

Spontaneous activity in the piriform cortex extends the dynamic range of cortical odor coding

Malinda L. S. Tantirigama^a, Helena H.-Y. Huang^a, and John M. Bekkers^{a,1}

^aEccles Institute of Neuroscience, John Curtin School of Medical Research, The Australian National University, Canberra, ACT 2601, Australia

Edited by Charles F. Stevens, The Salk Institute for Biological Studies, La Jolla, CA, and approved January 26, 2017 (received for review December 20, 2016)

Neurons in the neocortex exhibit spontaneous spiking activity in the absence of external stimuli, but the origin and functions of this activity remain uncertain. Here, we show that spontaneous spiking is also prominent in a sensory paleocortex, the primary olfactory (piriform) cortex of mice. In the absence of applied odors, piriform neurons exhibit spontaneous firing at mean rates that vary systematically among neuronal classes. This activity requires the participation of NMDA receptors and is entirely driven by bottom-up spontaneous input from the olfactory bulb. Odor stimulation produces two types of spatially dispersed, odor-distinctive patterns of responses in piriform cortex layer 2 principal cells: Approximately 15% of cells are excited by odor, and another approximately 15% have their spontaneous activity suppressed. Our results show that, by allowing odor-evoked suppression as well as excitation, the responsiveness of piriform neurons is at least twofold less sparse than currently believed. Hence, by enabling bidirectional changes in spiking around an elevated baseline, spontaneous activity in the piriform cortex extends the dynamic range of odor representation and enriches the coding space for the representation of complex olfactory stimuli.

olfaction | in vivo | calcium imaging | two-photon | anesthetic

The brain remains intensely active even under conditions when overt external stimuli are absent (1). At first glance this behavior can seem puzzling. Pyramidal neurons in primary sensory neocortex, for example, typically fire spontaneously at approximately 1 Hz when the relevant stimuli are not present (2), raising the possibility that this background activity is noise that must be mitigated by signal averaging (3). On the other hand, spontaneous cortical activity is often structured (4) and can reflect the global state of the animal (5), suggesting that it is not simply an impediment but may serve a useful computational role. Indeed, such activity may be important for gating sensory inputs (4, 6), establishing and maintaining sensory maps (7–9), and extending the dynamic range of sensory-evoked responses (10, 11). The significance of spontaneous activity is also supported by theoretical studies showing that it emerges naturally from balanced neural networks with desirable information-processing characteristics (9, 12).

Given that spontaneous spiking is likely to be a key element of cortical function, it is important to study its prevalence and properties in different cortical areas. The piriform cortex (PC) is a sensory paleocortex in mammals that is critical for recognizing and remembering odors (13-17). The PC offers a number of advantages for the study of neural information processing, including a simple trilaminar architecture and well-defined afferent input from the olfactory bulb (OB) (16, 17). As a phylogenetically "old" structure, the PC is also likely to express canonical cortical circuits that have been conserved across evolution (18). Spontaneous spiking has been reported in the PC of rodents breathing odor-free air (19-23), but these studies used unit recordings, which provide a biased estimate of firing rates (2), and none of them identified the drivers or roles of this activity. Here we use two-photon calcium imaging and whole-cell patch clamping from identified PC neurons in vivo to measure the properties of stimulus-decoupled spiking. We find that this ongoing spiking allows a broader dynamic range of odor responsiveness, implying that the representation of odor

information in the PC is considerably less sparse than is currently thought (20, 24).

Results

Spontaneous Spiking Occurs at Different Rates in Different Classes of Piriform Neurons. Using mice, we imaged somatic intracellular calcium (as a proxy for spiking) in layer 2 neurons of the anterior PC while the animals breathed clean air. Layer 2 of the PC contains the densely packed somas of glutamatergic cells of different types (Fig. S1A). The more superficial lamina (layer 2a) is enriched in semilunar (SL) cells, whereas the deeper lamina (layer 2b) mainly contains superficial pyramidal (SP) cells (Fig. S1A). Because SL and SP cells are likely to perform different functions in the PC circuit (Fig. S1B) (25, 26), we took care to study them separately by focusing on cells at different depths below the cortical surface (160–210 μ m for SL cells and 250–300 μ m for SP cells).

Using the indicator Cal-520, we found that both SL and SP cells exhibited ongoing spontaneous somatic calcium transients in the absence of applied odors (Fig. 1*A* and Movie S1), and we obtained similar results with the genetically encoded calcium indicator GCaMP6f. Control experiments showed that these transients were completely blocked by tetrodotoxin (TTX; 10 μ M) applied to the cortical surface (n=3 animals), indicating that they are sodium spike-driven. The transients occurred randomly, as was confirmed by the exponential distribution of interevent time intervals (linear fit to the logarithm of interval histogram; SL: $r^2 = 0.98$, n=5 mice, 5,609 events in 725 cells; SP: $r^2 = 0.98$, n=5 mice, 1,408 events in 275 cells).

To convert the calcium transient rate to an action potential rate, we calibrated one of our indicators (Cal-520) by measuring spontaneous calcium fluorescence transients in a single SL or SP cell in vivo while simultaneously recording spikes in that cell via a patch electrode (Fig. S2). Applying this calibration to our imaging

Significance

Sensory neurons are often spontaneously active even in the absence of the relevant stimuli, raising the question, Is this background activity a bug or a feature of neural circuits? Using in vivo calcium imaging in mice, we show that neurons in the primary olfactory (piriform) cortex are spontaneously active when the animal is breathing unodorized air and this activity is entirely driven by spontaneous firing in the upstream olfactory bulb. When the animal smells an odor, some piriform neurons fire more strongly, and others have their spontaneous activity suppressed. By allowing two-way changes in firing, spontaneous background activity serves as a mechanism to extend the range of odor coding.

Author contributions: M.L.S.T. and J.M.B. designed research; M.L.S.T. and H.H.-Y.H. performed research; M.L.S.T. and J.M.B. analyzed data; and M.L.S.T. and J.M.B. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission

¹To whom correspondence should be addressed. Email: John.Bekkers@anu.edu.au.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10. 1073/pnas.1620939114/-/DCSupplemental.

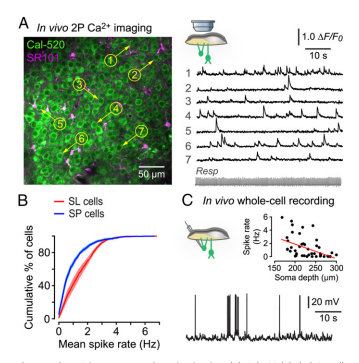


Fig. 1. The PC is spontaneously active in vivo. (A) Cal-520–labeled SL cell somas (Left) and corresponding $\Delta F/F_0$ fluorescence traces from the numbered cells (Right) showing spontaneous activity. Resp, simultaneously recorded respiration. SR 101 labels astrocytes. (B) Averaged cumulative histograms showing that the mean spontaneous spike rate in SL cells (n=7 mice, 1,928 cells) and SP cells (n=6 mice, 1,005 cells; shaded regions are \pm SEM) are significantly different (P<0.0001, Kolmogorov–Smirnov test). (C) A typical in vivo whole-cell recording showing spontaneous spiking in an SL cell (depth $166~\mu m$). (Inset) The spontaneous spike rate was higher in the more superficial SL cells (soma depth $160-230~\mu m$, 1.7 ± 0.3 Hz; mean \pm SEM) than in the deeper SP cells ($230-300~\mu m$, 0.7 ± 0.2 Hz; mean \pm SEM, P=0.005, Mann–Whitney test) and was negatively correlated with recording depth (r=0.47, P=0.0006, n=50 cells from 48 mice; F test on linear regression).

dataset, we found that the mean spontaneous spike rate was significantly higher in SL cells than in SP cells (SL: 1.24 ± 0.04 Hz, mean \pm SEM, n=7 mice, total 1,928 cells; SP: 0.74 ± 0.05 Hz, n=6 mice, 1,005 cells; significantly different, P < 0.0001, Kolmogorov–Smirnov test) (Fig. 1B).

To confirm that the majority of cells imaged were indeed glutamatergic, we repeated these experiments using the red-shifted calcium indicator Cal-590 in GAD67-GFP mice, in which GABAergic interneurons express GFP (Fig. S3) (27). We confirmed that interneurons comprise a small fraction (approximately 4%) of layer 2 neurons in the PC, consistent with previous reports (28). Thus, most of the neurons imaged in layer 2 are glutamatergic (SL and SP cells). Interestingly, GFP⁺ interneurons also exhibited spontaneous spiking activity (Fig. S3 B and C).

In Vivo Patch-Clamp Experiments Confirm the Calcium Imaging Results. To validate the findings from the calcium imaging, we turned to in vivo whole-cell patch clamping. Spontaneous spiking was measured at the resting potential of each neuron while the animal breathed odor-free air (Fig. 1C). These experiments confirmed that the spontaneous spike rate was higher in the more superficial SL cells (soma depth 160– $230~\mu m$, $1.7 \pm 0.3~Hz$, mean \pm SEM, n = 28 cells) than in the deeper SP cells (230– $300~\mu m$, $0.7 \pm 0.2~Hz$, mean \pm SEM, n = 22~cells; P = 0.005, Mann–Whitney test). In addition, the spontaneous spike rate was negatively correlated with recording depth (P = 0.0006, n = 50~cells from 48~mice; F test on linear regression) (Fig. 1C, Inset). Thus, both patch clamping and imaging

show that SL and SP cells are spontaneously active, with a higher spontaneous firing rate in SL cells than in SP cells.

Spontaneous Spiking Requires NMDA Receptors. Because the difficult surgical access to the PC precludes the use of awake animals, we sought to validate our findings under different anesthetics. Spontaneous spiking was similar under urethane and fentanyl/ medetomidine anesthesia [SL cells in fentanyl: 1.13 ± 0.05 Hz, mean \pm SEM, n=2 mice, 612 cells; not significantly different from corresponding control (urethane) values, P > 0.1, one-way ANOVA with Tukey's post hoc test; only SL cells were imaged in fentanyl] (Fig. 24). Fentanyl/medetomidine is thought to produce a more awake-like brain state (29). Surprisingly, spiking was significantly reduced under ketamine/xylazine anesthesia (combined SL and SP data: 0.26 ± 0.04 Hz, mean \pm SEM, n = 2 mice, total 1,328 cells; significantly smaller than corresponding control values, P < 0.0001, one-way ANOVA with Tukey's post hoc test) (Fig. 24). This result may help explain why spontaneous activity was not seen in the only previous in vivo calcium imaging study in the PC, which used ketamine/xylazine (24).

Given that ketamine blocks NMDA channels, we asked whether local application of the NMDA channel blocker MK-801 (1 mM) to the PC would also suppress spontaneous spiking. We found that it did so (SL: 0.05 ± 0.01 Hz, mean \pm SEM, n=3 mice, 1,728 cells; SP: 0.14 ± 0.01 Hz, n=2 mice, 352 cells; both significantly smaller than corresponding controls, P < 0.0001, one-way ANOVA with Tukey's post hoc test) (Fig. 24). Thus, spontaneous spiking in the PC depends critically on NMDA receptor-mediated synaptic transmission.

Spontaneous Activity in the PC Is Driven Entirely by the OB. What drives spontaneous spiking in the PC? Mechanosensitive effects of nasal airflow across the olfactory epithelium (30) are not involved, because tracheotomizing mice to bypass the nasal airway did not alter the activity (SL: 1.10 ± 0.06 Hz, mean \pm SEM, n = 3mice, 1,659 cells; SP: 0.77 ± 0.06 Hz, n = 3 mice, 406 cells; not significantly different from corresponding controls, P > 0.1, oneway ANOVA with Tukey's post hoc test) (Fig. 2A). However, silencing the afferent input from the ipsilateral OB, either surgically or by locally applying TTX (10 µM) to the OB, reduced spontaneous spiking in the PC (combined SL and SP data: $0.25 \pm$ 0.05 Hz, mean \pm SEM, n = 5 mice, 2,453 cells; significantly smaller than control, P < 0.0001, one-way ANOVA with Tukey's post hoc test in Fig. 2A, Kolmogorov–Smirnov test in Fig. 2B). Removal of both OBs abolished all activity in the PC (combined SL and SP data: 0.01 ± 0.00 Hz, mean \pm SEM, n = 3 mice, 2,162 cells; significantly smaller than control, P < 0.0001, one-way ANOVA with Tukey's post hoc test) (Fig. 24). These data indicate that input from the OB is required for spontaneous spiking activity in the PC.

Our findings suggest that the projection cells in the OB, the mitral/tufted (M/T) cells, are spontaneously active in the absence of applied odors. Imaging in the bulb confirmed this suggestion (Fig. S4) (31). Interestingly, local application of TTX to the PC had no effect on spontaneous spiking in M/T cells (n = 93 M/T cells, P = 0.11, Kolmogorov–Smirnov test) (Fig. S4 B and C). This result suggests that only OB \rightarrow PC connections are important for spontaneous spiking in the PC and that connections in the reverse direction (PC \rightarrow OB) are not involved (32–34).

Odors both Excite and Suppress Spontaneous Activity. How does this spontaneous spiking activity influence the way exogenous odors are detected by the PC? To address this question, we used two-photon calcium imaging to record the effect of prolonged (60-s) steps of odorant on SL and SP cell activity. The responses were heterogeneous: Some cells showed transient excitation at odor onset, others were persistently activated throughout the odor application, and still others were suppressed (Fig. 34 using Cal-520

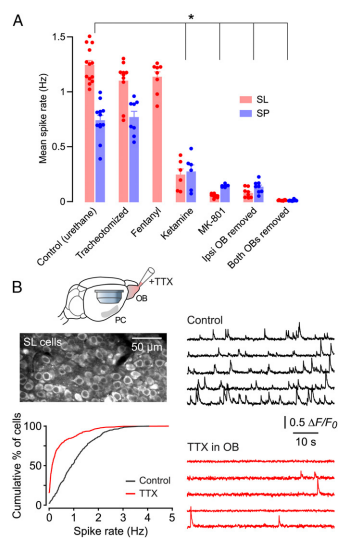


Fig. 2. Properties of spontaneous spiking in SL and SP cells under different experimental conditions. (A) Each data point is the averaged firing rate for all cells (n approximately 60–230 cells) in a field of view in a single experiment. Bars show mean \pm SEM of the points. *P < 0.05 (one-way ANOVA with Tukey's post hoc test, comparing control with other conditions in SL cells and SP cells separately). SP cells were not imaged in fentanyl. (B, $Upper\ Left$) Imaged field of SL cells in this experiment (using Cal-520). (Right) ΔFIF_0 traces for the same five SL cells recorded before (control, Upper) and during (Lower) the superfusion of TTX over the ipsilateral OB. ($Lower\ Left$) Cumulative histograms of spontaneous spike rates recorded in this experiment before (black trace) and during (red trace) TTX application. P < 0.0001 (n = 368 cells, Kolmogorov–Smirnov test).

and Movie S2 using GCaMP6f). The variety of responses in a typical experiment is shown in Fig. 3B, which is a color raster plot of activity in a sample of SL cells ordered by their activity in the first 4 s after odor onset (filled circle in Fig. 3B). A gradient of responsiveness is apparent, ranging from excitation (warm colors just after odor onset; top) to inhibition (cool colors just after onset; bottom), with a subset of cells showing no response (middle). Similar distinctions were seen at odor offset (open circle in Fig. 3B).

These data were quantified by using a statistical criterion to measure the percentage of cells either excited or suppressed at odor onset or offset (Fig. S5). This analysis indicated that a similar percentage of SL and SP cells was responsive to odor onset (SL: $34.1 \pm 4.5\%$ of cells, mean \pm SEM, n = 10 imaging fields tested with six odors; SP: $23.3 \pm 2.3\%$, n = 8 fields and five odors; P = 0.063;

two-sample t test) (Fig. 3C). In contrast, fewer SP cells responded at odor offset (SL: 35.6 \pm 3.0%, mean \pm SEM; SP: 16.1 \pm 1.6%; P < 0.0001; two-sample t test) (Fig. 3C). Similar results were obtained if excitation and suppression were considered separately (Fig. S6A) and if the odor concentration was varied over two orders of magnitude (Fig. S6B and C). Furthermore, we found no significant differences between any of these measures when comparing responses to different odors (one-way ANOVA, all P > 0.05). In summary, spontaneous activity allows odor-driven bidirectional changes in spiking around an elevated baseline. Although there are some differences between SL and SP cells, the percentage of responsive cells is independent of odor identity and concentration.

Balanced Excitation and Suppression. Given that odors both excite and suppress firing in PC neurons, we wondered if the net effect was to keep firing in equilibrium. Indeed, when averaged across all SL cells or all SP cells in our dataset, the mean spike rate during the odor was not significantly different from that in unodorized air, except at odor onset and offset (Fig. S7). Thus, following a perturbation of the olfactory stimulus, excitation and suppression are quickly balanced to keep the net population activity constant (35).

Relationship Between Spontaneous Firing Rate and Odor Responsiveness.

We next asked whether cells with a higher spontaneous firing rate also exhibited a stronger odor response, given that both types of activity are driven by bulbar input. We tested this notion by plotting the peak odor-evoked $\Delta F/F_0$ in each odor-responsive cell versus the mean spontaneous rate for that cell (Fig. 4A). SP cells exhibited a significant correlation, but SL cells did not (SL: P=0.20, n=321 cells; SP: P<0.001, n=83 cells; F test on linear regression). Thus, in SP cells the input that drives odor responsiveness is closely related to the input that drives spontaneous activity, but this association is absent in SL cells.

Excited and Suppressed Neurons Form Spatially Dispersed, Odor-Distinctive Patterns. As noted above (Fig. 3C and Fig. S6), different odors could not be distinguished by the mean responsiveness of SL or SP cells (i.e., the fraction of cells excited or suppressed). On the other hand, it is known that different odors produce distinctive, spatially dispersed patterns of excited principal neurons in the PC (red somas, Fig. 4B) (20, 24); these patterns are thought to form the basis of odor identification by the PC (15–17). We found a similar effect for suppression: Odors also produced distinctive, spatially dispersed patterns of suppression of SL and SP cells at both odor onset and offset (blue somas in Fig. 4B; similar results were obtained in 10 experiments). We quantified this effect by calculating, separately for excitation and suppression, the mean distance between each odor-responsive neuron in the imaged field and its five nearest odor-responsive neighbors. These mean distances then were averaged across all odor-excited or odor-suppressed neurons in each imaging experiment (red and blue points, respectively, in Fig. 4C). The mean distance between odorexcited and odor-suppressed neurons was not significantly different across all experiments (SL: $60.3 \pm 5.7 \mu m$ for excitation, $63.7 \pm$ 7.7 μ m for suppression, mean \pm SEM, both n=10 experiments, P=0.73; SP: 116.1 \pm 15.2 μ m for excitation, n = 8, 98.5 \pm 24.3 μ m for suppression, n = 5, P = 0.53; two-sample t test) (Fig. 4C). Similar results were obtained when we measured the mean distance to all (not just to the nearest five) odor-responsive neighbors in the imaged field. We also asked if excited cells tended to be located near suppressed cells, or if the two types of cells were randomly intermingled in a salt-and-pepper arrangement. An analysis similar to that described above confirmed the salt-and-pepper model (Fig. S8). Thus, odors both excite and suppress layer 2 principal cells in a manner that lacks any obvious spatial pattern.

Discussion

Spontaneous firing in the absence of overt input is commonly observed in sensory neocortex, as well as in many other brain

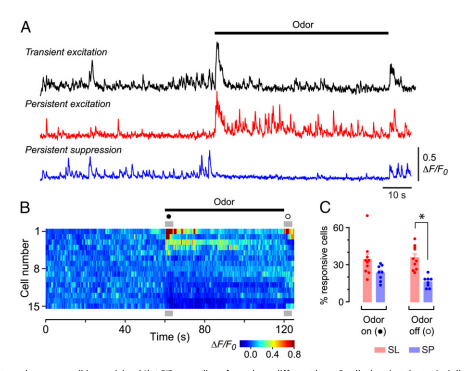


Fig. 3. Odors both excite and suppress spiking activity. (A) $\Delta F/F_0$ recordings from three different layer 2 cells showing the typical diversity of odor responses. The indicator was Cal-520. (B) Color raster plot of $\Delta F/F_0$ in a selected subset of SL cells in one experiment, ranked by their peak $\Delta F/F_0$ during the first 4 s after odor application. Gray bars designate 4 s after odor onset (\blacksquare) and offset (O). (C) Percentages of SL and SP cells that were responsive (i.e., were either excited or suppressed) during odor onset or offset. Bars show mean \pm SEM of the individual experiments (points). *P < 0.001 (n = 10 SL and n = 8 SP experiments, unpaired two-tailed t test).

regions, and its prevalence may suggest that it is a feature, not a bug, of neural circuits (2–4). Here we confirm that spontaneous firing is also a feature of a sensory paleocortex, the PC. We find that spontaneous activity in the anterior PC strictly requires bottom-up input from the OB (Fig. 24) and differs significantly between classes of principal cells (i.e., SL and SP cells) in its main input layer, layer 2 (Figs. 1B, 2A, 3C, and 4A). Importantly, we also find that spontaneous spiking activity increases the dynamic range of odor representations in the anterior PC by defining a set point around which odor stimulation can produce either excitation or suppression of spiking activity (Fig. 3B). By expanding the definition of odor responsiveness in this way, we find that at least twice as many layer 2 neurons respond to odors than previously thought; i.e., approximately 25–35% respond at odor onset (Fig. 3C), rather than the approximately 3–15% reported previously (24).

Before proceeding, we address two potential weaknesses of our study. First, the use of anesthesia can be problematic (36), but it was required in our experiments because of the invasive surgery needed to access the surface of the PC (24). To address this issue, we confirmed that two different anesthetics (urethane/chlorprothixene and fentanyl/medetomidine) gave identical results. Previous work using unit recordings in awake mice has reported spontaneous firing in layer 2 PC neurons at approximately 0.6–3.5 Hz (21), compared with approximately 0.7–1.2 Hz here (Fig. 24). Keeping in mind that unit recordings are biased toward more active cells, we believe that our results are in line with what occurs in awake animals. Given our finding that ketamine/xylazine inhibits spontaneous activity (Fig. 24), we recommend avoiding this anesthetic for PC experiments (see also ref. 34).

A second potential concern is the use of calcium imaging as an imperfect proxy for spikes. We mitigated this concern by using the calcium indicators Cal-520 and GCaMP6f, which are at least 10-fold more sensitive than Oregon Green BAPTA-1 (OGB-1) (37) used in the one previous calcium imaging study in the PC (24). We confirmed that OGB-1 could not detect spontaneous activity in

the PC (n=7 imaging fields in two mice). Despite our improved sensitivity, single spikes were not reliably detected in our imaging experiments (Fig. S2), implying that the firing rate is underestimated. The patch-clamp experiments (Fig. 1C) were not subject to this limitation and confirmed the difference in the spontaneous firing rate in SL and SP cells (Fig. 1B). Hence, imaging may underestimate absolute spike rates, but relative comparisons remain valid.

The higher spontaneous firing rate in SL cells is consistent with previous reports that SL cells receive stronger afferent input from the OB (25, 26), which we have shown here is the primary driver of spontaneous activity. SP cells receive much stronger associational input than SL cells, and some of this input comes from the contralateral PC (25). However, removal of the ipsilateral OB almost completely abolished spontaneous activity (Fig. 2A), implying that little spontaneous input comes via associational fibers from the contralateral PC. Another difference between SL and SP cells is that only SP cells show a correlation between spontaneous and odor-evoked activity (Fig. 4A). This correlation may reflect the stronger associational connections received by SP cells that might amplify, in parallel, the spontaneous and odor-evoked excitation received from the OB. A functional consequence may be that SP cells, but not SL cells, maintain a stable signal-to-noise ratio over a range of inputs (38).

We were surprised to find that spontaneous activity in the PC was driven entirely by feed-forward excitation from the OB (Fig. 2A, both OBs removed). The anterior PC is the recipient of excitatory input from a number of other brain areas, including the anterior olfactory nucleus, orbitofrontal cortex, entorhinal cortex, and amygdala (39, 40), but we found no sign of residual spontaneous drive after OB removal. However, it remains possible that intracortical and descending inputs could contribute to spontaneous activity in the awake state. The anterior PC also sends corticobulbar axons back to the OB (32, 33, 41), but, again, we found no evidence for the contribution of a reciprocal OB \leftrightarrow PC



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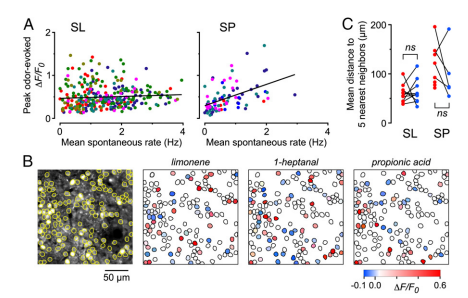


Fig. 4. Properties of odor-evoked excitation and suppression. (A) Each point shows the peak odor-evoked response in an odor-excited neuron plotted versus the mean spontaneous spike rate measured in the same neuron. Different colors indicate different odors. Superimposed straight lines indicate a correlation for SP cells (slope = 0.25, correlation coefficient = 0.48, n = 83 cells) but not for SL cells (slope = 0.02, correlation coefficient = 0.07, n = 321 cells). (B, Left) Imaged field of SL cells with somatic ROIs outlined. The other three panels show the same ROIs with color-coded odor responses to the three odors indicated. The indicator was GCaMP6f. (C) Each red point indicates the mean distance between each excited cell in an imaged field and its five nearest neighbors that also were excited, averaged across all excited cells in that field. Blue points show the same measure for suppressed cells. Lines connect data from the same experiment (n = 10 experiments for SL cells, n = 8 experiments for SP cells). In three SP cell experiments (red points without connected blue points) there were fewer than six suppressed cells in the imaged field. ns, not significant (P > 0.5, unpaired two-tailed t test).

loop in the generation of spontaneous excitation in the PC (Fig. S4). Two recent reports have described spontaneous calcium transients in boutons on corticobulbar axons in the OB (33, 34). Their presence implies that the sources of those axons, SP cells in the anterior PC (42), are also spontaneously active, consistent with our findings.

Turning to odor-evoked responses, we observed a heterogeneity of response patterns (Fig. 3 and Fig. S5). However, when averaging across all imaged neurons in all experiments, the mean spike rate increased only at odor onset and offset, suggesting that global excitation and suppression are normally balanced in an unchanging olfactory environment (Fig. S7). These properties are suited to a mechanism enabling animals to follow odor plumes or scent trails (43). Gradients in odor concentration at the edges of the plume may be efficiently encoded by "on" and "off" responses, whereas a more stable concentration within the plume may be represented by the neurons that exhibit persistent excitation or suppression. Interestingly, a similar diversity of odor-evoked responses was recently reported in the lateral entorhinal cortex (44), suggesting that it may be a general feature of odor-processing circuits.

How might the enhancement or suppression of spontaneous spiking be incorporated into a mechanism for encoding odors? As in other cortical regions, the rate of spontaneous spiking in the PC is low, approximately 1 Hz (Fig. 1B). However, the PC is thought to use a rate code (22, 45) and is able to identify odors quickly (46). Thus, if changes in the low rate of spontaneous firing are to represent information on a fast timescale, the PC or higher areas need to compute averaged firing rates across many neurons simultaneously. This computation must be achieved without erasing information about odor identity that is thought to be encoded in the spatially distinctive patterns of activity elicited by different odors (Fig. 4B). How this task might be implemented (47, 48) is an intriguing question for future work.

In summary, we have found that spontaneous activity in the anterior PC is inherited from the OB, differs between types of principal cells, and can be suppressed by odors. Our results suggest that spontaneous spiking can extend the dynamic range

of odor representations in the PC, enhancing the ability of this sensory paleocortex to encode complex olfactory information.

Materials and Methods

Animals and Anesthesia. All procedures involving mice were approved by the Animal Experimentation Ethics Committee of the Australian National University and conform to the guidelines of the National Health and Medical Research Council of Australia. Experiments used male or female, 50- to 70-d-old C57BL6/J mice. Mice were anesthetized with urethane + chlorprothixene (0.7 g/kg s.c. and 5 mg/kg s.c., respectively), ketamine + xylazine (100 mg/kg and 10 mg/kg, respectively), or fentanyl + medetomidine (0.1 mg/kg and 1 mg/kg, respectively).

Surgery. To access the anterior PC, overlying muscle and bone were sequentially dissected, and then a craniotomy (approximately 1.5 × 1.5 mm) was opened over the region where the middle cerebral artery (MCA) and the dorsal aspect of the lateral olfactory tract (LOT) intersect. The dura was removed also. A water-tight chamber was constructed around the surgical site to accommodate the water-immersion objective. The chamber was filled with Ringer's solution containing (in mM) 135 NaCl, 5.4 KCl, 1.8 CaCl₂, 1 MgCl₂, 5 Hepes at pH 7.4, and drugs were applied by perfusing this chamber.

Calcium Indicators. We used three types of calcium indicators in this study: the green dye Cal-520 AM, its red-shifted variant Cal-590 AM, and the green genetically encoded calcium indicator GCaMP6f. Cal-520 AM or Cal-590 AM (1 mM; AAT Bioquest) was pressure-injected into the PC at a depth of 200–300 μm with a micropipette. Sulforhodamine (SR) 101 (50 μM) was added to the Cal-520 solution to label astrocytes. Calcium imaging was performed at least 30 min after dye loading. For imaging with GCaMP6f, virus (AAV1.Syn. Flex.GCaMP6f.WPRE.SV40 + AAV1.hSyn.Cre.WPRE.hGH, from the University of Pennsylvania Vector Core) was stereotaxically injected into the PC or OB in mice anesthetized with isoflurane. Imaging was performed 2–3 wk later.

Two-Photon Imaging. Imaging was done using a customized Thorlabs two-photon microscope with a 16× water-immersion objective (0.8 NA; Nikon), resonant galvanometer scanners, and a Ti:Sapphire laser (Chameleon Ultra; Coherent). Time-series movies were captured at 30 or 60 frames/s.

Odor Stimulation. To measure spontaneous activity, a constant flow (0.5 L/min) of charcoal-filtered and humidified medical-grade air was delivered to the nares of the mouse. Odor stimulation used a custom-built 16-channel flow dilution

olfactometer to deliver odorized air into the clean air stream. The performance of the olfactometer was confirmed using a photo-ionization detector (miniPID; Aurora Scientific) (Fig. S6B).

Whole-Cell Recordings. Blind whole-cell current clamp recordings were obtained from layer 2 neurons using a patch pipette (5–7 $M\Omega$) filled with an intracellular solution containing (in mM) 135 KMeSO₄, 7 NaCl, 2 MgCl₂, 0.1 EGTA, 2 Na₂-ATP, 0.3 Na₃-GTP, 10 Hepes, adjusted to pH 7.2 with KOH and an osmolarity of 290 mOsm/kg.

Analysis. Analysis was done with ImageJ and custom MATLAB code (Math-Works). The raw fluorescence intensity of each region of interest (ROI) was

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calculated and expressed as $\Delta F/F_0 = (F - F_0)/F_0$, where F_0 is the median of the lower 80% of values during the clean air period. Transients in the $\Delta F/F_0$ trace were detected using a sliding template (49). Excitation and suppression were determined using a bootstrapping procedure to determine appropriate thresholds. False-positive rates were estimated to be <1.7%.

A full description of experimental procedures is found in *SI Materials* and *Methods*.

ACKNOWLEDGMENTS. We thank Brian Billups, Mark McDonnell, and Greg Stuart for helpful comments on the manuscript. This work was supported by Project Grants 1009382 and 1050832 from the National Health and Medical Research Council of Australia.

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