

Connective field modeling

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ARTICLE INFO

Article history:

Accepted 6 October 2012

Available online 27 October 2012

Keywords:

Visual cortex

fMRI

Functional connectivity

Connective field

Population receptive field

ABSTRACT

The traditional way to study the properties of visual neurons is to measure their responses to visually presented stimuli. A second way to understand visual neurons is to characterize their responses in terms of activity elsewhere in the brain. Understanding the relationships between responses in distinct locations in the visual system is essential to clarify this network of cortical signaling pathways. Here, we describe and validate connective field modeling, a model-based analysis for estimating the dependence between signals in distinct cortical regions using functional magnetic resonance imaging (fMRI). Just as the receptive field of a visual neuron predicts its response as a function of stimulus position, the connective field of a neuron predicts its response as a function of activity in another part of the brain. Connective field modeling opens up a wide range of research opportunities to study information processing in the visual system and other topographically organized cortices.

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Introduction

The interpretation of visual neuroscience measurements made in different parts of the brain is unified by the receptive field concept. A measurement at any point in the visual pathway is usually summarized by referring to the stimulus properties (location, contrast, color, motion) that are most effective at driving a neural response. Stimulus-referred receptive fields provide a common framework for understanding the sequence of visual signal processing. The classic receptive field construct summarizes the entire set of signal processing steps from the stimulus to the point of measurement. This sequence of signal processing can be made explicit by modeling how the activity of one set of neurons predicts the responses in a distinct set of neurons. Characterizing the responses of a cortical neuron in terms of the activity of neurons in other parts of cortex can provide insights into the computational architecture of visual cortex. Such measurements are exceptionally difficult to achieve with single-unit recordings. The relatively large field of view in functional magnetic resonance imaging (fMRI) offers an opportunity to measure responses in multiple brain regions simultaneously, and thus to derive neural-referred properties of the cortical responses. These cortical

response properties provide important information about how neuronal signals are transformed along the visual processing pathways. For example, stimulus-referred measurements in cortex show that visual space is sampled according to a compressive function (i.e., the V1 cortical magnification factor corresponds to a logarithmic compression of cortical space with eccentricity). Neural-referred measurements show that this compression is established at the earliest stages of vision; later visual field maps sample early maps uniformly and inherit the early compressive representation (Harvey and Dumoulin, 2011; Kumano and Uka, 2010; Motter, 2009).

A limitation in developing models of how fMRI responses in two parts of cortex relate to each other is that the problem is under-constrained. For example, there are many voxels in visual area V1, and there are many ways in which these responses could be combined to predict the response in a voxel in V2. Hence, any estimate requires imposing some kind of prior constraint on the set of possible solutions. Heinze and colleagues (Heinze et al., 2011), for example, used a support vector machine approach to reduce the dimensionality of the solution of V1 signals and predict responses in extrastriate cortex. Here, we take a different approach based on the idea that in retinotopic cortex connections are generally spatially localized. We build on a model-based population receptive field (pRF) analysis that was developed to estimate the stimulus-referred visual receptive field of a voxel (Dumoulin and Wandell, 2008). In the pRF analysis, the receptive field is modeled and fit to the fMRI signals elicited by visual field mapping stimuli. This is done by generating fMRI signal predictions from a combination of the receptive field model and the experimental stimuli. In the present

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analysis, fMRI signal predictions are generated from fMRI signals originating from the regions of cortex covered by a model of the inter-areal connective field (Angelucci et al., 2002; Lehy and Sejnowski, 1988; Sholl, 1953). Conceptually, this means that the localized activity in one cortical region acts as a stimulus for voxels in another region. We model the connective field as a two-dimensional, circular symmetric Gaussian that is folded to follow the cortical surface (Fig. 1). The assumption of a Gaussian connective field model is motivated by findings that the receptive fields of two extrastriate areas in the macaque, V4 and MT, can be described as two-dimensional, circularly symmetric, Gaussian sampling from the V1 map (Kumano and Uka, 2010; Motter, 2009). The Gaussian width parameter provides crucial information about the connective field, namely its size. Because the inter-areal connective field size is a measure of spatial integration, the analysis can be used to trace the extent of spatial integration as information moves from the primary visual cortex to higher visual areas.

Methods

Participants

Cortical responses were measured using 7 Tesla fMRI in subjects S1 and S2 with 1.6, 2.0 as well as 2.5 mm isotropic voxel sizes. S1 also participated in a 3 Tesla fMRI experiment with a 2.5 mm isotropic resolution. During all experimental sessions, the participants

viewed high-contrast drifting bar stimuli interposed with mean luminance periods. Both subjects had normal visual acuity. All experiments were performed with the informed written consent of the subjects and approved by the UMCU Medical Ethics Board.

Stimulus presentation

The visual stimuli were generated in the Matlab programming environment using the Psychtoolbox extensions (Brainard, 1997; Pelli, 1997). Stimuli were displayed in one of two configurations. In both configurations, the participants viewed the display through an angled mirror. The first display configuration consisted of an LCD projecting the stimuli on a translucent display at the back of the magnet bore with a maximum stimulus radius of 5.5 degrees of visual angle. This configuration was used during the 7 T experiments. The second display configuration consisted of an LCD with a maximum stimulus radius of 6.25 degrees of visual angle. This configuration was used during the 3 T experiment.

Stimulus description

In both the 7 T and 3 T experiments, we measured responses to drifting bar apertures at various orientations that exposed a high-contrast checkerboard pattern (Dumoulin and Wandell, 2008; Harvey and Dumoulin, 2011; Zuiderbaan et al., 2012). Parallel to the

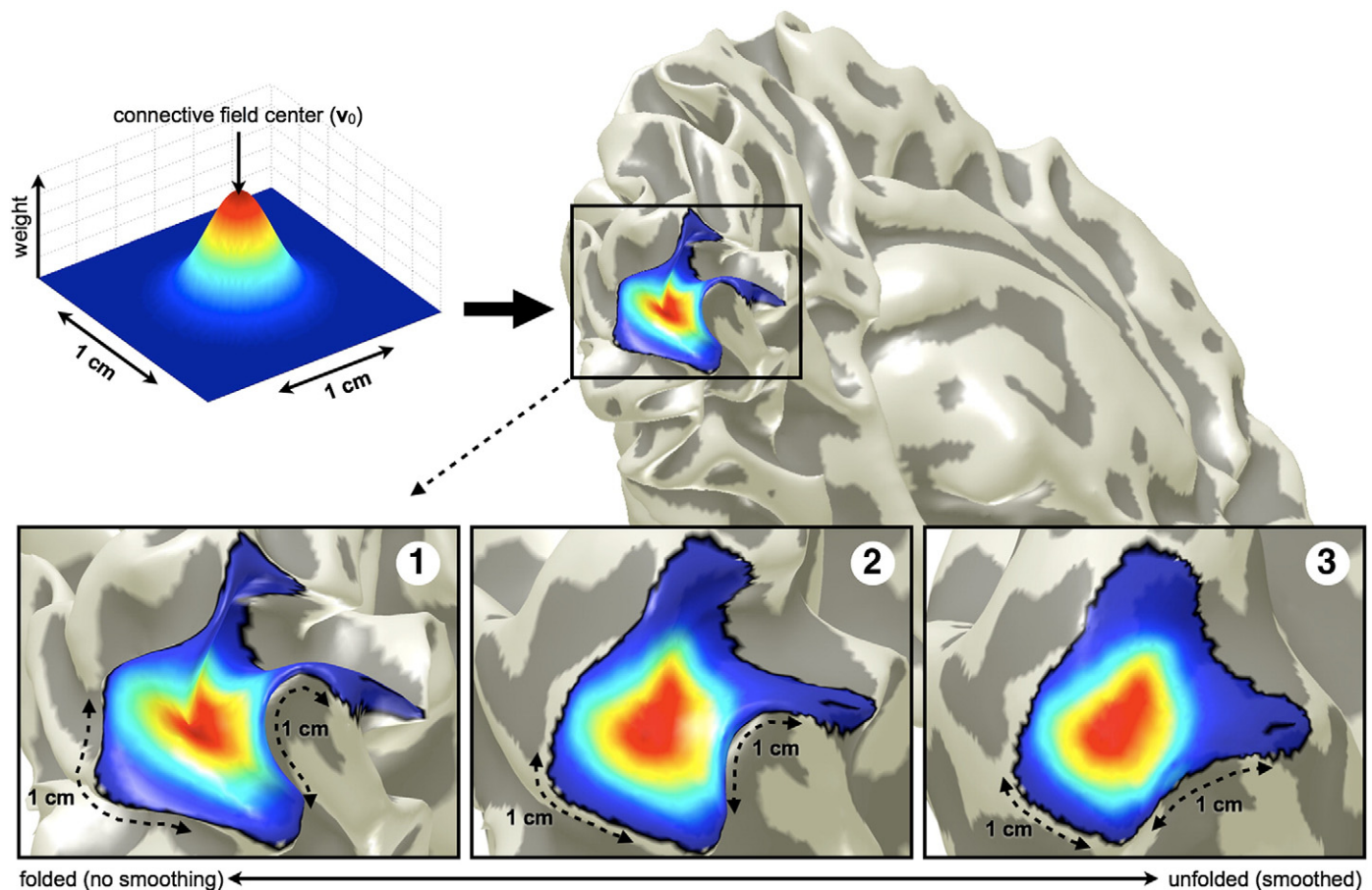


Fig. 1. Connective field models follow the curvature of the cortex. A two-dimensional, Gaussian connective field model (top-left) is defined as a function of Dijkstra's shortest path distance between pairs of vertices in a three-dimensional mesh representation of the original, folded cortical surface (top-right). The advantage of this approach is that the measurement of cortical distance avoids the distortions introduced if the Gaussian were projected onto a flattened, two-dimensional cortical surface representation. Panels 1, 2 and 3 (bottom) further illustrate the connective field model projection when the surface mesh is unfolded (smoothed).

bar orientation, alternating rows of checks moved on opposite directions. This motion reversed at random intervals of at least 4 s. The bar width subtended 1/4th of the maximum stimulus radius. The bar moved across the stimulus window in 20 equally spaced steps. Four bar orientations and two different motion directions for each bar were used, giving a total of 8 different bar configurations within a given scan (up, down, left, right, and the four diagonals). After each horizontal and vertical pass, a 30 s zero contrast, mean luminance stimulus was presented.

Magnetic resonance imaging

Magnetic resonance images were acquired with 3 T and 7 T Philips MRI scanners equipped with sixteen-channel SENSE head coils. Foam padding was used to minimize head motion. Functional T2* weighted echo-planar images were acquired at both field-strengths. For the 3 T runs, images were acquired at an isotropic resolution of 2.5 mm, 24 slices. The TR was 1500 ms, the TE was 30 ms, and the flip-angle was 70°. For the 7 T runs, images were acquired at isotropic resolutions of 1.6 mm, 2.0 mm, and 2.5 mm. The TR was 1500 ms, the TE was 25 ms, and the flip-angle was 80°. The functional runs each were 248 time frames (372 s). The first eight time-frames (12 s) were discarded. At 7 T, eight functional runs were performed using 1.6 mm isotropic voxels, 5 functional runs were performed using 2.0 mm isotropic voxels, and 5 functional runs were performed using 2.5 mm isotropic voxels. At 3 T, 9 functional runs were performed. In addition to the functional runs, high-resolution T1-weighted whole-brain anatomical MR images were acquired at 3 T for both subjects.

Preprocessing of MR images

The T1-weighted anatomical MRI data sets were re-sampled to a 1 mm isotropic resolution. Gray and white matter were automatically segmented from the whole-brain anatomical data set using FSL (Smith et al., 2004) and subsequently hand-edited to minimize segmentation errors (Teo et al., 1997). The cortical surface was reconstructed at the white/gray matter border and rendered as a smoothed 3D surface (Wandell et al., 2000). Motion correction within and between scans was applied (Nestares and Heeger, 2000). Finally, functional images were aligned with the whole-brain anatomical segmentation.

Population receptive field analysis

Population receptive field (pRF) parameters were estimated according to procedures described by Dumoulin and Wandell (Dumoulin and Wandell, 2008). Briefly, fMRI time-series predictions were generated by varying the parameters (x , y and σ) of a circular symmetric Gaussian pRF model across a wide range of plausible values. The optimal pRF parameters were found by minimizing the residual sum of squares (RSS) using a coarse-to-fine search. First, the fMRI data were re-sampled to an 1 mm isotropic resolution within the identified gray matter. The fMRI data were then smoothed along the cortical surface using a diffusion smoothing process that approximated a 5 mm full-width at half-maximum Gaussian kernel, after which the pRF parameters were estimated for a sub-sample of the voxels and interpolated for the remaining voxels. Subsequently, an optimization algorithm (Fletcher and Powell, 1963) was applied for every voxel whose initial estimates exceeded 10% of the variance explained, so that the pRF model predictions were fitted to fMRI time courses without any spatial smoothing. As in previous work, eccentricity, polar angle, and pRF size maps were derived from the best pRF fits that exceeded 15% of the variance explained (Baseler et al., 2011; Haak et al., 2012; Winawer et al., 2010).

Connective field modeling

As in the population receptive field analysis, the connective field parameters were estimated from the time-series data using a linear spatiotemporal model of the fMRI response:

$$y(t) = p(t)\beta + \varepsilon \quad (1)$$

where $p(t)$ is the predicted fMRI signal, β is a scaling factor that accounts for the unknown units of the fMRI signal, and ε accounts for measurement noise. In the present analysis, $p(t)$ is calculated using a parametrized model of the underlying neuronal population and the spatial distribution of its inputs laid out across the cortical surface. The model is estimated by finding the parameters that best predict the observed fMRI time course $y(t)$.

The current implementation of the analysis uses a circular symmetric Gaussian connective field model. The two-dimensional circular symmetric Gaussian connective field of voxel \mathbf{v} , $g(\mathbf{v})$, is defined by two parameters: \mathbf{v}_0 and σ :

$$g(\mathbf{v}) = \exp\left[-d(\mathbf{v}, \mathbf{v}_0)^2 / 2\sigma^2\right] \quad (2)$$

where $d(\mathbf{v}, \mathbf{v}_0)$ is the shortest three-dimensional distance along the cortical manifold between voxel \mathbf{v} and the connective field center \mathbf{v}_0 , and σ is the Gaussian spread (mm) across the cortical surface. The distance $d(\mathbf{v}, \mathbf{v}_0)$ was computed using Dijkstra's algorithm (Dijkstra, 1959) on a triangular mesh representation of the gray/white matter border. The calculation of $g(\mathbf{v})$ is done for each gray-matter voxel \mathbf{v} directly adjacent to the white-matter in a predefined region-of-interest; V1 for example. Distances were calculated separately for each hemisphere: hence, a connective field model solution for any given voxel comprised voxels either in the ipsilateral hemisphere or the contralateral hemisphere, but not both.

The neuronal population inputs, $a(\mathbf{v}, t)$, are defined as the percent BOLD signal change ($\Delta\%$) time course for voxels \mathbf{v} . Low-frequency signals were removed from these time-courses using a discrete cosine transform (DCT) high-pass filter. The time-series prediction is then obtained by calculating the overlap between the connective field and the neuronal population inputs (note that there is no need to do a convolution with the hemodynamic response function):

$$p(t) = \sum_{\mathbf{v}} [a(\mathbf{v}, t) \cdot g(\mathbf{v})] \quad (3)$$

Finally, the optimal connective field parameters were found by minimizing the residual sum of squares (RSS) between the prediction, $p(t)$, and the observed time-series, $y(t)$. To do this, we generated various different fMRI time-series predictions by varying the connective field parameters \mathbf{v}_0 , and σ , across all existing voxel positions on the V1 surface (both hemispheres) and 50 sigma values ranging from 0 to 25 mm. Neither spatial smoothing nor interpolation was performed; all connective field models were fitted to the observed time-series. Best models were retained if the explained variance in the fMRI time-series exceeded 15%.

Computing the V1 sampling extent

We obtained the V1 sampling extent by first finding the linear relationship between the pRF laterality index (λ), which indicates the extent to which a pRF overlaps with the ipsilateral visual field (0 represents no overlap, 0.5 represents 50% overlap), and the connective field size (σ):

$$\sigma(\mathbf{v}) = m \cdot \lambda(\mathbf{v}) + b \quad (4)$$

where m is the slope of the line and b is the intercept. We then computed the V1 sampling extent (r) for each voxel \mathbf{v} using the following formula:

$$r(\mathbf{v}) = \sigma(\mathbf{v}) + [2 \cdot \lambda(\mathbf{v}) \cdot b] \quad (5)$$

Statistical analyses

We derived the percent variance explained to specify how well the pRF and connective field models fit the fMRI time series. These r^2 values were calculated from the total sum of squares of the observed time series and the residual sum of squares of the predicted versus observed time series. Given that the time series consisted of 240 samples, the 15% variance explained threshold that was applied to all further analyses corresponds to $p < 0.001$, corrected for testing ~100,000 different models per voxel (Bandettini et al., 1993). Furthermore, the correlation coefficients that were derived to quantify the agreement between the polar angle maps were circular–circular correlation coefficients to appropriately assess the association between these two angular variables (Behrens, 2009; Jammalamadaka and Sengupta, 2001). Finally, all ranges reported in text represent 95% confidence intervals for the bootstrapped weighted means ($N = 1000$) using Student's t -distribution. Where appropriate, Bonferroni correction was applied to the confidence intervals – as reported in both the text and in the figures.

Results

We first employed the conventional model-based pRF method (Dumoulin and Wandell, 2008) to derive estimates of the population receptive field for each voxel in visual cortex. These pRF estimates were used to delineate visual maps V1, V2, V3, and hV4 (Amano et al., 2009; Brewer et al., 2005; Dougherty et al., 2003; Dumoulin and Wandell, 2008; Harvey and Dumoulin, 2011; Wandell and Winawer, 2011; Wandell et al., 2007; Winawer et al., 2010; Zuiderbaan et al., 2012). We then employed the new analysis (Fig. 2) to derive several inter-areal connective field models for each voxel within the delineated visual areas. For the sake of brevity, we refer to these models in a compact manner; a connective field model m for voxel v can be specified by $S \rightarrow R$ (“ S projects on R ”), if m has been defined on cortical surface S , and v falls in cortical region R (note that if S represents the visual field, the same notation can also be used to describe conventional population receptive fields). In this notation, we derived the following connective field models: $V1 \rightarrow \{V2, V3, hV4\}$. Across the two subjects and the three resolution sets, the best-fitting models explained on average 76%, 66%, and 46% of the time-series variance in V2, V3, and hV4, respectively.

Fig. 3 further shows two examples of the connective field model fit to the fMRI time-series. A comparison of the connective field model prediction with the conventional pRF model prediction suggests that the connective field model captures more of the time-series variance than the pRF model. Indeed, across subjects and voxel sizes, we found an average difference (connective field – pRF) of ~23%, ~14% and ~10% in variance explained for visual areas V2, V3 and hV4 respectively. This improvement is particularly evident during the mean luminance periods when there was no stimulus. During the mean luminance periods, the conventional pRF predicts a uniform signal: in contrast, the connective field model can capture some of the time-varying signals. Note, however, that the standard pRF prediction could be improved by adding extra model parameters. For example, one could add a second Gaussian spread parameter to model the pRF's suppressive surround and explain more of the negative trenches around the peaks (Zuiderbaan et al., 2012). In addition, some of the time-series variance during the mean luminance periods could be non-neuronal physiological noise (although the time series were averaged across runs).

Connective field modeling links a voxel in one brain region to many voxels in another region. The voxels in the two regions should respond to overlapping regions of visual space (i.e., they should have similar pRFs). This is because voxels that have similar patterns of stimulus-evoked responses will also have similar time-series. Hence, once the connective fields are known it should be possible to derive the visual field map in one area from the visual field map in another area. Qualitatively, Fig. 4 shows that this is indeed the case. Panels a and b depict the eccentricity and polar angle maps for visual areas V1–hV4 derived with conventional pRF mapping. In the same figure, panels d and e show the result of deriving the V2–hV4 maps from V1 using the connective field models. To quantify the agreement between the conventional (pRF based) and derived (connective field based) visual field maps in these areas, we computed the correlation between the visual field positions of the V2–hV4 voxels' standard pRFs, and the pRF locations of the V1 voxels corresponding to the V2–hV4 voxels' connective field centers. The visual field map

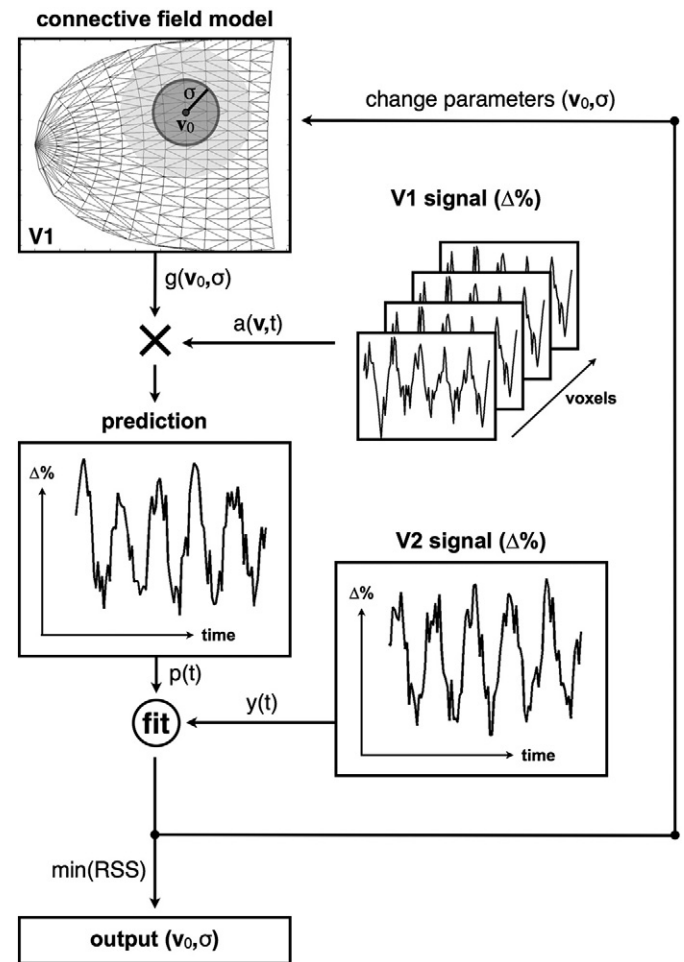


Fig. 2. Estimating the $V1 \rightarrow V2$ connective field for a V2 voxel. Assuming a linear relationship between the blood-oxygenation levels and the fMRI signals, the observed blood-oxygenation level-dependent (BOLD) time course, $y(t)$, can be described in terms of the predicted BOLD signal, $p(t)$. The prediction, $p(t)$, is calculated using a parametrized model of the connective field. The parameters of the connective field model are its center location, v_0 , in voxel coordinates, and the Gaussian spread, σ , laid out across the folded cortical surface in millimeters cortex. The definition of this circular symmetric Gaussian model is achieved by its projection on a three-dimensional mesh representation of the boundary between the gray and white matter of the brain. The predicted BOLD time course, $p(t)$, for a V2 voxel is then obtained by calculating the overlap between the connective field, $g(v_0, \sigma)$, and the fMRI signals in V1, $a(v, t)$. Finally, the optimal connective field model parameters are found by minimizing the residual sum of squares (RSS) between the prediction, $p(t)$, and the observed time series, $y(t)$.

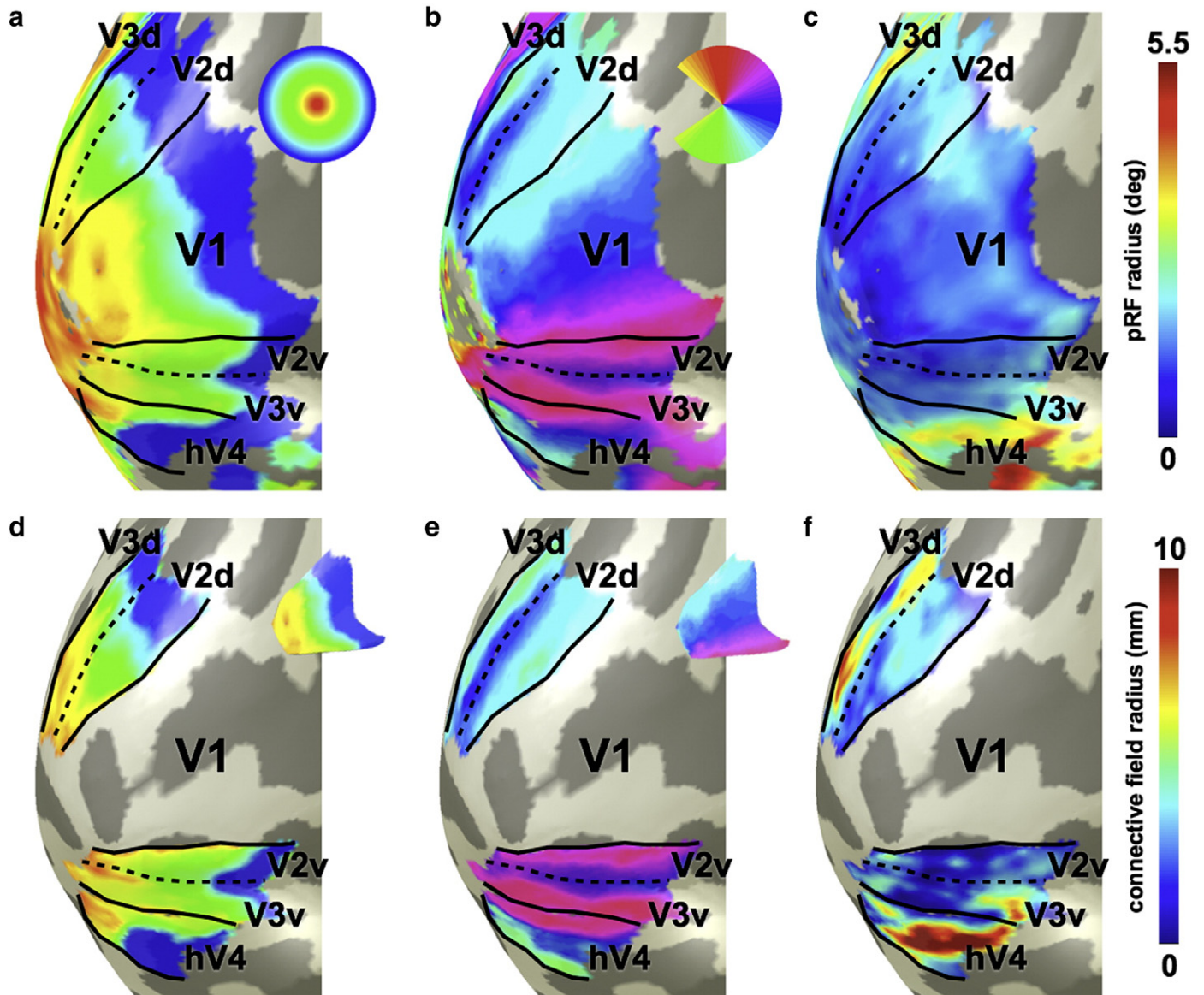


Fig. 4. Stimulus- and neural-referred maps on the posterior medial surface of the occipital lobe of the left cerebral hemisphere at 7 Tesla. (a, b) Stimulus-referred eccentricity and polar angle maps revealed using conventional pRF modeling. The pRF eccentricity and pRF polar angle were used to delineate visual areas V1–hV4. Insets indicate the color maps that define the visual field locations. (c) The stimulus-referred pRF size estimates, as indicated by the colors shown in the color bar. The pRF size increases with eccentricity for all visual areas shown. (d, e) Neural-referred eccentricity and polar angle maps derived from the best-fitting V1 > {V2, V3, hV4} connective field models in visual areas V2–hV4. The insets indicate the color maps that define the cortical locations, which are the V1 maps shown panels a and b. (f) The neural-referred connective field size, as indicated by the colors shown in the color bar.

factor, and also found a constant V1 sampling extent across eccentricity. Thus, in agreement with several past studies, the results are consistent with the idea that cortical magnification in extrastriate cortical areas (V2–hV4) is inherited from V1, and that there is no further magnification in the pooling of signals from V1.

From Fig. 5 it is also clear that the connective field size increases systematically between different visual field maps. This feature is expected on the basis that visual information converges up the visual processing hierarchy. If the connective field size corresponds to the radius of sampling from V1, then the sampling area is $\sim 30 \text{ mm}^2$ for V2, $\sim 90 \text{ mm}^2$ for V3, and $\sim 300 \text{ mm}^2$ for hV4. These values correspond to approximately 1/100, 1/25 and 1/8 of the total V1 hemispheric surface area (Andrews et al., 1997).

Finally, there are two important instrument-related factors that could influence the spatial specificity of the connective field estimate. The first reflects the fact that coarser fMRI resolutions result in a poorer

ability to estimate small changes in the connective field position and size. The second captures the feature that data from lower magnetic fields normally have a lower spatial specificity due to the increased intra-vascular contribution of draining veins (Logothetis, 2008; Ogawa et al., 1998). Therefore, we asked whether the connective field method is robust to changing these two parameters. Table 1 summarizes the effect of changing the resolution and field-strength on the correlation between the visual field maps derived using conventional pRF modeling and the connective field method. It is clear that increasing the voxel size from $\sim 4 \text{ mm}^3$ to $\sim 16 \text{ mm}^3$ and then decreasing the magnetic field strength from 7 to 3 Tesla does not systematically influence the accuracy by which the connective field method accurately links voxels with overlapping receptive fields. Fig. 6 further summarizes the effect of changing these two instrument-related features on the estimates of the V1 sampling extent. This figure indicates that the connective field size estimate is also robust to increasing the voxel size and decreasing

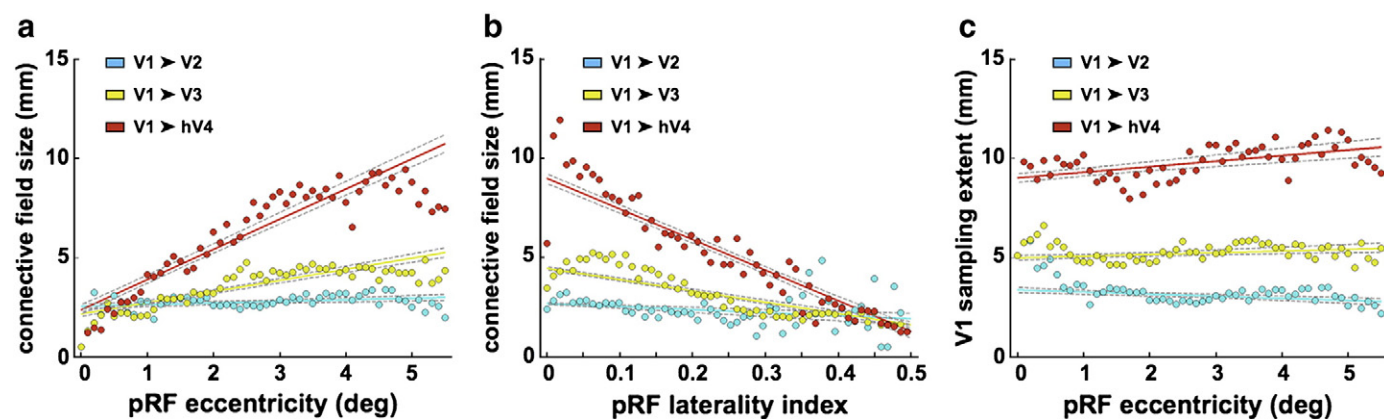


Fig. 5. The relationship between eccentricity and V1-referred connective field size in visual areas V2–hV4, grouped from both participants and all voxel sizes. (a) The connective field size increases up the visual processing hierarchy and is dependent on eccentricity. (b) The connective field size decreases as a function of the pRF laterality index, which indicates the extent to which a pRF overlaps with the ipsilateral visual field (0 represents no overlap, 0.5 represents 50% overlap). (c) Adjusting the graph in a for pRF laterality yields the V1 sampling extent, which appears roughly constant across eccentricities. Colored lines represent a linear fit to the bins (dots). The bins were bootstrapped and linear fits repeated to give the 95% confidence intervals (dashed gray lines).

the magnetic field-strength. These results show that the connective field method yields similar quantitative estimates from 3 T and 7 T data using a wide range of fMRI resolutions.

Discussion

Just as the receptive field of a visual neuron describes its response as a function of visual field position, the connective field of a neuron predicts its response as a function of activity in another part of the brain. Here, we have shown how fMRI can be used to estimate the connective field of a population of neurons. The analysis is based on a model of neuronal responses, accurately traces the fine-grained topographic connectivity between visual areas, and provides a quantitative estimate of the connective field size. The method is non-invasive and robust to changes in fMRI resolution as well as field-strength.

Connective field modeling represents a fundamental departure from the existing approaches to estimating fMRI connectivity in the human brain. One reason is that it emphasizes the spatial profile of the functional connectivity between brain areas: connective field modeling harnesses the core strengths of fMRI – a large field of view and high spatial resolution – to make inferences about the spatial coupling among brain areas. While some of the existing methods such as seed-voxel correlation mapping (Biswal et al., 1995) and independent component analysis (Arfanakis et al., 2000) are capable of producing spatial connectivity maps, these methods have not yet provided the level of spatial detail associated with connective field modeling. Another important aspect that is unique to connective field modeling is that it informs about the

direction of information flow in terms of converging versus divergent connections. For example, if the connective field size for V1 > V2 is larger than for V2 > V1, this would indicate that visual information converges from V1 to V2. To the best of our knowledge, existing methods for fMRI connectivity analysis only deal with the question of directionality by framing cortical information processing in terms of temporal causation (Buchel and Friston, 1997; Friston et al., 1995, 1997, 2003; Goebel et al., 2003; Harrison et al., 2003), which is not a trivial thing to do with fMRI due to its poor temporal resolution.

The connective field modeling method depends on some but not all of the unwanted factors that also influence the conventional pRF estimate (Dumoulin and Wandell, 2008; Smith et al., 2004). Common factors include eye and head movements, brain pulsations, and BOLD spread. These factors create a bias towards larger connective field size estimates, and add noise but no bias towards the connective field location estimates (Levin et al., 2010). Also like the pRF estimate, the connective field estimate is a statistical summary of the neuronal properties within the sampled voxel. Therefore, the connective field model parameters depend on the size and intrinsic properties of the sampled

Table 1
Correlation between the visual field maps derived using pRF and connective field modeling.

	7 T/1.6 ³ mm ³		7 T/2.0 ³ mm ³		7 T/2.5 ³ mm ³		3 T/2.5 ³ mm ³	
	<i>r</i>	ϑ	<i>r</i>	ϑ	<i>r</i>	ϑ	<i>r</i>	ϑ
V2	0.96	0.93	0.95	0.96	0.92	0.96	0.95	0.98
V3	0.93	0.89	0.92	0.93	0.87	0.96	0.90	0.92
hV4	0.83	0.82	0.82	0.86	0.78	0.84	0.85	0.64

The correlation coefficients compare the eccentricity (*r*) and polar-angle (ϑ) maps in V2–hV4 for subject S1 derived using conventional pRF modeling to those derived using the connective field method. The correlation coefficients were computed for all voxels in the regions of interest for which the best-fitting connective field models explained more than 15% of the time-series variance. Columns indicate different combinations of magnetic field-strength and the voxel size. All correlation coefficients were highly significant ($p < 0.0001$).

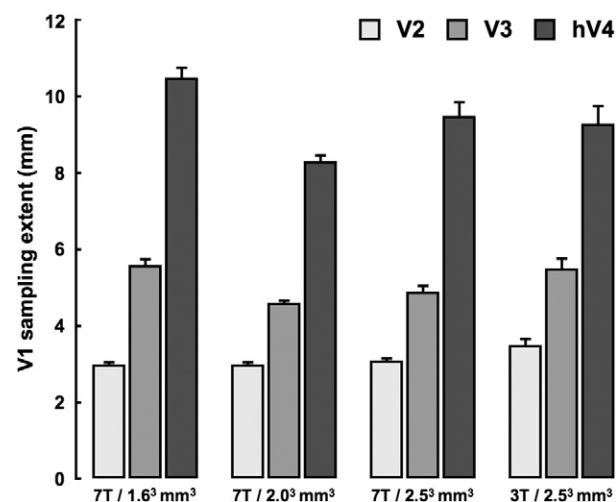


Fig. 6. Estimates of the V1 sampling extent for three different voxel sizes and two different field-strengths for visual areas V2–hV4 in subject S1. Increasing the voxel size from 1.6³ mm³ to 2.5³ mm³ and then decreasing the magnetic field-strength from 7 T to 3 T reveals that the connective field modeling method is robust to changing these instrumental parameters; there is noise but no bias. Error-bars indicate the 95% bootstrapped confidence intervals.

neuronal population. Different neuronal populations, for example in different cortical layers, will likely have different connective fields (Ress et al., 2007). Finally, pRF fits extending outside the maximum stimulus radius get noisy because they are based on less information than the fits that lie entirely within the stimulus area. The same is true of connective fields. If connective fields extended beyond the stimulated area of V1, then part of the connective field would be determined by the activity of the unstimulated part of V1. This unstimulated part will have lower amplitude responses than the stimulated area, so estimates here will be noisier; connective field model solutions will generally not be great for voxels near the edge of the stimulus representation.

In the present implementation of the analysis, we used a single circular symmetric, two-dimensional Gaussian connective field model. This model provides a compact description of the connective field using only two parameters. Other models, however, may also be used. The single isotropic Gaussian connective field model could be readily replaced with sums and differences of Gaussians, an anisotropic Gaussian, or any other type of mathematical function to describe the connective field. Such models may be suitable to examine connective fields in other topographically organized cortices. In addition, determining what connective field forms best explain the fMRI time-series in the different visual areas could be a very fruitful approach for understanding the different types of computation across the visual pathways.

While different stimuli may alter the connective field estimate, an important feature of connective field modeling is that the analysis itself is stimulus-independent. Consequently, the connective field models also capture some of the spontaneous signal fluctuations during periods when there is no stimulus. Using connective field modeling, therefore, it should also be possible to extract the intrinsic properties of sensory information processing based on resting-state fMRI. This idea is supported by Heinze and colleagues' work, who showed that "non-invasive imaging techniques such as fMRI are applicable to study detailed spatial interactions between topographically organized cortical regions in humans even in the absence of inputs driving the system under investigation" (Heinze et al., 2011). It should be noted, however, that if connective field modeling were applied to resting-state rather than task-evoked responses, it would be important to adopt a physiological noise removal strategy, such as for example, global signal regression (Birn et al., 2006), retroicor (Glover et al., 2000), or drifter (Sarkka et al., 2012).

In conclusion, we have described and validated connective field modeling, a new model-based fMRI data-analysis that can be used to make inferences about how the spatial coupling among retinotopically organized brain regions is influenced by changes in experimental context, development, ageing, and disease. An important methodological difference between this and previous work is the use of a two-dimensional circular symmetric Gaussian connective field model. This is a valuable improvement because it is more interpretable biologically, and it allows for calculations on straightforward parameters such as the connective field size, a measure of spatial integration. Because the method is stimulus-agnostic, it should also be possible to employ the method to non-visual topographically organized brain regions as well as resting-state responses.

Acknowledgments

KVH and FWC were supported by a grant from Stichting Nederlands Oogheekkundig Onderzoek (SNOO) and European Union grants #043157 (Syntex) and #043261 (Percept). KVH was also supported by the Groningen Research School for Behavioral and Cognitive Neurosciences (BCN) and The Netherlands Society for Biophysics and Biomedical Engineering (VVB-BMT). FWC and SOD were supported by The Netherlands Organization for Scientific Research (NWO grant #433-09-251). JW was supported by National Institutes of Health grant EY022116. BAW was supported by National Eye Institute grant EY03164. The software

is freely available and will be distributed as part of the VISTA software package (<http://white.stanford.edu/software/>).

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