Developmental divergence of sensory stimulus representation in cortical interneurons

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Vasocative-intestinal-peptide (VIP+) and somatostatin (SST+) interneurons are involved in modulating barrel cortex activity and perception during active whisking. Here we identify a developmental transition point of structural and functional rearrangements onto these interneurons around the start of active sensation at P14. Using in vivo two-photon Ca2+ imaging, we find that before P14, both interneuron types respond stronger to a multi-whisker stimulus, whereas after P14 their responses diverge, with VIP+ cells losing their multi-whisker preference and SST+ neurons enhancing theirs. Additionally, we find that Ca2+ signaling dynamics increase in precision as the cells and network mature. Rabies virus tracings followed by tissue clearing, as well as photostimulation-coupled electrophysiology reveal that SST+ cells receive higher cross-barrel inputs compared to VIP+ neurons at both time points. In addition, whereas prior to P14 both cell types receive direct input from the sensory thalamus, after P14 VIP+ cells show reduced inputs and SST+ cells largely shift to motor-related thalamic nuclei.

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Postnatal development and maturation of neuronal circuits responsible for sensory processing is fundamental for accurate representation of the environment as animals transition into actively interacting with the external world. Unravelling the mechanisms of neocortical development is key for understanding the emergence of purposeful and goal-directed motor actions informed by sensory cues. It is therefore of great importance to study the alterations that sensory cortical networks undergo in responding to diverse sensory stimuli upon developmental behavioural transitions.

The whisker primary somatosensory cortex (wS1) of rodents is a well-suited model to study sensory processing throughout development due to the somatotopic way in which the information is transmitted from the whiskers to the cortex, as well as its role in spatial navigation. Although the whiskers of mice are already present at the time of birth, during the first postnatal weeks the whisker pad only exhibits spontaneous muscle twitches, which coincide with activity in wS1. Around postnatal day 14 (P14), mice start displaying bilateral rhythmic movements of the whiskers (“active whisking”), enabling them to explore their environment and extract more detailed information from their surrounding.

Although the wS1 has been a subject of research for many decades, it is only recently that the role of inhibition in sensory processing in the adult cortex has started to be explored. Several studies showed that layer 2/3 (L2/3) vasoactive intestinal peptide-expressing (VIP+) interneurons (INs) in the wS1 can inhibit the inhibitory somatostatin-expressing (SST+) INs, thus leading to a net excitation of pyramidal cells (disinhibition). The same disinhibitory connectivity motif has also been described in other cortical areas, suggesting a general mechanism by which cortico-cortical loops can influence cortical processing. In the wS1, this VIP+-SST+ disinhibitory loop can be recruited in a top-down fashion by the whisker primary motor cortex (wM1), which directly innervates the wS1 VIP+ INs in L2/3. In addition, it has been shown that VIP+ INs can be activated by bottom-up thalamic inputs, potentially also engaging in disinhibition through SST+ cells. Nevertheless, a recent study reports opposite results, with SST+ cells being strongly activated by a whisker-driven sensory stimulus, compared to VIP+ cells which are found to be silenced. We hypothesized that the main reason behind this discrepancy between the aforementioned results is the stimulus paradigm used. Researchers either stimulated one single whisker or multiple whiskers at the same time, two paradigms that would engage the adult barrel cortex in very distinct ways. We further hypothesized that these INs would be differently engaged by these stimulation paradigms prior to P14, when top-down modulation is absent and discrimination of the two cell types in response to this stimulation paradigm. We found that at P21+- the peak ΔF/F of the average Ca2+ response in SST+ INs was lower compared to that of VIP+ INs, matching the published data. Nevertheless, when looking at the integral of ΔF/F average response (over 8 s following whisker stimulation) there was no significant difference between the two cell types, and the same was true before P14.

As the response to a single-whisker stimulus in adult L2/3 VIP+ and SST+ INs has been described before by cell-attached electrophysiological recordings, we first compared the activity of the two cell types in response to this stimulation paradigm. We found that at P21+- the peak ΔF/F of the average Ca2+ response in SST+ INs was lower compared to that of VIP+ INs, matching the published data. Nevertheless, when looking at the integral of ΔF/F average response (over 8 s following whisker stimulation) there was no significant difference between the two cell types, and the same was true before P14.

We then assessed how IN responses to multi- compared to single-whisker stimulation may change during development by comparing the ΔF/F integral as an overall measure of direct or indirect activation of the cells by the sensory stimuli. The analysis showed that before the onset of active whisking (P8–12), both VIP+ and SST+ INs responded significantly stronger to multi- compared to single-whisker stimulation (Fig. 1d, f). For the SST+ INs this continued to be the case and even enhanced after the onset of active whisking (P21+), while for the VIP+ INs the ΔF/F integral became similar between multi- versus single-whisker stimulation (Fig. 1f).

Since the Ca2+ transients recorded at P8–12 had a clear late component and lower peak amplitude than the mature cells, we sought to better understand how they relate to action potentials calcium (Ca2+) imaging under light-anesthesia at two developmental time points; prior to the onset of active whisking (age: P8–12) and after the beginning of active whisking (age: P21–41, denoted as P21+). To visualize VIP+ and SST+ IN activity, we used animals expressing tdTomato in either VIP+ or SST+ INs (VIPCre-Ail4 and SSTCre-Ail4 lines, respectively) and injected the membrane-permeable AM-ester form of the Ca2+ indicator OGB-1 into wS1. Two-photon Ca2+ imaging was performed during spontaneous activity and upon stimulation of either the principal C2 whisker one time alone (single-whisker stimulation) or both the principal whisker and the majority of the macro vibrissae once together (multi-whisker stimulation) (Fig. 1b).

We first tested whether VIP+ and SST+ INs are functionally integrated before P14 by examining their spontaneous activity. We observed prominent Ca2+ transients in both IN types that correlated with the activity of the surrounding cells (Figs. 1c and 1a), indicating that they receive input capable of driving action potentials before the onset of active whisking. Previous studies have shown that cortical pyramidal neurons and superficial 5HT3aR+ INs display highly correlated spontaneous activity within their population in early postnatal development and then undergo a de-correlation upon maturation of the circuit. To test if the same is true for VIP+ and SST+ INs, we compared the correlations of spontaneous activity within each IN population before and after P14 and found a significant drop of co-activation with time (Fig. 1e). Interestingly, at P21+, spontaneous activity of SST+ cells is more correlated than that of VIP+ cells, a phenomenon that has also been described in the visual cortex of adult mice. In the wS1, the single-whisker deactivation of VIP+ INs was missing and SST+ INs were more strongly activated by a whisker-driven sensory stimulus, compared to VIP+ cells. We therefore assessed how IN responses to multi- compared to single-whisker stimulation may change during development by comparing the ΔF/F integral as an overall measure of direct or indirect activation of the cells by the sensory stimuli. The analysis showed that before the onset of active whisking (P8–12), both VIP+ and SST+ INs responded significantly stronger to multi- compared to single-whisker stimulation (Fig. 1d, f). For the SST+ INs this continued to be the case and even enhanced after the onset of active whisking (P21+), while for the VIP+ INs the ΔF/F integral became similar between multi- versus single-whisker stimulation (Fig. 1f).

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across development. We therefore filled SST$^{+}$ INs with OGB-1 in vitro and evoked a set number of action potentials (APs), while recording the Ca$^{2+}$ transients in the cell bodies using two-photon microscopy. We found, that when comparing the ΔF/F responses between the two age groups, there was no significant difference, indicating that the integral of the Ca$^{2+}$ responses could provide a good estimate for the functional activation of the cell types across development (Fig. S1d). Nonetheless, it is worth noting that at both ages the time constant of the decay (tau) increases with the number of APs elicited, with the Ca$^{2+}$ response to 10 APs.
Fig. 1 Divergent sensory stimulus responses in superficial cortical VIP+ and SST+ interneurons during development. a Acute in vivo two-photon Ca²⁺ imaging was performed before (P8–12) and after (P21+) the start of active whisking. b Schematic representation of the whisker stimulation protocol. c Top: average intensity projection of Ca²⁺ imaging regions after bulk loading of OGB-1 (green). Interneurons are labeled with tdTomato using reporter mouse lines; scale bar 35 µm. Bottom: representative examples of raw ΔF/F traces of spontaneous activity of two interneurons (INI/2) and two surrounding cells (Cl2). (N = 3 animals per group, VIP P8–12: 109 cells, SST P8-12: 38 cells, VIP P21+: 138 cells, SST P21+: 51 cells). d Top: ΔF/F signal over time of all recorded VIP and SST cells after whisker stimulation. Ca²⁺ responses are baseline corrected and aligned to whisker stimulus onset (dashed line). Cells are sorted by their cumulative activity following multi-whisker stimulation. Same number of cells are included as in c. Bottom: Average ΔF/F responses with SEM. c Pearson’s correlations of spontaneous neuronal activity within each interneuron type before- and after P14. Same number of cells are included as in c, but the data are plotted by imaging spot. Statistics: two-sided Mann-Whitney U test, p(VIP P8–12) = 6.298 × 10⁻⁵, p(SST) = 0.047. f Distribution of average values of the evoked ΔF/F integral after single and multi whisker stimulation for all cells analyzed. Pie charts show the fraction of cells that increase (coloured) or decrease (white) in activity by multi- vs single whisker stimulation. Same number of cells as in c. Statistics: two-sided paired Wilcoxon signed-rank test, p(VIP P8–12) = 1.368 × 10⁻⁵, p(SST P8–12) = 0.031, p(SST P21+) = 5.576 × 10⁻⁷. g Average ΔF/F traces with SEM of evoked and spontaneous events. Fitted curve of decay time constant is plotted in black (continuous line: evoked, dotted line: spontaneous). Insets show the cumulative distribution of the decay tau. Statistics: two-sample Kolmogorov-Smirnov test; VIP P8–12: p(spont vs. single) = 1.257 × 10⁻⁴, p(spont vs.multi) = 2.611 × 10⁻⁵; SST P8-12: p(spont vs. single) = 0.034, p(spont vs. multi) = 2.615 × 10⁻⁴. Source data are provided as Source Data file.

returning to baseline two-times slower than the response to 1 AP (Fig. S1d). This non-linearity could either be caused by saturation of Ca²⁺ extrusion mechanisms when the cell is repeatedly activated or due to the diffusion of Ca²⁺ buffer between the cell soma and the patching pipet during the experiment. Furthermore, even though the prolongation in decay tau after multiple APs is about two-fold at both ages, at P8–12 the ΔF/F integral increases much more than that (Fig. S1d), likely due to the immature state of the neurons.

To further investigate developmental changes in Ca²⁺ signaling, we analyzed the decay of spontaneous activity at P8–12 and P21+ and found that both in VIP+ and SST+ INs Ca²⁺ dynamics sharpen as the animal matures, with spontaneous transients dissipating faster at P21+ than at P8–12 (Fig. S1e). We then compared the decay tau of spontaneous and evoked activity in VIP+ and SST+ at the two different time points. Our results show that at P21+ the decay of Ca²⁺ signal is very similar, regardless of the cell type and of how the activity was elicited (spontaneous, single whisker, multi whisker) (Fig. 1g). At P8–12 on the other hand, the decay of spontaneous activity is much faster than that of evoked activity (Fig. 1g). Furthermore, spontaneous events show a larger amplitude and shorter duration than evoked ones (Fig. 1g). The data suggest that the pronounced late component of the evoked Ca²⁺ signal at P8–12 is not due to immature Ca²⁺ buffering capacity, but rather due to the way the network activates the cells. Finally, to address if the sensory-evoked Ca²⁺ responses we observed for the cell types and stimuli across development carry information that can make them discriminatory, we trained three different decoders to quantify differences in response profiles. Each decoder was trained on a subset of Ca²⁺ responses, evoked through either multi- or single-whisker stimulation, for each IN type and age. All of the classifiers showed above chance level values with VIPs displaying a reduction with age, whereas the SSTs an increase. This finding lends further support for the developmental divergence of these two cell type’s activation and function (Fig. S1f).

Superficial VIP+ and SST+ interneurons show distinct barrel-field afferent connectivity motifs. Increased responsiveness to multi- over single-whisker stimulation at both ages could be due to the cortico-cortical inputs originating from surrounding barrel columns, which would provide convergent excitation originating from multiple whiskers. We therefore aimed at investigating the presynaptic inputs onto L2/3 VIP+ and SST+ INs at the two developmental time points, using a monosynaptic rabies virus approach, which allowed us to assess both local and long-range connectivity.

By utilizing compound mouse genetics (VIPCre-HTB or SSTCre-HTB), in combination with the pseudo-typed rabies virus, direct presynaptic partners of L2/3 VIP+ and SST+ INs were labeled with mCherry. The primary infected cells (starter cells) could be identified by double labeling of mCherry and eGFP (provided by the HTB line). The viral injections were done at either P5 or P15 and after 7 days the brains were collected (at P12 or P22 respectively). A CLARITY-based tissue clearing protocol was performed on the brains and they were subsequently imaged in their entirety using a custom-built light-sheet microscope (mesoSPIM) (Fig. 2a). The auto-fluorescence of the barrels allowed for the detection of the barrel field and enabled us to accurately localize our injection site within the wS1 (Figs. 2b–e and S2). A second batch of injected brains was processed using classical histology, followed by eGFP immunostaining and widefield imaging. Both the cleared whole brain and the histology dataset were used to quantify the number and laminar position of the starter and the presynaptic cells located around the wS1 injection site. This dual approach was performed to safeguard that no information was lost while clearing the brains. For the histology dataset, a subset of all cut sections was used to identify the cells manually, whereas in the cleared brains a deep-neural network was trained to detect and count cells. For both methods, the results were highly comparable (Fig. S3).

The analysis showed that although there are starter cells in L2-6 at both time points we get robust primary injection in L2/3 for both VIP+ and SST+ cells (Fig. S3a, b). The distribution of the presynaptic cells across the layers mirrored that of the starter cells, with the VIP+ INs receiving more inputs from the upper cortical layers while the SST+ neurons more from the deeper layers (Fig. S3c, d).

To assess if L2/3 SST+ and VIP+ INs receive a significant proportion of their inputs from neighboring barrels, the cleared 3D brain images were used to localize both starter and presynaptic cells within the barrel field (Figs. 2b–e and S2a, b). Only L2/3 presynaptic neurons located within wS1 were included in the analysis, since it is most likely that these inter-columnar inputs would originate from the upper layers. As it is not possible to determine which presynaptic partner connects to which starter cell, a probabilistic approach was taken, where the distance between every starter to every presynaptic cell within 800 µm distance was calculated and compared between the two cell types (Fig. S4). The analysis revealed that SST+, compared to VIP+ INs, receive on average more distant inputs, both before and after P14 (Fig. 2f). To further validate the results and overcome the variability observed in the data we also calculated the cumulative distance distribution of randomly selected starter and presynaptic cells (Fig. S2c) (Online Methods). This anatomical data can help explain why the SST+ cells are able to report multi- over single-whisker activation both before and after...
However, it cannot explain why the VIP⁺ INs are able to differentiate between multi- and single-whisker stimulation before P14 and not after, unless there is a change in the functional connectivity.

Distinct functional connectivity motifs onto superficial VIP⁺ and SST⁺ interneurons. After revealing the anatomical connections of the wS1 cortical inputs onto superficial VIP⁺ and SST⁺ INs, we assessed their functionality and strength. Specifically, we investigated if after P14, SST⁺ INs receive stronger functional input from lateral sources compared to VIP⁺ INs, while before P14 the two cell types display more functionally similar input.

In order to study the developmental trajectory of VIP⁺ and SST⁺ IN intrinsic properties we performed whole-cell current-clamp recordings of L2/3 VIP⁺ or SST⁺ INs and analyzed their passive and active electrophysiological membrane properties (Figs. S5a–e). As expected, we found evidence of cell maturation in both cell types, with the membrane resistance decreasing and the action potentials becoming faster (Fig. S5a, d, e). Nevertheless, the threshold for action potential generation dropped and action potential amplitude increased only in VIP⁺ INs, (Fig. S5b, c), suggesting that the SST⁺ cells are closer to their mature state at P8–12.

In order to functionally analyze the distribution of neurons providing synaptic input to the two IN types, we performed glutamate uncaging. This was carried out using light
Fig. 3 Distinct functional connectivity motifs onto superficial VIP$^+$ and SST$^+$ cortical interneurons. a Left: a graphic representation of photostimulation calibration with a patched putative pre-synaptic neuron within a neocortical slice indicated with layer boundaries (L1-6). Illumination calibration were done so that the paradigm successfully evoked a short train of APs in the putative pre-synaptic partners. AP discharge is displayed to the right, with the peak of the first AP until after the last AP creating a time-window used for future analysis of evoked currents. Right: schematic representation of photostimulation-based mapping of incoming currents onto interneurons through whole-cell patch-clamp recordings. Overlaid is a schematic of the 20 × 20 grid, indicating the quadrants that can be photostimulated. Two of the AP discharges of the interneuron is shown to the right, with one not eliciting (1) and one eliciting (2) a post-synaptic response. The time-window acquired during calibration is used to register and analyse the currents. b Top: example traces of voltage responses to square hyperpolarizing and depolarizing currents pulses, $\Delta \pm 20 \text{pA}$. Bottom: Heatplot representations of normalized evoked excitatory current integral (in pC), recorded over development (P8-12 and P21+) from VIP$^+$ (left) and SST$^+$ (right) IN while perfoming glutamate uncaging in a grid pattern (VIP P8-12: 11 cells, VIP P21+ and SST P8-12: 10 cells, SST P21+: 9 cells). c plots of excitatory input onto VIP$^+$ and SST$^+$ cells (individual dots) averaged per lamina and normalized to average overall excitation within the field of view. The grand average of all cells per group is depicted as a continuous filled wave and compared within age groups. One data point has been excluded for display purposes only. Same number of cells as in b. Statistics: two-sided Mann-Whitney U test, $p$(L2/3 P8-12) = 4.909 × 10^{-4}, $p$(L4 P8-12) = 0.018, $p$(L2/3 P21+) = 0.0015. d Mean evoked excitation originating from within L1-3 and plotted as a function of lateral distance from either side of the recorded cells somata. Same number of cells as in b. Statistics: two tailed t-test ($^*p<0.05$, $^{**}p<0.01$, $^{***}p<0.001$). Source data are provided as a Source data files.

stimulation in a grid pattern in wS1-containing brain slices while measuring evoked postsynaptic excitatory and inhibitory currents in L2/3 VIP$^+$ or SST$^+$ INs through whole-cell voltage-clamp recordings at −45 mV (Fig. 3a). With this approach, we mapped out the origin as well as the strength of incoming cortical connections at the two developmental stages investigated (P8-12 and P21+). We found that L2/3 SST$^+$ INs receive the majority of their excitatory input from within their own layer and that this is the case both before and after P14. These results fit well with the laminar patterns of excitatory inputs described for the L2/3 SST/CR$^+$ cells in adult mice. The VIP$^+$ cells on the other hand receive almost equal amounts of excitatory inputs from all layers, with a decrease in L5/6 (Fig. 3c). Because the photostimulation was carried out over a large field of view we were able to assess lateral activity originating from up to 500 µm away from either side of the IN cell body and therefore inputs coming from outside the resident column. To compare our glutamate uncaging results to the rabies virus mapping, inputs originating in the superficial layers (L1-3) were plotted in relation to their lateral distance from the recorded cell. In agreement with the rabies virus findings, the analysis showed that prior to P14 SST$^+$ INs receive excitatory inputs from more lateral sources compared to VIP$^+$ cells (Fig. 3d). Even though distal inputs onto both cell types seem to have undergone pruning at P21+ (Fig. 3d compare upper and lower panel), SST$^+$ INs still receive a higher proportion of excitatory inputs from further away compared to VIP$^+$ INs (Fig. 3d). Interestingly, the VIP$^+$ INs receive more inhibition than the SST$^+$ INs from L2/3 both before and after P14, acting almost in an opposite fashion to the mapped incoming excitation (Fig. S5f, g). Because glutamate uncaging could activate the receptors on the patched cell directly, and holding the cell at −45 mV does not allow for a proper estimation of purely excitatory responses, a new set of photostimulation experiments was carried out in voltage-clamp mode at −70 mV, first without and then in
the presence of tetrodotoxin (TTX). By subtracting the direct responses (in the presence of TTX) from the evoked excitatory ones, recorded before the addition of the drug, currents evoked exclusively from pre-synaptic neurons could be calculated (Fig. S5h, i). The analysis of this new dataset showed that the SST+ INs do indeed receive excitation from more distant sources compared to the VIP+ cells within the superficial layers (Fig. 3d).

The anatomical tracing data together with the functional incoming excitation data suggest that SST+ INs receive more lateral excitation compared to VIP+ cells at both developmental time points. This provides a plausible mechanism for the stronger activation of SST+ INs by multi-whisker stimulation both before and after P14. Although VIP+ cells only respond stronger to multi- compared to single-whisker stimuli before P14, they do not show striking changes in cross-columnar inputs across the two ages. Nonetheless, small changes in connectivity can have a large functional effect given the slow membrane time constant (Fig. 3b) and high membrane resistance of immature VIP+ cells (Fig. S5a). Therefore the developmental changes observed in VIP+ cells for multi-whisker activation could be enabled by small changes in local cortical connectivity motifs. However, this does not preclude that VIP+ INs might also undergo changes in long-range thalamic connectivity.

Rearrangement of thalamic inputs onto superficial VIP+ and SST+ interneurons during development. Since the observed inter-barrel connectivity motifs are not able to fully explain the developmental changes in single- versus multi-whisker responses seen in the in vivo Ca2+ imaging data, we turned our attention to bottom-up inputs coming from the thalamus, labeled by our rabies tracings. The analysis was facilitated by the preservation of the 3D architecture of the thalamus in the cleared whole-brain dataset. Interestingly, we found that before P14, both VIP+ and SST+ INs had many presynaptic cells in the thalamus (Fig. 4a, c and Movies S1 and S3), mainly located in the ventral posteromedial nucleus of the thalamus (VPM), but also in the posterior complex (PO), a higher-order nucleus primarily innervating VIP+ cells (Fig. S5a). Therefore the developmental changes observed in VIP+ cells for multi-whisker activation could be enabled by small changes in local cortical connectivity motifs. However, this does not preclude that VIP+ INs might also undergo changes in long-range thalamic connectivity.

Discussion

This study investigates the engagement of VIP+ and SST+ INs by sensory stimuli, and the underlying circuits they are embedded in before and after the onset of active whisking (P14).

We initially hypothesized that in the juvenile mouse wS1, VIP+ and SST+ cells would be differentially activated by a single-versus a multi-whisker deflection based also on published reports[11,20]. By testing the activation pattern of the same population of INs for the two types of stimuli, we found support for our hypothesis. VIP+ INs at P21+ responded in the same manner whether single or multiple whiskers were deflected, even though prior to P14 they responded much stronger to the multi-whisker stimulus. SST+ cells at P21+ on the other hand responded stronger to the multi-whisker stimulus, which was also the case before P14. Even though in this study we have not directly examined the connectivity between these two interneuron types, recently published work in the adult mice showed strong layer-dependent effects of channelrhodopsin-based activation of VIP+ neurons[20]. Specifically, the authors reported that L2-4 excitatory neurons did not increase their firing rate upon VIP+ neuron activation, suggesting that the recruitment of the VIP-SST inhibitory loop is layer- and most likely also stimulus-dependent. On the other hand, the same study suggests that upon active whisking and stimulation of multiple whiskers, SST+ cells are strongly activated, whereas VIP+ INs are inhibited with a delay that matches the activation of SST+ neurons. These results are in line with the reciprocity of the connections between the two IN types[12], and together with our data suggest that the di-synaptic disinhibitory VIP-SST connection is engaged in a dynamic manner, with one cell inhibiting the other, the directionality of which is context-dependent.

Developmentally, the robust activation of both VIP+ and SST+ INs prior to P14 by the multi- compared to the single-whisker stimulus, suggests that even if the connectivity between VIP+ and SST+ cells is present at this time point, it seems to be overridden by either local or long-range thalamic inputs onto both IN types.

Interestingly, we found striking developmental changes in the Ca2+ dynamics of the two INs, suggesting that VIP+ and SST+ cells undergo significant maturation around the time the mouse starts to actively whisk. First, we found that before P14 the amplitude of whisker-stimulation-evoked Ca2+ activity is much lower than that of spontaneous events, suggesting that evoked responses have a different origin to spontaneous ones or that the former are being actively decreased upon sensory stimulation during this developmental period. Both VIP+ and SST+ cells receive stronger and more widespread inhibitory inputs at P8–12 which could potentially decrease the amplitude of somatic Ca2+ transients through GABAA and/or GABAB receptor-mediated mechanisms. Furthermore, evoked Ca2+ transients recorded before P14 are slower and more prolonged compared to both spontaneous activity

support the data obtained with the rabies-based mapping, and suggest that upon the onset of active whisking thalamic input onto VIP+ INs is strongly reduced.

Overall, our data suggest a model by which the VIP+ IN preference for multi-whisker stimulation before P14 is mainly supported by strong thalamo-cortical inputs that come from both VPM and PO. These inputs are significantly reduced after the onset of active whisking, together with the VIP+ INs response to multi- versus single-whisker stimulation (Fig. 5). For the SST+ INs on the other hand, the preference for multi-whisker stimulation, both before and after P14 would be supported by the strong lateral inter-barrel connectivity, with a contribution of direct VPM and PO-derived excitation prior to P14 (Fig. 5).
Fig. 4 Rearrangement of thalamic inputs onto superficial VIP\textsuperscript{+} and SST\textsuperscript{+} interneurons during development. a–d Top panel: representative examples of cleared rabies injected brains seen from an oblique, top-down and side angle. The thalamus is highlighted in yellow. The red square indicates the area projected in bottom panel. Bottom panel: Coronal view of maximum intensity projection of the thalamus. Left: overview. Right: zoom-in with VPM (green) and PO (blue) highlighted. N = 3 brains for SST; N = 4 brains for VIP P12). e Quantification of pre-synaptic cells in different thalamic nuclei normalized to the total number of cells in the thalamus. VIP P22 is not included because no cells were found in the thalamus. Same number of brains as in a–e. f Example of VGlut2 staining (green) in L2/3 of a wS1 section from a VIPCre-tdTomato animal at P9. The dotted square indicates the zoomed-in part on the right, which shows a close-up of puncta appositions in xy and yz direction. The graph shows quantification of VGlut2 puncta on tdTomato-positive dendrites. The number of appositions in every picture is normalized to the area of tdTomato positive dendrites in the same picture. (N = 3 brains per group; VIP P8–12: 23 images, VIP P21+: 27 images, SST P8–12: 21 images SST P21+: 28 images. Statistics: two-sided Mann–Whitney U test, \( p(\text{VIP}) = 2.492 \times 10^{-7} \). Source data are provided as a Source data file.
at the same age and evoked activity after P14. There are a number of potential mechanisms that could explain changes in evoked transients over time such as developmental changes in action potential speed leading to a slower intracellular Ca\textsuperscript{2+} rise in the cytoplasm or altered buffering and/or extrusion capacity of the cells. Nevertheless, the fact that spontaneous Ca\textsuperscript{2+} signals before P14 do not show the same dynamics as the evoked suggests that the prolongation of the signal is not due to a cell-intrinsic mechanism, but rather to the network activity engaged upon sensory stimulation. For example, reverberating networks could underlie the generation of multiple action potentials spread across time and lead to slower Ca\textsuperscript{2+} transients. This may be especially the case when multiple whiskers are deflected. In this case, the signal would activate the VPM, which through direct connections activate the VIP\textsuperscript{+} and SST\textsuperscript{+} INs. In addition to this early activation, there would be a later wave of activity onto INs via the engagement of a parallel pathway, consisting of a loop between these INs in L1 and PO\textsuperscript{37,38}, leading to the prolonged Ca\textsuperscript{2+} transient component. Furthermore, our sW1 distance-dependent anatomical and functional connectivity analysis show that in neonatal mice there is strong widespread cortical input in both cell types which could also contribute to the prolonged Ca\textsuperscript{2+} transients before P14. Nevertheless, excitation coming from lateral cross-barrel domains is stronger onto SST\textsuperscript{+} compared to VIP\textsuperscript{+} cells both before and after P14, whereas inhibition from within layers 2/3 is stronger onto VIP\textsuperscript{+} cells. This higher inhibition onto the latter cells, could be originating from multiple types of presynaptic GABAergic cells, including other VIP\textsuperscript{+} cells, reelin-expressing INs\textsuperscript{39} or SST\textsuperscript{+} cells, as suggested by Yu et al.\textsuperscript{20}.

At the same time, we find a striking developmental change in bottom-up thalamic inputs onto SST\textsuperscript{+} INs. Whereas prior to active whisking they receive the majority of inputs from the sensory-related VPM and PO nuclei, after the onset of active whisking their thalamic inputs shift to motor-related nuclei, such as the VM and the VAL. A study by Wall et al., that combined AAV helper virus and monosynaptic rabies virus to trace the inputs onto inhibitory INs in the adult S1, showed that the VPM and PO strongly project to SST\textsuperscript{+} INs, while the motor-related nuclei only showed few presynaptic cells\textsuperscript{40}. However, it is important to note that in our experiments we get very few starter cells in the main thalamo-recipient layer (L4) compared to the abovementioned study. In addition, due to the usage of whole-brain clearing and the autofluorescence of the thalamic barreloids (VPM), we have been able to very precisely assign cells to different thalamic nuclei, which could have been mis-assigned to more rostral VPM using histological approaches.

Along the same lines, another study that reported the in vivo activity of SST\textsuperscript{+} neurons in adult animals showed that they do not receive strong inputs from the VPM\textsuperscript{29,32}. The authors suggested that SST\textsuperscript{+} INs may rather be recruited by local excitatory inputs, which fits well with our cortical connectivity results onto these interneurons. However, we can certainly not exclude that some of the SST\textsuperscript{+} activation comes from different thalamic nuclei still to VM and VAL. These nuclei are in fact known to project to the striatum, L1 of motor-related cortical areas, the cingulate as well as the prefrontal cortex\textsuperscript{41–45}. A proportion of VM cells also have axonal collaterals to S1, the target specificity of which was until this study unknown\textsuperscript{46,47}. AAV tracings from the Allen Mouse Brain Connectivity Atlas (http://connectivity.brain-map.org/projection) show that axonal projections from the VM exclusively target L1 of the somatosensory cortex. This strongly suggests that the connection we uncover with the rhabies tracing is formed between VM and L2/3 SST\textsuperscript{+} cells as they are known to extend their dendrites into L1\textsuperscript{48}. Furthermore, the VM is activated during whisking events, as assessed by in vivo photometry with GCaMP6\textsuperscript{49}, suggesting that it is involved in reporting whisker-related information, which it could potentially pass on to the superficial SST\textsuperscript{+} INs.

Notwithstanding the underlying mechanisms, both VIP\textsuperscript{+} and SST\textsuperscript{+} INs show greater responses upon multi- compared to single-whisker stimulation before P14. Together with our anatomical data, this indicates that the functional impact of bottom-up input onto VIP\textsuperscript{+} cells decreases and becomes more specific after P14. We would speculate that this decrease coincides with the emergence of top-down modulation from wM1 to allow these cells to act as coincident detectors between active whisking and a touch event of an individual whisker. The SST\textsuperscript{+} INs on the other hand would be preferentially activated if multiple whiskers are deflected simultaneously. We would argue that the ability of SST\textsuperscript{+} INs to differentiate between single- and multi-whisker stimuli in the juvenile sW1, suggests that these cells act antagonistically to the activation of VIP\textsuperscript{+} cells, depending on how each whisker is deflected in relation to their neighboring ones. We would speculate that upon complex navigation or object exploration, in some parts of the sW1 SST\textsuperscript{+} activation would take over, while in others VIP\textsuperscript{+} cells would engage more. Therefore, our results support a model that upon the onset of active whisking, VIP\textsuperscript{+} and SST\textsuperscript{+} INs diversify their functions to be able to register multiple aspects of the environment and hence help facilitate appropriate tactile perception.

**Methods**

**Animals.** All animal experiments were approved by the Cantonal Veterinary Office Zurich and the University of Zurich. Animals were housed in a 12-h reverse dark-light cycle (7 a.m. to 7 p.m. dark) at 24 °C and variable humidity. Animal lines used in this study are: VIP-IREs-Cre (Viptm1zjh/J)\textsuperscript{50}, SST-IREs-Cre (Ssttm2.1zjh/J)\textsuperscript{50}, AAV11 Cre (B6.129S7-Gt(Rosa)26Sortm14(CAG-tdTomato)Hit/Hpr)\textsuperscript{50} and HTB Gt (ROSA26Sortm1(CAG-loxP-tdTomato)Hr)\textsuperscript{52}.

**Animal surgery and preparation for in vivo imaging experiments.** We used 6 VIPCre-tdTomato and 6 SSTCre-tdTomato mice (7 males, 5 females) at ages
ranging from P8 to P41 for two-photon Ca\textsuperscript{2+} imaging. As described previously\cite{13}, mice were sedated with chlorprothixene (0.1 g/kg, intraperitoneal (i.p.), Sigma- Aldrich Chemie GmbH, Buchs, Switzerland) and urethane (10 mg/kg, Sigma-Aldrich Chemie GmbH, Buchs, Switzerland) and used to guide the location of the neuron of interest with (without excluding any interesting neighboring neuronal ROI). Neuronal correction was performed as \( F_{\text{correction}} = F_{\text{neuron}} / \alpha \cdot F_{\text{fluorescent}} \). \( \alpha \) is estimated for each imaging spot separately using the formula \( F_{\text{blood vessel/surrounding}} = F_{\text{correction}} \). For each stimulus, the evoked responses of 10 trials were analyzed and the response magnitude expressed as \( \Delta F/F \). The magnitude of the evoked \( \Delta F/F \) responses (starting at stimulus onset). Pearson’s correlation coefficients of sensory-evoked responses for any two neurons at zero lag were calculated for each single trial evoked calcium traces in a 17.4 s window starting from stimulus onset. Spontaneous intrinsic image fluctuations were obtained by averaging correlations of 1000 randomly segmented 17.4 s long pieces from a 300 s’s spontaneous recording, to get rid of the trace length-dependent correlation fluctuations for the comparison of spontaneous and evoked correlations. To detect spontaneous events, we first ran a peak finding algorithm (in MATLAB) and afterwards manually checked each peak to verify if it was an actual event. Our definition of an event was a peak followed by an (bi-) exponential decay with a 1 ± 2 s flat baseline. When the signal-to-noise (SNR) in the data is relatively high, it is easier to detect such events, however, it gets harder as the SNR gets lower. In the juvenile dataset, SNR was significantly higher than the neonatal dataset, making the detection of the events in the latter more difficult. In order to fit the calcium responses properly in all groups we had to use a double exponential fit, after which we calculated the weighted decay time:

\[ t = (a_f \cdot t_f^2 + b_f \cdot t_f + c_f) / (a_f + b_f + c_f) \]

where \( a_f \), \( b_f \), and \( c_f \) are the coefficients for the decay.

Decoders. Three different decoders were trained either for the multi- or single-whisker stimulation paradigm, to quantify the decoding capacities of the STSs and the VIPs at P8-P12 and P21+. The decoders were used random forest, naïve Bayes and Error-correcting output codes (ECOC) that classified a cell type into one out of two groups based on baseline-subtracted average stimulus response trials\cite{32}. Each classifier was trained 1000 times, with randomly setting the random number generator for the training (Matlab function TreeBagger, with parameters ntree = 50, minleaf = 5; fliccet, and ficcoc, respectively). For each trained set, 30% of the remaining data was used to calculate classification accuracy.

Acute slice electrophysiology. Whole-cell patch-clamp electrophysiological recordings were performed on either SfC PREF-tomato or VIPPREF-tomato cells in L2/3 of the wS1, in acute slices prepared from P8–12 or P21+ animals. Animals were anesthetized, decapitated, the brain extracted and transferred to 4°C physiological Ringer’s solution (aCSF), of the following composition (mM): 125 NaCl, 5 KCl, 25 NaHCO\textsubscript{3}, 1.25 Na\textsubscript{2}HPO\textsubscript{4}, 1 MgCl\textsubscript{2}, 2 CaCl\textsubscript{2}, and 20 glucose. The brain was then glued to a stage and cut into 300 μm-thick coronal slices using a vibratome (VT 1200S, Leica). The slices recovered in room temperature aCSF for 30 min before recording. The slices were then placed in the recording chamber of an upright microscope (AxioScope 2 FS, Zeiss) and superfused with 32°C oxygenated (95% O2 and 5% CO2) aCSF at a rate of 2–3 ml/min. The microscope was equipped with immersion differential interference contrast (DIA) and the following objectives were used to visualize the cells (10x/0.3, Olympus and 40x/0.8, Zeiss). A CMOS camera (optiMOS, QImaging) was attached to the scope to visualize the slice and cells through a computer screen. A white-light source (HAL 1000, Dage) and a LED based lighting source (Polychrome source) in combination with a tdTomato filter set (34 HE, Zeiss, Excitation 550/25, Emission 605/70) were used to locate the fluorescent INs. Patch pipettes were pulled from borosilicate glass capillaries (1.5 OD × 0.86 ID × 7.5 mm, Harvard Apparatus) at a resistance of 4–6 MΩ.

For recordings of intrinsic electrophysiological properties and photon-stimulation-evoked currents, Clampex was used (v10.7.0.3, Molecular Devices 2016). The recording pipettes were filled with a solution containing the following (mM): 135 potassium D-glucosate, 4 NaCl, 0.3 Na-GTP, 5 Mg-ATP, 12 phosphocreatine-di-tris, 10 HEPS, 0.0001 CaCl\textsubscript{2} (pH 7.25, mOu 290). In all cases, 1% biocytin (Tocris Bioscience) was added in the recording solution and no extra pharmacology was added. Access resistance was constantly monitored to ensure the quality and stability of the recording. The recorded data were accepted only if the initial series resistance was less than or equal to 25 MΩ and did not change by more than 20% throughout the recording period. Compensation was made for the pipette and cell capacitance.

We used a similar stimulation protocol as previously reported\cite{33,34}. For each analyzed cell, passive and active membrane properties were recorded in current-clamp mode at −65 mV by applying a series of linearly increasing hyperpolarizing and depolarizing sub- and supra-threshold current steps (300 ms, ΔV ± 20 pA). The analysis of intracellular properties was performed in Clampfit (v10.7.0.3, Molecular Devices 2016). Using this protocol, we analyzed the cells for five different parameters: For calculating the membrane resistance, we used the \( R = \Delta V / \Delta I \) on hyperpolarizing pulses that did not lead to an activation of voltage-dependent conductances; Spike threshold was obtained as the inflection point of a rapid change in dV/dt of the first spike after the step protocol. Spike height was calculated as the amplitude of the spike to the maximum value of the peak. Spike width was calculated as the width at half of the maximum amplitude; After hyperpolarization amplitude was measured
from spike threshold to the lowest occurrence point after the repolarization phase of the spike.

All intrinsic electrophysiology data presented in the paper are average.

**In vitro calcium transient and AP correlation.** Whole-cell patch-clamping of SSTCre-ttdTomato<sup>+</sup> neurons of L2/3 in wt was performed at P8–12 and P21+. Patch pipette was filled with 100 μM OGB-1 and right after breaking into patched cell, an incubation period of 25 min took place to let the OGB-1 equilibrate within the cell body. A square pulse current was injected giving rise to a set number of APs (1, 3, 5, 7, and 10), while simultaneously recording the evoked Ca<sup>2+</sup> transient from presynaptic terminals. The data were analyzed using the software Scanipix and a laser system at 900 nm excitation. Two-channel fluorescent images of 256 × 128 pixels at 11.25 Hz (HyperScope galvo-mode) were collected with a 40X water-immersion objective lens. The ΔF/F of the gathered Ca<sup>2+</sup> responses per a given number of APs were analyzed in terms of the Ca<sup>2+</sup> transient max amplitude, decay tau, and integral. The Averages of these parameters were taken and compared between the two age groups, P8–12 and P21+.

**Glutamate uncaging and data analysis.** For photostimulation experiments, whole-cell patch-clamp recordings were performed on randomly selected INs in L2/3 and using a 10× objective lens (Olympus, NA 0.3, 1048 × 1960 µm field of view). For light stimulation, the tissue was digitally divided into 400 subregions making up a grid pattern, spanning all cortical layers. The tissue was submerged in oxygenated aCSF containing caged glutamate (295 µM RuBi-Glutamate, Tocris) and photostimulation was performed at least three times at each spot, in a pseudorandom manner, using a high-power LED light source (470 nm, 3300 mW, Mightex Systems). Photostimulation parameters were calibrated beforehand so that the experiment paradigm successfully evoked a short train of ~4 APs in randomly selected INs. This set of synaptic partners of the recorded cells. The amplitude of the evoked APs in putative pre-synaptic neurons were used as a window of analysis for the responses evoked within the patched INs (Fig. 3a). This time-window was used to capture majority of mono-synaptic activity and to minimize the presence of poly-synaptic. Although, when performing the glutamate uncaging, the activation of presynaptic terminal through photostimulation never resulted in AP elicited currents within the postsynaptic patched cell, suggesting that the contribution of polysynaptic activity would be small. Recordings of synaptic activity were performed in voltage-clamp mode at ~45 mV (to register both excitatory and inhibitory currents), with a sampling rate of 10 kHz.<sup>45</sup> Custom-written Matlab script was used to analyze the photostimulation evoked current data. Within the data gathered at ~45 mV, the excitatory currents were extracted by taking all activity registered below baseline (average of the signal within 500 ms before illumination) within the analysis window, while inhibitory currents were all activity above baseline.

For each recording, the field of view was segmented per layer and the average activity within each layer was normalized to the overall average activity within the whole field of view. Mann–Whitney U test was carried out for all data points within each layer and comparison was made between lamina.

For the lateral activity analysis, the average activity within L2/3 was calculated per given distance from each recorded cells soma. The average was then calculated for all recordings within a given group. Two-tailed t-test was used to look for significance between groups and points with same given distance from soma.

The second set of photostimulation experiments were performed on SSTCre-ttdTomato or VIPCre-ttdTomato cells in L2/3 of the wt, in acute slices prepared from P12–14 wt animals. The set of glutamate uncaging experiments was performed in the same manner as described above but at ~70 mV, to allow to selectively detect only incoming excitatory currents to our recorded cells. When a first round of photostimulation had been carried out, 1 μM TTX (Tetrodotoxin, Tocris) was added to the circulating aCSF and 12 min after, the same glutamate uncaging experiment was performed. Through custom-written Matlab script we then subtracted the values obtained before from after TTX to calculate the extent of the synaptically evoked responses originating from pre-synaptic partners alone, rather than direct stimulation of the recorded cell. When performing this analysis only evoked activity originating from the superficial layers were considered.

**Immunohistochemistry.** For the immunohistochemistry experiments, the Ai14 reporter mouse line was combined with either VIPCre or SSTCre. The offspring were then bred with VGlut2<sup>−/−</sup> (before onset of active whisking) to P12+ (then on the slide overnight (>17 h) at 4 °C while the secondary, donkey a Rabbit488 (1:1000, Thermo Fisher A21206), was left for 2 h at RT. The slides were then coverslipped using VectaShield Gold and stored at 4 °C for imaging.

**Confocal imaging and image processing of immunohistochemical samples.** The slides were imaged using a Confocal Microscope (Olympus FV1000 with Fluoview software). All images were taken from L2/3 of the barrel cortex and of each of the processed brain 8–10 stacks were generated. To image the VGlut2 puncta a 60x oil-immersion objective was used and stacks were taken at 0.47-µm distance between.

A custom MATLAB script was used to analyze the VGlut2-ttdTomato colocalization. The algorithm worked as follow: Image de-noising was performed using Wiener filtering followed by 2-D bilateral filtering. Wiener filter performs 2-D adaptive noise-removal resulting in a low pass filtered version of an intensity image. The stationary additive noise component is then isolated based on statistics at a local neighborhood of each pixel. The bilateral filter is an edge-preserving nonlinear filter that smoothens a signal while preserving strong edges.<sup>65</sup> After de-noising, Z-stacks were binarized using multilevel Otsu thresholding for each xy-, xz-, and yz- slices separately, resulting in 3 binary 3D-stacks combined using logical AND operator (This very stringent binarisation operation makes sure the detected signal is noise free). Finally 3D binarized stacks for channel 1 (VGlut2) and 2 (VIP- or SST-ttdTomato) were combined with another AND operator and 3D connected components analysis was used to count number of co-localizations. Number of co-localizations was normalized by total ttdTomato signal for each image.

**Viruses injections.** For these experiments the HTB reporter line was crossed with either VIPCre or SSTCre. The pups were injected with SLAV-Y envelope glyco-protein (Envα)-pseudotyped, glycoprotein-deleted rabies virus SADG-mCherry (Envα) at either P4/P5 or P15. All pups were injected with 100 ml of the virus, 120–170 µm deep into the barrel field of the primary somatosensory cortex. The injections were done using a glass microprobe attached to a Nanolitre 2010 pressure injection apparatus (World Precision Instruments). After the surgeries, the animals were returned to their home-cage for 7 days to allow for adequate viral expression.

For the control rabies viruses experiments VIPCre pups aged P15 were injected with 100 nl of a 1:1 mix of pAAV-hSyn-FLEX-TVA-P2A-EGFP-2A-oG (gift from John Naughton, Addgene plasmid # 85225) and rabies virus (see above). The injections were done at the same depth and using the same equipment as described above. The animals were sacrificed 7 days after the injection and all the brains were cleared as described below.

**Tissue processing after viral infections.** Seven days after the rabies virus injection the animals were transcardially perfused using 1x PBS and 4% PFA. Following that, the brain were either processed for immunohistochemistry as described above or for tissue clearing. For immunohistochemistry the whole brain was cut at 20 μm using a cryostat (Leica, CM3050 S) and collected on slide. Every third of the collected sections was stained with chicken-aGFP antibodies (abcam, ab13970, 1:1000) to enhance the HTB signal.

The method used for hydrogel-based tissue clearing is explained in detail elsewhere.<sup>66,70</sup> The animals were transcardially perfused using 4% PFA and the brains post-fixed for 48 h in a Hydrogel solution (1% PFA, 4% Acrylamide, 0.05% Biot<sup>68,69</sup>) before the Hydrogel polymerization was induced at 37 °C. Following the polymerization, the brains were in 40 ml of 5% Biot for 7 days. While taking off room temperature until the tissue was cleared sufficiently (10–40 days depending on the age of the animals). Finally, after 2–4 washes in PBS, the brains were put into a self-made refractive index matching solution (RIMS)<sup>71</sup> for the last clearing step. They were left to equilibrate in 5 ml of RIMS for at least 4 days at RT before being imaged.

**Imaging of rabies injected brains.** A Slidescanner (Zeiss, AxioScan Z1) was used to image the stained sections. Mosaic images of the injected hemisphere were taken using a 20x objective. The obtained pictures were processed for analysis using the ZEN Software and Fiji.

After clearing, brains were attached to a small weight and loaded into a quartz cuvette, then submerged in RIMS and imaged using a home-built mesoscope beam (105 µm diameter, 1.5 m-think brain sections that contained the barrel sections that contained the barrel cortex were cut and collected on-slide (wS1) using a cryostat (Leica, CM3050 S). The slides were then stored at ~80 °C until further processing. For labeling tachykinin input onto the ttdTomato expressing cells an antibody staining was performed. For blocking 0.5% Nonidet P-40 (Sigma), 0.2% TTX (Tetrodotoxin) and 1.5% NaCl into the recording experiment was performed on the slide and left for 1 h at RT. Both primary and secondary antibodies were diluted in the same blocking solution. The primary, rabbit aVGLUT2 (1:500, Syssy 135 402), was left on the slide overnight (>17 h) at 4 °C while the secondary, donkey a Titan888 (1:1000, Thermo Fisher A21206), was left for 2 h at RT. The slides were then coverslipped using VectaShield Gold and stored at 4 °C for imaging.
the field-of-view of 5–10 µm (depending on zoom and wavelength). Field of views ranged from 10.79 mm at ×1.25 magnification (Pixel size: 5.27 µm) for overview datasets to 3.27 mm at ×4 (Pixel size: 1.6 µm). Further technical details of the mesoSPIM are described elsewhere (www.mesoSPIM.org). The images generated with the mesoSPIM were further processed using the Fiji and the Imaris software.

**Analysis of cleared tissue imaging data.** Close-up stacks taken from the barrel cortex of rabies injected and cleared SSTCre-HTB and VIPCre-HTB brains were initially processed through Fiji and z-stacked samples of super cortex of rabies injected and cleared SSTCre-HTB and VIPCre-HTB brains were further processed using the Fiji and the Imaris software.

**Code availability**

**Data availability** Data are available upon request. Source data are provided with this paper.

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**Author contributions**

R.K. performed the anatomical mapping, tissue clearing, immunohistochemistry and wrote the manuscript; R.V. performed functional electrophysiology experiments, in vitro experiments; A.I. developed the deep neural network-based neuron detector pipeline and wrote the manuscript; R.K. performed the anatomical mapping, tissue clearing, immunohistochemistry and wrote the manuscript; T.K. conceptualized the study, designed the experiments and wrote the paper; A.v.d.B. performed in vivo Ca2+ imaging and wrote the paper; A.d.B. performed in vivo Ca2+ imaging experiments and initial virus infections; A.O.A. performed the analysis of the in vivo Ca2+ imaging data, puncta co-localization in histology, analysis of the in vitro electrophysiology data and decoder analysis; A.I. developed the deep neural network-based neuron detector pipeline (DeNeRiD), did the analysis of the rabies-traced cells and the their distribution in the cortex; F.F.V. built, provided access to and guidance for the light-sheet microscope, F.H. provided access to the Light-Sheet Microscope and a two-photon microscope. D.K. and A.A. provided initial guidance on tissue clearing, T.K. conceptualized the study, designed the experiments and wrote the paper.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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