Modular network between postrhinal visual cortex, amygdala, and entorhinal cortex

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**Recommended Citation**

Meier, Andrew M.; Wang, Quanxin; Ji, Weiqing; Ganachaud, Jehan; and Burkhalter, Andreas, "Modular network between postrhinal visual cortex, amygdala, and entorhinal cortex." *The Journal of Neuroscience*. 41, 22. 4809 - 4825. (2021).  
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The postrhinal area (POR) is a known center for integrating spatial with nonspatial visual information and a possible hub for influencing landmark navigation by affective input from the amygdala. This may involve specific circuits within muscarinic acetylcholine receptor 2 (M2)-positive (M2⁺) or M2⁻ modules of POR that associate inputs from the thalamus, cortex, and amygdala, and send outputs to the entorhinal cortex. Using anterograde and retrograde labeling with conventional and viral tracers in male and female mice, we found that all higher visual areas of the ventral cortical stream project to the amygdala, while such inputs are absent from primary visual cortex and dorsal stream areas. Unexpectedly for the presumed salt-and-pepper organization of mouse extrastriate cortex, tracing results show that inputs from the dorsal lateral geniculate nucleus and lateral posterior nucleus were spatially clustered in layer 1 (L1) and overlapped with M2⁺ patches of POR. In contrast, input from the amygdala to L1 of POR terminated in M2⁻ interpatches. Importantly, the amygdalocortical input to M2⁻ interpatches in L1 overlapped preferentially with spatially clustered apical dendrites of POR neurons projecting to amygdala and entorhinal area lateral, medial (ENTm). The results suggest that subnetworks in POR, used to build spatial maps for navigation, do not receive direct thalamocortical M2⁺ patch-targeting inputs. Instead, they involve local networks of M2⁻ interpatches, which are influenced by affective information from the amygdala and project to ENTm, whose cells respond to visual landmark cues for navigation.

Key words: amygdala; entorhinal cortex; navigation; postrhinal cortex; thalamocortical connections; visual cortex

Significance Statement

A central purpose of visual object recognition is identifying the salience of objects and approaching or avoiding them. However, it is not currently known how the visual cortex integrates the multiple streams of information, including affective and navigational cues, which are required to accomplish this task. We find that in a higher visual area, the postrhinal cortex, the cortical sheet is divided into interdigitating modules receiving distinct inputs from visual and emotion-related sources. One of these modules is preferentially connected with the amygdala and provides outputs to entorhinal cortex, constituting a processing stream that may assign emotional salience to objects and landmarks for the guidance of goal-directed navigation.

Introduction

The chief concern for survival is to detect danger and react to it (Price, 2005). In rodents, defensive strategies include the reflexive startle response, freezing when danger is distant and escaping when it is close. To do this, it is important to know the spatial layout of the environment. For this, animals rely on neural circuits connecting multiple brain structures including the amygdala, which drives flight, freezing, and approach (Janak and Tye, 2015); and areas of the limbic and sensory cortex (LeDoux, 2012). In rodents, a critical structure is the postrhinal area (POR; Burwell, 2001), which is part of the parahippocampal cortex (Burwell et al., 1995) and is a center for visual scene processing (Epstein et al., 1999). Studies in rats have shown that POR integrates visual inputs to extract contextual information (Furtak et al., 2012). It also references head-centered information to the visual scene, which is thought to form the basis for the grid-cell metric in entorhinal cortex (Furtak et al., 2012; Gofman et al., 2019; LaChance et al., 2019). The network for spatial processing is often contrasted with a pathway that carries nonspatial information through perirhinal and lateral entorhinal cortices to the hippocampus (Deshmukh et al., 2012; Knierim et al., 2014; Nilssen et al., 2019). The second function is the association of visual objects with reward and fear memory via a circuit with the amygdala (Agster et al., 2016; Burgess et al., 2016; Ramesh et al., 2019).
2018). This pathway is known as the “cortical route” for visual fear response, as opposed to the “subcortical route” for aversive responses (Pessoa and Adolphs, 2011; Agster et al., 2016). A source of the subcortical input channel is the superior colliculus (SC), which connects through the lateral posterior nucleus to the amygdala, where it triggers defensive responses to visual threats (Wei et al., 2015). This information may be used in the circuit with POR to associate affective significance to objects and places (Furtak et al., 2012; Burgess et al., 2016; Ramesh et al., 2018). A second branch of the pulvinocortical pathway sends input directly to POR and provides for its high sensitivity to moving stimuli (Nishio et al., 2018; Beltramo and Scanziani, 2019). Calcium recordings in V1 have shown that lateral posterior nucleus (LP) terminals carry signals for unexpected visual motion that are incongruent with the running speed of the animal (Roth et al., 2016). These responses suggest that locomotion-related cues may be integrated in POR, enabling the detection of moving objects during self-motion. Direct input to POR may also come from the dorsal lateral geniculate nucleus (dLGN; experiment 479670988, Allen Brain Atlas), whose terminals in layer 1 (L1) of primary visual cortex (V1) are most active when the running speed matches the optic flow (Roth et al., 2016), providing cues for determining the position relative to landmarks.

We have shown in mouse POR that muscarinic acetylcholine receptor 2 (M2) expression in L1 is patchy (Ji et al., 2015). Recordings in V1 have further shown that L2/3 neurons aligned with M2+ patches are more sensitive to object features, while cells in M2− interpatches are more sharply tuned to moving stimuli (Ji et al., 2015). This pattern is reflected in the organization of dLGN inputs to M2+ patches and LP projections to M2− interpatches (D’Souza et al., 2019). The findings raised the question whether the modular organization of thalamocortical inputs is preserved in POR, and inputs from the amygdala overlap with neurons projecting to the amygdala and ENTm. To address these questions, we have used anterograde and retrograde tracers of connections to and from POR. The results show that POR receives inputs mainly from areas of the ventral cortical processing stream (Wang et al., 2012). dLGN and LP projections to L1 overlap in M2+ patches, whereas inputs from the amygdala terminate in M2− interpatches. The overlap of amygdalocortical inputs with the apical dendritic tufts of POR neurons projecting to the amygdala and ENTm suggests that M2 modules provide a substrate for the affective modulation of POR output used for navigation.

Materials and Methods

Animals

Experiments were performed in 5- to 10-week-old male and female C57BL/6J, Ai9 (Gt[ROSA]26Sortm1(CAG-tdTomato)Hze])/J, PV-Cre (B6.129P2-Pdhltm1(CAG-Cre)Abhr/J], x Ai9 (Gt[ROSA]26Sortm1(CAG-tdTomato)Hze])/J, and ChrM2 tdt-D knock-in (B6.Cg-Chrm2tm20(Aai)1Hze)J mice. All experimental procedures were performed in accordance with the National Institutes of Health guidelines and under the approval of the Washington University Institutional Animal Care and Use Committee.

Surgical procedures

Mice were anesthetized with a ketamine/xylazine mixture (86/13 mg/kg, i.p.). Analgesia was achieved by presurgical injections of buprenorphine SR (0.1 mg/kg, s.c.). Ocular ointment was applied to protect the corneas. Mice were head fixed in a stereotaxic frame. Body temperature was maintained at 37°C. For each injection site, a small craniotomy was made over the target. Tracer injections were performed with glass micro-pipets (tip diameter, 15−20 μm) and, depending on the tracer used, attached to a Picospritzer [for bisbenzimide and diamidino yellow (DY)], an iontophoresis current source [for biotinylated dextran amine (BDA)], or a Nanoject II pump [for adeno-associated viruses (AAVs)]. At each cortical site, two injections were made: one at 0.3 mm and the other at 0.5 mm below the pial surface. After injections were complete, pipettes remained in place for 3 min before they were retracted. The scalp was stapled closed with wound clips.

Combined anterograde tracing with BDA and retrograde tracing with bisbenzimide

For mapping axonal projections from visual cortex to the amygdala in C57BL/6J mice, BDA (10,000 molecular weight, 5% in H2O; Thermo Fisher Scientific) was injected through a glass micropipet using iontophoresis (7 s duty cycle for 7 min, 3 μA; Midgard, Stoelting). Injections were made into the left hemisphere (in mm, anterior to transverse sinus/ lateral to mid sagittal suture), as follows: V1 (1.1/2.8), lateral medial area (LM; 1.4/4.0), posterior area (P; 1.0/4.2), lateral intermediate area (LI; 1.4/4.2), POR (1.15/4.3), anterolateral area (AL; 2.4/3.7), rostrolateral area (RL; 2.8/3.3), posterior medial area (PM; 1.9/1.6), anteromedial area (AM; 3.0/1.7), and anterior area (A; 3.4/2.4). In the same animal, callosal connections were retrogradely labeled by blanketing the right visual cortex with 20–30 injections (20 nl each) of bisbenzimide (5% in H2O; Sigma-Aldrich). After 3 d of survival, mice were killed with an overdose of ketamine/xylazine (500/50 mg/kg, i.p.) and perfused with 4% paraformaldehyde [PFA; 0.1 M phosphate buffer (PB), pH 7.4]. The brain was extracted from the cranium and cryoprotected in 30% sucrose (PB). The bisbenzimide-labeled callosal projection pattern was imaged in situ under a stereomicroscope (model MZ16F, Leica) equipped with fluorescence optics (excitation, 338 nm; emission, 505 nm). Injection sites were visible in situ (and validated later in sections) as weakly fluorescent spots (see Fig. 3A–I), whose locations relative to the callosal labeling pattern were used for areal identification (Wang and Burkhalter, 2007). Coronal sections (50 μm) were cut on a cryostat and stained for BDA using an avidin–biotin complex (ABC) reaction (Vectastain ABC Elite System). Sections containing the brainstem, diencephalon, and the amygdala were mounted on glass slides, and after dehydration in ethanol and clearing in xylene, the diaminobenzidine reaction product was intensified with AgNO3 and H2O2 (Wang et al., 2012). Representative sections were stained with cresyl violet for Nissl substance. The slides were coverslipped with mounting medium (DPX, Sigma-Aldrich) and imaged under a microscope equipped with dark-field optics. Images were captured with a CCD camera (CoolSnap EZ, Roper Scientific). For analyses of the termination pattern in each nucleus, coronal sections were imaged under 10× magnification, and grayscale images were recorded with Meta Morph NX2.0 acquisition software (Molecular Devices). The optical density (OD) of BDA-labeled axons was determined by blurring the raw image and measuring pixel values in regions of interest (ROIs) with ImageJ.

Retrograde tracing with diamidino yellow

To determine the sources of cortical and subcortical inputs to POR, neurons were retrogradely labeled with the fluorescent tracer diamidino yellow (2% DY in H2O, 50 nl; EMS-Chemie). Injections into POR (1.15/4.3 mm anterior to transverse sinus/lateral to mid sagittal suture) were made in PVDt (parvalbumin-tdt) mice whose pattern of tdt (tdTomato) fluorescence was used to identify cortical areas and subcortical nuclei (Gâmanu et al., 2018). After 3 d of survival, mice were perfused with 1% PFA. Brains were extracted, and the cortex including the hippocampus was detached from the rest of the brain and flattened (Ji et al., 2015). Next, the tissue was postfixed in 4% PFA (overnight, at 4°C), followed by overnight immersion in 30% sucrose (in PB, at 4°C). Tangential sections were cut at 40 μm on a freezing microtome, wet mounted on glass slides, and coverslipped in PB. Images were captured with a CCD camera (Lumenera Infinity3-URM, Teledyne) and Meta Morph NX2.0 acquisition software (Molecular Devices) using a stereomicroscope (model MZ16F, Leica) or an epifluorescence microscope (model 80i, Nikon), respectively, equipped with UV fluorescence (excitation, 355–425 nm; emission, 470 nm) and tdt (excitation, 520–600 nm; emission, 570–720 nm) optics. For storage, sections were mounted on glass slides and coverslipped with DPX. To reveal DY-
labeled cells in the amygdala, the neocortex was separated along the rhi-
nal fissure from the rest of the brain, which was then sectioned at 40 μm in the
coronal plane.

Anterograde tracing with AAVs
For retinotopic mapping of POR in C57BL/6j mice, side-by-side injec-
tions with AAV2/1.hSynapsin.EGFP.WPRE.bGH (46 nl; Penn Vector Core) and AAV2/1.hSynapsin.tdTomato.WPRE (46 nl; Penn Vector Core) were made at select combinations of retinotopic locations in V1. The injections were confined to a space within (in mm, anterior to trans-
verse sinus/lateral to midsagittal suture) 1.0–1.6/2.7–1.4/3.3. The same
viral tracers and injection volumes were used to trace inputs from the subcortical structures (in mm, posterior of bregma/lateral of midline/
below pial surface) lateral amygdala (LA; 1.8/3.55/3.7), dLGn (2.35/2.15/
2.55), and LP (1.85/1.25/2.65) in C57BL/6j or Chrm2-tdT mice. After
14–18 d of survival mice were killed and perfused with 1% PFA. The cor-
tex of V1–, dLGn-, and LP-injected mice was flattened, postfixed in
4% PFA, cryoprotected in 30% sucrose, and stored overnight with the
rest of the brain. In amygdala-injected mice, neocortex was separated
from the limbic region along the rhinal fissure, flattened, postfixed,
cryoprotected, and sectioned at 40 μm in the tangential plane on a freez-
ing microtome. The amygdala was sliced in the coronal plane. For visual-
mization of M2 expression in C57BL/6j mice, tangential, and coronal
sections through cortex were immunostained with an antibody against
M2. This procedure involved preincubation in blocking solution (0.1% Triton X-100, 10% normal goat serum, PB), followed by incubation in
primary rat anti-M2 antibody (1:500, for 48 h at 4°C; catalog #MAB367,
Millipore) and reaction with Alexa Fluor 647-labeled goat anti-rat sec-
ondary antibody (1:500; catalog #A21247, Thermo Fisher Scientific). M2
labeling patterns in Chrm2-tdT mice were identified by tdT fluores-
cence. Selected sections through cortex and amygdala were counter-
stained with Invitrogen NeuroTrace 435/455 (Thermo Fisher Scientific).
Anterograde labeling axons and M2 patches were imaged at 4–40×
under an epifluorescence microscope equipped with extended green flu-
orescent protein (EGFP), tdT, and infrared (IR; excitation, 650 nm;
emission, 665 nm) fluorescence optics, a CCD camera, and acquisition
software. In cortical sections, POR and PORa (anterior POR) were out-
lined based on M2 expression. Above-background EGFP or tdT fluores-
ence in POR, LM, and LI was determined in series of 2–12
tangential sections.

To disambiguate laminar identification based on cytoarchitecture
(i.e., counterstaining with NeuroTrace 435/455 (Thermo Fisher Scientific).
Anterograde labeled axons and M2 patches were imaged at 4–40×
under an epifluorescence microscope equipped with extended green flu-
orescent protein (EGFP), tdT, and infrared (IR; excitation, 650 nm;
emission, 665 nm) fluorescence optics, a CCD camera, and acquisition
software. In cortical sections, POR and PORa (anterior POR) were out-
lined based on M2 expression. Above-background EGFP or tdT fluores-
ence in POR, LM, and LI was determined in series of 2–12
tangential sections.

To investigate the tangential distribution of POR neurons that project to
the amygdala and the entorhinal cortex, we used retrograde tracing with
rAAV2-Retro-CAG-Cre (University of North Carolina Vector Core) in
A19 mice. Injections (46 nl) were made into the LA (in mm, posterior of
bregma/lateral of midline/below pial surface, 1.8/3.55/3.7) and the
ENTm (in mm, anterior to transverse sinus/lateral to midline/below the
pial surface, 0.7/5.25/3.55) angled 40° from vertical. After 14–18 d of sur-
face, perfusion with 1% PFA, flattening of cortex, postfixation in 4% PFA,
cryoprotection, and cutting in the tangential plane, sections were
immunostained with rat anti-M2 and Alexa Fluor 647-labeled goat anti-rat
antibodies. Sections were wet mounted on glass slides and imaged
under IR and tdT illumination at 4–40× magnification to identify M2
patches, cell bodies, dendrites, spines, and local axons (Vaughan
and Peters, 1973; Karimi et al., 2020) of retrogradely labeled cells in POR.

Patchy M2 expression
M2+ patches and M2− interpatches were delineated from images of
imunolabeling in C57BL/6j mice or the expression of Chrm2-tdT
using custom MATLAB scripts. M2 expression images were spatially
normalized by dividing the intensity of every pixel by the mean intensity
within a 100 μm radius. Images were blurred with the MATLAB “nan-
conv” function, a circular (pillbox) filter with 29 μm radius (see Fig. 6D),
and then divided into six quantiles based on pixel intensity (see Fig. 6E).
These borders were overlaid onto images of virally anterogradely labeled
axons or retrogradely labeled cell bodies and dendrites. Comparison of
M2 immunostaining and Chrm2-tdT expression yielded nearly identical
borders of M2+ patches and areal boundaries; the same analyses were
therefore used for both labels. This similarity was verified by performing
M2 immunostaining in Chrm2-tdT mice and overlaying M2 expression
patterns from both markers in the same section.

Experimental design and analyses
Data were obtained from 44 C57BL/6j, 7 Chrm2tdT, 2 PVTdT, and 8 A19
adult male (N = 36) and female (N = 25) mice.

Optical density of BDA labeling
We determined the weight of an-	erogradely BDA-labeled projections by using ImageJ to measure the pixel
values and compute the mean background-subtracted OD of axons
(three sections/target/mouse). The OD in a specific nucleus was normal-
ized to the sum of ODs measured in all of the 31 identified subcortical
projection targets and expressed as averages across injection sites and
mice. The relative strength of inputs from different visual areas to sub-
nuclei of the amygdala was determined by computing the OD in the tar-
get relative to the maximal OD of the injection site. The data were
plotted as the mean percentage OD of projections. The t test was used
for statistical comparisons.

Distribution and shape and of M2 + patches. For analysis of M2 patch
distribution and shape, patch borders were drawn around the highest
20% intensity pixels and lowest 80% intensity pixels in blurred, high-
pass-filtered M2 expression images of tangentially sectioned L1. A high-
intensity threshold for pixel inclusion was used to ensure the separation
of individual patches and to prevent the formation of large continuous
units. To find out whether patches were anisotropic and aligned with the
asymmetry of visual areas (see Fig. 2CD), we first drew the outline of
each area containing M2+ patches (V1, LM, LI, POR) based on M2 expres-
sion patterns in L4. These outlines were then used to determine the
major and minor axes of visual areas, using custom scripts and the
MATLAB “regroup” function. Patch anisotropy was quantified by the
major/minor axis ratio. The spatial density of patches was calculated as
units per square millimeter. Patches per square degree of visual space were
determined by multiplying spatial densities with magnification fac-
tor of respective area (Garrett et al., 2014, their Fig. 7). The Tukey’s test
was used for statistical comparisons of patches in different areas.

Projection strength.
For the quantification of the distribution of an-	erogradely labeled axonal projections in M2+ patches and M2− inter-
patches, M2 expression images of tangential sections through L1 were
high pass filtered, blurred, and divided into 6 intensity quantiles of equal
area (see Fig. 6D–F). The lower three quantiles were treated as M2+
interpatches and the upper three quantiles treated as M2+ patches. The
strengths of anterogradely labeled LA/basolateral amygdala (BLA) →
POR (N = 5), dLGn → POR (N = 5), and LP → POR (N = 2) projections
in M2+ patches and M2− interpatches of C57BL/6j and Chrm2tdT mice
were determined using a custom-written MATLAB script to measure the
pixel values of fluorescence intensity in ROIs (360–630 × 10 μm2) and
subtract the unlabeled background. For each projection target (POR,
LM, LI, V1), the background-subtracted mean pixel values in M2+
patches and M2− interpatches within the ROI were computed from one
to two sections through L1. The laminar distribution of inputs to
POR was compiled from sections through L2 to L6 in which registration
to the patch/interpatch pattern of L1 was not available. Mean intensities
in patches and interpatches and patch/interpatch intensity ratios were
compared across quantiles using Pearson’s correlation and one-sample t
test, respectively.

Distribution of dendrites. Images of dendrites labeled by retrograde
tracing with rAAV2-Retro-CAG-Cre in A19 mice and M2-immunola-
beled patches in L1 were acquired with a Leica Thunder Tissue Imager
(model DMB 6) equipped with a camera (model DFC 9000 GT, Leica)
and divided into six quantiles based on M2 expression intensity (see
above). Apical dendrites in L1/2 of retrogradely labeled pyramidal cells

patch/interpatch pattern of L1 was not available. Mean intensities
in patches and interpatches and patch/interpatch intensity ratios were
compared across quantiles using Pearson’s correlation and one-sample t
test, respectively.
in POR were distinguished from axons by greater thickness and were positively identified by their spiny morphology (Vaughn and Peters, 1973; Karimi et al., 2020). The dendrites were reconstructed at 40× magnification using NeuroLucida (MBF Bioscience), and their total length within each quantile of M2 expression was computed. For each quantile in each mouse, the proportion of dendrite length of all dendrites within that image was determined, and the Pearson correlation coefficient between M2 quantile level and the proportion of dendrites was computed. Images from four mice were analyzed for POR → ENTm projections and POR → LA projections.

**Laminar distribution of retrogradely labeled cells.** Series of flat-mounted tangential sections were taken from A9 mice in which rAAV2-Retro/CAG-Cre had been injected into either ENTm (N = 4 mice) or LA (N = 2 mice). Images of M2 expression and retrogradely tdT-labeled cells across layers were taken at 4× and 10× magnification and overlaid. In each image, borders for LM, LI, P, POR, and PORa were manually drawn based on M2 immunolabeling visualized with Alexa Fluor 647 IgG, and all tdT-labeled cells were marked. Custom MATLAB scripts were used to tabulate counts of cells within each area. Cell counts for unusable sections were interpolated using Akima piecewise cubic interpolation with the MATLAB `interp1` function. To compensate for variations in labeling effectiveness across cases, cell counts were normalized for each mouse by dividing by the maximum number of cells in any one area in any one section of that mouse. For each mouse, the ratio of total cells labeled within a group within deep layers (L5 and L6 pooled together) to total cells labeled within L2/3 was computed.

**Tangential clustering of retrogradely labeled cells.** To quantify the clustering of cells in M2- interpatches and M2- interpatches of POR, retrogradely labeled from LA (N = 2) and ENTm (N = 4), we analyzed in tangential sections through L2/3 aligned with M2-immunostained patches in L1. Images were taken at 10× magnification (ROI, 160 × 103 μm2) and divided into six quantiles based on M2 expression intensity. Cells were plotted manually and custom MATLAB scripts were used to count the number of cells in each M2 quantile. Cell counts were normalized within each image by dividing the cell count per quantile by the total number of cells labeled in that image. The Pearson correlation coefficient was then computed between quantile levels and normalized across multiple mice. Only images that contained >10 cells were analyzed.

**Data availability**

Code use for analysis in this study is available at https://github.com/ammueier/Meier-et-al._2021.

**Results**

**POR is contained within a M2+ region of parahippocampal cortex**

The borders of POR have been outlined previously by mapping of retinotopic inputs from V1 (Wang and Burkhalter, 2007). However, it remained unknown whether its borders align with the expression of M2, an effective marker used to identify cortical areas (Gâmãnuţ et al., 2018). Additionally, the original description of POR in the studies by Burwell (2001) and Beaudin et al. (2013) was derived from cytoarchitectonic features observed in coronal sections. This raised the question whether the borders seen in coronal sections and flatmounted cortex are aligned with one another. To find out, we combined multicolor viral tracing of retinotopic projections from V1 with the expression patterns of M2 and PVtdT. We then analyzed the axonal projections in tangential sections through the flattened cortical hemisphere. Reconstructions of the retinotopic map were done from serial tangential sections through the cortical mantle, eliminating the challenging task of aligning sequential coronal sections. In tangential sections through L4 stained with an antibody against M2, we identified V1 as an intensely immunopositive triangular area surrounded by the more weakly fluorescent extrastriate cortex (Fig. 1A, inset). Lateral to V1, M2 stained a boot-shaped region that overlapped with the ventral stream areas LM (lateromedial), LI (laterointermediate), POR and PORa (Gâmãnuţ et al., 2018). M2 expression dropped sharply at a M2+ strip along the rhinal fissure, which contained ectorhinal area (ECT) and the posterior area (P), and was continuous with the posterior temporal area (TEp), on the anterior–dorsal side of POR and PORa. A similar boot-shaped pattern was seen in the expression of PVtdT in PVcre x Ai9 mice (Fig. 1D), which overlapped with M2 (data not shown; Gâmãnuţ et al., 2018).

To outline the borders of areas within the M2+ boost, we simultaneously injected C57BL/6j mice (N = 5) with two antero-grade viral tracers, AAV.2/1.hSynapsin.EGFP (AAV.EGFP) and AAV.2/1.hSynapsin.tdTomo (AAV.tdT), at different visuo-topic locations of V1. The result of one such case in which AAV. EGFP was injected near the intersection of the vertical and horizontal meridian (Garrett et al., 2014) and AAV.tdT was targeted to the upper temporal visual field (Wang and Burkhalter, 2007) is illustrated in Figure 1A–C. It shows that axonal projections terminated in multiple pairs of red and green patches distributed across the surrounding extrastriate cortex (Fig. 1B,C). Anterodorsal to the rhinal fissure, we found three pairs of patches: two within the M2+ boot and one in the M2- strip. Reading the map from posterior to anterior starting at the border of V1, the colors of the projections switch from red to green, reverse back to red, and then reverse again to label a fainter patch of more interspersed red and green fibers. Because map reversals have been interpreted as areal borders (Sereno and Allman, 1991; Garrett et al., 2014), we concluded that the red-to-green/green-to-red and the green-to-red/red-to-green transitions mark the P/POR and the POR/PORa areal borders, respectively (Wang and Burkhalter, 2007; Wang et al., 2011). The systematic topographic shifts observed in five different mice suggest that P, POR, and PORa contain nearly complete maps of the visual field. The maps observed in two representative cases are depicted graphically in the inset of Figure 1B, in which the blue arrows indicate the progression from lower/nasal to upper/peripheral fields extracted from five different experiments. Together with previous findings that P, POR, and PORa have different connectivity profiles (Gâmãnuţ et al., 2018), these data suggest that these are distinct visual areas. We next asked whether each of these areas was contained within M2+ parahippocampal cortex or whether they extend across the border with ECT. We found that V1 injections at higher retinotopic altitudes shifted the anterogradely labeled patches in both POR and PORa closer to the borders with LM and LI, respectively (Fig. 1B, inset). In contrast, injections below the horizontal meridian pushed the projections backward, but never out of the M2+ region into the M2- strip. This topographic pattern shows that the POR/ECT and PORa/ECT borders coincide with the posterior margin of the M2+ region, and indicates that PORa belongs to M2- cortex and is not part of the agranular perirhinal area 36p, as suggested previously (Wang and Burkhalter, 2007).

According to this scheme, postrhinal cortex is completely contained within the M2+ boot. This raised the question of whether the agranular ventral POR (PORv), identified in coronal sections by Beaudin et al. (2013), is part of postrhinal cortex. To find out, we looked at the cytoarchitecture of postrhinal cortex in Chrm2-tdT mice (N = 2). Figure 1, E and F, shows a coronal section from the approximate location indicated by the green line on the flatmap in Figure 1A. It shows that the multilayered M2 expression in the dorsal part is interrupted suddenly by an unstained gap and then reappears as a simpler pattern in more ventral parts of the section. In tangential sections, the unstained gap coincides precisely with the M2- band aligned with the
rhinal fissure and overlaps with agranular cortex that lacks L4 and shows a much thinner L5 than in M2 cortex (Fig. 1F). This cytoarchitecture suggest that the M2 gap represents ECT (Dong, 2008) and that in coronal sections the gap corresponds to the M2 band seen in tangential slices. Thus, agranular PORv lies in ECT and is not part of V1-recipient, M2 postrhinal cortex.

**M2** patches in higher visual areas are linked to cortical magnification

We have shown previously that alternating M2 and M2 patches are not unique to V1, but is also observed throughout the thickness of L1, most notably at the L1/2 border in LM, LI, POR, PORa, auditory (AUD), and retrosplenial (RSP) cortex (Ji et al., 2015). In addition, we noted that in each of these areas M2 expression is more uniform in layers below the L1/2 border (Fig. 2A,B). In V1, M2 patches parcellate the plexiform L1 in the tangential plane into constant size domains, which extend to the cellular L2/3, and in groups of four to eight represent the visual point image (Ji et al., 2015). This organization raised the question of whether the shape, density, and cortical magnification factor of M2 patches is related to the layout of the visuotopic maps in LM, LI, and POR. We determined the aspect ratio of M2 patches along the major and minor axes of V1, LM, LI, and POR measured from the study by Garrett et al. (2014, their Fig. 7; see Materials and Methods). We found that in all four areas, average patch anisotropy was aligned with the axes of the
containing area (Fig. 2D). In areas V1, LI, and LM, whose major axes represent altitude, patches were stretched along elevation. In POR, with the major axis mapping azimuth, average patch elongation occurred along latitude. In V1, LM, and LI, patch dimensions closely matched those of their containing area [with major:minor length for patches; V1, 1.21 ± 0.03 (12 mice, 1354 patches); LM, 1:1.4 ± 0.05 (7 mice, 122 patches); LI, 1:1.42 ± 0.08 (7 mice, 78 patches); POR, 1:2.1 ± 0.03 (16 mice, 278 patches)] and areas (V1, 1:1.22; LM, 1:1.53; LI, 1:1.53; POR, 1:1.59; Garrett et al., 2014). These results suggest that anisotropies in the representation of retinotopic space are shared by the M2⁺ patches within the areas we have examined.

Next, we determined the spatial distribution of M2⁺ patches and found that the density was similar in V1, LM, LI, and POR (p > 0.005, Tukey’s test; Fig. 2E). These results indicate that the patch size (top 20% of M2⁺ pixels) is constant across areas, as was the case across different quadrants of V1 (Ji et al., 2015). How much of the visual field is represented by a M2⁺ patch in a given area? To find out, we multiplied patch density (patches per square millimeter) by the cortical magnification factor (square millimeters per square degree; Garrett et al., 2014) and obtained a quantity patches per square millimeter in V1, LM, LI, and POR. We found that a 10 × 10⁴-wide field of visual space, which accounts for about the size of the point image in V1 (12 mice)
and LM (7 mice; D’Souza et al., 2020), contains 3.93 ± 0.08 and 3.46 ± 0.23 patches/mm², respectively. In LI (7 mice) and POR (16 mice), a similar 10 × 10°-wide field contained about half the number of patches, indicating that a single LI or POR patch represents a proportionately larger amount of visual space than a patch in V1 of LM (p < 0.001, Tukey’s test; Fig. 2F).

Anterogradely labeled outputs from visual cortex to amygdala, thalamus, brainstem, and striatum

Despite massive efforts, there are no detailed descriptions of the connectivity of visual cortical areas with the amygdala in mice, except for POR (Burgess et al., 2016). In previous studies (Oh et al., 2014; Zingg et al., 2014; Hintiryan et al., 2019), injections involved multiple areas, and areal assignments were based on the computed majority stake, which is known to underestimate the connectome density (Gâmanuț et al., 2018). We have taken a different approach in which we only accepted clean hits of the source area for anterograde tracing with BDA. To do this, we relied on the complementary patterns of M2 expression and callosal connections (Wang et al., 2011). Callosal connections were labeled by retrograde tracing with bisbenzimide and visualized in situ in fixed brains imaged under a fluorescence stereomicroscope. This procedure also revealed the injection site as a bright spot, which was based on the distinctive callosal projection pattern, readily assigned to V1, PM (posteromedial), AM (anteromedial), A (anterior), RL (rostromedial), AL (anterolateral), LM, LI, P, or POR (Wang and Burkhalter, 2007; Fig. 3A–J). The cortex dorsal to the rhinal fissure was then separated from the

Figure 3. Axonal projections of visual cortex → amygdala pathways. A–J, In situ ex vivo images of callosal connections traced by retrograde transport of bisbenzimide from the posterior third of the right (contralateral) hemisphere. The arrows point to bright spots that represent the BDA injection sites in V1, LM, P, LI, POR, AL, RL, PM, AM, and A of the left (ipsilateral) hemisphere. A′–J′, Dark-field images of coronal sections showing anterogradely BDA-labeled axonal projections (yellow) from V1, ventral (LM, P, LI, POR), and dorsal (AL, RL, PM, AM, A) stream areas terminating in different nuclei of the amygdala. Note that the white structures represent unlabeled, highly reflective myelinated fibers of the internal capsule (int). Scale bars: A–J, 1 mm; A′–J′, 0.5 mm.
brain, flattened, cut tangentially, reacted for M2 immunofluorescence, and stained with an ABC reaction to reveal the BDA injection site. Injections were considered clean hits if they were confined within the areal boundaries delineated by Gămanuț et al. (2018). The rest of the brain was cut coronally, and sections containing the amygdala were stained to reveal BDA-labeled axonal projections. We found that the projections from V1 and the dorsal stream areas, AL, RL, PM, AM, and A, were extremely weak and too sparse for determining their optical density (Fig. 3F′–J′). In sharp contrast, the projections from the ventral stream areas, LM, P, LI, and POR, were much denser (Fig. 3A′–E′). The vast majority (98–100%) of inputs from LM, LI, and P to the amygdala terminated in LA with only minor offshoots to BLA (basolateral amygdala; Fig. 3B′–D′). Projections from POR were percentage-wise (percentage of total input to amygdala and percentage of optical density relative to injection site) the strongest in LA, but, compared with LM, LI, and P, a larger fraction terminated in BLA and central amygdala (CeA; Fig. 4B–E, inset).

Compared with the total output from 10 visual cortical areas to 31 subcortical targets, the weight of amygdala projections was weak, and even the strongest combined input from POR to LA and BLA was only ~10% of the total strength (Fig. 4A–J). Notably, inputs from ventral areas to LA were significantly stronger than from dorsal areas (Fig. 4K). POR input to another emotion regulatory center for fear, the dorsal lateral periaqueductal gray (DLPAG), which responds to cortical stressors and, unlike the amygdala, does not receive visceral input (Floyd et al., 2000), was only ~2% (Fig. 4E). Similarly sparse projections to DLPAG were found from P, AM, and PM (Fig. 4D,H,I).

Bennett et al. (2019) have shown that dorsal stream areas are more weakly connected to posterior LP than areas of the ventral stream. We found a similar rostrocaudal distribution by anterograde BDA labeling of outputs from nine higher visual areas. In our data, however, ventral stream projections include the rostral LP, the primary target of the dorsal stream. The strong preference for posterior LP [LP, latero-caudal (LPLC); LP, mediocaudal (LPMC)] was in the present inputs from ventral stream areas LM, LI, P, and POR (Fig. 4B–E), which was in striking contrast to the absence of input from the dorsal stream, AL, RL, AM, PM, and A (Fig. 4F–J). Inputs to the middle and anterior part of LP [LP, laterostral (LPLR); LP, mediorstral (LPMR)], including the widely connected lateral dorosomal nucleus, dorsomedial, and laterodor sal nucleus, ventrolateral (lateraldor sal thalamic nuclei; Juavine tt et al., 2020), originated from both ventral and dor sal stream areas (Fig. 4B–J). Unexpectedly, we found that POR sent weak outputs to dLGN, from which it also received weak inputs (Figs. 4E, 5D,E). Notably all of the retrogradely DY-labeled cells found in the dLGN were located in the shell (Fig. 5D,E). This suggests that intracortical communication through transthalamic circuits (Sherman, 2016) may involve the higher-order LP as well as the first-order dLGN. Weak POR projections were also found to the auditory medial geniculate (MGN), and marginal zone of medial geniculate (Fig. 4E). Unlike dorsal stream areas, which more strongly project to oculomotor and motor coordination centers than ventral stream areas, such inputs were extremely weak or absent from POR (Fig. 4E). A notable exception was the strong POR inputs to the dorsomedial caudoputamen (Fig. 4E), which plays a role in goal-directed behavior (Hunnicutt et al., 2016).

Retrogradely labeled sources of inputs to POR
To map the sources of inputs to POR, we retrogradely traced neurons in PVδ+ mice with DY (N = 2). Unsurprisingly, given the results from anterograde tracing (Fig. 4E), we found DY-labeled cells in LA, LP, and dLGN (Fig. 5D–G). Of a total of 2454 DY-labeled neurons in the primary and secondary visual thalamus, the vast majority were found in different subdivisions of the LP (LPLC, 54.2 ± 6.1%; LPMC, 6.4 ± 1.9%; LPLM, 25.3 ± 3.4%; LPMR, 11.7 ± 2.0%). Only a small percentage of labeled cells was encountered in the dLGN (2.4 ± 1.8%), notably all of them in the dLGN shell (Fig. 5D,E). This shows that POR receives retinal input via a direct dLGN → POR and an indirect SC → LP → POR (Beltramo and Scanziani, 2019) thalamocortical pathway. In addition, we found DY-labeled POR projecting thalamic neurons in the zona incerta, suprageniculate, dorsal anteromedial, central medial thalamic nucleus, paratethalamic nucleus, and MGV (data not shown). Limbic regions contained DY-labeled cells in LA; medial (ENTm) and lateral (ENTi) entorhinal cortex (Fig. 5B,C,G). Each of these structures received strong inputs from POR (Figs. 3E, 4E; Wang et al., 2012), suggesting bidirectional communications among POR, amygdala (confirming the findings of Burgess et al., 2016), and entorhinal cortex (Fig. 6A–C; also see Figs. 8A–C,E–G, 9A–C,E–G). DY-labeled POR projecting neurons were also encountered in multiple neocortical areas including the following: V1; PM; MM (mediomedial area), RSP (retrosplenial area); AM; RL; AL; LM; LI; P; ECT; TEP; TEA (anterior temporal area); DP (dorsal posterior area), AUD, Alv (ventral anterior insula); GU (gustatory area), ORBl (lateral orbitofrontal area), ILA (anterior infralimbic area), and ACAv (ventral anterior cingulate area) (Fig. 5A–C).

Anterogradely labeled projections from amygdala to POR terminate in M2 interpatches
It has been known from studies in rats and mice that projections from LA and BLA to POR terminate in L1–3 with weaker input to the layers below (Pikkarainen and Pitkänen, 2001; Burgess et al., 2016). To find out whether these inputs are modularly organized, we traced inputs from LA and BLA (Fig. 6b, inset) with AAV,EGFP or AAV-tdTom in C57BL/6 and Chr12-tdTomice. We found that the laminar distribution of inputs from both sources was similar, which let us pool (LA/BLA, N = 5) the baseline-subtracted fluorescence intensity of axonal projections. Inputs were densest in L1, weaker in L3/4, and sparse in L2, L5, and L6 (Fig. 7E). To study whether inputs specifically target M2 patches or M2’ interpatches, we removed the cortex dorsal to the rhinal fissure, flattened the tissue, cut tangential sections, and sliced the amygdala in the standard coronal plane. Projections to L1 of POR were strikingly nonuniform, preferentially targeted M2’ interpatches, and largely avoided M2’ patches (Fig. 6A–C). We performed an automated delineation of M2’ patch borders with custom MATLAB scripts, using a procedure similar to that used by Sinich and Horton (2005). M2 intensity images were high-pass filtered and blurred, then partitioned into six quantiles based on fluorescence intensity (Fig. 6D,E). To test for different input strengths to patches and interpatches, EGFP intensity was compared across M2-defined quantiles (Fig. 6F). EGFP intensity showed a strong negative correlation with M2 expression (R = −0.86, p = 10−8, N = 5; Pearson correlation), and projection strength in the highest quantile (6) was a mere 17% of that in the lower quantile (1), indicating that M2’ interpatches received significantly stronger input from the LA/BLA than M2’ patches (Figs. 6F, 7F).
Anterogradely labeled projections from dLGN and LP to POR terminate in M2 patches

We have found recently that M2 expression in V1 divides L1 into interdigitating modules receiving dLGN input to M2$^+$ patches and LP input to M2$^-$ interpatches (Ji et al., 2015; D’Souza et al., 2019). Both of these thalamic nuclei are also known to project to extrastriate visual cortex (Oh et al., 2014), but only LP has been shown unequivocally to project to POR (Zhou et al., 2017; Beltramo and Scanziani, 2019; Bennett et al., 2019; Juavinett et al., 2020). Here, we investigated whether
dLGN and LP project to POR and whether inputs to L1 show an interdigitating pattern similar to the one we have found in V1 (D’Souza et al., 2019). For this purpose, we injected the anterograde viral tracers (AAV.EGFP, AAV.tdTomato) into either dLGN or LP (Fig. 7A-D, inset). We found that input from dLGN (N = 5) and LP (N = 2) terminated in POR and PORa (Fig. 7A–D). In two cases with a complete series of sections across cortex, we found that dLGN input was densest in L1 and L5, moderate in L4 and L6, and weakest in L2/3 (Fig. 7E). LP input was dense in L1 and L4, moderate in L5 and L6, and weakest in L2/3 (Fig. 7E). To determine the patch/interpatch ratio of input to L1, the top three M2 quantiles were taken as M2+ patches and the bottom three quantiles as M2− interpatches. The results show that the tangential distribution of dLGN and LP inputs to L1 of POR and PORa overlapped with M2+ patches (patch/interpatch ratios: dLGN → POR, 3.04 ± 0.67, N = 5, p < 0.01, t test; LP → POR, 3.51 ± 0.42, N = 2, p < 0.01, t test; Fig. 7F). The targeting in POR and PORa to M2+ patches distinguished dLGN and LP input from amygdala inputs, which terminated in M2− interpatches (Fig. 6C,F). EGFP intensity positively correlated with M2 quantile (LP → POR, p < 0.001, r = 0.83, N = mice; dLGN → POR, p < 10−11, r = 0.91, N = 5 mice, Pearson correlation). dLGN and LP inputs to LM and LI likewise targeted M2+ patches and avoided M2− interpatches (Fig. 7F; patch/interpatch ratios: dLGN → LM, 2.45 ± 0.59, N = 4, p < 0.01, t test; LP → LM, 2.11 ± 0.71, N = 3, p < 0.05, t test; dLGN → LI, 2.09 ± 0.49, N = 4, p < 0.01, t test; LP → LI, 2.1 ± 0.75, N = 3, p < 0.05, t test). In agreement with previous studies (D’Souza et al., 2019), dLGN inputs to V1 were preferentially targeted to M2+ patches, whereas LP inputs preferred M2− interpatches (data not shown). Inputs from V1 → POR, while topographical (Fig. 1B,C), showed no clear preference for either compartment (patch/interpatch ratio: 1 ± 0.03, N = 4, p = 0.88, t test; Fig. 7F).

Cell bodies and apical dendrites of POR → amygdala and POR → ENTm projecting neurons are aligned with M2+ interpatches

The striking specificity of amygdala → POR axonal projections for M2+ interpatches suggested that these inputs may target apical dendrites of long-range projecting neurons in L1. Of particular interest were POR neurons projecting to downstream targets such as the amygdala and ENTm, which are known for their roles in affective processing (Phelps and LeDoux, 2005), memory, and spatial navigation (Buzsáki and Moser, 2013), respectively. To label the complete dendritic arbor of such neurons, we performed retrograde viral tracing from LA/BLA and ENTm in Ai9 mice with AAV2retro-CAG-Cre. M2+ patches were identified by immunostaining for M2 and dividing fluorescence intensity into quantiles. The results show that LA/BLA and ENTm injections retrogradely labeled pyramidal cell dendrites, somata, and occasionally their local axonal projections in POR (Fig. 8A–G), PORa, P, ECT, LM, and LI (Fig. 8I,F).

Dendrites were readily distinguished from axons by the greater thickness and positively identified based on the presence of spines (Vaughan and Peters, 1973; Karimi et al., 2020). Module preference of apical dendrites in L1 was quantified by reconstructing branches in three to four ROIs of the top two sections through POR. An example is shown in Figure 8D. Most strikingly, we found that apical dendrites of POR → LA/BLA projecting cells branched preferentially in M2+ interpatches of L1 (Fig. 8E–G). Similar results were obtained for POR → ENTm projecting neurons.
Figure 6. Axonal projections of lateral amygdala → POR pathway. A, Tangential section through L1 of flattened POR in Cnr2 tdTom mouse showing nonuniform pattern of M2+ patches (magenta) and M2- interpatches (black). B, Patchy axonal projections to L1 labeled by anterograde tracing with AAV2/1.HSyn.EGFP from the LA. Coronal section of injection site in LA (B, inset). C, Overlay of A and B shows that LA → POR projection to L1 terminates preferentially in M2+ interpatches. D, Heat map of M2 expression in POR. E, Partitioning of heat map (D) into six equal-area quantiles (different shades of gray) for the delineation of M2+ patches and M2- interpatches. F, Normalized average ± SEM (N = 5 mice) EGFP fluorescence intensity in each of the six quantiles shows that LA → POR inputs are preferentially associated with M2+ interpatches (R = −0.86, p < 10−5, Pearson correlation).

In cases in which complete z-stacks of sections were available, somas and retrogradely labeled POR → ENTm (N = 2) and POR → LA/BLA (N = 2) cells were found in L2–6 of POR (Fig. 8I,J). Similar size injections into ENTm and LA consistently labeled more cells in POR than in any other area. LM → ENTm and L1 → ENTm neurons were mostly confined to L2/3. P → ENTm, PORa → ENTm, and POR → ENTm cells were biased to deep layers with a preference for the L5/6 border (Fig. 8I). The ratio of total labeled POR → ENTm L2/3 cells to L5/6 cells within a given case was 1:4.8 ± 2.5 (N = 2, p < 0.042, t test). Deep layers were also the preferred source of cells projecting to LA/BLA (L2/3-to-L5/6 ratio, 1:2.8 ± 0.1 (N = 2, p < 10−3, t test).

Unlike POR → ENTm cells, POR → LA/BLA neurons were distributed across the thickness of L5 (Fig. 8I,J), suggesting that the projections originate from distinct cell types. We quantified the alignment of retrogradely labeled L2/3 neurons (sections across L4–6 were not available for analysis) with M2+ patches and M2- interpatches by overlaying M2 expression images with labeled cells and determining cell counts in each of six M2 quantiles. Cell counts were normalized by dividing the cell count within each quantile by the total number of labeled cells in that image. POR → ENTm cells showed a strong preference for aligning with M2+ interpatches (Fig. 9A–D; R = −0.86, p < 10−5, N = 4 mice, 132 total cells, Pearson correlation). POR → LA/BLA cells also showed a significant preference for aligning with M2+ interpatches, though with less specificity than POR → ENTm cells (Fig. 9E–H; R = −0.63, p < 0.027, N = 2 mice, 53 cells, Pearson correlation).

Input from POR to hippocampus strongly biased to dorsal CA1
We have shown previously that POR provides input to CA1 of the hippocampal formation (HPF) but have not described the topographic organization of this pathway (Wang et al., 2012). Here, we have retraced the connections anterogradely with AAV2/1.HSyn.EGFP from POR in two mice. The case illustrated in Figure 10 shows a tangential section through flattened cortex. The injection site was assigned to POR by location and clearly spares PORa (Fig. 10A). To physically flatten the HPF (for complete graphical flattening, see Swanson and Hahn, 2020), it was separated from the external capsule along the alveus, flipped backward so that the dorsomedial pole came to lie at the posterior edge of the section (Fig. 10B, inset). The green-labeled axonal projection was densest in the dorsal part of CA1 and gradually lost strength toward the ventral pole (Fig. 10B). A similar nonuniform distribution of projections was found when the injection involved POR and PORa, but was confined to the M2+ region (data not shown). This suggests that either the projections from POR and PORa are similar or that input from PORa was undetectable. Regardless, the results show that POR input is strongly biased to the dorsomedial part of the HPF. The results further show that POR → CA1 input can either be direct, bypassing ENTm, or indirect, by way of ENTm (Wang et al., 2012). Both of these pathways provide visuospatial information (Fanselow and Dong, 2010), which through the reciprocal POR–amygdala network (Figs. 6, 8E–H) receives affective input to influence spatial navigation.
Discussion

Postrhinal cortex contains two modularly organized areas

We have found that postrhinal cortex contains two separate areas, POR and PORa, with distinct visuotopic maps and connection profiles (Gământă et al., 2018). Both areas have granular cytoarchitectures, are contained within an M21 region, and have borders with the surrounding M2 areas of ECT and TEp. We found no support for a ventral PORv (Beaudin et al., 2013), which based on its agranular cytoarchitecture and lack of M2 expression may correspond to rodent ECT or primate parahippocampal cortex TH (Burwell, 2001). The POR/PORA border was revealed by a transition from the orderly map in POR to the more dispersed but spatially clustered connectivity in PORa. A similar map reversal exists between primate parahippocampal cortices TFO and TF (Ungerleider et al., 2008), suggesting that the areal organization of the PV- and M2-expressing parahippocampal cortex (Saleem et al., 2007) in rodents and primates may be homologous.

A striking feature of POR, PORa, LM, and LI is the nonuniform tangential distribution of M21 patches and M2–interpatches in L1. Patchiness is a known attribute of primate extrastriate cortex (Kaas, 2012) and has also been shown to exist in the intrinsic connectivity of TF (Lavenex et al., 2004). The patchiness in POR, PORa, LM, and LI resembles the pattern we have seen in mouse V1 (Ji et al., 2015). However, unlike the quasi-isotropic modules in V1, patches in higher visual areas

Figure 7. Axonal projections of dLGN → POR and LP → POR pathways. A, C, Immunostained M21 patches in L1 of flatmounted tangential section through POR and PORa. B, dLGN → POR axonal projections traced anterogradely with AAV2/1.hSyn.EGFP. B, inset, Coronal section of injection site spreading across core and shell of dLGN. White contours indicate borders between fourth and fifth intensity quantities (of six) of M2 expression (A). Overlay of contours shows overlapping patterns of M21 patches and dLGN → POR projections (B). D, AAV2/1.hSyn.EGFP-labeled LP → POR projections to M21 patches (C) in L1 of POR. D, inset, Coronal section of injection site in LP. E, Laminar distribution of anterogradely labeled LA/BLA → POR, dLGN → POR, and LP → POR projections. F, Mean ± SEM patch: interpatch EGFP intensity ratio of dLGN → POR, LP → POR, dLGN → LM, LP → LM, dLGN → LI, LP → LI, LA/BLA → POR, and V1 → POR axonal labeling in L1 of the projection target. **p < 0.01; *p < 0.05; ns, not significant (one-sample t test).
were anisotropic and showed aspect ratios similar to the overall shape of the area (Garrett et al., 2014) in which they reside. Notably, we found a reduced (relative to V1) magnification factor per patch in LM (12.5% smaller), POR (37.5% smaller), and LI (50% smaller), indicating that a module in each of these areas represents a larger portion of visual space than in V1. The results show that the magnification factor decreases as the receptive field size increases from V1 → LM → LI → POR (D’Souza et al., 2020; Siegle et al., 2021), indicating that at the apex the product (i.e., point image) is represented by the smallest number of modules and remains constant. The findings further indicate that inputs from diverse retinotopic locations converge and that individual patches are processing units that integrate a broader set of stimulus attributes inherited from intracortical and subcortical sources (Beltramo and Scanziani, 2019; Harris et al., 2019; D’Souza et al., 2020).

POR contains separate module-specific circuits with dLGN, LP, amygdala, and ENTm

Our anterograde tracings with BDA have shown that only the ventral stream areas LM, LI, P, and POR provide input to the amygdala. No such connections were found from V1 and the dorsal stream areas AL, RL, A, AM, and PM. Conceptually similar results were reported in rat (McDonald and Mascagni, 1996). Together, the results suggest that only higher visual areas involved in encoding of objects and the context in which they appear have direct connections to the amygdala, while areas that play a role in visually guided actions (Jin and Glickfeld, 2020) do not. From this simplified perspective, it is easy to overlook that POR is highly interconnected with dorsal and ventral stream areas (Wang et al., 2012; Gâmnăuț et al., 2018). Through these connections, POR combines task-specific information across streams,
Figure 9. Modular distribution of retrogradely labeled cell bodies of POR → LA/BLA and POR → ENTm projecting neurons in L2/3 of POR. A, B, Tangential section through flatmounted POR showing nonuniform distribution of somata (false-colored green) of POR → ENTm projecting L2/3 neurons labeled by retrograde tracing with rAAV2-Retro/CAG-Cre in Ai 9 mice (A) overlaid with patchy expression of M2 immunostaining in L1 (B). C, Overlay of A and B shows that cell bodies and basal dendrites are preferentially aligned with M2− interpatches. D, Normalized cell counts (mean ± SEM) of POR → ENTm in M2 quantiles (shades of gray) of POR. The fraction of total cell bodies shows a strong preference for M2− interpatches. E–G, Tangential section through flatmounted POR showing nonuniform distribution of somata (false-colored green) of POR → LA/BLA projecting L2/3 neurons labeled by retrograde tracing with rAAV2-Retro/CAG-Cre in Ai 9 mice (E) overlaid with patchy expression of M2 immunostaining in L1 (F). G, Overlay of E and F shows that cell bodies are preferentially aligned with M2− interpatches. H, Normalized cell counts (mean ± SEM) of POR → LA/BLA in M2 quantiles of POR. The fraction of total cell bodies shows a preference for M2− interpatches.
associates objects with locations (i.e., context; Furtak et al., 2012), and, through interactions with the amygdala, assigns affective credits to neurons that guide behavior (Bucci and Burwell, 2004; Burgess et al., 2016). Notably, only POR sends significant input to the BLA nucleus, which projects onward to the CeA (Pitkänen et al., 1995) and is the major driver of behavioral responses from the amygdala (Tye et al., 2011). Thus, it appears that POR has more direct control over visually related affective behaviors than areas LM, LI, and P, whose projections are weak and largely confined to LA.

The patchy pattern of M2 expression we have found in L1 of areas LM, LI, POR, and PORa reveals that the modular organization of V1 (Ji et al., 2015) is preserved across visual areas of the ventral stream (Wang et al., 2012). Modularity, in the sense used here, refers to the columnar organization of stimulus-selective responses and the spatial clustering of projection neurons within an area (da Costa and Martin, 2010). That such constraints emerge from the segmentation of the plexiform sheet of L1 has only been shown in V1, where a patchy pattern of M2 expression label sites of thalamocortical and intracortical synaptic input to apical dendrites of pyramidal neurons (Ji et al., 2015; D'Souza et al., 2019). We have found that similar patchy patterns in L1 organize the modularity of amygdalocortical and thalamocortical inputs to POR, PORa, LM, and LI. The results show that of all layers the LA innervates L1 most strongly and preferentially targets M2 patches. Thalamic inputs from dLGN and LP to POR, PORa, LM, and LI not only differ from amygdalocortical afferents in their laminar distribution, but also in their preferential targeting of M2 patches in L1. The distinctive connectivity patterns suggest that amygdalocortical and thalamocortical inputs to L1 target apical dendrites of separate populations of POR and PORa neurons, whose cell bodies are located in the layers below. Spatial clustering of pyramidal cell dendritic tufts in L1 are known features of rat V1 and retrosplenial cortex (Ichinohe and Rockland, 2002; Ichinohe et al., 2003). Here, we demonstrate directly that the apical dendrites of POR → amygdala projecting neurons preferentially terminate in M2 interpatches. Thus, it is tempting to speculate that pyramidal cells in POR with dendrites in M2 interpatches may be modulated by affective information from the amygdala (Bucci et al., 2000;
Burgess et al., 2016), which is driven directly by visual input from LP (Wei et al., 2015). In contrast, responses of POR neurons with dendrites in M2+ patches may be influenced preferentially by direct locomotion-modulated inputs from the dLGN shell and LP (Marshall et al., 2012; Roth et al., 2016; Beltramo and Scanziani, 2019; Bennett et al., 2019) rather than by inputs ascending through the hierarchical cortical network originating in V1 (Beltramo and Scanziani, 2019; D’Souza et al., 2020).

The overlap between dendrites of POR output neurons with inputs from the amygdala suggests that the amygdala and POR are interconnected through a reciprocal loop, as has been shown directly by Burgess et al. (2016). It seems likely that these reward-modulated responses are mediated by synaptic contacts in L1. Tagging sensory stimuli with affective significance imported from the amygdala is not the only influence on the multimodal response profile of POR neurons, which are also sensitive to the spatial context in which objects appear (Furtak et al., 2012). This information may derive from dorsal stream inputs (Wang et al., 2012; Gâmânûţ et al., 2018) and is sent to multiple downstream targets (Agster et al., 2016), including the ENTm, with which we have found POR to be reciprocally connected. Similar to POR → amygdala projecting cells, the apical dendrites of POR → ENTm neurons terminate preferentially in M2+ interpatches. These dendrites may belong to POR → LA/BLA and POR → ENTm cells or belong to the same neurons with branching projections to both the amygdala and ENTm. Regardless, the results suggest that some POR neurons with dendrites in M2+ interpatches send affect-modulated outputs to ENTm, where they may enhance the emotional salience of landmark information and unexpected objects in the external world (Knierim et al., 2014), and guide defensive responses to visual threats (Bucci and Burwell, 2004; Wei et al., 2015). In addition to this indirect pathway, POR also provides direct input to the dorsal HPF, suggesting that it serves visuospatial processing for environmental exploration (Fanselow and Dong, 2010). Although the modular origin of POR → HPF neurons remains unknown, the sparse innervation of intermediate and ventral HPF suggests that the input may carry little affective information (Fanselow and Dong, 2010).

In contrast to amygdala → POR projections, we found that dLGN → POR and LP → POR inputs overlap with M2+ patches of L1. Retrograde labeling from POR confirmed that dLGN inputs originate from shell neurons, which are known to receive afferents from the retina and the SC (Dhande and Huberman, 2014; Morin and Studholme, 2014; Bickford et al., 2015; Martersteck et al., 2017) and, like the SC → LP → POR pathway, convey visual motion information to its target (Cruz-Martín et al., 2014; Beltramo and Scanziani, 2019; Bennett et al., 2019). Inputs from matrix-type thalamic neurons to dendrites in L1 have been shown to elicit spiking in pyramidal cells of rat barrel cortex (Cruikshank et al., 2010) and may possess similar synaptic strength in POR. Both pathways carry locomotion signals, in which running speed is either synchronized or desynchronized with optic flow motion (Roth et al., 2016). Synchronous convergent thalamocortical inputs, heavily biased from LP, to M2+ patches in POR may mostly signal moving objects at speeds different from the motion of the animals. These direct thalamic inputs to apical dendrites in M2+ patches of POR may increase the sensitivity of POR neurons (Takahashi et al., 2016) to nonthreatening moving stimuli independent of the hierarchical cortical circuitry (Beltramo and Scanziani, 2019). Future studies are necessary to determine to which circuit M2+ patch-aligned POR cells belong (Fig. 11).

References


