

# Top-down modulation of sensory processing and mismatch in the mouse posterior parietal cortex

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## **ABSTRACT (141 words)**

An important function of the neocortex is to compare sensory feedback stimuli with internal predictions of the outside world and evoke mismatch responses to deviations, thus allowing expectations to be updated. The mechanisms behind sensory feedback mismatch and prediction formation however remain unclear. Here we created a learned association of an auditory-tactile stimulus sequence in awake head-fixed mice, where a sound predicted an up-coming whisker stimulus and introduced mismatches by omitting or altering the whisker stimulus intensity. We showed that layer 2/3 posterior parietal cortex (PPC) neurons could report stimulus sequence mismatches, as well as display neural correlates of expectation. Inhibition of PPC-projecting secondary motor cortex (M2) neurons suppressed these correlates, along with population mismatch responses. Hence, M2 can influence sensory processing in the PPC and potentially provide the prediction of sensory feedback from learned relationships within sequences of sensory stimuli.

## **KEY WORDS**

2-photon calcium imaging, in vivo awake, sensory association, predictive processing

## **INTRODUCTION**

The experience of different sensory stimuli enables us to make associations between them, especially when presented as part of a sensory sequence. For example, we learn to anticipate the sound of thunder after a flash of lightning. We would be surprised in the absence of the expected stimulus because there was a mismatch between what we anticipated and what we actually experienced. The predictive processing framework postulates that the brain constantly predicts sensory feedback based on an internal representation of the outside world built upon prior sensory experience<sup>1,2</sup>. Mismatch responses, or prediction errors, that are generated during deviations between what was predicted and the actual sensory input are then used for updating stimulus expectations<sup>3</sup>. This is an essential feature at both the physiological and behavioral level as it facilitates the detection of unexpected, potentially dangerous events<sup>4</sup>, and is thus crucial to survival.

Sensory feedback mismatch responses have been described in the primary visual cortex (V1) in mice<sup>5,6</sup>, as well as in the primate primary auditory cortex<sup>7</sup> and the primary auditory pallium of songbirds<sup>8</sup>. In mice, local cortical circuits implicated in generating mismatch responses have been investigated mainly during locomotion in a virtual reality setting. There, the predicted sensory feedback from self-generated motion was compared against the actual sensory flow, when coupled with closed-loop visual, auditory or tactile stimuli<sup>9-11</sup>. Conversely it was recently shown that neurons within the posterior parietal cortex (PPC) of awake head-fixed mice can report mismatches in sensory sequences in the absence of locomotion/motor-output-related sensory feedback predictions, such as during the omission of a previously experienced passive touch to the whiskers<sup>12</sup>. The PPC, a sensory associative area, receives bottom-up sensory input from the primary sensory cortices<sup>13-15</sup>, and has reciprocal connections with the higher order cortical areas (secondary motor cortex M2, anterior cingulate cortex A24, midcingulate cortex A24', and orbitofrontal cortex) that can provide top-down feedback information<sup>16-18</sup>. It could therefore compare incoming sensory stimuli with the expectations of these stimuli constructed from prior experience, and report mismatches between the two, analogous to what has been described in visual processing in mouse V1<sup>19</sup>. However, little is known about how such learned associations formed from experiencing sensory sequences, where one stimulus predicts another, can influence sensory processing as well as the generation of mismatch responses.

Here, we presented awake head-fixed mice with an auditory stimulus via a loudspeaker followed by tactile stimuli through whisker deflection with a magnetic coil. Hence, we created a sensory stimulus sequence where a sound predicts the up-coming whisker stimulus. We could then introduce a mismatch in this sequence by varying the intensity of the whisker stimuli, or omitting its presentation, while recording neuronal activity of layer 2/3 PPC neurons at single-cell resolution with in vivo 2-photon calcium imaging. We demonstrate that the PPC can represent associations of sequences of sensory stimuli and reliably generate mismatch responses to deviations in these sequences. We further reveal that this dynamic representation of sensory stimuli is modulated by top-down feedback from M2 and can be interpreted within the framework of predictive processing.

# RESULTS

## Creating a sensory association at the PPC and reporting mismatch

To create an association between two sensory stimuli, we first presented awake head-fixed mice with an auditory tone (“sound” session), either looming or non-looming in intensity (Fig. 1a). This was followed by trials with only a whisker stimulus (“whisker” session). We then presented the whisker stimulus immediately after the auditory stimulus (“pairing” session), in this case the looming sound, so that the sound would predict the up-coming whisker stimulus. In subsequent trials (“interleaved” session), we created a mismatch (mismatch trials) in the experienced auditory-tactile sequence by randomly omitting the whisker stimulus in 20% of the trials. Hence the interleaved session comprised of randomized paired (sound followed by whisker stimulus) and mismatch (sound followed by whisker stimulus omission) trials (Fig. 1a). These sessions were presented sequentially within the same imaging session. Neuronal activity within the PPC was measured using 2-photon calcium imaging at single-cell resolution in layer 2/3 neurons of mice expressing the genetically encoded calcium indicator (GECI) RCaMP1.07<sup>20</sup>, with a chronic cranial window in place. Expression of this viral construct was previously shown to be mainly in pyramidal neurons<sup>11</sup>. Using intrinsic optical signal (IOS) imaging (see Methods and Supplementary Fig. 1) we localized area A of the PPC via exclusion, where mismatch responses to the omission of tactile stimuli were previously reported<sup>12</sup>.

We first identified neurons (6 mice, 1 field of view, FOV, per mouse) that responded to one of the presented stimuli (sound or whisker stimulus) in any of the four sequentially presented sessions and classified them as significantly responsive when compared to the shuffled data (see Methods). We subsequently based all our analysis on these shuffle-corrected neurons and followed their activity through the different sessions described below. Here, we present the mean  $\Delta F/F$  computed during a 1 s window from stimulus onset minus the baseline mean  $\Delta F/F$  1 s prior to stimulus. We focus primarily on the whisker stimulus window responsive neurons in the whisker, pairing and interleaved sessions, and describe the sound responsive neurons in Supplementary Fig. 2. During

whisker stimulation (“whisker” session) 40.0% of PPC neurons (331 of 834 neurons, 37% response probability) were responsive to whisker deflection (Fig. 1d, whisker-responsive neurons). The pairing of auditory and whisker stimuli (“pairing” session) recruited a new population of neurons that were previously weakly or non-responsive to the whisker stimulus alone (Fig. 1d, pairing-responsive neurons). Whisker stimulus evoked responses were larger in the pairing sessions compared to whisker sessions ( $p = 2.9 \times 10^{-7}$ , 331 whisker-responsive neurons, pairing-responsive neurons, with 35 overlapping neurons or 7.8%, Wilcoxon-Mann-Whitney test) (Fig. 1e).

In the subsequent interleaved session, we found that 8.6% of neurons within the PPC (72 of 834 neurons, 34.7% response probability) could report the omission of the previously experienced whisker stimulus (Fig. 1d, mismatch-responsive neurons). These mismatch responses were observed from the very first mismatch trial and were overall stable over time (Fig. 1f). Among the mismatch-responsive neurons, only three neurons were previously identified as looming offset-responsive in the sound session. This confirms that the mismatch neurons represent a newly recruited group of cells that report whisker stimulus omission and are not merely sound offset-responsive neurons. The mismatch response was also not associated with whisker movements (Fig. 1b, d, interleaved mismatch), consistent with previous studies<sup>12</sup>. We also observed mismatch responses of comparable size when the non-looming sound was paired with the whisker stimulus (Supplementary Fig. 2). In the interleaved session, we found that a larger fraction of neurons (271 of 834, 32.5%, with 31.7% response probability) were responsive to the interleaved paired trials, but with a significantly smaller response compared to the mismatch response ( $p = 2.5 \times 10^{-6}$ , 72 mismatch-responsive neurons, 271 interleaved paired-responsive neurons, Wilcoxon-Mann-Whitney test). Also, the interleaved paired-responsive neurons in the PPC displayed an increase in the mean  $\Delta F/F$  0.5 s before whisker stimulus onset (shaded window in Fig. 1d) from the pairing to interleaved session. This pre-stimulus increase was absent in the pairing-responsive neurons (Fig. 1g). Hence, the PPC can represent sensory stimuli in a stimulus sequence and report mismatches in the experienced sequence, with mismatch responses being larger than that of the predicted stimulus.

## Neural correlates of expectation in the PPC

We then questioned if the pre-stimulus increase in the mean  $\Delta F/F$  during the predicting sound window (Fig. 1g) could represent a neural correlate of expectation. In a separate set of experiments, we presented mice (6 mice, 6 FOV in total) to a first interleaved session where the whisker stimulus was delivered immediately after the sound (as in Fig.1a). This was followed immediately by a second interleaved session where we introduced a 1 s delay between the sound and the onset of the whisker stimulus, to test if the pre-stimulus increase in neural response could be prolonged and/or enhanced. We observed that in the interleaved paired trials, the pre-stimulus response was indeed prolonged as well as enhanced with the delay, along with a decrease in the post-stimulus whisker response (Fig. 2a, b). In the mismatch trials, there was no corresponding prolongation, and the pre-stimulus response was comparable to that without the delay, while the mismatch response itself was reduced in size (Fig. 2c, d). Hence the pre-stimulus response in the interleaved paired-responsive neurons could potentially represent a top-down driven neural correlate of expectation in the PPC, as mismatch trials are interleaved with the paired trials.

## PPC can report different types of sensory mismatch

So far, we employed the omission of the whisker stimulus as the mismatch in the interleaved trials. Can the PPC also report mismatches in stimulus intensity? We designed two different types of interleaved sessions that were presented separately to the mice (Fig. 2e). In one session, we interleaved the paired trials with mismatch trials that had a smaller whisker stimulus intensity. These decreased-intensity trials would then signal a negative mismatch. In contrast, to signal a positive mismatch in another session, we interleaved the paired trials with mismatch trials of a higher whisker stimulus intensity (increased-intensity trials). This would allow for the identification of potential negative and positive mismatch neurons as described in the canonical microcircuit for predictive processing<sup>21</sup>. Hence three separate interleaved sessions with either omission, decreased or increased whisker stimulus intensities as a mismatch were performed sequentially in a single imaging session. The PPC was able to effectively report mismatches in stimulus intensity in both directions, where the mismatch response was larger than the

corresponding interleaved paired response (Fig. 2f, g). The mismatch responses were also encoded by largely separate groups of neurons (Fig. 2h and Supplemental Fig. 3b), indicating the presence of distinct negative and positive mismatch neurons. We further observed that the pairing-responsive neurons showed a response in the increased-intensity mismatch trials, although comparable in size to their interleaved paired trial response (Supplemental Fig. 3b). Conversely the increased-intensity mismatch neurons were responsive in the prior pairing trials. Both groups displayed smaller responses to the whisker stimulus in the interleaved session with the omission-mismatch trials (Supplementary Fig. 4c, d). This suggests that the positive mismatch neurons may be part of a subgroup within the pairing-responsive neurons (see Discussion). Thus, the PPC can report positive and negative mismatches with distinct groups of neurons whose response dynamics can match the canonical microcircuit for predictive processing.

## **M2 modulates sensory processing in the PPC**

Next, we investigated the source of the top-down feedback to the PPC that could drive the pre-stimulus response/expectation. In addition to the primary sensory cortices, the PPC has strong reciprocal connections with higher order cortical areas, including M2<sup>12,18</sup>. These cortices have been shown to exchange sensory and motor information, respectively<sup>22</sup>, with the PPC. To test the potential contribution of M2 as the source of top-down feedback, we expressed the inhibitory DREADD hM4Di in ipsilateral PPC-projecting M2 neurons using a retro-Cre based viral expression strategy in mice that expressed RCaMP1.07 in PPC neurons (3 mice, 6 FOVs with and without CNO) (Fig. 3a and see Methods). We then administered CNO intraperitoneally (i.p.) to the mice to inhibit the activity of the PPC-projecting M2 neurons and presented the auditory-tactile sequence as described in Fig. 1a. We compared their neuronal response to that of control mice that were similarly administered CNO i.p. and did not express the inhibitory DREADD (4 mice, 6 FOVs). We observed that with the reduction of top-down feedback from ipsilateral M2 (hM4Di+CNO), the whisker stimulus responses of pairing-responsive neurons were reduced to a lesser extent between the pairing and interleaved paired session, as compared to the +CNO control experiments (Fig. 3b, c). The whisker stimulus response of the interleaved paired-responsive neurons on the other hand was equally enhanced in

the interleaved paired session, for both the hM4Di+CNO and +CNO control experiments (Fig. 3d, e). The reduction of M2 feedback also reduced the pre-stimulus response of the interleaved paired-responsive neurons in the interleaved paired session, compared to the +CNO control experiments (Fig. 3d, f). For the interleaved mismatch trials, we found that neurons could still report the omission mismatch, with a response size comparable to +CNO control experiments in the absence of M2 suppression (Fig. 4a, b). There were, however, less neurons recruited to report the mismatch when M2 was inhibited (4.1%, 42/1014 vs 8.8%, 78/882 for hM4Di+CNO and +CNO control experiments respectively). When we analyzed the impact of M2 inhibition on mismatch responses at the population level, we observed that the mismatch response was greatly suppressed compared to that in the +CNO control experiments (Fig. 4c, d). Taken together, these findings show that M2 can modulate sensory processing in the PPC, selectively suppressing bottom-up inputs while driving the pre-stimulus response/expectancy in different groups of neurons. Furthermore, M2 also contributes to the top-down prediction to the PPC that drives the omission mismatch neurons (Supplementary Fig. 5).

## DISCUSSION

Using an auditory tactile stimulus sequence, we were able to follow the formation of a sensory association in mice, along with the updating of stimulus expectations while we introduced mismatches in the sequence, as represented by the neural activity of layer 2/3 neurons in the PPC. We have shown that the PPC is able to dynamically represent sensory sequences with different groups of neurons, in contrast to S1. One of the features of learned sensory associations is the inhibition of the expected bottom-up input<sup>23,24</sup>. This became evident in the size of the whisker stimulus response, which was significantly reduced in the transition from pairing to interleaved trials. Indeed, both the whisker- and pairing-responsive neurons were suppressed outside of their respective sessions. This was in part modulated by top-down feedback from M2, which has been shown to suppress<sup>25</sup>, as well as enhance<sup>26-28</sup> sensory processing in primary sensory cortices, as part of a widespread cortical circuit motif<sup>26,29</sup>. Effectively, the paired-responsive neurons



increased their response to the whisker stimulus in the interleaved session, indicating that local microcircuits can be modulated differentially by top-down feedback at the same time. As both parvalbumin (PV) and vasoactive intestinal peptide (VIP) expressing inhibitory neurons can be directly activated by top-down inputs<sup>25,26</sup>, it would be important to determine their role in our findings.

In our experiments we used a sound presentation to predict an up-coming whisker stimulus, and subsequently show the appearance of a pre-stimulus response that could represent a neural correlate of expectation (Fig. 2e, h). This pre-stimulus response was present in the interleaved paired-responsive neurons and could be prolonged and enhanced temporally upon introducing a delay between the predicting sound and the expected whisker stimulus (Fig. 3a, b). We were able to suppress the pre-stimulus response with the inhibition of M2 feedback to the PPC, demonstrating that M2 can provide the top-down expectation/prediction to the PPC, which could also be interpreted as an attentional signal<sup>26</sup>. This top-down driven prediction can also drive inhibition, as observed in the suppression of the pairing-responsive neurons. Neurons in the PPC can then compare this prediction with the actual bottom-up sensory information and report mismatches.

Interestingly, the CNO injection itself in the absence of DREADD expression led to an enhancement of the whisker stimulus response (Supplementary Fig. 4b-e), as well as an increase in the overlap of whisker stimulus responsive neurons across the sessions (Supplementary Fig. 4f), compared to control experiments in the absence of CNO injection. Mismatch responses were also enhanced by CNO (Supplementary Fig. 4h), while the pre-stimulus response remained unchanged. CNO can be reverse metabolized to clozapine and can potentially exert physiological and behavioral changes associated with antipsychotic treatment<sup>30</sup>. This might in part explain the enhanced neuronal responses observed during the presentation of the audio-tactile stimulus sequence. Nonetheless decreasing M2 feedback reduced the size of the pre-stimulus response, as well as mismatch generation at the level of the population.

We have demonstrated that the PPC can report mismatches in the form of omission as well as deviations from the predicted strength of the whisker stimulus. Based on the canonical microcircuit for predictive coding<sup>10,21</sup>, these mismatch neurons could be categorized into negative and positive error neurons that would report mismatches that are less (omission and decreased intensity mismatch) or more (increased intensity mismatch) than what is expected. In our experimental paradigm the positive error neurons could form part of the pairing-responsive neurons, as they become suppressed by top-down feedback from M2 (Supplementary Fig. 5), as the prediction is formed in the interleaved sessions. Indeed, when the bottom-up input exceeded this prediction, these neurons could report this positive mismatch (Supplementary Fig. 3b). The paired-responsive neurons on the other hand showed mixed responses, integrating both top-down predictions and bottom-up stimulus information, in the form of the pre- and post-stimulus response. Finally, the omission and decreased-intensity mismatch-responsive neurons could function as negative error neurons as they are driven largely by the top-down prediction. Notably, we observed little overlap between the omission and decreased-intensity mismatch neurons, with mismatch responses of comparable size. Differences in the degree of bottom-up driven inhibition for the two mismatch types could potentially level out eventual expected differences in prediction error magnitude.

While the silencing of ipsilateral M2 feedback to the PPC reduced the number of mismatch neurons, it did not affect the size of single neuron mismatch responses. This suggests that multiple higher order areas, in addition to ipsilateral M2, could contribute to the top-down prediction. Nonetheless, the mismatch response was suppressed at the net population average. Also, a response to the predicting sound in the population average of all neurons can be observed prior to the mismatch response (Fig. 4c). This is likely mediated by neurons responding to the predicting sound within the population, that are independent of mismatch neurons, and furthermore not recruited when M2 feedback is silenced. This demonstrates a strong effect of M2 on the population code.

Based on these findings, we propose that during the interleaved paired trials, top-down inputs predicting the whisker stimulus are matched by bottom-up whisker stimuli that

recruit neighboring inhibitory neurons, such as the somatostatin (SST) expressing inhibitory neurons. These neurons are strongly driven by local excitation and could inhibit the dendrites of layer 2/3 neurons in the PPC<sup>10,21</sup>. In the interleaved mismatch trials, the top-down prediction can directly activate the dendrites of the PPC neurons in the absence of this locally recruited inhibition to drive the mismatch response. This would be in accordance with the framework of predictive coding during sensorimotor transformations and brings our study in line with the hierarchical visual processing described in mouse V1<sup>10,19</sup>.

## EXPERIMENTAL MODEL AND SUBJECT DETAILS

All experimental procedures followed the guidelines of the Veterinary Office of Switzerland and were approved by the Cantonal Veterinary Office in Bern. The data was collected from wild-type C57/BL6 mice (n = 15) and both males and females were used. All mice were at least 8 weeks old at the time of viral injection/head-post implantation.

## METHOD DETAILS

### Experimental design

This study did not involve randomization or blinding. No data or mice were excluded from the analysis.

### Surgery

Mice were anesthetized with isoflurane (3%) and subcutaneously injected with carprofen (5 mg/kg) prior to surgery for viral injections. For calcium imaging, the genetically encoded calcium indicator (GECI) RCaMP1.07 was injected into the PPC at 1.7 mm lateral and 2.0 mm posterior of bregma, to target layer 2/3 (~250 µm below the pial surface). For the silencing of M2 neurons, the DREADD hMD4i was injected into M2 at 0.5 mm lateral and 0.2 mm anterior of bregma, to target layers 2/3 and 5 (~250 and 500 µm below the pial surface). For long-term in vivo calcium imaging, a cranial window was implanted 24 h to 1 week after virus injections over the PPC as described previously<sup>31</sup>. Mice were

anesthetized with isoflurane and subcutaneously injected with buprenorphine (0.1 mg/kg) prior to surgery for window implantation. Briefly, a craniotomy was performed at the injection site. A cover glass (4 mm diameter) was placed directly over the exposed dura mater and sealed to the skull with dental acrylic. A metal post was fixed to the skull with dental acrylic, posterior to the cranial window, to allow for subsequent head fixation. One week after chronic window implantation, mice were handled daily for another week, and gradually habituated to head fixation. Intrinsic optical signal (IOS) imaging was performed on the mice to identify the location of PPC by exclusion as previously described<sup>12</sup>. In brief, to avoid the activation of surrounding whiskers, all whiskers except the  $\gamma$ -barrel-column whisker on the right whisker pad of the mice were trimmed and their locations were mapped using IOS on the exposed skull during whisker stimulation (rostrocaudal deflections at 10 Hz). The location of the primary visual cortex (V1) was similarly mapped using full-field stimulation with a green LED placed 5 mm in front of the contralateral eye. The non-activated region between these two identified sites was delineated as PPC for subsequent imaging sessions.

### **Viral constructs**

For calcium imaging, AAV2/1-hEF1 $\alpha$ -RCaMP1.07-WPRE-hGHp(A) (300 nL,  $\sim 5.0 \times 10^{12}$   $\mu$ g/mL) was injected into the PPC of wild-type mice targeting layer 2/3 to induce expression of the GEC1 RCaMP1.07 in neurons. For the silencing of M2 neurons, AAVretro2-hSyn-EBFP-iCre was injected in layer 2/3 of the PPC, along with RCaMP1.07 (for calcium imaging), and AAV2/1-hSyn-dlox-HA-hM4D-Gi-mCitrine was injected in M2 to selectively target PPC projecting M2 neurons.

### **DREADD inhibition and CNO control experiments**

The inhibitory DREADD hM4Di was used to chemogenetically silence M2 neurons projecting to the PPC. An i.p. injection of clozapine N-oxide (CNO dihydrochloride, 1 mg/kg, Tocris cat. no. 4936), the ligand that activates hM4Di, was done 30 minutes before presentation of the sensory stimulus sequence. We performed multiple imaging sessions with the presence of CNO first over several days, before repeating them in the absence of CNO at least 48 hours after the last CNO session. In the CNO control experiments,

wild-type mice without any DREADD expression were injected i.p. with CNO (1 mg/kg) 30 minutes before presentation of the sensory stimulus sequence.

## **Presentation of sound and tactile stimuli**

To create a sensory association of the stimulus sequence, each mouse was presented with, in the following order, the sound session (approximately 25 randomly interleaved trials each of the looming and non-looming sound), the whisker session (approximately 50 whisker stimulus trials), the pairing session (approximately 20 trials with looming sound followed by the whisker stimulus), the interleaved session (approximately 120 paired trials of looming sound followed by the whisker stimulus, randomly interleaved with approximately 30 mismatch trials). The looming (increasing intensity) and non-looming (constant intensity) sounds were based on previously recorded soundtracks<sup>12</sup> and recreated in MATLAB, that consisted of a cloud of tones (0.1–8 kHz). Each sound was 1 s long and was delivered via a loudspeaker at 75 dB placed contralateral to the imaging site. We used the looming sound to predict the whisker stimulus in the pairing session as mismatch responses to the absence of a tactile stimulus cued by a looming sound could be generated in the PPC<sup>12</sup>. Mismatch responses could also be evoked in the PPC when the non-looming sound was used in the pairing session (Supplementary Fig. 2), suggesting that a range of auditory cues could be potentially used in the sensory sequence. Tactile stimuli were delivered as deflections ( $3 \times 10$  Hz, 1 ms pulse) to multiple whiskers. This was achieved by attaching small metal particles to the whiskers and subsequently moving them via a brief magnetic field generated by a coil placed beneath the head of the mouse<sup>32</sup>. The whisker stimulus intensity was set to 80% for all experiments except for those performed during the intensity mismatch experiments (Fig. 3e, f). Here, the interleaved paired stimulus was set to 60%, and the decreased and increased intensity mismatch stimuli were set to 40% and 80%, respectively. Each trial began with a 2 s baseline, followed by a brief sound cue (50 ms, 85 dB) to signal trial start to the mouse. In the pairing and interleaved trials, the looming sound was played 1 s after the sound cue and was followed immediately by the whisker stimulus. In a subset of control experiments, the non-looming sound was played instead of the looming sound (see

Supplementary Fig. 2). Each trial was 8 to 9 s long and trials were presented with an inter-trial interval of 2 to 5 s, to render them irregular in presentation timing. White noise (65 dB) was played during the entire duration of the imaging session. Sound and whisker stimulus delivery was controlled by custom-written software in C.

### **Whisker tracking**

The whisker field was illuminated with a 940-nm infrared LED light and movies were acquired at 100 Hz (500 × 500 pixels) using a pixy camera system coupled with an Arduino board to track the position of the whiskers in real time<sup>33</sup>.

### **Two-photon calcium imaging**

We used a custom-built 2-photon microscope controlled by ScanImage 2019 equipped with a fixed wavelength fiber laser at 1064 nm (Fidelity; Coherent), a water-immersion objective (16×LWDPF, 0.8 NA; Nikon), resonant scan mirrors (model 6210; Cambridge Technology), and a Pockel's Cell (Conoptics) for laser intensity modulation. For calcium imaging, RCaMP1.07 was excited at 1064 nm with the Fidelity. Emitted fluorescence was collected with red (617/73 nm) and green (520/60 nm) emission filters. Images were acquired at 30 Hz with 512 x 512-pixel resolution.

### **Histology**

After the calcium imaging recordings, mice were deeply anaesthetized by intraperitoneal injection of 80/10 ketamine/xylazine mixture and transcardially perfused with 4% paraformaldehyde (PFA). Brains were removed and post-fixed in PFA for 24–48h at 4°C and were subsequently washed in phosphate-buffered saline and sliced at 100 µm. Brain slices were mounted using Mowiol® 4-88 prior to imaging on a LEICA m205 FCA fluorescence stereo microscope to confirm the location of the PPC.

## **QUANTIFICATION AND STATISTICAL ANALYSIS**

### **Two-photon calcium data processing**

Somatic calcium signals were automatically detected using the Python-based CalmAn analysis pipeline, which performed motion correction, source extraction, and component registration<sup>34</sup>. Raw fluorescence traces ( $F$ ) of the calcium signals are presented here as  $\Delta F/F = (F - F_0)/F_0$ , where  $F_0$  was calculated for each trial as the mean of the 1 s window prior to the sound cue that signalled trial start. The responsiveness of a detected neuron to a given stimulus (sound, sound-offset, whisker, mismatch) was determined by comparing the distribution of its single trial responses (single trial mean  $\Delta F/F$  calculated in a 1 s window from stimulus onset minus the mean baseline window 1 s before stimulus onset) against the distribution of 1000 randomly selected events from its same session (random baseline corrected mean  $\Delta F/F$  calculated in a 1 s window as above), hence taking into account the random noise of each neuron. Significance was determined with a two-sided Mann-Whitney-U test ( $p < 0.05$ ). We thus identified populations of neurons that were positively or negatively modulated by a given stimuli. We subsequently classified the positively modulated shuffle-corrected neurons as looming-, non-looming-, looming offset-, non-looming offset-, whisker-, pairing-, paired- or mismatch-responsive, based on the session in which they were significantly responsive compared to their shuffled data. The mismatch- and interleaved paired-responsive populations were not mutually exclusive and can show overlaps. For example, 6 neurons in Fig.1 were classified as both mismatch and interleaved paired-responsive. Response probabilities of neurons were calculated by dividing the number of responsive trials by the total number of trials presented for a particular stimulus in a session. Single trial  $\Delta F/F$  traces were first smoothed with a 1<sup>st</sup>-order Savitsky-Golay filter, 150 ms window. A neuron was considered responsive in a trial when its smoothed  $\Delta F/F$  trace in a 1 s window from stimulus onset was significantly larger than its baseline, calculated in a 1 s window preceding the stimulus onset (comparison of 30 frames before and after the stimulus onset, Wilcoxon-Mann-Whitney test,  $p < 0.05$ ). Population average responses presented here were determined from the averaged traces of each neuron that has been identified as being session responsive in its respective stimulus window. The population average neuronal traces were baseline-corrected by subtracting the mean of the baseline (1 s prior to stimulus onset, corresponding to the sound presentation window) from the entire averaged trace. For the experiments in Fig. 3 with a 1 s delay, the sound presentation



window was used as the baseline. Pre- and post-whisker stimulus response difference between interleaved paired and pairing session was computed as the difference between the baseline corrected mean  $\Delta F/F$  calculated in the respective 1 s window (pre-stimulus or post-stimulus) from the interleaved-paired and pairing session, of pairing-responsive and interleaved paired-responsive neurons. Hence a positive value would signal an increase in the response, and a negative value a decrease in the response. Single trial  $\Delta F/F$  traces in Fig. 1b were smoothed with a 1<sup>st</sup>-order Savitsky-Golay filter, 150 ms window. No smoothing was performed for the other population average neuronal traces. We present the mean  $\Delta F/F$  as a first measure of neuronal activity.

## Statistical analysis

All neuronal traces are presented as mean  $\pm$  s.e.m. unless stated otherwise. In each box plot the central line indicates the median, the bottom and top edges of the box indicate the 25<sup>th</sup> and 75<sup>th</sup> percentiles respectively, the whiskers extend to the extreme data points (maximum and minimum points within 1.5 standard deviation), and outliers are marked with crosses. The non-parametric Wilcoxon signed-rank paired test and Wilcoxon-Mann-Whitney test for paired and unpaired group comparisons were performed respectively. All tests were two-sided. We did not test for a normal distribution of the data.

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## AUTHOR CONTRIBUTIONS

C.R., S.K., B.L and S.S. performed the experiments and analyzed the data. T.N. provided materials and resources. S.K. setup the IOS, whisker tracking, and data analysis pipeline. C.R. and S.S designed the experiments. S.S. built the microscope, secured the funding



and supervised the project. C.R. and S.S. wrote the manuscript with comments from all authors.

## **DECLARATION OF INTERESTS**

The authors declare no competing interests.

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## Figure legends

**Fig. 1 Creating a sensory association at the PPC.** **a** Schematic of the sequence of stimulus presentation carried out in a single imaging session. All trials begin with a brief sound cue (not shown). Looming and non-looming sounds are randomly interleaved in the sound trials. The whisker stimulus (orange dot in **b**) delivered by a magnetic coil is presented alone in the whisker session. The looming sound (black bar in **b**) is paired with the whisker stimulus in pairing session. In the interleaved session, the paired trials (looming sound followed by whisker stimulus) are randomly interleaved with the mismatch trials where the whisker stimulus is omitted. **b** Example stimulus-responsive neurons classified as whisker-, pairing-, interleaved paired-, and interleaved mismatch-responsive neurons from layer 2/3 of the PPC, expressing RCaMP1.07. Average (black) and single trial (grey)  $\Delta F/F$  calcium traces are shown, along with the corresponding average whisker position (green). **c** Heat maps representing the average stimulus responses of whisker- (331 neurons), pairing- (153 neurons), interleaved paired- (271 neurons) and interleaved mismatch-responsive neurons (72 neurons) from 6 wild-type mice. Mean pre-stimulus activity (1 s) was subtracted from the calcium transients and neurons were sorted to their mean  $\Delta F/F$  response times in the stimulus window. **d** Population averages ( $\pm$  s.e.m.) of  $\Delta F/F$  traces of the neurons in **c**, along with their corresponding population average for the other presented stimuli. The average whisker position from the above sessions is shown below in green. **e** Box plot of average population responses of whisker-, pairing-, paired- and mismatch-responsive neurons. **f** Average population responses of mismatch-responsive neurons over mismatch trials. **g** Boxplot of pre-whisker stimulus response of pairing- and interleaved paired-responsive neurons, during their pairing and interleaved-paired sessions. Pre-stimulus response corresponds to shaded area in **d**. Outliers are shown in Supplementary Fig. 2f. Data are represented as mean  $\pm$  s.e.m. in **f**. Statistical significance is indicated by \*\* for  $p < 0.01$  and \*\*\* for  $p < 0.001$ , with two-sided Wilcoxon-Mann-Whitney test.

**Fig. 2 PPC can reliably update the mismatch response and report different types of mismatch.** **A** Heat maps and corresponding population averages ( $\pm$  s.e.m.) of  $\Delta F/F$

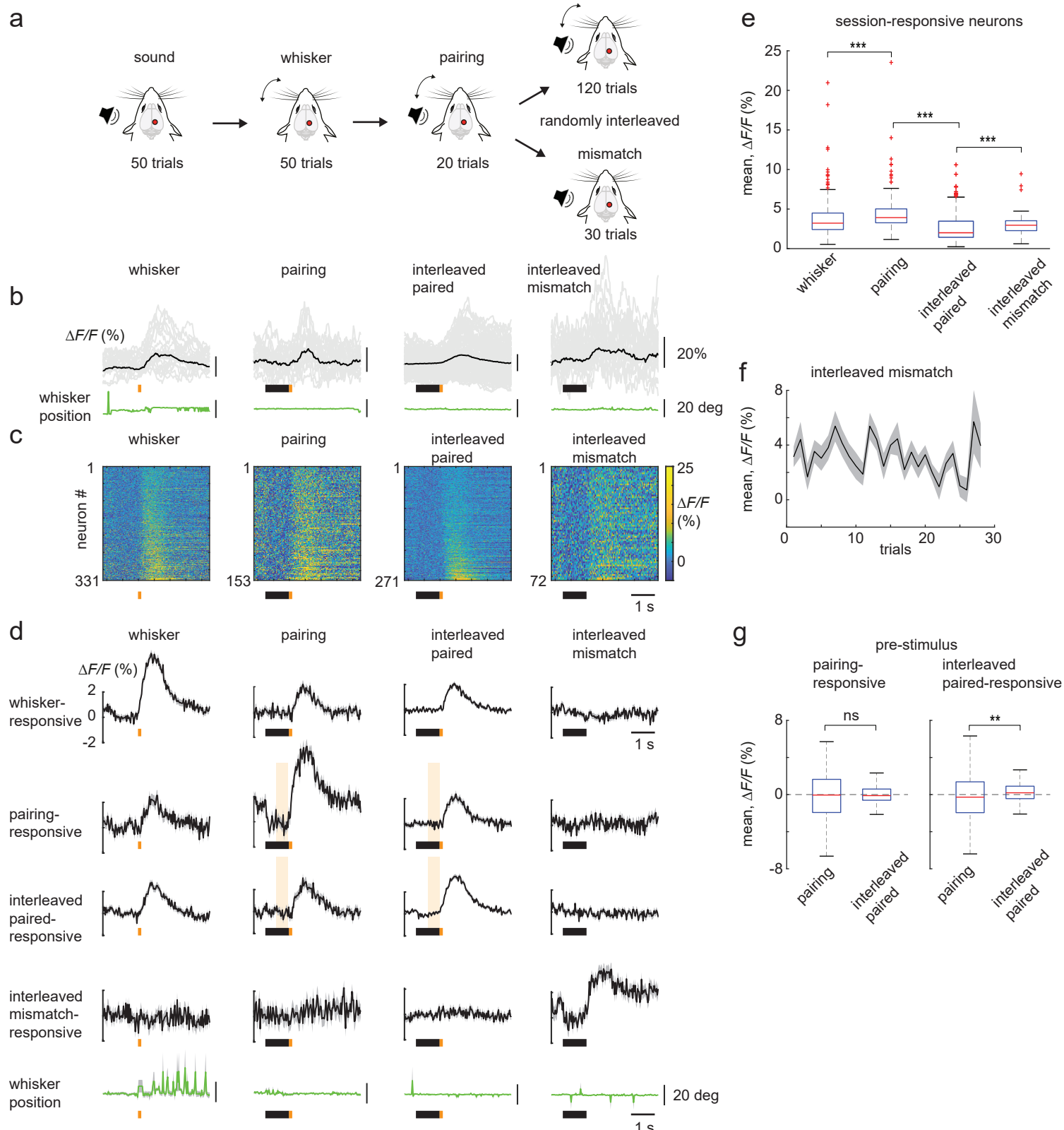
traces of interleaved paired-responsive neurons with no delay (0 ms) and a 1000 ms delay between the looming sound and whisker stimulus presentation (without delay, 279 neurons; with delay, 369 neurons; 6 mice; 6 FOVs). The average whisker position from the above sessions is shown below in green. **b** Box plot of average population pre-stimulus and post-stimulus responses of interleaved paired-responsive neurons, without and with delay. **c–d** same as **a–b**, but for mismatch-responsive neurons (without delay, 79 neurons; with delay, 94 neurons). **e** Schematic of the presentation of different mismatch trial types in the form of whisker stimulus omission, decreased and increased whisker stimulus intensity. **f** Population averages ( $\pm$  s.e.m.) of  $\Delta F/F$  traces of interleaved mismatch-responsive neurons and paired-responsive neurons, for the interleaved sessions with omission (124 mismatch neurons, 307 paired neurons), decreased (140 mismatch neurons, 217 paired neurons) and increased (116 mismatch neurons, 316 paired neurons) whisker stimulus mismatches, with their corresponding paired (red) and mismatch (blue) trial averages. **g** Box plot of average population responses of mismatch-responsive and paired-responsive neurons, for the interleaved sessions with omission, decreased and increased whisker stimulus mismatches. **h** Venn diagram of the different mismatch-responsive neurons in **f**, their respective overlaps between mismatch sessions. wh stim: whisker stimulus. Data are represented as mean  $\pm$  s.e.m. Statistical significance is indicated by \*\* for  $p < 0.01$ , and \*\*\* for  $p < 0.001$ , with two-sided Wilcoxon-Mann-Whitney test.

**Fig. 3 M2 modulates sensory processing in the PPC.** **a** Injection scheme to express the inhibitory DREADD hM4Di in M2 neurons projecting to the PPC (sagittal view), along with a coronal section showing their expression in M2 neurons as represented by mCitrine fluorescence. **b** Population averages ( $\pm$  s.e.m.) of  $\Delta F/F$  traces of pairing-responsive neurons, in the pairing and interleaved paired sessions, for the hM4Di+CNO (3 mice, 6 FOVs, 80 neurons) and +CNO control experiments (4 mice, 6 FOVs, 49 neurons). **c** Box plot of post-whisker stimulus response difference between interleaved paired and pairing sessions in the pairing-responsive neurons in **b**. **d** Population averages ( $\pm$  s.e.m.) of  $\Delta F/F$  traces of interleaved paired-responsive neurons, in the pairing and interleaved paired sessions, for the hM4Di+CNO (195 neurons) and +CNO control experiments (90

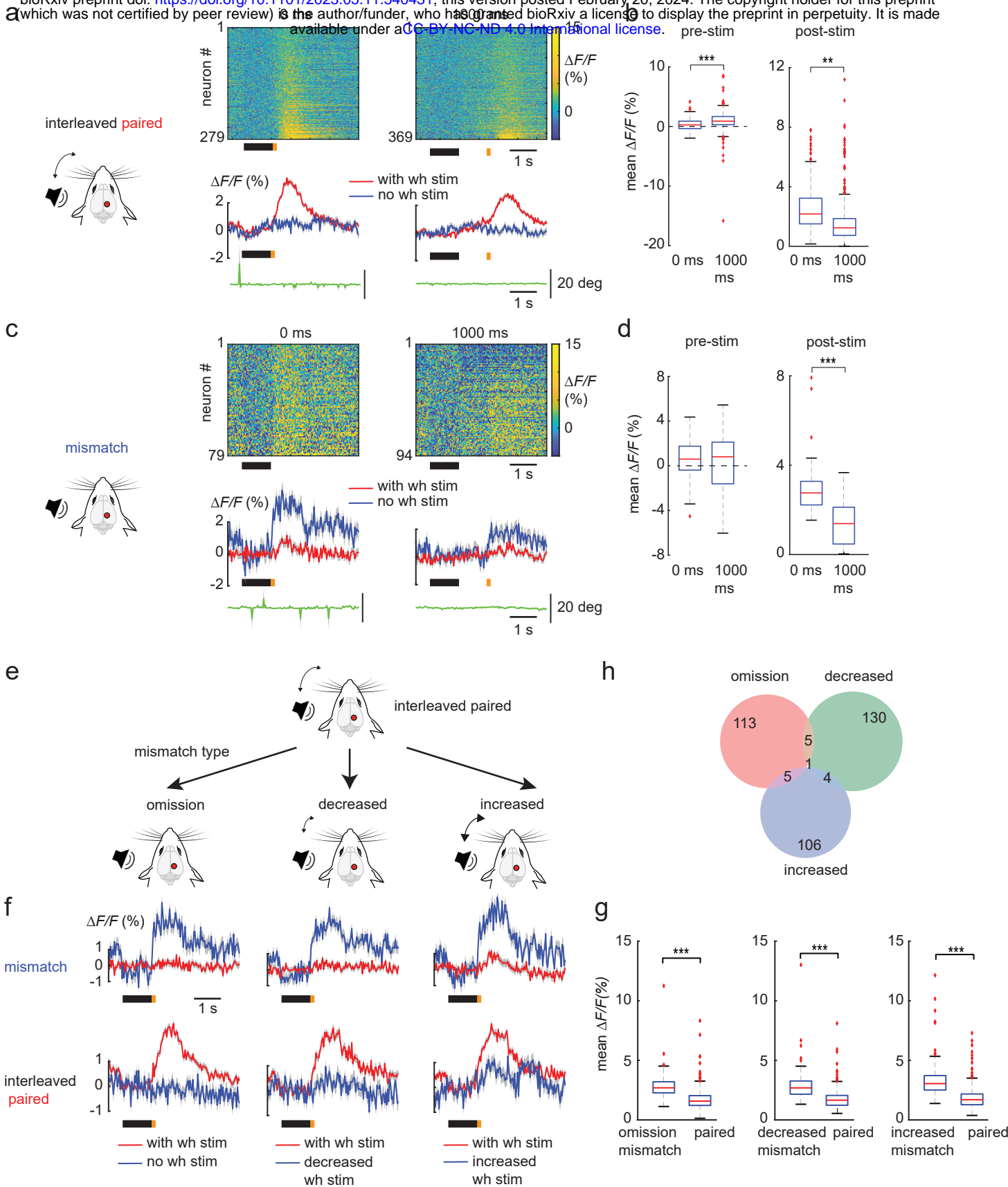
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**Fig. 4 M2 contributes to the top-down prediction to the PPC.** **a** Population averages ( $\pm$  s.e.m.) of  $\Delta F/F$  traces of interleaved mismatch-responsive neurons in layer 2/3 neurons in the PPC with M2 suppression using hM4Di+CNO (42 neurons) and with +CNO control experiments (78 neurons) (same mice as in Fig. 4). **b** Box plot of average population responses of the interleaved mismatch-responsive neurons in **a**. **c** Population averages ( $\pm$  s.e.m.) of  $\Delta F/F$  traces of all neurons during the mismatch trials in hM4Di+CNO ( $n = 1014$  neurons) and +CNO control experiments ( $n = 882$  neurons). **d** Box plot of average population responses of the neurons in **c** during the mismatch trials. Data are represented as mean  $\pm$  s.e.m. Statistical significance is indicated by \*\*\* for  $p < 0.001$ , with two-sided Wilcoxon-Mann-Whitney test.

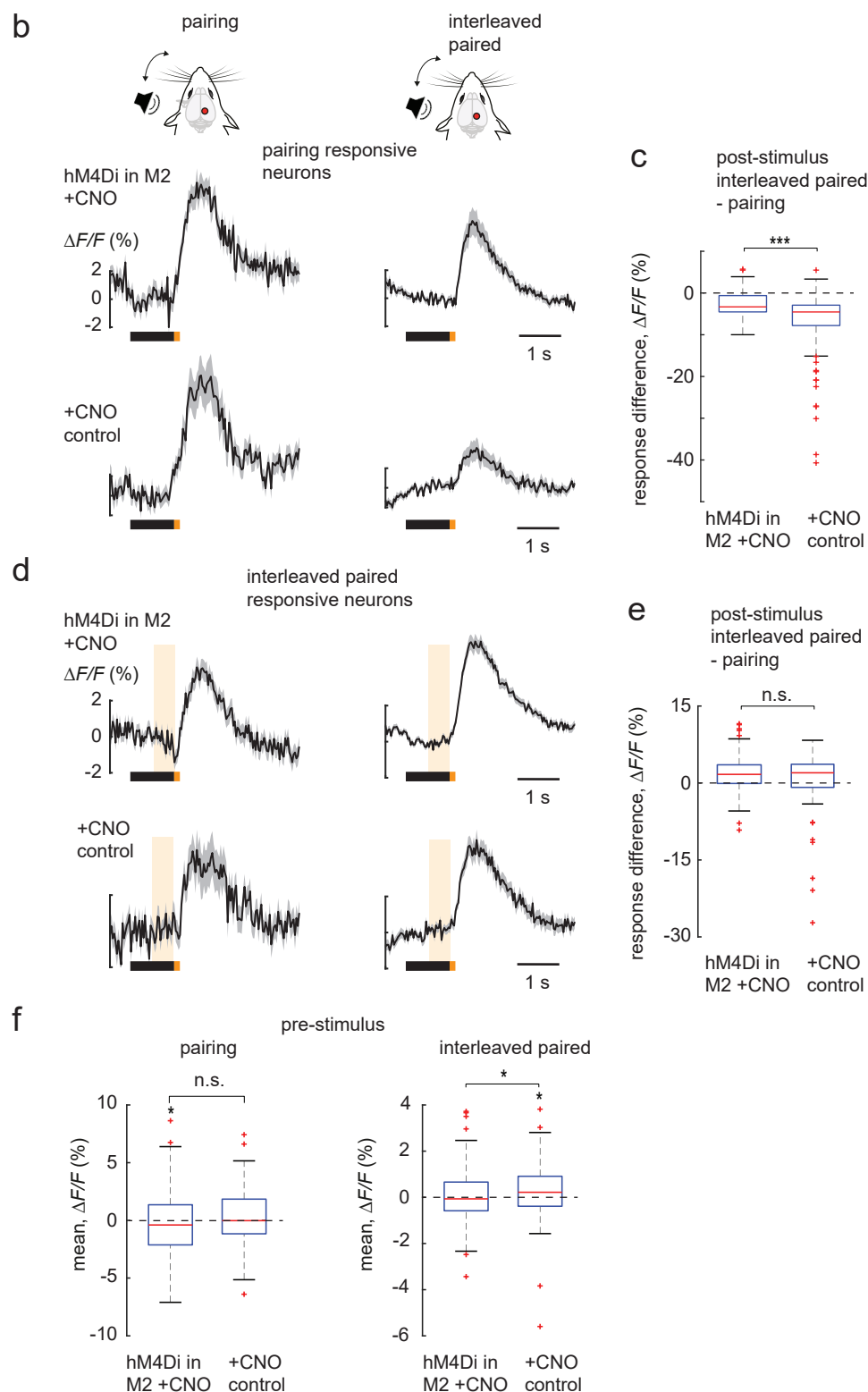
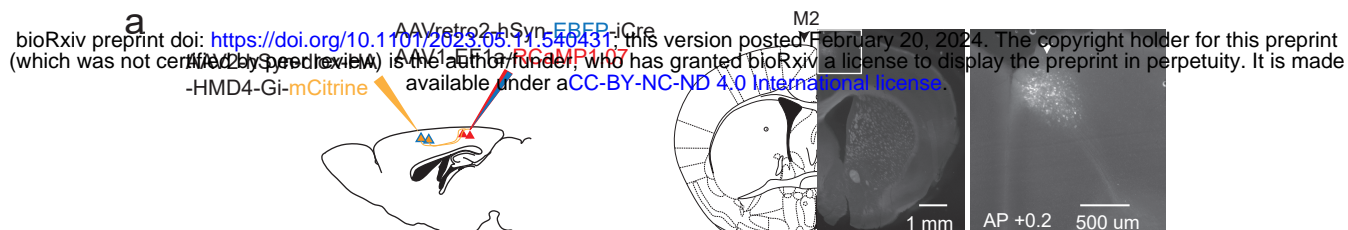




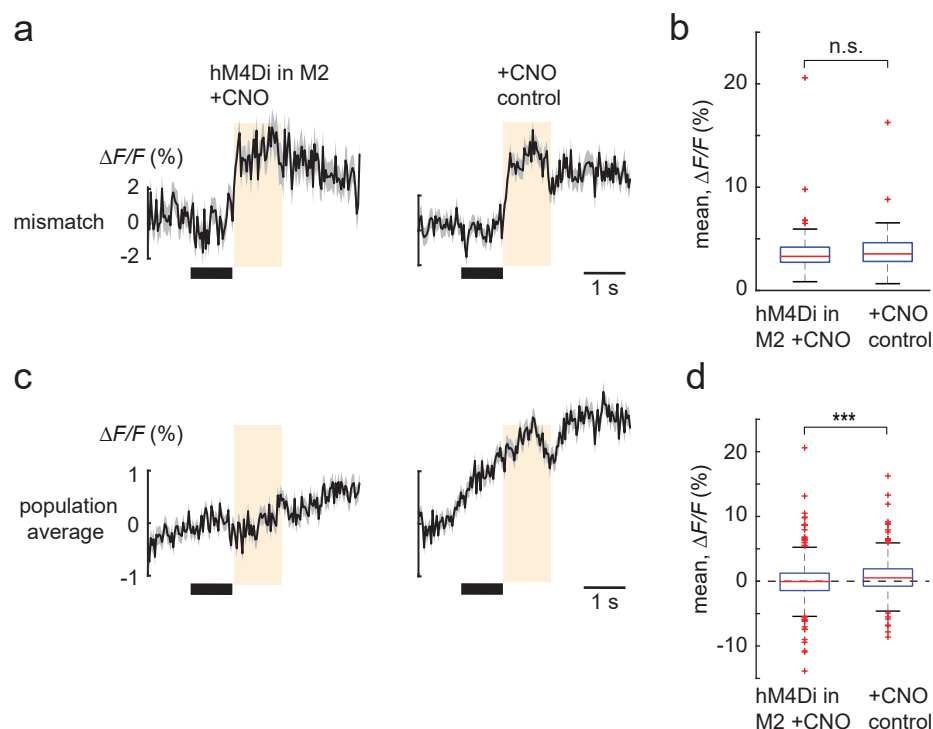
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