Neural correlates of single-vessel haemodynamic responses in vivo

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Neural activation increases blood flow locally. This vascular signal is used by functional imaging techniques to infer the location and strength of neural activity. However, the precise spatial scale over which neural and vascular signals are correlated is unknown. Furthermore, the relative role of synaptic and spiking activity in driving haemodynamic signals is controversial. Previous studies recorded local field potentials as a measure of synaptic activity together with spiking activity and low-resolution haemodynamic imaging. Here we used two-photon microscopy to measure sensory-evoked responses of individual blood vessels (dilation, blood velocity) while imaging synaptic and spiking activity in the surrounding tissue using fluorescent glutamate and calcium sensors. In cat primary visual cortex, where neurons are clustered by their preference for stimulus orientation, we discovered new maps for excitatory synaptic activity, which were organized similarly to those for spiking activity but were less selective for stimulus orientation and direction. We generated tuning curves for individual vessel responses for the first time and found that parenchymal vessels in cortical layer 2/3 were orientation selective. Neighbouring penetrating arterioles had different orientation preferences. Pial surface arteries in cats, as well as surface arterioles and penetrating arterioles in rat visual cortex (where orientation maps do not exist), responded to visual stimuli but had no orientation selectivity. We integrated synaptic or spiking responses around individual parenchymal vessels in cats and established that the vascular and neural responses had the same orientation preference. However, synaptic and spiking responses were more selective than vascular responses—vessels frequently responded robustly to stimuli that evoked little to no neural activity in the surrounding tissue. Thus, local neural and haemodynamic signals were partly decoupled. Together, these results indicate that intrinsic cortical properties, such as propagation of vascular dilation between neighbouring columns, need to be accounted for when decoding haemodynamic signals.

To determine how neural activity leads to changes in cerebral blood flow, the haemodynamic responses of individual vessels need to be compared to neural activity in the surrounding tissue. While sensory-evoked responses of individual vessels have been measured in the somatosensory cortex and olfactory bulb of rodents, these studies have not measured vessel responses to the full range of stimuli for which the neighbouring neural tissue is responsive. Thus, the degree to which vascular signals match local neural activity has been difficult to assess. Here we compare neural and vascular responses to a full range of stimulus orientations in cat primary visual cortex to determine if vascular responses can be predicted from local neural activity. Additionally, the primary visual cortex of the cat, similar to that of primates including humans, is organized into precise maps such that different stimulus orientations in neural tissue are optimally activated by different stimulus orientations. Therefore the orientation selectivity of vessel responses can be linked to the spatial scale of neurovascular coupling. For example, if blood flow in a single cortical vessel is sensitive to neural activity over a large spatial scale covering many orientation columns, then the vessel should dilate to a broad range of stimulus orientations. By contrast, if the vascular response is controlled very locally, that is, within the scale of an orientation column, then individual vessels may be highly orientation selective.

We first labelled blood vessels in the cat primary visual cortex with the fluorescent indicators Texas Red Dextran or Alexa 633 (see Methods), and measured the dilation responses to drifting grating stimuli of different orientations. Veins and capillaries, which were distinguished from arteries by a number of means (see Methods), were not included in this initial analysis because they rarely exhibit rapid sensory-evoked dilation 12–14. Our data set included all other blood vessels, providing that they were sufficiently labelled and imaged in tissue with minimal movements from respiration. All blood vessels in this data set dilated in response to drifting grating visual stimuli (P < 0.05 analysis of variance (ANOVA)). Specifically, we found that parenchymal arterioles in layer 2/3 typically dilated more strongly in response to one or two of the stimulus orientations presented (Fig. 1b), whereas pial surface arteries dilated to all orientations nearly equally (Fig. 1c). For each vessel, we computed the orientation selectivity index (OSI; see Methods), such that when a vessel dilates equally to all stimulus orientations the OSI = 0 and when a vessel responds only to a single orientation the OSI = 1. The OSI was much greater for parenchymal arterioles than for pial surface arteries (OSI parenchymal arteriole mean ± standard error of the mean (s.e.m.) = 0.21 ± 0.01; n = 79 vessels and OSI surface artery mean ± s.e.m. = 0.06 ± 0.01; n = 24 vessels; P < 10−10, Mann–Whitney test; Fig. 1d).

To illustrate further the role of an organized map of neocortical neurons in generating tuned parenchymal vessel responses, we also measured dilation changes in rat primary visual cortex. Because cortical neurons in rats are not organized in an orientation map15, each parenchymal vessel is surrounded by neurons displaying a variety of orientation preferences (Fig. 1e). In rats, we found no orientation selectivity in cortical layer 2/3 parenchymal arterioles (Fig. 1f; OSI mean ± s.e.m. = 0.06 ± 0.01; n = 16 vessels) or pial surface arteries (Fig. 1g; OSI mean ± s.e.m. = 0.05 ± 0.01; n = 21 vessels) (Fig. 1h).

To compare the orientation selectivity of cat parenchymal vessels to spiking activity in the surrounding tissue, we performed calcium imaging using Oregon Green BAPTA-1 AM (OGB-1 AM) or GCaMP6s, along with vascular imaging from the same sites (see Methods). Figure 2a shows a penetrating arteriole that dilates most strongly in response to a noise stimulus as compared to the preferred stimulus orientation (0.30) (Fig. 1a). Therefore this vessel exhibited high orientation selectivity (OSI = 0.82). The OSI was calculated using the following formula: OSI = 2 × (P - OS0.5)/(1 + OS0.5), where P is the percentage of neural spiking activity. To illustrate further the role of an organized map of neocortical neurons in generating tuned parenchymal vessel responses, we also measured dilation changes in rat primary visual cortex. Because cortical neurons in rats are not organized in an orientation map15, each parenchymal vessel is surrounded by neurons displaying a variety of orientation preferences (Fig. 1e). In rats, we found no orientation selectivity in cortical layer 2/3 parenchymal arterioles (Fig. 1f; OSI mean ± s.e.m. = 0.06 ± 0.01; n = 16 vessels) or pial surface arteries (Fig. 1g; OSI mean ± s.e.m. = 0.05 ± 0.01; n = 21 vessels) (Fig. 1h).

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Figure 1 | Selectivity of blood vessel dilation to sensory stimuli in species with and without cortical orientation maps. a, Schematic of cat visual cortex showing the columnar organization of neurons by orientation preference and a pial surface artery with multiple branches penetrating the parenchyma. Different colours of neuronal cell bodies represent their different preferred stimulus orientations. b, Time courses and polar plots (averages of six (top) and eight (bottom) trials) of the changes in dilation of two layer 2/3 arterioles in cat visual cortex to visual stimulation. Error bars represent s.e.m. and grey bars represent the periods of visual stimulation. In this and subsequent figures, stimuli were gratings that drifted in eight different directions of motion and polar plots are normalized to the maximum response. c, Time course and polar plot of responses from a surface artery in cat (average of four trials). d, Population distribution and median OSI for parenchymal (n = 79 vessels in 18 cats) and surface (n = 24 vessels in 9 cats) vessels. e, Schematic of rat visual cortex where neurons with different orientation preferences are intermingled. f, g, Time courses and polar plots of responses from two parenchymal arterioles (averages of seven (top) and eight (bottom) trials) and a surface artery (average of five trials) in rat visual cortex. h, Population distribution and median OSI for parenchymal (n = 16 vessels in 6 rats) and surface (n = 21 vessels in 7 rats) vessels.

A larger region might explain the broad orientation selectivity of the parenchymal vessels. Previous work has shown that occlusion of a single penetrating arteriole in the neocortex leads to tissue death in regions with approximately 400-μm diameter15, suggesting that this is the region of tissue that an individual penetrating arteriole supplies (see also Supplementary Information). Therefore we compared vascular responses to calcium signals integrated over 400-μm-diameter windows around each parenchymal artery (Fig. 2b, c). The orientation preference of these regions of spiking activity matched those of the arteries at their centres (Fig. 2d; R = 0.94, n = 19 pairs, P < 10^-6). However, the orientation selectivity of the spiking activity was higher than the corresponding artery in all regions examined (Fig. 2e; P < 10^-5, paired t-test). Because the spacing of penetrating arterioles is heterogeneous (see Supplementary Information), we also examined the selectivity of neural responses in a wide range of window sizes around each vessel (100–600 μm diameter). We found that for all window sizes the spiking activity OSI was still at least 60% higher than the vessel dilation OSI (Fig. 2f; P < 10^-5 at each window size).

Our calcium imaging results suggest that additional sources of neural activity (besides spiking in the local tissue around the vessel) may be contributing to sensory-evoked vasodilation. Experimental and theoretical work has implicated synaptic glutamate release as a driver of haemodynamic responses16. In particular, calculation of the energy budget of the neocortex estimated that, of all the cellular processes performed, excitatory synaptic activity has the largest metabolic demand17. Therefore, to measure directly excitatory synaptic activity over different spatial scales and compare it to single-vessel responses, we labelled neurons in the cat visual cortex with a glutamate sensor (iGluSnFR; see Methods). We found that glutamate activity (like neuronal spiking) is organized in direction and orientation maps (Fig. 3 and Extended Data Fig. 1). However, glutamate signals were generally less orientation selective than spiking activity. Integrating over 400-μm-diameter windows, the OSI for calcium (mean ± s.e.m. = 0.59 ± 0.02; n = 19 regions) was sharper than that for glutamate (mean ± s.e.m. = 0.44 ± 0.02; n = 37 regions; P < 0.001, Mann–Whitney test). To determine if excitatory synaptic activity alone could account for single-vessel haemodynamic responses, we integrated the glutamate signals over 400-μm-diameter regions around individual arteries (Fig. 3a, b) and compared these to vessel responses from the same sites (Fig. 3b–e). We found that the visual stimulus that produced the largest glutamate signal in a 400-μm window matched the visual stimulus that resulted in the largest vessel dilation (Fig. 3c; R = 0.90, n = 37 pairs, P < 10^-13). However, the 400-μm region of synaptic activity was always more selective than its corresponding penetrating arteriole (Fig. 3d; P < 10^-11, paired t-test). The mismatch between orientation selectivity in individual blood vessels and synaptic activity was confirmed for a range of glutamate response window sizes (100–600 μm diameter; Fig. 3e; P < 10^-5 at each window size).

The broader tuning of the vascular response relative to synaptic and spiking activity (Figs 2 and 3) suggests that vessels can respond to sensory stimuli that evoke little to no concomitant neural activity in the surrounding tissue. This phenomenon can be directly observed by comparing the response amplitudes of vessel dilation and neural activity to individual stimulus conditions. Extended Data Figure 2a, b shows an example in which two visual stimuli (135° and 180°) evoked robust dilations in a penetrating arteriole but essentially no glutamate release in the region around the blood vessel. Across the data sets of synaptic and spiking activity, we compared the amplitude of each vessel's response to each sensory stimulus against the neural response around the vessel to the same stimulus (Extended Data Fig. 2c). Our analysis confirmed that there are many instances where there is a non-existent (or very small) synaptic or spiking response to a visual stimulus despite a robust dilation response. In general, there are very few instances where a stimulus failed to evoke a dilation response of some magnitude.

Like orientation tuning, direction selectivity is a hallmark feature of the primary visual cortex and represents the capacity of a neuron to respond preferentially to one direction of stimulus motion at the optimal stimulus orientation. We found direction maps for excitatory synaptic activity (Fig. 3) that were qualitatively similar to direction maps of spiking activity (Fig. 2). However, the directionality index18 (DI) over 400-μm-diameter windows was greater for spiking activity than for synaptic responses (DI spiking mean ± s.e.m. = 0.59 ± 0.07; n = 19 regions; DI synaptic mean ± s.e.m. = 0.33 ± 0.03; n = 37 regions; P < 0.01; Extended Data Fig. 3a). Blood vessel responses appeared to have little direction selectivity, even when surrounded by iso-direction territories of spiking activity, for example, Fig. 2b vessels 2 and 4. Indeed, across the population, the direction selectivity of vessels was smaller than that of regions of spiking activity (DI vessel mean ± s.e.m. = 0.30 ± 0.02; n = 79 vessels; P < 0.0005). The population mean DI of vessels and synaptic activity were similar (Extended Data Fig. 3a; P = 0.70) although
Figure 2 | Stimulus selectivity of single vessels and of spiking activity in the surrounding tissue. a, In vivo anatomical image of a small region of layer 2/3 cat visual cortex labelled with OGB-1 AM (green) and an arteriole labelled with Alexa 633 (red). Polar plots show the sensory-evoked calcium responses from six neurons and the dilation of the arteriole. b, Pixel-based direction map and polar plots of responses from another cat labelled with OGB-1 AM and Alexa 633. The pixels are colour-coded by their preferred stimulus, with the brightness indicating the response strength. Red polar plots show the dilation responses of the five vessels indicated by solid white circles on the direction map. Green polar plots show the calcium responses pooled in 400-μm-diameter windows around four of these vessels (dashed circles). No calcium responses are shown for the region around vessel number 1 because this vessel was near the edge of the imaging field. c, Direction map and polar plots of responses from cat visual cortex labelled with GCaMPs and blood vessels labelled with Texas Red Dextran. Left, in vivo anatomical images of a large region of layer 2/3 where the positions of seven penetrating arterioles are numbered. Right, pixel-based direction maps of neural responses. The polar plots at the bottom show the vessel dilation responses (red) and the calcium responses (pooled over 400-μm-diameter regions) around five of these vessels (green). d, Correlation between the preferred orientation of the vessel dilation and the preferred orientation of calcium responses in 400-μm-diameter windows around each vessel (R = 0.94; P < 10^{-10}; n = 19 windows in 8 cats; regression line shown in red). e, No significant correlation between calcium and vessel OSI (R = 0.41; P = 0.08). f, Distribution of OSI for calcium responses across tissue regions of different window sizes (n = 11 cats) and for the population of dilation responses (n = 18 cats). Solid bars are medians and boxes indicate the interquartile range. For all window sizes, calcium responses were more selective than the vessel dilation (P < 0.0001, Mann–Whitney test). Pixel maps shown are averages of five to six trials. Scale bars, 25 μm (a) and 100 μm (b, c).

There was no correlation between the direction selectivity of a particular vessel and the glutamate signals in the surrounding tissue (Extended Data Fig. 3b; R = 0.20; n = 37 pairs; P = 0.23).

While vessel dilation responses over the population of parenchymal arterioles did not match neural orientation selectivity, we tested the possibility that the smallest vessels would show similar selectivity to neural responses. Larger penetrating arterioles (with baseline diameter >15 μm) may perfuse larger regions of tissue than small penetrating arterioles and their finer branches. Therefore, these smaller vessels (typically 8–15-μm baseline diameter) may be sensitive to vasodilators from smaller regions of neural tissue and thus have sharper orientation tuning. Indeed we found that OSI was inversely correlated with baseline vessel diameter in cat layer 2/3 (Extended Data Fig. 4a; R = 0.37, P < 0.001). We compared the dilation responses of these small vessels (baseline diameter ≤15 μm, mean = 11.8 μm) to those with baseline diameter >15 μm (based on consistency of Alexa 633 labelling12; see Methods). We found that the small vessels were slightly more tuned (OSI mean ± s.e.m. = 0.24 ± 0.02; n = 35) than the larger ones (OSI mean ± s.e.m. = 0.18 ± 0.01; n = 44; P < 0.05; Extended Data Fig. 4a, b). Importantly, however, the OSI of these small vessels was still lower than synaptic and spiking activity over the full range of window sizes (P < 0.005; see Extended Data Fig. 4b, c).

Capillaries are the smallest vessels in the neocortex and therefore may be even more tuned for stimulus orientation than small arterioles. However, whether capillaries have the capacity to dilate in vivo to sensory stimuli is controversial13,14,18,19. This is probably due to inconsistent criteria for defining capillaries and distinguishing them from pre-capillary arterioles as well as to the difficulty of detecting dilation in very small vessels even with two-photon microscopy resolution13,14,18,19. However, a small dilation in a capillary that is undetectable with two-photon microscopy would still lead to easily detectable changes in red blood cell (RBC) velocity. Because RBC size is slightly larger than the capillary lumen diameter, a very small dilation in a capillary could produce a dramatic reduction in the resistance to flow (see figure 2e in ref. 20). Therefore, we measured the stimulus-evoked changes in RBC velocity in a set of micro-vessels that would probably be classified as capillaries based on their high tortuosity and small diameter (4–7 μm; see Methods)21,22. We found that the orientation selectivity on the basis of blood velocity in these capillaries (OSI mean ± s.e.m. = 0.30 ± 0.04; n = 15 vessels) was no different from what was found for dilation of the ≤15-μm-diameter vessels (P = 0.16; Extended Data Fig. 4a, b). To determine if the tuning of capillaries was due to these vessels being in unusually broadly tuned windows of neural activity, we compared the OSI of neural activity around capillaries with what was found around parenchymal arterioles. The OSIs of 400-μm-diameter windows of spiking activity around capillaries (mean ± s.e.m. = 0.60 ± 0.03; n = 13) and around arterioles (mean ± s.e.m. = 0.59 ± 0.02; n = 19) were indistinguishable (P = 0.94, Mann–Whitney test). Thus, stimulus-evoked changes even in capillaries were still not as selective as the responses in adjacent neural tissue.

Our results suggest that blood flow increases in parenchymal vessels are partially driven by local neural activity (which would generate the match in orientation preference) and also by an additional global component arising from adjacent functional columns (which would induce the dilation to non-preferred orientations). One possibility is that this global component is due to the propagation of dilatory signals along vessel walls. Specifically, the lack of orientation tuning in surface arteries could result from the dilation of penetrating arterioles from many different orientation domains propagating back to a surface artery. Then the propagation of dilation along the surface artery and down into multiple penetrating arterioles could broaden the locally driven dilation signal, leading to dilation in adjacent regions of tissue that have no concomitant neural activity. Previous studies in rodents have demonstrated the propagation of dilation from the parenchyma up to the cortical surface23 and along the surface over distances of at least 1 mm (refs 24–26) at rapid speeds27, but these have not been linked to the stimulus selectivity of vessel or neural responses. Consistent with this propagation of dilation hypothesis, we found that in cat visual cortex parenchymal vessels dilate before the surface vessels and that the dilation to the preferred orientation came before the dilation to the stimulus that was oriented orthogonal to the preferred orientation, that is, at the null orientation (see Supplementary Information and Extended Data Fig. 5). Alternative
Figure 3 | Stimulus selectivity of single vessels and of excitatory synaptic activity in the surrounding tissue. a, Bright-field image of the surface of cat visual cortex showing the location of six penetrating arterioles and the regions targeted for two-photon imaging. b, Direction maps and polar plots of glutamate responses from cortical neurons labelled with iGluSnFR and dilation from blood vessels labelled with Texas Red Dextran. The positions of the arterioles and the 400-μm-diameter windows of pooled glutamate responses in cortical layer 2/3 are indicated by solid white and dashed circles, respectively. Red polar plots show the dilation responses and blue plots show the pooled glutamate activity in the windows around each arteriole. c, Correlation between the preferred orientation of the dilation responses and the preferred orientation of glutamate activity in 400-μm-diameter windows around each vessel for all cat data (R = 0.90, P < 10^{-11}; n = 37 windows in 5 cats). d, No significant correlation between glutamate and vessel OSI (R = 0.28, P = 0.68; linear regression line shown in red). e, Distribution of OSI for glutamate activity across windows of different sizes (n = 5 cats) and for the population of dilation responses (n = 18 cats). Solid bars are medians and boxes indicate the interquartile range. For all window sizes, the glutamate responses were more selective than vessel dilation (P < 10^{-5}; Mann–Whitney test). Pixel maps shown are averages of 8–10 trials. Scale bars, 500 μm (a) and 100 μm (b).

hypotheses on the origin of the selectivity of vessel dilation are discussed in Supplementary Information.

In conclusion, our results have a number of implications for the interpretation of haemodynamic signals in relation to neural activity. We provide direct single-vessel evidence for the untuned global signal in the pial vasculature that has been found in low-resolution haemodynamic imaging studies. After subtracting the global signal, these earlier studies often suggested that the residual tuned vascular responses were of capillary origin. We show that individual penetrating arteries also display stimulus-specific responses. Furthermore, the orientation selectivity of these parenchymal vessels is an order of magnitude higher than what is obtained with intrinsic signal optical imaging. We also demonstrate that an organized functional map of neural responses is required for attaining tuned haemodynamic signals (see also ref. 30). Furthermore, by sampling responses over the full range of a stimulus parameter and by directly measuring synaptic and spiking activity along with single-vessel responses in precisely defined spatial regions of tissue, we overcome many of the technical limitations of earlier studies that examined neurovascular coupling. The difficulties inherent in correlating low-resolution vascular signals with electrophysiological metrics of neural activity and in interpreting glutamate pharmacology has led to controversy regarding the spatial scale over which synaptic versus spiking activity matches vascular signals (see Supplementary Information). Here we establish that the sensory stimulus that elicits the largest synaptic or spiking response also produces the largest haemodynamic signal. However, the complete selectivity profile of neither synaptic nor spiking activity in the local tissue around a vessel can be inferred from the tuning curves of haemodynamic signals. Thus, vascular signals are partially decoupled with local neural signals, over distances of at least 300 μm.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Author Contributions P.K. conceived and supervised the project. All authors collected data. P.O’H. and P.Y.C. analysed data. P.O’H., M.L. and P.K. wrote the paper. All authors commented on and approved the final manuscript.

Author Information Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to P.K. (kara@musc.edu).
METHODS

Animals and surgery. All surgical and experimental procedures were approved by the Institutional Animal Care and Use Committee at Medical University of South Carolina. Cats (∼25 of either sex; postnatal day 28 to >2.5 kg adult) were anaesthetized with isoflurane (1–2% during surgery, 0.5–1.0% during imaging) and paralyzed with a continuous infusion of vecuronium bromide (0.2 mg kg⁻¹ h⁻¹, intravenously). Cats were artificially ventilated through a tracheal cannula, and the end tidal CO₂ was regulated at 3.5–4.5%. Heart rate, respiration rate, temperature and electrocardiogram were monitored. Evoked responses were offset at ∼10 min after postnatal days 31–45 and C57BL/6J mice (∼1 male, postnatal day 63) were initially anaesthetized with a bolus injection of fentanyl citrate (0.04–0.06 mg kg⁻¹), midazolam (3.75–6.25 mg kg⁻¹), and dexmedetomidine (0.19–0.31 mg kg⁻¹). The one mouse was used for a control experiment to confirm that the iGluSnFR sensor was not being saturated during sensory stimulation (see later). During two-photon imaging, continuous intrapetaleral infusion with a lower concentration mixture (fentanyl citrate: 0.02–0.03 mg kg⁻¹; midazolam: 1.50–2.50 mg kg⁻¹; and dexmedetomidine: 0.10–0.25 mg kg⁻¹) was administered using a catheter connected to a syringe pump.

For all animals, craniotomies (2–3 mm square) were opened over the primary visual cortex (area 18), the dura was reflected, and the craniotomies were sealed with agarose (1.5–3% dissolved in artificial cerebral spinal fluid (ACSF)) and a glass coverslip. When the calcium indicator OGB-1 AM was used, before the placement of the coversglass, a pipette was inserted into the craniotomy and the dye was injected with air pressure puffs. The dye loading procedure has been described in detail⁴⁻⁸.

In cats, we also used the genetically encoded indicators GCaMP6s⁹–¹¹ and iGluSnFR¹² to measure calcium and glutamate activity respectively. Two- to four-weeks before the imaging session, viral injections of AAV2/9.Hsyn.GCaMP6s, WPRE.SV40-eGFP.AAV2/1.hSyn.iGluSnFR.SV40 were performed under sterile surgery conditions. Cats were anaesthetized with 1–2% isoflurane and visible signs were monitored. One to three craniotomies were performed over the primary visual cortex (area 18) and small holes were made in the dura. Aliquots of virus (5 μl) were diluted in PBS and mannitol (5:9:6 ratio of virus:PBS:mannitol) to titres of ~10¹⁲ genomes ml⁻¹ with 50–200 nl of Fast Green dye (Sigma) added to visualize the injection. Glass pipettes containing the virus solution were lowered 500–800 μm into the cortex and pressure puffs were administered over 15–20 min until approximately 1 μl had been injected. After 10 min, the pipettes were slowly retracted, the craniotomies were sealed with agarose (3% dissolved in ACSF), the scalp was sutured closed and the animals were recovered and returned to their housing. All animals were treated similarly and so randomization and blinding were not required. No statistical methods were used to predetermine sample size.

Two-photon imaging. For vascular imaging, three fluorescent dyes were used as described previously¹⁻³, Alexa 633 fluor hydrazide selectively labels artery walls, while Texas Red dextran (70 kDa) and fluorescein dextran (2,000 kDa) were not required. No statistical methods were used to predetermine sample size. Visual stimulation and size of imaged region.

Fluorescence was monitored with a custom-built microscope (Prairie Technologies) coupled with a Mai Tai (Newport Spectra-Physics) mode-locked Ti:sapphire laser (810 nm or 920 nm) with DeepSee dispersion compensation. Excitation light was focused by ×40 (NA 0.8, Olympus), ×20 (NA 1.0, Olympus) or ×16 (NA 0.8, Nikon) water immersion objective and beam expansion optics. Full-field two-photon excitation imaging and the stimulus presentation were controlled using custom software written in LabView (National Instruments) and driven by a tethered Windows computer. Stimulus presentation was accomplished through a pair of 10-bit drivers (National Instruments). The average power delivered to the sample plane was 100–200 mW. The system was calibrated to control the average delivery of 10–20 mW to the sample plane. The temporal stability and wavefront error of the laser were monitored using a large area photodiode (New Focus 4050-12) and an interferometric wavefront sensor (New Focus 8004). Images were usually oversampled by interpolating between pixels from 2 to 20 times to allow the algorithm to compute diameter values with a spatial resolution that was finer than the pixel size in the raw data images. To compute the vascular response to each condition, a stimulus response window was defined. Because of the slow onset and offset of the vascular response, we could not simply assign the response period to correspond to the period when the stimulus was displayed on the monitor. Instead, for each vessel we selected the response period by examining the average response across all stimulus conditions and then selected the imaging frames that best approximated this response interval. Shifting this time window by addition of 50 ms did not significantly change the responses. The mean response across this time window was divided by the baseline level for each condition to get the percentage change in diameter. Responsive vessels were defined by ANOVA across baseline and eight directions over multiple trials (P < 0.05). The orientation selectivity index (OSI) was defined as:

\[
\text{OSI} = \frac{\sum \left[ \frac{r_k - \bar{r}}{\sigma} \right]^2}{N} \sum_{k=1}^{N} r_k
\]

where \( \bar{r} \) is the orientation of each stimulus and \( r_k \) is the mean response across trials to that stimulus⁴⁻⁸. Note that OSI = 1.0 for circular variance. The preferred orientation was defined as: \( \text{arctan} \left( \frac{\Sigma r_k \cos(2\theta)}{\Sigma r_k \sin(2\theta)} \right) \). The directionality index (DI) was computed as 1.0 – rnull/rpeak, where rnull is the response amplitude to the preferred stimulus and rpeak is the response to the stimulus with the same orientation.
drifting in the opposite direction. Computing the OSI based on flow rather than diameter by scaling the diameter values to the 4th power (Poiseuille's law) did not affect our results.

Measuring onset latency to dilation. To compare the latencies of pial arteries and parenchymal arterioles, we fit a linear regression line to the rising phase of the dilation (20–80% of the peak response) of each vessel. For parenchymal vessels, we used only the response to the preferred orientation because of potential latency differences between dilation to the preferred and other stimulus orientations (see later). For the pial vessels, we pooled the response to all stimulus conditions because these vessels are untuned to stimulus orientation. We used the time at which the regression line crossed the pre-stimulus baseline level as the onset latency. This regression line metric on the average response is applicable when responses are large and relatively stable from trial to trial—as is the case for pial vessels to any stimulus orientation and for parenchymal vessels to the preferred stimulus orientation.

Since parenchymal vessels are orientation selective (Fig. 1b, d), responses to the null orientation are the weakest and, by definition, smallest in amplitude and more noisy from trial to trial. Thus, to compare the latency between the response to the preferred and null orientations in parenchymal vessels, we used a statistical test, the standardized mean difference (SMD, specifically Hedge's g; refs 36, 37), in which vessels are weighted by the trial-by-trial variance in latency values (see Extended Data Fig. 5a). We first smoothed each trial's time course with a three-frame running average. We then performed linear regression on the same interval as earlier (20–80% of the peak). We took the difference in the average onset latency across trials between the responses to preferred and null stimuli and standardized this difference by the pooled variability across the two conditions. The population summary SMD was obtained by using a random-effect model. This model weighs each vessel by the inverse variance of its SMD and factors in the heterogeneity present across the individual vessel data. As a control for spurious effects, the preferred and null responses were assigned randomly for each trial and the analysis was repeated (Extended Data Fig. 5b).

Quantifying blood velocity. Velocity data was analysed as described previously. Briefly, line scans were first pooled into blocks of 250, 500, or 1,000 lines. The angle of the RBC streaks in each image was used to determine the velocity of that block and a time course of velocity measurements was extracted. Baseline and stimulus windows were defined similarly to the dilation data and equivalent OSI and statistical analyses were performed.

Analysis of calcium and glutamate responses. Calcium and glutamate signals were analysed the same way. Raw images were first smoothed with a 4 μm Gaussian filter. The mean fluorescence of each pixel within a given 100–600-μm-diameter window around a vessel was computed for each blank and stimulus epoch. A t-test was performed on the difference in stimulus and baseline fluorescence for each condition in each trial and if the distribution was significantly higher than zero (P < 0.05), the pixel was included in the integration window. We also performed this analysis without excluding the unresponsive pixels and the responses were indistinguishable. In some data sets, part of the 100–600-μm-diameter analysis window fell outside of the image boundary and so there would be fewer pixels from those domains contributing to the overall response. Therefore, to avoid biasing the overall response of the integrated region, we divided the 100–600-μm-diameter analysis window into wedges before averaging the data over the full window. Each wedge was 1/16 of the circle and was further divided into sections of 50 μm radial length. Thus, a 100-μm-diameter window had 16 sections whereas a 400-μm-diameter window had 64. The pixels with significant responses within each section were averaged together to create a time course. The time course was then normalized by a sliding baseline of the mean fluorescence of each blank interval (ΔF/F). Each time course was then weighted by the total number of pixels represented by its section, because sections farther from the vessel contain more pixels. Finally, the time courses of all the sections were averaged together to obtain the time course of the entire region. For inclusion in the population data set, responses from the 100–600-μm-diameter analysis windows had to pass the following criteria. First, each wedge had to have at least 30% of the imaged pixels passing the initial t-test to ensure that windows with wedges having no response and/or weak labelling would be removed. Second, at least 80% of the circular area of the window had to be within the image to ensure that a sizeable region of tissue whose orientation preference could dramatically affect the overall response was not being missed. In addition, at least 10% of each wedge had to be within the image to ensure that each wedge had some representation. For data that passed all these criteria, the responses to each condition were computed by averaging the imaging frames during stimulus presentation and across trials. Before the OSI was computed, if any conditions showed a negative response (below the baseline level), then the absolute value of the minimum response was added to all responses (to make the minimum equal zero). We have recently published a mechanistic rationale for applying such a correction in fluorescence imaging of neural responses—stimulus-evoked dilation of surface arteries can block fluorescence from the underlying tissue and make a very small response actually appear negative. We also analysed the data without this correction and, in addition, when only including the first 1 s of the response (to avoid the slower surface artery interference). Although there were small changes in the OSI values of individual windows, the overall results did not change in either case. The neural response amplitudes, OSI and DI were all computed using the same formulae as for the vessel data. Population distribution statistics on OSI and DI measurements used the Mann-Whitney test.

Additional control for calcium imaging. Spiking activity in the neuropil should also contribute to metabolic needs and hence neurovascular coupling. Therefore, when integrating the calcium signals in the tissue surrounding each artery, we included all pixels that passed a signal-to-noise criterion (see earlier) and not only those corresponding to cell bodies. However, the neuropil may include a mixture of calcium signals from synaptic events in dendritic spines and spiking in axons arriving from regions outside of the integration window we selected. Therefore, as a control, we compared the orientation selectivity in 400-μm-diameter windows with and without including the neuropil. Masks excluding the neuropil were generated in the same way as described earlier except that the pixels within each wedge were constrained to the cell bodies. Cell body masks were first created using an automated algorithm that applied a series of morphological filters to identify the contours of cell bodies based on intensity, size and shape. Cell outlines were visually inspected and errors were corrected manually. Then a t-test was performed on each pixel of these masks and the wedges were created in the same manner as before. Because of the sparse distribution of cell bodies, we did not enforce the 30% significantly responding imaged pixels criterion but all other criteria applied. The orientation selectivity with the two mask types was indistinguishable (Extended Data Fig. 9).

Control to show that visual stimulation was not saturating the iGluSnFR sensor. With visual stimulation, the glutamate signals peaked at <10% ΔF/F. To determine if the iGluSnFR sensor responded linearly and responded over a greater range than that obtained with visual stimulation, we used iontophoresis to apply large doses of exogenous glutamate. We lowered a pipette containing 0.5 M glutamate into layer 2/3 of the visual cortex of a mouse that was labelled with iGluSnFR. We applied a range of currents (10, 20, 40, 60, 80 nA) and found that the fluorescence signals increased linearly (R > 0.99; P < 0.0001) and peaked at ~60% ΔF/F (data not shown). Thus, our in vivo imaging with iGluSnFR for (example, Extended Data Fig. 10) is probably revealing the true spatial profile of glutamate direction maps (Fig. 3) and orientation maps (Extended Data Fig. 1).

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Extended Data Figure 1 | Glutamate release is organized into orientation maps. a, Region of cat visual cortex labelled with iGluSnFR. Pixels are colour-coded by preferred orientation with the brightness indicating the response strength. Time courses and polar plots (averages of four trials) are shown for three regions of tissue with different orientation preferences. b, Orientation maps of iGluSnFR responses from a different cat. Time courses and polar plots are averages of ten trials.
Extended Data Figure 2 | Arteriole dilation in the absence of glutamate signalling or local spiking. a, Time courses and polar plots of arteriole dilation (red) and the release of glutamate in a 400-μm-diameter window surrounding an arteriole (blue). Averages of eight trials are shown for vessel dilation and ten trials for glutamate responses. In time courses, error bands represent s.e.m. and grey bars represent the periods of visual stimulation. The responses to the 135° and 180° stimuli (outlined by the black box over the time courses) are large for the vessel dilation but virtually non-existent for the glutamate activity. b, Quantifying the relative amplitude of the vessel and neural responses to each of the eight stimulus directions for the single cat experiment shown in a. Each data point in the scatterplot represents the average response of the vessel and of the neural tissue surrounding it to a single direction of visual stimulation, normalized by the response to the best direction. c, Quantifying the relative amplitude of vessel and neural responses across all cat experiments. Top panel shows glutamate versus dilation data (n = 37 windows and vessels in 5 cats) and the bottom panel shows calcium versus dilation responses (n = 19 windows and vessels in 8 cats). Each data point in the scatterplot is as described in b. The histograms at the top and right show the distributions of neural and dilation responses, respectively. In both population scatterplots, there are many data points in the top left quadrant, indicating stimuli that drove robust dilation responses but minimal glutamate or calcium responses. All data are from cat visual cortex layer 2/3.
Extended Data Figure 3 | Direction selectivity of parenchymal vessels and of local spiking and synaptic activity. a, Population distributions of the direction index of calcium (green, $n = 19$ windows in 8 cats), glutamate (blue, $n = 37$ windows in 5 cats) and vessel dilation (red, $n = 79$ vessels in 18 cats) responses. All data were obtained from cat visual cortex and neural responses were pooled over 400-μm-diameter windows. The DI of spiking activity was greater than the DI of synaptic responses ($P < 0.01$, Mann–Whitney test) and the DI of vessel dilation ($P < 0.0005$, Mann–Whitney test). The DI of synaptic activity was not different from the DI of vessel dilation ($P = 0.70$, Mann–Whitney test). Solid bars are medians and boxes show the interquartile range. b, For each vessel that had a corresponding 400-μm-diameter window of calcium or glutamate activity, the vessel direction index is plotted against the corresponding neural direction index. There was no significant correlation for calcium ($R = 0.2$, $P = 0.43$, $n = 19$ pairs) or glutamate ($R = 0.2$, $P = 0.23$, $n = 37$ pairs).
Extended Data Figure 4 | Dilation and velocity responses in parenchymal blood vessels with different baseline diameters. a, The diameter of all vessels and their OSI values from cat visual cortex layer 2/3. For arterioles, OSI was determined based on dilation (n = 79 vessels in 18 cats) whereas for capillaries, OSI was calculated from blood velocity measurements (n = 15 vessels in 7 cats). b, The distribution of OSI for the three subgroups of layer 2/3 vessels analysed in our study (>15 μm, n = 44 vessels in 15 cats; ≤15 μm, n = 35 vessels in 14 cats; capillaries, n = 15 vessels in 7 cats). The OSI of the ≤15 μm vessels was greater than the OSI of the >15 μm vessels (P < 0.05, Mann–Whitney test). The OSI of the ≤15 μm vessels was not different from the OSI of the capillaries (P = 0.16, Mann–Whitney test). Solid bars are medians and boxes indicate the interquartile range. c, The OSI distribution of dilating vessels and 400 μm-diameter windows of calcium and glutamate responses.
Extended Data Figure 5 | Onset latency of dilation in parenchymal vessels. a, Vessel-by-vessel comparison of the onset latency difference between the response to preferred and orthogonal (null) stimulus orientations. Each whisker diagram represents a single vessel with the circle position indicating the standardized mean difference (SMD; calculated as Hedge’s g) in latency. The whisker length represents the 95% confidence interval (CI) of the SMD. The size of the circle represents the weight given to the vessel when calculating the population summary SMD. 

The population summary SMD is shown by the solid square with the error bands giving the 95% CI. The population average shows that parenchymal vessels responded significantly faster for the preferred than the null stimulus orientation. b, As a control, the analysis shown in a was repeated after randomizing the assignment of preferred and null on individual trials for each vessel. All data are from cat visual cortex layer 2/3 (n = 79 vessels in 18 cats).
Extended Data Figure 6 | Dilation measurements with circle fitting.

**a**, The steps of the circle fitting algorithm are illustrated for a blank and a stimulus frame corresponding to the penetrating arteriole shown in the bottom panel of Fig. 1b. The raw image data (first panel) is oversampled by linear interpolation between pixels (second panel). Then a luminance threshold (a fraction of the gradient between the brightest and darkest pixel of the image) is applied (third panel). Finally, a two-dimensional sobel filter is applied to the thresholded pixels to detect the edge of the vessel (fourth panel). The circle fit is only applied to the pixels in the fourth panel but it is overlaid on all the panels for illustration purposes.

**b**, As the threshold is increased, fewer pixels pass the threshold and therefore the baseline diameter changes. However, the percentage change in diameter across baseline and stimulus presentations (the response amplitude) and the response selectivity remain the same. Note that for vessel geometries needing an elliptical fit rather than a circular fit (see Extended Data Fig. 8c and Methods), the shorter axis of the fitted ellipse was used to estimate the vessel diameter.
Extended Data Figure 7 | Dilation measurements using the cross-section algorithm do not depend on the precise location and angle of the selected cross-section. a, Example cat pial artery (from Fig. 1c) labelled with Texas Red Dextran. b, Another pial artery from a different cat labelled with the artery-specific dye Alexa 633. Both arteries show similar tuning for cross-sections drawn >100 μm apart and also drawn perpendicular and obliquely relative to the vessel walls.
Extended Data Figure 8 | Dilation measurements in small arterioles and comparison of dilation measurement techniques. a, A penetrating artery (#1, the responses of which are shown in the top panel of Fig. 1b) and its daughter branch (#2) in cat layer 2/3 labelled with Texas Red Dextran. Red lines indicate the position of the laser scan path across the vessels for line-scan diameter measurements. b, Individual line-scans are stacked next to each other to create X-time (XT) images. The four large rectangular panels are XT images of a blank and stimulus frame for each of two vessels shown in a. The small panels to the right are the average across the image (~0.96 s) for each of the four frames. The computed diameter values are also shown. These images were oversampled by interpolating between pixels (by 5 times for vessel 1 and by 20 times for vessel 2) before the diameter was calculated. c, The time courses and polar plots of the responses for three different diameter measurements are shown for vessel 1—as a line-scan, a cross-section from a full-frame imaging run (seven trials), and the circle fit from the full-frame imaging run. In this particular example we used an ellipse rather than a circle because of the elongation of the vessel due to its diving obliquely to the imaging plane. d, Time courses of the vessel responses to preferred stimulus orientations for the three groups of vessels shown in Extended Data Fig. 4b. The responses for each vessel were aligned by stimulus onset and binned in 400-ms bins. The population average was then smoothed with a three-frame running average. Mean responses in dark colours and light bands indicate s.e.m. Note that the similar error bands and temporal profiles indicate that the smallest vessels had a similar quality of responses to the larger ones.
Extended Data Figure 9 | Comparison of orientation selectivity in regions of calcium responses with and without neuropil. a, In vivo anatomical image of cells labelled with OGB-1 AM in cat visual cortex and selection of two different masks for quantitative analysis of orientation selectivity. Left, a 400 μm-diameter mask comprising soma pixels only. Right, a 400 μm-diameter mask comprising all significantly responding pixels (see Methods). b, The time courses of calcium responses computed from the two masks. Time courses are averages of five trials, error bands represent s.e.m. and grey bars represent the periods of visual stimulation. c, For a population of 16 imaged regions (from 7 cats), the OSI was computed with the two masks and found to be indistinguishable (cell bodies only OSI mean ± s.e.m. = 0.46 ± 0.04; cell bodies and neuropil OSI mean ± s.e.m. = 0.47 ± 0.04; \( P = 0.12 \), paired \( t \)-test).
Extended Data Figure 10 | Orientation-selective responses in layer 1 neurons and synapses. a, Region of cat visual cortex labelled with OGB-1 AM (to measure spiking activity) and SR101 (to distinguish astrocytes). Note the much sparser density of neuronal cell bodies in layer 1 (left) compared with the higher density of cells deeper in layer 2/3 (right). The polar plots are the responses of the two layer 1 neurons labelled in the image. 

b, Region of cat visual cortex labelled with iGluSnFR (to measure synaptic activity). Again the density of cell bodies (the small black holes) in layer 1 (left) is much lower than in layer 2/3 (right). The polar plots are the responses of a 400-μm- and 100-μm-diameter window of layer 1 glutamate activity.