RESEARCH ARTICLE

In-situ recording of ionic currents in projection neurons and Kenyon cells in the olfactory pathway of the honeybee

Jan Kropf1,2*, Wolfgang Rössler1

1 Behavioral Physiology and Sociobiology (Zoology II), Biozentrum, University of Würzburg, Würzburg, Germany, 2 Centre for Neural Circuits and Behaviour, University of Oxford, Oxford, United Kingdom

* jan.kropf@cncb.ox.ac.uk

Abstract

The honeybee olfactory pathway comprises an intriguing pattern of convergence and divergence: ~60,000 olfactory sensory neurons (OSN) convey olfactory information on ~900 projection neurons (PN) in the antennal lobe (AL). To transmit this information reliably, PNs employ relatively high spiking frequencies with complex patterns. PNs project via a dual olfactory pathway to the mushroom bodies (MB). This pathway comprises the medial (m-ALT) and the lateral antennal lobe tract (l-ALT). PNs from both tracts transmit information from a wide range of similar odors, but with distinct differences in coding properties. In the MBs, PNs form synapses with many Kenyon cells (KC) that encode odors in a spatially and temporally sparse way. The transformation from complex information coding to sparse coding is a well-known phenomenon in insect olfactory coding. Intrinsic neuronal properties as well as GABAergic inhibition are thought to contribute to this change in odor representation. In the present study, we identified intrinsic neuronal properties promoting coding differences between PNs and KCs using in-situ patch-clamp recordings in the intact brain. We found very prominent K+ currents in KCs clearly differing from the PN currents. This suggests that odor coding differences between PNs and KCs may be caused by differences in their specific ion channel properties. Comparison of ionic currents of m- and l-ALT PNs did not reveal any differences at a qualitative level.

Introduction

Olfaction is a crucial sense for almost all animal species, and olfactory systems show striking similarities across a wide range of taxa—like the odorant receptor proteins or the glomerular neuronal architecture of the first relay station in the brain (for example reviewed by [1]). Honeybees need a powerful olfactory system for the location and evaluation of food sources, social (pheromone) communication, nestmate recognition, and for finding mating partners. Odorants are received by odorant receptors located in olfactory sensory neurons (OSNs) that are housed in different types of olfactory sensilla on the antennae (reviewed in [2]). OSN axons...
form two nerve bundles running along the antenna to the antennal lobe (AL) entrance and diverge into four distinct tracts (T1-T4) within the AL [3]. Here, OSN axons synapse on projection neurons (PNs) and local interneurons (LNs) forming spheroidal structures that are commonly termed olfactory glomeruli [2]. Glomeruli represent functional units of the AL and encode odor information in a spatially combinatorial fashion—theoretically allowing for coding an almost infinite number of odors (for example [4]). The exact synaptic connectivity in the honeybee AL between OSNs, LNs and PNs is not yet known. In the honeybee, three output tracts are formed by distinct groups of PNs: the medial (m-ALT), lateral (l-ALT) and medio-lateral antennal lobe tract (ml-ALT) [3,5,6]. They convey the olfactory information to the mushroom bodies (MBs) and the lateral horn (LH). The m-ALT is formed mainly by uniglomerular PNs and proceeds medially to the MB calyces and finally reaches the LH. The l-ALT, formed by another group of mainly uniglomerular PNs, runs just in the opposite direction and innervates the LH first before the axons target the MBs (reviewed in [7]). Both m- and l-ALT PNs possess axonal arborizations in the MBs and form distinct synaptic complexes, so called microglomeruli, with MB intrinsic neurons, the Kenyon cells (KC) [8,9]. The ml-ALT comprises multiglomerular PNs that do not innervate the MBs but project directly to the LH. Whereas multiple PN tracts can be found in various insect species including *Drosophila melanogaster* [10] and *Manduca sexta* [11], symmetrical dual olfactory pathways formed by uniglomerular PNs proceeding in parallel to the MBs, so far, have only been found in Hymenoptera [2,7,12]. The function of the dual olfactory pathway in parallel olfactory coding is still under debate. Both tracts convey information about a largely similar range of odorants [13–17] with only queen mandibular pheromone components and brood pheromone components as rare exceptions [18]. PNs of both tracts exhibit various response patterns, including tonic response patterns, phasic-tonic response patterns, bursts with a post burst hyperpolarization phase response patterns, and even inhibitory responses [5,13,17]. The reason for this general complexity in odorant response patterns may be that PNs need to convey a large amount of information from a very high number of OSNs (bottleneck). Approximately 60,000 OSNs [19] synapse on only ~900 PNs [20], which have to transmit the odor information reliably to a large number of KCs (~183,000 in total on each side of the brain) in the MBs [21,22].

In contrast to PNs, KCs are far more numerous and can be expected to employ less complex response dynamics. Indeed, calcium imaging experiments revealed that only a few honeybee KCs respond to the same odor. Furthermore, KC odor responses were shown to be shorter than both the actual odor stimulation and the PN odor response. KCs thus employ a temporally as well as spatially sparse information code [23]. The temporal response pattern of KCs was recently shown to be unaffected by GABAergic inhibition [24] and, therefore, can be assumed to be mainly caused by intrinsic electrical properties.

In the honeybee, intrinsic electrical properties of PNs and KCs, so far, have mostly been studied in primary cell cultures. PNs were shown to possess Na⁺, K⁺ and Ca²⁺ currents that are commonly known to be involved in action potential (AP) generation, and to contain Ca²⁺ dependent K⁺ channels, which could serve as an intrinsic self-inhibitory mechanism [25,26]. The whole-cell currents of KCs drastically differ from the ones found in PNs: KCs have very prominent A-type K⁺ currents [25,27–29] and most likely Ca²⁺ dependent K⁺ currents [30]. These were shown to be involved in sparse coding in cockroaches [31].

The aim of this study was to identify neuronal electrical properties supporting the complex odor response patterns of PNs [13] and the sparse coding properties of KCs [23,24] using *in-situ* patch-clamp recording techniques in the intact adult brain. Furthermore, using live tracing of PN cell bodies for proper identification of PN subtypes we were able to investigate, for the first time, m- and l-ALT PNs separately.
Material and methods

Animals

All honeybees (*Apis mellifera carnica*) were taken from colonies of our departmental bee station at the University of Würzburg. Only adult bees were used, but we did not control for the exact age of the bees. The bees were cooled in a refrigerator (4°C) and harnessed in custom-built plastic holders. Immediately after the bees had recovered from cooling, they were fed with a ~50% sucrose solution in distilled water.

Pre-experimental PN staining

To distinguish PN cell bodies from LN cell bodies, we stained PNs 10 to 20 hours before recording. A small window was cut by removing the cuticle above the MB calyces, and glands and trachea covering the calyces were removed manually using fine forceps. Thin-walled glass micropipettes (1B100F-3, WPI, Sarasota, USA) were pulled with a Zeitz Puller (Zeitz-Instruments, Martinsried, Germany) and coated with Microruby™ (tetramethylrhodamine dextran with biotin, 3,000 MW, lysine-fixable, D-7162; Molecular Probes, Eugene, Oregon, USA) dissolved (3–5%) in distilled water. The dye-coated micropipettes were used to puncture the calyces of both MBs, to have the dye subsequently taken up by the neurons. After the staining procedure, a two component tissue glue (Kwik-Sil™ World Precision Instruments, Sarasota, USA) was used to seal the window and to prevent desiccation of the brain.

Preparation for recording

All glands and trachea covering the brain were manually removed using fine forceps. Then, the heads were severed from the thorax. The head capsule was fixed in the recording chamber (RC-22C, Warner Instruments, LLC, Hamden, Connecticut, USA). With the head oriented in an upright fashion, this preparation allowed access to l-ALT PN and KC cell bodies. In order to gain access to m-ALT PN cell bodies, the complete preparation was turned upside down. The proboscis and the cuticle beneath the brain were removed using microscissors. Glands, muscles and trachea covering the brain were gently removed. Additionally, the SEG was displaced by pushing it into the direction of the MBs. To identify stained cell bodies, an Olympus imaging system (200×-800×, upright microscope: BX51WI, filter set: excitation 560/40 DCTX 590 emission 610 LP, objective: XLUMP, NA 0.95, light source: MT20, software: Cell R v2.5, all Olympus Imaging Europa, camera: model 8484-03G Hamamatsu Photonics) was used. The only neurons connecting the MBs with the AL are uniglomerular PNs—therefore, stained cell bodies in the AL could be clearly identified as l- and m-ALT PNs. KC cell bodies are numerous and can be easily identified. Differentiation between clawed (type II) and spiny (type I) KCs is possible according to their anatomical location. Only type II (clawed) KCs with cell bodies outside the calyx cup [22] were recorded in our experiments. This type of KCs is likely to receive convergent input from both PN tracts [7,32].

Patch-clamp recordings

Patch-clamp electrodes were pulled from thick-walled borosilicate glass with filament (GB150F-8P, Science Products, Hofheim, Germany) with a Zeitz Puller. Electrodes had tip resistances of 4–6 MΩhm (PN-recording) and 6–9 MΩhm (KC-recording) measured in the extracellular solution. The patch-clamp electrodes were moved to the cell bodies by using an electrical micromanipulator (Junior Unit, Luigs & Neumann, Ratingen, Germany). For patch-clamp recordings, we used an Axopatch 200b amplifier, a Digidata 1440A acquisition board and the ClampEx software (all Molecular devices, Sunnyvale, California, USA).
extracellular solution comprised in mmol: NaCl (140), KCl (5), MgCl₂ (1), CaCl₂ (2.5), NaHCO₃ (4), NaH₂PO₄ (1.2), HEPES (6) and glucose (14), adjusted to pH 7.4 with NaOH. The intracellular solution contained in mmol: K-glucuronate (110), HEPES (25), KCl (10), MgCl₂ (5), Mg-ATP (3), Na-GTP (0.5) and EGTA (0.5), pH 7.2. Currents were isolated by blocking Na⁺ currents and Na⁺ dependent currents with TTX (10⁻⁷ mol) and by blocking Ca²⁺ currents and Ca²⁺ dependent currents with CdCl₂ (5x10⁻⁵ mol) in the extracellular solution. All chemicals were purchased at Sigma-Aldrich Chemie Gmbh (Munich, Germany) or at Carl Roth GmbH + Co. KG (Karlsruhe, Germany).

Data analyses and statistics
Prior to voltage-clamp experiments, recordings were allowed to stabilize for approximately one minute. Voltage clamp protocols under standard conditions; TTX conditions and TTX/CdCl₂ conditions were completed within 10 minutes after break-in. Leakage currents were subtracted using a p/5 protocol in ClampEx. The series resistance did not significantly change between recording under standard conditions and under TTX+CdCl₂ conditions in l-ALT PNs (paired Wilcoxon test: p>0.05). All patch clamp raw data traces were analyzed with the "statistics" function implemented in pClamp (Molecular devices, Sunnyvale, California, USA). The positive and negative maximal values at the transient peak and during the persistent phase were determined. Membrane voltages were directly measured with the digital meter on the amplifier and corrected by subtracting the liquid junction potential (13mV) of our solutions. Cell capacitances were obtained with the electrode test function implemented in ClampEx. The obtained data sets were further processed with Excel (Microsoft Corporation Redmond, Washington, USA), IV plots were also generated with Excel. All statistics as well as box plots were made with R (R Foundation for Statistical Computing, Vienna, Austria).

Neuronal anatomical analyses and image composition
Brains stained with Microruby™ were put into fixative solution (4% formaldehyde) overnight and rinsed five times for 10 min in PBS (phosphate-buffered saline, pH 7.2) the next day. Afterwards, the brains were dehydrated in an ascending ethanol series (50, 70, 90, 95, 2x 100%, each 10 min) and cleared in methyl salicylate (M2047, Sigma-Aldrich Chemie Gmbh, Munich, Germany). The brains were then mounted in methyl salicylate in custom made microscopy slides and scanned with a confocal laser-scanning microscope (Leica TCS SP2; Leica Microsystems, Wetzlar, Germany). Image stacks were processed with ImageJ 1.46j (Wayne Rasband, National Institutes of Health, Befesta, Md., USA). All images were finally arranged with CorelDrawX6 (Corel, Ottawa, ON, Canada).

Results
Pre-experimental identification of neurons
One important prerequisite for selectively recording from PNs was the identification of the respective neuronal cell bodies in the AL tissue, as they are embedded in many cell bodies of AL LN's [3]. The injection of dextran coupled dyes into the brain leads to the uptake of the dye in neurons with either pre- or post-synaptic profiles in the area where the dye was injected. The dye then is transported retrogradely as well as anterogradely along the respective neurites. Confocal microscopy analyses after the injection of Microruby™ into the MBs revealed that the dye was taken up by KCs, PNs from the optic ganglia, PNs from the subesophageal (SEG) tract (potentially gustatory and/or mechanosensory PNs), and olfactory PNs from the antennal lobe. In a broad overview, KCs proceeding from the MB calyces to the vertical lobe,
visual commissures connecting the optic lobes, and tracts from SEG PNs and olfactory PNs ascending from the AL to the MBs could be distinguished (Fig 1A). However, the only cells with axonal arborizations in the MBs as well as cell bodies in the AL are uniglomerular olfactory PNs [3]. Therefore, cell bodies in the AL stained after dye injection into the MBs could clearly be identified as PNs ascending to the MBs. PN ramifications within glomeruli as well as stained cell bodies clustered around the AL were visible and can be identified as l- and m-PNs according to anatomical data from Fig 2 in [3] (Fig 1B). We approached these PN cell bodies in situ using a fluorescence light microscope (Fig 1C).

**Recording under standard conditions**

As this represents the first study that comparatively investigated the composition of ionic currents of honeybee PNs as well as KCs by in situ recordings in the intact brain, we started by analyzing and comparing basic neuronal properties of the three neuronal classes under...
standard conditions. The membrane voltages of l-ALT PNs under our recording conditions ranged from -30 to -50 mV—the mean membrane voltage of all cells was -42.0 mV (N = 12). In contrast, KC membrane voltages were more negative and ranged from -42 to -61 mV (mean = -52.0 mV, N = 7). As the whole-cell patch clamp technique can induce leakage currents, which potentially depolarize the cells, we assumed even lower membrane voltages in intact cells. Therefore, during current clamp recordings, a minimal negative current was injected to achieve resting potentials of -60 mV in PNs and -80 mV in KCs.

Because l-ALT PN cell bodies are accessible without turning the preparation around and removing parts of the SEG l-ALT PNs recordings proved to be much easier than recording from m-ALT PNs. Therefore, we first focused on l-ALT PN recordings for the comparison between PNs and KCs. PNs recorded in current clamp started to fire action potentials at membrane voltages from -50 to -40 mV, and KCs started to fire action potentials at -30 mV (Fig 2A). In voltage-clamp recordings under standard conditions, all recorded neurons showed transient inward, transient outward and persistent outward currents (Fig 2B). Considering the ion concentrations we used for recording under standard conditions, we conclude that the inward currents were mainly Na\(^+\) currents and only partly Ca\(^{2+}\) currents, whereas all outward currents likely were K\(^+\) currents. The currents that were activated first during the voltage step protocol (-70 to 70 mV in PNs, -90 to 70 mV in KCs) were inward currents. The inward currents of PNs were activated at approximately -50 mV, the inward currents of KCs at -30 mV (Fig 2B). The transient K\(^+\) currents were activated right after the inward currents at -40 mV in both PNs and KCs (Fig 2B). These currents deactivated very fast in PNs, and they were only slightly higher than the persistent K\(^+\) currents. In contrast, KCs exhibited a very prominent transient outward current with about twice the amplitude of the persistent K\(^+\) current. This current activated and deactivated significantly slower than the transient current in PNs (Fig 2B). Persistent currents activated at -30 mV in both neuron types, yet they showed different IV relations in PNs and KCs. PN persistent currents increased with increasing membrane voltages in a linear IV relationship (Fig 2C). Conversely, the persistent K\(^+\) current of KCs showed a nonlinear, distinct N-shape in the IV plot, which increased until 30 mV to then decrease until reaching 50 mV (Fig 2C). This N-shape hints at the presence of Ca\(^{2+}\)-dependent K\(^+\) currents, which decrease once the driving force for Ca\(^{2+}\) declines (for example, [31]).

l-ALT PN currents under the influence of TTX and CdCl\(_2\)

As the IV relationships we observed under standard conditions indicated Na\(^+\) currents, K\(^+\) currents as well as Ca\(^{2+}\)-sensitive K\(^+\) currents (Fig 2B and 2C), we used tetrodotoxin (TTX) and CdCl\(_2\) as primary pharmacological agents. TTX (10\(^{-7}\) mol), a toxin which is well known from pufferfish, blocks voltage dependent Na\(^+\) channels and Na\(^+\) dependent K\(^+\) channels. CdCl\(_2\) (5x10\(^{-5}\) mol) blocks Ca\(^{2+}\) channels and, therefore, also affects Ca\(^{2+}\) dependent K\(^+\) channels. In the IV plots obtained after adding TTX to the extracellular solution, the fast inward currents were almost completely abolished in PNs (Fig 3A). Thus, we conclude that the inward currents are dominated by Na\(^+\) currents. Additionally, both the transient and the persistent K\(^+\) current were strongly reduced after adding TTX, which leads to the conclusion that the outward current partly consists of Na\(^+\) dependent K\(^+\) currents (K\(_{Na}\)). To be able to identify purely TTX sensitive currents, we subtracted the IV curves obtained with TTX in the extracellular solution as a control from the curves obtained under standard conditions (Fig 3B). The purely TTX sensitive traces revealed a very fast voltage dependent Na\(^+\) current, a very fast peak K\(_{Na}\) and a sustained K\(_{Na}\). The two TTX sensitive K\(^+\) currents, the very fast transient, and the persistent K\(^+\) current are all K\(_{Na}\) currents activated by the fast Na\(^+\) inwards current and the sustained Na\(^+\). The transient K\(_{Na}\) current activated at a slightly higher membrane voltage than the Na\(^+\).
Fig 2. A: Action potentials (AP) of a honeybee lateral antennal-lobe tract projection (l-ALT PN) and a Kenyon cell (KC) elicited in current-clamp mode. The PN started to generate APs at ~ -50mV, the KC at -40 mV. B: Representative voltage clamp recordings of an l-ALT PN and a KC. I-ALT and m-ALT PNs have similar currents, but KCs differ drastically as they have only small $I(\text{Na}^+\text{,Ca}^{2+})$ and a prominent $I(K^+)$ with about twice the size compared to the $I(K^+)$ in KCs. PNs were kept at -70 mV, KCs at -90 mV. In both PNs and KCs, a quick hyperpolarizing step of -20 mV was used, then the membrane voltage was increased in 10mV increments. C: I-V plots of the mean currents of the two neuronal types. Note the persistent $K^+$ current decreasing above 30 mV hinting Ca$^{2+}$ dependent $K^+$ currents in the $I(K^+)$ in KCs.

https://doi.org/10.1371/journal.pone.0191425.g002
inward current and increased drastically until 0 mV. It increased only marginally above 0 mV. (Fig 4A). In a further step, CdCl$_2$ was added to the extracellular solution to block all Ca$^{2+}$ currents and, therefore, also the K$^+$ currents depending on extracellular Ca$^{2+}$ (K$_{Ca}$). These currents were analyzed by subtracting the IV traces obtained after adding CdCl$_2$ to the extracellular solution from the traces obtained under standard conditions (Fig 3C). The current subtraction revealed a slight peak of inward currents between -30 and -20 mV indicating the presence of Ca$^{2+}$ currents. Furthermore, two K$_{Ca}$ currents, a transient current (K$_{Ca}$t) and a persistent current (K$_{Ca}$p), could be observed. Both K$_{Ca}$t current and K$_{Ca}$p current activated between -40 and -30 mV. The K$_{Ca}$t current increased until 0 mV and reached a plateau there, which is, similar to the decreasing transient K$_{Na}$, most likely due to a decreasing driving force for Ca$^{2+}$ ions. Again, K$_{Ca}$p did not reach plateau levels (Fig 4B).

In summary, l-ALT PNs exhibit transient Na$^+$ currents (Na$_t$), transient Na$^+$ dependent K$^+$ currents (K$_{Na}$t), Ca$^{2+}$ currents, transient Ca$^{2+}$ dependent K$^+$ currents (K$_{Ca}$t), persistent Ca$^{2+}$ dependent K$^+$ currents (K$_{Ca}$p).
Qualitative comparison between m-ALT and l-ALT PNs

In a set of pilot experiments, we selectively recorded from m-ALT PNs using the head preparation with fluorescently labeled PNs in an upside down position and after removal of the SEG. Recording of m-ALT PNs, therefore, was much more difficult compared to l-ALT PN recordings. The mean resting membrane voltage of m-ALT PNs was -39.4 mV (N = 4) and close to the l-ALT PN resting membrane voltage -42.0 mV (N = 12). Pharmacological isolation of currents did not reveal any qualitative differences between m- and l-ALT PNs (Fig 4). M-ALT

---

Fig 4. Comparison after pharmacological blocking of Na\(^+\), Ca\(^{2+}\), Na\(^+\) dependent and Ca\(^{2+}\) dependent currents in lateral and medial antennal-lobe tract projection neurons (l-ALT PNs, m-ALT PNs). A, B: Mean current-voltage plots of TTX sensitive (A) and CdCl\(_2\) sensitive (B) isolated currents in l-ALT PNs. Transient Na\(^+\) and K\(^+\) currents as well as persistent K\(^+\) currents can be observed. C, D: Mean current-voltage plots of TTX sensitive (C) and CdCl\(_2\) sensitive (D) isolated currents in m-ALT PNs. Transient Na\(^+\) and K\(^+\) currents as well as persistent K\(^+\) currents can be observed.

https://doi.org/10.1371/journal.pone.0191425.g004

**Qualitative comparison between m-ALT and l-ALT PNs**

In a set of pilot experiments, we selectively recorded from m-ALT PNs using the head preparation with fluorescently labeled PNs in an upside down position and after removal of the SEG. Recording of m-ALT PNs, therefore, was much more difficult compared to l-ALT PN recordings. The mean resting membrane voltage of m-ALT PNs was -39.4 mV (N = 4) and close to the l-ALT PN resting membrane voltage -42.0 mV (N = 12). Pharmacological isolation of currents did not reveal any qualitative differences between m- and l-ALT PNs (Fig 4).
PNs also exhibit transient Na$^+$ currents (Na$t$), transient Na$^+$ dependent K$^+$ currents (K$_{Na,t}$), Ca$^{2+}$ currents, transient Ca$^{2+}$ dependent K$^+$ currents (K$_{Ca,t}$), and persistent Ca$^{2+}$ dependent K$^+$ currents (K$_{Ca,p}$) (Fig 4C and 4D). Although, due to the difficulties of the upside down preparation, recordings in this experiment could only be obtained from a limited number of m-ALT PNs, the results, at a qualitative level, indicate a high degree of similarity in the electrical properties of l-ALT and m-ALT PNs under standard conditions.

### Comparison of transient currents in PNs and KCs

The transient K$^+$ currents very clearly differed between PNs and KCs. Our pharmacological analysis revealed that PN transient K$^+$ currents are mostly dependent on Na$^+$ or Ca$^{2+}$ influx (Fig 3). Furthermore, in contrast to the persistent K$^+$ current, the transient K$^+$ current in KCs rose linearly and, therefore, hints at a mostly voltage-dependent transient K$^+$ current (Fig 2C). By looking at an expanded time scale, it becomes obvious that PN transient K$^+$ currents activate and deactivate faster than KC transient K$^+$ currents (Fig 5A and 5B). To analyze the timing of the activation of the transient K$^+$ current, we determined its latency, which is significantly shorter in l-ALT PNs than in KCs (Fig 5C, Wilcoxon test l-ALT vs KC: p<0.05). As the transient K$^+$ currents are likely to be Na$^+$ or Ca$^{2+}$ dependent, we also analyzed the latency of the transient currents after pharmacological blocking (Fig 5D). The TTX-sensitive transient K$^+$ current was significantly faster than CdCl$_2$-sensitive transient K$^+$ currents (TTX vs CdCl$_2$: p<0.05).

In addition to the different timing between PN and KC transient K$^+$ currents, there are also differences in its maximal amplitude. Generally, PNs are larger than KCs (mean cell capacitance m-ALT: 17.27 pF, N = 4, l-ALT: 13.12 pF, N = 12; KC: 6.18 pF, N = 7). To adjust for the differences in their cell size, we calculated the current densities of PNs and KCs (Fig 5E). The transient Na$^+$/Ca$^{2+}$ currents of KCs are especially small (mean maximal current: 44pA) whereas the transient K$^+$ current is relatively large (mean maximal current: 1192pA) (Fig 5E).

### Discussion

We performed, for the first time, selective in situ whole-cell voltage clamp recordings from l- and m-ALT PNs and from KCs in the intact brain to compare their electrical properties. The differences between KCs and PNs were prominent: PNs possess strong transient Na$^+$ currents as well as Na$^+$ dependent transient K$^+$ currents and Ca$^{2+}$ dependent transient and persistent K$^+$ currents. KCs, in contrast, have a relatively small transient Na$^+$ current, a prominent transient K$^+$ current, and a most likely Ca$^{2+}$ dependent persistent N-shaped K$^+$ current.

### Electrophysiological properties of PNs

In our recordings, PNs fired action potentials starting at membrane voltages of around -40 mV without any spike frequency adaptation. PNs had cell capacitances of about 13–17 pF and a mean membrane resting potential of around -40 mV. The capacitances indicate that the cells measured in situ were slightly larger than cultured PNs that were reported around 11 pF [25]. This could be due to the fact that neurons were primary cultures from pupal stages. Although the PNs in primary cultures did grow dendritic processes, we assume that the difference at least partly is due to substantially larger dendritic and axonal arborizations in in situ measured PNs compared to cultured neurons. A resting membrane potential of -40 mV appears rather depolarized. We assume that the resting potential of PNs approximately ranges between -50 and -60 mV, which resembles the lowest measured resting potentials. This may be due to damage either during the preparation, staining or rupturing of the membrane patch. No significant differences between m- and l-ALT PNs regarding their capacitances or their resting membrane potentials were found.
PNs possess prominent Na\(^+\) currents, K\(_{\text{Na}}\), and K\(_{\text{Ca}}\) currents. The IV relations observed under standard conditions partly resemble the ones recorded in primary cell cultures [25,26]. In both in vitro studies and in our in-situ study, PNs exhibit a fast transient Na\(^+\) current, a fast activating and deactivating transient K\(^+\) current and a persistent K\(^+\) current. However, Grünwald [25] described a non-linear IV relation for the persistent K\(^+\) current leading to an N-shape of the IV plot for cultured PNs. Perk and Mercer [26] recorded two types of cultured AL neurons, and only one of the two types displayed IV relationships similar to our recordings. The other type did not display a fast transient K\(^+\) current, but instead had a non-linear N-shaped persistent K\(^+\) current. We conclude from our recordings that the first type measured by Perk and Mercer [26] was a PN type, whereas the second type may be a LN type. The only explanation for the differences between our recordings and the primary cell culture PN

Fig 5. Transient current analyses. A, B: Representative voltage clamp recordings from a lateral antennal-lobe tract projection neuron (l-ALT PN) (A) and a Kenyon cell (KC) (B). C: Delay of the transient K\(^+\) currents. The delay was significantly shorter in l-ALT PNs compared to KCs (p<0.05). D: Delay of l-ALT TTX and CdCl\(_2\) sensitive transient K\(^+\) currents (p<0.05). E: Mean current densities of l-ALT and KCs. Note the high current density of the transient K\(^+\) current and the low current densities of the Na\(^+\), Ca\(^{2+}\) currents in KCs.

https://doi.org/10.1371/journal.pone.0191425.g005
recordings from Grünewald [25] are differences between in situ (adult) and in vitro (pupal phase) ion channel expression.

The fast Na\(^+\) current resembles the typical Na\(^+\) current necessary for fast depolarization during action potential generation. This current was sensitive to TTX, which blocks a wide range of Na\(^+\) channels and prohibits action potential generation.

We observed a fast transient K\(_{\text{Na}}\) current that activated at -40 mV and increased until -10 mV to reach a maximal current plateau. Additionally, the plateau value was reached at the voltage level when the Na\(^+\) current declined. Concerning their timing, K\(_{\text{Na}}\) currents seem to be the fastest transient K\(^+\) currents in PNs. K\(_{\text{Na}}\) currents have been described in DUM neurons in cockroaches [33] and are thought to contribute to the limitation of action potential duration [34].

In our study, Ca\(^{2+}\) currents were measured after blocking Na\(^+\) currents with TTX. As K\(^+\) currents were not additionally blocked, the full magnitude of the Ca\(^{2+}\) currents could not be investigated in our studies. Nevertheless, the results strongly indicate that honeybee AL PNs possess a transient Ca\(^{2+}\) current, which appeared at -30 to -20 mV. To precisely determine activation thresholds and dynamics of Ca\(^{2+}\) currents, further experiments are necessary.

In addition to the K\(_{\text{Na}}\) current, we observed a transient and a persistent K\(_{\text{Ca}}\) current. Both K\(_{\text{Ca}}\) currents seem to be not purely Ca\(^{2+}\) dependent, but also voltage dependent. The activation membrane voltage, the existence of a transient and a persistent component, as well as the plateau reached by the transient current together with the linear IV relation of the persistent component are all properties of the pSlo subunit of the BK channel described in cockroach DUM neurons [35] and also cockroach PNs [36]. A similar current was observed in cultured AL neurons recorded by Perk and Mercer [26]. However, in cultured honeybee PN recordings from Grünewald [25] no such current was observed. Instead, the K\(_{\text{Ca}}\) current exhibited a descriptive N-shape in the IV plot [25], which is typical for SK K\(_{\text{Ca}}\) currents that are purely Ca\(^{2+}\) dependent and not voltage-dependent (reviewed by [37]). We did not observe an N-shape of the K\(^+\) current. Apart from the general difference to the in-situ situation and potential developmental aspects, we have no comprehensive explanation for these differences with recordings of cell cultured PNs by Grünewald [25]. To finally determine the nature of the K\(_{\text{Ca}}\) channels, specific pharmacological blocking with Apamin, a selective SK channel blocker, and Iberiotoxin, a selective BK channel blocker [38], is necessary. Transient K\(^+\) currents influence AP shape, the total AP duration [39], and the magnitude of the post AP hyperpolarization [35].

Based on our qualitative comparison with a limited number of m-ALT PNs, we assume that the ionic current properties do not strongly differ from l-ALT PNs (Fig 5). However, for a more comprehensive understanding of the ionic currents in both types of neurons, further pharmacological experiments are necessary. To be able to do this under in-situ conditions, further improvements of the preparation and recording techniques are necessary.

**Electrophysiological properties of KCs**

KCs had mean membrane resting potentials between -42 and -61 mV, and a mean capacitance of 6.18 pF. Similar as in PNs, we suppose that membrane potentials may be lower in unmanipulated neurons, and we expect membrane potentials of -60 to -65mV, which would be consistent with the mean potential (-62.1 mV) in in situ recordings of honeybee KCs in isolated brains [30]. The capacitance measured by Palmer et al. [30] amounted 3.2 pf. In current clamp, KCs showed spike frequency adaptation after pronounced depolarization, which was also observed by Palmer et al. [30].

Under standard conditions, KCs exhibited a fast inward current, which is likely to be a Na\(^+\) current. The outward currents can roughly be divided into a transient, A-shaped K\(^+\) current
and a persistent current with a non-linear IV relation leading to an N-shaped IV plot. The transient K$^+$ current activates at -45 mV, the Na$^+$ and the persistent K$^+$ current activate at ~-40 mV. The activation voltages of the currents as well as the basic shape of the IV plots are consistent with the IV plots obtained from KCs in cell culture [25,27,28] and recordings in isolated brain preparations [30]. The only prominent difference is that no N-shape was observed in cell cultured KCs [25]. The most prominent current observed in KC recordings was the transient K$^+$ current. This current activated at ~ -45 mV and increased linearly in a voltage dependent manner. In a single recording, it remained almost unaffected by blocking of Na$^+$ channels with TTX and blocking of Ca$^{2+}$ channels with CdCl$_2$. This points towards an A-type K$^+$ current. Additionally, its activation threshold and dynamics are similar to A-type K$^+$ currents in all studies on primary KC cultures [25,27,28].

The N-shape of the persistent K$^+$ current is normally caused by K$_{Ca}$ currents. The Ca$^{2+}$ influx after depolarization diminishes once the driving force for Ca$^{2+}$ ions is reduced. Therefore, the K$_{Ca}$ current will also decrease leading to an N-shaped IV plot. The typical shape was reported both in honeybee KC primary cultures [27] and in the isolated brain preparation [30] but was not found in another cell-culture patch clamp study. Similar K$_{Ca}$ currents were found in cockroaches, yet their amplitude was about twice as large as the amplitude of the A-type K$^+$ current [31].

Potential influence of ionic currents on odor coding

The subset of currents we found in PNs is relatively similar to currents observed in cockroach DUM neurons (reviewed by [34]). Especially K$_{Ca}$ currents closely resembling K$_{Ca}$ currents recorded in our study were shown to affect action potential waveform and excitability [36,39]. These currents allow PNs to be well suited as neurons with relatively high response probabilities and spontaneous AP frequencies producing reliable responses to synaptic input. On a qualitative level, we did not find any obvious differences between the currents of l- and m-ALT PNs. This may indicate that coding differences between PNs of the two tracts in terms of spontaneous activity and odor response frequencies producing reliable responses to synaptic input. On a qualitative level, we did not find any obvious differences between the currents of l- and m-ALT PNs. This may indicate that coding differences between PNs of the two tracts in terms of spontaneous activity and odor response frequencies were most likely caused by differences in synaptic connectivity between OSNs, LNs and PNs. In addition, differences in synaptic currents may contribute to coding differences. For example, OSNs of a specific type of olfactory sensilla (Sensillum basiconicum) preferentially project to m-ALT associated glomeruli [40] indicating that the OSN-PN connectivity varies between the two sets of PNs. Furthermore, differences in LN connectivity between the two AL hemilobes may as well affect odor coding in m- and l-ALT PNs [41].

KCs do have a completely different set of ionic currents than PNs. Instead of the very fast transient K$_{Ca}$ and K$_{Na}$, KCs have slower transient A-type and persistent N-shaped K$^+$ currents similar to cockroach KCs [31]. Generally, KCs are thought to play a role in coincidence detection and express synaptic plasticity [7,32,42–44]. For coincidence detection and spatial separation of information, neurons need to encode information in a spatially and temporally sparse fashion. Calcium-imaging studies support this temporal sparseness in KCs [23,24], even though responses in presynaptic PN axonal boutons outlasted the entire odor stimulus [23]. Such a drastic change in stimulus representation can be caused by strong transient K$^+$ currents in combination with K$_{Ca}$ currents or by GABAergic inhibition. Honeybee KCs do receive GABAergic input in the MB calyx [45,46], which could lead to self-inhibition after odor stimulation [47], yet Froese et al. [24] show that blocking of the fast ionotropic part of the GABA response did not affect the fast termination of odor responses in KCs. Therefore, it is more likely that ionic currents cause the change in stimulus representation, as shown in the cockroach [31]. Furthermore, SK channels in Manduca PNs are thought to mediate an after
hyperpolarization following pheromone responses [48,49]. Both the N-shaped $K_{Ca}$ currents in cockroaches (31) and the SK channels in Manduca PNs [48,49] are likely to shorten odor responses. Both the transient and the persistent $K^+$ currents we observed in honeybee KCs are relatively large compared to the inward currents and can therefore support the generation of temporally sparse, phasic responses. Whether the dominant transient $K^+$ currents or the persistent $K_{Ca}$ currents are more important for temporally sparse coding remains to be investigated. In addition to the temporal sparseness of stimulus representation, the number of excited KCs (spatial sparseness) responding to a given stimulus is also important. In Drosophila, blocking of GABAergic feedback neurons in the MBs resulted in activation of higher KC numbers (less spatial sparseness) [50]. Considering both morphological and physiological features of these neurons, we propose that feedback neurons inhibiting KCs most likely promote spatial sparseness in the representation of odor stimuli in the MBs, whereas the temporal sparseness may be caused by intrinsic KC neuronal properties.

Acknowledgments

We thank Robert Kittel for expert help with setting up the patch-clamp setup, Julie Carcaud for advice on the m-ALT preparation, and Dirk Ahrens for beekeeping.

Author Contributions

Conceptualization: Jan Kropf, Wolfgang Rössler.
Data curation: Jan Kropf.
Formal analysis: Jan Kropf.
Funding acquisition: Wolfgang Rössler.
Methodology: Jan Kropf.
Project administration: Wolfgang Rössler.
Supervision: Wolfgang Rössler.
Writing – original draft: Jan Kropf.
Writing – review & editing: Jan Kropf, Wolfgang Rössler.

References

1. Leinwand SG, Chalasani SH. Olfactory networks: From sensation to perception. Curr Opin Genet Dev. Elsevier Ltd; 2011; 21(6):806–11. https://doi.org/10.1016/j.gde.2011.07.006 PMID: 21889328
2. Galizia CG, Rössler W. Parallel olfactory systems in insects: anatomy and function. Annu Rev Entomol. 2010; 55:399–420. https://doi.org/10.1146/annurev-ento-112408-085442 PMID: 19737085
3. Kirschner S, Kleineidam CJ, Zube C, Rybak J, Grünewald B, Rössler W. Dual olfactory pathway in the honeybee, Apis mellifera. J Comp Neurol. 2006 Dec 20; 499(6):933–52. https://doi.org/10.1002/cne.21158 PMID: 17072827
4. Galizia CG, Menzel R. The role of glomeruli in the neural representation of odours: Results from optical recording studies. J Insect Physiol. 2001; 47(2):115–30. PMID: 11064019
5. Abel R, Rybak J, Menzel R. Structure and response patterns of olfactory interneurons in the honeybee, Apis mellifera. J Comp Neurol. 2001; 437(3):363–83. PMID: 11494262
6. Ito K, Shinomiya K, Ito M, Armstrong JD, Boyan G, Hartenstein V, Harzsch S, Heisenberg M, Homberg U, Jenett A, Keshishian H, Restiflo LL, Rössler W, Simpson JH, Strausfeld NJ, Strauss R, Vosshall LB. A systematic nomenclature for the insect brain. Neuron. 2014; 81(4):755–65. https://doi.org/10.1016/j.neuron.2013.12.017 PMID: 24559571
7. Rössler W, Brill MF. Parallel processing in the honeybee olfactory pathway: Structure, function, and evolution. J Comp Physiol A Neuroethol Sensory, Neural, Behav Physiol. 2013; 199(11):981–96.
8. Homberg U, Christensen TA, Hildebrand JG. Structure and Function of the Deutocerebrum in Insects. Annu Rev Entomol. 1989; 34(1):477–501.

9. Groh C, Tautz J, Rössler W. Synaptic organization in the adult honey bee brain is influenced by brood-temperature control during pupal development. Proc Natl Acad Sci U S A. 2004; 101(12):4268–73. https://doi.org/10.1073/pnas.0400773101 PMID: 15024125

10. Tanaka NK, Suzuki E, Dye L, Ejima A, Stopfer M. Dye fills reveal additional olfactory tracts in the protocerebrum of wild-type Drosophila. J Comp Neurol. 2012; 520(18):4131–40. https://doi.org/10.1002/cne.23149 PMID: 22592823

11. Homberg U, Montague RA, Hildebrand JG. Anatomy of antennocerebral pathways in the brain of the sphinx moth Manduca sexta. Cell Tissue Res. 1988; 254(2):255–81. PMID: 3197087

12. Rössler W, Zube C. Dual olfactory pathway in Hymenoptera: Evolutionary insights from comparative studies. Arthropod Struct Dev. Elsevier Ltd; 2011; 40(4):349–57. https://doi.org/10.1016/j.asd.2010.12.001 PMID: 21167312

13. Müller D, Abel R, Brandt R, Zöckler M, Menzel R. Differential parallel processing of olfactory information in the honeybee, Apis mellifera L. J Comp Physiol A Neuroethol Sensor y, Neural, Behav Physiol. 2002; 188(5):359–70.

14. Krofczik S, Menzel R, Nawrot MP. Rapid odor processing in the honeybee antennal lobe network. Front Comput Neurosci. 2008; 2:9. https://doi.org/10.3389/neuro.10.009.2008 PMID: 19221584

15. Carcaud J, Hill T, Giurfa M, Sandoz J-C. Differential coding by two olfactory subsystems in the honeybee brain. J Neurophysiol. 2012; 108(4):1106–21. https://doi.org/10.1152/jn.01034.2011 PMID: 22572948

16. Galizia CG, Franke T, Menzel R, Sandoz J-C. Optical imaging of concealed brain activity using a gold mirror in honeybees. J Insect Physiol. Elsevier Ltd; 2012; 58(5):743–9. https://doi.org/10.1016/j.jinsphys.2012.02.010 PMID: 22414536

17. Brill MF, Rosenbaum T, Reus I, Kleineidam CJ, Nawrot MP, Rössler W. Parallel processing via a dual olfactory pathway in the honeybee. J Neurosci. 2013; 33(6):2443–56. https://doi.org/10.1523/JNEUROSCI.4268-12.2013 PMID: 23392673

18. Carcaud J, Giurfa M, Sandoz J-C. Differential Combinatorial Coding of Pheromones in Two Olfactory Subsystems of the Honey Bee Brain. J Neurosci. 2015; 35(10):4157–67. https://doi.org/10.1523/JNEUROSCI.0734-14.2015 PMID: 25762663

19. Esslen J, Kaissling KE. Zahl und Verteilung antennaler Sensillen bei der Honigbiene (Apis mellifera L.). Zoomorphologie. 1976; 83(3):227–51.

20. Rybak J. The digital bee brain: integrating and managing neurons in a common 3D reference system. Front Syst Neurosci. 2010; 4:15.

21. Wittthöft W. Absolute Anzahl und Verteilung der Zellen im Hirn der Honigbiene. Zeitschrift für Morphol der Tiere. 1967; 61(1):160–84.

22. Strausfeld NJ. Organization of the honey bee mushroom body: Representation of the calyx within the vertical and gamma lobes. J Comp Neurol. 2002; 450(1):4–33. https://doi.org/10.1002/cne.10285 PMID: 12124764

23. Szyszka P, Ditzen M, Galkin A, Galizia CG, Menzel R. Spursening and temporal sharpening of olfactory representations in the honeybee mushroom bodies. J Neurophysiol. 2005; 94(5):3303–13. https://doi.org/10.1152/jn.00397.2005 PMID: 16014792

24. Fros e A, Szyszka P, Menzel R. Effect of GABAergic inhibition on odorant concentration coding in mushroom body intrinsic neurons of the honeybee. J Comp Physiol A Neuroethol Sensory, Neural, Behav Physiol. 2014; 200(3):183–95.

25. Grünewald B. Differential expression of voltage-sensitive K+ and Ca2+ currents in neurons of the honeybee olfactory pathway. J Exp Biol. 2003; 206(1):117–29.

26. Perk CG, Mercer AR. Dopamine modulation of honey bee (Apis mellifera) antennal-lobe neurons. J Neurophysiol. 2006; 95(2):1147–57. https://doi.org/10.1152/jn.01220.2004 PMID: 16262199

27. Schäfer S, Rosenboom H, Menzel R. Ionic currents of Kenyon cells from the mushroom body of the honeybee. J Neurosci. 1994; 14(8):4600–12. PMID: 7519255

28. Pelz C, Jander J, Rosenboom H, Hammer M, Menzel R. IA in Kenyon cells of the mushroom body of honeybees resembles shaker currents: kinetics, modulation by K+, and simulation. J Neurophysiol. 1999; 81(4):1749–59. https://doi.org/10.1152/jn.1999.81.4.1749 PMID: 10200210

29. Wüstenberg DG, Boycheva M, Grünewald B, Byrne JH, Menzel R, Baxter DA. Current- and voltage-clamp recordings and computer simulations of Kenyon cells in the honeybee. J Neurophysiol. 2004; 92(4):2589–603. https://doi.org/10.1152/jn.01259.2003 PMID: 15190098
30. Palmer MJ, Moffat C, Saranzewa N, Harvey J, Wright GA, Connolly CN. Cholinergic pesticides cause mushroom body neuronal inactivation in honeybees. Nat Commun. Nature Publishing Group; 2013; 4:1634. https://doi.org/10.1038/ncomms2648 PMID: 23535655

31. Demmer H, Kloppenburg P. Intrinsic membrane properties and inhibitory synaptic input of Kenyon cells as mechanisms for sparse coding? J Neurophysiol. 2009; 102(3):1538–50. https://doi.org/10.1152/jn.00183.2009 PMID: 19553491

32. Brill MF, Meyer A, Rössler W. It takes two—coincidence coding within the dual olfactory pathway of the honeybee. Front Physiol. 2015; 6:1–14.

33. Grolleau F, Lapied B. Transient Na(+)-activated K+ current in beating pacemaker-isolated adult insect neurosecretory cells (dum neurones). Neurosci Lett. 1994; 167(1–2):46–50. PMID: 8177528

34. Derst C, Messutat S, Walther C, Eckert M, Heinemann SH, Wicher D. The large conductance Ca2+-activated potassium channel (pSlo) of the cockroach Periplaneta americana: Structure, localization in neurons and electrophysiology. Eur J Neurosci. 2003; 17(6):1197–212. PMID: 12670308

35. Rhadler C, Warren B, Bardos V, Schleicher S, Klein A, Kloppenburg P. Properties and physiological function of Ca 2+ -dependent K+ currents in uniglomerular olfactory projection neurons. J Neurophysiol. 2016; 115(5):2330–40. https://doi.org/10.1152/jn.00840.2015 PMID: 26823514

36. Adelman JP, Maylie J, Sah P. Small-Co nductance Ca 2+ -Activated K + Channels: Form and Function. Annu Rev Neurosci. 2012; 35(1):245–69.

37. Derst C, Messutat S, Walther C, Eckert M, Heinemann SH, Wicher D. The large conductance Ca2+ -activated potassium channel (pSlo) of the cockroach Periplaneta americana: Structure, localization in neurons and electrophysiology. Eur J Neurosci. 2003; 17(6):1197–212. PMID: 12670308

38. Bradler C, Warren B, Bardos V, Schleicher S, Klein A, Kloppenburg P. Properties and physiological function of Ca 2+ -dependent K+ currents in uniglomerular olfactory projection neurons. J Neurophysiol. 2016; 115(5):2330–40. https://doi.org/10.1152/jn.00840.2015 PMID: 26823514

39. Adelman JP, Maylie J, Sah P. Small-Co nductance Ca 2+ -Activated K + Channels: Form and Function. Annu Rev Neurosci. 2012; 35(1):245–69.

40. Meyer A, Galizia CG. Elemental and configural olfactory coding by antennal lobe neurons of the honeybee (Apis mellifera). J Comp Physiol A Neuroethol Sensory, Neural, Behav Physiol. 2012; 198(2):159–71.

41. Perez-Orive J, Bazhenov M, Laurent G. Intrinsic and circuit properties favor coincidence detection for decoding oscillatory input. J Neurosci. 2004; 24(26):6037–47. https://doi.org/10.1523/JNEUROSCI.1084-04.2004 PMID: 15229251

42. Gyula N, Stopfer M. Insect olfactory coding and memory at multiple timescales. Curr Opin Neurobiol. Elsevier Ltd; 2011; 21(5):768–73. https://doi.org/10.1016/j.conb.2011.05.005 PMID: 21632235

43. Rybak J, Menzel R. Anatomy of the mushroom bodies in the honey bee brain: the neuronal connections of the alpha-lobe. J Comp Neurol. 1993; 334(3):444–65. https://doi.org/10.1002/cne.903340309 PMID: 8376627

44. Grünewald B. Morphology of feedback neurons in the mushroom body of the honeybee, Apis mellifera. J Comp Neurol. 1999; 404(1):114–26. PMID: 9886029

45. Palmer MJ, Harvey J. Honeybee Kenyon cells are regulated by a tonic GABA receptor conductance. J Neurophysiol. 2014; 44(0):2026–35.

46. Lei H, Riffell JA, Gage SL, Hildebrand JG. Contrast enhancement of stimulus intermittency in a primary olfactory network and its behavioral significance. J Biol. 2009; 8(2):21. https://doi.org/10.1186/jbiol120 PMID: 19232128

47. Lei H, Yu Y, Zhu S, Rangan A V, Anton S. Intrinsic and Network Mechanisms Constrain Neural Synchrony in the Moth Antennal Lobe. Front Physiol. 2016; 8:7 1–24.

48. Lin AC, Bygrave AM, de Calignon A, Lee T, Miesenböck G. Sparse, decorrelated odor coding in the mushroom body enhances learned odor discrimination. Nat Neurosci. 2014; 17(4):559–68. https://doi.org/10.1038/nn.3660 PMID: 24561998