

High resolution quantitative and functional MRI indicate lower myelination of thin and thick stripes in human secondary visual cortex

Daniel Haenelt^{1,2*}, Robert Trampel¹, Shahin Nasr^{3,4}, Jonathan R. Polimeni^{3,4,5}, Roger B. H. Tootell^{3,4}, Martin I. Sereno⁶, Kerrin J. Pine¹, Luke J. Edwards¹, Saskia Helbling^{1,7}, Nikolaus Weiskopf^{1,8}

*For correspondence:
haenelt@cbs.mpg.de

¹Department of Neurophysics, Max Planck Institute for Human Cognitive and Brain Sciences, 04103 Leipzig, Germany; ²International Max Planck Research School on Neuroscience of Communication: Function, Structure, and Plasticity, 04103 Leipzig, Germany; ³Athinoula A. Martinos Center for Biomedical Imaging, Massachusetts General Hospital, Boston, MA 02129; ⁴Department of Radiology, Harvard Medical School, Boston, MA 02114; ⁵Harvard-MIT Division of Health Sciences and Technology, Massachusetts Institute of Technology, Boston, MA 02139; ⁶Department of Psychology, College of Sciences, San Diego State University, San Diego, CA 92182; ⁷Poeppel Lab, Ernst Strüngmann Institute (ESI) for Neuroscience in Cooperation with Max Planck Society, 60528 Frankfurt am Main, Germany; ⁸Felix Bloch Institute for Solid State Physics, Faculty of Physics and Earth Sciences, Leipzig University, 04103 Leipzig, Germany

Abstract The characterization of cortical myelination is essential for the study of structure-function relationships in the human brain. However, knowledge about cortical myelination is largely based on post mortem histology, which generally renders direct comparison to function impossible. The repeating pattern of pale-thin-pale-thick stripes of cytochrome oxidase (CO) activity in the primate secondary visual cortex (V2) is a prominent columnar system, in which histology also indicates different myelination of thin/thick versus pale stripes. We used quantitative magnetic resonance imaging (qMRI) in conjunction with functional magnetic resonance imaging (fMRI) at ultra-high field strength (7 T) to localize and study myelination of stripes in four human participants at sub-millimeter resolution *in vivo*. Thin and thick stripes were functionally localized by exploiting their sensitivity to color and binocular disparity, respectively. Resulting functional activation maps showed robust stripe patterns in V2 which enabled further comparison of quantitative relaxation parameters between stripe types. Thereby, we found lower longitudinal relaxation rates (R_1) of thin and thick stripes compared to surrounding gray matter in the order of 1–2%, indicating higher myelination of pale stripes. No consistent differences were found for effective transverse relaxation rates (R_2^*). The study demonstrates the feasibility to investigate structure-function relationships in living humans within one cortical area at the level of columnar systems using qMRI.

Introduction

In primates, visual information sent from the primary visual cortex (V1) to the secondary visual cortex (V2) is segregated into distinct modules known as thin, thick and pale stripes (*Hubel and Livingstone, 1987; Livingstone and Hubel, 1987*). These stripes form a columnar system in the sense that their functional properties extend roughly through cortical depth (*Tootell and Hamilton, 1989*). Functional properties include the sensitivity to different visual features like color, orientation, binocular disparity and motion, which are largely processed in different stripe types and sent to distinct cortical areas. For example, thin stripes are sensitive to color content and project to functional area V4, whereas thick stripes are more sensitive to binocular disparity and project to area MT (V5) (*Shipp and Zeki, 1985; Hubel and Livingstone, 1987; Livingstone and Hubel, 1987; Ts'o et al., 2001*).

Using cytochrome oxidase (CO) staining, these stripes were found first in squirrel monkeys and macaques as dark and pale patches organized in repeating pale-thin-pale-thick cycles, running through V2 and oriented approximately perpendicular to the V1/V2 border (*Livingstone and Hubel, 1982; Tootell et al., 1983*). In macaques, stripes of the same type have a center-to-center distance of around 4.0 mm and a width ranging from 0.7 mm to 1.3 mm (*Shipp and Zeki, 1985; Tootell and Hamilton, 1989*). In humans, these widths are approximately doubled in size (*Hockfield et al., 1990; Tootell and Taylor, 1995; Adams et al., 2007*).

Histological studies also showed a stripe pattern in V2 of post-mortem brain specimens when techniques for the staining of myelin were used (*Tootell et al., 1983; Krubitzer and Kaas, 1989; Horton and Hocking, 1997*). However, these studies gave an inconsistent picture of the correspondence between stripes defined by CO activity and myelin density. Staining with Luxol fast blue indicated stronger myelination in thin/thick (*Tootell et al., 1983*) stripes, while Gallyas silver staining showed pale (*Krubitzer and Kaas, 1989*) stripes being more myelinated. This discrepancy between myelin staining methods was replicated in another study in which several methods were compared to each other (*Horton and Hocking, 1997*). In addition to inconsistencies across staining methods, all standard histochemical methods are highly sensitive to the condition of the brain specimen (e.g. post-mortem delay time), variations in fixation and staining procedures, and exposure time (*Savaskan et al., 2009*).

Magnetic resonance imaging (MRI) provides an alternative view to study tissue microstructure in living humans with the possibility to generate a multitude of image contrasts which can specifically be sensitized to myelin (*Edwards et al., 2018; Weiskopf et al., 2021*). Quantitative MRI (qMRI) is a collective term of techniques that aims to isolate one source of image contrast and represent it as a single quantitative parameter map, e.g., maps of longitudinal relaxation rate (R_1), effective transverse relaxation rate (R_2^*), proton density (PD) or magnetic susceptibility. Parameter maps are created by combining multiple “weighted” MR images in a model-based fashion yielding reproducible and standardized measures in physical units, which are less dependent on the acquisition (*Edwards et al., 2018; Trampel et al., 2019; Weiskopf et al., 2021*). Furthermore, by separating different sources of image contrast, qMRI provides a closer relationship to specific microstructural components, e.g. myelin and iron (*Edwards et al., 2018; Weiskopf et al., 2021*). In this regard, it should be kept in mind that single qMRI parameter maps do not have the specificity to directly infer the abundance of single microstructural tissue components by being sensitive to multiple tissue components to a varying extent and therefore can only serve as a surrogate, e.g. in the study of cortical myelination (*Lazari and Lipp, 2021*). Nevertheless, concerning R_1 , it is known that cortical R_1 is largely affected by myelin content (*Stüber et al., 2014; Leuze et al., 2017*) and, hence, was often used as surrogate to study the macroscopic distribution of cortical myelination (*Sigalovsky et al., 2006; Dick et al., 2012; Sereno et al., 2013; Lutti et al., 2014*). In addition, it might be possible to exploit the complementary multi-modal information from different qMRI parameter maps using appropriate biophysical models to directly quantify the underlying tissue microstructure with higher specificity, which would open the way to in vivo histology (*Weiskopf et al., 2021*).

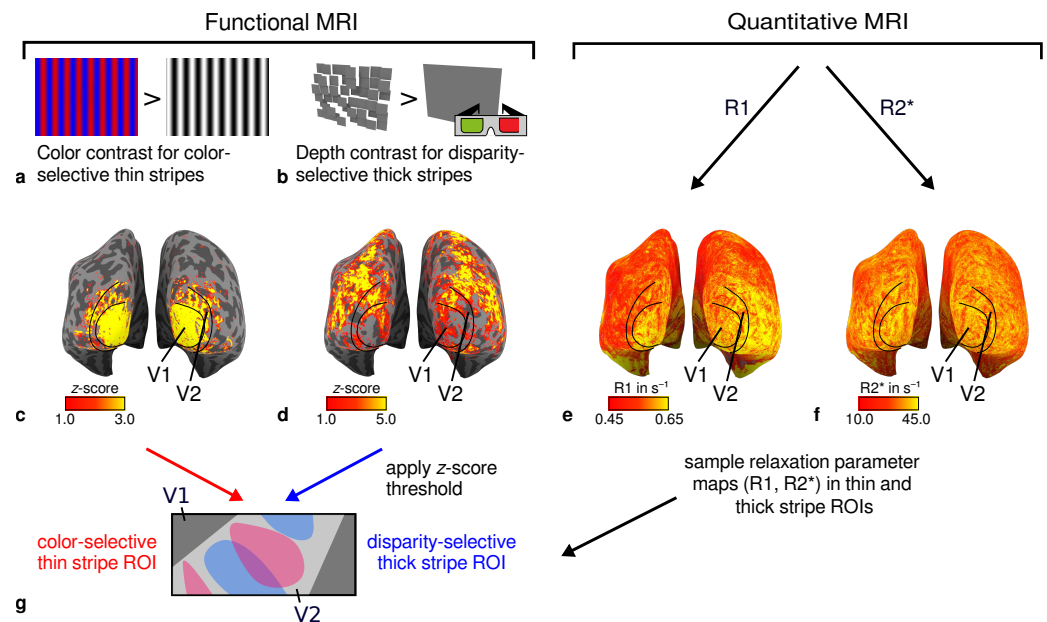


Figure 1. General overview of acquired MR data and their use in the analysis. (a) Example of chromatic and achromatic stimuli used to map color-selective thin stripes. (b) Schematic illustration of stimuli when viewed through anaglyph spectacles used for mapping disparity-selective thick stripes. These stimuli consisted of a disparity-defined checkerboard and a plane intersecting at zero depth, respectively. Exemplary activation maps from thin stripe (contrast: color > luminance) and thick stripe (contrast: depth > no depth) mapping sessions are shown for a representative participant (subject 3) in (c) and (d), respectively. Quantitative R_1 and R_2^* maps from the same participant are shown in (e)–(f). (g) Activation maps from (c) and (d) were used to define regions of interest (ROIs) for thin- and thick-type stripes in V2 by applying a z-score threshold. R_1 and R_2^* values were sampled in these ROIs for further analysis. Borders in (c)–(f) were manually defined on the basis of a separate retinotopy measurement.

Functional MRI (fMRI) allows in vivo localization of functional architecture. Recent developments in ultra-high field MRI enabled the functional localization of thin and thick stripes using high resolution fMRI (Nasr et al., 2016; Dumoulin et al., 2017; Navarro et al., 2021) by, e.g., exploiting their different sensitivity to color (Tootell et al., 1983, 2004) and binocular disparity (Peterhans and von der Heydt, 1993; Chen et al., 2008), respectively (Nasr et al., 2016). This allows MR-based investigations of mesoscale structure-function relationships in the same living participant, i.e., at the spatial scale of cortical columns and layers.

We combined the localization of V2 stripes using high resolution fMRI with qMRI measurements to infer myelination differences between stripe types. We robustly show lower R_1 values in color-selective thin and disparity-selective thick stripes in comparison to locations containing pale stripe contributions, which points towards higher myelin density in pale stripes. Whereas recent studies have explored cortical myelination in V2 in macaques (Li et al., 2019) and humans (Dumoulin et al., 2017) using non-quantitative, weighted MR images, to the best of our knowledge, we showed for the first time myelination differences using MRI on a quantitative basis at the spatial scale of columnar systems. This shows the feasibility to use high resolution qMRI in conjunction with high resolution fMRI to study the relationship between functional and structural properties of the brain in living humans, which is a fundamental goal in neuroscience.

Results

Participants ($n = 4$) were invited for multiple fMRI and qMRI sessions at 7 T (see Figure 1). On different days, we measured high resolution (0.8 mm isotropic) fMRI responses to stimuli varying in color and binocular disparity content, respectively, to locate color-selective thin stripes (color

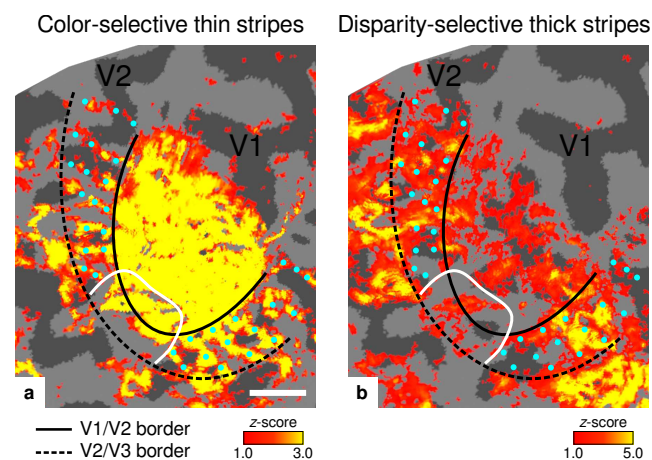


Figure 2. Activation maps for color-selective thin and disparity-selective thick stripes. Thin stripes (contrast: color > luminance) and thick stripes (contrast: depth > no depth) are shown as thresholded activation maps in (a) and (b), respectively. Both maps were averaged across sessions, sampled at mid-cortical depth and are illustrated on the flattened surface of the right hemisphere for one representative participant (subject 3). Surfaces were flattened using FreeSurfer (6.0.0, <http://surfer.nmr.mgh.harvard.edu/>) after cutting out a region on the surface mesh which included all stimulated portions of V1 and V2. Data from all participants can be found in **Appendix 1**. In V2, patchy stripes can be identified, which run through V2 oriented perpendicular to the V1/V2 border. Borders were manually defined on the basis of a separate retinotopy measurement, which is illustrated in **Figure 2-Figure Supplement 1** (see **Appendix 1** for retinotopy data from all participants). The white border illustrates a region around the foveal representation with missing activation. This region was excluded in the supporting analysis shown in **Figure 5-Figure Supplement 2**. For the main analysis, all data points in V2 were used. Manually drawn cyan dots mark activated regions in (a) to illustrate the alternating activation pattern between (a) and (b). Cortical curvature is shown in gray (sulcal cortex is dark gray and gyral cortex light gray). Another example can be found in **Figure 2-Figure Supplement 2**. Scale bar: 1 cm.

Figure 2-Figure supplement 1. Eccentricity and polar angle maps.

Figure 2-Figure supplement 2. Exemplary activation maps from another participant.

stripes) and disparity-selective thick stripes (disparity stripes) in V2 (Nasr et al., 2016). We use the terms color-selective thin and disparity-selective thick stripes acknowledging the close relationship between color processing and CO thin stripes (Tootell et al., 2004) and disparity processing and CO thick stripes (Chen et al., 2008), respectively, despite not having directly measured CO content in this study. In a separate session, we used the multi-parameter mapping (MPM) protocol (Vaculčíková et al., 2022) to acquire high resolution anatomical images with 0.5 mm isotropic resolution from which quantitative parameter maps (R_1 , R_2^* , PD) were derived.

Functional mapping of color-selective and disparity-selective stripes

Color- and disparity-selective stripes were identified in each individual in separate scanning sessions. **Figure 2** shows activation maps averaged over two sessions and sampled at mid-cortical depth from one representative participant (see **Appendix 1** for activation maps from all participants).

Color-selective thin stripes can be identified in **Figure 2a** with expected topography (Tootell et al., 1983; Nasr et al., 2016), i.e., they start at the V1/V2 border, radiate outwards in parallel and are confined to area V2. **Figure 2b** shows locations selective for binocular disparity. Activation maps for binocular disparity showed a less pronounced stripe pattern in V2. It should be noted that color-selective stripes are known to be largely confined to CO thin stripes (Xiao et al., 2003; Tootell et al., 2004), whereas selectivity for binocular disparity is found in all stripe types but most frequently in CO thick stripes (Peterhans and von der Heydt, 1993; Chen et al., 2008). In V1, no activation was found for binocular disparity in **Figure 2b** which is consistent with findings by Tsao

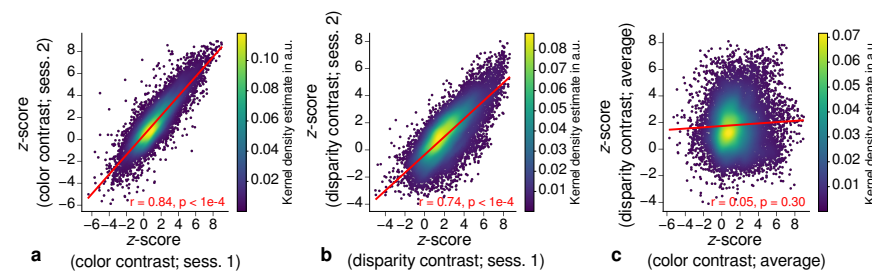


Figure 3. Repeatability of fMRI activation maps across scanning sessions. Scatter plots with kernel density estimation illustrate the consistency of activation maps across scanning sessions for one representative participant (subject 3). Sessions were carried out on different days and activation maps were sampled at mid-cortical depth. **(a)** shows correspondences of z-scores in V2 between single color-selective thin stripe mapping sessions (contrast: color > luminance). **(b)** shows the same for single disparity-selective thick stripe mapping sessions (contrast: depth > no depth). In **(c)**, correspondences of average z-scores (across sessions) between thin and thick stripe sessions are shown. Regression lines are indicated as red lines. Spearman's rank correlation coefficients r and p -values determined by permutation analysis (see Materials and methods) are annotated inside the plots and demonstrate high repeatability of color-selective thin and disparity-selective thick stripe scanning sessions. Note that the comparison between thin and thick stripe sessions shows no statistically significant correlation as expected from the interdigitated nature of both stripe types. Plots for all participants can be found in **Figure 3–Figure Supplement 1** and **Figure 3–Figure Supplement 2**.

Figure 3–Figure supplement 1. Correlation plots for single participants (subjects 1–2).

Figure 3–Figure supplement 2. Correlation plots for single participants (subjects 3–4).

et al. (2003); Nasr et al. (2016) while large V1 activation was found for color contrast in **Figure 2a** as also shown by Nasr et al. (2016).

Cyan dots were added in **Figure 2** to qualitatively illustrate the alternation of activation clusters between stripe types as expected from the thin/thick stripe arrangement. We note that, as **Figure 2a** and **Figure 2b** show results from two independent experiments, the alternating stripe pattern is not an intrinsic outcome of the experimental design.

Each stripe type was localized in two independent scanning sessions and activation maps were consistent between sessions of color and disparity stripe measurements, respectively. This is illustrated in **Figure 3**, which shows statistically significant correlations of activation maps between sessions for one representative participant.

Consistent qMRI maps across cortical regions and cortical depth

Figures 4a–b show an R_1 map sampled at mid-cortical depth for one representative participant. Primary motor and primary sensory cortical areas have higher R_1 values, congruent with higher myelin density in these areas (Flechsig, 1920; Glasser and Van Essen, 2011; Sereno et al., 2013). To further check the consistency of our data with literature, we qualitatively compared cortical mean R_1 parameters between several cortical regions of interest (ROIs) with known myelination differences. ROIs were defined by probabilistic FreeSurfer (6.0.0, <http://surfer.nmr.mgh.harvard.edu/>) labels for each participant. First, we used the FreeSurfer Brodmann area maps of V1, V2 and MT (V1_exvivo.thresh.label, V2_exvivo.thresh.label and MT_exvivo.thresh.label) (Fischl et al., 2008; Hinds et al., 2008). Second, we defined an angular gyrus label from the FreeSurfer parcellation (Destrieux et al., 2010).

Figure 4c shows systematic R_1 variations with highest values in V1 for each participant, which is in line with Fig. 1(b) in Sereno et al. (2013). **Figure 4–Figure Supplement 1** and **Figure 4–Figure Supplement 2** illustrate the same comparison for R_2^* and PD values. Whereas R_2^* values showed similar results, PD lacked a consistent trend across participants. This might be due to remaining receiver bias in final PD maps, which is challenging to remove especially at high magnetic field strengths (Volz et al., 2012). We therefore did not consider PD parameter maps for the main

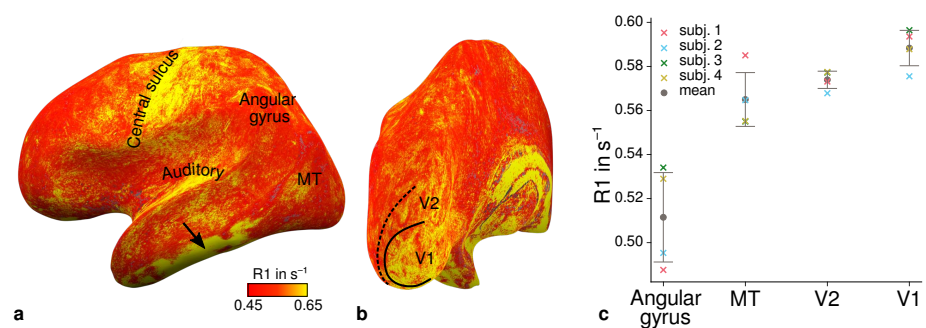


Figure 4. Illustration of quantitative R_1 across cortical areas. Cortical R_1 values are shown at mid-cortical depth of the left hemisphere on an inflated surface from a representative participant (subject 3) in lateral (a) and posterior (b) view. Higher R_1 values can be qualitatively identified in primary motor and sensory areas, which reflects known cortical myeloarchitecture (Flechsig, 1920; Glasser and Van Essen, 2011). The arrow in (a) points to an artifact outside of V2 caused by magnetic field inhomogeneities. Data from all participants can be found in Appendix 1. (c) Mean R_1 values are shown for different cortical regions (angular gyrus, MT, V2, V1) defined by corresponding FreeSurfer labels (Fischl et al., 2008; Hinds et al., 2008; Destrieux et al., 2010) of each participant (similar to Fig. 1(b) in Sereno et al. (2013)). All participants show increased R_1 values in V1. Corresponding source data is given in Figure 4-source data 1. A corresponding illustration for quantitative R_2^* can be found in Figure 4-Figure Supplement 1. Figure 4-Figure Supplement 2 shows mean R_1 (based on a separate whole-brain MP2RAGE acquisition) and PD values across cortical areas. Higher R_1 values in V1 as shown in (c) could be confirmed with the independent R_1 estimate from the MP2RAGE acquisition. Mean across participants is shown in gray. Vertical error bars indicate 1 standard deviation across participants.

Figure 4-Figure supplement 1. Illustration of quantitative R_2^* across cortical areas.

Figure 4-Figure supplement 2. Mean quantitative R_1 (MP2RAGE) and PD values across cortical areas.

Figure 4-source data 1. Source data for mean qMRI values (R_1 , R_2^* , PD) across cortical areas (angular gyrus, MT, V2, V1).

analysis. We also checked cortical profiles of mean parameters in V2 by sampling data on surfaces defined at different cortical depths (see Appendix 2). In all participants, we confirmed the expected decrease of R_1 , R_2^* and $MTVF = 100\% - PD$ (macromolecular tissue volume fraction (Mezer et al., 2013)) values towards the pial surface since all three parameters are sensitive to myelin (Marques et al., 2017; Carey et al., 2018; Kirilina et al., 2020).

Higher myelination of pale stripes

Using R_1 and R_2^* as surrogates for cortical myelination, we tested whether different stripe types are differentially myelinated by comparing R_1 and R_2^* parameter values between stripe types following a similar procedure described in Li et al. (2019). In brief, color-selective thin and disparity-selective thick stripe ROIs were demarcated by applying a z-score threshold to the corresponding functional contrasts (see Appendix 3 for a depiction of ROIs for each participant defined at $z = 1.96$). Mean R_1 and R_2^* from one stripe type were then tested against the mean value within V2 excluding data belonging to the other stripe type (see Materials and methods). This enabled us to indirectly demarcate pale stripes assuming a strict tripartite stripe division of V2. Since the definition of ROIs solely based on z-score thresholds is inevitably arbitrary, we performed the above analysis for several thresholds. Figure 5 shows the pooled R_1 and R_2^* for $z \in \{0, 0.5, \dots, 4.5\}$ across participants. Quantitative parameter values are shown as deviation from the mean within V2 after regressing out variations due to local curvature. For each z-score threshold level, we tested the difference for statistical significance using permutation testing. Figures 5a-b show statistically significant differences of R_1 between thin or thick stripes and mean of V2 excluding the other stripe type, which points towards higher myelin density in pale stripes. These results were confirmed by an independent data set using R_1 values estimated from the MP2RAGE sequence (Marques et al., 2010), which is shown in Figure 5-Figure Supplement 2. The maximum z-score threshold was chosen arbitrarily and was limited by the resulting ROI size. ROI sizes for all threshold levels and participants

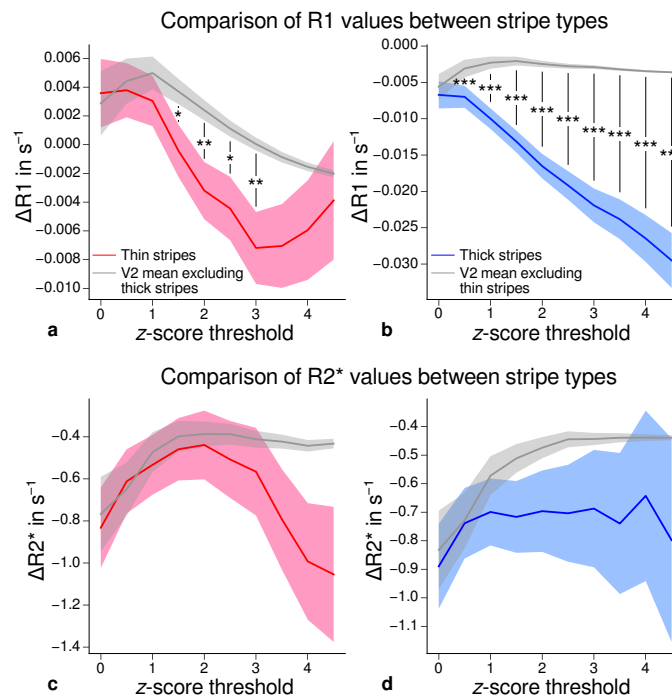


Figure 5. Comparison of quantitative R_1 and R_2^* values between V2 stripe types. Cortical R_1 (a)–(b) and R_2^* (c)–(d) values in thin stripes (red), thick stripes (blue) and whole V2 excluding the other stripe type (gray; and therefore containing contributions from pale stripes) are shown for various z-score threshold levels, which were used to define thin and thick stripe ROIs. Quantitative values are illustrated as deviation from the mean within V2 after removing variance from local curvature. Values were pooled across participants and hemispheres. Differences between data in thin/thick stripes and whole V2 without thick/thin stripes were tested for statistical significance at $z \in \{0, 0.5, \dots, 4.5\}$. Statistical significance was assessed by permutation testing (see Materials and methods). R_1 in both thin and thick stripes is lower than surrounding gray matter, which points towards higher myelination of pale stripes assuming a strict tripartite stripe division in human V2. No effects were found for R_2^* . A supporting analysis confirmed these results when excluding the region around the foveal representation from the analysis. This addresses a potential bias due to missing activation around the foveal representation in functional activation maps, which were used to define stripe ROIs. This is shown in **Figure 5–Figure Supplement 1**. The results for R_1 values were further confirmed using an independent estimate of cortical R_1 based on separately acquired whole-brain MP2RAGE scans, which is shown in **Figure 5–Figure Supplement 2** and **Figure 5–Figure Supplement 3**. Corresponding source data is given in **Figure 5–source data 1**. Statistically significant differences are marked by asterisks, *: $p < 0.05$, **: $p < 0.01$, ***: $p < 0.001$. Shaded areas indicate 1 standard deviation of the generated null distribution used for permutation testing.

Figure 5–Figure supplement 1. Comparison of quantitative R_1 and R_2^* values between V2 stripe types excluding foveal parts.

Figure 5–Figure supplement 2. Comparison of quantitative R_1 values (MP2RAGE) between V2 stripe types.

Figure 5–Figure supplement 3. Comparison of quantitative R_1 values (MP2RAGE) between V2 stripe types excluding foveal parts.

Figure 5–source data 1. Source data for comparisons of R_1 and R_2^* between V2 stripe types.

are illustrated in **Appendix 4**. Note that higher thresholds lead to an expansion of the pale stripe ROI and contamination from other stripe types. In **Figure 5**, shaded areas denote the standard deviation of the generated null distribution used for permutation testing. This illustrates the enlargement of pale stripe ROIs at high threshold levels since larger ROIs lead to less variation across permutations. For an intermediate threshold level of $z = 1.96$ ($p < 0.05$, two-sided), R_1 values in thin and thick stripes differ from pale stripes by 0.005 s^{-1} and 0.014 s^{-1} , respectively, which corresponds to a deviation of around 1–2% assuming a mean longitudinal relaxation rate of 0.58 s^{-1} in V2 (see **Figure 4c**). No statistically significant effects were found for R_2^* as shown in **Figures 5c–d**.

From functional contrasts shown in **Figure 2** (see also **Appendix 1** and **Appendix 3**), it is evident that the region around the foveal representation of V2 largely did not show functional activation. Since we used all portions of V2 in the analysis which were covered by retinotopy measurements, this potentially could have biased our results. To check if missing activation around the foveal representation affected our results, we performed an additional analysis, in which we excluded that foveal region. More precisely, we used the phase responses from retinotopy measurements (eccentricity) and constructed masks to exclude the inner one third of the full phase cycle in individual activation maps (see white lines in **Figure 2** and **Appendix 1**). The excluded eccentricity range was a compromise between masking out a sufficient region around the foveal representation with no functional activation and leaving enough data points for analysis. Comparison of qMRI parameters yielded basically the same results both for MPM and MP2RAGE data shown in **Figure 5–Figure Supplement 1** and **Figure 5–Figure Supplement 3**, respectively. Note that **Figure 5–Figure Supplement 1** also shows significant differences for R_2^* which also points into the direction of higher myelin content of pale stripes without being as conclusive as R_1 .

Discussion

The secondary visual cortex of the primate contains a repeating pale-thin-pale-thick stripe pattern of CO activity. It is known that components of visual information like color, orientation and binocular disparity are largely segregated into separate pathways and processed in different stripe types (*Hubel and Livingstone, 1987; Livingstone and Hubel, 1987*). We robustly mapped color-selective thin and disparity-selective thick stripes in humans using high resolution fMRI. Furthermore, we demonstrated that R_1 parameter maps reflect the expected macroscopic features of cortical myelination in primary motor and sensory areas and across cortical depth. By combining data from fMRI and qMRI, we then showed that locations in V2 have higher R_1 values which neither correspond to functionally defined thin nor thick stripes. Because myelin content is a major contrast mechanism for cortical R_1 (*Stüber et al., 2014*) and we expect a tripartite stripe architecture, we interpret this as an indication of higher myelination of pale stripes than thin and thick stripes in V2.

It is important to note that R_1 is used as a surrogate for cortical myelination and histological findings of myelin content in V2 are not conclusive so far. On the one hand, *Tootell et al. (1983)* showed higher myelination in dark CO stripes in monkey V2 using the Luxol fast blue staining technique and using wet unstained brain tissue sections. On the other hand, *Krubitzer and Kaas (1989)* showed higher myelination in pale CO stripes using the Gallyas staining technique. These conflicting findings were reproduced in a later study by *Horton and Hocking (1997)*. This study also examined myelination in monkey V1 after monocular enucleation. After monocular enucleation, CO activity is down-regulated in layer 4c neurons located in ocular dominance columns (ODCs) of the deafferented eye. This enables the visualization of ODCs of the missing eye, which appear as light columns after CO staining. Monocular enucleation further leads to atrophy of the optic nerve and loss of myelin through the process of Wallerian degeneration. Unexpectedly, *Horton and Hocking (1997)* showed higher myelination in ODCs of the missing eye in wet unstained tissue and using myelin basic protein (MBP) immunohistochemistry. Furthermore, wet unstained sections showed higher myelination in V2 dark CO stripes of normal monkeys. Both findings would be in conflict with each other if a relationship between patches/stripes of CO activity and myelination is assumed. Both classical myelin stains (Luxol fast blue and Gallyas) showed the expected higher myelination in ODCs of the intact eye. However, it is known that Luxol fast blue, which stained CO-rich regions in both V1 and V2, is not only sensitive to myelin content but also shows an affinity to mitochondria where CO is mostly found. This dual character might have influenced the staining results in CO-rich cortical regions of V1 and V2. The Gallyas staining showed the expected higher myelination in ODCs of the normal eye after monocular enucleation and higher myelination in V2 pale stripes. While this does not reveal a complete and conclusive picture, we reproduced the results obtained with the Gallyas staining method using qMRI. Our results were further corroborated by a recent

239 MRI study in macaques (*Li et al., 2019*), which used a similar approach to define thin and thick
240 stripes as in the present study.

241 However, our results do not align with a recent human MRI study by *Dumoulin et al. (2017)*,
242 which found higher myelin density in thick stripes using a T_1 -weighted imaging sequence (MPRAGE)
243 to infer myelination differences. Although the reason for this discrepancy cannot be conclusively
244 determined, two aspects that differ between studies are worth mentioning. First, the functional lo-
245 calization of stripes was different. In *Dumoulin et al. (2017)*, parvo- and magnocellular dominated
246 pathways were targeted by exploiting known differences in the processing of slow and fast tempo-
247 ral frequencies in the visual stimulus, respectively. However, the assignment of parvo- and mag-
248 nocellular streams to particular stripes in V2 is still controversial (*Sincich and Horton, 2005*). Thus,
249 their relation to the tripartite stripe architecture is less clear than for color content and binocular
250 disparity as used in our study (*Hubel and Livingstone, 1987; Tootell et al., 2004; Chen et al., 2008*).
251 Second, in contrast to our study, myelin density was inferred from weighted MR images, which are
252 known to be more affected by technical biases.

253 The observed differences of myelin density between stripe types were based on R_1 estimates
254 using the MPM protocol in the main analysis. We further confirmed these results with an inde-
255 pendent data set using R_1 estimates from the MP2RAGE acquisition (see *Figure 5–Figure Supple-*
256 *ment 2*). This further demonstrates the generalizability of our results across acquisition methods.
257 Nevertheless, although previous studies show the validity of R_1 as a myelin marker in this type of
258 healthy participant group with no known pathologies (*Stüber et al., 2014; Leuze et al., 2017*) and
259 differences were found with two different R_1 mapping methods, we cannot fully exclude that some
260 unknown factors may systematically bias the indirect measures.

261 We did not find any significant differences of R_2^* between stripe types in the main analysis as
262 shown in *Figures 5c–d*. Whereas R_1 in the normal cortex is largely influenced by myelination lev-
263 els, R_2^* is sensitive to paramagnetic iron and diamagnetic myelin (*Stüber et al., 2014; Kirilina et al.,*
264 *2020*). Other factors like vasculature and the orientation of the cortex to the static magnetic field
265 of the MR system have an influence on R_2^* , which might have obscured the underlying dependency
266 on myelin content (*Cohen-Adad et al., 2012*). The dark appearance of thin and thick stripes after
267 CO staining is a marker for increased oxidative metabolism compared to pale stripes. This favors
268 the hypothesis of richer vascularization in thin and thick stripes, which potentially could counter-
269 act reductions in R_2^* due to lower myelination. Indeed, higher vessel densities were found in blobs
270 (another CO rich structure in V1) and stripes of squirrel and macaque monkeys (*Zheng et al., 1991;*
271 *Keller et al., 2011*). However, this was later disputed by another study, which showed no differ-
272 ences in vascular supply between blobs and inter-blobs in V1 (*Adams et al., 2015*).

273 *Figures 5a–b* show that R_1 differences are in the range of around 1–2%. This is smaller but
274 comparable to R_1 differences between cortical areas, which are in the range of a few percent at
275 7 T (*Marques et al., 2017*). The required high spatial resolution and the measured small effect sizes
276 limited the contrast-to-noise ratio in our measurements. This most probably hindered a direct
277 visualization of stripes at the voxel level in R_1 maps as illustrated in *Figure 4* (see *Appendix 1* for
278 visualization of cortical R_1 and R_2^* in V1 and V2 for each participant) and required pooling of data
279 within stripe types defined by fMRI followed by a statistical analysis. The coefficient of variation
280 in R_1 maps was 11.3 ± 0.7 (mean \pm standard deviation across participants) in V2. However, when
281 data were pooled across participants, we could robustly detect significant differences between
282 stripe types. We believe that the usage of qMRI parameters at this spatial scale in living humans is
283 favorable as these parameters are less affected by technical biases and are in physical units and
284 thus more accessible for biophysical modeling, which will facilitate studies of tissue microstructure
285 with higher specificity.

286 The following considerations are related to the definition of the stripe ROIs. We could only
287 functionally delineate thin and thick stripes but not pale stripes. Thus, the definition of pale stripes
288 was indirect and relied on the assumption of a tripartite architecture.

289 In the analysis, the whole stimulated part of V2, which could be defined by retinotopy, was con-

sidered. However, the paradigms used for localization of color and disparity stripes did not show pronounced activation at the representation of the central fovea (see **Figure 2** and **Appendix 1**). First, the color stimulus with red/blue gratings (see **Figure 1a**) is expected to have a different effect in the central fovea than in parafoveal regions due to the macula lutea (yellow pigmented spot of the retina) and absence of blue cones in the central fovea, which might have hindered the detection of color stripes there (**Nasr and Tootell, 2018**). Second, missing activation at the representation of the central fovea for the disparity stimulus could be due to eccentricity dependence of disparity tuning. Other possible explanations might be a smaller fusion limit in central fovea or a global attention mechanism, which might preferentially activate peripheral representations (**Tsao et al., 2003**). Using conventional fMRI with lower resolution, **Tsao et al. (2003)** found an overall similar eccentricity-dependent activation pattern for a stimulation with the same maximal disparity ($\pm 0.22^\circ$). To address the possibility that our results were biased by eccentricity, we repeated the analysis shown in **Figure 5** but excluded the foveal representation of V2, which yielded basically the same results (see **Figure 5–Figure Supplement 1** and **Figure 5–Figure Supplement 3**).

Figure 2 shows that activation maps for color-selective thin and disparity-selective thick stripes partly overlap, which might complicate the definition of separate ROIs for thin and thick stripes. It should be kept in mind that spatial overlap is expected to some degree and mainly driven by the limiting physiological point spread function of the measured blood oxygenation level-dependent (BOLD) signal in fMRI (**Polimeni et al., 2010; Chaimow et al., 2018**). This did not interfere with our analysis, since all data points with overlapping activation were excluded in ROI definitions. On the one hand, it is expected that the degree of overlap depends on the chosen z -score threshold level (**Nasr et al., 2016**) assuming higher thresholds to increase the probability of solely sampling within one stripe type. On the other hand, high z -score thresholds bear the risk to predominantly sample from large veins (**Boxerman et al., 1995**), which degrades the accuracy of the ROI due to blurring and displacement of the functional signal (**Olman et al., 2007**). We based the ROI definition on activation maps from differential contrasts between two experimental conditions as illustrated in **Figures 1a–b**, which are known to be less affected by unspecific macrovascular contributions and draining veins. Furthermore, we would have expected any venous bias to be reflected in R_2^* maps (**Yacoub et al., 2001; Peters et al., 2007**), e.g., by uneven sampling of veins between stripe types which is not the case. For these reasons, we conclude that venous bias did not drive our results.

The regular compartmentalization of V2 into distinct stripe types leads to the expectation of specific coverage of cortical area by thin, thick and pale stripes. For example, it is expected that thick stripes are slightly larger than thin stripes as their name suggests, and that pale stripes cover around 50% of V2 (**Shipp and Zeki, 1985; Tootell and Hamilton, 1989**). Using fMRI for ROI definitions, the coverage depends on the chosen z -score threshold as stated further above. For $z = 1.96$ ($p < 0.05$, two-sided), the relative V2 coverage of non-overlapping portions of thin and thick stripes is $14.1\% \pm 3.4\%$ and $24.5\% \pm 6.9\%$ (mean \pm standard deviation across participants and hemispheres; see **Appendix 4** for absolute coverage of stripe ROIs at different threshold levels). This sums up to a pale stripes coverage of 61.4%.

An alternative hypothesis for different myelination of V2 stripes is that only the borders between pale and dark CO stripes are more strongly myelinated. **Pistorio et al. (2006)** showed in monkey V1 that the edges of blobs are more myelinated rather than blobs or inter-blob regions itself using a modified Gallyas stain. We note that the used methods in the present study neither allow a functional determination of a definite border between stripe types nor are able to spatially resolve the border region between stripes. Therefore, we cannot rule out this alternative hypothesis. However, to our knowledge, no previous study showed evidence for higher myelination of stripe borders in V2.

Measurements with high resolution are vulnerable to head movements during image acquisition, especially for the long anatomical scans. Therefore, we used an optical tracking system to prospectively correct head movements during anatomical scans (see **Materials and methods**).

With this system, head movements could be robustly detected and corrected for at the length scale of movements induced by respiration and heart beat. Exemplary motion traces are shown in **Appendix 5**.

The packing density of myelinated fibers in the cerebral cortex varies with cortical depth (*Flechsig, 1920; Glasser and Van Essen, 2011*) and is also dependent on the cortical folding (*Smart and McSherry, 1986*). The correct and consistent sampling of data within cortex is therefore critical for our study. We used the equi-volume model to sample at mid-cortical depth. This model has been shown to be less affected by curvature biases than other models (e.g. equi-distant sampling) (*Waehnert et al., 2014*). The validity of the depth model also depends on accurate cortex segmentation. We visually inspected the cortical segmentation carefully in each participant (see **Figure 6**). Remaining curvature contributions were regressed out as in other studies (*Glasser and Van Essen, 2011; Sereno et al., 2013; Dumoulin et al., 2017*).

Our findings point towards higher myelination of pale stripes which exhibit lower oxidative metabolic activity according to staining with CO. V2 receives most of its input from V1 and the pulvinar (*Tootell et al., 1983; Sincich and Horton, 2005*). Pulvinar projections, however, only terminate in layer 3 and 5, whereas the alternating myelin pattern is most obvious in layer 4, which receives input almost exclusively from V1 (*Tootell et al., 1983; Sincich and Horton, 2005*). An anterograde tracer study in macaques by *Sincich and Horton (2002)* showed that [^3H]proline injections into V1 preferentially targeted V2 pale stripes. Although we cannot exclude that systematic differences in terminal axonal arborization between stripe types could explain this observation, we speculate that the results of that study correspond to higher axonal density of V1 to V2 projections in layer 4 of pale stripes. This would lead to higher myelination in pale stripes, which is in line with our measurements.

By comparing myelin-sensitive longitudinal relaxation rates (R_1) between stripe types in V2 defined by high resolution fMRI, we found significant quantitative evidence for myelination differences in living humans at the level of columnar systems. This shows the feasibility to use high resolution quantitative R_1 values to study cortical myelination, which is known to be less biased by technical artifacts and are thus better comparable amongst participants and scanner sites. Moreover, it is well known that the myelination of cortical areas and structures affects their functional properties, i.e. the propagation of action potentials (*Sanders and Whitteridge, 1946*), and correlates with postnatal development (*Glasser and Van Essen, 2011*). Therefore, the estimation of myelin content of specific structures in the human brain *in vivo* may increase our knowledge about its relationship to functional properties of the brain in particular and the ontogeny of the human brain in general. Our study shows that with the use of qMRI at ultra-high magnetic field strength, this is possible at the spatial scale of thin, thick and pale stripes. We therefore believe that the current study shows the applicability of qMRI to further advance our knowledge of cortical myelination and tissue microstructure for exploration of structure-function relationships in the living human brain at mesoscopic scale.

Materials and methods

Participants

Four healthy participants (1 female, age = 27.50 ± 4.39 , mean \pm standard deviation) gave written informed consent to participate in this study. The study was approved by the local ethics committee of the University of Leipzig. All participants had normal or corrected-to-normal visual acuity, normal color vision (based on Ishihara and Farnsworth D15 tests) and normal stereoscopic vision (based on Lang I test).

General procedures

Each participant was scanned multiple times on different days in an ultra-high field MR scanner (7 T). The first session was used to acquire a high resolution anatomical reference scan and retino-

389 topy data (*Sereno et al., 1995; Engel et al., 1997*) to functionally locate area V2 in each individual.
390 Additionally, a baseline fMRI scan without task was acquired to aid between-session registrations
391 (see below). Color-selective thin stripes (two sessions) and disparity-selective thick stripes (two
392 sessions) were mapped in subsequent scanning sessions. For two participants, we had time to ac-
393 quire a third thin and thick stripe session, respectively. However, we restricted the data analysis to
394 the use of data from two sessions for consistency. Furthermore, high resolution anatomical scans
395 (one session) were acquired in a separate scanning session to estimate whole-brain quantitative
396 MR relaxation parameters. A subset of acquired quantitative MR and fMRI retinotopy data was al-
397 ready used in other experiments (*Attar et al., 2020; McColgan et al., 2021*) but was independently
398 processed for this study.

399 Visual stimulation

400 For the presentation of visual stimuli, we used an LCD projector (Sanyo PLC-XT20L with custom-
401 built focusing objective, refresh rate: 60 Hz, pixel resolution: 1024 × 768), which was positioned
402 inside the magnet room. To suppress interference with the MR scanner, the projector was placed
403 inside a custom-built Faraday cage. Stimuli were projected onto a rear-projection screen mounted
404 above the participants' chest inside the bore and viewed through a mirror attached to the head coil.
405 This setup allowed the visual stimulation of around 22° × 13° visual angle. Black felt was put around
406 the screen and all lights were turned off during experiments to mitigate scattered light reaching
407 the participants' eyes. Experimental stimuli were written in GNU Octave (4.0.0, <http://www.gnu.org/software/octave/>) using the Psychophysics Toolbox (*Brainard, 1997; Pelli, 1997; Kleiner et al., 2007*) (3.0.14). A block design consisting of two experimental conditions was used for mapping
410 color-selective thin stripes and disparity-selective thick stripes in V2, which was reported in detail
411 previously (*Nasr et al., 2016*) and was only changed marginally for this experiment.

412 *Experiment 1: Color-selective thin stripes* Stimuli consisted of isoluminant sinusoidal color-varying
413 (red/blue) or luminance-varying (black/white) gratings as illustrated in *Figure 1a*. Gratings moved
414 perpendicular to one of four orientations (0°, 45°, 90°, 135°) with direction reversals every 5 s and
415 a drift velocity of 5°/s. Orientations were pseudorandomized between blocks. A low spatial fre-
416 quency (0.4 cpd) was used to mitigate linear chromatic aberration at color borders and exploit
417 the relatively higher selectivity to color relative to luminance at this spatial scale (*Tootell and Nasr, 2017*). We point out that the appropriateness to use red and blue colors to stimulate color-selective
418 thin stripes has already been demonstrated for macaques (*Tootell et al., 2004; Li et al., 2019*) and
419 humans (*Nasr et al., 2016*). In one run, color and luminance stimuli were both shown four times
420 in separate blocks with a length of 30 s. Each run started and ended with 15 s of uniform gray. Ten
421 runs were conducted in one session. During runs, participants were asked to fix their gaze on a
422 central point (0.1° × 0.1°) and respond on a keypad when the fixation point changed its color (light
423 green, dark green). To measure functional activation related to color, it is important to control for
424 luminance variations across stimuli. Furthermore, isoluminance points between colors are known
425 to change with eccentricity (*Livingstone and Hubel, 1987; Bilodeau and Faubert, 1997*). We used
426 a flicker photometry (*Ives, 1907; Bone and Landrum, 2004*) paradigm to get isoluminance ratios
427 between stimuli for each participant. In brief, the luminance of blue was set to 17.3 cd/m² (cf. *Li et al. (2019)*). Before scanning, each participant performed a behavioral task inside the scanner in
428 which they viewed a uniform blue flickering in temporal counter-phase with gray (30 Hz). Partici-
429 pants were asked to adjust the luminance of gray so that the perceived flickering was minimized
430 using a keypad. This procedure was repeated to adjust the luminance for red and conducted at
431 three different eccentricities (0°-1.7°, 1.7°-4.1°, 4.1°-8.3°). As expected, isoluminance ratios changed
432 with eccentricity, which is illustrated in *Appendix 6*.

433 *Experiment 2: Disparity-selective thick stripes* Stimuli consisted of two overlaid random dot stere-
434 ograms (RDSs) (*Julesz, 1971*) made of red and green dots on a black background (dot size: 0.1°, dot
435 density: ~ 17%), respectively. Participants viewed stimuli through custom-built anaglyph spectacles
436 using Kodak Wratten filters No. 25 (red) and 44A (cyan). In one condition, red and green RDSs per-
437
438

439 formed a horizontal sinusoidal movement with temporal frequency of 0.25 Hz. Phases of red and
440 green dots were 180° out of phase and initialized to create the perception of a 8 × 6 checkerboard
441 moving periodically in depth (away and towards the participant), which is schematically illustrated
442 in **Figure 1b**. Maximal disparity was set to $\pm 0.22^\circ$ (cf. *Tsao et al. (2003)*). In the other condition, static
443 dots were presented, which were perceived as a plane at depth of the fixation point. In one run,
444 both conditions were shown four times in separate blocks with a length of 30 s. Each run started
445 and ended with 15 s of black background. 10 runs were conducted in one session. During runs,
446 participants were asked to fix their gaze on a central point ($0.2^\circ \times 0.2^\circ$) and respond on a keypad
447 when the fixation point changed its form (square, circle). The luminance of red and green dots
448 was kept low to decrease cross-talk between eyes (red dots through red filter: 3.1 cd/m^2 , red dots
449 through cyan filter: 0.07 cd/m^2 , green dots through green filter: 5.7 cd/m^2 , green dots through cyan
450 filter: 0.09 cd/m^2). Luminance of green dots was doubled to approximately excite the same amount
451 of cone photoreceptors with both colors (*Dobkins et al., 2000*).

452 **Retinotopic mapping** A standard phase-encoded paradigm (*Sereno et al., 1995; Engel et al., 1997*)
453 was used to locate the stimulated portion of V2. Stimuli consisted of a flickering (4 Hz) black-and-
454 white radial checkerboard restricted to a clockwise/anticlockwise rotating wedge (angle: 30° , pe-
455 riod: 64 s) or expanding/contracting ring (period: 32 s) presented in separate runs to reveal polar
456 angle and eccentricity maps, respectively. 8.25 cycles were shown in each run. Each run started
457 and ended with 12 s of uniform gray background. Mean luminance was set to 44 cd/m^2 . Partici-
458 pants were asked to fix their gaze on a central point during visual stimulation. No explicit task was
459 given.

460 Imaging

461 All experiments were performed on a 7 T whole-body MR scanner (MAGNETOM 7 T, Siemens Health-
462 ineers, Erlangen, Germany) equipped with SC72 body gradients (maximum gradient strength: 70 mT/m ;
463 maximum slew rate: 200 mT/m/s). For radio frequency (RF) signal transmission and reception, a
464 single-channel transmit/32-channel receive head coil (Nova Medical, Wilmington, USA) was used.
465 At the beginning of each scanning session, a low resolution transmit field map was acquired to
466 optimize the transmit voltage over the occipital lobe.

467 Functional data was acquired with a 2D single-shot gradient-echo (GE) echo-planar imaging
468 (EPI) sequence (*Feinberg et al., 2010; Moeller et al., 2010*). A coronal-oblique slab was imaged,
469 which covered all stimulated portions of V2. The following parameters were used for the mapping
470 of color-selective thin stripes, disparity-selective thick stripes and the baseline fMRI scan without
471 task: nominal voxel size = 0.8 mm isotropic, repetition time (TR) = 3000 ms , echo time (TE) = 24 ms ,
472 excitation flip angle (FA) = 77° , field of view (FOV) = $148 \times 148 \text{ mm}^2$, 50 slices, readout bandwidth (rBW)
473 = 1182 Hz/px , echo spacing = 1 ms , partial Fourier = $6/8$ and generalized autocalibrating partially
474 parallel acquisition (GRAPPA) (*Griswold et al., 2002*) = 3. A slightly modified protocol was used for
475 retinotopy measurements with the following parameter changes: voxel size = 1.0 mm isotropic, TR
476 = 2000 ms , TE = 21 ms , FA = 68° , 40 slices and rBW = 1164 Hz/px .

477 MR relaxation parameters (R_1 , R_2^* , PD) were measured with a multi-echo variable flip angle
478 (VFA) protocol for multi-parameter mapping (MPM) (*Vaculčíková et al., 2022*). The protocol was
479 adapted for whole-brain coverage with 0.5 mm isotropic voxel size and consisted of two multi-echo
480 3D fast low angle shot (FLASH) scans with T_1 - and PD -weighting (T_1w , PDw) plus maps of B_1^+ and
481 B_0 . For T_1w and PDw , the following parameters were used: TR = 25 ms , TE = $2.8\text{--}16.1 \text{ ms}$ (6 equidis-
482 tant echoes with bipolar readout), FA(PDw/T_1w) = $5^\circ/24^\circ$, FOV = $248 \times 217 \times 176 \text{ mm}^3$ (read × phase
483 × partition), rBW = 420 Hz/px and GRAPPA = 2×2 in both phase-encoding directions. Head move-
484 ments during the scan were corrected prospectively using an optical tracking system (KinetiCor,
485 USA). For motion detection, a mouth guard assembly with attached markers was manufactured
486 for each participant by the Department of Cariology, Endodontology and Periodontology of the
487 University of Leipzig Medical Center (*Vaculčíková et al., 2022*). No prospective motion correction
488 was used during functional scans because the camera system and the projection screen did not fit

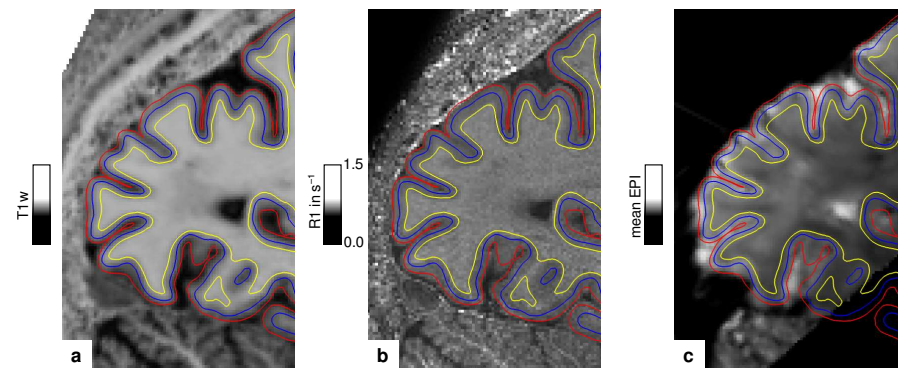


Figure 6. Illustration of segmentation and registration quality. (a) Posterior part of the MP2RAGE UNI image in sagittal orientation of one representative participant (subject 3), which was used for segmentation of the cerebral cortex. Overlaid contour lines show the reconstructed white matter/gray matter boundary surface (yellow), the pial boundary surface (red) and a surface at mid-cortical depth (blue). The computed R_1 map from the MPM acquisition and the temporal mean from a representative fMRI session are shown in (b) and (c), respectively. Overlaid contour lines are identical to (a) to visualize the segmentation and registration quality. Other illustrations of registration quality are provided for each participant in video format, see [Figure 6–video 1](#), [Figure 6–video 2](#), [Figure 6–video 3](#), and [Figure 6–video 4](#). A more quantitative analysis of the achieved registration performance is shown in [Appendix 7](#).

Figure 6–video 1. Illustration of registration quality (subject 1). The video shows the posterior part of the R_1 map from the MPM protocol in temporal counter-phase with registered data from other measurements (temporal mean BOLD acquisitions from color-selective thin stripe measurement, disparity-selective thick stripe measurement, baseline measurement without task, retinotopy measurement and the R_1 map from the MP2RAGE acquisition). 20 slices are shown around the midsagittal plane. A red grid is overlaid for visual guidance.

Figure 6–video 2. Illustration of registration quality (subject 2). Other details as in [Figure 6–video 1](#).

Figure 6–video 3. Illustration of registration quality (subject 3). Other details as in [Figure 6–video 1](#).

Figure 6–video 4. Illustration of registration quality (subject 4). Other details as in [Figure 6–video 1](#).

together in the bore. Note that functional scans are also less sensitive to motion due to the short acquisition time per volume.

For the correction of RF transmit field (B_1^+) inhomogeneities in relaxation parameter maps (R_1 , PD), we followed the procedure detailed in ([Lutti et al., 2010, 2012](#)), acquiring spin-echo (SE) and stimulated echo (STE) images with a 3D EPI readout. The total scanning time of the MPM protocol was approximately 45 minutes.

For cortex segmentation and image registration, a whole-brain anatomy was acquired using a 3D T1-weighted MP2RAGE sequence ([Marques et al., 2010](#)) with the following parameters: voxel size = 0.7 mm isotropic, TR = 5000 ms, TE = 2.45 ms, inversion times (T11/T12) = 900 ms/2750 ms with FA = 5°/3° for T1w and PDw images, respectively, FOV = 224×224×168 mm³ (read × phase × partition), rBW = 250 Hz/px, partial Fourier = 6/8 and GRAPPA = 2 (primary phase-encoding direction; outer loop). From both inversion times, a uniform T_1 -weighted image (UNI) and a T_1 -map were created in the online image reconstruction on the scanner.

Data analysis

Functional time series from color-selective and disparity-selective stripe mapping sessions were corrected for within-run and between-run motion using SPM12 (v6906, <https://www.fil.ion.ucl.ac.uk/spm/>) with Matlab R2019b (MathWorks, Natick, USA). Motion corrected time-series were high-pass filtered (cutoff frequency: 1/270 Hz) and voxel-wise statistical analyses were performed for each session using a general linear model (GLM) as implemented in SPM12 with both experimental conditions as regressors.

For retinotopy measurements, time series were first converted to percent signal change by division by their temporal mean. Furthermore, slice timing correction was added before motion

correction by voxel-wise temporal interpolation to a common time grid using *3drefit* from Analysis of Function NeuroImages software (Cox, 1996) (AFNI, 19.1.05). Motion corrected time-series were high-pass filtered (cutoff frequency: $1/(3 \times \text{stimulus cycle period})$ Hz) and data from the first quarter of the stimulus cycle was discarded from further processing. A voxel-wise Fourier transform was computed and real and imaginary parts at stimulus frequency were averaged from runs with opposite stimulus direction to compensate for the hemodynamic lag. A phase map from averaged polar angle real and imaginary parts was computed to delineate the borders of V2.

For one participant (subject 3), we acquired a second set of retinotopy data in a separate session due to low functional responses in comparison to other participants. Motion corrected time series from the second session were registered nonlinearly to the first session using ANTs, 2.3.1, <http://stnava.github.io/ANTs/>). Time series from both session were then averaged before further processing.

Quantitative parameter maps (R_1 , R_2^* , PD) were computed using the hMRI toolbox (Tabelow et al., 2019) (0.2.2, <http://hmri.info>) implemented in SPM12 (v7487). In brief, T1w and PDw images from the MPM protocol were averaged across echoes and used to compute a registration between both contrasts using SPM12. All available echoes from both contrasts were then used to compute an R_2^* map by ordinary least squares regression using the ESTATICS model (Weiskopf et al., 2014). For the calculation of R_1 and PD maps, the extrapolation of T1w and PD to $TE = 0$ (to remove any R_2^* -weighting bias from resulting maps) was fit to an approximation of the Ernst equation for short-TR dual flip angle measurements using the FLASH signal (Helms et al., 2008; Edwards et al., 2021). The $B1^+$ field map was corrected for off-resonance effects using the acquired $B0$ map. A flip angle map was computed from the resulting $B1^+$ map to correct the apparent flip angles for inhomogeneities of the RF transmit field in the fitting procedure. For PD map calculations, the resulting map was corrected for the receiver coil sensitivity profile using the adapted data-driven UNICORT method, which applies the bias field correction implemented in the segmentation module of SPM12 (Weiskopf et al., 2011), and calibrated such that the mean PD over a white matter mask $PD(WM) = 69$ percent units (pu) (Tofts, 2018). Final maps (R_1 , R_2^* , PD) were corrected for spatial gradient nonlinearity distortions using the gradunwarp toolbox (Glasser et al., 2013) (1.0.2, <https://github.com/Washington-University/gradunwarp>) and spherical harmonic coefficients provided by the manufacturer.

Cortex segmentation was based on the MP2RAGE UNI image. First, the UNI image was corrected for gradient nonlinearities with the gradunwarp toolbox and remaining bias fields using SPM12. The resulting image was input to the *recon-all* pipeline in FreeSurfer (Dale et al., 1999; Fischl et al., 1999) (6.0.0, <http://surfer.nmr.mgh.harvard.edu/>) with the *hires* flag to segment at the original voxel resolution (Zaretskaya et al., 2018). The brain mask used during segmentation was computed from the second inversion image of the MP2RAGE using SPM12 and was defined by excluding all voxels that exceeded the tissue class threshold of 10% in non-WM and non-GM tissue classes. Final gray matter/white matter and pial boundary surfaces were corrected manually. Extra care was applied to correct the pial surface around the sagittal sinus. The resulting gray matter/white matter surface was shifted 0.5 mm inwards to counteract a potential segmentation bias using FreeSurfer with MP2RAGE (Fujimoto et al., 2014). Final surface meshes were slightly smoothed and upsampled to an average edge length of around 0.3 mm. A surface mesh at mid-cortical depth was computed using the equi-volume model (Waehnert et al., 2014; Wagstyl et al., 2018).

All images were registered to the space of the qMRI maps. For the registration of MP2RAGE and MPM, we used R_1 maps from both acquisitions. Just for the purpose of registration, both images were corrected for potentially remaining bias fields (SPM12) and a brain mask was applied. Images were then transformed into the same space via the scanner coordinate system and a rigid registration was computed using *flirt* (Jenkinson et al., 2002) (6.0) from the FMRIB Software Library (5.0.11; <https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/>). A nonlinear transformation was computed to register activation maps and qMRI data in several steps. First, the baseline fMRI scan from the the first session was registered to the MP2RAGE using the Symmetric Normalization (SyN) algorithm (Avants et al.,

2008) from Advanced Normalization Tools (ANTs, 2.3.1, <http://stnava.github.io/ANTs/>). A nonlinear registration was chosen to account for geometric distortions in functional images resulting from the low bandwidth in phase-encoding direction. Since both images were acquired in the same session, the registration between modalities was robust. Both images were prepared by removing any bias fields (Tustison et al., 2010) and applying a brain mask. Functional data from other sessions were then registered nonlinearly to the baseline EPI using the same procedure. The final transform was computed by concatenating transforms from all steps (EPI → baseline EPI → MP2RAGE → MPM). An exemplary illustration of the registration and segmentation quality can be seen in Figure 6. A more detailed analysis on achieved registration accuracy is given in Appendix 7.

Generated surfaces from cortex segmentation were transformed to MPM space using linear interpolation. For data sampling, images were transformed to MPM space using linear interpolation before sampling onto the surface mesh using nearest neighbor interpolation.

Reliability analysis of fMRI sessions The consistency of activation maps was analyzed by computing the vertex-wise correlation of activities within V2 between sessions acquired on different days. Spearman's rank correlation coefficient r was computed. A p -value was determined by permutation testing. A null distribution was created by computing correlation coefficients between data from the first session and spatially shuffled data from the second session n times ($n = 10,000$). We paid attention to preserve the spatial autocorrelation in spatially shuffled maps using the BrainSMASH package (Burt et al., 2020) (0.10.0) to consider the non-independence of data from neighboring locations. The p -value was then defined as the fraction of the null distribution which is greater or smaller than r . We corrected the estimate of the p -value for the variability resulting from the finite sample size of the null distribution. The variability was described by the variance of the binomial distribution $\sigma^2 = np(1 - p)$. Here, we used an upper bound of 3σ , which was added to the number of samples exceeding the test statistics (Burt et al., 2020). A p -value of < 0.05 was considered as statistically significant.

Quantitative comparison of qMRI parameters between stripe types We tested the hypothesis that pale stripes are differentially myelinated in comparison to color-selective thin and disparity-selective thick stripes. Activation maps from color and disparity stripe measurements were averaged across sessions, respectively. For participants with more than two acquired sessions, we chose to use the two sessions with highest between-session correlation of activities within V2. Color and disparity stripes were demarcated by thresholding activation maps at a selected threshold level $z \in \{0, 0.5, \dots, 4.5\}$. Data points that did not exclusively belong to one stripe type were discarded. Similar to a procedure described in Li et al. (2019), mean qMRI parameter values across participants sampled in color/disparity stripes were tested against the mean throughout V2 excluding values sampled in disparity/color stripes to correct for effects from the other stripe type. This allowed us to indirectly infer effects in pale stripes assuming a tripartite stripe division of V2. For each participant, we subtracted the mean within V2 from qMRI parameter values to account for inter-subject variability (e.g., see variability between participants in Appendix 2). We considered the covariance of qMRI parameter values with local curvature of the cortical sheet (Sereno et al., 2013) by regressing out any linear curvature dependencies. Note that partial volume effects induced by cortical folding are themselves linear, which justifies the use of linear regression. The mean was computed across participants and statistical significance was determined by permutation testing. A null distribution was created by repeating the same procedure n times ($n = 10,000$) with ROIs generated from spatially shuffled activation maps. The spatial autocorrelation in shuffled maps was preserved using the BrainSMASH package (Burt et al., 2020) and the p -value was computed as stated further above for the fMRI reliability analysis.

Acknowledgments

The research leading to these results has received funding from the European Research Council under the European Union's Seventh Framework Program (FP7/2007-2013) / ERC grant agreement n° 616905. Nikolaus Weiskopf has received funding from the European Union's Horizon

2020 research and innovation program under the grant agreement n° 681094 and from the BMBF (01EW1711A & B) in the framework of ERA-NET NEURON. We thank the University of Minnesota Center for Magnetic Resonance Research for the provision of the multiband EPI sequence software. We thank Roland Mueller for the help with building the anaglyph spectacles.

Author Contributions

Daniel Haenelt, Conceptualization, Methodology, Software, Formal analysis, Investigation, Data curation, Writing - original draft preparation, Writing - review & editing, Visualization; Robert Trampel, Investigation, Writing - review & editing, Supervision; Shahin Nasr, Jonathan R. Polimeni, Roger B. H. Tootell, Martin I. Sereno, Methodology, Software, Writing - review & editing; Kerrin J. Pine, Methodology, Investigation, Writing - review & editing; Luke J. Edwards, Methodology, Writing - review & editing; Saskia Helbling, Formal analysis, Writing - review & editing; Nikolaus Weiskopf, Conceptualization, Resources, Writing - review & editing, Supervision, Project administration, Funding acquisition

Competing Interests

The authors declare the following competing interests: The Max Planck Institute for Human Cognitive and Brain Sciences has an institutional research agreement with Siemens Healthcare. Nikolaus Weiskopf holds a patent on MRI data acquisition during spoiler gradients (United States Patent 10,401,453). Nikolaus Weiskopf was a speaker at an event organized by Siemens Healthcare and was reimbursed for the travel expenses.

Data Availability

Anonymized and defaced MRI data used in the present study are openly accessible at: <https://osf.io/624cz/>.

References

- Adams DL, Piserchia V, Economides JR, Horton JC. Vascular supply of the cerebral cortex is specialized for cell layers but not columns. *Cereb Cortex*. 2015; 25:3673–3681. doi: <https://doi.org/10.1093/cercor/bhu221>.
- Adams DL, Sincich LC, Horton JC. Complete pattern of ocular dominance columns in human primary visual cortex. *J Neurosci*. 2007; 27:10391–10403. doi: <https://doi.org/10.1523/JNEUROSCI.2923-07.2007>.
- Attar F, Kirilina E, Haenelt D, Pine KJ, Trampel R, Edwards LJ, Weiskopf N. Mapping short association fibers in the early cortical visual processing stream using in vivo diffusion tractography. *Cereb Cortex*. 2020; 30:4496–4514. doi: <https://doi.org/10.1093/cercor/bhaa049>.
- Avants BB, Epstein CL, Grossman M, Gee JC. Symmetric diffeomorphic image registration with cross-correlation: evaluating automated labeling of elderly and neurodegenerative brain. *Med Image Anal*. 2008; 12:26–41. doi: <https://doi.org/10.1016/j.media.2007.06.004>.
- Bilodeau L, Faubert J. Isoluminance and chromatic motion perception throughout the visual field. *Vis Res*. 1997; 37:2073–2081. doi: [https://doi.org/10.1016/s0042-6989\(97\)00012-6](https://doi.org/10.1016/s0042-6989(97)00012-6).
- Bone RA, Landrum JT. Heterochromatic flicker photometry. *Arch Biochem Biophys*. 2004; 430:137–142. doi: <https://doi.org/10.1016/j.abb.2004.04.003>.
- Boxerman JL, Hamberg LM, Rosen BR, Weisskoff RM. MR contrast due to intravascular magnetic susceptibility perturbations. *Magn Reson Med*. 1995; 34:555–566. doi: <https://doi.org/10.1002/mrm.1910340412>.
- Brainard DH. The psychophysics toolbox. *Spat Vis*. 1997; 10:433–436. doi: <https://doi.org/10.1163/156856897X00357>.
- Burt JB, Helmer M, Shinn M, Anticevic A, Murray JD. Generative modeling of brain maps with spatial autocorrelation. *Neuroimage*. 2020; 220:117038. doi: <https://doi.org/10.1016/j.neuroimage.2020.117038>.

- 655 **Carey D**, Caprini F, Allen M, Lutti A, Weiskopf N, Rees G, Callaghan MF, Dick F. Quantitative MRI provides markers
656 of intra-, inter-regional, and age-related differences in young adult cortical microstructure. *Neuroimage*.
657 2018; 182:429–440. doi: <https://doi.org/10.1016/j.neuroimage.2017.11.066>.
- 658 **Chaimow D**, Yacoub E, Uğurbil K, Shmuel A. Spatial specificity of the functional MRI blood
659 oxygenation response relative to neuronal activity. *Neuroimage*. 2018; 164:32–47. doi:
660 <https://doi.org/10.1016/j.neuroimage.2017.08.077>.
- 661 **Chen G**, Lu HD, Roe AW. A map for horizontal disparity in monkey V2. *Neuron*. 2008; 58:442–450. doi:
662 <https://doi.org/10.1016/j.neuron.2008.02.032>.
- 663 **Cohen-Adad J**, Polimeni JR, Helmer KG, Benner T, McNab JA, Wald LL, Rosen BR, Mainero C. T_2^* mapping and B_0
664 orientation-dependence at 7 T reveal cyto- and myeloarchitecture organization of the human cortex. *Neu-*
665 *roimage*. 2012; 60:1006–1014. doi: <https://doi.org/10.1016/j.neuroimage.2012.01.053>.
- 666 **Cox RW**. AFNI: software for analysis and visualization of functional magnetic resonance neuroimages. *Comput*
667 *Biomed Res*. 1996; 29:162–173. doi: <https://doi.org/10.1006/cbmr.1996.0014>.
- 668 **Dale AM**, Fischl B, Sereno MI. Cortical surface-based analysis: I. segmentation and surface reconstruction.
669 *Neuroimage*. 1999; 9:179–194. doi: <https://doi.org/10.1006/nimg.1998.0395>.
- 670 **Destrieux C**, Fischl B, Dale A, Haglren E. Automatic parcellation of human cortical gyri
671 and sulci using standard anatomical nomenclature. *Neuroimage*. 2010; 53:1–15. doi:
672 <https://doi.org/10.1016/j.neuroimage.2010.06.010>.
- 673 **Dick F**, Tierney AT, Lutti A, Josephs O, Sereno MI, Weiskopf N. In vivo functional and myeloar-
674 chitectonic mapping of human primary auditory areas. *J Neurosci*. 2012; 32:16095–16105. doi:
675 <https://doi.org/10.1523/JNEUROSCI.1712-12.2012>.
- 676 **Dobkins KR**, Thiele A, Albright TD. Comparison of red-green equiluminance points in humans and
677 macaques: evidence for different L:M cone ratios between species. *J Opt Soc Am*. 2000; 17:545–556. doi:
678 <https://doi.org/10.1364/josaa.17.000545>.
- 679 **Dumoulin SO**, Harvey BM, Fracasso A, Zuiderbaan W, Luijten PR, Wandell BA, Petridou N. In vivo evidence
680 of functional and anatomical stripe-based subdivisions in human V2 and V3. *Sci Rep*. 2017; 7:733. doi:
681 <https://doi.org/10.1038/s41598-017-00634-6>.
- 682 **Edwards LJ**, Kirilina E, Mohammadi S, Weiskopf N. Microstructural imaging of human neocortex in vivo. *Neu-*
683 *roimage*. 2018; 182:184–206. doi: <https://doi.org/10.1016/j.neuroimage.2018.02.055>.
- 684 **Edwards LJ**, Pine KJ, Helms G, Weiskopf N. Rational approximation of the Ernst equation for dual angle
685 R_1 mapping revisited: beyond the small flip-angle assumption. In: *Book of Abstracts ESMRMB 2021 On-*
686 *line 38th Annual Scientific Meeting 7–9 October 2021. Magn Reson Mater Phy*, vol. 34; 2021. p. S45–S46. doi:
687 <https://doi.org/10.1007/s10334-021-00947-8>.
- 688 **Engel SA**, Glover GH, Wandell BA. Retinotopic organization in human visual cortex and the spatial precision of
689 functional MRI. *Cereb Cortex*. 1997; 7:181–192. doi: <https://doi.org/10.1093/cercor/7.2.181>.
- 690 **Feinberg DA**, Moeller S, Smith SM, Auerbach E, Ramanna S, Glasser MF, Miller KL, Uğurbil K, Yacoub E. Mul-
691 tiplexed echo planar imaging for sub-second whole brain fMRI and fast diffusion imaging. *PLoS ONE*. 2010;
692 5:e15710. doi: <https://doi.org/10.1371/journal.pone.0015710>.
- 693 **Fischl B**, Rajendran N, Busa E, Augustinack J, Hinds O, Yeo BTT, Mohlberg H, Amunts K, Zilles K. Cor-
694 tical folding patterns and predicting cytoarchitecture. *Cereb Cortex*. 2008; 18:1973–1980. doi:
695 <https://doi.org/10.1093/cercor/bhm225>.
- 696 **Fischl B**, Sereno MI, Dale AM. Cortical surface-based analysis. II: inflation, flattening, and a surface-based
697 coordinate system. *Neuroimage*. 1999; 9:195–207. doi: <https://doi.org/10.1006/nimg.1998.0396>.
- 698 **Flechsig P**. Anatomie des menschlichen Gehirns und Rückenmarks auf myelogenetischer Grundlage. Leipzig:
699 Georg Thieme; 1920.
- 700 **Fujimoto K**, Polimeni JR, van der Kouwe AJW, Reuter M, Kober T, Benner T, Fischl B, Wald LL. Quantitative
701 comparison of cortical surface reconstructions from MP2RAGE and multi-echo MP2RAGE data at 3 and 7T.
702 *Neuroimage*. 2014; 90:60–73. doi: <https://doi.org/10.1016/j.neuroimage.2013.12.012>.

703 **Glasser MF**, Sotiropoulos SN, Wilson JA, Coalson TS, Fischl B, Andersson JL, Xu J, Jbabdi S, Webster M, Polimeni
704 JR, Van Essen DC, Jenkinson M. The minimal preprocessing pipelines for the Human Connectome Project.
705 *Neuroimage*. 2013; 80:105–124. doi: <https://doi.org/10.1016/j.neuroimage.2013.04.127>.

706 **Glasser MF**, Van Essen DC. Mapping human cortical areas *in vivo* based on myelin content as revealed by
707 T1- and T2-weighted MRI. *J Neurosci*. 2011; 31:11597–11616. doi: <https://doi.org/10.1523/JNEUROSCI.2180-11.2011>.
708

709 **Greve DN**, Fischl B. Accurate and robust brain image alignment using boundary-based registration. *Neuroim-*
710 *age*. 2009; 48:63–72. doi: <https://doi.org/10.1016/j.neuroimage.2009.06.060>.

711 **Griswold MA**, Jakob PM, Heidemann RM, Nittka M, Jellus V, Wang J, Kiefer B, Haase A. Generalized
712 autocalibrating partially parallel acquisitions (GRAPPA). *Magn Reson Med*. 2002; 47:1202–1210. doi:
713 <https://doi.org/10.1002/mrm.10171>.

714 **Helms G**, Dathe H, Dechent P. Quantitative FLASH MRI at 3T using a rational approximation of the Ernst equa-
715 tion. *Magn Reson Med*. 2008; 59:667–672. doi: <https://doi.org/10.1002/mrm.21542>.

716 **Hinds OP**, Rajendran N, Polimeni JR, Augustinack JC, Wiggins G, Wald LL, Rosas D, Potthast A, Schwartz EL, Fischl
717 B. Accurate prediction of V1 location from cortical folds in a surface coordinate system. *Neuroimage*. 2008;
718 39:1585–1599. doi: <https://doi.org/10.1016/j.neuroimage.2007.10.033>.

719 **Hockfield S**, Tootell RBH, Zaremba S. Molecular differences among neurons reveal an organization of human
720 visual cortex. *Proc Natl Acad Sci USA*. 1990; 87:3027–3031. doi: <https://doi.org/10.1073/pnas.87.8.3027>.

721 **Horton JC**, Hocking DR. Myelin patterns in V1 and V2 of normal and monocularly enucleated monkeys. *Cereb*
722 *Cortex*. 1997; 7:166–177. doi: <https://doi.org/10.1093/cercor/7.2.166>.

723 **Hubel DH**, Livingstone MS. Segregation of form, color, and stereopsis in primate area 18. *J Neurosci*. 1987;
724 7:3378–3415. doi: <https://doi.org/10.1523/JNEUROSCI.07-11-03378.1987>.

725 **Ives FE**. A new color meter. *J Franklin Inst*. 1907; 164:47–56. doi: [https://doi.org/10.1016/S0016-0032\(07\)90164-](https://doi.org/10.1016/S0016-0032(07)90164-7)
726 [7](https://doi.org/10.1016/S0016-0032(07)90164-7).

727 **Jenkinson M**, Bannister P, Brady M, Stephen S. Improved optimization for the robust and accurate
728 linear registration and motion correction of brain images. *Neuroimage*. 2002; 17:825–841. doi:
729 [https://doi.org/10.1016/s1053-8119\(02\)91132-8](https://doi.org/10.1016/s1053-8119(02)91132-8).

730 **Julesz B**. *Foundations of Cyclopean Perception*. Chicago: University of Chicago Press; 1971.

731 **Keller AL**, Schüz A, Logothetis NK, Weber B. Vascularization of cytochrome oxidase-rich blobs in the
732 primary visual cortex of squirrel and macaque monkeys. *J Neurosci*. 2011; 31:1246–1253. doi:
733 <https://doi.org/10.1523/JNEUROSCI.2765-10.2011>.

734 **Kirilina E**, Helbling S, Morawski M, Pine K, Reimann K, Jankuhn S, Dinse J, Deistung A, Reichenbach JR,
735 Trampel R, Geyer S, Müller L, Jakubowski N, Arendt T, Bazin PL, Weiskopf N. Superficial white mat-
736 ter imaging: contrast mechanisms and whole-brain *in vivo* mapping. *Sci Adv*. 2020; 6:eaaz9281. doi:
737 <https://doi.org/10.1126/sciadv.aaz9281>.

738 **Kleiner M**, Brainard DH, Pelli D, Ingling A, Murray R, Broussard C. What's new in psychtoolbox-3. *Perception*.
739 2007; 36:1–16. doi: <https://doi.org/10.1177/030100660703605101>.

740 **Krubitzer LA**, Kaas JH. Cortical integration of parallel pathways in the visual system of primates. *Brain Res*.
741 1989; 478:161–165. doi: [https://doi.org/10.1016/0006-8993\(89\)91490-x](https://doi.org/10.1016/0006-8993(89)91490-x).

742 **Lazari A**, Lipp I. Can MRI measure myelin? Systematic review, qualitative assessment, and meta-analysis
743 of studies validating microstructural imaging with myelin histology. *Neuroimage*. 2021; 230:117744. doi:
744 <https://doi.org/10.1016/j.neuroimage.2021.117744>.

745 **Leuze C**, Aswendt M, Ferenczi E, Liu CW, Hsueh B, Goubran M, Tian Q, Steinberg G, Zeineh MM, Deisseroth K,
746 McNab JA. The separate effects of lipids and proteins on brain MRI contrast revealed through tissue clearing.
747 *Neuroimage*. 2017; 156:412–422. doi: <https://doi.org/10.1016/j.neuroimage.2017.04.021>.

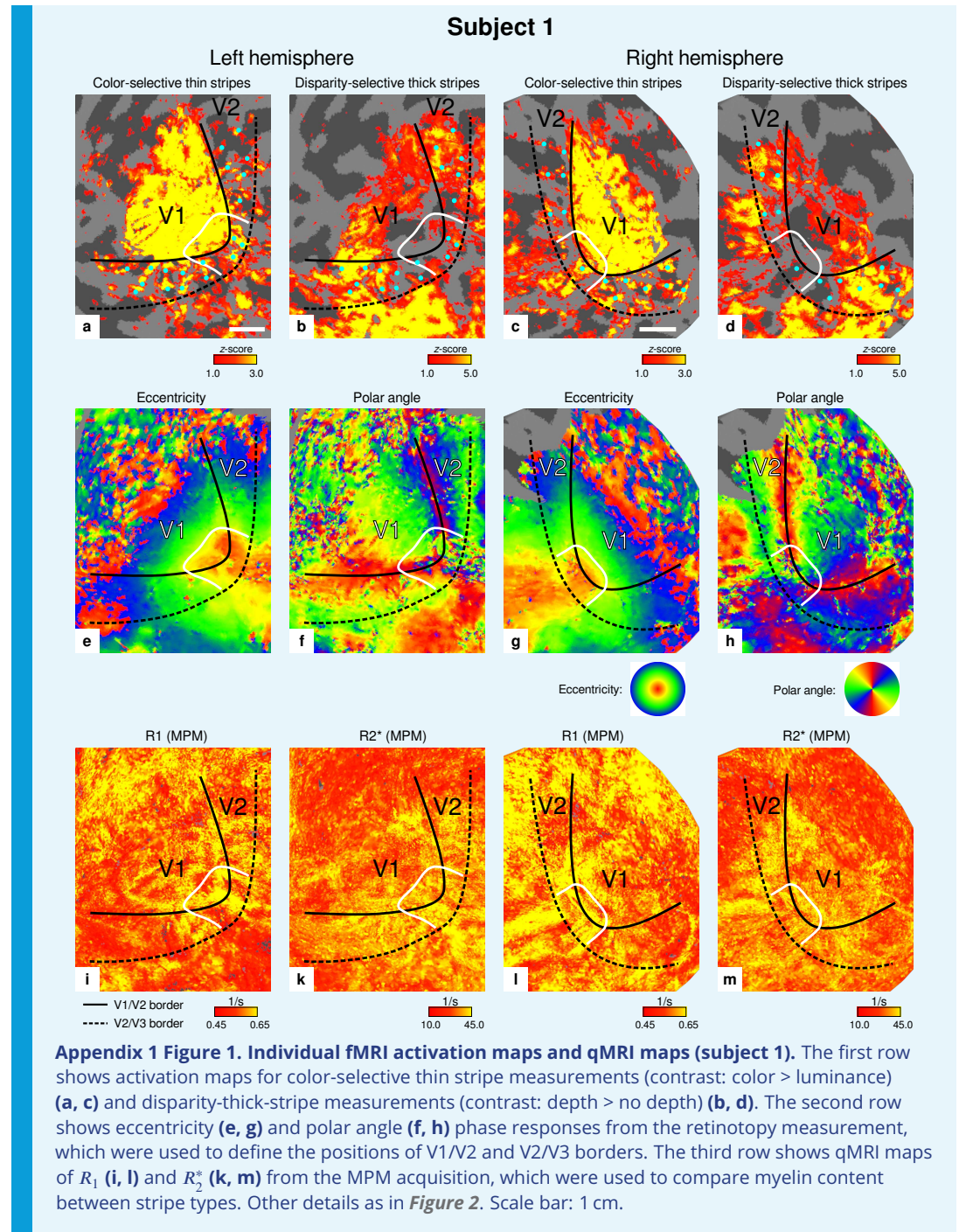
748 **Li X**, Zhu Q, Janssens T, Arsenault JT, Vanduffel W. *In vivo* identification of thick, thin, and pale stripes of
749 macaque area V2 using submillimeter resolution (f)MRI at 3 T. *Cereb Cortex*. 2019; 29:544–560. doi:
750 <https://doi.org/10.1093/cercor/bhx337>.

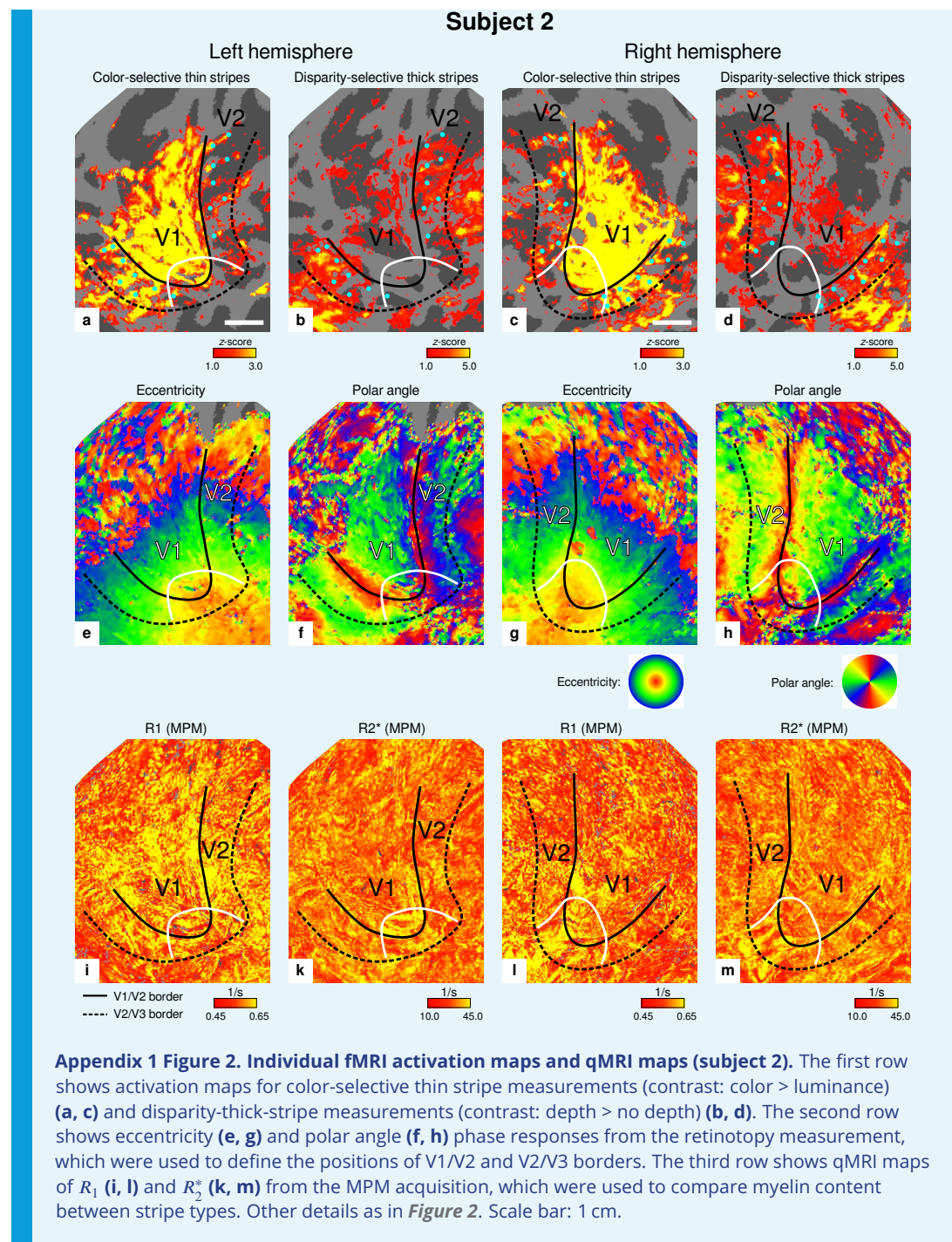
- 751 **Livingstone MS**, Hubel DH. Thalamic inputs to cytochrome oxidase-rich regions in monkey visual cortex. *Proc*
752 *Natl Acad Sci USA*. 1982; 79:6098–6101. doi: <https://doi.org/10.1073/pnas.79.19.6098>.
- 753 **Livingstone MS**, Hubel DH. Psychophysical evidence for separate channels for the perception of form, color,
754 movement, and depth. *J Neurosci*. 1987; 7:3416–3468. doi: [https://doi.org/10.1523/JNEUROSCI.07-11-](https://doi.org/10.1523/JNEUROSCI.07-11-03416.1987)
755 [03416.1987](https://doi.org/10.1523/JNEUROSCI.07-11-03416.1987).
- 756 **Lutti A**, Dick F, Sereno MI, Weiskopf N. Using high-resolution quantitative mapping of R1 as an index of cortical
757 myelination. *Neuroimage*. 2014; 93:176–188. doi: <https://doi.org/10.1016/j.neuroimage.2013.06.005>.
- 758 **Lutti A**, Hutton C, Finsterbusch J, Helms G, Weiskopf N. Optimization and validation of methods for
759 mapping of the radiofrequency transmit field at 3T. *Magn Reson Med*. 2010; 64:229–238. doi:
760 <https://doi.org/10.1002/mrm.22421>.
- 761 **Lutti A**, Stadler J, Josephs O, Windischberger C, Speck O, Bernarding J, Hutton C, Weiskopf N. Robust
762 and fast whole brain mapping of the RF transmit field B1 at 7T. *PloS ONE*. 2012; 7:e32379. doi:
763 <https://doi.org/10.1371/journal.pone.0032379>.
- 764 **Marques JP**, Khabipova D, Gruetter R. Studying cyto and myeloarchitecture of the human cortex at ultra-high
765 field with quantitative imaging: R_1 , R_2^* and magnetic susceptibility. *Neuroimage*. 2017; 147:152–163. doi:
766 <https://doi.org/10.1016/j.neuroimage.2016.12.009>.
- 767 **Marques JP**, Kober T, Krueger G, van der Zwaag W, Van de Moortele PF, Gruetter R. MP2RAGE, a self bias-field
768 corrected sequence for improved segmentation and T_1 -mapping at high field. *Neuroimage*. 2010; 49:1271–
769 1281. doi: <https://doi.org/10.1016/j.neuroimage.2009.10.002>.
- 770 **McColgan P**, Helbling S, Vaculčíaková L, Pine K, Wagstyl K, Attar FM, Edwards L, Papoutsis M, Wei Y, Van den
771 Heuvel MP, Tabrizi S, Rees G, Weiskopf N. Relating quantitative 7T MRI across cortical depths to cy-
772 toarchitectonics, gene expression and connectomics. *Hum Brain Mapp*. 2021; 42:4996–5009. doi:
773 <https://doi.org/10.1002/hbm.25595>.
- 774 **Mezer A**, Yeatman JD, Stikov N, Kay KN, Cho NJ, Dougherty RF, Perry ML, Parvizi J, Hua LH, Butts-Pauly K, Wan-
775 dell BA. Quantifying the local tissue volume and composition in individual brains with magnetic resonance
776 imaging. *Nat Med*. 2013; 19:1667–1672. doi: <https://doi.org/10.1038/nm.3390>.
- 777 **Moeller S**, Yacoub E, Olman CA, Auerbach E, Strupp J, Harel N, Uğurbil K. Multiband multislice GE-EPI at 7
778 Tesla, With 16-fold acceleration using partial parallel imaging with application to high spatial and temporal
779 whole-brain fMRI. *Magn Reson Med*. 2010; 63:1144–1153. doi: <https://doi.org/10.1002/mrm.22361>.
- 780 **Nasr S**, Polimeni JR, Tootell RBH. Interdigitated color- and disparity-selective columns within human visual
781 cortical areas V2 and V3. *J Neurosci*. 2016; 36:1841–1857. doi: [https://doi.org/10.1523/JNEUROSCI.3518-](https://doi.org/10.1523/JNEUROSCI.3518-15.2016)
782 [15.2016](https://doi.org/10.1523/JNEUROSCI.3518-15.2016).
- 783 **Nasr S**, Tootell RBH. Columnar organization of mid-spectral and end-spectral hue preferences in human visual
784 cortex. *Neuroimage*. 2018; 181:748–759. doi: <https://doi.org/10.1016/j.neuroimage.2018.07.053>.
- 785 **Navarro KT**, Sanchez MJ, Engel SA, Olman CA, Weldon KB. Depth-dependent functional MRI responses
786 to chromatic and achromatic stimuli throughout V1 and V2. *Neuroimage*. 2021; 226:117520. doi:
787 <https://doi.org/10.1016/j.neuroimage.2020.117520>.
- 788 **Olman CA**, Inati S, Heeger DJ. The effect of large veins on spatial localization with GE BOLD at 3 T: displacement,
789 not blurring. *Neuroimage*. 2007; 34:1126–1135. doi: <https://doi.org/10.1016/j.neuroimage.2006.08.045>.
- 790 **Pelli DG**. The VideoToolbox software for visual psychophysics: transforming numbers into movies. *Spat Vis*.
791 1997; 10:437–442. doi: <https://doi.org/10.1163/156856897X00366>.
- 792 **Peterhans E**, von der Heydt R. Functional organization of area V2 in the alert macaque. *Eur J Neurosci*. 1993;
793 5:509–524. doi: <https://doi.org/10.1111/j.1460-9568.1993.tb00517.x>.
- 794 **Peters AM**, Brookes MJ, Hoogenraad FG, Gowland PA, Francis ST, Morris PG, Bowtell R. T_2^* mea-
795 surements in human brain at 1.5, 3 and 7 T. *Magn Reson Med*. 2007; 25:748–753. doi:
796 <https://doi.org/10.1016/j.mri.2007.02.014>.
- 797 **Pistorio AL**, Hendry SH, Wang X. A modified technique for high-resolution staining of myelin. *J Neurosci*
798 *Methods*. 2006; 153:135–146. doi: <https://doi.org/10.1016/j.jneumeth.2005.10.014>.

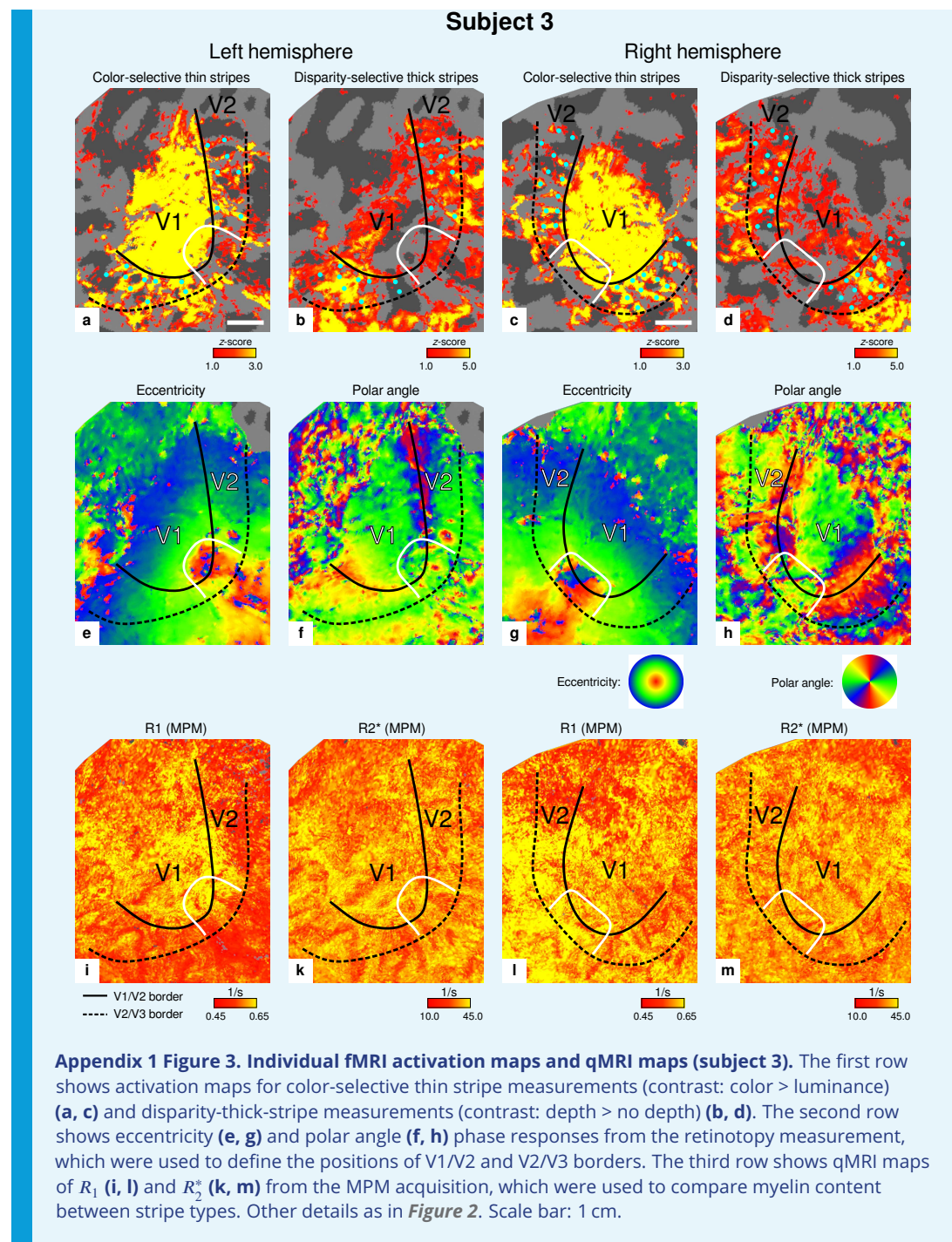
- 799 **Pluim JPW**, Muenzing SEA, Eppenhof KAJ, Murphy K. The truth is hard to make: validation of medical image
800 registration. 2016 23rd International Conference on Pattern Recognition (ICPR). 2016; p. 2294–2300. doi:
801 <https://doi.org/10.1109/ICPR.2016.7899978>.
- 802 **Polimeni JR**, Fischl B, Greve DN, Wald LL. Laminar analysis of 7T BOLD using an im-
803 posed spatial activation pattern in human V1. Neuroimage. 2010; 52:1334–1346. doi:
804 <https://doi.org/10.1016/j.neuroimage.2010.05.005>.
- 805 **Sanders FK**, Whitteridge D. Conduction velocity and myelin thickness in regenerating nerve fibres. J Physiol.
806 1946; 105:152–174. doi: <https://doi.org/10.1113/jphysiol.1946.sp004160>.
- 807 **Savaskan NE**, Weinmann O, Heimrich B, Eyupoglu IY. High resolution neurochemical gold staining method for
808 myelin in peripheral and central nervous system at the light- and electron-microscopic level. Cell Tissue Res.
809 2009; 337:213–221. doi: <https://doi.org/10.1007/s00441-009-0815-9>.
- 810 **Sereno MI**, Dale AM, Reppas JB, Kwong KK, Belliveau JW, Brady TJ, Rosen BR, Tootell RBH. Borders of multiple
811 visual areas in humans revealed by functional magnetic resonance imaging. Science. 1995; 268:889–893. doi:
812 <https://doi.org/10.1126/science.7754376>.
- 813 **Sereno MI**, Lutti A, Weiskopf N, Dick F. Mapping the human cortical surface by combining quantitative T1 with
814 retinotopy. Cereb Cortex. 2013; 23:2261–2268. doi: <https://doi.org/10.1093/cercor/bhs213>.
- 815 **Shipp S**, Zeki S. Segregation of pathways leading from area V2 to areas V4 and V5 of macaque monkey visual
816 cortex. Nature. 1985; 315:322–324. doi: <https://doi.org/10.1038/315322a0>.
- 817 **Sigalovsky IS**, Fischl B, Melcher JR. Mapping an intrinsic MR property of gray matter in auditory cortex of living
818 humans: a possible marker for primary cortex and hemispheric differences. Neuroimage. 2006; 32:1524–
819 1537. doi: <https://doi.org/10.1016/j.neuroimage.2006.05.023>.
- 820 **Sincich LC**, Horton JC. Pale cytochrome oxidase stripes in V2 receive the richest projection from macaque
821 striate cortex. J Comp Neurol. 2002; 447:18–33. doi: <https://doi.org/10.1002/cne.10174>.
- 822 **Sincich LC**, Horton JC. The circuitry of V1 and V2: integration of color, form, and motion. Annu Rev Neurosci.
823 2005; 28:303–346. doi: <https://doi.org/10.1146/annurev.neuro.28.061604.135731>.
- 824 **Smart IHM**, McSherry GM. Gyrus formation in the cerebral cortex of the ferret. II. description of the internal
825 histological changes. J Anat. 1986; 147:27–43.
- 826 **Stüber C**, Morawski M, Schäfer A, Labadie C, Wähnert M, Leuze C, Streicher M, Barapatre N, Reimann K, Geyer
827 S, Spemann D, Turner R. Myelin and iron concentration in the human brain: a quantitative study of MRI
828 contrast. Neuroimage. 2014; 93:95–106. doi: <https://doi.org/10.1016/j.neuroimage.2014.02.026>.
- 829 **Tabelow K**, Balteau E, Ashburner J, Callaghan MF, Draganski B, Helms G, Kherif F, Leutritz T, Lutti A,
830 Phillips C, Reimer E, Ruthotto L, Seif M, Weiskopf N, Ziegler G, Mohammadi S. hMRI — a toolbox
831 for quantitative MRI in neuroscience and clinical research. Neuroimage. 2019; 194:191–210. doi:
832 <https://doi.org/10.1016/j.neuroimage.2019.01.029>.
- 833 **Tofts PS**. PD: proton density of tissue water. In: *Quantitative MRI of the brain: principles of physical measurement*
834 Boca Raton: CRC Press; 2018. p. 55–71. doi: <https://doi.org/10.1201/b21837>.
- 835 **Tootell RBH**, Hamilton SL. Functional anatomy of the second visual area (V2) in the macaque. J Neurosci. 1989;
836 9:2620–2644. doi: <https://doi.org/10.1523/JNEUROSCI.09-08-02620.1989>.
- 837 **Tootell RBH**, Nasr S. Columnar segregation of magnocellular and parvocellular streams in human extrastriate
838 cortex. J Neurosci. 2017; 37:8014–8032. doi: <https://doi.org/10.1523/JNEUROSCI.0690-17.2017>.
- 839 **Tootell RBH**, Nelissen K, Vanduffel W, Orban GA. Search for color ‘center(s)’ in macaque visual cortex. Cereb
840 Cortex. 2004; 14:353–363. doi: <https://doi.org/10.1093/cercor/bhh001>.
- 841 **Tootell RBH**, Silverman MS, De Valois RL, Jacobs GH. Functional organization of the second cortical visual area
842 in primates. Science. 1983; 220:737–739. doi: <https://doi.org/10.1126/science.6301017>.
- 843 **Tootell RBH**, Taylor JB. Anatomical evidence for MT and additional cortical visual areas in humans. Cereb
844 Cortex. 1995; 5:39–55. doi: <https://doi.org/10.1093/cercor/5.1.39>.
- 845 **Trampel R**, Bazin PL, Pine K, Weiskopf N. In-vivo magnetic resonance imaging (MRI) of laminae in the human
846 cortex. Neuroimage. 2019; 197:707–715. doi: <https://doi.org/10.1016/j.neuroimage.2017.09.037>.

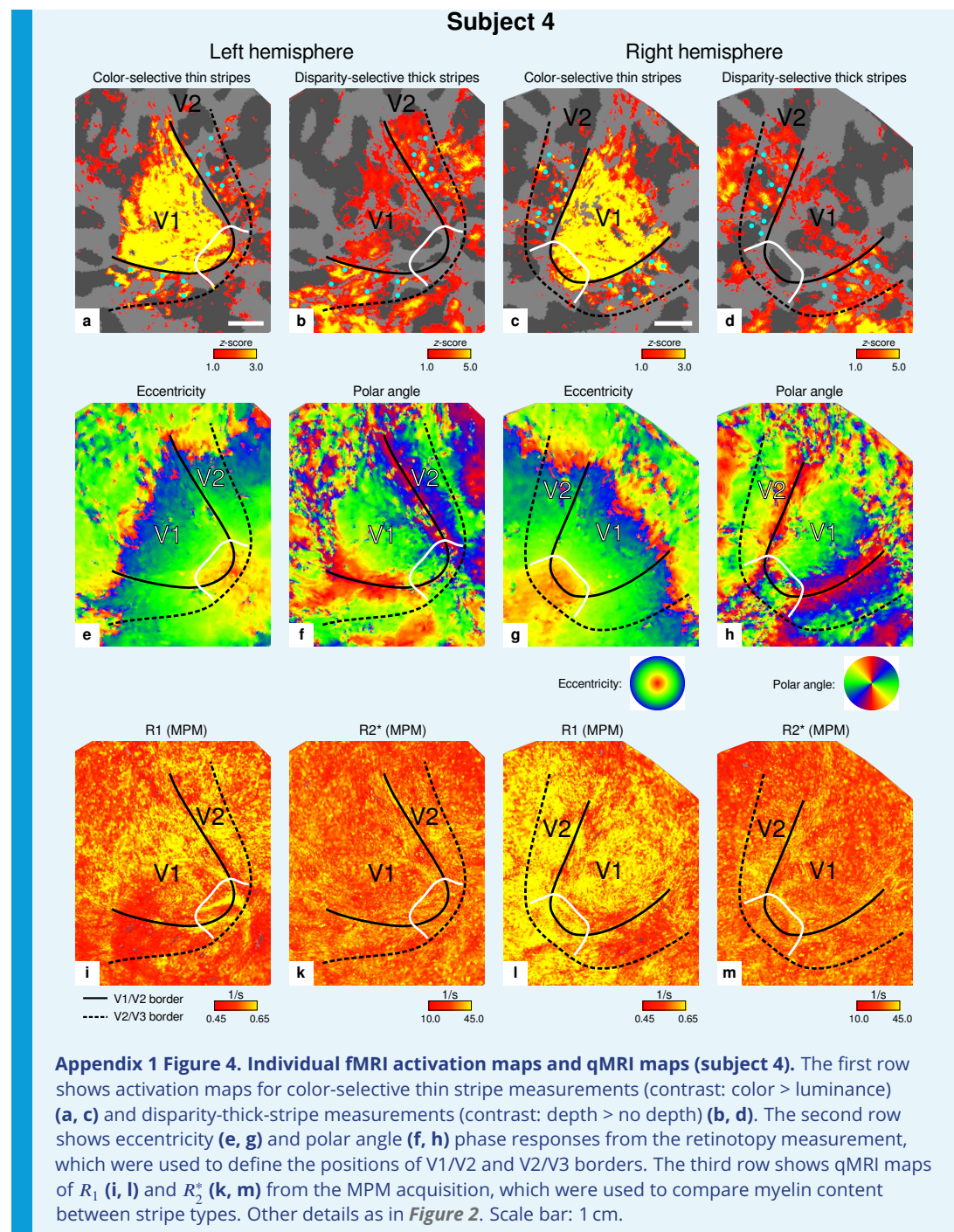
- 847 **Tsao DY**, Vanduffel W, Sasaki Y, Fize D, Knutsen TA, Mandeville JB, Wald LL, Dale AM, Rosen BR, Van Essen DC,
848 Livingstone MS, Orban GA, Tootell RBH. Stereopsis activates V3A and caudal intraparietal areas in macaques
849 and humans. *Neuron*. 2003; 39:555–568. doi: [https://doi.org/10.1016/s0896-6273\(03\)00459-8](https://doi.org/10.1016/s0896-6273(03)00459-8).
- 850 **Ts'o DY**, Roe AW, Gilbert CD. A hierarchy of the functional organization for color, form and disparity in primate
851 visual area V2. *Vis Res*. 2001; 41:1333–1349. doi: [https://doi.org/10.1016/s0042-6989\(01\)00076-1](https://doi.org/10.1016/s0042-6989(01)00076-1).
- 852 **Tustison NJ**, Avants BB, Cook PA, Zheng Y, Egan A, Yushkevich PA, Gee JC. N4ITK: improved N3 bias correction.
853 *IEEE Trans Med Imaging*. 2010; 29:1310–1320. doi: <https://doi.org/10.1109/TMI.2010.2046908>.
- 854 **Vaculčíaková L**, Podranski K, Edwards LJ, Ocal D, Veale T, Fox NC, Haak R, Ehses P, Callaghan MF, Pine
855 KJ, Weiskopf N. Combining navigator and optical prospective motion correction for high-quality 500 µm
856 resolution quantitative multi-parameter mapping at 7T. *Magn Reson Med*. 2022; 88:787–801. doi:
857 <https://doi.org/10.1002/mrm.29253>.
- 858 **Volz S**, Nöth U, Deichmann R. Correction of systematic errors in quantitative proton density mapping. *Magn*
859 *Reson Med*. 2012; 68:74–85. doi: <https://doi.org/10.1002/mrm.23206>.
- 860 **Waehnert MD**, Dinse J, Weiss M, Streicher MN, Waehnert P, Geyer S, Turner R, Bazin PL.
861 Anatomically motivated modeling of cortical laminae. *Neuroimage*. 2014; 93:210–220. doi:
862 <https://doi.org/10.1016/j.neuroimage.2013.03.078>.
- 863 **Wagstyl K**, Paquola C, Bethlehem R, Evans AC, Huth A, Equivolumetric layering for mesh surfaces. Zenodo;
864 2018. doi: <https://doi.org/10.5281/ZENODO.1412054>, last access: 2021-11-05.
- 865 **Weiskopf N**, Callaghan MF, Josephs O, Lutti A, Mohammadi S. Estimating the apparent transverse relaxation
866 time (R_2^*) from images with different contrasts (ESTATICS) reduces motion artifacts. *Front Neurosci*. 2014;
867 8:278. doi: <https://doi.org/10.3389/fnins.2014.00278>.
- 868 **Weiskopf N**, Edwards LJ, Helms G, Mohammadi S, Kirilina E. Quantitative magnetic resonance imaging of brain
869 anatomy and in-vivo histology. *Nat Rev Phys*. 2021; 3:570–588. doi: <https://doi.org/10.1038/s42254-021-00326-1>.
- 870
871 **Weiskopf N**, Lutti A, Helms G, Novak M, Ashburner J, Hutton C. Unified segmentation based correction of
872 R_1 brain maps for RF transmit field inhomogeneities (UNICORT). *Neuroimage*. 2011; 54:2116–2124. doi:
873 <https://doi.org/10.1016/j.neuroimage.2010.10.023>.
- 874 **Xiao Y**, Wang Y, Felleman DJ. A spatially organized representation of colour in macaque cortical area V2. *Nature*.
875 2003; 421:535–539. doi: <https://doi.org/10.1038/nature01372>.
- 876 **Yacoub E**, Shmuel A, Pfeuffer J, Van De Moortele PF, Adriany G, Andersen P, Vaughan JT, Merkle H, Uğur-
877 bil K, Hu X. Imaging brain function in humans at 7 Tesla. *Magn Reson Med*. 2001; 45:588–594. doi:
878 <https://doi.org/10.1002/mrm.1080>.
- 879 **Zaretskaya N**, Fischl B, Reuter M, Renvall V, Polimeni JR. Advantages of cortical surface re-
880 construction using submillimeter 7 T MEMPRAGE. *Neuroimage*. 2018; 165:11–26. doi:
881 <https://doi.org/10.1016/j.neuroimage.2017.09.060>.
- 882 **Zheng D**, LaMantia AS, Purves D. Specialized vascularization of the primate visual cortex. *J Neurosci*. 1991;
883 11:2622–2629. doi: <https://doi.org/10.1523/JNEUROSCI.11-08-02622.1991>.

Appendix 1

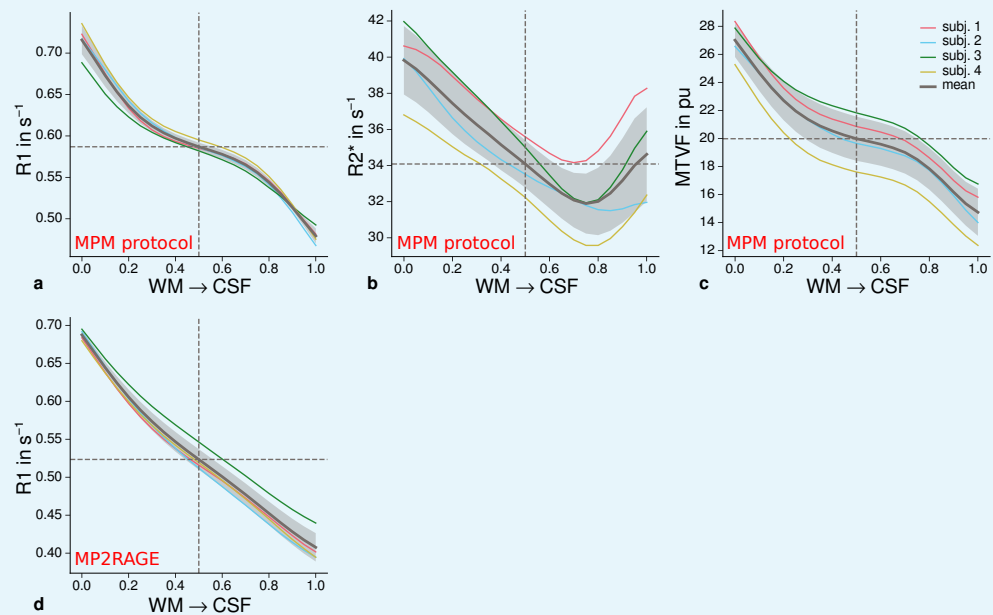






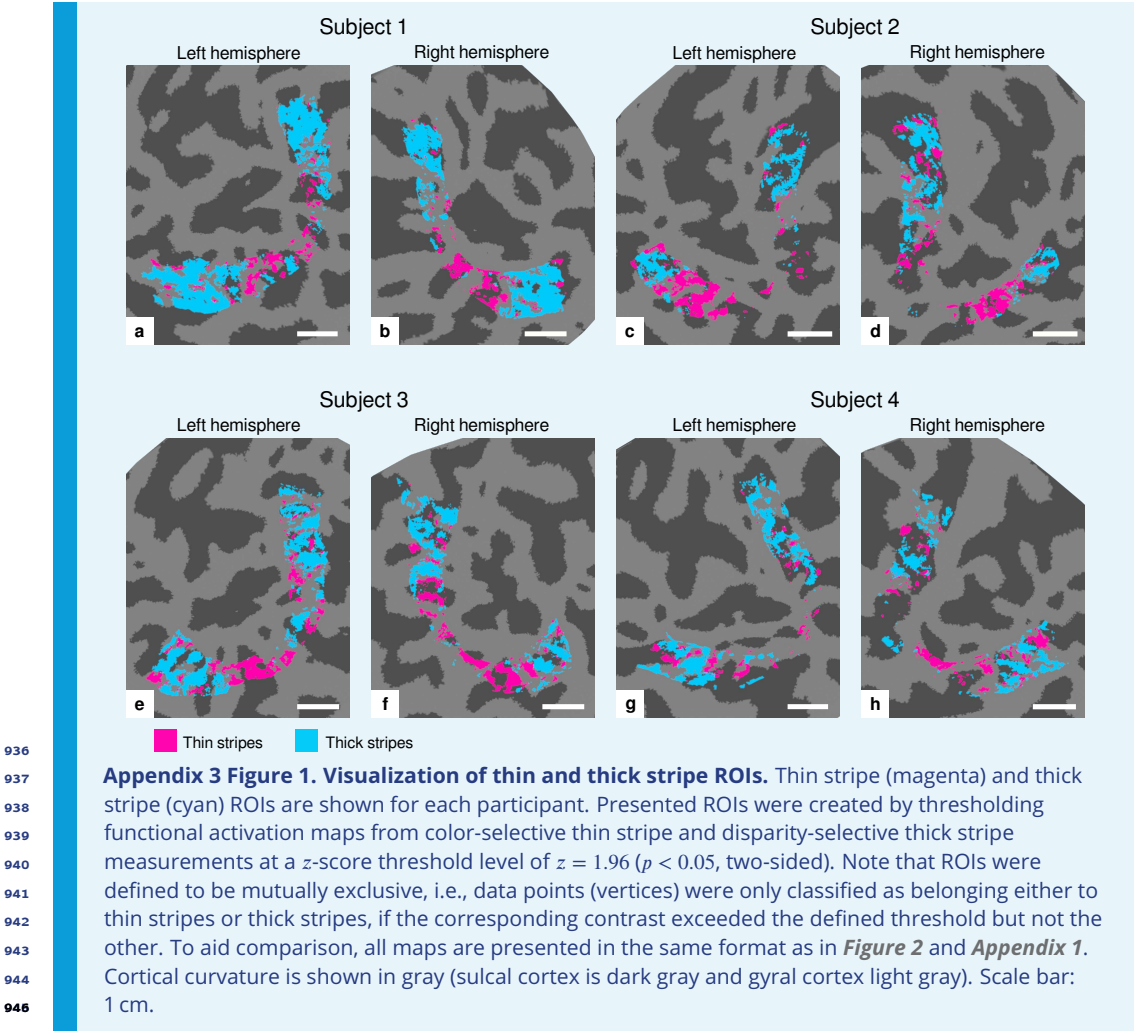


Appendix 2

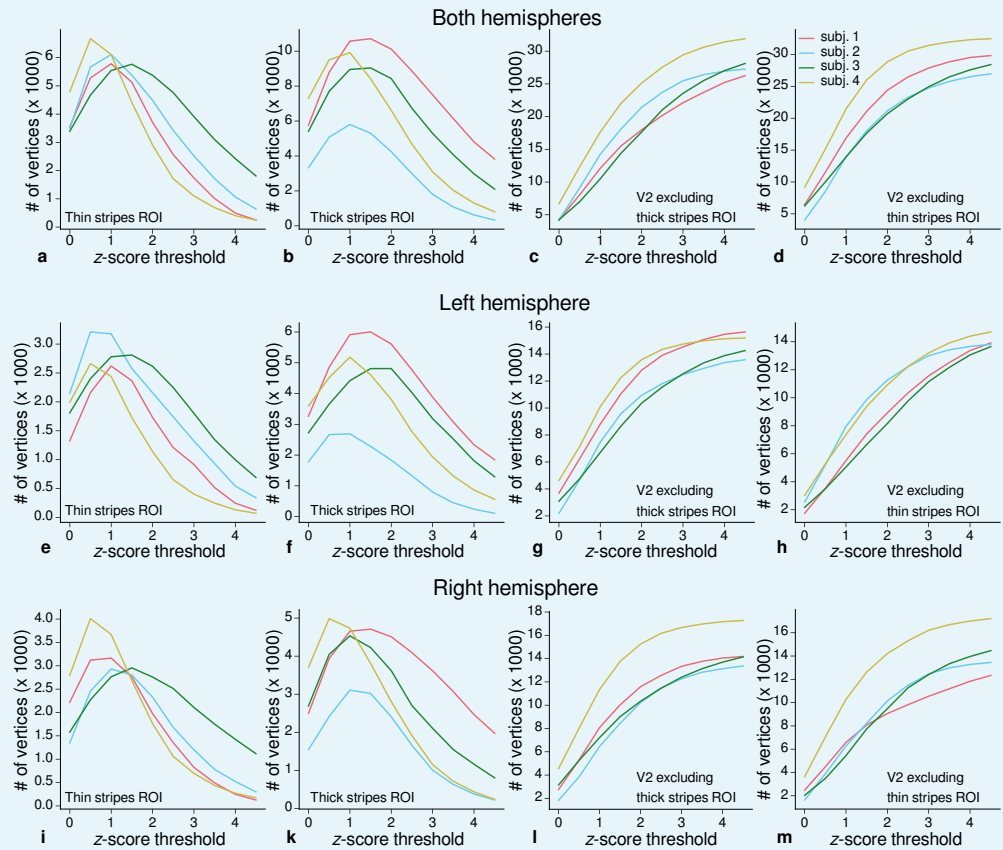


Appendix 2 Figure 1. Cortical profiles of mean qMRI parameters within V2. Mean R_1 (a), R_2^* (b) and $MTVF = 100\% - PD$ (macromolecular tissue volume fraction (Mezer et al., 2013)) (c) values based on the MPM protocol were sampled at 21 cortical depths between the gray matter/white matter boundary surface and pial surface. The expected decrease of quantitative parameters towards the pial surface can be seen in all plots. (d) shows mean R_1 across cortical depth based on a separate data set using the MP2RAGE sequence. A similar decrease as shown in (a) can be seen. However, in comparison to (a), a plateau at middle depths is less visible. This might be attributable to the larger voxel size in (d) and the larger point spread function along the phase-encoding direction (Marques et al., 2010) of MP2RAGE acquisitions. The mean across participants is shown as gray line. Dashed lines mark the mean at mid-cortical depth. Shaded areas indicate 1 standard deviation across participants.

935 **Appendix 3**

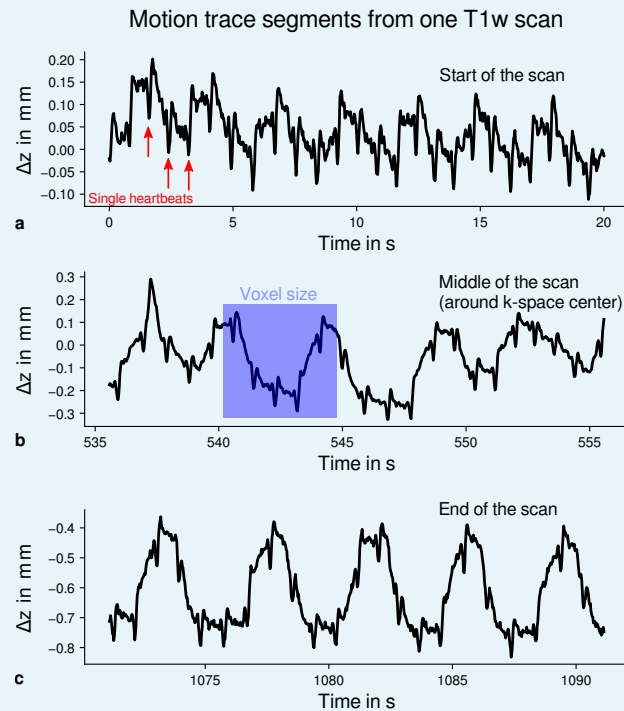


Appendix 4



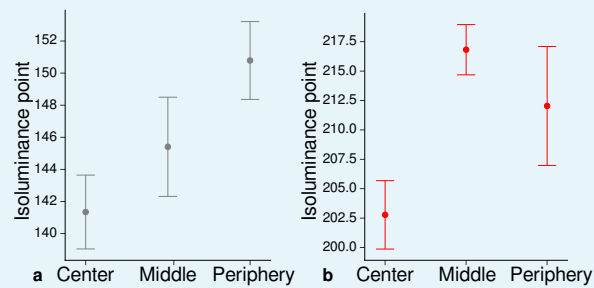
Appendix 4 Figure 1. Sizes of stripe ROIs for each threshold level. ROI sizes (number of vertices) across hemispheres for color-selective thin stripes (a), disparity-selective thick stripes (b) and whole V2 excluding one stripe type (c)–(d) are shown for several z-score threshold levels and all participants (upper row). These ROIs were used for comparison of qMRI parameters (R_1 , R_2^*) between stripe types as shown in Figure 5. Voxels activated by both color and disparity contrasts were excluded from thin and thick stripe ROIs, explaining the non-monotonous dependence on the z-score threshold. Middle and lower rows show number of vertices from single hemispheres.

957 Appendix 5



Appendix 5 Figure 1. Exemplary motion data from one MPM measurement. For one representative participant (subject 3), head movements are shown from the 3D multi-echo fast low angle shot (FLASH) scan with T_1 -weighting (T1w). Movements were measured during acquisition using an optical tracking system and used for prospective motion correction. From the start (**a**), the middle (time of k -space center sampling) (**b**) and the end of the scan (**c**), excerpts of 20 s depict translational movements for each TR = 25 ms in z -direction (inferior-superior) relative to start of the scan. In all plots, the participant's breathing pattern can be seen as slow oscillation. On top of this oscillation, displacements at a faster rate related to the cardiac cycle can be identified. In (**a**), a subset of cardiac beats are marked by red arrows. This demonstrates the ability to detect small movements, which were prospectively corrected during high resolution anatomical scans. Note that motion was overall on the order of the $(0.5 \text{ mm})^3$ voxel size (even for the short excerpts shown), which makes an accurate correction for subject motion necessary during the long scan time (acquisition time around 18 min). This is qualitatively illustrated by showing the 0.5 mm voxel edge length as blue square in (**b**). It can be further noticed that movements were getting larger towards the end of the scan as expected due to the long scan time.

Appendix 6



Appendix 6 Figure 1. Measured mean isoluminance points across participants and sessions at different eccentricities. For each participant, isoluminance points were measured before scanning within the scanner at three different eccentricities (Center: 0°-1.7°, Middle: 1.7°-4.1°, Periphery: 4.1°-8.3°). Luminance points for gray (**a**) and red (**b**) were adjusted to match blue (RGB: 0, 0, 255), see Materials and methods. Across participants and scanning sessions, we found a statistically significant effect of eccentricity for gray ($F(2, 93) = 3.25$, $p = 0.04$, $\eta_p^2 = 0.07$) and red ($F(2, 93) = 3.96$, $p = 0.02$, $\eta_p^2 = 0.08$) with the application of a one-way ANOVA. Error bars indicate 1 standard error of the mean.

Appendix 7

Analysis of registration accuracy

Accurate registration between qMRI and fMRI data is of utmost importance for the current study. However, knowledge about the quality of registration beyond visual inspection, which is subjective, qualitative and not usable for large quantities of data, is difficult to achieve (*Pluim et al., 2016*).

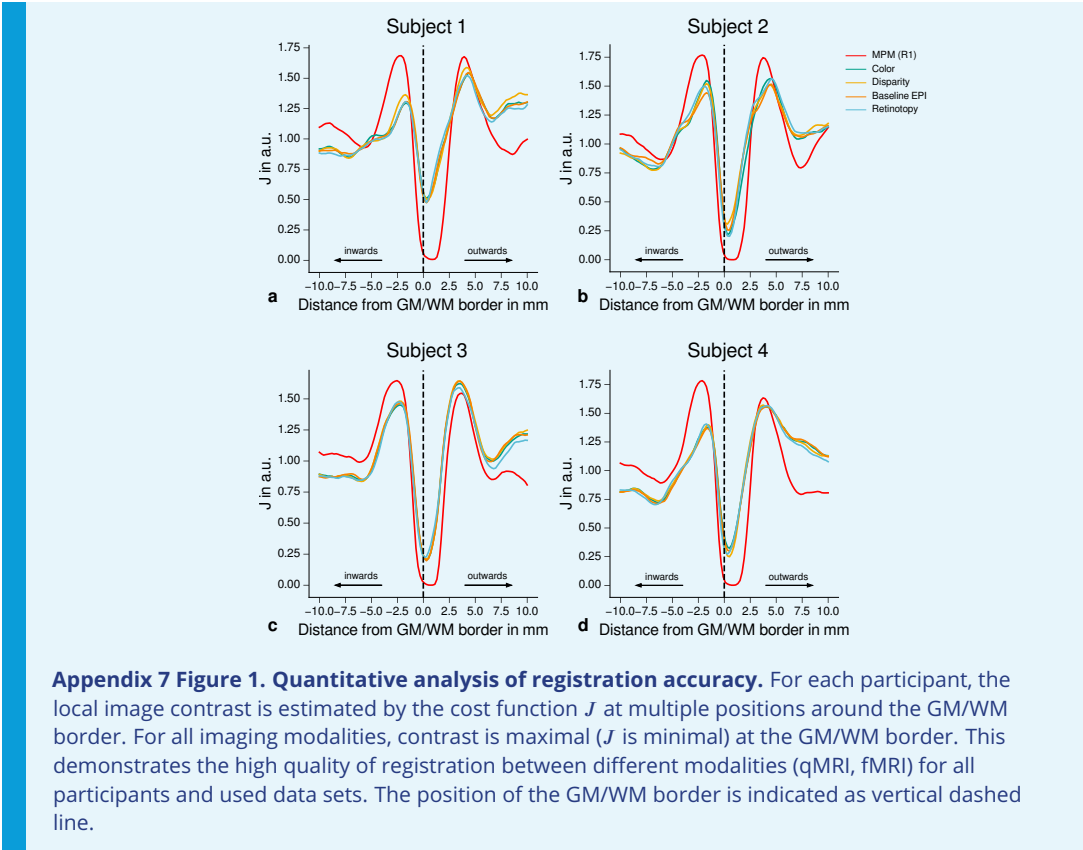
In the following analysis, we quantitatively assessed the quality of registration by exploiting that both qMRI and fMRI data were registered to the same whole-brain anatomy (MP2RAGE). Knowing that in all acquired MR images, gray matter (GM) and white matter (WM) can easily be differentiated, we estimated registration accuracy by the analysis of local image contrast of the registered image around the GM/WM border, which was defined by cortex segmentation based on the MP2RAGE. We expect that the local image contrast is higher at the WM/GM border than in nearby locations in the registered image if registration between whole-brain anatomy and qMRI and fMRI is accurate, respectively. We restricted the analysis to vertices v in V2. For each vertex v , we computed a normal vector and defined 100 points along the normal in equidistant steps around the GM/WM border (± 10 mm). Let λ be the parameter defining the position on the normal line, then $v(\lambda)$ is the coordinate of vertex v shifted along the normal direction by λ . For each $v(\lambda)$, we sampled data from two points with 1 mm distance along the normal vector in both positive v_+ and negative v_- direction using trilinear interpolation. To evaluate the local image contrast, we used the cost function J , which was introduced for the boundary-based registration (BBR) method (*Greve and Fischl, 2009*). J has the property to approach zero if local image contrast increases. Note that we did not use the BBR method for actual registration. To derive J , a contrast measure $Q(\lambda)$ is first computed for each $v(\lambda)$ in V2.

$$Q(\lambda) = 100 \frac{v_+(\lambda) - v_-(\lambda)}{0.5 \cdot (v_+(\lambda) + v_-(\lambda))} \quad (1)$$

The sign of the contrast depends on the imaging modality, i.e., if WM is darker or brighter than GM. This was considered by choosing the direction of the normal vector to point towards the brighter tissue component. J was then computed based on all contrast measures in V2.

$$J(\lambda) = \frac{1}{N} \sum_{v(\lambda) \in V2} [1 + \tanh(0.5 \cdot Q(\lambda))] \quad (2)$$

where N is the number of vertices in V2. **Appendix 7 Figure 1** shows for each participant and imaging modality the size of J around the GM/WM border. It can be seen that the position of minimum J generally coincides with the GM/WM border for each data set. This enables us to conclude that accurate registration could be achieved for both qMRI and fMRI data, respectively. However, a small shift was also observed in all data sets relative to the GM/WM border. While partial volume effects could have contributed to this bias, we note that we applied an inward shift (0.5 mm) to the GM/WM boundary surface during segmentation (see **Materials and methods**). This was done to counteract a potential bias in the segmentation pipeline using the flat image from the MP2RAGE acquisition which yielded visually better results. The current analysis reveals that this step was probably not necessary. However, note that this did not affect any conclusions drawn in further analyses, since the shift is negligible small when compared to changes in local image contrast as shown in **Appendix 7 Figure 1**.



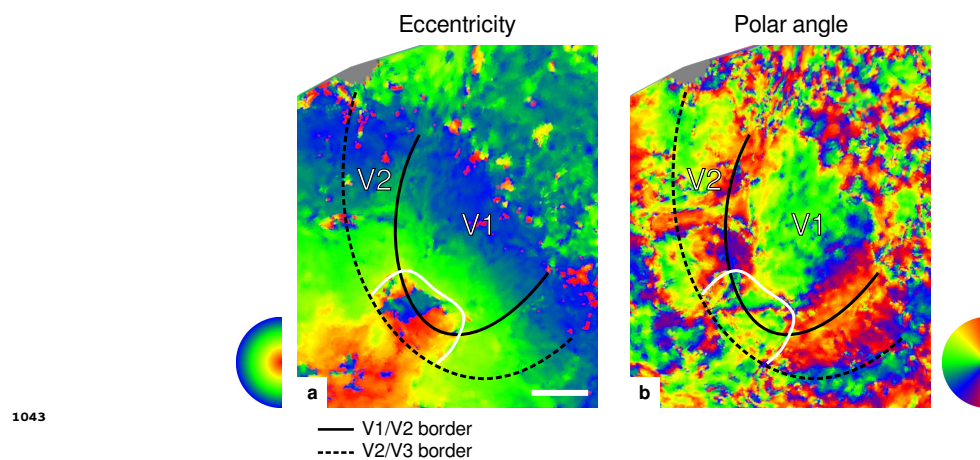


Figure 2-Figure supplement 1. Eccentricity and polar angle maps. Maps in (a) and (b) show eccentricity and polar angle phase responses, respectively, for the participant (subject 3) shown in Figure 2. The white border encircles a region around the foveal representation in V1 and V2 and was defined by manually drawing a line around the region of eccentricity values $\varphi < 2\pi/3$ ($\varphi \in [0, 2\pi]$). Borders between V1/V2 and V2/V3 were based on the polar angle data and defined by the location of vertical and horizontal meridians around the calcarine fissure, respectively. Other details as in Figure 2. Maps from all participants can be found in Appendix 1. In comparison to all other participants, retinotopy data for this participant was very noisy. However, visual borders could still clearly be identified. Note that, in contrast to other functional measurements, we did not control for subject attention during retinotopy measurements. Scale bar: 1 cm.

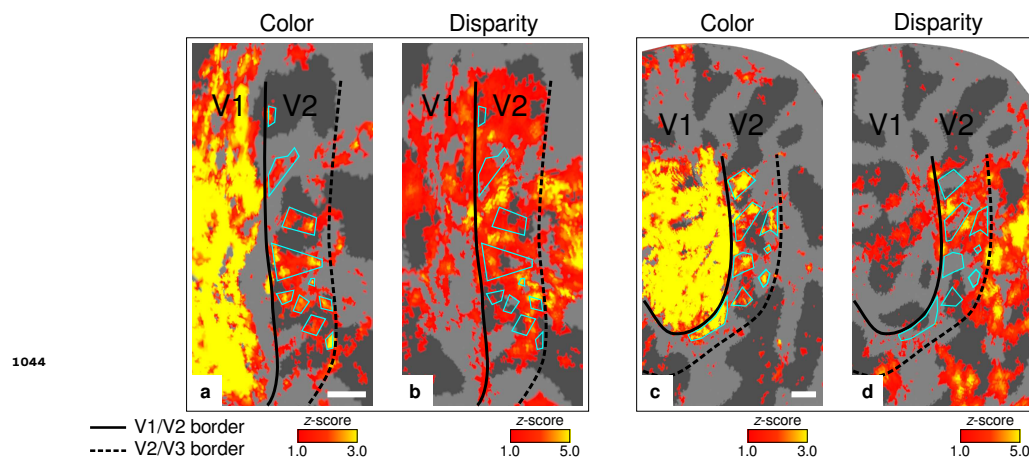
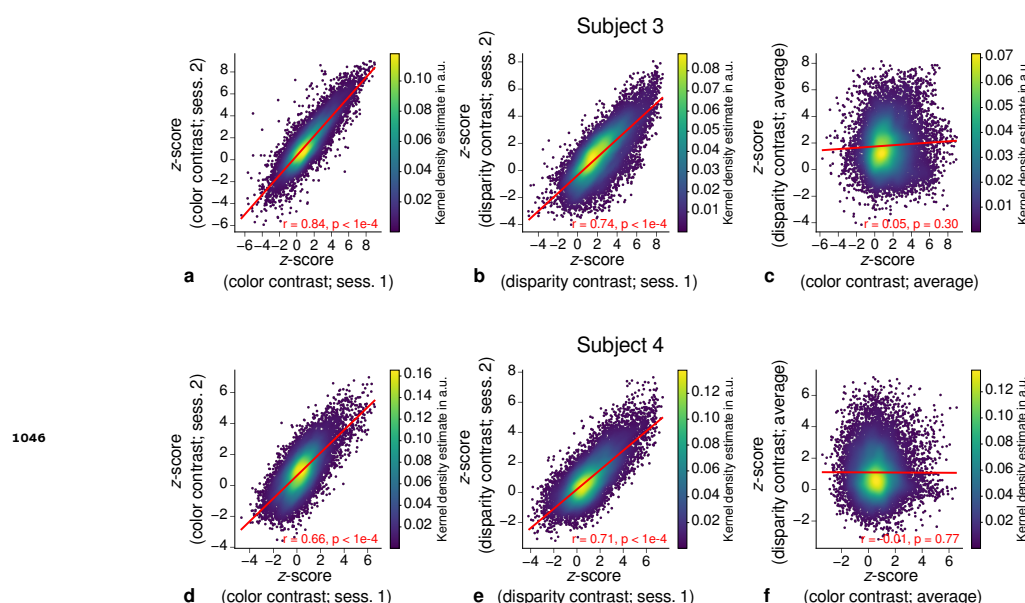
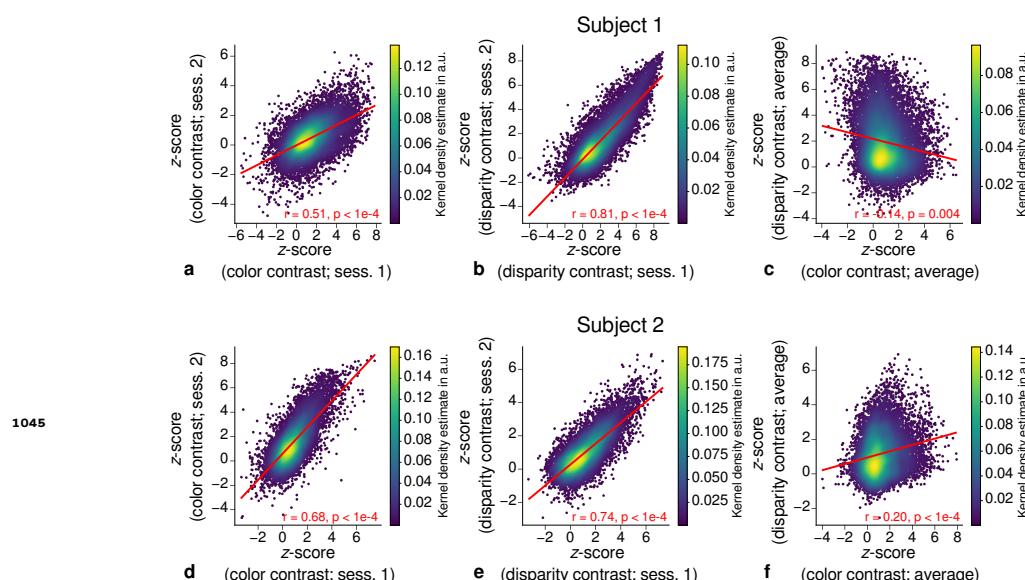


Figure 2-Figure supplement 2. Exemplary activation maps from another participant. Activation maps are shown for another representative participant (subject 4) to further illustrate the alternating nature of color- and disparity-selective stripes. (a)–(b) show color- and disparity-selective stripes for the ventral aspect of V2 on the left hemisphere, respectively. (c)–(d) show corresponding maps for the dorsal aspect of V2 on the right hemisphere. Cyan outlines emphasize activation clusters in (a) and (c) to illustrate the alternating pattern between color-selective thin and disparity-selective thick stripes in (b) and (d), respectively. Other details as in Figure 2. Scale bar: 5 mm.



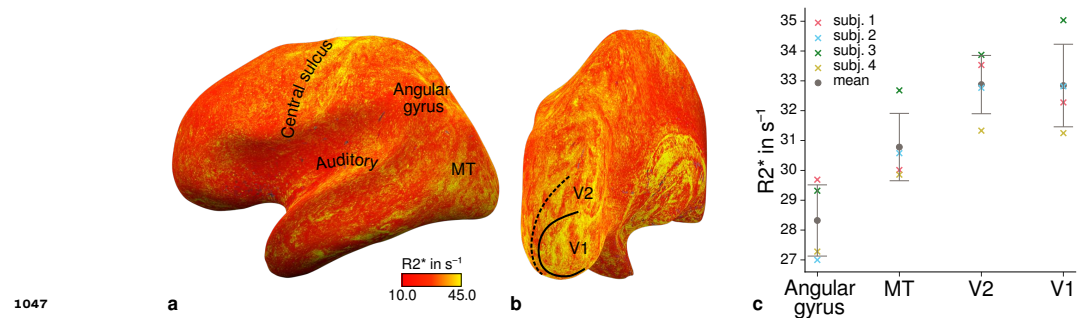


Figure 4-Figure supplement 1. Illustration of quantitative R_2^* across cortical areas. Cortical R_2^* values are shown at mid-cortical depth of the left hemisphere on an inflated surface from a representative participant (subject 3) in lateral (a) and posterior (b) view. Higher values in primary motor and sensory areas are less pronounced than for R_1 shown in Figure 4. Data from all participants can be found in Appendix 1. (c) Mean R_2^* values are shown for different cortical regions (angular gyrus, MT, V2, V1). Overall, higher R_2^* in early visual cortex can be qualitatively seen similar to Figure 4c. Corresponding source data is given in Figure 4-source data 1. Other details as in Figure 4c.

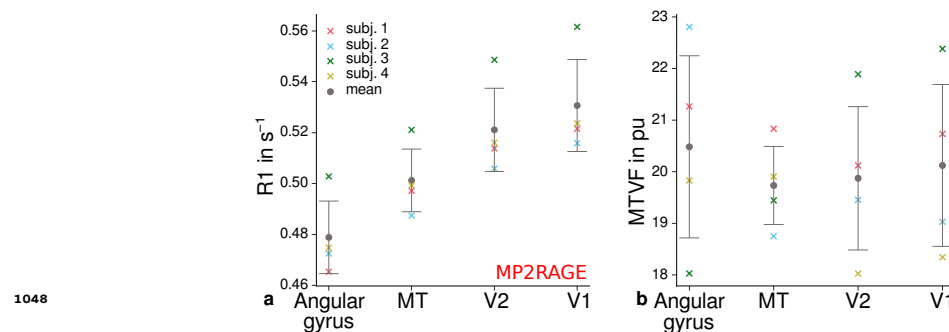


Figure 4-Figure supplement 2. Mean quantitative R_1 (MP2RAGE) and PD values across cortical areas. Mean R_1 (MP2RAGE) (a) and $MTVF = 100\% - PD$ (macromolecular tissue volume fraction (Mezer et al., 2013)) (b) values are shown for different cortical regions (angular gyrus, MT, V2, V1). (a) shows higher R_1 values in early visual cortex and especially V1 and therefore qualitatively reproduces the results shown in Figure 4c. No consistent trend across participants was found for MTVF as shown in (b), which was the reason to exclude PD from further analyses. Corresponding source data is given in Figure 4-source data 1. Other details as in Figure 4c.

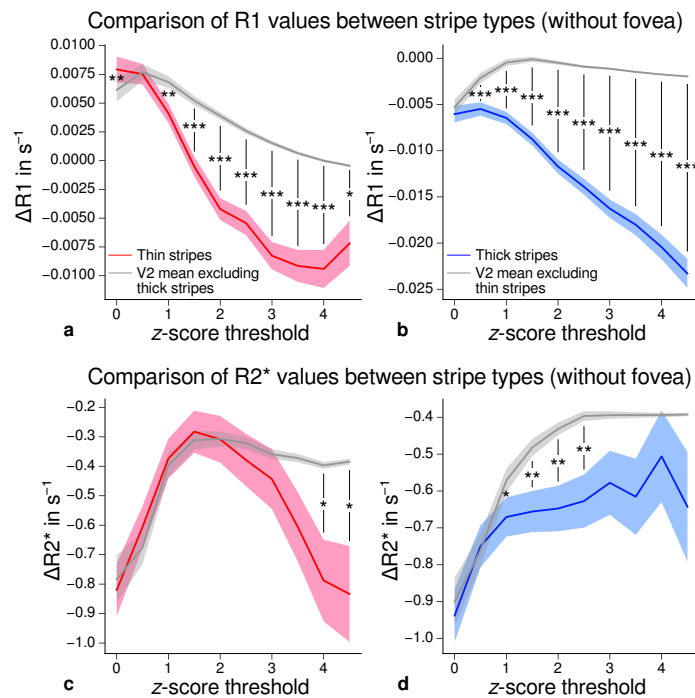


Figure 5-Figure supplement 1. Comparison of quantitative R_1 and R_2^* values between V2 stripe types excluding foveal parts. Cortical R_1 (a)-(b) and R_2^* (c)-(d) values in thin stripes (red), thick stripes (blue) and V2 excluding the other stripe type plus the region around the foveal representation (gray; and therefore containing contributions from pale stripes) are shown for various z-score threshold levels, which were used to define thin and thick stripe ROIs. R_1 is lower in both thin and thick stripes than surrounding gray matter, which confirms the results of the main analysis illustrated in **Figure 5**. This supporting analysis was done to exclude the possibility of bias in the main analysis coming from the stripe ROI definition, which was based on functional activation maps that generally showed absence of activation around the foveal representation. Statistically significant differences are marked by asterisks, *: $p < 0.05$, **: $p < 0.01$, ***: $p < 0.001$. Shaded areas indicate 1 standard deviation of the generated null distribution used for permutation testing. Other details as in **Figure 5**.

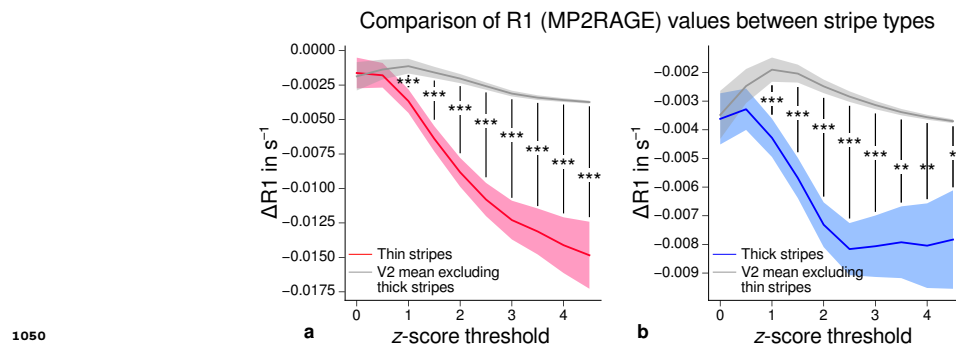


Figure 5-Figure supplement 2. Comparison of quantitative R_1 values (MP2RAGE) between V2 stripe types. Cortical R_1 values in thin stripes (red), thick stripes (blue) and whole V2 excluding the other stripe type (gray; and therefore containing contributions from pale stripes) are shown for various z-score threshold levels, which were used to define thin and thick stripe ROIs. R_1 values are based on a separate data set using the MP2RAGE sequence. Lower values were found in thin (**a**) and thick stripes (**b**) when compared to surrounding gray matter, which confirms the results of the main analysis illustrated in **Figure 5**. Statistically significant differences are marked by asterisks, *: $p < 0.05$, **: $p < 0.01$, ***: $p < 0.001$. Shaded areas indicate 1 standard deviation of the generated null distribution used for permutation testing. Other details as in **Figure 5**.

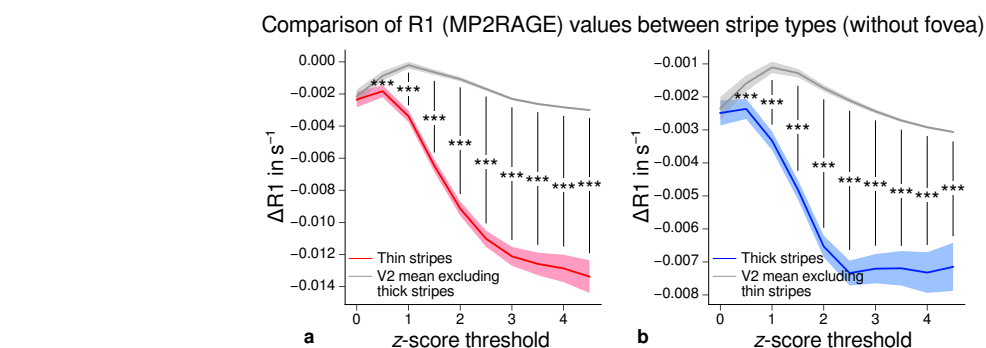


Figure 5-Figure supplement 3. Comparison of quantitative R_1 values (MP2RAGE) between V2 stripe types excluding foveal parts. Cortical R_1 values in thin stripes (red), thick stripes (blue) and V2 excluding the other stripe type plus the region around the foveal representation (gray; and therefore containing contributions from pale stripes) are shown for various z-score threshold levels, which were used to define thin and thick stripe ROIs. R_1 values are based on a separate data set using the MP2RAGE sequence. Lower values were found in thin (**a**) and thick stripes (**b**) when compared to surrounding gray matter, which confirms the results of the main analysis illustrated in **Figure 5** and the supporting analysis which covered whole V2 shown in **Figure 5-Figure Supplement 2**. Statistically significant differences are marked by asterisks, *: $p < 0.05$, **: $p < 0.01$, ***: $p < 0.001$. Shaded areas indicate 1 standard deviation of the generated null distribution used for permutation testing. Other details as in **Figure 5**.