge in the developing cochlea†‡

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Prior work supports the hypothesis that ATP release through connexin hemichannels drives spontaneous Ca\textsuperscript{2+} signaling in non-sensory cells of the greater epithelial ridge (GER) in the developing cochlea; however, direct proof is lacking. To address this issue, we plated cochlear organotypic cultures (COCs) and whole cell-based biosensors with nM ATP sensitivity (ATP-WCBs) at the bottom and top of an ad hoc designed transparent microfluidic chamber, respectively. By performing dual multiphoton Ca\textsuperscript{2+} imaging, we monitored the propagation of intercellular Ca\textsuperscript{2+} waves in the GER of COCs and ATP-dependent Ca\textsuperscript{2+} responses in overlying ATP-WCBs. Ca\textsuperscript{2+} signals in both COCs and ATP-WCBs were inhibited by supplementing the extracellular medium with ATP diphosphohydrolase (apyrase). Spontaneous Ca\textsuperscript{2+} signals were strongly depressed in the presence of G\textsubscript{b6}−/− COCs, in which connexin 30 (Cx30) is absent and connexin 26 (Cx26) is strongly downregulated. In contrast, spontaneous Ca\textsuperscript{2+} signals were not affected by replacement of Panx1−/− with Panx1+/− COCs in the microfluidic chamber. Similar results were obtained by estimating ATP release from COCs using a classical luciferin–luciferase bioluminescence assay. Therefore, connexin hemichannels and not pannexin 1 channels mediate the release of ATP that is responsible for Ca\textsuperscript{2+} wave propagation in the developing mouse cochlea. The technological advances presented here have the potential to shed light on a plethora of unrelated open issues that involve paracrine signaling in physiology and pathology and cannot be addressed with standard methods.

Introduction

Extracellular ATP modulates the function of both sensory and non-sensory cells in the inner ear (reviewed in ref. 1–3). In the developing cochlea, ATP triggers cytosolic Ca\textsuperscript{2+} concentration oscillations and propagation of intercellular Ca\textsuperscript{2+} waves that carry crucial biochemical information to the cochlear sensory epithelium. 4–6 The latter is subdivided into two embryologically distinct regions, the lesser epithelial ridge (LER) and the greater epithelial ridge (GER), which are separated by a pair of immature inner and outer pillar cells. 7–9 The GER encompasses the nascent inner sulcus, the tall columnar cells of the Kölliker organ,10,11 the inner hair cells (IHCs) and their abutting supporting cells,12,13 Ca\textsuperscript{2+} waves arise spontaneously in the GER14–19 and appear to play a crucial role in normal development of the cochlear sensory epithelium, hearing acquisition, functional maturation of hair
cells, redox homeostasis and age-related hearing loss. A wealth of experimental data and modeling studies support the notion that Ca²⁺ wave propagation in the developing cochlea is sustained by the release of ATP from the cytosol of non-sensory cells to the endolymph, in which baseline ATP concentration is in the low nM range. Genetic interference experiments support a critical role for Cx26 and Cx30, the two major cochlear connexins, in intercellular Ca²⁺ wave propagation in the developing cochlea (reviewed in ref. 2 and 37). However, direct proof that ATP release through connexin hemichannels drives spontaneous Ca²⁺ wave propagation in the GER is lacking.

To address this critical issue, we used two global knock out (KO) mouse strains, namely connexin 30 KO (Gjb6−/−; MGI: 2447863) and pannexin 1 KO (Panx1−/−; MGI: 5310802). Gjb6−/− mice fail to acquire hearing and do not develop endocochlear potential. Furthermore, they show defects of the endothelial barrier in capillaries of the stria vascularis (30) and impaired gap junction-mediated transfer of the fluorescent D-glucose derivative 2-NBDG. Accordingly, not only is Cx30 not expressed, but also Cx26 levels are severely reduced in the developing cochlea of this KO strain. In contrast, Panx1−/− mice have an increased susceptibility for atrial fibrillation and show a QT-prolongation cardiac phenotype. However, their hearing sensitivity, outer hair cell-based “cochlear amplifier” and cochlear nerve function, analyzed by auditory brainstem response (ABR) and distortion product otoacoustic emission (DPOAE) recordings, are normal, together with connexin expression and gap-junction coupling in the developing organ of Corti.

As detailed below, we developed an innovative organ-on-chip model which, by the coupling of microfluidic technology, whole-cell based biosensors (WCBs) with nM sensitivity to extracellular ATP and multiphoton imaging, allowed systematic addressing of purinergic signaling in COCs obtained from the above mentioned KO mice. Our results provide direct evidence that ATP release through connexin hemichannels drives spontaneous Ca²⁺ wave propagation in the GER.

Methods

Animals

All animals (Mus musculus) used in this study (see Table 1) were bred at the National Research Council-Institute of Biochemistry and Cell Biology (CNR-IBBC), Infranfrontier/ESFRI-European Mouse Mutant Archive (EMMA), Specific Pathogen-Free (SPF) barrier unit (Monterotondo Scalo, Rome, Italy). Mice were housed in individually ventilated caging systems (Tecniplast, Gazzada, Italy) at a temperature (T) of 21 ± 2 °C, relative humidity (RH) of 55 ± 15% with 50–70 air changes per hour (ACH) and under controlled (12:12 hour) light–dark cycles (7 am–7 pm). Mice had ad libitum access to water and a standard rodent diet (Emma 23, Mucedola, Settimo Milanese, Italy). Both male and female homozygous Gjb6−/− [EMMA ID (EM): 00323] or Panx1−/− (EM: 11476) pups at post-natal day 5 (P5) and their wild type P5 siblings (Gjb6+/−, Panx1+/−) were used. The background strains of these mice were C57BL/6J for Gjb6−/− and C57BL/6N for Panx1−/− mice. Note that there is no discernible difference in the auditory phenotype of the closely related C57BL/6J and C57BL/6N mouse strains.

Experimental animals were culled by trained personnel using gaseous anaesthesia followed by a rising concentration of CO₂. All the experimental procedures were agreed upon, reviewed and approved by local animal welfare oversight bodies and were performed with the approval and direct supervision of the CNR-IBBC/Infranfrontier—Animal Welfare and Ethical Review Body (AWERB), in accordance with several guidelines regarding animal experimentation, approved by the Italian Ministry of Health, in compliance with the Legislative Decree 26/2014 (ref. Project licence no. 68/2016-PR), transposing the 2010/63/EU Directive on protection of animals used in research. This work was also conducted based on recommendations from both ARRIVE and PREPARE guidelines.

Genotyping

Panx1 mice were genotyped according to published protocols by standard PCR on extracted mouse tail tips using the following primers:

Panx1 f: 5′-GGAAGTCAAGAGGTACCC-3′
Panx1 r: 5′-CTTGCCACGGAGTATGTGTTT-3′
LacZ: 5′-GTCCCCTCACCACCTTTTCCAG-3′

Mice with normal Panx1 alleles (Panx1+/+) were targeted by the above f and r primers and identified by a 330 bp band, whereas Panx1−/− were targeted by primers Panx1 f and LacZ, and were identified by a 630 bp band. Panx1−/− mice were identified by the simultaneous presence of a 330 bp and a 630 bp band. Primer pairs for Gjb6−/− mice were specific for the wild type alleles:

Gjb6 f: 5′-GGTACCTTTCTACTAATTAGCTTGG-3′
Gjb6 r: 5′-AGGTTGACATCCCATTGGAGGA-3′

To visualize the deletion, primers specific for the lacZ region (that flanks the deleted allele) were used in combination with the corresponding wild type forward primer:

Gjb6lac f: 5′-AGCGAGTACAAACCCGTGCAGATC-3′

Mice with normal Gjb6 alleles (Gjb6+/+) were identified by a 544 bp band, whereas Gjb6−/− mice were identified by a 460 bp band. See also Table 1.

Study design

In order to construct the optimal experimental design and estimate the minimum number of animals necessary for the experiments (sample size of the groups), for each type of

| Genotype | Gjb6+/+ | Gjb6−/− | Panx1+/+ | Panx−/− | C57BL6/N |
|----------|---------|---------|----------|---------|----------|
| Male     | 5       | 5       | 7        | 7       | 2        |
| Female   | 5       | 5       | 6        | 6       | 1        |
| Total    | 10      | 10      | 13       | 13      | 3        |
experiment and for each genetically modified and control strain (Gjb6<sup>−/−</sup>, Gjb6<sup>+/−</sup>, Panx1<sup>−/−</sup>, Panx1<sup>+/−</sup>), we set probability \( \alpha = 0.05 \) for the type I error in the \( t \) test. Then, fixing \( \beta = 4 \alpha = 0.20 \) to obtain a test power of \( 1 - \beta = 80\% = 0.8 \), we computed the number \( n \) of each of the two samples to be compared using the formula:

\[
n > \frac{(z_{\alpha/2} + z_{\beta})^2}{\Delta^2}
\]

with \( z_{\alpha/2} = 1.96 \) and \( z_{\beta} = 0.842 \). Based on experiments of the same type carried out in prior work, we quantified the variability of the data (variance, \( \sigma^2 \)) and established the minimum difference \( \Delta = \mu_1 - \mu_2 \) between averages that had a biological significance. By expressing both parameters as percentages and setting \( \sigma = 12.5\% = 0.125 \) and \( \Delta = 16\% = 0.16 \), we obtained \( n > 9.58 \). The actual number of mice is provided in Table 1. To minimize subjective bias, sample identity (e.g. genotypes) was randomized by associating an identification number to each sample before processing. No sample was excluded from the analysis.

**Preparation of COCs**

P5 mouse pups were humanely euthanized, both cochlea were carefully removed, quickly dissected in ice-cold HEPES-buffered (pH 7.2) HBSS (Cat. No. 14025050, Thermo Fisher Scientific, Waltham, MA, USA) and placed onto 5 mm round glass coverslips coated with Cell-Tak (Cat. No. 354240, Corning, Corning, NY, USA). COCs were incubated overnight in DMEM/F12 (Cat. No. 11320-074, Thermo Fisher Scientific) supplemented with 5% FBS (Cat. No. 10270-106, Thermo Fisher Scientific) and 100 \( \mu \)g mL<sup>−1</sup> ampicillin (Cat. No. A0166, Merck, St. Louis, MO, USA) at 37 °C in a humid air atmosphere enriched with 5% CO<sub>2</sub>. The tectorial membrane was carefully removed prior to each experiment.

**Generation and culture of ATP-WCBs**

To generate ATP-WCBs, we transduced HEK293-T cells (CRL-3216™, ATCC, Manassas, VI, USA) with a lentivector encoding P2Y<sub>2</sub> purinoceptor (P2Y<sub>2R</sub>), a high affinity G-protein-coupled receptor that mobilizes Ca<sup>2+</sup> from intracellular stores upon binding extracellular ATP.\(^{48,49}\) By using the Gibson assembly technique,\(^{50}\) we inserted the P2Y<sub>2R</sub> complementary DNA (a gift of Marta Fumagalli, Laboratory of Molecular and Cellular Pharmacology of Purinergic Transmission, University of Milan) into a 3rd generation backbone vector (pUltra-hot, Addgene plasmid #24130, Addgene, Watertown, MA, USA; a gift from Malcolm Moore) that uses 2A self-cleaving peptide technology\(^{51}\) to express the protein of interest (P2Y<sub>2R</sub>) and the mCherry fluorescent reporter\(^{52}\) in a 1:1 ratio under the UbC promoter. Viral particles based on this transfer vector were produced using a standard protocol,\(^{53}\) thereafter the virus-containing supernatant was harvested and stored into cyriovials in 1 ml aliquots at −80 °C.

Prior to viral transduction, parental HEK293-T cells were maintained in DMEM/F12 (Cat. No. 11320-074, Thermo Fisher Scientific) containing 10% heat inactivated FBS (Cat. No. 10270-106, Thermo Fisher Scientific), 1% penicillin/streptomycin (Cat. No. 15070063, Merck), 1% L-glutamine (Cat. No. 25300-024, Merck) and 1% sodium pyruvate (Cat. No. 11360070, Merck). Twenty-four hours before viral transduction (day 1), HEK293-T cells were trypsinized and plated onto 35 mm Petri dishes such that they would be 30–50% confluent at the time of transduction. On the day of transduction (day 2), the culture medium was removed and replaced with the thawed virus-containing supernatant. Sequa-brene (8 \( \mu \)g mL<sup>−1</sup>, Cat. No. S2667, Merck; synonym: hexadimethrine bromide, polybrene) was added to each Petri dish, followed by gentle swirl to enhance transduction efficiency. HEK293-T cells were incubated overnight, at 37 °C under a controlled atmosphere of CO<sub>2</sub> (5%), in the virus-containing supernatant. The following day (day 3), the transduction medium was removed and replaced with 2 ml of complete culture medium. The percentage of HEK293-T transduced cells (>95%), henceforth referred to as WBCs, was estimated 48 hours post-transduction by fluorescence microscopy of the mCherry reporter.

**Construction of the ATP dose–response curve for ATP-WCBs**

ATP-WCBs were re-plated in black 96-well plates (Cat No. 655090, Greiner Bio-One Italia S.r.l, Cassina de Pecchi, Italy) and cultured for 24 hours as detailed above. Thereafter, ATP-WCBs were incubated for 2 hours at 37 °C and 5% CO<sub>2</sub> with the loading solution of the FLIPR Calcium 6 Assay Kit (Cat. No. R8190, Molecular Devices, San Jose, CA, USA). Ca<sup>2+</sup>-Dependent fluorescence changes promoted by ATP binding to the P2Y<sub>2R</sub> were measured using a Varioskan LUX multimode microplate reader (Cat. No. VL0L00D0, Thermo Fisher Scientific) after injection of known amounts of ATP (from 0.1 nM to 1 \( \mu \)M) in the wells. To achieve the desired [ATP], a 1 \( \mu \)M stock solution was prepared by dissolving ATP (Cat. No. A1852, Merck) in PBS and diluting the stock in the loading solution of the FLIPR Calcium 6 Assay Kit.

**Ca<sup>2+</sup> imaging in individual ATP-WCBs**

To visualize Ca<sup>2+</sup> responses evoked by ATP in individual ATP-WCBs, the latter were plated on 12 mm Ø coverslips and incubated for 30 min at 37 °C in serum-free DMEM (Cat. No. 10-017-CV, Corning, Corning, NY, USA) supplemented with the Ca<sup>2+</sup>-selective fluorescent probe Fluo 8H™ AM (5 \( \mu \)M, dissociation constant \( K_d = 232 \) nM, Cat. No. 21091, AAT Bioquest, Sunnyvale, CA, USA). The incubation medium also contained Pluronic F-127 (0.1% w/v, Cat. No. P2443, Merck) and sulfinpyrazone (250 \( \mu \)M, Cat. No. S9509, Merck) to prevent dye sequestration and secretion.\(^{54}\) To allow for de-esterification, coverslips with loaded ATP-WCBs were transferred to the stage of a custom-made spinning disk microscope\(^{55}\) and perfused for 10 min at a flow rate of 0.1 mL min<sup>−1</sup> with extracellular medium (EXM) containing (in mM): NaCl 135, KCl 5.8, CaCl<sub>2</sub> 1.3,
NaH₂PO₄ 0.7, MgCl₂ 0.9, HEPES–NaOH 10, n-glucose 6, pyruvate 2, amino acids, and vitamins (pH 7.48, 307 mOsm). ATP dissolved in EXM was applied by pressure (5 pounds per square inch, PSI) using a PV820 pneumatic pico-pump (World Precision Instruments Inc., Sarasota, FL, USA) coupled to a glass microcapillary with an opening diameter of 4 µm placed under a 60× water immersion microscope objective (NIKON FLUOR, NA = 1.0, Nikon Corporation, Tokyo, Japan) near the center of the field of view and in close proximity to ATP-WCBs. For Ca²⁺ imaging, Fluo 8H™ fluorescence was excited using a 488 nm diode laser (Cat. No. COMPACT-150G-488-SM, World Star Tech, Markham, Ontario, Canada) by 45° reflection off a triband dichroic mirror (Cat. No. FF395/527/610-Di01, Semrock, Rochester, NY, USA). Fluo 8H™ fluorescence emission was filtered through a triband filter (Cat. No. FF425/527/685-25, Semrock) followed by a green band pass filter (Cat. No. ET535/30M, Chroma Technology Corp., Bellows Falls, VT, USA) located in front of a cooled s-CMOS camera (pco.edge, PCO AG, Kelheim, Germany) coupled to the microscope. Suramin (150 µM; Cat. No. S2671, Merck) was used to reversibly inhibit the ATP-evoked Ca²⁺ responses (see Results).

Dual focal plane multiphoton Ca²⁺ imaging in COCs and ATP-WCBs

The day before the experiment, ATP-WCBs were plated on 12 mm ∅ coverslips at a density of 2 × 10⁴ cells per cm². At experiment time, they were loaded with Fluo 8H™ AM as described above. COCs were plated on 5 mm ∅ coverslips the day before the experiment, and incubated for 45 min at 37 °C in serum-free DMEM supplemented with 20 µM Fluo 8H™ AM, Pluronic F-127 (0.1% w/v) and sulfinpyrazone (250 µM) at experiment time. To allow for de-esterification, both cultures were transferred into an ad hoc designed microfluidic chamber (Fig. 1), perfused for 10 min with EXM and placed on the stage of a custom-made two-photon microscope (Fig. 2) based on a Bergamo II architecture (Thorlabs Imaging System, Sterling, VI, USA). The polydimethylsiloxane (PDMS) insert of the chamber (Fig. 1b–d) included microfluidic channels connected by 25 mm stainless-steel tubing (ASI 316L, 0.25/0.12 mm, Unimed SA, CH-1007 Lausanne, Switzerland) and flexible external tubing (Tygon ST R-3603/R3607, Harvard Apparatus, Holliston, MA 01746 United States) to a proximally located peristaltic pump (P-70, Harvard Apparatus). This arrangement reduced the effective chamber volume to 10 µl and the total volume in the fluid path to 200 µl. The flow rate was kept ≤100 µl min⁻¹ to minimize the effects of fluid shearing stress, and perfusion was stopped during the image-recording session (which generally lasted 1 min). EXM (containing 2 mM Ca²⁺) was replaced by LCS, a medium with an endolymph-like Ca²⁺ concentration (20 µM) and an otherwise identical composition. LCS was supplemented either with the ATP analogue 6-N,N-diethyl-o-beta-gamma-dibromomethylene adenosine triphosphate (ARL67156, 100 µM, Cat. No. A265, Merck), or apyrase (40 U ml⁻¹, Cat. No. A6535, Merck), depending on the type of experiment (see Results).

The multiphoton system was equipped with two scanning heads, one with resonant-galvo (RG) mirrors and the other with galvo-galvo (GG) mirrors, and was coupled to a mode-
locked titanium–sapphire (Ti:Sa) fs pulsed laser (Chameleon Vision II Laser, Coherent, Inc., Santa Clara, CA, USA). The RG scanner was used for imaging, whereas the GG scanner was used to focally photodamage a pre-defined spot in the GER of preparations). Multiphoton excitation of Fluo 8H™ (37 °C, 5% CO₂), COCs plated onto 5 mm glass coverslips were transferred into a black 96-well plate containing 200 μl serum-free DMEM/F12 in each well. Before starting the ATP release-stimulation protocol, COCs were washed once with a solution (NCS) containing a normal (1.8 mM) extracellular Ca²⁺ concentration and (in mM): 137 NaCl, 5.36 KCl, 0.81 MgSO₄, 0.44 KH₂PO₄, 0.18 Na₂HPO₄, 25 HEPES and 5.55 μ-glucose (pH 7.3). COCs were then washed a second time either with NCS or with a zero Ca²⁺ solution (ZCS) containing (in mM): 137 NaCl, 5.36 KCl, 0.44 KH₂PO₄, 0.18 Na₂HPO₄, 0.1 EGTA, 25 HEPES and 5.55 μ-glucose (pH 7.3).

To quantify ATP release, COCs were incubated in either NCS or ZCS for 20 minutes at 37 °C, 5% CO₂. To limit degradation of the released ATP, both solutions were supplemented with ARL67156 at a final concentration of 100 μM. The amount of ATP released under these conditions was measured with a bioluminescent ATP assay kit (Cat. No. A22066, ThermoFisher Scientific) using a luminometer (Victor Light 1420, Perkin Elmer, Waltham, MA, USA). All bioluminescence measurements reported in this article fell within the linearity range of the ATP standard curve generated according to the manufacturer’s instructions. All experiments were performed at room temperature (22–25 °C).

Luciferin–luciferase ATP bioluminescence assay

After overnight incubation under standard culture conditions (37 °C, 5% CO₂), COCs plated onto 5 mm glass coverslips were transferred into a black 96-well plate containing 200 μl serum-free DMEM/F12 in each well. Before starting the ATP release-stimulation protocol, COCs were washed once with a solution (NCS) containing a normal (1.8 mM) extracellular Ca²⁺ concentration and (in mM): 137 NaCl, 5.36 KCl, 0.81 MgSO₄, 0.44 KH₂PO₄, 0.18 Na₂HPO₄, 25 HEPES and 5.55 μ-glucose (pH 7.3). COCs were then washed a second time either with NCS or with a zero Ca²⁺ solution (ZCS) containing (in mM): 137 NaCl, 5.36 KCl, 0.44 KH₂PO₄, 0.18 Na₂HPO₄, 0.1 EGTA, 25 HEPES and 5.55 μ-glucose (pH 7.3).

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Image processing and data analysis

Image processing and data analysis were carried out using the open source ImageJ software and MATLAB (R2019, The MathWorks, Inc., Natick, MA, USA). Ca²⁺ signals were quantified either as pixel-by-pixel relative changes of fluorescence emission intensity ΔF(t), where t is time, F(t) is fluorescence at time t, ΔF(t) = F(t) − F₀ and F₀ is the pre-stimulus fluorescence, or as 100×ΔF(t)/(Fₘₐₓ − F₀) where Fₘₐₓ = max[F(t)] during the time interval of the recording.

Statistics

For statistical data analysis, the normality of distribution was assessed using the Kolmogorov–Smirnov test and statistical comparisons of means were made using the Student t test. The Mann–Whitney U test was used for data which were not normally distributed and/or had dissimilar variance. All statistical analyses were performed using MATLAB (R2019). Mean values are quoted ± standard error of the mean (s.e.m.)
where \( p = p\text{-value} < 0.05 \) was assumed as statistically significant.

Results

Sensitivity of ATP-WCBs to ATP

To characterize the sensitivity of ATP-WCBs to ATP, we loaded cells with Ca\(^{2+}\) selective dyes and measured fluorescence intensity changes (\( \Delta F \)) evoked by different [ATP] in the range from 0.1 nM to 1 \( \mu \)M (see Methods). Fig. 3a shows dose-response data fitted with a Hill function

\[
\frac{\Delta F}{\Delta F_{\text{sat}}} = \left( 1 + \frac{\text{EC}_{50}}{\text{[ATP]}} \right)^{-1}
\]

(1)

where \( \Delta F_{\text{sat}} \) is the value attained by \( \Delta F \) at saturating values of the [ATP]. The fit yielded a half-maximal effective concentration (EC\(_{50} = 6.2\) nM) very close to the baseline [ATP] in the endolymph.\(^{28,29}\) Ca\(^{2+}\) signals evoked by 100 nM [ATP] (a nearly saturating concentration, see Fig. 3a and b) were dramatically reduced after 30 min of incubation with suramin (150 \( \mu \)M, Fig. 3c), a reversible P2Y\(_{12}\)R antagonist.\(^{39-41}\) The ATP-evoked \( \Delta F \) signal, integrated over time and averaged over the population of ATP-WCBs, increased 30-fold after drug washout (Fig. 3c and d). We note in passing that parental HEK293-T cells, from which these ATP-WCBs were derived, are very weakly coupled through gap junction channels composed of Cx43 subunits;\(^{62}\) therefore the observed Ca\(^{2+}\) signals were independent of the confluence state and reflected individual cell responses to the proximal [ATP].

Solving eqn (1) for [ATP] with EC\(_{50} = 6.2\) nM yields

\[
[\text{ATP}] = (6.2\text{ nM}) \left( \frac{\Delta F_{\text{sat}}}{\Delta F} - 1 \right)^{-1}
\]

(2)

which can be used to estimate [ATP] in the proximity of the ATP-WCBs from the measurement of \( \Delta F \) and \( \Delta F_{\text{sat}} \).

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Fig. 3  Ca\(^{2+}\) responses of ATP-WCBs evoked by exogenously applied ATP. (a) Each data point is the mean Ca\(^{2+}\)-dependent fluorescence change \( \Delta F/F_{\text{sat}} \pm \text{s.e.m.} \) for \( n = 8 \) wells of a 96-well plate. Each well contained ATP-WCBs seeded at a density of \( 1 \times 10^5 \) cells per cm\(^2\) and loaded with the FLIPR Calcium 6 Assay Kit (see Methods). The solid line is a fit with the Hill function described in the main text. (b) Representative traces showing ATP-evoked Ca\(^{2+}\) responses in individual ATP-WCBs loaded with Fluo8\(^{TM}\); each trace was generated as the pixel average of \( \Delta F \) signals in a region of interest (ROI) encompassing a responding cell. (c) Shown are pseudo-color images acquired using the spinning disk confocal fluorescence microscope (see Methods) and representing percent fluorescence changes, encoded as shown by the color scale bar, evoked by 100 nM ATP; each image is the maximal projection rendering of all frames in a sequence of 490 fluorescence images acquired at 4 frames per second (f.p.s.) after 30 minutes of incubation with suramin (150 \( \mu \)M, left) and after washout (right); scale bars: 30 \( \mu \)m. (d) Box plots represent distributions of \( \Delta F \) signals integrated over time in \( n = 49 \) ATP-WCBs from the experiments in (c) (***, \( p < 0.001 \); Mann–Whitney \( U \) test; a.u., arbitrary units).
Together, these experiments suggested that ATP-WCBs, if placed sufficiently close to the ATP source in a suitable environment, had the required sensitivity to detect physiologically relevant levels of ATP.

Design, construction and test of a microfluidic chamber suitable to detect ATP release from COCs

To determine whether ATP-WCBs responded to the ATP released in the extracellular medium by COCs, we exploited the intrinsic optical sectioning capabilities of multiphoton microscopy and performed dual focal plane Ca²⁺ imaging in an ad hoc designed, closed and transparent micro-fluidic chamber hosting ATP-WCBs at the top and a proximally located COC at the bottom (see Methods and Fig. 1 and 2). To increase detection sensitivity, hydrolysis of ATP was reduced by supplementing the extracellular medium with ARL67156 (100 μM), a widely used ectonucleotidase inhibitor.

Initially, we used the second scanning head and fast beam shuttering of the multiphoton microscope (Fig. 2) to rapidly photodamage a single GER cell (see Methods) in COCs from wild type C57BL6/N pups, resulting in the time-controlled the intrinsic optical sectioning capabilities of multiphoton microscope and performed dual focal plane Ca²⁺ imaging in an ad hoc designed, closed and transparent micro-fluidic chamber hosting ATP-WCBs at the top and a proximally located COC at the bottom (see Methods and Fig. 1 and 2). To increase detection sensitivity, hydrolysis of ATP was reduced by supplementing the extracellular medium with ARL67156 (100 μM), a widely used ectonucleotidase inhibitor.

Replacing ARL67156 in the extracellular medium with apyrase (40 U ml⁻¹), an ATP-hydrolyzing enzyme, caused a dramatic reduction of Ca²⁺ wave amplitude and extent in COCs. Ca²⁺ responses in the overlying ATP-WCBs were correspondingly reduced, confirming that they were due to the diffusion of the released ATP in the narrow fluid gap (20–100 μm) intercalated between the two cellular systems. For the representative experiment in Fig. 4b, the responses, integrated over time and summed over all ATP-WCBs within the field of view, were 3.36 × 10⁶ A.U. in the presence of ARL67156 vs. 7.00 × 10⁴ A.U. in the presence of apyrase (a 48-fold reduction). By using eqn (2) to estimate the average ATP that reached ATP-WCBs in the presence of ARL67156, we obtained values in the range from 5 nM to 48 nM. Similar results were obtained in two other independent experiments.

Together, these results suggested that ATP-WCBs in the microfluidic chamber could detect ATP release during stimulated Ca²⁺ wave propagation in the GER.

Analysis of spontaneous Ca²⁺ wave propagation and ATP release from the GER of genetically modified mice by dual multiphoton imaging in the microfluidic chamber

Based on the positive outcomes of the aforementioned experiments, we decided to investigate spontaneous Ca²⁺ wave propagation in the GER of pups with global deletion of Panx1 (Fig. 5 and Video S2 in the ESI‡) or Gjb6 (Fig. 6 and Video S3 in the ESI‡) using the microfluidic chamber. The distributions of ATP-WCB Ca²⁺ responses, integrated over time and driven by spontaneous Ca²⁺ wave propagation in the underlying GER from Panx1⁻/⁻ and Panx1⁺/+ COCs, were statistically indistinguishable (Fig. 7a: Panx1⁻/⁻: mean = 2.38 × 10⁴ A.U., median = 7.33 × 10³ A.U., I.Q.R. = 2.19 × 10⁴ A.U.; Panx1⁺/+: mean = 2.07 × 10⁴ A.U., median = 3.88 × 10³ A.U., I.Q.R. = 1.74 × 10⁴ A.U.; m = 1695 ATP-WCBs in each case; p = 0.105, Mann–Whitney U test; pooled results from n = 7 pups

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of each genotype). By using eqn (2) to estimate the average [ATP] near ATP-WCBs, we obtained values in the range from 7 nM to 30 nM for Panx1−/− as well as Panx1+/+ COCs.

In prior work, the frequency of GER spontaneous Ca2+ waves was dramatically reduced in Gjb6−/− animals compared to their siblings with normal Gjb6 alleles (Gjb6+/+).18,20 Here, in addition, we report significantly different distributions of Ca2+ responses obtained from ATP-WCBs overlying Gjb6−/− COCs in the microfluidic chamber (Fig. 7b; mean = 2.90 x 10^3, median = 460, I.Q.R. = 1.97 x 10^3 A.U.) vs. Gjb6+/+ COCs (mean = 2.30 x 10^4, median = 2.59 x 10^3, I.Q.R. = 1.67 x 10^4 A.U.; m = 606 ATP-WCBs in each case; p = 3.04 x 10^-27, Mann-Whitney U test; pooled results from n = 4 pups of each genotype, aged P5).

Independent estimate of ATP release from Panx1−/− and Cx30−/− cochlear organotypic cultures

As an additional control, we used also a standard luciferin–luciferase ATP bioluminescence assay67 to measure ATP release from COCs in 96-well plates (see Methods). In COCs obtained from Panx1+/+ and Panx1−/− P5 pups, ATP release was equally enhanced by lowering the concentration of extracellular Ca2+ from 1.8 mM (NCS) to 0 mM (ZCS), a procedure that increases the open probability of connexin hemichannels68 without affecting pannexin channels.69–71 Specifically, the amount of ATP released in the extracellular medium in ZCS was 205% ± 30% of the NCS control in Panx1−/− COCs (p = 0.015, t test) and 268% ± 54% in Panx1+/+ COCs (p = 0.011, t test). The ATP released in the extracellular medium was not significantly different between the two genotypes, either in ZCS (p = 0.265, t test) or in NCS (p = 0.93, t test, n = 6 cultures of either genotype; Fig. 8a). In contrast, the ATP released in the extracellular medium was not significantly different between the two genotypes, either in ZCS (p = 0.0001, t test) and 145% ± 16% in Gjb6+/+ COCs (p = 0.074). In this case, the amount of released ATP in ZCS was significantly different between the two genotypes (p = 0.001, t test; Fig. 8b).

Together, the results of Fig. 4–8 indicate that: (i) ATP-WCBs in the microfluidic environment had the required
sensitivity to detect the ATP released by stimulated as well as spontaneous Ca\(^{2+}\) wave propagation in the GER; (ii) ATP release was unaffected by Panx1 genetic ablation (iii) but strongly reduced by Gjb6 ablation.
Discussion and conclusions

Prior work has unveiled a critical role for connexin hemichannels in Ca2+ wave propagation in both the LER and the GER.17,18,22 The goal of this work was to determine whether ATP release through connexin hemichannels underlies Ca2+ wave propagation also in the GER of the developing cochlea (rather than only in the LER, as abundantly documented). To address this key issue, we used microfluidics, which is capable of manipulating cells and organ models in a highly controllable microenvironment (Fig. 1), coupled with multiphoton imaging (Fig. 2). This disruptive combination of key enabling technologies is creating breakthroughs in current understanding of cell and developmental biology and is providing insights into disease diagnosis in numerous fields.72

WCBs were the third essential element of our design (Fig. 3). For more than a quarter century, WCBs have provided critical insight into the physiological effect of numerous analytes (reviewed in ref. 43, 73 and 74). To create ATP-WCBs, we infected HEK-293T cells with a lentiviral vector encoding the highly sensitive P2Y2R, one of the two major P2Y receptors expressed in the developing cochlea.3,6 The binding of ATP to P2Y2R promotes the generation of diacylglycerol and InsP3 from PLC-dependent hydrolysis of PIP2.18 Intracellular diffusion of InsP3 and subsequent binding to InsP3 receptors triggers Ca2+ release from the endoplasmic reticulum (ER), increasing the cytosolic Ca2+ concentration.75 By optically monitoring these Ca2+ fluctuations with Ca2+-selective dyes,58 we determined that ATP release in other cellular systems,76 our ATP-WCBs had the required sensitivity to detect [ATP] fluctuations in ATP-WCBs to stimulated (Fig. 4) as well as spontaneous Ca2+ wave propagation in the GER of genetically modified mice (Fig. 5 and 6). Ca2+ signals where inhibited by apyrase, as previously reported for ATP-dependent Ca2+ wave propagation in the LER of wild type mice.4 Importantly, spontaneous Ca2+ wave propagation in the GER resulted in time-delayed ATP-WCB responses that were strongly reduced by global deletion of Gjb6, but unaffected by global deletion of Panx1 (Fig. 7). Quantification of ATP release from COCs by means of a standard luciferin-luciferase bioluminescence assay corroborated this conclusion (Fig. 8). However, the luciferin-luciferase approach, the main strategy to measure ATP nowadays, which focuses on supernatants, falls short of providing the kind of information that is made available by our organ-on-chip/ATP-WCBs system and optical technology.

Although pannexin 1 channels71 are thought to mediate ATP release in other cellular systems,76-81 the function of cochlear pannexins remains obscure.34,82-84 Prior work with the Panx1−/− strain used here confirmed (i) successful ablation of Panx1 both in the cochlea and in the brain; (ii) normal hearing sensitivity, normal function of the outer hair cell-based “cochlear amplifier” and absence of cochlear nerve defects; (iii) normal expression of inner ear connexins and gap junction communication in the organ of Corti.74 These published results and the data presented here together rule out that pannexin 1 channels contribute to the purinergic signaling that underlies Ca2+ wave propagation in the developing cochlea.2

Gjb6−/− mice, in which Cx30 is deleted globally38 and Cx26 is dramatically down-regulated in the sensory epithelium of the developing cochlea,33 are a model of non-syndromic hearing loss and deafness (DFNB1, reviewed in ref. 37 and 85). Prior work revealed impaired Ca2+ wave propagation in the developing cochlea of these mice,18,33 which could be
restored by viral transduction with a bovine adeno-associated vector (BAAV) encoding Cx30. The expression of connexin hemichannels at the endolympathic surface of the cochlear sensory epithelium has been confirmed by immunolabeling with antibodies that bind the extracellular loops of Cx26 and Cx30. In addition, spontaneous Ca\(^{2+}\) wave propagation at the surface of the GER was inhibited by (i) flufenamic acid and (ii) the abEC1.1 antibody, which, unlike carbenoxolone, is ineffective on pannexin 1 channels at concentrations <300 \(\mu\)M (ref. 80 and 91). ATP release during Ca\(^{2+}\) wave propagation in the GER has been proposed to trigger Ca\(^{2+}\) action potential (AP) activity in immature IHCs, driving bursts of APs in the auditory nerve fibers. However, subsequent work has shown AP firing in the complete absence of Ca\(^{2+}\) activity in the GER. Therefore, it remains to be established if and how these two types of spontaneous activities correlate with each other. This stands out a key open question for cochlear physiology and pathology and cannot be addressed with technological advances presented here have the potential to shed light on this as well as a plethora of other unrelated open issues that concern the role of paracrine signaling in physiology and pathology and cannot be addressed with standard methods.

Author contributions

Conceptualization, F. M. (Fabio Mammano); methodology, F. M., Fl. M. (Flavia Mazzarda), A. D'E., R. M., A. D. N., F. R. B., A. T. M., L. B., F. C.; resources, F. M., F. R. B., L. B., A. S., M. R., F. S.; investigation, F. M., C. P., G. Z., V. Z., C. N.; formal analysis, data curation, visualization and writing—original draft preparation, Fl. M.; supervision, writing—review and editing, project administration and funding acquisition A. M. S., M. R., J. Y., F. M. All the authors have read and agreed to the published version of the manuscript.

Conflicts of interest

The authors declare no conflict of interest. The funders had no role in the design of the study, in the collection, analyses, or interpretation of data, in the writing of the manuscript, or in the decision to publish the results.

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