

Molecular, Cellular, and Developmental Organization of the Mouse Vomeronasal Organ at Single Cell Resolution

Max Hills Jr¹., Limei Ma¹, Ai Fang¹, Thelma Chiremba¹, Seth Malloy¹, Allison Scott¹, Anoja Perera¹, and C. Ron Yu^{1, 2, *}

- 1 Stowers Institute for Medical Research, 1000 E. 50th Street, Kansas City, MO 64110, USA
 - 2 Department of Cell Biology and Physiology, University of Kansas Medical Center, 3901 Rainbow Boulevard, Kansas City, KS 66160, USA.
- * Correspondence: ryu@stowers.org

SUMMARY

We have generated single cell transcriptomic atlases of vomeronasal organs (VNO) from juvenile and adult mice. Combined with spatial molecular imaging, we uncover a distinct, previously unidentified class of cells that express the vomeronasal receptors and a population of canonical olfactory sensory neurons in the VNO. High resolution trajectory and cluster analyses reveal the lineage relationship, spatial distribution of cell types, and a putative cascade of molecular events that specify the V1r, V2r, and OR lineages from a common stem cell population. The expression of vomeronasal and olfactory receptors follow power law distributions, but there is high variability in average expression levels between individual receptor and cell types. Substantial co-expression is found between receptors across clades, from different classes, and between olfactory and vomeronasal receptors, with nearly half from pairs located on the same chromosome. Interestingly, the expression of V2r, but not V1r, genes is associated with various transcription factors, suggesting distinct mechanisms of receptor choice associated with the two cell types. We identify association between transcription factors, surface axon guidance molecules, and individual VRs, thereby uncovering a molecular code that guides the specification of the vomeronasal circuitry. Our study provides a wealth of data on the development and organization of the accessory olfactory system at both cellular and molecular levels to enable a deeper understanding of vomeronasal system function.

INTRODUCTION

In many terrestrial species, the vomeronasal organ (VNO) is dedicated to the detection of inter- and intra-species chemosensory cues¹⁻⁵. Detection of these cues triggers innate neuroendocrine responses and elicits stereotypic social and reproductive behaviors^{1,2,6-16}. The VNO shares a developmental origin with the main olfactory epithelium (MOE), which detects the odor world at large and allows associative learning to take place. Both develop from the olfactory placode during early embryogenesis^{17,18}, but the two sensory organs follow different developmental trajectories to establish distinct characteristics in morphology, cellular composition, and molecular features. Single cell atlases of the MOE have been generated to reveal astonishing details in its molecular composition and developmental trajectories¹⁹⁻²³. In contrast, transcriptomic information of the VNO is rather limited^{24,25}. In this study, we generate single cell atlases of developing and adult mouse VNO to answer fundamental questions about the VNO.

The vomeronasal sensory neurons (VSNs) express three families of G-protein coupled receptors, the V1rs, V2rs, and the formyl peptide receptors (Fprs)²⁶⁻³². With more than 400 members, the vomeronasal receptors are among the fastest evolving genes³³⁻³⁸. Signaling pathways in the VNO include the G₁₂ and G_o proteins, and a combination of Trpc2, Girk1, Sk3, and Tmem16a ion channels, to transduce activation of the vomeronasal receptors³⁹⁻⁵⁹. The spatially segregated expression of G₁₂ and G_o, as well as the VR genes suggests two major classes of neurons. Our study reveals new classes of sensory neurons, including a class of canonical olfactory sensory neurons (OSNs) in the VNO. We further determine the developmental trajectories of the separate lineages and the transcriptional events that specify the lineages.

Pheromones are highly specific in activating the VSNs^{5,60-65}. Previous studies have shown that VSNs residing in the apical layer of the VNO express V1R (Vmn1r) genes in a monoallelic manner^{63,66,67}, whereas the basal VSNs express one of the few broadly expressed V2R (Vmn2r) genes, plus a unique V2R gene belonging to different clades²⁶⁻³¹. Although these results suggest that receptor expression in the VNO conforms to the “one neuron one receptor” pattern as found in the MOE, the mechanisms that control receptor expression are unknown. Here, we find substantial co-expression of vomeronasal receptors, and of vomeronasal and odorant receptors. Moreover, our analyses indicate that selection of V1R expression likely results from stochastic regulation as in the OSNs, but V2R expression likely result from deterministic regulation.

Finally, we address the molecular underpinning of how VSNs establish anatomical connections to transmit sensory information. VSNs expressing the same receptor project to dozens of glomeruli in the AOB^{67,68}. Individual stimuli activate broad areas in the AOB⁶⁹. Moreover, the dendrites of the mitral cells in the AOB innervate multiple glomeruli⁷⁰⁻⁷². This multi-glomerular innervation pattern is in stark contrast with the main olfactory system, where olfactory sensory neurons (OSNs) expressing the same odorant receptor converge their axons into mostly a single glomerulus in each hemisphere of the main olfactory bulb (MOB)⁷³. The anatomical arrangement in the AOB has strong implications as to how species-specific cues are encoded and how the information is processed. In the MOB, when the convergent glomerular innervation is experimentally perturbed to become divergent, it does not affect detection or discrimination of

odorants but diminishes behavioral responses to innately recognized odors^{74,75}. Thus, stereotypic projection patterns provide a basis for genetically specified connections in the neural circuitry to enable innate behaviors. Consistent with this notion, it has been shown that mitral cell dendrites innervate glomeruli containing the same vomeronasal receptor (VR) type such that the divergent projection pattern of the VSNs is rendered convergent by the mitral cells⁷⁰. This homotypic convergence suggests that rather than using the spatial position of the glomeruli, the connection between VSNs and mitral cells in the AOB may rely on molecular cues to enable innate, stereotypical responses across different individuals. To a lesser extent, heterotypic convergence, i.e., axons expressing different receptors innervating the same glomeruli, is also observed⁷¹. In either case, expression of the molecular cues is likely genetically specified and tied to individual VRs, but little is known about how this specification is determined. Our analyses revealed the stereotypic association between transcription factors, axon guidance molecules, and the VRs to suggest a molecular code for circuit specification.

RESULTS

Cell types in the VNO

We dissected mouse VNOs from postnatal day 14 (P14) juveniles and P56 adults. Cells were dissociated in the presence of actinomycin D to prevent procedure-induced transcription. From four adult (P56) and four juvenile (P14) mice (equal representation of sexes), we obtained sequence reads from 34,519 cells. The samples and replicates were integrated for cell clustering. In two-dimensional UMAP space, 18 cell clusters can be clearly identified (**Fig. 1A**). These clusters were curated using known cell markers (**Fig. 1B**). There was no obvious difference in the cell clusters between juvenile and adult VNOs (**Fig. 1C**), even though there were slight differences in gene expression profile between the samples (see below). We also did not detect a significant difference between female and male samples (**Fig. 1D**).

Clustering confirmed all previously identified cell types but also revealed some surprises. The largest portion of cells belonged to the neuronal lineage, including the globose basal cells (GBC), immediate neuronal progenitors (INPs), the immature and mature VSNs, and a population of cells that do not co-cluster with either VSN type (see below). There were substantial numbers of sustentacular cells (SCs), horizontal basal cells (HBCs), and ms4-expressing microvillus cells (MVs). Cells engaged in adaptive immune responses, including microglia and T-cells, were also detected. A population of Fpr-1 expressing cells that were distinctive from the VSNs expressing the Fpr family of genes formed a separate class. These were likely resident cells mediating innate immune responses. We also identified the olfactory ensheathing cells (OECs) and a population of lamina propria (LP) cells, which share molecular characteristics with what we have found in the MOE⁷⁶.

To obtain the spatial location of the various cell types, we selected 100 target genes based on the scRNA-seq results. Using probes for these genes, we used the Molecular Cartography® platform to perform spatial molecular imaging (**Fig. 1E**). Based on molecular clouds and DAPI nuclei staining, we segmented the cells and quantified gene expression profiles to cluster the cells. We then map individual cell clusters onto their spatial locations in VNO slices. Unlike previous

studies that relied on a few markers to identify cell types, our approach relied on the spatial transcriptome to calculate the probability that a cell belongs to a specific class. This analysis revealed that the VSNs and supporting cells are located in the pseudostratified neuroepithelium (**Fig. 1F**). The LP cells are located along the lamina propria underlying the neuroepithelium as found in the MOE (**Figs. 1F and 1G**). Surprisingly, however, there are few HBCs located along the basal lamina, in direct contrast to the MOE (**Figs. 1F and 1G**). Most of the HBCs are found in the non-neuronal epithelium surrounding the blood vessel, with few near the marginal zone. The marginal zone is thought to be the neurogenic region⁷⁷⁻⁷⁹. We found both GBCs and INPs in this area. A previous study suggested neurogenic activity in the medial zone⁷⁸, but we did not find evidence in support of this conclusion, which was largely based on BrdU staining of mitotic cells without lineage-specific information. Based on our transcriptomic analysis, we conclude that neurogenic activity is restricted to the marginal zone.

Novel classes of sensory neurons in the VNO

To better understand the developmental trajectory of the VSNs, we segregated the cells in the neuronal lineage from the whole dataset (**Fig. 2A**). The neuronal lineage consists of the GBCs, INPs, immature neurons as determined by the expression of *Gap43* and *Stmn2* genes (**Figs. S1A and S1B**), and the mature VSNs. Cells expressing *Xist*, which is expressed in female cells, were intermingled with those from males (**Fig. S1C**). This observation is consistent with our previous studies using bulk sequencing indicating that the cell types are not sexually dimorphic²⁵. It is also consistent with physiological responses of the VSNs to various stimuli^{5,62,64,80,81}. For further analyses, therefore, we did not segregate the samples according to sex.

The VSNs are clearly separated into the V1R and V2R clusters as distinguished by the expression of *Gnai2* and *Gnao1*, respectively (**Fig. 2B**). Surprisingly, we observed that a portion of the V2R cells also expressed *Gnai2*. These cells were primarily from adult, but not juvenile male mice (**Fig. S1D**). Past studies based on *Gnai2* and *Gnao1* expression would have identified this group of cells as V1R VSNs, but the transcriptome-based classification places them as V2R VSNs. The signaling mechanism of this group of cells may be different from the canonical V2R VSNs.

We identify a distinct set of cells expressing the odorant receptors that comprise ~2.3% of the total neurons (**Fig. 2A**). Previous studies have found odorant receptor (OR) expressing cells that project to the AOB^{82,83}. It was not clear how prevalent OR expression was in the VNO, nor whether the cells were VSNs or OSNs. Our results show that these neurons lack the V1R or V2R markers (**Fig. 2C**). Instead, they express *Gnal* and *Cnga2*, which are the canonical markers of OSNs in the main olfactory epithelium, marking them canonical OSNs. Spatial mapping reveals that the OSNs are mainly in the neuroepithelium, with some cells concentrated in the marginal zone (**Fig. 2D**).

We also found a major group of cells that expressed the prototypical VSN markers but formed a cluster distinct from the mature V1R and V2R cells (**Figs. 1A and 2A**). Within the cluster, there was an apparent segregation among the cells into V1R and V2R groups based on marker gene expression (**Fig. 2B**). A small group of OR-expressing neurons also belonged to this cluster. These cells expressed fewer overall genes and with lower total counts when compared with the

mature V1R and V2R cells (**Figs. S1E and S1F**). The lower ribosomal gene expression suggests that these neurons are less active in protein translation (**Fig. S1G**). This group of cells are scattered within the epithelium (**Fig. 2E**).

There are 503 differentially expressed genes in this cluster when compared with other mature neurons ($p_{adj} < 0.001$; $FC > 1.5$; **Figs. 2F, S1H**). Gene ontology (GO) analysis reveals that differentially expressed genes enriched in this group are associated with odorant binding proteins (**Fig. 2G**). Correspondingly, mucin 2 (Muc2), odorant binding protein 2a (Obp2a), Obp2b, and lipocalin 3 (Lcn3) genes, usually enriched in non-neuronal supporting or secretory gland cells, are expressed at higher levels in these neurons than the canonical VSNs (**Figs. 2H, S2A-D**). They also have higher expression of Wnk1, which is involved in ion transport, the lncRNA Neat1, the purinergic receptor P2ry14, the zinc finger protein Zfp738, and the centrosome and spindle pole associated Csppl (**Fig. S2E-I**). On the other hand, these cells exhibit lower expression of Omp and JunD (**Figs. 2F, S2M-N**). The lower level of Omp suggests that these cells do not have the full characteristics of mature VSNs (mVSNs), but they are also distinct from the immature VSNs (iVSNs) as they do not express higher levels of immature markers such as Gap43 and Stmn2 (**Figs. S1A and S1B**). The lower expression of Ndua2, Ndua7, Cox6a1, and Cox7a2, which are involved in mitochondrial activities, suggest that these cells are less metabolically active than the canonical VSNs (**Figs. 2F, S2O-R**). Taken together, the gene expression profile suggests that this new class of neurons may not only sense environmental stimulus, but also may provide proteins to facilitate the clearing of the chemicals upon stimulation. We, therefore, name these cells as putative secretory VSNs (sVSNs).

Developmental trajectories of the neuronal lineage

The vomeronasal organ develops from the olfactory pit during the embryonic period and continues to develop into postnatal stages^{79,84,85}. Neurons regenerate in adult animals^{86,87}. Cell types in the vomeronasal lineage have been shown to be specified by BMP and Notch signaling^{88,89}, and coordinated by transcription factors (TFs) including Bcl11b/Ctip2, C/EBP γ , ATF5, Gli3, Meis2, and Tfap2e⁹⁰⁻⁹⁵. However, the transcriptomic landscape that specifies the lineages is not known.

To explore VSN development, we performed pseudotime inference analysis of the V1R and V2R lineages for P14 and P56 mice using *Slingshot*⁹⁶ (**Fig. 3A**). We set GBCs as the starting cluster and mVSNs as terminal clusters. A minimum spanning tree through the centroids of each cluster was calculated using the first fifty principal components and fit with a smooth representation to assign pseudotime values along the principal curve of each lineage. Cell density plots for both the V1R and V2R lineage reveal a higher portion of immature VSNs at P14 than at P56. For the mature VSNs, the P14 samples have peaks at an earlier pseudotime than the P56 mice (**Fig. 3B**). This result indicates the VSNs in juvenile mice are developmentally less mature than their counterparts in adults, but these differences do not distinguish them in obvious ways (**Fig. 1C**).

On the other hand, we have identified dynamic changes in transcriptomes associated with the V1R and V2R lineages. There were 2,037 significantly differentially expressed genes between V1R and V2R lineages (**Fig. 3C**). While V1R and V2R lineages shared some of the genes in the early stages of development, most show distinct expression patterns between the two. To

determine the transcriptional program associated with the lineages, we examined the expression sequence of individual transcription factors and signaling molecules from the gene list.

The transcription factor *Ascl1* is expressed by a subset of the GBCs that appears to be the earliest in developmental stage whereas *Sox2* is expressed by a broader set of GBCs (**Fig 3D-F**). The early INPs are defined by the expression of *Sp8*, *Neurog1* and *NeuroD1*. *Neurog1* is mostly restricted in the early INPs whereas *NeuroD1* and *Sp8* are also expressed by the late INPs (**Fig. 3G-I, S3A-B**). *NeuroD1* is expressed by iVSNs and iOSNs, but *Sp8* is only expressed by the iVSNs.

To obtain a refined view of the transition from INPs to the immature sensory neurons, we took the subset and re-clustered them (**Fig. 3J**). The late INPs are separated into two clusters that share marker genes with the V1R and V2R iVSNs, respectively, indicating that commitment to the two lineages begins at the late INP stage. Consistent with previous findings, the homeobox protein *Meis2* was specific to the V1R lineage (**Fig. 3K**). Concomitant with *Meis2*, there are a number of other genes expressed by the late INPs committed to the V1R fate, including secretin (*Sct*), *Foxj1* target gene *Fam183b*, the microRNA *Mir100hg*, acyl-CoA dehydrogenase *Acad10*, and *Keratin7* (*Krt7*; **Fig. 3L-O, S3C**). Notably, *Bcl11b* is exclusively absent from INPs committed to the V1R fate. This observation is consistent with the observation by Katreddi and colleagues that *Bcl11b* is lost in basal INPs in Notch knockout⁸⁹. Together with the observation that loss of *Bcl11b* results in increased number of V1R VSNs⁹⁰, these results indicate that transient downregulation of *Bcl11b* is required for the V1R lineage (**Fig. 3U**).

The transcription factor *Tfap2e*, which is required to maintain the V2R VSNs⁹¹, is expressed by the iVSNs but not by the late INPs committing to the V2R fate (**Fig. 3P**). *Sp8* is the only transcription factor specifically expressed in the late INPs committed to the V2R but not the V1R fate (**Fig. 3I**). *Emx2* is expressed by all neuronal lineage cells (**Fig. S3E**). *Krt8* is also found throughout the V2R lineage, but its expression is diminished in the V1R cells (**Fig. S3D**).

The OSN lineage is marked by the expression of *Olig2*, *Fezf1*, *Tshz2*, and *Nfib* (**Fig. 3Q-T**). The expression of *Olig2* and *Fezf1* is exclusive to the OSN fate. *Tshz2* is expressed at a late stage of the iVSNs, but not in the INPs. There are multiple genes that may not directly be engaged in cell fate determination but are clearly markers of the cell types (**Fig. S3F-R**). Although *Notch1* and *Dll4* are not identified as significantly differentially expressed, feature plots show clear distinction in their expression in the V2R and V1R lineage, respectively, as an earlier study showed⁸⁹.

Based on these patterns, we propose a model of molecular cascades that specify the neuronal lineages in the VNO (**Fig. 3V**). *Ascl1* and *Sox2* specify the GBCs. The down regulation of *Ascl1*, *Sox2*, and subsequent upregulation of *Neurog1*, *NeuroD1*, and *Sox8* commits the cells to the early INPs. The VSN and OSN lineages diverge after the early INP stage. The downregulation of *Sp8*, *Nfib*, and *Bcl11b* and the expression of *Meis2* promote the V1R fate. The downregulation of *Sp8* and the expression of *Fezf1*, *Tshz2*, and *Olig2* are required for commitment to the OSN lineage. Downregulation of *Sp8* is not required for the V2R lineage, which begin expressing *Tfap2e*. *NeuroD1* is expressed in all INPs. These patterns of expression

suggest that both the OSN and V1R lineages required the expression of specific transcription factors. The V2R lineage, on the other hand, appears to rely on factors inherited from the early INPs. This suggests the possibility that the V2R lineage is a default path for the VSNs.

Receptor expression in the VSNs

We quantified the expression of V1Rs, V2Rs, and ORs to gain insights into how chemosensory cues may be encoded by the VNO. For comparison, we also included OR expression from the MOE ⁷⁶. Within each class of receptors, the probability of expression of a gene follows a power law distribution except for the lower ranked genes (**Fig. S4A**). The sharp deviation from the power law curve for the lower ranked receptors is likely from technical dropout of genes expressed at low levels as they cannot be effectively captured by the scRNA-seq platforms. We plotted the relationship between total reads and the number of cells expressing a given receptor, and the average reads per cell for the receptors (**Fig. 4A-H**). We observed weak correlations between the total reads and the cell number expressing a given V1R or a V2R (**Fig. 4A and B**). This non-uniform distribution of VR genes is consistent with our observation from bulk sequencing results ²⁵.

Most V1Rs and V2Rs are expressed by less than 1500 cells (out of 34,519). Most VRs are expressed at less than 100 counts per cell (**Fig. 4E and F**). Several V1Rs, including *Vmn1r184*, *Vmn1r89*, *Vmn1r196*, *Vmn1r43*, and *Vmn1r37*, are highly expressed in individual cells (**Fig. 4A**). *Vmn1r196*, *Vmn1r43*, and *Vmn1r37* are also expressed at the highest level per cell. Some others, including *Vmn1r183*, *Vmn1r81*, and *Vmn1r13*, are expressed in large numbers of cells but have low expression in individual cells. Notably, the two highest expressed receptors recognize female pheromone cues. *Vmn1r89*, also known as *V1rj2*, is one of the receptors that detects female estrus signals. *Vmn1r184* is one of the receptors for the female identity pheromone ^{4,61,97}. The functions of *Vmn1r196*, *Vmn1r43*, and *Vmn1r37*, however, are unknown. We did not detect the expression of 45 V1Rs, which may be the result of technical dropout (**Fig. 4E**).

We do not observe obvious correlations between expression level and chromosomal locations. Both *Vmn1r89* and *Vmn1r184* are located on Chromosome 7, in a region enriched in V1R genes. Similarly, *Vmn1r37* and *Vmn1r43* are in a V1R-rich region on Chr. 6. However, *Vmn1r183*, *Vmn1r13*, and *Vmn1r81*, which are in the same clusters, are expressed by many cells but at some of the lowest levels.

All V2R genes are detected in the VNO (**Fig. 4B and F**). *Vmn2r53*, which has been shown to mediate intermale aggression through a dedicated circuit, has the highest level of expression and is expressed by the second most cells ⁹⁸. Among the highly expressed V2Rs, *Vmn2r1* and *Vmn2r7* are co-receptors for other V2Rs. *Vmn2r59* has been shown to detect predator signals ⁴. *Vmn2r115*, a receptor for ESP22 that is secreted by juveniles ⁹⁹, is expressed by the highest number of cells. However, *Vmn2r116* (*V2rp5*), which recognizes ESP1 and induces lordosis behavior in females, is a close homolog of *Vmn2r115* but not among the highly expressed genes ^{99,100}. Notably, *Vmn2r114*, close homolog of *Vmn2r115* and *Vmn2r116*, is also expressed by

large numbers of cells. *Vmn2r88*, a hemoglobin receptor¹⁰¹, was not identified as a highly expressed gene. The functions of other highly expressed receptors are not known.

In contrast to the VR genes, total counts for ORs in the VNO exhibit a tight relationship with the number of cells (**Fig. 4C and G**). Except for *Olfr124*, most of the OR genes align well with the linear regression curve. This relationship is different from the VR genes and is also distinct from the single cell data from the MOE, which exhibits a similar distribution as the VRs⁷⁶ (**Fig. 4D and H**). Out of the 686 OR genes detected in the VNO, only 80 are expressed by more than 10 copies per cells, indicating that a majority of the ORs do not contribute to meaningful signaling of chemosensory cues.

To comprehensively survey receptor expression, we also included VR and OR pseudogenes in our analysis (**Fig S4B-G**). We did not detect a significant expression of pseudogenized V1Rs (**Fig. S4B and E**), but *Vmn2r-ps87* has the highest count/cell in the V2R population (**Fig. S4F**). *Olfr709-ps1*, and *Olfr1372-ps1* are the two highest expressed genes in terms of total count and total number of cells in the VNO (**Fig. S4D**).

We next examined transcription factors associated with individual receptor types. In the MOE, ORs are monoallelically expressed¹⁰². The unique expression of an OR gene is mediated by chromosomal repression and de-repression, coordinated by transcription factors and genes involved in epigenetic modification¹⁰³⁻¹⁰⁸. Monoallelic V1R gene expression is also observed⁶⁷. Epigenetic modification takes place at V1R gene clusters^{109,110}, but the repression of receptor genes appears to permit transcriptional stability rather than receptor choice¹¹¹. How VR genes are selected is not known. To explore our dataset for clues of transcriptional activities associated with VR expression, we plotted the cross-correlation between VRs and their TF profiles. We observed correlations among receptors (**Figs. 4I and 4J**). We did not find an obvious association between TF profiles and VR sequence similarity of pairs (**Fig. S5A**).

There is a high correlation among a large portion of V1Rs (**Fig. 4I**). By analyzing the correlation between individual TFs with the VRs, we found that V1R expression is overwhelmingly associated with *Meis2* (**Fig. S5B**). The few receptor types that do not show high correlation with *Meis2* are associated with *Egr1* or *c-Fos*. It is not clear whether these prototypical immediate early genes are involved in specifying receptor choice. This result indicates that once the V1R lineage is specified by *Meis2*, receptor choice is not determined by specific combinations of transcription factors. This scenario is similar to receptor choice by the OSNs in the MOE.

Different from the V1Rs, we observed islands of high similarity of TF expression among the V2Rs, indicating that receptors in these islands share the same set of TFs (**Fig. 4J**). We identify correlation between individual TFs with V2Rs (**Fig. S5C**). Whereas *Tfap2e* is involved in specifying the V2R fate, it is only associated with the expression of a subset of V2Rs. *Pou2f1* and *Atf5* are more strongly associated with other subsets of receptors. Unlike the V1Rs, there is a disparate set of TFs associated with the V2Rs, suggesting that the V2R choice may be determined by combinations of TFs. Based on the prevalence of individual transcription factors in association with the VRs, we propose a model of transcriptional cascade that may be involved in receptor choice (**Fig. 4K**). For the V2Rs, *Tfap2e+* cells can be further specified by *Ikzf4*,

Tcf4, and Trps1. In *Ikzf4* negative cells, *Batf3*, *Atf5*, *Pbx2*, and *Pou2f1* can specify receptor types, respectively. In the *Tfap2e*- cells, receptors can be specified by *Pou2f1*, *Rlf*, and *Batf3*.

Co-expression of chemosensory receptors

We next investigated co-expression of receptors in individual cells. Since we applied SoupX to limit ambient RNA contamination and Scrublet to remove doublet cells, we set a stringent criterion in counting receptors expressed by single cells¹¹². *V1R* and *V2R* genes on average constitute ~2% of total reads per cell, and the ORs constitute less than 1% of the total reads per cell (**Fig. S6A-C**). On average, the *V2Rs* have significantly more than one receptor per cell (**Fig. S6B**). Using Shannon Index to measure uniqueness of receptor expression in individual cells, we found that the mature VSN and OSNs have relatively high specificity, but many cells show significantly higher index values, indicating that they expressed multiple receptors (**Fig. 5A**). In support of this observation, there are significant representations of the second and third highest expressed receptors in all three types of neurons (**Fig. S6D-G**).

To further reduce contributions of spurious, random low-level expression, we only consider those receptors with at least 10 raw counts per cell and are found in at least 5 cells to evaluate co-expression among receptors. We split cells into groups according to cell-type, age, and neuronal lineage and plot the percentage of cells expressing zero, one, two, and three or more species (**Fig. 5B-C**). This analysis shows that receptor expression specificity increases as the lineages progressed from progenitors into mature neurons. Immature neurons have more cells co-expressing receptors than mature cells. More co-expressions are observed in the younger animals than the older ones. This line of evidence indicates that the co-expression we have found is not from experimental artifacts but reflects real biological events. Contamination would not be selectively enriched in immature cells and not present in the INP cells.

We performed Fisher's exact test using contingency tables for every pairing of expressed receptor genes. For the pairs that pass the test, we generated Circos plots to show the genomic loci for all significant *V1R*, *V2R*, and interclass pairs (**Fig. 5D-F**). This analysis showed that 47.7% of co-expressed receptors are co-localized on the same chromosome.

We found a few sets of co-expressed VRs that would have strong implications for how pheromone signals are detected. *Vmn1r85* (*V1rj3*) and *Vmn1r86* are two receptors located next to each other on Chr. 7, sharing a high level of homology, and belonging to the *V1rj* clade. They have a high level of co-expression but are not co-expressed with *Vmn1r89* (*V1rj2*), located ~100Kb away, even though both *Vmn1r85* and *Vmn1r89* receptors are activated by sulfated estrogen and carry information about the estrus status of mature female mice^{61,65,97} (**Fig. 5D**). Another set of receptors that recognize female-specific pheromone cues are the *V1re* clade receptors. We found that *Vmn1r185* (*V1re12*), which recognizes female identity pheromones, was co-expressed with its close homolog *Vmn1r184* gene, which is about 350kb away on Chr 7. They are not co-expressed with *Vmn1r69* (*V1re9*), which also recognizes female pheromones, but is located 16Mb apart on Chr. 7^{61,97,113,114} (**Fig. 5D**).

On Chr. 7 there are two other major clusters of *V1R* genes that show co-expression. One cluster includes *Vmn1r55*-*Vmn1r64*, 10 genes belonging to the *V1rd* clade and located within a 600Kb

region. Another one includes Vmn1r167, Vmn1r168, and Vmn1r169, which appear to be specifically paired with Vmn1r175, Vmn1r177, and Vmn1r176, respectively. These receptor pairs are arranged in a 300Kb region with a head-head orientation. Outside of Chr. 7, several small clusters on Chr. 6 and one large cluster on Chr. 3 exhibit significant co-expression of V1Rs.

The V2R neurons coordinately express one common V2R and one specific receptor^{115,116}. Our co-expression analysis confirms the broad association of Vmn2r1-7, which are located on Chr. 3, with other receptors across various chromosomal locations (**Fig. 5E**). In addition, we also identified co-expression patterns among the specifically expressed V2Rs. Among these receptor genes, intra-chromosomal co-expressions are observed for receptors residing on Chr. 7, Chr. 17, and Chr. 5. There is also inter-chromosomal co-expression between one locus on Chr. 17 with a cluster on Chr. 7 (**Fig. 5E**).

Lastly, we have observed co-expression of receptors across different classes of receptors (**Fig. 5F**). Notably, Vmn2r1-3 are co-expressed with several Vmn1r receptors on Chr. 3. Vmn2r7 is co-expressed with a group of Vmn1r genes on Chr. 13. The odorant receptor Olfr344 is co-expressed with several V1R and V2R genes.

In past studies, VR expression was examined using *in situ* hybridization, immunostaining, or genetic labeling. The traditional histological methods were not sensitive enough to quantitatively measure signal strength. Moreover, pairwise double *in situ* is too laborious to capture co-expression of two or more receptors. To verify co-expression of VR genes by individual VSNs, we selected 30 VR genes based on scRNA-seq data and used Molecular Cartography© to examine their expression patterns *in situ*.

Given the high incidents of co-expression between receptors in the Vmn1r55-64 cluster on Chr. 7 (**Fig. 5D**), we included 5 probes for this set of genes and confirmed colocalization of pairs in the VSNs (**Fig. 5G**). Interestingly, we did not find cells expressing more than two receptor genes for this set. We also confirmed the co-expression between Vmn1r85 and Vmn1r86 as indicated by single cell data (**Fig. 5D and G**).

We confirmed the co-expression among genes from the V2R genes (**Fig. 5H**)^{117,118}. Consistent with previous reports^{117,119}, Vmn2r2, Vmn2r3, Vmn2r6, and Vmn2r7 are comingled in several cells, but Vmn2r1 is not co-expressed with these 4 broadly expressed V2Rs (**Fig. 5H**). Outside of the Vmn2r1-7 group, we find that Vmn2r81 is co-expressed with Vmn2r20 or Vmn2r24 but without any of the Vmn2r1-7 transcripts (**Fig. 5H**). We detected more incidents of co-expression between Fpr3 and Fpr-rs4, and between Fpr3, Fpr-rs3, Fpr-rs4 with V1Rs than with V2Rs (**Fig. 5I and J**). We also found co-expressions that are not predicted by the single cell analysis (**Fig. 5K**). The discrepancy likely can be attributed to the relatively low-level expression of one of the receptor genes. This type of co-expression may not pass the strict criteria we set for the single cell analysis.

A surface molecule code for individual receptor types

VSNs expressing a given receptor type project to the AOB to innervate glomeruli distributed in quasi-stereotypical positions^{67,68,100}. The high number of glomeruli innervated by a given VSN

type raises the question about mechanisms that specify the projection patterns and the connection between the VSNs and the mitral cells. For a genetically specified circuit that transmits pheromone and other information to trigger innate behavioral and endocrine responses, there must be molecules that instruct specific connections among neurons. Several studies have revealed the requirement of Kirrel2, Kirrel3, Neuropilin2 (Nrp2), EphA, and Robo/Slit in vomeronasal axon targeting to the AOB¹²⁰⁻¹²⁶. However, how individual guidance molecules or their combinations specify connectivity of individual VSN types is completely unknown. Here we leverage the unbiased dataset to identify surface molecules that may serve as code for circuit specification.

We identified 307 putative axon guidance (AG) molecules, including known cell surface molecules involved in transcellular interactions and some involved in modulating axon growth. Using this panel, we calculated pairwise similarity between two VR genes, and the similarity in their guidance molecule expression. Analysis of the relationship indicates there is a general increase in guidance molecule similarity associated with VR similarity (**Fig. 6A**). Consistent with this observation, when we plotted the similarity of surface molecule expression among cells expressing different receptors, we found islands of similarity among the receptor pairs (**Fig. 6B**). We then conducted correlation analysis between the guidance molecules with the V1Rs (**Fig. 6C**) and V2Rs (**Fig. 6D**). Consistent with previous reporting, we found that Kirrel2 was associated with nearly half of the V1Rs and Kirrel3 was mainly associated with V2Rs that project to the caudal AOB^{120,121}. Robo2 is associated with nearly all V2Rs. We found that Teneurin (Tenm2 and Tenm4) and protocadherin (Pcdh9, Pcdh10, and Pcdh17) genes were associated with specific receptors^{127,128}. EphA5, Pcdh10, Tenm2, Nrp2 are also strongly associated with V1Rs with partial overlap with each other. Pcdh9, Tenm4, Cntn4, EphrinA3, Pcdh17, as well as Kirrel2 and Kirrel3 all show association with specific V2Rs. Numerous guidance molecules that are not broadly expressed are also associated with individual VRs.

Based on these associations, and supported by existing literature, we propose a model of the specification of separate groups of VRs. In this model, Robo2 separates the rostral vs. caudal AOB. Robo2⁺ V2R neurons project to the caudal AOB whereas Robo2⁻ V1R cells project to the rostral AOB^{125,126,129}. Among V1Rs, Kirrel2 expression distinguishes between two groups of cells^{120,130}. The Kirrel2⁺ population can be further separated into EphA5⁺ and EphA5⁻ population^{122,131}. The EphA5⁺ population can be separated further by Pcdh10 expression. In the Kirrel2⁻ cells, EphA5, Pcdh10, Tenm4, and Tenm2 mark separate groups. Ncam1, EphA5, and Cntn4 may contribute to specifying small sets of cells. For the V2Rs, Pcdh9, Cntn4, Tenm4, Tenm2, and Pcdh17 have a decreasing range of expression, which may be used to specify increasingly refined connection. Notably, even though Kirrel2 and Pcdh10 are mostly detected in the V1Rs, they are also expressed by small sets of the V2Rs.

Transcriptional regulation of receptor and axon guidance cues

The specification of a vomeronasal circuit needs to be tied to receptor expression. We next address whether a transcriptional code is associated with both VR and AG molecule expression. We calculated pairwise similarity among the receptors according to their expression of TF and AG genes (**Fig. 7A**). The result shows that V1R and V2R are distinctively separated. Besides a

small group of broadly expressed V2Rs, all VR types are unique in their gene expression. Fpr types are more similar in their expression profile with the V1Rs, but the OR types are intermingled with both VR types.

To further determine the TF/AG associations that are specific to individual receptor types, we applied stringent criteria to calculate the Jaccard Index for each pair of TF/AG (**Fig. 7B**), which reflect the statistical probability of co-expression within the same cell. The analysis reveals unique combinations for every receptor type (**Fig. 7C-G**). Even though the only TF associated with V1R expression is *Meis2*, other TFs are involved in specifying AG gene expression. For example, cells expressing V1Rs with high sequence homology and in chromosomal proximity share a similar set of TF and AG genes, but the expression patterns are distinct from each other (**Fig. 7D**). For broadly expressed V2Rs, *Vmn2r3* and *Vmn2r7*, which are co-expressed by the same set of cells, share nearly identical TF/AGs (**Fig. 7E**). In contrast, *Vmn2r1*, which does not co-express with either *Vmn2r3* or *Vmn2r7*, lacks the expression of *Pou2f1* and *Tenn2* despite sharing all other genes. Other V2R types also show distinct TF/AG associations (**Fig. 7F-G**).

DISCUSSION

Single cell RNA-seq analysis provides an unprecedented opportunity to identify cell types and determine genes associated with individual cells, but it is not without pitfalls. At the current state of the art, the depth of sequencing only allows sampling of transcripts that are expressed at relatively high levels. Sequencing dropouts and potential contamination also can complicate the analysis. Using the current state of the art tools, and applying conservative criteria, we provide an in-depth look at the molecular and cellular organization of the mouse VNO. The analyses reveal new cell types, specific co-expression of receptors, and transcriptional regulation of lineage specification. Moreover, our analyses uncover specific associations between transcription factors, surface guidance molecules, and individual receptor types that may determine the wiring specificity in the vomeronasal circuitry.

The molecular distinction between the sVSNs and the classic VSNs indicates that they may serve a specific function. They are different from the solitary chemosensory cells that are trigeminal in nature¹³¹. We speculate that these cells may secrete olfactory binding proteins and mucins in response to VNO activation. The VNO is a semi-blind tubular structure. Chemical cues are actively transported into the VNO and can only be cleared by active transport systems, which are thought to be carried out by the lipocalin family of proteins, or by degradation¹³¹⁻¹³⁴. These proteins are important to protect the integrity of neuroepithelia. They are generally produced by the sustentacular cells or the Bowman's gland¹³⁴. It is plausible that the sVSNs can produce specific lipocalin proteins based on the ligand detected by the VRs they express. These cells may also convey chemosensory information to the AOB, but we do not know if they project to the central brain. We also have identified a class of canonical OSNs in the VNO. Previous reports show that these neurons project to AOB. It is plausible that the OSNs can detect a set of volatile odors that carry species-specific information and directly convey it to brain areas that regulate innate responses. Our list of these ORs could direct effort to identify these odors to reveal their ethological relevance.

The co-expression of multiple VRs in individual VSNs is intriguing, as a previous analysis of the MOE detected minimal co-expression of ORs²¹. Importantly, there is a higher propensity for VR co-expression between certain receptor pairs. Notably, there is co-expression of receptors sharing common ligands, or that are similar in their sequences. These observations indicate that co-expressed receptors may serve redundant function detecting the cues. For example, the V1rj receptors are cognate receptors for sulfates estrogen and estrus signals. Their co-expression indicates that the neurons detect the same class of molecules redundantly. Moreover, co-expression of similarly tuned receptors makes it plausible for heterotypic convergence, when these neurons converge into glomeruli that express one or the other receptors.

How specificity of neuronal connections in the olfactory system is determined remains unknown. In the main olfactory system, glomerular positions are coarsely specified along the anterior-posterior and dorsal-ventral axes by gradients of axon guidance molecules whereas the sorting of axons according to the odorant receptors is mediated by homophilic attraction and heterotypic repulsion using a different set of guidance molecules¹³⁵. Spontaneous neural activities determine the expression of both sets of molecules. It is not known whether VSNs utilize the same mechanism. Given the multi-glomerular innervation patterns by the VSNs, it is exceedingly difficult to determine the contribution of individual guidance molecules to specifying VSN innervation.

We have identified guidance molecules associated with individual VRs that potentially constitute a code set that specifies VSN axon projections and their connection with postsynaptic cells. Each receptor type has a unique combination of guidance molecules expressed, which provides a basis for axon segregation and convergence. There are a few molecules that are shared broadly by various VSN types. These can be used to instruct general spatial locations of the VSN axons. For example, *Robo2* separates the anterior vs. posterior AOB. Knockout of *Robo2* causes mistargeting of V2R neurons to the rostral AOB. Our models also indicate that *Kirrel2* and *Kirrel3* are expressed by nearly half of the VR types in partially overlapping patterns. Deletion of *Kirrel2* or *Kirrel3* leads to disorganization of glomeruli in the posterior AOB. Protocadherins and tenurins add new dimensions to this code. We also identified several guidance molecules that are more specifically associated with individual VRs. They could provide additional cues to separate axons that share broadly expressed guidance molecules.

We have identified lineage relationships among cells in the VNO and a dynamic transcriptional cascade that likely specifies cell types during development. We confirm *Meis2* and *Tfap2e* as transcription factors that maintain the V1R and V2R fate. We also identify several transcription factors that likely play key roles in specifying these fates. For example, the down regulation of *Neurog1*, but not *NeuroD1*, is associated with a transition from early INPs to late INPs. The downregulation of *Sp8*, *Nfib*, and *Bcl11b* is likely important for committing to the V1R lineage for late INPs. On the other hand, downregulation of *Sp8* and upregulation of *Fezf1*, *Olig2*, and *Tshz2* likely set up commitment to the OSN fate. We also find that in all three lineages, the expression of *Tshz2* is associated with transition to the immature neuronal fate from the late INPs.

We observed a striking difference between V1R and V2R VSNs in the transcription factors associated with receptor choice. There is no overt association between V1R with specific transcription factors. This observation is reminiscent of OSNs in the olfactory epithelium, where OR expression is stochastic and mediated by de-repression of epigenetically silenced OR loci¹⁰³⁻¹⁰⁸. The absence of specific transcription factors with individual V1R choice suggests that a similar mechanism may operate in the V1R VSNs. Monoallelic expression of V1Rb2 supports this notion⁶⁷. On the other hand, we observed that for individual V1R types, there are specific associations between transcription factors with guidance molecules. This observation implies that the expression of guidance molecules is determined by combinations of transcription factors even though these transcription factors may not determine V1R expression. This is also reminiscent of the OSNs, where the expression of guidance molecules is determined by spontaneous neural activities¹³⁶⁻¹³⁸. That is, once the receptor choice is made, the specific receptor being expressed determines the guidance molecules to specify their projection patterns.

In direct contrast, V2R VSNs likely use combinations of transcription factors to specify receptor expression as well as guidance molecules. Some of the transcription factors that we observe to be associated with V2R expression are also associated with guidance molecule expression. For example, Pou2f1, Atf5, and Zfp268 are involved in both processes.

SUPPLEMENTAL INFORMATION

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AUTHOR CONTRIBUTIONS

L.M. and C.R.Y. conceived the study. A.F. and L.M. performed single cell RNA-seq experiments. T.C. and S.M. prepared samples for Molecular Cartography analysis. A.P. and A.S. facilitated single cell library preparation and RNA sequencing. M.H. performed bioinformatic analysis. M.H. and T.C. created figures and tables. C.R.Y. acquired funding, supervised the project, and wrote the final manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

STAR METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Software and Algorithms		
Fiji ImageJ	NA	https://imagej.net/software/fiji/
QuPath	NA	https://qupath.github.io
MGI	NA	https://www.informatics.jax.org/vocab/gene_ontology/
Seurat	R Package	https://satijalab.org/seurat/
kallisto bustools	Pachter Lab	https://www.kallistobus.tools
DropletUtils	R Package	https://bioconductor.org/packages/release/bioc/html/DropletUtils.html
Illustrator	Adobe	https://www.adobe.com/illustrator
SoupX	R Package	https://cran.r-project.org/web/packages/SoupX/index.html
BUSpaRse	R Package	https://bioconductor.org/packages/release/bioc/html/BUSpaRse.html
kit	R Package	https://cran.r-project.org/web/packages/kit/index.html
tidyverse	R Package	https://cran.r-project.org/web/packages/tidyverse/index.html
stringr	R Package	https://cran.r-project.org/web/packages/stringr/index.html

clustree	R Package	https://cran.r-project.org/web/packages/clustree/index.html
ggplot2	R Package	https://cran.r-project.org/web/packages/ggplot2/index.html
data.table	R Package	https://cran.r-project.org/web/packages/data.table/index.html
org.Mm.eg.db	R Package	https://bioconductor.org/packages/release/data/annotation/html/org.Mm.eg.db.html
dplyr	R Package	https://cran.r-project.org/web/packages/dplyr/index.html
DBI	R Package	https://cran.r-project.org/web/packages/DBI/index.html
glmGamPoi	R Package	https://bioconductor.org/packages/release/bioc/html/glmGamPoi.html
ggpubr	R Package	https://cran.r-project.org/web/packages/ggpubr/index.html
svglite	R Package	https://cran.r-project.org/web/packages/svglite/index.html
vegan	R Package	https://cran.r-project.org/web/packages/vegan/index.html
scales	R Package	https://cran.r-project.org/web/packages/scales/index.html
Scrublet	Python Package	https://github.com/swolock/scrublet
reticulate	R Package	https://cran.r-project.org/web/packages/reticulate/index.html
plotly	R Package	https://plotly.com/r/
Matrix	R Package	https://cran.r-project.org/web/packages/Matrix/index.html
tibble	R Package	https://cran.r-project.org/web/packages/tibble/index.html
stats	R Package	https://stat.ethz.ch/R-manual/R-devel/library/stats/html/00Index.html
patchwork	R Package	https://cran.r-project.org/web/packages/patchwork/index.html
openxlsx	R Package	https://cran.r-project.org/web/packages/openxlsx/index.html
gridExtra	R Package	https://cran.r-project.org/web/packages/gridExtra/index.html
ggrepel	R Package	https://cran.r-project.org/web/packages/ggrepel/index.html
GeneOverlap	R Package	https://bioconductor.org/packages/release/bioc/html/GeneOverlap.html
biomaRt	R Package	https://bioconductor.org/packages/release/bioc/html/biomaRt.html
circlize	R Package	https://cran.r-project.org/web/packages/circlize/index.html
slingshot	R Package	https://www.bioconductor.org/packages/release/bioc/html/slingshot.html
tradeSeq	R Package	https://www.bioconductor.org/packages/release/bioc/html/tradeSeq.html
SingleCellExperiment	R Package	https://bioconductor.org/packages/release/bioc/html/SingleCellExperiment.html
viridis	R Package	https://cran.r-project.org/web/packages/viridis/index.html
pheatmap	R Package	https://cran.r-project.org/web/packages/pheatmap/index.html
msigdbr	R Package	https://cran.r-project.org/web/packages/msigdbr/vignettes/msigdbr-intro.html
fgsea	R Package	https://bioconductor.org/packages/release/bioc/html/fgsea.html
IDPmisc	R Package	https://cran.r-project.org/web/packages/IDPmisc/index.html
msa	R Package	https://bioconductor.org/packages/release/bioc/html/msa.html
ggpointdensity	R Package	https://cran.r-project.org/web/packages/ggpointdensity/index.html
eulerr	R Package	https://cran.r-project.org/web/packages/eulerr/index.html
seqinr	R Package	https://cran.r-project.org/web/packages/seqinr/index.html
geomtextpath	R Package	https://cran.r-project.org/web/packages/geomtextpath/index.html
Rmisc	R Package	https://cran.r-project.org/web/packages/Rmisc/index.html
ggplotify	R Package	https://cran.r-project.org/web/packages/ggplotify/index.html

Spatial
Transcriptomics

Molecular Resolve <https://resolvebiosciences.com/>
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RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by C. Ron Yu (ryu@stowers.org).

Data and code availability

All RNA-seq data are available from the NCBI GEO server (GSE252365). All original data generated in this study will be available for download at Stowers original data repository upon publication. No custom generated computer code was used for analysis.

An HTML file containing relevant figures and statistics from the study, as well as tables showing co-expression and differential expression results, can be accessed in the following public folder: https://ftp.stowers.org/#/public/VNO_Atlas/.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Wildtype CD1 postnatal day 14 (P14) pups and adults (P56) were used for the experiment. Both sexes were randomly assigned to the experiment. All animals were maintained in Stowers LASF with a 14:10 light cycle and provided with food and water *ad libitum*. Experimental protocols were approved by the Institutional Animal Care and Use Committee at Stowers Institute and in compliance with the NIH Guide for Care and Use of Animals.

METHODS

scRNA library preparation & sequencing

Mice VNOs were dissected in cold oxygenated ACSF following Ma et al, 2011¹³⁹. Dissected epitheliums were dissociated in papain solution (20mg/mL papain and 3mg/mL L-cysteine in HBSS) with RNase-free DNase I (10unit) at 37°C for 15-20 minutes. 0.01% BSA in PBS was added to the digestion solution before filtering with 70µm and 30µm filters (pluriSelect). Dissociated cells were washed twice in 0.01% BSA with final volume 1mL, followed by Draq5 (25µM) and DAPI (0.5µg/mL) staining 5min on ice. Draq5+/DAPI- cells (live/nucleated cells) were sorted on BD Influx cytometer (BD Bioscience) with 100µm nozzle. Dissociated sorted cells were assessed for concentration and viability via Luna-FL cell counter (Logos Biosystems). Cells deemed to be at least 90% viable were loaded on a Chromium Single Cell Controller (10x Genomics), based on live cell concentration. Libraries were prepared using the Chromium Next GEM Single Cell 3' Reagent Kits v3.1 (10X Genomics) according to manufacturer's directions. Resulting cDNA and short fragment libraries were checked for quality and quantity using a 2100 Bioanalyzer (Agilent Technologies) and Qubit Fluorometer (Thermo Fisher Scientific). With cells captured estimated at ~5,500-8,000 cells per sample, libraries were pooled and sequenced to

a depth necessary to achieve at least 40,000 mean reads per cell on an Illumina NovaSeq 6000 instrument utilizing RTA and instrument software versions current at the time of processing with the following paired read lengths: 28*8*91bp.

scRNA-Seq pre-processing and QC-filtering

Gene-by-cell barcode count matrices were generated from raw FASTQ files using the kallisto|bustools (v0.48.0|v0.41.0)^{140,141} workflow with the *Mus musculus* genome assembly GRCm39 (mm39) and GTF gene annotation files retrieved from ENSEMBL release 104¹⁴². All downstream QC-filtering and analysis was performed in an R environment (v4.0.3)¹⁴³. Empty droplets were estimated and filtered from the data using the *barcodeRanks* function of the DropletUtils package (v1.10.3)¹⁴⁴ with the lower bound of UMI-counts set to 100. The count data was then imported into R (v4.0.3) using the Seurat package (v4.3.0)¹⁴⁵ and ambient RNA contamination was automatically estimated with the *autoEstCont* function and removed with the *adjustCounts* function from the SoupX package (v1.5.2)¹¹². Cell barcodes representing multiplets were identified and removed with Scrublet (v0.2.3)¹⁴⁶ interfacing with Python using reticulate (v1.30)¹⁴⁷. Cell barcodes expressing <750 genes, >2.5 standard deviations above the mean number of genes or counts, or >5% of reads originating from mitochondrial genes were removed from the downstream analysis.

scRNA-Seq integration and clustering

To cluster cells from multiple samples, raw gene counts for each sample were normalized with the *SCTransform* function (v0.3.2)¹⁴⁸ using the *glmGamPoi* method from the glmGamPoi package (v1.6.0)¹⁴⁹, samples were then integrated using the Seurat integration pipeline. After principal component analysis (PCA) the dimensionality of the whole integrated VNO dataset was estimated to be 25 based on visual identification of an ‘elbow’ using the output from the *ElbowPlot* function. The shared nearest neighbor graph was constructed with the *FindNeighbors* function. Then, using the *FindClusters* function, an optimal resolution of 0.7 was chosen for the discovery of broad cell types in the VNO by running the *clustree* function from the clustree package (v0.30)¹⁵⁰ to evaluate cluster-level mean expression for a curated list of VNO cell-type marker-genes in a range of resolutions between 0 and 1, with intervals of 0.1; optimal resolution was considered the lowest resolution at which the expression of *Neurod1* and *Ascl1* diverge into two clusters, as these genes are unique marker-genes for immediate neural progenitors (INPs) and globose basal cells (GBCs), respectively. At 0.7 resolution 31 clusters were observed.

Differential gene expression analysis and cell labeling

To label VNO cell-types the *FindAllMarkers* function was run on clusters using log normalized counts to determine the top significantly differentially expressed genes (DEGs) present in $\geq 50\%$ of each cluster’s cells with an absolute value \log_2 fold-change ≥ 0.5 ; additionally, we used the *FeaturePlot* function to plot expression of the marker-genes in 2D UMAP space; then, clusters were assigned to broad cell-types with a dual approach of considering each cluster’s DEG overlap with the curated list of marker-genes, and visually correlating marker-gene expression in UMAP space with cluster identity.

Neuronal Lineage

Cells labelled as GBC, INP, iVSN, iOSN, mVSN, mOSN, or sVSN were subset from the integrated whole VNO dataset, split by animal sample, then re-integrated and re-clustered, as

described above. The *FindNeighbors* function was run using 12 PCs and the *FindClusters* function was run with a resolution of 6.0 which resulted in 84 clusters. Clusters were assigned to cell-types as described previously, while a further distinction between early and late INPs was inferred from the expression of *Ascl1*, *Neurod1*, *Neurog1*, and *Gap43* in 2D UMAP space. Differential gene expression analysis was performed between the novel sVSN cluster and the mature V1R, V2R, and OSN clusters, independently, running the *FindMarkers* function with the *logfc.threshold* and *min.pct* parameters set to 0 to accommodate downstream gene set enrichment analysis (GSEA).

Gene set enrichment analysis

Gene ontology (GO) terms from the biological process, molecular function, and cellular compartment categories along with their associated gene sets were retrieved with the *msigdb* function from the *msigdb* package (v7.5.1)¹⁵¹. Ranked Wald test differential expression results between sVSNs and V1R/V2R mVSNs, respectively, were input into the *stats* parameter of the *fgsea* function from the *fgsea* package (v1.20.0)^{152,153} along with the GO term gene sets. The top significant GO terms were plotted using the *ggplot* function from the *ggplot2* package.

Immediate neural progenitors and immature vomeronasal sensory neurons

Cells previously identified as early and late INP, iVSN, iOSN, or mOSN, were subset from the neuronal dataset, split and reintegrated, as above, using 15 PCs and a resolution of 2.5, resulting in 30 clusters. Differential gene expression analysis was performed with the *FindMarkers* function between two clusters showing either V1R or V2R like properties but previously identified uniformly as early INPs in the whole neuronal lineage dataset. The *FeaturePlot* function, from the Seurat package, was used to show normalized expressions for genes of interest.

Trajectory inference and differential gene expression analysis

To explore transcriptional differences over pseudotime between V1R and V2R VSNs, we subset GBCs, early and late INPs, V1R and V2R iVSNs, and mVSNs from the neuronal dataset, split the data by sample and reintegrated, then performed PCA. Trajectory inference analysis was performed with the *slingshot* function from the *slingshot* package (v1.8.0)⁹⁶ using the first 12 PCs and cell type labels from the neuronal dataset, with an input starting cluster of GBCs and two input end clusters for V1R and V2R mVSNs. To determine the *nknots* parameter for the *fitGAM* function from the *tradeSeq* package (v1.4.0)¹⁵⁴, we ran the *evaluateK* function with the raw count matrix, and the pseudotime and cell weight values output from *slingshot*. We then ran the *fitGAM* with *nknots*=5. We then ran the *patternTest* function to test for differences in gene expression patterns over pseudotime between the V1R and V2R lineages.

Gene co-expression analysis

VR co-expression. To investigate cell-level diversity of vomeronasal receptor species across all cells in the neuronal dataset we calculated the Shannon diversity index for the raw gene counts for all *Vmn1r*, *Vmn2r*, *Olfir*, and *Fpr* genes using the *diversity* function from the *vegan* R package (v2.6-4)¹⁵⁵ with default parameters. To determine what proportion of cells in the neuronal lineage had one, two, or three or more receptor species, we set a threshold of ≥ 10 raw counts for a receptor to be considered 'present'. To test whether co-expressing vomeronasal

receptor species were significant, using the same raw-count threshold of ≥ 10 , we gathered a list of all cell barcodes where a receptor was observed, for all receptors. Using the lists of cell barcodes associated with the receptors, we ran the *newGOM* function from the GeneOverlap R package (v1.26.0) ¹⁵⁶, which calculates p-values using Fisher's exact test on a contingency table. P-values were then corrected for multiple testing using the Benjamini-Hochberg procedure. Using the circlize R package (v0.4.15) ¹⁵⁷, we plotted all co-expressed receptor pairs on circos plots showing each receptors genetic location and the number of cells expressing the pair, for all significant pairings ($p_{\text{adj}} \leq 0.05$).

VR co-expression with axon guidance (AG) and transcription factor (TF) genes. To ascertain vomeronasal receptor co-expression with AG genes expressed at the plasma membrane, and with DNA binding TF genes, respectively, we used the Mouse Genome Informatics database to find all genes associated with the biological process gene ontology (GO) term 'axon guidance', or with the molecular function GO term 'DNA-binding transcription factor activity'. For the axon guidance gene set, we subset genes that were expressed at the plasma membrane. We set the VR raw count threshold to ≥ 10 counts and the AG and TF gene raw count threshold to ≥ 3 . Then, using the cell barcodes associated with each VR and the cell barcodes associated with candidate AG and TF genes, we ran the *newGOM* function to find significant co-expression ($p_{\text{adj}} \leq 0.05$) for all VRxAG and VRxTF pairs.

VR-specific co-expression of AG and TF genes. To test whether AGs and TFs co-expressed for a given VR, we gathered all cell-barcodes where there was significant VRxAG or VRxTF co-expression. Then, using the same contingency table scheme as above, we looked for significant co-expression ($p_{\text{adj}} \leq 0.05$) between all AGs and TFs previously found to co-express with a given VR.

Spatial transcriptomics

Samples. VNO tissues were dissected from 7-8 weeks old C57BL/6J mice. Briefly, mice were anesthetized with urethane at a dose of 2000 mg/kg body weight. Following general anesthesia, mice heads were decapitated, and the lower jaw was removed by cutting the mandible bone with scissors. The ridged upper palate tissue was peeled off to expose the nasal cavity. A surgical blade was inserted between the two upper incisors to expose the VNO. The whole VNO was carefully extracted by holding onto the tail bone and slowly lifting it up from the nasal cavity. The dissected VNO was immediately transferred to cold 1X PBS on ice, and subsequently embedded in frozen section media (Leica Surgipath FSC 22, Ref # 3801481) and frozen on liquid nitrogen. Frozen samples were sectioned at 10 μm thickness using the Thermo Scientific CryoStar NX70 cryostat. VNO sections were placed within capture area of cold slides that were provided by Resolve Biosciences. Slides were sent to Resolve Biosciences on dry ice for spatial transcriptomics analysis. The probe design, tissue processing, imaging, spot segmentation, and image preprocessing were all performed using the Resolve Biosciences platform. Names and ENSEMBL IDs for genes probed are available in this study's public repository at https://ftp.stowers.org/#!/public/VNO_Atlas/.

Analysis. Regions of interest in the VNO were selected on brightfield images provided by Resolve Biosciences. Cell segmentation on final images was performed in QuPath ¹⁵⁸. Detected gene transcripts were then assigned to the segmented cells, thereby creating a gene-count matrix for each sample. To predict cell types, count matrices for all samples were imported into R then

normalized using the *SCTransform* function with the *glmGamPoi* method. The samples were then integrated using the Seurat integration pipeline. Both the integrated whole VNO scRNA-seq dataset and the integrated Resolve molecular cartography dataset were renormalized with *SCTransform* with the default method using `ncells=3000`, then *RunPCA* was called on the renormalized data. Using the whole VNO scRNA-seq dataset as a reference and the molecular cartography dataset as a query, we ran the *FindTransferAnchors* function, then we ran the *TransferData* function to create a table of prediction score values for each cell in the spatial dataset. Cells were then labeled by type using the maximum prediction score for each cell. Images showing co-localization of vomeronasal receptors were obtained in ImageJ using genexyz Polylux (v1.9.0) tool plugin from Resolve Biosciences.

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FIGURES:

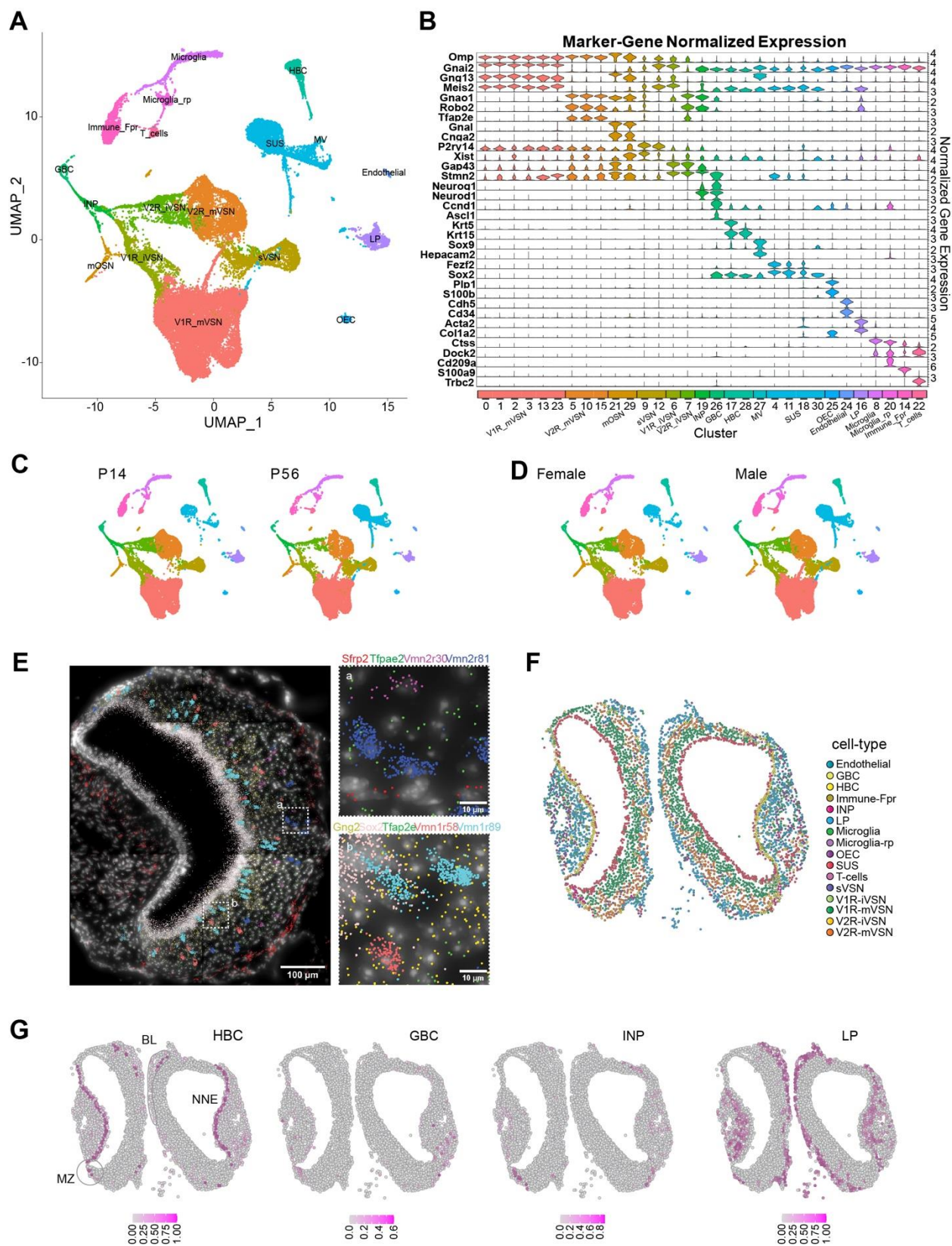


Fig. 1. Single cell transcriptomic profile of the whole vomeronasal organ

A. UMAP visualization of integrated cell-type clusters for whole-VNO single-cell RNA-seq. **B.** Cell-type marker-gene normalized expression across the cell clusters. **C.** UMAP of cell-type clusters split by age. **D.** UMAP of cell-type clusters split by sex. **E.** A representative image of transcript distribution for 9 genes in a VNO slice using the Molecular Cartography© platform Resolve Biosciences. Insets (a and b) show the magnified image of areas identified in the main panel. Individual cell shapes can be determined from the transcript clouds. **F.** Spatial location of individual VNO cells color-coded according to cell type prediction based on the spatial transcriptomic analysis. **G.** Location of cell belonging to HBC, GBC, INP, and LP cell types, respectively. Heat indicates confidence of predicted values. BL: basal lamina; MZ: marginal zone.

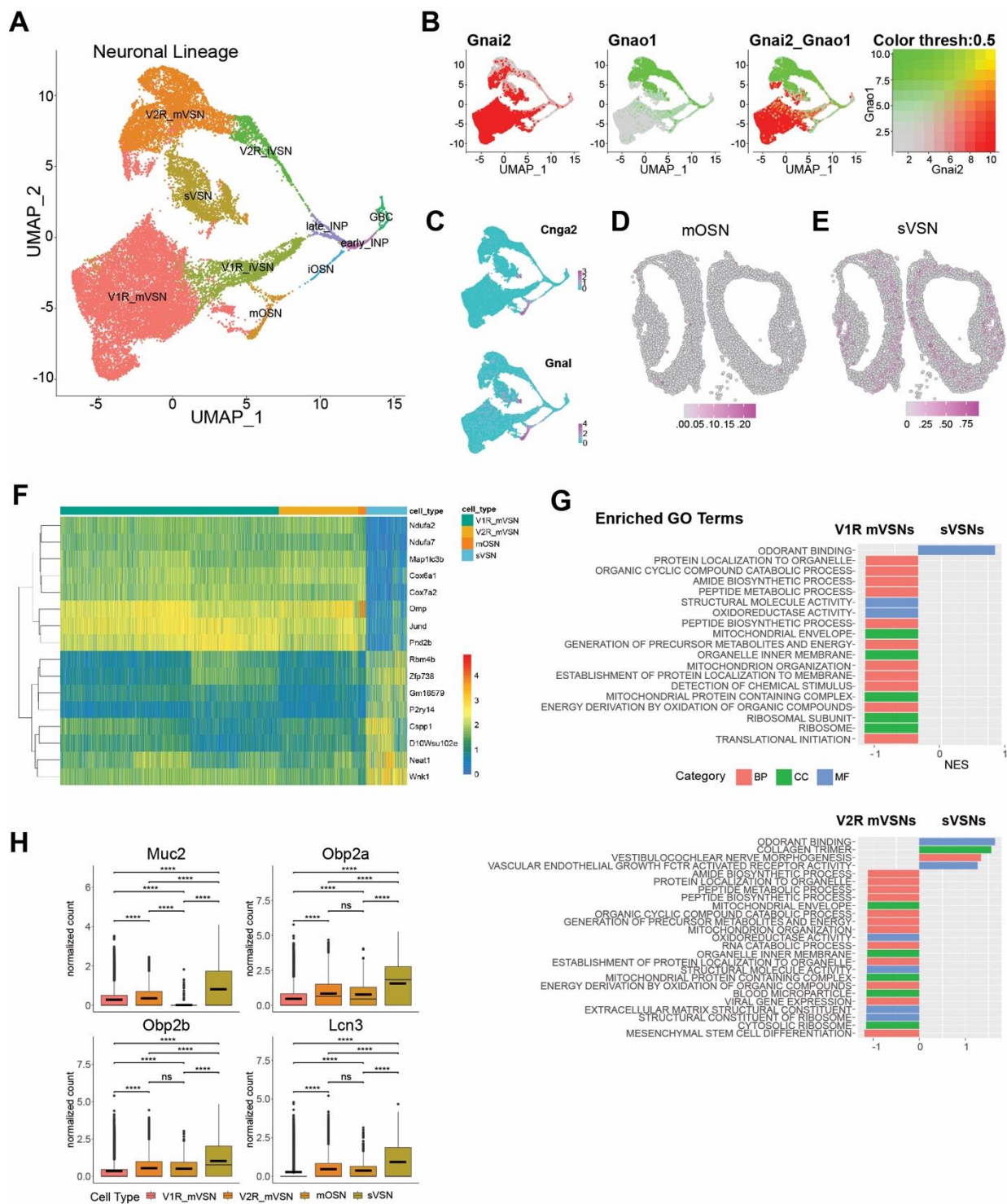


Fig. 2. Novel neuronal lineage

A. UMAP visualization of cell-type clusters for the neuronal lineage. **B.** Expression of *Gnai2* and *Gnao1* in the neuronal lineage. **C.** Expression of *Cnga2* and *Gnal* in the OSN lineage. **D-E.** Location of mOSNs (D) and sVSNs (E) in a VNO slice. Color indicates prediction confidence. **F.** Heatmap of normalized expression for a select set of mutually differentially expressed genes between sVSNs and mature V1Rs, V2Rs, and OSNs. **G.** Enriched gene ontology (GO) terms in sVSNs when compared with V1R and V2R VSNs, respectively. **H.** Box plots of normalized expression for *Muc2*, *Obp2a*, *Obp2b*, and *Lcn3* across mature sensory neurons.

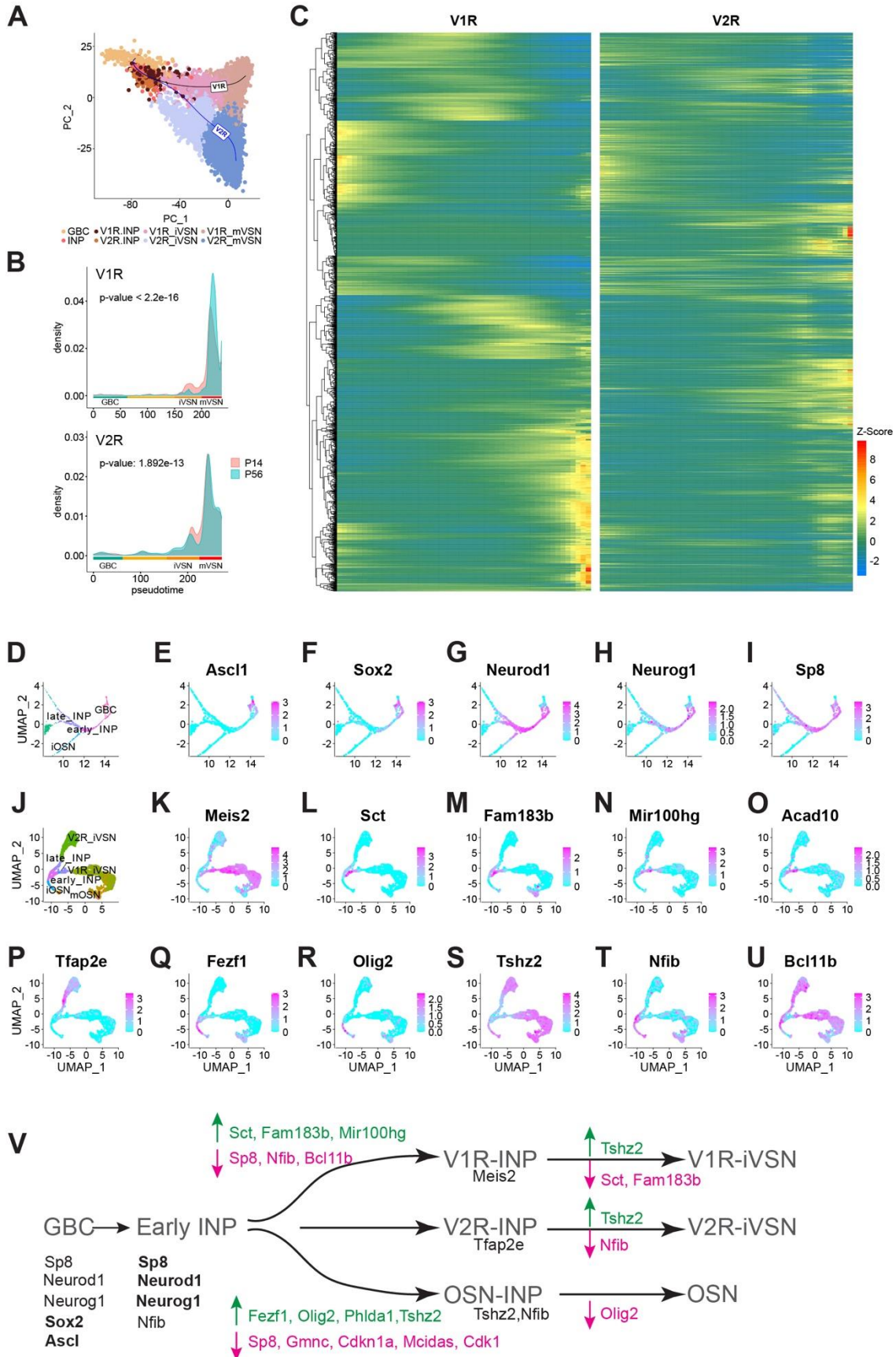


Fig. 3. Molecular cascades separating the neuronal lineages

A. PCA plot of V1R and V2R pseudotime principal curves across cell types. **B.** Cell density plots across pseudotime for P14 and P56 mice. **C.** Heatmap showing expression in pseudotime for genes that differentially expressed between the V1R and V2R lineages. Heat indicates Z-score values. **D.** Zoomed in UMAP of cell types early in the neuronal lineage. **E-I.** Feature plots for select genes expressed early in the neuronal lineage. **J.** UMAP of INP, iVSN, iOSN, and mOSN cell types. **K-U.** Normalized expression of candidate genes for V1R/V2R/OSN lineage determination. **V.** A simplified model for lineage determination by transcription factors in VNO sensory neurons.

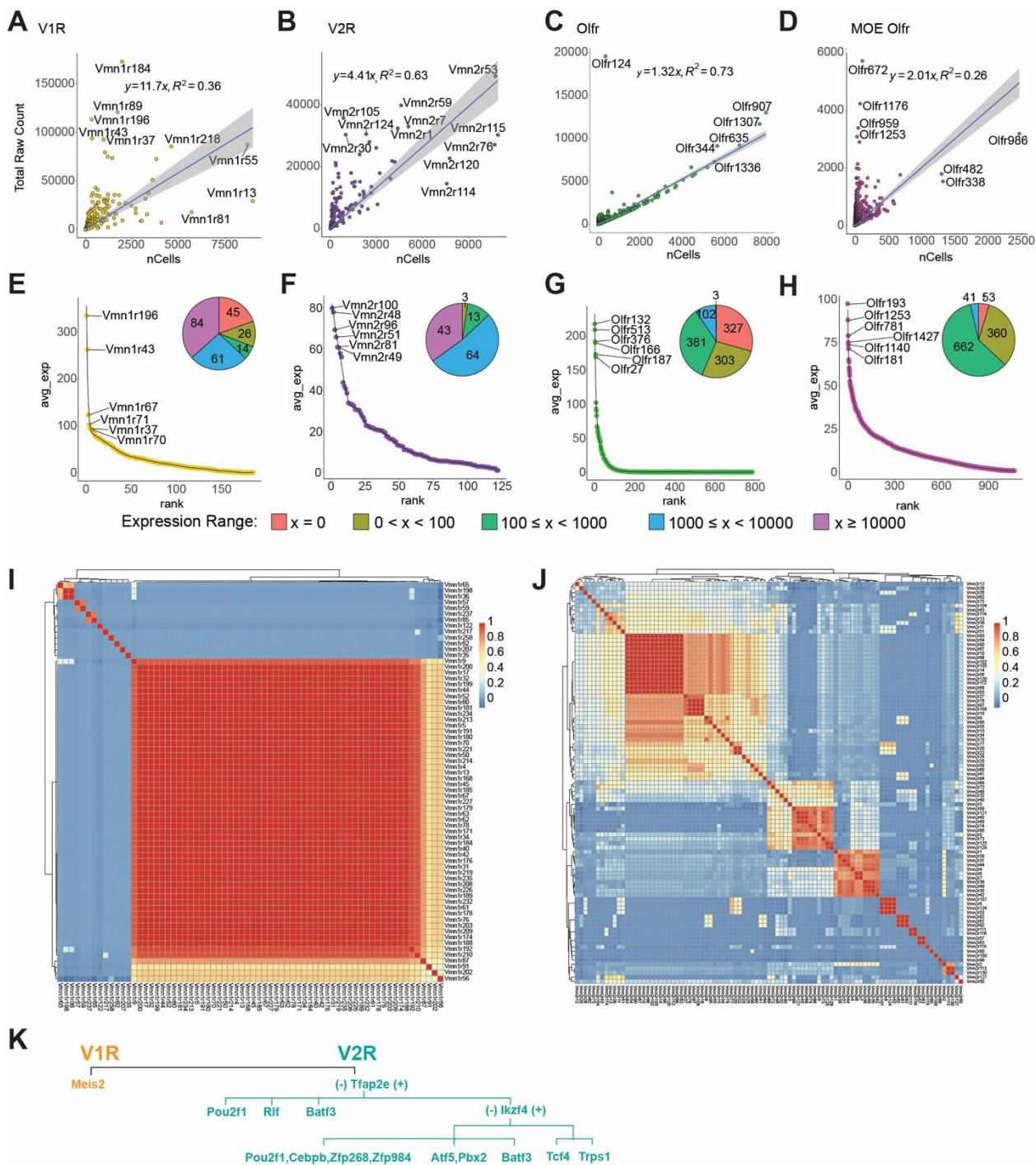


Fig. 4. Receptor expression in the VNO

A-D. Expression level of individual receptor (total raw counts) plot against the number of cells expressing the receptor for V1R (A), V2R (B), VNO-OlfR (C), and MOE-OlfR (D). **E-H.** Ranked distribution of average expression per cell for the four receptor classes. Inset pie charts show the number of cells expressing a receptor at the specified range. **I and J.** Heatmaps showing the correlation of transcription factor expression among the V1Rs (I) or V2Rs (J). **K.** A simplified model of transcription factor selection in mVSNs.

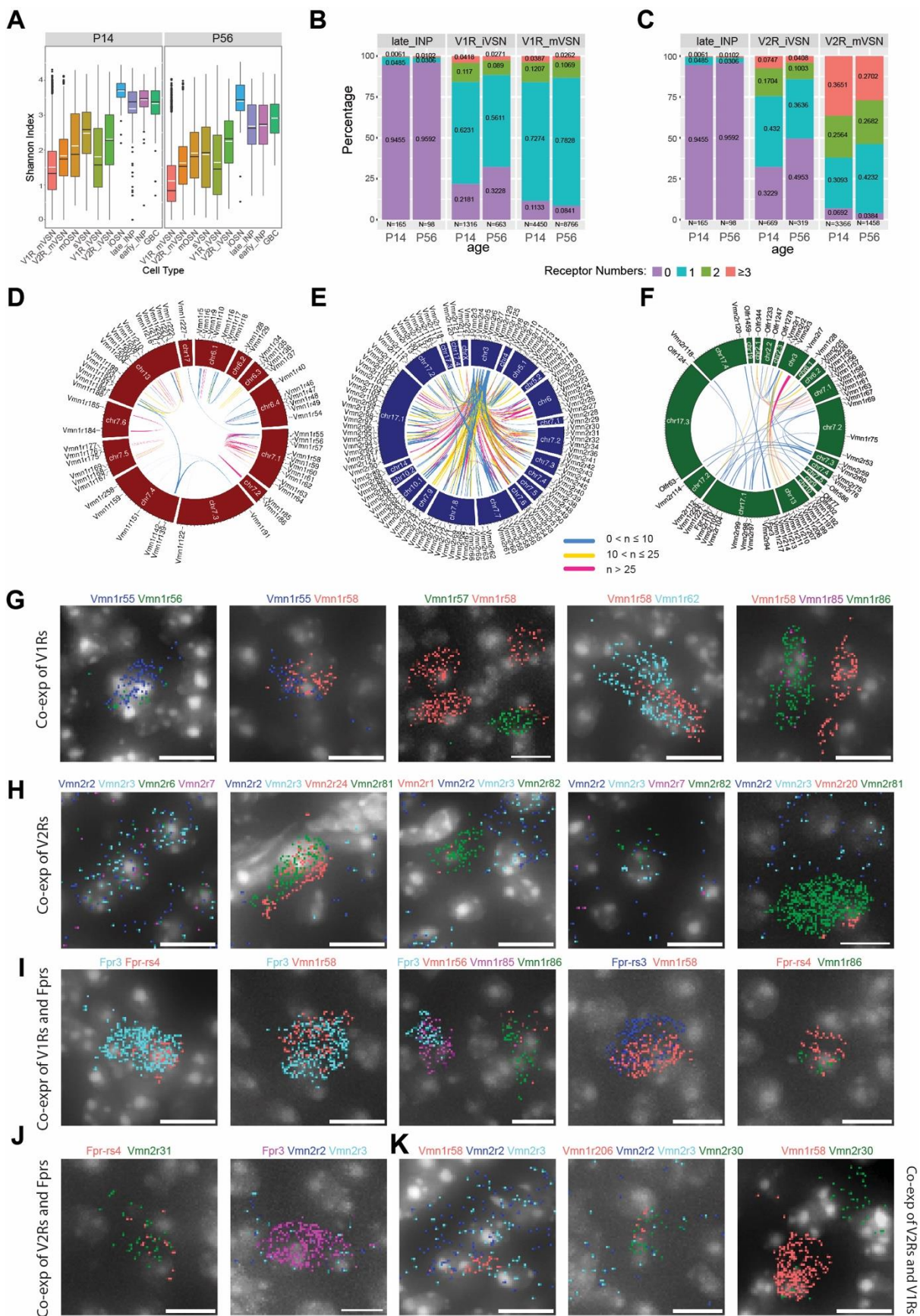


Fig. 5. Co-expression among VNO receptor classes

A. Shannon Indices showing the specificity of receptor expression. High values indicate more co-expressions. **B-C.** Prevalence and level of receptor co-expression by age and cell-type for the V1R (B) and V2R (C) lineages, respectively. **D-F.** Circos plot of genomic loci for significantly co-expressed receptor pairs in the V1R (D), V2R (E), and across-type populations (F). **G-K.** Detection of receptor gene co-expression using Molecular Cartography©. Individual dots represent single molecules. Colors represent different receptor genes. DAPI stain is shown as gray. Scale bar: 10 μm .

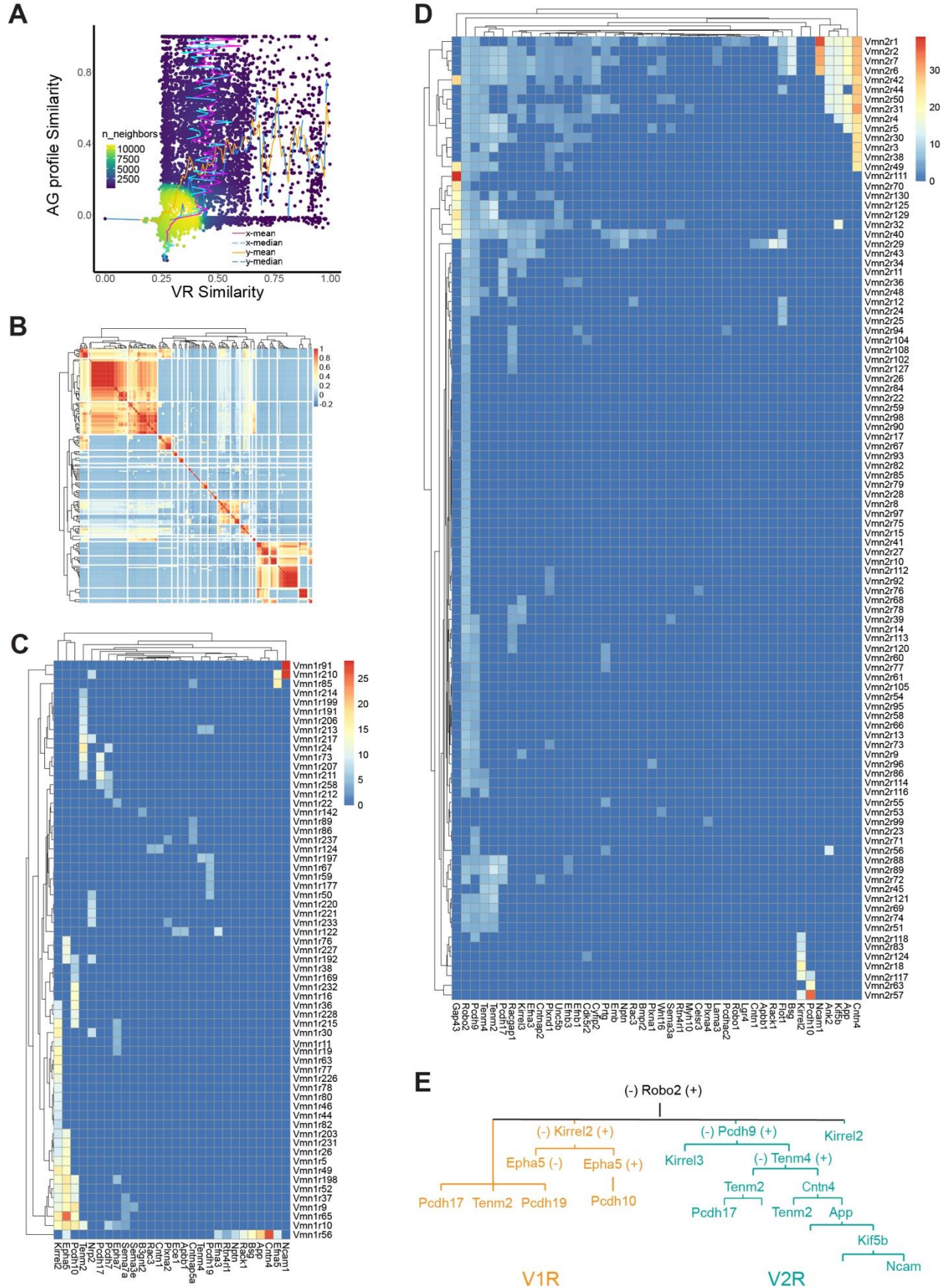


Fig. 6. Axon guidance molecules associated with receptor genes.

A. Distribution density plot showing relationship between similarity of axon guidance (AG) gene expression profiles and receptor sequence similarity. The distribution of receptor similarity (x-mean and x-median) remains constant over the range of AG similarity. The AG similarity (y-mean and y-median) as a function of receptor similarity shows a strong correlation at the dense part of the curve. **B.** Heatmap showing the correlation among VRs in their AG expression. **C** and **D.** Heatmaps showing the V1R-AG (C) and V2R-AG (D) associations. Heat shows average expression level for AG genes for a given receptor type. **E.** A simplified model of hierarchical distribution of AG in the mVSNs.

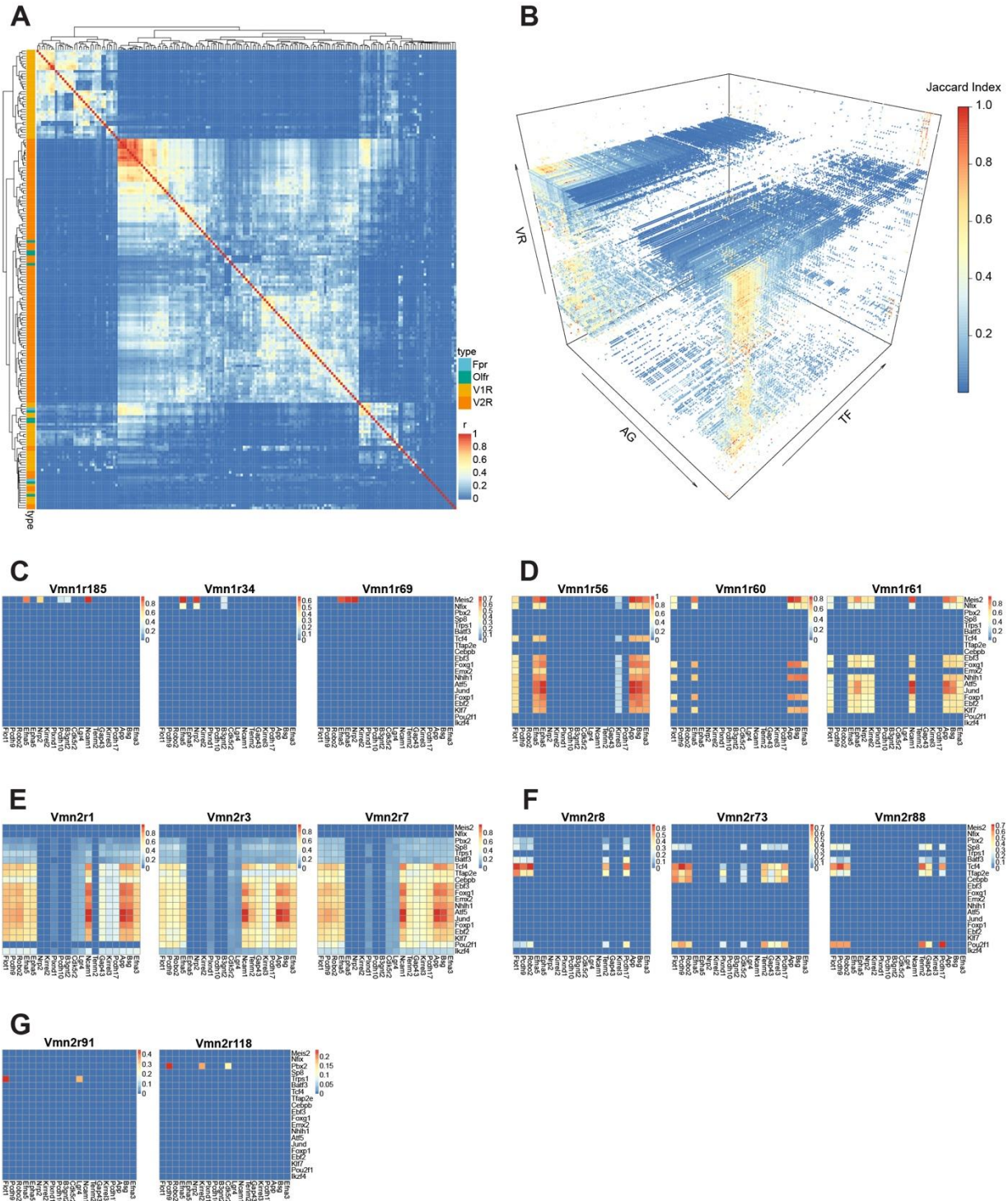


Fig. 7. Transcriptional determinants of axon guidance molecules for individual receptor types.

A. Correlation heatmap between receptor types calculated from co-expressed AG and TF genes. Receptor types are color-coded. **B.** 3-D heatmap showing the Jaccard Indices between AG and TF genes for each of the VRs in the dataset. **C-G.** Heatmaps showing Jaccard Indices of TF-AG associations for V1R (C and D) and V2R types (E-G). The lists of TF and AG genes here are abridged from the full list to enhance visualization. (C), these receptors share Meis2 expression but different AG genes. Note that Vmn1r185 and vmn1r69 both recognize female identify pheromones. (D), similarity and distinction of AG/TF expression for three V1R types that are located in the same genomic location and with high sequence homology. (E and F), shared TFs and AG genes for broadly (E) and uniquely (F) expressed V2R types. (G), distinct TFs and AGGs for uniquely expressed V2R types.