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# Revisiting horizontal connectivity rules in V1: from like-to-like towards like-to-all

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## Abstract

Horizontal connections in the primary visual cortex of carnivores, ungulates and primates organize on a near-regular lattice. Given the similar length scale for the regularity found in cortical orientation maps, the currently accepted theoretical standpoint is that these maps are underpinned by a like-to-like connectivity rule: horizontal axons connect preferentially to neurons with similar preferred orientation. However, there is reason to doubt the rule's explanatory power, since a growing number of quantitative studies show that the like-to-like connectivity preference and bias mostly observed at short-range scale, are highly variable on a neuron-to-neuron level and depend on the origin of the presynaptic neuron. Despite the wide availability of published data, the accepted model of visual processing has never been revised. Here, we review three lines of independent evidence supporting a much-needed revision of the like-to-like connectivity rule, ranging from anatomy to population functional measures, computational models and to theoretical approaches. We advocate an alternative, distance-dependent connectivity rule that is consistent with new structural and functional evidence: from like-to-like bias at short horizontal distance to like-to-all at long horizontal distance. This generic rule accounts for the observed high heterogeneity in interactions between the orientation and retinotopic domains, that we argue is necessary to process non-trivial stimuli in a task-dependent manner.

**Keywords** Primary visual cortex · Horizontal intra-cortical axons · Orientation and retinotopic maps · Structural advanced anatomy · Functional optical imaging · Computational and theoretical neuroscience · Connectivity rules

## Introduction

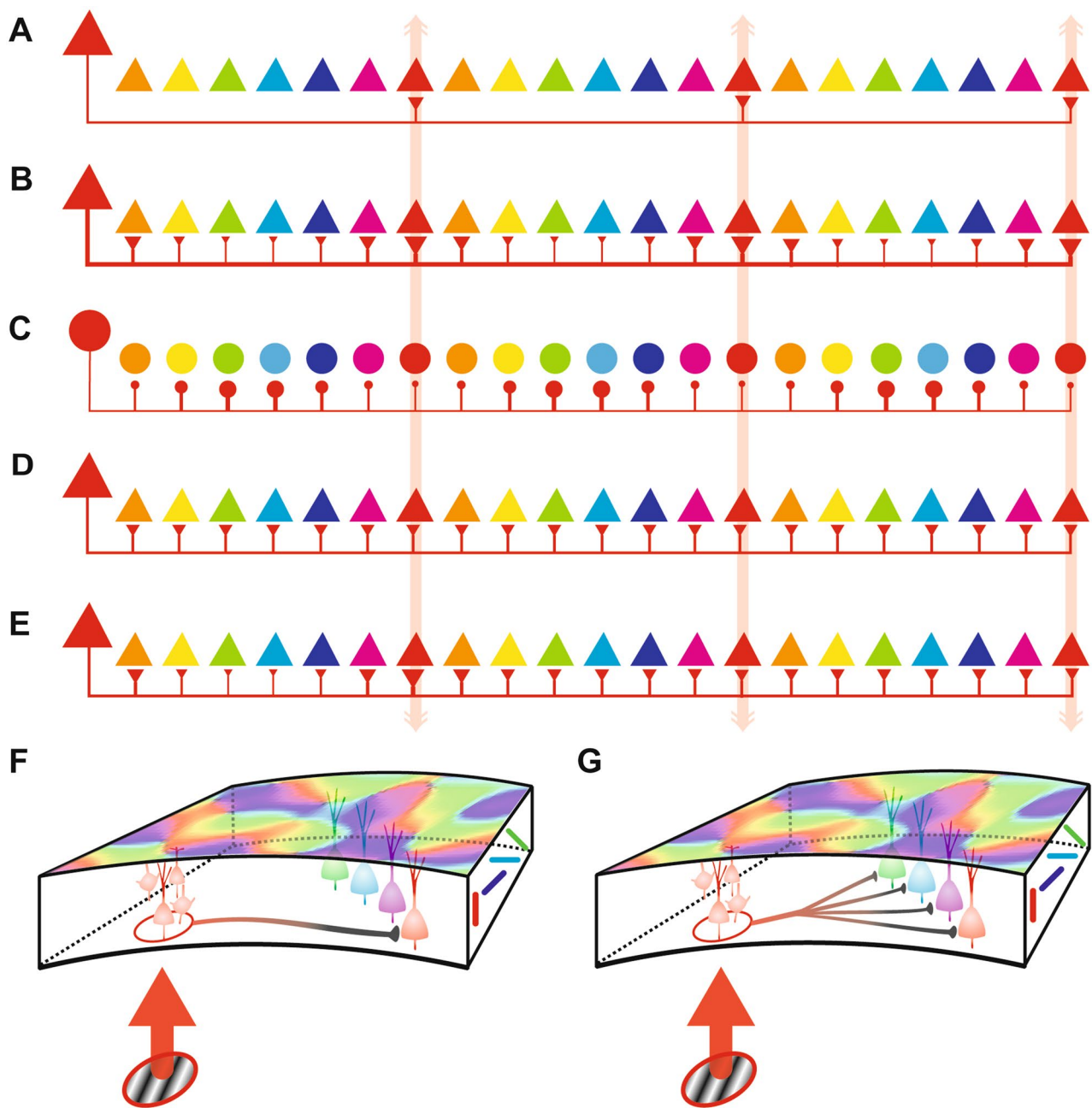
Retinotopy and orientation are two of the main features processed and topographically organized into maps in primary visual cortex (V1) of carnivores, ungulates and primates. Anatomical connections between neurons separated on the cortical sheet, through the so-called *intrinsic*, *intra-cortical* or *horizontal* axons, have a crucial theoretical importance for understanding the computational operations that V1 can perform. Indeed, these axons connect different points in the retinotopic and orientation maps and thereby generate a set of

possible topological interactions within a multidimensional representation of space, orientation and time. It is therefore critical to characterize structural horizontal interactions in order to understand their functional relevance. A vast majority of presynaptic contacts in cortex originate from neurons located in the same area as the postsynaptic target (> 80% in macaque V1, Markov et al. 2011), thus forming an important intra-cortical network. In the primary visual cortex, the feedforward thalamocortical inputs drive the cortical network, which in turn strongly shapes the evoked response through excitatory and inhibitory recurrent circuits within the column (Douglas et al. 1991), a canonical circuit that constitutes nearly 2/3 of intra-cortical connectivity (Markov et al. 2011). The rest of the intra-cortical network connects neurons in adjacent columns separated laterally over distances up to several millimeters, the horizontal network. Early anatomical observations reported that the horizontal connectivity of carnivores, ungulates and primates is spatially distributed into regular clusters (Fig. 1A; Braitenberg 1962; Fiskens et al. 1975; Creutzfeldt et al. 1977; Gilbert and

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**Fig. 1** Illustration of different connectivity rules from literature and possible outcomes for functional activation. In A–E the local neuron (large on left) connects to neighbours in a radially approximated schema spanning outwards over three hypercolumns (where the same preference is encountered, as indicated by the vertical red arrows). Colours indicate the orientation preference of neurons. **A** Strict like-to-like connectivity (extends to long distances). **B** Modulated like-to-like bias (extends to long distances). **C** Like-to-unlike bias as exhibited by inhibitory interneuron. **D** Like-to-all as exhibited by neurons in layers 4 and 6. **E** Like-to-like bias that reduces with distance

resulting in like-to-all at distances beyond adjacent hypercolumns. **F**, **G** Two extreme hypotheses for the net outcome of functionally driven connectivity rule at long-range distance. In response to a local oriented stimulus, all neurons that have a receptive field in overlap with the stimulus will be activated, for excitatory and inhibitory neurons, different lamina and positions in the orientation map. Such functional activation can lead either to a strict iso-orientation activation of neighbouring neurons through the horizontal network (like-to-like rule, **F**) or omni-orientation activation (like-to-all rule, **G**)

Wiesel 1979; Rockland et al. 1982) forming a radially projecting pattern that resembles a daisy's petals (Douglas and Martin 2004). Since orientation maps are also regular with

comparable spatial frequency, the currently accepted theoretical standpoint is that these maps are underpinned by a like-to-like connectivity rule: cortical columns are connected by

horizontal connections only if they share similar orientation preference (Fig. 1A), a hypothesis originally put forward by Mitchison and Crick (1982). Correlative studies, comparing bouton labelling with autoradiography, or with optical imaging maps, qualitatively supported the like-to-like rule (Gilbert and Wiesel 1989). Later combined quantitative anatomy with optical imaging studies confirmed the existence of an orientation preference bias (Fig. 1B, in the range of 1.5–2 times greater than chance, Bosking et al. 1997, Kisvárdy 1997; Schmidt et al. 1997; Malach et al. 1993; Rochefort et al. 2009), with high cell-to-cell variability. Probably due to its simplicity and its elegant topological implications, the highlighted iso-orientation biases have led to a general acceptance of the hypothesis of a simplified and unique like-to-like connectivity. One consequence is that theoretical and computational models have implemented it as a strict rule, not as a bias (e.g. Bressloff et al. 2001; Raizada and Grossberg 2003; Rangan et al. 2005; Sarti et al. 2008; Baker and Cowan 2009; Kaschube et al. 2010; Rubin et al. 2015; Carroll and Bressloff 2016). However, we believe such an oversimplified schema may impair the development of our theoretical understanding of the primary visual cortex function.

Actually, there are reasons to doubt the explanatory power of a global and strict like-to-like connectivity rule. First, a growing number of quantitative studies show that there is a wide variety of connectivity biases (like-to-like bias, no bias, like-to-unlike bias) depending on cell type (Fig. 1C, excitatory vs inhibitory neurons, see Kisvárdy et al. 1994; Buzás et al. 2001), layer origin (Fig. 1D, no bias in layer 4 or layer 6, see Yousef et al. 1999; Karube and Kisvárdy 2010; Karube et al. 2017) and position in the orientation map (Yousef et al. 2001, iso-orientation domain vs pinwheels). Second, the effect is mostly observed at short range where most of the connectivity arises (< 1–1.5 mm), but connections can connect neurons over distances of a few millimeters. The rare analyses over larger cortical distances (more difficult because far fewer boutons are present) showed a global tendency for the iso-orientation bias to reduce with distance (Fig. 1E) due to wider selectivity or deviation from the iso-orientation bias, as observed in Buzás et al. (2006, Fig. 8C), Kisvárdy et al. (1997, Fig. 9—area 17) and Bosking et al. (1997, Fig. 5); however, see a counter example for area 18 in Kisvárdy et al. (1997, Fig. 10—area 18). As a consequence, the effective functional selectivity of horizontal axons beyond the short-range distance is not very clear. Lastly, the functional impact of the structural organization, as described by anatomy, is far from trivial to predict. Indeed, any visual stimulation will activate a neuronal mass encompassing all layers, both excitatory and inhibitory neurons and at least a full hypercolumn composed of pinwheels and iso-orientation domains (see Fig. 1D in Chavane et al. 2011). Furthermore, not only neurons with preferred orientation matching the orientation of the stimulus will be

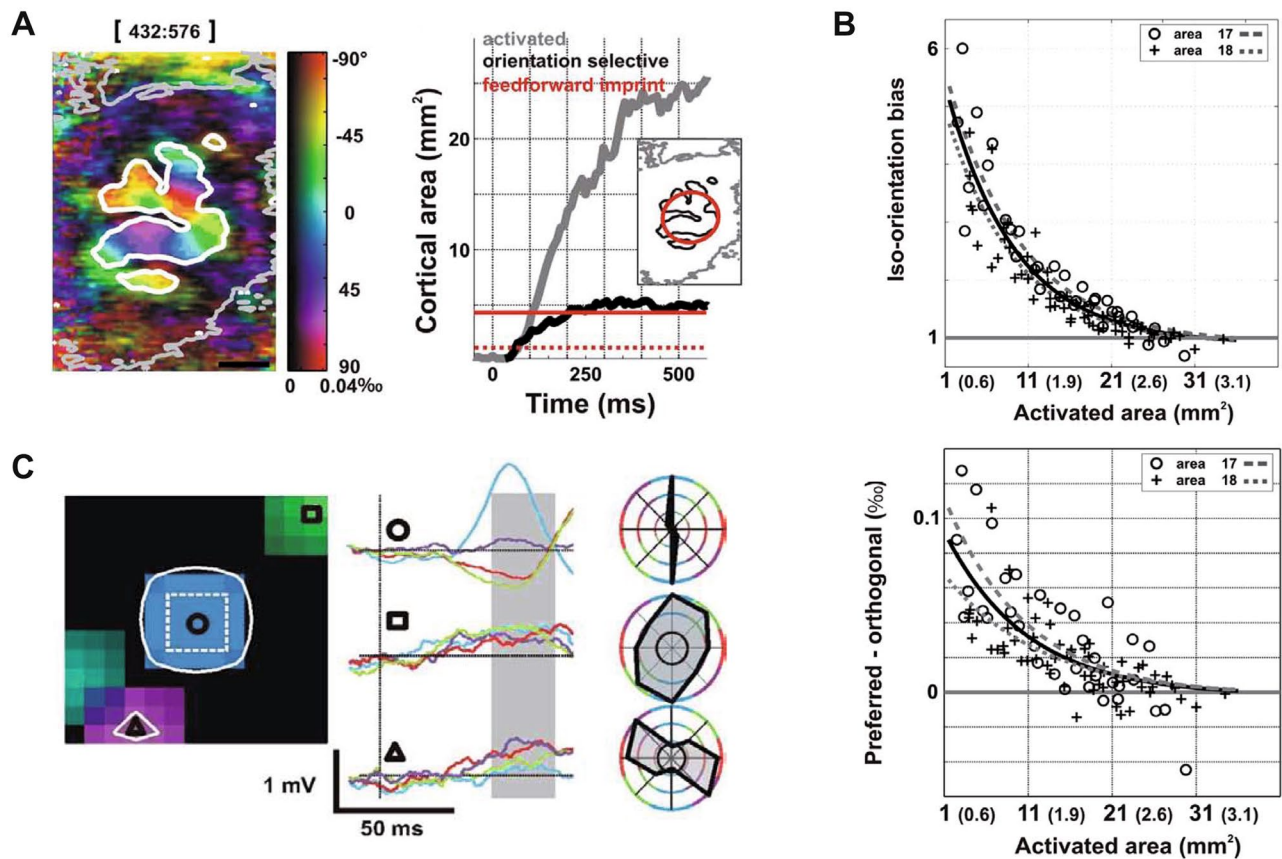
significantly activated, but also a distribution of neurons with say,  $\pm 15^\circ$  around the stimulus orientation. The intra-cortical horizontal network triggered by this functionally activated neuronal mass will forcibly contact a diversity of orientation tuned neurons (ranging from an iso-orientation, Fig. 1F, to an omni-orientation interaction, Fig. 1G) with an overall net effect beyond short-range distance that is particularly difficult to predict without a computational approach.

In this review, we present a body of recent evidence from anatomy, physiology and computational modelling, leading to the conclusion that horizontal interactions do not forcibly conform with a like-to-like orientation preference. In the last decade, structural (Hunt et al. 2011; Martin et al. 2014, see Kisvárdy 2016 for review) and functional (Chavane et al. 2011; Huang et al. 2014) studies have shown that the rule is not valid for long-distance connections. Chavane et al. (2011) proposed revisiting the connectivity rule as a function of horizontal distance: from like-to-like at short distance towards like-to-all and long distances (Fig. 1E; see discussion in Alonso and Kremkow 2014a, b). In their computational modelling study, Rankin and Chavane (2017) showed that this behaviour is in fact to be expected based on the anatomical observations made by Buzás et al. (2006). The functional implications of such evidence is further discussed in the framework of natural scenes analysis (Perrinet and Bednar 2015; Boutin et al. 2021). In light of converging evidence from a range of approaches, this review argues for a timely, in-depth revision of V1 horizontal connectivity rules. Revisiting this textbook mindset is an important prerequisite to better understand the relationship between structure and function in the visual cortex.

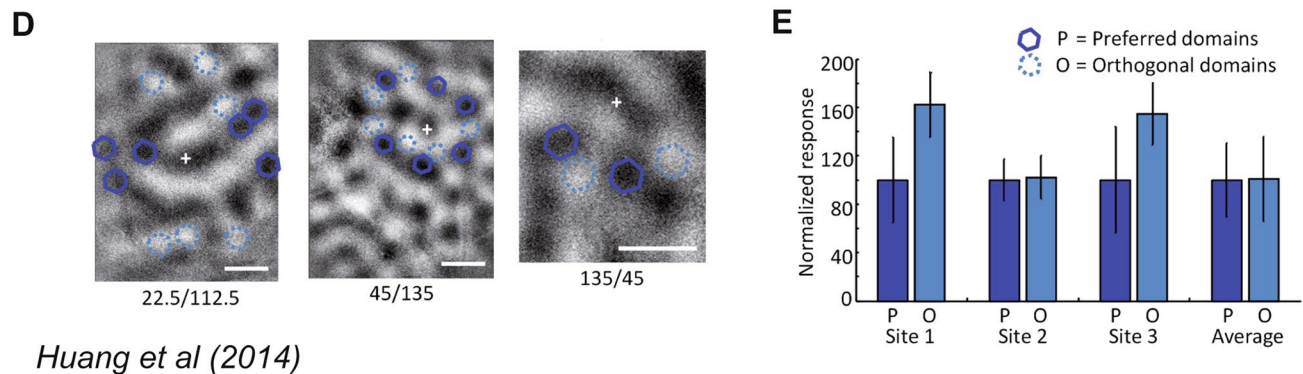
## New physiological evidence

### Neuronal population activity measures

Here, we review more recent evidence for versatile connectivity rules reported in different species and with different recording techniques. Importantly, one should keep in mind that long-range horizontal axons only have a subthreshold influence on their postsynaptic targets (Bringuier et al. 1999). In order to study the selectivity of the postsynaptic target of these axons, it is therefore important to use methods that are sensitive to subthreshold membrane potential changes. Indeed, methods that only record spiking activity necessitate experimental protocols that co-activate the presynaptic source and postsynaptic target of the horizontal network, for instance to study cross-correlation between neurons (Michalski et al. 1983; Ts'o et al. 1986; Schwarz and Bolz 1991; Das and Gilbert 1999). Under these conditions, it is hard to tease apart the direct effects of the horizontal



Chavane et al (2011)



Huang et al (2014)

axons from secondary activation of recurrent columnar circuits.

Chavane et al. (2011) used complementary recording tools that specifically record the subthreshold activity of a mesoscopic population (voltage-sensitive dye imaging, VSDI), and of individual neurons (intracellular recordings) in areas 17 and 18 of the anaesthetized cat. The first method allowed us to visualize and quantify the orientation selectivity of the laterally spreading activity evoked by a local stimulus (Jancke et al. 2004). The second method enabled a precise measurement of the impact of this subthreshold

spread of activity on individual neurons. Using VSDI in the cat areas 17 and 18, the authors showed that a local oriented stimulus evokes a spread of activity along the horizontal dimension, extending up to 3 mm laterally (see also Bringuier et al. 1999; Reynaud et al. 2012; Muller et al. 2014, 2018). It is to be noted that the spread of activity did not show any patchiness, contrary to the anatomical observations. We believe that this is to be expected considering the large variability in the patches that will be activated from different neurons, varying as a function of a neuron's type, layer and position in the orientation map. As a consequence,



**Fig. 2** Probing for the orientation selectivity of the horizontal network with functional imaging. **A–C:** Adapted from Chavane et al. (2011) with CC-BY permission and **D–E:** modified from Huang et al. (2014) with CC-BY permission. **A** Voltage-sensitive dye imaging of the orientation-selective response evoked by local oriented gratings, example from area 17 of an anaesthetized cat. (Left) Polar orientation map averaged over the final 145 ms of the response (time stamps indicated above the frame). Colour hue and brightness code, respectively, for the preferred orientation and the strength of the orientation tuning. Contours delineate the outer border of the cortical domain within which significant activation level (thin grey contour) or significant orientation selective response (thick white contour) are observed. (Right) Spatial extent of the activated area (grey) and of its orientation selective component (black) as a function of time. A red line indicates the expected limit of the feedforward imprint, defined and estimated from Albus (2004) as the population of neurons directly or partially activated by the feedforward stream. Dotted red line indicates the retinotopic area of the stimulus representation. Inset: the spatial extent of the activation spread (grey) and the orientation-selective activation (black) are shown in comparison with the expected limit of the feedforward imprint (red). **B** Population analysis over nine hemispheres [three in area 17 (o) and six in area 18 (+)] of the horizontal distance-dependent decrease of orientation selectivity. (Top) Iso-orientation bias as a function of the spatial eccentricity of the lateral spread. The first point corresponds to the area of the initial cortical activation. Exponential fit is shown in black. (Bottom) Decrease in condition-wise modulation depth with lateral propagation distance. **C** Visuotopic orientation polar map of an intracellular subthreshold response; colour hue and brightness code, respectively, for the preferred orientation and the strength of the membrane potential's orientation tuning. The white contours delineate the significant responsive regions when combining both amplitude and orientation selectivity criteria. Middle: averaged subthreshold responses to four different orientations (same colour code) presented for particular locations (circle, triangle and square); scale bars: 50 ms and 1 mV; right: Normalized orientation tuning curves, integrated within a fixed temporal window (shaded area of middle panel). The black circle indicates the spontaneous level for the depolarizing integral measure. **D** Three orientation maps measured with optical imaging of intrinsic signals (Huang et al. 2014) with the extracellular recording site (white "+"). Optogenetic layout stimulation sites that were centred over orientation domains with the same orientation preference (blue solid hexagons) as the recording site (+), and stimulation sites that were centred over domains with the orthogonal preference (light blue dashed hexagons). Scale bars: 500  $\mu$ m. **E** Extracellular responses to optogenetic stimulation of preferred domains (dark blue) and orthogonal domains (light blue) in the three example cases shown in **D**, and the average responses to stimulation of preferred or orthogonal domains across all cases examined ( $n = 10$ )

and in sharp contrast to the extended horizontal activation, the orientation selective component of this spread remains confined to the cortical feedforward imprint of the stimulus (Fig. 2A). The feedforward imprint being defined in Chavane et al. (2011) as the population of neurons directly or partially activated by the feedforward stream. This effect was systematically observed in both areas 17 and 18, and quantified using complementary methods to quantify decrease of the orientation selectivity with horizontal distance. Both at the level of orientation preference and orientation selective response, the bias towards like-to-like activation (and therefore functional connectivity) decreases exponentially

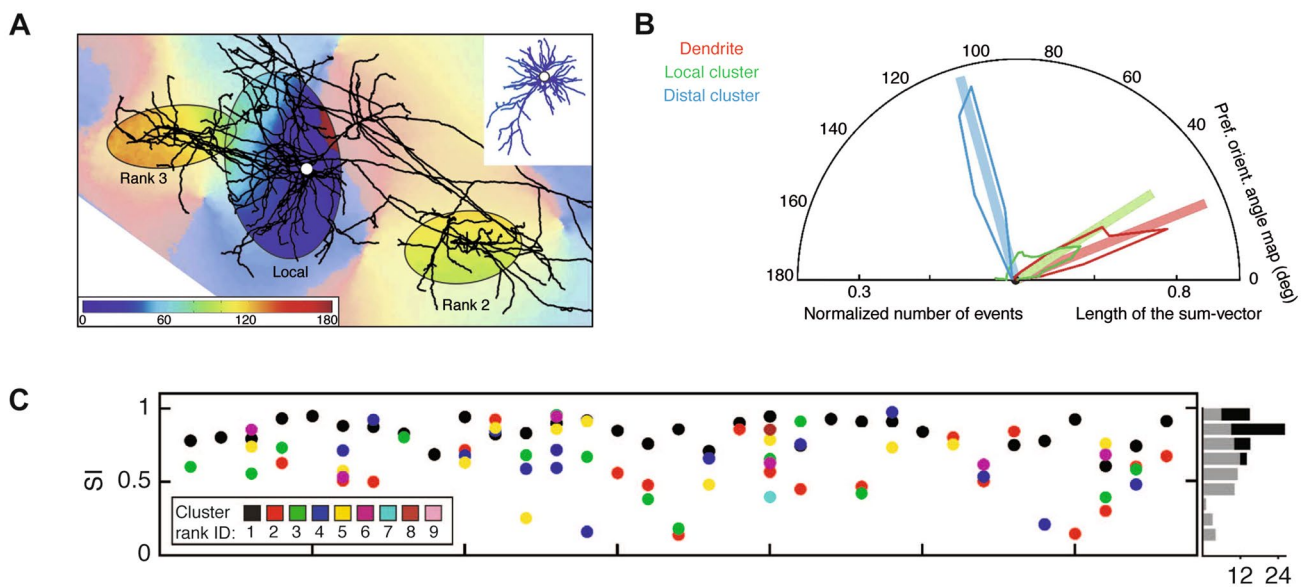
with horizontal distance with a similar characteristic cortical space constant of about one mm or one hypercolumn (Fig. 2B). Importantly, this signifies, that, for a lateral radius of about 1.5 mm, the iso-orientation bias (Fig. 2B) was in the same range as that observed in the anatomy for similar lateral distance (Bosking et al. 1997; Kisvárdy 1997; Schmidt et al. 1997; Malach et al. 1993; Rochefort et al. 2009). However, VSDI is a population measure of the subthreshold activation that pools activity from all neurons (excitatory and inhibitory), all compartments (dendrite, soma and axons) and mostly the upper layer (see Chemla et al. 2017). VSDI offers a unique population view of the functional activation but it is less precise than anatomical studies: it is for instance possible that the lack of overall bias comes from the mix of tuned and untuned subpopulations (see Kisvárdy 2016 for further discussion). Chavane et al. (2011) therefore used intracellular recordings to confirm the VSDI observations and further showed that this loss of orientation selectivity actually arises from the diversity of converging synaptic inputs originating from outside the classical RF (Fig. 2C). The conclusion from this work is that the lateral spread of cortical activity gradually loses its orientation iso-preference at a distance of around one hypercolumn and that there exists a range of strategies for different postsynaptic neurons.

A more recent work, Huang et al. (2014) provided complementary results in a different species, V1 of the tree shrew and using a different methodological approach. The authors used optical imaging of intrinsic signals to monitor the impact of intra-cortical optogenetic stimulation under various stimulation configurations. In particular, their results show that the optogenetic stimulation of excitatory neurons within a set of orientation domains in the cortex generated the same response amplitude for either iso- or orthogonal domain stimulation (Fig. 2D, E). The responses actually depended primarily on intra-cortical distance (similar to the results obtained via cross-correlation in Das and Gilbert (1999)). Using their innovative approach, the authors also tested stimulation along an axis in the retinotopic map, either collinear with the preferred orientation or orthogonal to it. The authors found no bias in either direction. Huang et al. (2014) therefore provides independent and complementary evidence that the horizontal network, when probed at the population level with functional measures, does not show a bias for iso-orientation preference in V1. It should be noted, however, that using optogenetic stimulation of excitatory neurons may drive complex dynamical activation of the cortex (Li et al. 2019), mixing excitatory and inhibitory recruitment of the lateral network with different dynamics. Since the authors have used intrinsic optical imaging, they could not access to the dynamics of the lateral activation that would be averaged out in the observed activation maps (see Kisvárdy 2016 for further discussion).

## Anatomical measures

In a recent anatomical study, Martin et al. (2014) carefully re-evaluated the orientation bias of horizontal boutons from upper layer pyramidal neurons in cat area 17 using single cell intracellular labelling, optical imaging to reveal the orientation map and advanced cluster-by-cluster analysis of synaptic boutons. In their analysis, Martin et al. (2014) compared the distribution of the preferred orientations spanned by the neuron's dendritic arbors (used to estimate the neuron's preferred orientation) and the preferred orientation covered by axonal clusters of the neuron (Fig. 3A). In the example of Fig. 3A, B, the preferred orientation of the dendrite (red) matched the one of the local cluster (green) but not of the distal cluster (blue). Over 33 neurons, their results revealed a very large variability in the orientation selectivity of their distal clusters (coloured in Fig. 3C), as estimated by their *Similarity Index* (1 corresponding to the same orientation preference distribution with respect to the neuron's preferred orientation, 0 to an orthogonal orientation preference). Their results demonstrated the existence of a very large variance of SI (0.13–0.96) out of all 51 clusters they observed over the 25 neurons. To test whether the clusters positions within the orientation maps occur by chance,

the authors made a detailed bootstrap statistical analysis of all 51 clusters, taking into account the bias that is introduced by the orientation map layout, the cluster size and position relative to the soma. Using this analysis, they found that a quarter of their clusters (14/51 clusters recently updated to 17/65, personal communication from Ruesch and Martin) were not positioned randomly in the map. Interestingly, only 9% (6/65) of these clusters (see their suppl Fig. 10I) had a significantly high SI, above the upper bound (hence iso-oriented), and 5% (3/65) below the lower bound (hence cross-oriented). In contrast 12% (8/65) were located in position of the orientation map unlikely to occur by chance whilst being neither iso- nor cross-oriented with the labelled cell. As a conclusion, only a small minority of clusters (9%) are significantly tuned to iso-orientation from non-random position in the orientation map. Furthermore, as shown by Buzás et al. (2006), this bias tends to decrease with lateral distance of the clusters, which is further in accordance with Chavane et al. (2011). Finally, as observed in Huang et al. (2014), Martin et al. (2014) did not find any specific alignment of the cluster distributions in the retinotopic map that could favour collinear vs orthogonal interactions with the cell's preferred orientation. At a more macroscopic level, diversity was also shown from animal to animal in tree shrew V1, specifically



Martin et al (2014)

**Fig. 3** Probing for the orientation selectivity of individual horizontal axons (modified from Martin et al. 2014 with CC-BY permission). **A** Axon of an intracellularly labelled neuron is displayed over the orientation map. Ellipses show clusters of boutons (not shown) for local and more distal positions. Dendritic tree (inset) was colour coded by the orientation value of their corresponding pixels (white dot indicates the soma). Scale bar, 0.5 mm. **B** Radial plots of the normalized distribution of the number of boutons counted within each local (green line) and distal clusters (blue line), but also the

dendrite (red line) over the preferred orientations. The individual vectors forming these hemispheric plots were summed up to generate one sum vector (bold vector). The length of this sum vector is termed as the “tuning” of the dendrite or cluster. **C** Similarity index (SI) values for individual clusters of 33 neurons sorted by normalized depth of soma. (Top) Neurons (x axis) can have clusters (colour coded by rank) with different SI values (y axis). The histogram on the right summarizes the SI across clusters of all neurons (grey = distal, black = local). Note the large variance within and across neurons

in the fine orientation/retinotopic arrangement of extracellular anatomical labelling (i.e. a population of neurons). Their detailed analysis of Hunt et al. 2011 showed that there is a diversity of co-circular connectivity rules across animals, some showing a significant bias towards co-circular rules, some towards anti-circular rules and others without biases. Thus, as stated by Martin et al. (2014), the horizontal axons thus cannot be treated as an homogeneous network with a net iso-oriented bias, but rather should be described as strongly heterogeneous, an heterogeneity that may be the core of its function (see also Kisvárdy 2016).

## Computational model linking structure to function

Population measures and anatomical data constrain connectivity in cortical space; however, the link between known anatomical details and the resulting functional expression (in terms of neural activity) is not obvious. Computational models provide a means to explore this relationship directly. Modelling studies of V1 consider a range of connectivity rules, and these frequently allow for the shaping of connection strengths based on the difference of orientation preference between connected sites. Abstracted models of single hypercolumns implement cross-orientation interactions in local circuits that further tune selectivity derived from weakly tuned LGN inputs (Ben-Yishai et al. 1995). Similar mechanisms for orientation selectivity in V1 have been explored in models with recurrent, lateral connections over short distances (between neighbouring hypercolumns in L4) (Somers et al. 1995; Kang et al. 2003; Chariker et al. 2016). Connections that extend over many mm of cortex (i.e. across multiple pinwheels) are considered in visual cortex modelling studies of contextual modulation (Rubin et al. 2015), motion illusions (Rangan et al. 2005), geometric visual patterns (Bressloff et al. 2001; Baker and Cowan 2009; Carroll and Bressloff 2016), travelling waves (Bressloff and Carroll 2015), and in a general setting (Raizada and Grossberg 2003). Whilst models do commonly feature a decay (e.g. exponential or Gaussian) in the strength of orientation-based connections with distance (Goldberg et al. 2004; Blumenfeld et al. 2006), the tuning strength is not distance dependent, is rarely systematically investigated and is not constrained by anatomical data. The function of patchy long-range connections has further been investigated in contexts not specific to orientation encoding (Voges et al. 2010; Voges and Perrinet 2012). In general, long-range connections feature a strong iso-orientation bias motivated by long-held assumptions that do not take into account the more recent functional and anatomical studies that motivate a modification of this rule.

Furthermore, a common modelling choice for local excitation-inhibition connectivity is the so-called Mexican hat

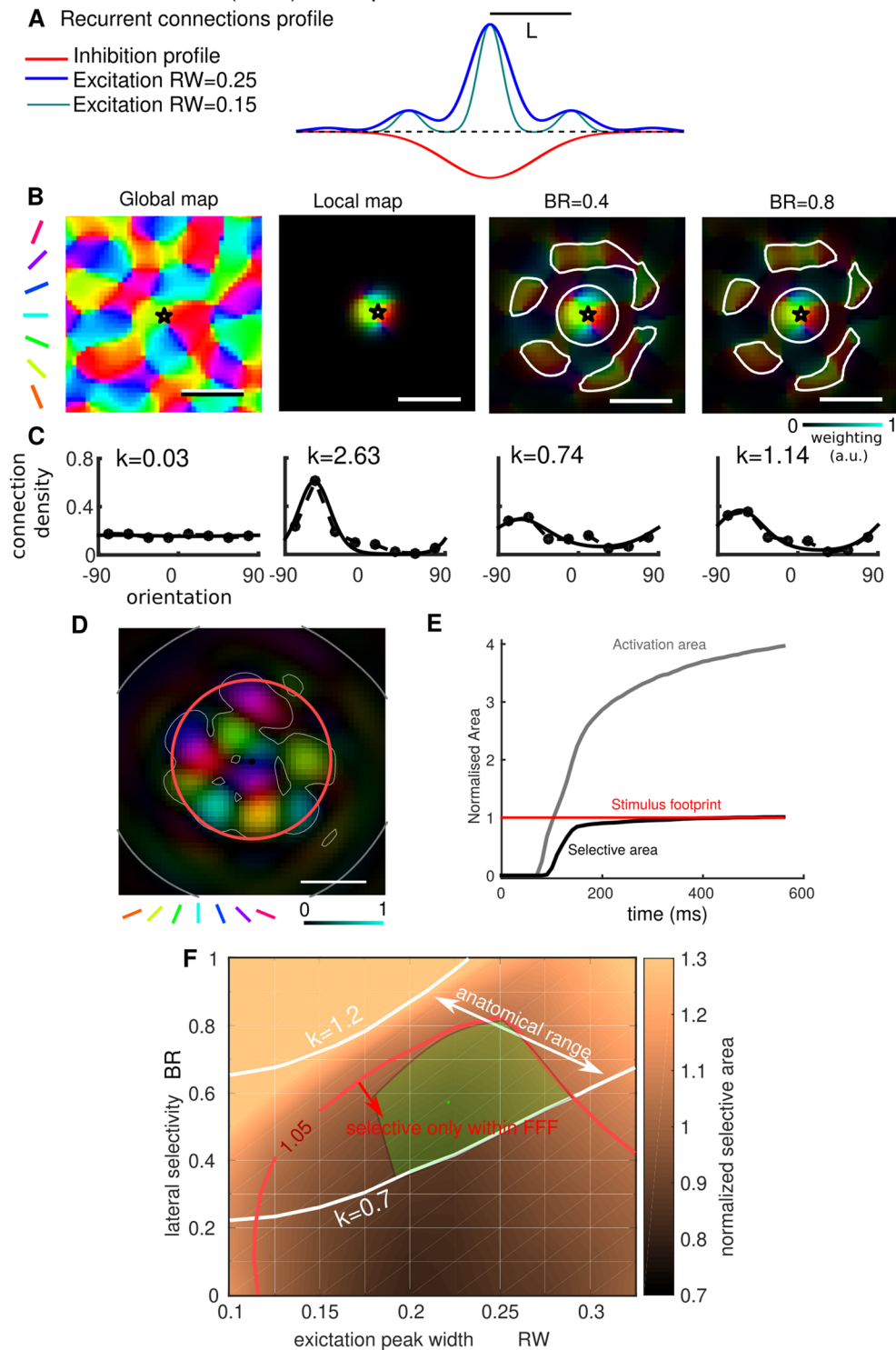
with inhibition extending further than excitation (Marr and Hildreth 1980; Grossberg 1983; Somers et al. 1995; Bressloff et al. 2001). This choice is known to generate stable localized patterns of activity (rather than spatial unstable dynamics that spreads across cortex) (Laing and Troy 2003); however, excitatory connections in V1 can extend many mm further than the local inhibitory footprint. In general, models that also feature long-range excitation are used to study unbounded patterns of activity rather than localized responses to inputs (Bressloff et al. 2001; Blumenfeld et al. 2006). Rankin et al. (2014) extended the results of Laing and Troy (2003) to demonstrate that localized inputs can generate stable localized activity patterns (rather than spreading activity) with a connectivity rule (as suggested in Buzás et al. 2001), and similar to Fig. 4A) that features long-range excitation, extending much further than the local inhibitory network.

Rankin and Chavane (2017) developed a planar spatial model of orientation selective activation in V1 L2/3 with the aim of bridging between known anatomical constraints on the tuning of long-range connections (Buzás et al. 2006) and the functional expression of laterally propagating activity driven by localized stimuli (Chavane et al. 2011). A neural field architecture with orientation-specific subpopulations provides a mesoscopic description of neural activity, ideal for comparison with the temporal and spatial resolution in VSDI imaging experiments. A novel connectivity function was flexibly parameterized to investigate clustering of connections, their orientation bias and balance between excitation and inhibition. We adopted the non-orientation-specific nature of local excitatory connections (Buzás et al. 2006) and inhibitory connections (Buzás et al. 2001); see also Koch et al. (2016) for a discussion of orientation specificity of excitatory and inhibitory connections. Taking motivation from Buzás et al. (2001), longer-range excitatory connections are proposed here to, although decaying with distance, form in rings at multiples of the hypercolumn separation  $L$  (Fig. 4A). This allows for the following important features to be captured: that excitatory connections (1) drop in number at a range of  $L/2$ , (2) have a peak at a range of  $L$  (and multiplies therefore) and (3) can extend several mm across cortex. Two parameters were tuned to agree with the available data from Buzás et al. (2006), the width of peaks in number of excitatory connections (RW; two values shown in Fig. 4A) and their orientation bias (BR; illustrated in Fig. 4B, C).

We found a significant overlap between the anatomically relevant parameter range and patterns of cortical activation consistent with imaging experiments (see Fig. 4D-F, Chavane et al. 2011). Hence, this computational approach allowed us to reconcile the imaging results with the reported level of orientation bias from anatomical studies. Specifically, Chavane et al. (2011) found a sharp decay of orientation selective activation at the stimulus



## Rankin &amp; Chavane (2017) - Computational model



retinotopic footprint border, resulting in peripheral activation that was not orientation selective (compare Fig. 2A with 3D-E). Our results demonstrate that this sharp decay is contingent on three factors: the diffuse clustering of long-range connections, the intermediate range (consistent

with anatomy) of their orientation bias and sufficient balance between excitation and inhibition. It is worth noting that orientation-biased long-range connections can recruit a local non-orientation-biased network at the target, resulting in non-orientation-specific activation (like

**Fig. 4** Neural field model to reconcile structure with function in primary visual cortex. Definition of model connectivity with anatomical constraints (A–C) and illustration of model behaviour with operating region in agreement with functional characteristics (D–F). **A** Radial connectivity profile for inhibition (Gaussian decay) and excitation (locally Gaussian decay, longer-range connections peak in number at distance  $L$  and multiples thereof). Ring width (RW) of peaks in excitatory connections illustrated for two values. **B** Example of local preference map and resulting lateral connectivity for two values of the orientation bias of recurrent connections (BR). **C** Orientation tuning for each panel in **B** above (circles) with tuning parameter  $k$  from a best-fit von Mises distribution (solid lines). Orientations are evenly represented in the global map but strongly biased at around  $-60^\circ$  for the local excitatory component (local map). The orientation bias of lateral connections increases to around  $k=1$  for  $BR>0.5$  (similar values reported in Buzás et al. 2006). **D** Model simulation snapshot at 600 ms showing the orientation-selective component within a thin white contour, confined to the feedforward footprint FFF of the stimulus in red; the much broader non-orientation-specific activity falls within a grey contour extending beyond the plot limits. **E** Time history of the area within the non-orientation-specific contour and the orientation-selective contour. **F** Colour map across range of RW and BR values showing the selective area as in **D** normalized by FFF. Within the red contour the selective activation is constrained to the FFF. White contours show the anatomically constrained range for the connectivity parameters RW and BR where  $k=0.7\text{--}1.2$ . In the green region other constraints on the correct orientation and the radial decay rate of orientation selectivity are also satisfied (modified from Rankin and Chavane 2017 with CC-BY permission)

in Huang et al. 2014). The modelling work illustrates that the observed levels of orientation bias in anatomical studies actually predict long-range activation beyond the retinotopic stimulus footprint with a sharply decaying orientation selectivity profile.

The model offers further insights into the mechanistic value of excitatory-inhibitory balance, and of intermediate levels of orientation bias in long-range connections. Long-range excitatory connections (reaching much further than the lateral inhibitory profile) could easily lead to destabilization of activity generated by localized visual stimuli. Our model was used to show that if the orientation bias of lateral connections is excessively strong, or if inhibition is particularly weak, the network operates close to an instability leading to unbounded cortical activation. This provides another line of evidence in favour of distance-decaying orientation bias in lateral connections. Diversity of long-range connections increasing with distance (i.e. decreasing orientation bias with distance) reflects a potential need to activate a broader range of orientations as we move further from a local stimulus with a specific orientation. Furthermore, the fact that, under particular circumstances, the preferred orientation of the horizontal propagation may be at odds with the underlying orientation preference map could unravel some new unexpected computational capacities of the horizontal network, which may be present in visual areas beyond V1. For instance, the ability to link information across position and orientation for non-co-circular filters, which is important

for processing objects with sharp angles. In line with this hypothesis, Chavane et al. (2011) showed that the spread of selective activity is not fixed but can increase with increasing spatial summation generated by annular stimuli.

## Functional advantages of such an organization

The insights we have reviewed at the physiological and modelling levels support a range of novel hypotheses for the organization of long-range lateral connectivity in the primary visual cortex. A functional approach, asking "*why should neurons in V1 be connected laterally?*" provides a complementary perspective. Indeed, a major argument is that the structure of V1 should fulfil its function and implement principles of perceptual organization, such as the principle of *good continuation* to bring a contour's constituent edges together into a unified global percept (Wertheimer 1923). How might these principles connect knowledge across anatomy, physiology, theory and modelling?

## Principles of perceptual organization in natural images

A major constraint for neurons in the primary visual cortex is that information is encoded locally and must be integrated globally across the visual field. Perceptual principles organizing the different fragments of an image can be directly extracted by analysing a database of natural images. One such principle is that pairs of edges in natural images are most likely organized along aligned contours, and more generally on a common circle (Sigman et al. 2001); the authors extracted edges from natural images and estimated the orientation of each edge. For each pair of active edges, they showed that the angle of maximum interaction corresponds to a configuration for which they are close to co-circularity. This long-range correlation is a marker of the structure of natural images and may provide strong prior knowledge for the perceptual organization of low-level features.

Such a structural prior can be described as a form of "association field" extending the concept of a neural receptive field to long-range local interactions. The seminal paper by Field et al. (1993) defines the association field as the set of local oriented elements (edges) in the visual field that facilitates the detection of a central oriented target. They showed that the association field obeyed a co-circular rule. In other words, if a common circle can pass through the central target and the peripheral element, they will facilitate each other's detection and generate suppression otherwise. This association field is invariant to translations or rotations. It extends the prior of collinearity (like-to-like) or co-circularity (Sigman et al. 2001) to a more generic description of

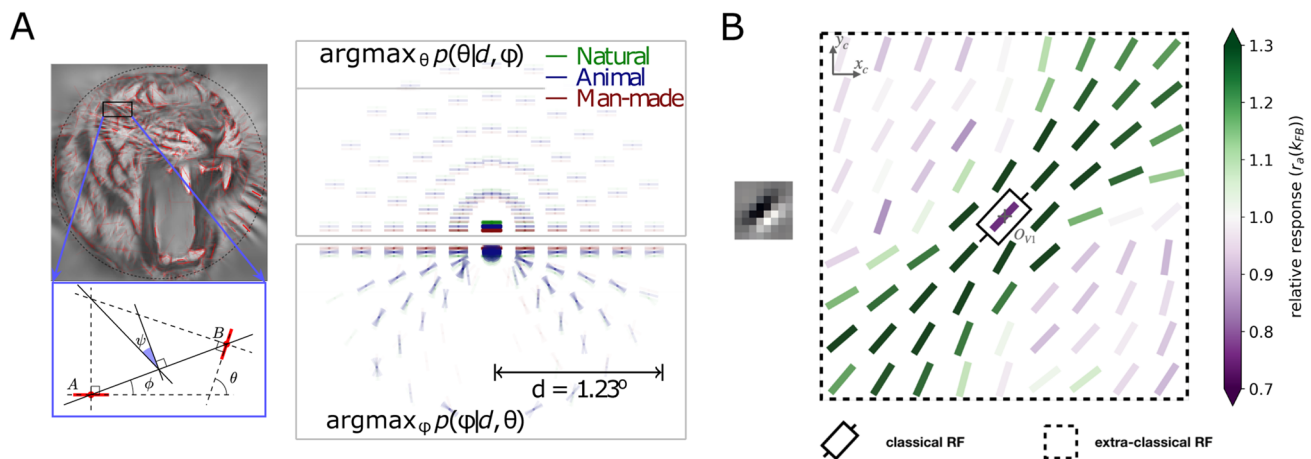
all possible co-occurrences. In particular, by exploring the interactions of edge pairs, they showed that these association fields explain the detection of paths embedded within a field of randomly oriented edges. The association field can then be understood in light of the computer vision problem of curve tracing. Parent and Zucker (1989) described it as a diffusion process over the tangent field of oriented edges, thus suggesting a principled and biologically realistic framework for association fields using long-range interactions.

This principle can be extended to explain psychophysical experiments in humans. Geisler et al. (2001) took a similar approach by reporting the full statistics of natural image edge co-occurrences. This yields a valuable model for the statistics of neighbouring edges. First, the edges are organized into parallel textures favouring parallel edges, and second, there is a bias for co-circular edges (see Fig. 5A). Using a Bayesian approach, the authors derived a clustering scheme for chaining edges into contours that was confirmed by psychophysical experiments. This approach was later extended to the high-level cognitive problem of image categorization (Perrinet and Bednar 2015). The authors showed that using supervised learning, one could derive a scheme using the association field in that image to categorize whether it contains an animal. This simple model achieved similar performance to humans and to a deep hierarchical model (Serre et al. 2007). Surprisingly, the model

made similar errors to humans. This illustrates first that association fields can be used to both group edges based on different tasks or to categorize images. This also shows that for the association field reflecting the statistics of edge co-occurrences in natural images, different datasets may lead to different association field structures (see Fig. 5A). As a consequence, it seems relevant at behavioural and ethological levels that mechanisms exist to tease apart the slight differences between the co-occurrence patterns present in different images, for instance, the surprising patterns of a perfect co-circularity, or that of a pair of rare but informative orthogonal edges forming a T-junction. This would then explain part of the variability in the association fields which can be involved in visual integration processes.

### How do these principles translate to the cortical space?

As Geisler et al. (2001) states, "*the obvious hypothesis for the local grouping is a neural population with the receptive field structure matched to the edge co-occurrence statistics*". Yet, the emergence of receptive field properties is a combination of anatomy and the dynamics of individual neurons. Can we link the statistics of natural images to the structure of processing in the primary visual cortex?



**Fig. 5** Function and diversity of association fields. **A** Following the work of Geisler et al. (2001), one could derive the association field from the statistics of natural images. This involves extracting edges from images (red segments) and computing for each pair the difference of angle  $\theta$  and the relative azimuth  $\phi$  of one edge compared to the other. This allows one to quantify the association field as a histogram, relative to a reference edge placed in the middle, for the most likely difference of angle—showing a prominent preference for parallel textures (top) or the relative azimuth, showing a prior for co-circular co-occurrences (bottom). The association field may vary for different databases with an excess of co-circularity in images containing animals, illustrating the variety of statistics faced by the visual system (Perrinet and Bednar 2015; modified with permission CC-BY).

**B** Boutin et al. (2021) described a biologically realistic multi-layer model of the visual cortex. The model is shown natural images and is optimized to represent images in the most efficient way. Edge-like filters emerge (see an example in the inset) and we show here the interaction of this edge with other edges outside the range of its classical receptive field. This pattern shows a large facilitatory (green) or inhibitory (purple) effect relative to a model without feedback. This functional modulation of the association field shows the importance of the activity in the whole network and we have further shown its shape could widely vary within the network and for different types of images, such as images of faces (Boutin et al. 2021; modified with permission CC-BY)

Olshausen and Field (1996) set out to show how the structure of V1 microcolumns can optimize the efficiency of the neural representation for natural images. Hyvärinen and Hoyer (2001) extended this to include a regularization of the representation with cortical topography. Franciosi et al. (2021) recently developed a biologically realistic, two-layered V1 sparse predictive coding model, including pooling mechanisms to impose a neighbourhood prior in cortical space, which includes by construction the possibility of representing as channels in each layer a variety of interaction patterns. Similarly, complex cells and topographic maps emerge, demonstrating the transfer of cortical connectivity in V1 to perceptual grouping principles. More surprisingly, depending on the density of neurons, different structures emerge to optimize cost efficiency: in addition to mammalian-specific features (such as topographical maps), a rodent-specific salt-and-pepper map emerges for models with a lower cell density. Interestingly, by focusing on this multi-channel convolutional architecture, the second layer showed a diversity of connectivities across channels, suggesting that differing anatomical constraints may induce different patterns of long-range lateral interactions.

A multi-layer sparse predictive coding model (Boutin et al. 2021) allows for the influence of an extrastriate cortical area, such as V2 on to V1 to be modelled. The activity in the layers of this model emerge from the recurrent interactions between neurons within and across layers (rather than a feed-forward pass as in convolutional networks). Convergence to an efficient representation of edge filters and interaction maps (resembling association fields) emerges after several processing iterations. However, training on different natural image datasets can produce different interaction maps, in accordance with Perrinet and Bednar (2015). For example, training on images of human faces generated features resembling mouths or eyes, resulting in more sparse and longer-range interactions. This suggests that instead of a simple similarity rule, lateral interactions between neurons reflect the variety of feature dependencies attached to the respective neurons. In addition, similarly to physiological observations (Gilbert and Li 2013), we observed that the interaction becomes sharper with stronger feedback (see Fig. 5B), which provides a synergy between the different pieces of information encoded by the network, as illustrated by improved performance for denoising natural images.

### Function and dynamics of long-range lateral interactions

Overall, these theoretical models propose, as an alternative to the like-to-like structure, that there should be a wide variety of long-range lateral interaction patterns. It should be noted that most of the models described above deal with

static natural images, whereas the visual world is characterized by a wealth of different dynamic scales, which raises the question of the role of neural dynamics in long-range lateral interactions.

If one imagines an edge moving in a direction parallel to its orientation, we can infer that we are following the tangent to a continuous contour. On the contrary, if the orientation of the edge is perpendicular to its direction, it is more likely that we are seeing a moving bar. This simple prototypical example shows that depending on the local intrinsic context, the optimal integration rule may change, as evidenced by intracellular recordings (Gerard-Mercier et al. 2016). If these interactions can be implemented via different contextual cues, such as recurrent or feedback connections, it is also possible that the multidimensional representation of information on the cortical surface is much more than a simple topographical orientation map.

In addition, there is physiological evidence that association maps can be dynamically influenced by the task at hand. In McManus et al. (2011), using a delayed-to-sample matching task, the authors trained monkeys to detect different patterns: a circle, a wiggle, or a line, which were embedded in a grid display of randomly oriented edges similar to that of Field et al. (1993). They found that depending on the pattern being searched, the recorded association field adapted to preferentially exploit collinearities (for lines) or co-circularities (for circles). Such a differential processing raises an implementation problem for the unsupervised schemes described above. This problem could be solved in a supervised learning scheme (Perrinet and Bednar 2015) but raises the question of how this supervised credit is assigned in V1. A similar problem is inherent in the backpropagation rule in generic deep learning paradigms which can be solved in a predictive coding framework (Boutin et al. 2021).

Lastly, the anatomical connectivity may be patchy for different functions than just connecting like-to-like orientation patches. Indeed, patchy connections likely play an important role in combining information from multiple visual cues beyond orientation, including context (Martin et al. 2017). Indeed, modelling work has shown that patch-based connectivity increases the versatility of the dynamic repertoire of neural states (Voges et al. 2010). That work compared networks of realistic conductance-based neurons with a range of connectivity rules. These rules had different complexities, from a completely random connectivity, to a neighbourhood-based local connectivity, and more interestingly, clustered networks, including a patch-based connectivity rule. This was extended in a further modelling study (Voges and Perrinet 2012) to include a comparison between a pure random patch-based connectivity and partially overlapping patches. As noted in Kisvárdy (2016), these patch-based connectivity rules were sufficient to induce a large dynamic repertoire, such as rhythms or travelling waves, and were for instance



characterized by enhanced variety in the shape of the power spectrum of population activity. In particular, such a range of dynamic behaviours is much richer when compared to those obtained with a random or local connectivity rule. Patchy connectivity rules introduce a heterogeneity in the lateral connections, which seems essential for building up an efficient population code (Martin et al. 2014). In particular, this would allow the propagation of combinations of contextual cues which would reflect the richness of visual information in natural scenes.

To conclude this section, the function studied in these theoretical models hints at a solution using a superposition of different long-range connectivity profiles. The diversity of patterns and their adaptability to the task or statistics should overall improve processing efficiency in the primary visual cortex. Yet there remain open questions regarding the richness of these like-to-all patterns. Theories suggest potential strategies for addressing these open questions explicitly in neurophysiology, for example, by synthesizing optimally responsive, model-driven dynamic stimuli (Walker et al. 2019).

## Discussion

In this review, we have documented convergent evidence from physiology, anatomy and computational models that the orientation selectivity of horizontal network connectivity in the primary visual cortex of carnivores, ungulates and primates is more versatile than initially proposed, which motivates the necessity to revisit the like-to-like connectivity rule (Mitchison and Crick 1982) still dominant today. At the anatomical level, there seems to be a diversity of connection rules between presynaptic source and postsynaptic target. At the individual cell level, anatomical studies have shown that the rule changes as a function of cell type (excitatory vs inhibitory) and layer/map locations (Yousef et al. 2001, 1999; Kisvárdy et al. 1994; Buzás et al. 2001; Karube and Kisvárdy 2010; Karube et al. 2017). Within one presynaptic origin, a large diversity exists with only moderate bias in the range of 1.5–2 times greater than chance (Bosking et al. 1997; Kisvárdy 1997; Schmidt et al. 1997; Malach et al. 1993; Rochefort et al. 2009). More recent work by Martin et al. (2014) on the upper layer pyramidal neurons of the cat V1, with a cluster-by-cluster analysis of horizontal boutons, has shown the existence of a very large diversity from like-to-like, like-to-any, like-to-all and like-to-unlike connectivity rules. Altogether no net significant bias towards one of these rules could be observed in their bootstrap statistical analysis. Taken together, these anatomical results show that there is diversity in the connectivity rules both within and between neuronal types and locations (see Kisvárdy 2016 for an extensive review). At a more macroscopic level, it is

interesting to note that Hunt et al. (2011) also observed that the co-circularity rule varies from animal to animal.

When probed with functional measures of neuronal activity in response to a local visual stimulus, using techniques sensitive to subthreshold membrane potential fluctuations (Chavane et al. 2011) or an optogenetic activation of specific orientation columns (Huang et al. 2014), the diversity revealed in anatomical studies leads not only to the absence of net bias towards like-to-like interactions along the horizontal network (see Alonso and Kremkow 2014a, b), but also the absence of patchy activation of the horizontal spread of activation. Importantly, VSDI measures demonstrate a clear exponential decay of the like-to-like connectivity bias with horizontal distance, an effect also observed in anatomy (Buzás et al. 2006; Martin et al. 2014). At short-range distances, similar iso-orientation biases, as reported in anatomical studies, were observed. However, after the equivalent of one hypercolumn, no significant bias could be observed (Chavane et al. 2011).

All these papers therefore demonstrate the existence of a connectivity rule that links neurons in the primary visual cortex depending on their preferred orientation, neuronal type, position (layer and orientation map) and intra-cortical distance. This multidimensional connectivity rule is also subject to large diversity not just from neuron to neuron but also from animal to animal. Due to the difficulty in making predictions from this complex pattern, it is necessary to use computational approaches to probe for the expected functional behaviour of such a network. In Rankin and Chavane (2017), we developed a neural field model and demonstrated that the functional results observed in Chavane et al. (2011) are indeed the expected mesoscopic behaviour of such a network when its connectivity is constrained to match the orientation bias of connections from anatomy (Buzás et al. 2006), thus demonstrating that the functional observations are to be expected given our understanding of anatomical characteristics.

In this review, we wish to update the accepted like-to-like connectivity rule widely assumed as the building block for connecting a local neuronal network from one position in the visual field to its postsynaptic targets. The connectivity rule should be revised to a distance-dependent formulation: from like-to-like bias at short horizontal distance to like-to-all at long horizontal distance (Fig. 1E). The space constant of the decrease of the like-to-like bias is about one hypercolumn distance. Functionally we can speculate that this translates to an iso-orientation bias for neurons with overlapping receptive fields and no net bias for neurons with non-overlapping receptive fields.

This may be at odds with the well-documented association field schema and co-circularity rules observed in natural scenes (Sigman et al. 2001; Geisler et al. 2001). However, it is important to differentiate the basic connectivity building

block, that specifies unidirectional rules from a presynaptic region to a postsynaptic target, from lateral interactions, that are evoked by more complex stimuli that co-activate both presynaptic and postsynaptic regions (e.g. as in cross-correlation studies). The like-to-all long-distance connectivity rule can be seen as generic and allows for a variety of interactions in the orientation and spatial domain. Importantly, this is possible if we take into account the large local diversity observed at the neuronal level (Chavane et al. 2011, Martin et al. 2014, see also Monier et al. 2003). Our proposition here is that such a rule could account for a wealth of interaction rules depending on the stimulus and/or the task. For instance, this would allow one to account for the interactions necessary to process orientation discontinuities, such as junctions or corners. Neurons in V1 have indeed been reported to be sensitive to orientation discontinuities, independent of the absolute orientation of the stimulus set (Sillito et al. 1995; Jones et al. 2001). This result could not be explained solely by iso-oriented lateral interactions. Such diversity could also contribute to shaping the orientation tuning of neurons away from the primary orientation preference (i.e. horizontal and vertical orientations, Vidyasagar and Eysel 2015). More generally, using natural, stationary scenes and/or contour integration tasks may indeed favour association field interactions. However, depending on the type of natural images, Perrinet and Bednar (2015) have shown that these interactions may already differ significantly (see also Boutin et al. 2021). Moreover, dynamic non-stationary visual stimuli, such as a simple moving object, and tasks that rely on motion integration for instance could lead to different associative rules for motion (Gerard-Mercier et al. 2016). In the case of integrating information along a coherent path, for instance, visual information should be transported in a direction of motion that can be in the cross-orientation dimension (Perrinet and Masson 2012).

Importantly, co-circularity rules that link orientation and position with respect to a central oriented feature are not found in the anatomy (Martin et al. 2014; Hunt et al. 2011), nor were they found by Huang et al. (2014) using optogenetic stimulation pattern in the horizontal network. This further supports a dynamic, context-dependent emergence of specific rules, such as co-circularity for contour integration in natural images, through higher-order network interactions. In that respect, Chavane et al. (2011) observed that increasing spatial summation of the stimulus increases the propagation of iso-orientation activity, even if the basic connectivity profile was shown to be not selective to orientation at long range. This means that from a basic unselective building block, selective interaction can occur (for a proposition of possible mechanisms see suppl FigA5 in Chavane et al. (2011)). This effect could result from the fact that inhibitory neurons tend to make more horizontal connections with cross-orientation neurons (Kisvárdy et al.

1994; Buzás et al. 2001). Increasing spatial summation could change the orientation dependence of excitatory/inhibitory balance and lead to the emergence of tuned activity at longer distance. More generally, the emergence of new selectivity depending on the stimulation pattern (or the task) is rendered possible by the existence of local diversity of orientation-selective connections at neuronal level (Monier et al. 2003; Chavane et al. 2011; Martin et al. 2014). Therefore, different stimulation patterns will lead to activation of different recurrent subnetworks and the emergence of a variety of selectivity characteristics. It is indeed now well documented that non-trivial, paradoxical effects can arise from recurrent balanced networks (Tsodyks et al. 1997; Ozeki et al. 2009; Pattadkal et al. 2018). In our model, we indeed observed that manipulating the balance between excitation and inhibition (i.e. reducing inhibition strength) predicts the emergence of spurious orientation selective activation through long-range lateral connections (Rankin and Chavane 2017).

Given the non-trivial effects that can arise with more complex stimuli, a number of avenues remain open to build on theoretical and modelling work. The model developed in Rankin and Chavane (2017) could also be used to investigate selective recruitment and spatial summation in regions between localized oriented stimuli (Chavane et al. 2011; Huang et al. 2014). Indeed, increasing spatial summation increases the slope of selectivity decay at the stimulus boundary, whilst selective propagation reaches further across cortex, a property easily explored in the model by a more diverse class of localized stimuli. More generally, the model could be used to make predictions to decipher the selective functional connectivity rules that link position and orientation in cortical space. Importantly, it could also be extended to differentiate inhibitory cell subclasses as reported in Buzás et al. (2001). As such it could generate functional predictions on, e.g. the role of long-range basket cell connections that preferentially target cross-orientations. Extending the framework further, a feature space including spatial frequency could be used to investigate lateral connections in light of recent work on interactions between orientation and spatial frequency maps (Romagnoni et al. 2015; Ribot et al. 2016).

In this review, we mostly focus on revisiting the connectivity rule of intra-cortical horizontal networks. However, it is important to consider that such connectivity patterns can also be influenced by feedback from higher cortical areas that provides a more diffuse and divergent input to the primary visual cortex (Salin et al. 1989, 1992). Anatomical studies in the cat suggest that area 17 and area 18 cells are preferentially connected when they share similar preferred orientations (Gilbert and Wiesel 1989). In the monkey, feedback from higher areas (V2 and V3) to V1 show the variable level of patchiness (Stettler et al. 2002; Angelucci et al. 2002), unselective to orientation (Stettler

et al. 2002). In the cat, electrophysiological and inactivation studies of various downstream areas seem to influence only response amplitude or tuning width of neurons in area 17 of the cat, but not orientation preference (Martinez-Conde et al. 1999; Wang et al. 2000, 2007; Huang et al. 2004, 2007; Liang et al. 2007; Shen et al. 2006, 2008; Galuske et al. 2002). However, it is important to consider that feedback will interact with the horizontal network as demonstrated in monkey visual cortex, with either specific interactions as suggested by Gilbert and Li (2013, for review), or contributing to centre-surround processing (Hupé et al. 1998; Roberts et al. 2007; Poort et al. 2012; Nurminen et al. 2018).

In conclusion, we believe that there are enough arguments today to accept a change to the connectivity rules for horizontal axons in V1, that is consistent with both new structural and new functional evidence. It remains to be established how this complex multidimensional rule (orientation  $\times$  distance  $\times$  neuron type  $\times$  neuron location) is expressed under different stimuli and task configurations. It would be important to understand what is the minimal stimulus design that can trigger particular tuned interactions for various spatial positions and whether it involves precisely the same neurons in the large-scale neural network. To test predictions that can arise from theoretical and computational approaches, new experimental tools to visualize large massive neural networks at neuronal level and sensitive to membrane potential fluctuations will be needed. Recent neuro-technological advances in awake animals, such as all-optical tools to measure and control a large set of neurons (Ju et al. 2018; Zhang et al. 2018), and the development of new genetically encoded voltage indicators that allow simultaneous two-photon microscopy of subthreshold activity recording from many cells (Villette et al. 2019); this provides the ideal experimental setting to probe the complex and dynamic network interactions underlying stimulus and task-dependent processing.

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