Reconstructing imagined letters from early visual cortex reveals tight topographic correspondence between visual mental imagery and perception

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Abstract
Visual mental imagery is the quasi-perceptual experience of “seeing in the mind’s eye”. While a tight correspondence between imagery and perception in terms of subjective experience is well established, their correspondence in terms of neural representations remains insufficiently understood. In the present study, we exploit the high spatial resolution of functional magnetic resonance imaging (fMRI) at 7T, the retinotopic organization of early visual cortex, and machine-learning techniques to investigate whether visual imagery of letter shapes preserves the topographic organization of perceived shapes. Sub-millimeter resolution fMRI images were obtained from early visual cortex in six subjects performing visual imagery of four different letter shapes. Predictions of imagery voxel activation patterns based on a population receptive field-encoding model and physical letter stimuli provided first evidence in favor of detailed topographic organization. Subsequent visual field reconstructions of imagery data based on the inversion of the encoding model further showed that visual imagery preserves the geometric profile of letter shapes. These results open new avenues for decoding, as we show that a denoising autoencoder can be used to pretrain a classifier purely based on perceptual data before fine-tuning it on imagery data. Finally, we show that the autoencoder can project imagery-related voxel activations onto their perceptual counterpart allowing for visually recognizable reconstructions even at the single-trial level. The latter may eventually be utilized for the development of content-based BCI letter-speller systems.

Introduction
Visual mental imagery refers to the fascinating phenomenon of quasi-perceptual experiences in the absence of external stimulation (Thomas 1999). The capacity to imagine has important cognitive implications and has been linked to working memory, problem solving, and creativity (Albers et al. 2013; Kozhevnikov et al. 2013). Yet, the nature of mental representations underlying imagery remains controversial. It has been argued that visual imagery is pictorial, with an intrinsic spatial organization resembling that of physical images (Kosslyn et al. 1997, 2006). Others have claimed that imagery resembles linguistic descriptions, lacking any inherent spatial properties (Pylyshyn 1973, 2003; Brogaard and Gatzia 2017). This debate has become increasingly informed by neuroimaging. For instance, several functional magnetic resonance imaging (fMRI) studies have indicated that imagery activates cortical networks that are also activated during corresponding perceptual tasks (Kosslyn et al. 1997; Goebel et al. 1998; Ishai et al. 2000; O’Craven and Kanwisher 2000; Ganis et al. 2004; Mechelli et al. 2004), lending credence to the notion that imagery resembles perception. Applying multi-voxel pattern analyses (MVPA), furthermore, enabled the decoding of feature-specific imagery content related to orientations (Harrison and Tong 2009; Albers et al. 2013), motion (Emmerling et al. 2013), and color (Goebel et al. 2006).
such as attention (Naselaris et al. 2015). To overcome this classification may have resulted from confounding factors of low-level visual features, leaving open the possibility that grounds that it does not rely on an explicit encoding model. Such correspondence opens new avenues for decoding. Specifically, they used the estimated encoding model to identify an imagined artwork from a set of candidates by comparing voxel activations empirically observed in response to imagery with those predicted from encoding each candidate (Naselaris et al. 2015).

While this study constitutes a major methodological advancement and largely defuses the aforementioned confounds, a complex encoding model allows only for limited inferences regarding the similarity of perception and imagery with respect to any particular feature. It is, for instance, conceivable that the largest contributor to image identification stemmed from an unspecified top–down modulation of salient regions in the imaged artwork with crude retinotopic organization. That is, activations in response to mental imagery might have been co-localized to highly salient regions of the image (without otherwise resembling it) and this might have been sufficient for image identification.

Indeed, results from studies reconstructing the visual field from fMRI data leveraging the retinotopic organization of early visual cortex give the impression that the retinotopic organization of mental imagery is rather diffuse. For instance, while seminal work has been conducted detailing the ability to obtain straightforwardly recognizable reconstructions of perceived physical stimuli (Thirion et al. 2006; Miyawaki et al. 2008; Schoenmakers et al. 2013); similar successes have not been repeated for imagery. Retinotopy-based reconstructions of imagined shapes have so far merely been co-localized with the region of the visual field, where they were imagined but bore no visual resemblance to their geometry (Thirion et al. 2006).

However, unless imagery of an object preserves the object’s geometry, it unlikely it would preserve any of its more fine-grained features. It is thus pivotal to empirically establish precise topographic correspondence between imagery and perception. Utilizing the high spatial resolution offered by 7T fMRI and the straightforwardly invertible population receptive field model (Dumoulin and Wandell 2008), we provide new evidence that imagery-based reconstructions of letter shapes are recognizable and preserve their physical geometry. This supports the notion of tight topographic correspondence in early visual cortex. Such correspondence opens new avenues for decoding. Specifically, we show that using a denoising autoencoder, it is possible to pretrain a classifier, intended to decode imagery content, purely based on easily obtainable perceptual data. Only fine-tuning of the classifier requires (a small amount of) additional imagery data. Finally, we show that an autoencoder can project imagery-related voxel activations onto their perceptual counterpart allowing for recognizable reconstructions even at a single-trial level. The latter could open new frontiers for brain–computer interfaces (BCIs).

Materials and methods

Participants

Six participants (2 female, age range = (21–49), mean age = 30.7) with normal or corrected-to-normal visual acuity took part in this study. All participants were experienced in undergoing high-field fMRI experiments, gave written informed consent, and were paid for participation. All procedures were conducted with approval from the local Ethical Committee of the Faculty of Psychology and Neuroscience at Maastricht University.

Stimuli and tasks

Each participant completed three training sessions to practice the controlled imagery of visual letters prior to a single scanning session which comprised four experimental (imagery) runs of ~ 11 min and one control (perception) run of ~ 9 min as well as one pRF mapping run of ~ 16 min.

Training session and task

Training sessions lasted ca. 45 min and were scheduled 1 week prior to scanning. Before the first training session, participants filled in the Vividness of Visual Imagery Questionnaire (VVIQ; Marks, 1973) and the Object–Spatial Imagery and Verbal Questionnaire (Blazhenkova and Kozhevnikov 2009). These questionnaires measure the subjective clearness and vividness of imagined objects and cognitive styles during mental imagery, respectively. In each training trial, participants saw one of four white letters (‘H’, ‘T’, ‘S’, or ‘C’) enclosed in a white square guide box (8° by 8° visual angle) on grey background and a red fixation dot in the center of the screen (see Fig. 1). With the onset of the visual stimulation, participants heard a pattern of three low tones (note C5) and one high tone (note G5) that lasted 1000 ms. This tone pattern was associated with the visually presented letter with specific patterns randomly assigned for each participant. After 3000 ms, the letter started to fade out until it completely disappeared at 5000 ms after trial onset. The fixation dot then turned orange and participants were...
instructed to maintain a vivid image of the presented letter. After an 18 s imagery period, the fixation dot turned white and probing started. With an inter-probe-interval of 1500 ms (jittered by ± 200 ms), three white probe dots appeared within the guide box. These dots were located within the letter shape or outside of the letter shape (however, always within the guide box). Participants were instructed to indicate by button press whether a probe was located inside or outside the imagined letter shape (Podgorny and Shepard 1978). Depending on the response, the fixation dot turned red (incorrect) or green (correct) before turning white again as soon as the next probe was shown. The positions of the probe dots were randomly chosen, such that they had a minimum distance of 0.16° and a maximum distance of 0.32° of visual angle from the edges of the letter (and the guide box), both for inside and outside probes. This ensured similar task difficulty across trials. A resting phase of 3000 ms or 6000 ms followed the three probes. At the beginning of a training run, all four letters were presented for 3000 ms each, alongside the associated tone pattern (reference phase). During one training run, each participant completed 16 pseudo-randomly presented trials. In each training session, participants completed two training runs during which reference letters were presented in each trial (described above) and two training runs without visual presentation (i.e., the tone pattern was the only cue for a letter). Participants were instructed to maintain central fixation throughout the entire run. After the training session, participants verbally reported the imagery strategies they used.

**Imagery runs**

Imagery runs were similar to the training task with changes to the probing phase and the timing of the trial phase. After the reference phase in the beginning of each run, there was no visual stimulation other than the fixation dot and the guide box. Imagery phases started when participants heard the tone pattern and the fixation dot turned orange. Imagery phases lasted 6 s. Participants were instructed to imagine the letter associated with the tone pattern as vividly and accurately as possible. The guide box aided the participant by acting as a reference for the physical dimensions of the letter. The resting phases that followed each imagery phase lasted 9 s or 12 s. There was no probing phase in normal trials. In each experimental run, there were 32 normal trials and two additional catch trials which entailed a probing phase of four probes. There was no visual feedback for the responses in the probing phase (the fixation dot remained white). Data from the catch trials were not included in the analyses.

**Perception run**

To measure brain activation patterns in visual areas during the perception of the letters used in the imagery runs, we recorded one perception run during the scanning session. The four letters were visually presented using the same trial timing parameters as in the experimental runs. There were neither reference nor probing phases. Letters were presented for the duration of the imagery phase (6 s) and their shape was filled with a flickering checkerboard pattern (10 Hz). No tone patterns were played during the perception run. The recorded responses were also used to train a denoising autoencoder (see below).

**pRF mapping**

A bar aperture (1.33° wide) revealing a flickering checkerboard pattern (10 Hz) was presented in four orientations. For each orientation, the bar covered the entire screen in 12 discrete steps (each step lasting 3 s). Within each orientation, the sequence of steps (and hence of the locations) was randomized (cf. Senden et al. 2014). Each orientation was presented six times.

**Stimulus presentation**

The bar stimulus used for pRF mapping was created using the open source stimulus presentation tool BrainStim (http://svengijsen.github.io/BrainStim/). Visual and auditory stimulation in the imagery and perception runs were controlled with PsychoPy (version 1.83.03; Peirce 2007). Visual stimuli were projected on a frosted screen at the top end of the scanner table by means of an LCD projector (Panasonic, No
PT- EZ57OEL; Newark, NJ, USA). Auditory stimulation was presented using MR-compatible insert earphones (Sennheiser, Model S14; Malden, MA, USA). Responses to the probes were recorded with MR-compatible button boxes (Current Designs, 8-button response device, HHSC-2×4-C; Philadelphia, USA).

**Magnetic resonance imaging**

We recorded anatomical and functional images with a Siemens Magnetom 7 T scanner (Siemens; Erlangen, Germany) and a 32-channel head-coil (Nova Medical Inc.; Wilmington, MA, USA). Prior to functional scans, we used a T1-weighted magnetization prepared rapid acquisition gradient echo (Marques et al. 2010) sequence [240 sagittal slices, matrix = 320 320, voxel size = 0.7 by 0.7 by 0.7 mm³, first inversion time TI1 = 900 ms, second inversion time TI2 = 2750 ms, echo time (TE) = 2.46 ms, repetition time (TR) = 5000 ms, first nominal flip angle = 5°, and second nominal flip angle = 3°] to acquire anatomical data. For all functional runs, we acquired high-resolution gradient echo (T2* weighted) echo-planar imaging (Moeller et al. 2010) data (TE = 26 ms, TR = 3000 ms, generalized auto-calibrating partially parallel acquisitions (GRAPPA) factor = 3, multi-band factor = 2, nominal flip angle = 55°, number of slices = 82, matrix = 186 by 186, and voxel size = 0.8 by 0.8 by 0.8 mm³). The field-of-view covered occipital, parietal, and temporal areas. In addition, before the first functional scan, we recorded five functional volumes with opposed phase encoding directions to correct for EPI distortions that occur at higher field strengths (Andersson et al. 2003).

**Processing of (f)MRI data**

We analyzed anatomical and functional images using BrainVoyager 20 (version 20.0; Brain Innovation; Maastricht, The Netherlands) and custom code in MATLAB (version 2017a; The Mathworks Inc.; Natick, MA, USA). We interpolated anatomical images to a nominal resolution of 0.8 mm isotropic to match the resolution of functional images. In the anatomical images, the grey/white matter boundary was detected and segmented using the advanced automatic segmentation tools of BrainVoyager 20 which are optimized for high-field MRI data. A region-growing approach analyzed local intensity histograms, corrected topological errors of the segmented grey/white matter border, and finally reconstructed meshes of the cortical surfaces (Kriegeskorte and Goebel 2001; Goebel et al. 2006). The functional images were corrected for motion artefacts using the 3D rigid body motion correction algorithm implemented in BrainVoyager 20 and all functional runs were aligned to the first volume of the first functional run. We corrected EPI distortions using the COPE (“Correction based on Opposite Phase Encoding”) plugin of BrainVoyager that implements a method similar to that described in Andersson, Skare, and Ashburner (Andersson et al. 2003) and the “topup” tool implemented in FSL (Smith et al. 2004). The pairs of reversed phase encoding images recorded in the beginning of the scanning session were used to estimate the susceptibility-induced off-resonance field and correct the distortions in the remaining functional runs. After this correction, functional data were high-pass filtered using a general linear model (GLM) Fourier basis set of three cycles sine/cosine, respectively. This filtering included a linear trend removal. Finally, functional runs were co-registered and aligned to the anatomical scan using an affine transformation (9 parameters) and z-normalized to eliminate signal offsets and inter-run variance.

**pRF mapping and region-of-interest definition**

For each subject, we fit location and size parameters of an isotropic Gaussian population receptive field model (Dumoulin and Wandell 2008) by performing a grid search. In terms of pRF location, the visual field was split into a circular grid of 100 by 100 points, whose density decays exponentially with eccentricity. Receptive field size exhibits a linear relationship with eccentricity with the exact slope depending on the visual area (Freeman and Simoncelli 2011). For this reason, we explored slopes in the range from 0.1 to 1 (step = 0.1), as this effectively allows for exploration of a greater range of receptive field sizes (10 for each unique eccentricity value). We used the pRF mapping tool from the publicly available Computational Neuroimaging Toolbox (https://github.com/MSend/CNI_toolbox). Polar angle maps resulting from pRF fitting were projected onto inflated cortical surface reconstructions and used to define regions-of-interest (ROIs) for bilateral visual areas V1, V2, and V3. The resulting surface patches from the left and right hemisphere were projected back into volume space (from −1 mm to +3 mm from the segmented grey/white matter boundary). Volume ROIs were then defined for V1, V2, V3, and a combined ROI (V1V2V3).

**Voxel patterns**

All our analyses and reconstructions are based on letter-specific spatial activation profiles of voxel co-activations; i.e., voxel patterns. Voxel patterns within each ROI were obtained for both perceptual and imagery runs. First, for each run, single-trial letter-specific voxel patterns were obtained by averaging BOLD activations in the range from +2 until +3 volumes following trial onset and z-normalizing the result. This lead to a total of eight (one per trial) perceptual and 32 (four runs with 8 trials each) imagery voxel patterns per letter. We, furthermore, computed perceptual and imagery average voxel patterns per letter by averaging.
over all single-trial patterns (and runs in case of imagery) of a letter and z-normalizing the result. Imagery average voxel patterns were used in an encoding analysis and for assessment of reconstruction quality, while perceptual average patterns were used for training a denoising autoencoder (Vincent et al. 2008).

Encoding analysis

To test the hypothesis that spatial activation profiles of visual mental imagery are geometry-preserving, we tested whether voxel activations predicted from the encoding model (one isotropic Gaussian per voxel) and a physical (binary) stimulus corresponding to the imagined letter provides a significantly better fit with measured voxel activations than predictions from the remaining binary letter stimuli. Specifically, for each participant and ROI, we predicted voxel activations for each of the four letters based on pRF estimates and physical letter stimuli.

Autoencoder

We trained an autoencoder with a single hidden layer $k = \lfloor 0.1 \cdot N_{\text{voxels}} \rfloor$ to reproduce average perceptual voxel patterns from noise-corrupted versions per subject and ROI. Since the values of voxel patterns follow a Gaussian distribution with a mean of zero and unit standard deviation, we opted for zero-mean additive Gaussian noise with a standard deviation $\sigma = 12$ for input corruption. Note that the exact value of $\sigma$ is not important as long as it sufficiently corrupts the data. We achieved similar results with values in the range $[8, 14]$. The hidden layer consisted of units with rectified linear activation functions. Output units activated linearly. Encoding weights (from input to hidden layer) and decoding weights (from hidden to output layer) were shared.

The input, hidden, and output layers were,

taken together, the input, hidden, and output layers were, respectively, given by

$$y_c = y + \epsilon, \text{ with } \epsilon \sim \mathcal{N}(0, \sigma)$$

$$h = \phi(W_c y_c + b_c), \text{ with } \phi(x) = \frac{1}{1 + e^{-x}}$$

$$y_r = W_d h + b_d, \text{ with } W_d = W_c^T$$

In Eq. 1, $y$ is a voxel pattern (of length $v$), $y_c$ its noise corruption, and $y_r$ its restoration. $W_c$ (k-by-v matrix) and $W_d$ (v-by-k matrix) are the tied encoding and decoding weights, respectively. Finally, $b_c$ (k-by-1 vector) and $b_d$ (v-by-1 vector) are the biases of the hidden and output layers, respectively. We used mean squared distances to measure loss between the input and its restoration and implemented the autoencoder in the TensorFlow library (Abadi et al. 2016) for Python (version 2.7, Python Software Foundation, https://www.python.org/). The autoencoder was trained using the Adam optimizer (Kingma and Ba 2014) with a learning rate of $1 \times 10^{-5}$ and a batch size of 100 for 2000 iterations. In addition to the four average perceptual voxel patterns, we also included an equal amount of noise-corrupted mean luminance images to additionally force reconstructions to zero if the input contained no actual signal. No imagery data were used for training the autoencoder.

Reconstruction

For each subject and ROI, we reconstructed the visual field from average perceptual and imagery voxel patterns. We obtained weights mapping the cortex to the visual field by inverting the mapping from visual field to cortex given by the population receptive fields. Since $W_{pRF}$, a v-by-p matrix (with v being the number of voxels and p the number of pixels) mapping a 150-by-150 pixel visual field to the cortex (i.e., $p = 22500$ pixels; after vectorizing the visual field) is not invertible, we minimize the error function:

$$E = (y - W_{pRF} x)^T (y - W_{pRF} x) + D ||x||_2^2$$

with respect to the input image $x$ (a vector of length p). The vector $y$ is of length v and reflects a measured voxel pattern. Finally, D is a diagonal matrix of the outdegree of each pixel in the visual field which provides pixel-specific scaling of the L2 regularization term $||x||_2^2$ and accounts for cortical magnification. Minimizing Eq. 2 leads to the expression:

$$x = (W_{pRF}^T W_{pRF} + D)^{-1} W_{pRF}^T y$$

with which we can reconstruct the visual field from voxel patterns. To minimize computational cost, we compute the projection matrix $W_{VF} = \left( W_{pRF}^T W_{pRF} + D \right)^{-1} W_{pRF}^T$ once per ROI and subject rather than performing costly matrix inversion for every reconstruction. Note that both raw voxel patterns ($y$) as well as restored voxel patterns ($y_r$) obtained from passing $y$ through the autoencoder, can be used for image reconstruction. In the former case, $x = W_{VF} y$. In the latter case, $x = W_{VF} y_r = W_{VF} \left[ W_d \phi(W_c y + b_c) + b_d \right]$.

For each letter, we assessed the quality of its reconstruction by calculating the correlation between the reconstruction and the corresponding binary letter stimulus. This constitutes a first-level correlation metric. However, since the four letters bear different visual similarities with each other (e.g., ‘S’ and ‘C’ might resemble each other more closely than either resemble ‘H’), we also defined a second-level correlation metric. Specifically, we obtained one vector of all pairwise correlations between physical letter stimuli and a second vector of pairwise correlations between corresponding reconstructions and correlated these two vectors.
Classification

We replaced the output layer of the pretrained autoencoder with a four-unit (one for each letter) softmax classifier. Weights from the hidden to the classification layer as well as the biases of output units were then trained to classify single-trial imagery voxel patterns using cross entropy as a measure of loss. Note that pretrained weights from input to hidden layer ($W_h$ in Eq. 1) as well as pretrained hidden unit biases ($b_h$ in Eq. 1) remained fixed throughout training of the classifier. These weights and biases were thus dependent purely on perceptual data. This procedure is equivalent to performing multinomial logistic regression on previously established hidden layer representations. Imagery runs were split into training and testing data sets in a leave-one-run-out procedure, such that the classifier was repeatedly trained on a total of 96 voxel patterns (8 trials per 4 letters for each of three runs) and tested on the remaining 32 voxel patterns. We again trained the network using the Adam optimizer. However, in this case, the learning rate was $1 \times 10^{-4}$, the batch size equal to 96, and training lasted merely 250 iterations.

Statistical analysis

Statistical analyses were performed using MATLAB (version 2017a; The Mathworks Inc.; Natick, MA, USA). We used a significance level of $\alpha = 0.05$ (adjusted for multiple comparisons where appropriate) for all statistical analyses.

Behavioral results were analyzed using repeated-measures ANOVA with task (visible or invisible runs) and time as within-subject factors.

For the encoding analysis, we performed a mixed-model regression for the average voxel activations of each imagined letter within each ROI with physical letter as fixed and participant as random factors, respectively. This was followed by a contrast analysis. For each imagined letter, the contrast was always between the corresponding physical stimulus and all remaining physical stimuli. For example, when considering voxel activations for the imagined letter ‘H’, a weight of 3 was placed on activations predicted from the physical letter ‘H’ and a weight of $-1$ was placed on activations predicted from each of the remaining three letters. Since we repeated the analysis for each imagined letter (4) and single region ROI (3), we performed a total of 12 tests and considered results significant at a corrected cutoff of $\alpha_c = 0.05/12 = 0.0042$.

To evaluate which factors contribute most to first-level reconstruction quality, we used the same approach omitting letter as a predictor.

To assess the significance of classification results, we evaluated average classification accuracy across the four runs against a Null distribution obtained from 1000 permutations of a leave-one-run-out procedure with randomly scrambled labels. We performed this analysis separately for each subject and ROI and consider accuracy results significant if they exceed the 95th percentile of the Null distribution. To statistically evaluate which factors contribute most to classification accuracy, we performed mixed-model regression with the VVIQ and the OSIVQ spatial and OSIVQ object scores, ROI (using dummy coding, V1 = reference), letter (dummy coding, ‘H’ = reference), and number of selected voxels (again grouped by ROI).

Results

Behavioral results

VVIQ and OSIVQ scores for each participant are shown in Fig. 2. The average score over participants for VVIQ was 4.07 (95% CI [3.71, 4.43]). For the object, spatial, and verbal sub-scales of OSIVQ, average scores were 2.88 (95% CI [2.48, 3.27]), 3.08 (95% CI [2.75, 3.41]), and 3.81 (95% CI [3.33, 4.29]), respectively. Participants reported that they tried to maintain the afterimage of the fading stimulus as a strategy to enforce vivid and accurate letter imagery. Furthermore, participants determined through button presses whether a probe was located inside or outside the letter shape, while the letter was either visible or imagined. A
repeated-measures ANOVA with task (visible or invisible runs) and time as within-subject factors revealed a statistically significant effect of time on probing accuracy \( (F_{(2,10)} = 19.84, p < 0.001) \), and no significant difference for task \( (F_{(1,5)} = 1.10, p = 0.341) \) (Table 1).

## Encoding

For each imagined letter (H, T, S, C) in each single-area ROI (V1, V2, V3), we investigated whether spatial voxel activation profiles can be predicted from a Prf-encoding model and the corresponding physical stimulus. That is, for each imagined letter–ROI combination, we ran a mixed-model regression with observed imagery voxel activations (averaged over trials and runs) as outcome variable, predicted voxel activations for each physical letter stimulus as predictors and participants as grouping variable. Since we were specifically interested in testing our hypothesis that the retinotopic organization of imagery voxel activations is sufficiently geometrically specific to distinguish among different imagined letters, we performed contrast analyses between the physical letter corresponding to the imagery and all the remaining letters (see “Methods” for details). Contrasts were significant after applying Bonferroni correction \( (\alpha_c = 0.0042) \) for each of the twelve letter-ROI combinations.

In other words, predictions based on a specific physical letter gave a better account of voxel activations observed for the imagery of that specific letter than those based on every other physical letter, as can be appreciated from Table 2. Differences between subjects might be due to differences in their ability to imagine shapes accurately and vividly as measured by the VVIQ and OSIVQ questionnaires. Differences between ROIs might be due to differences with respect to their retinotopy (most likely receptive field sizes) or due to different numbers of voxels included for analysis of each ROI. Only the former would be a true ROI effect. We investigate which factors account for observed correlations (transformed to Fisher z-scores for analyses) by performing a mixed-model regression with questionnaire scores, ROI (using dummy coding, V1 = reference), letter (dummy coding, ‘H’ = reference), and number of selected voxels as predictors. A number of voxels were grouped by ROI. Furthermore, the regression model included the VVIQ and the OSIVQ spatial and object scores. However, the VVIQ score was not included since it correlated highly with the OSIVQ verbal score (leading to collinearity). To further prevent collinearity, we also only included single-area ROIs in this analysis and not the combined ROI. A number of voxels \( [t_{(62)} = 2.59, p = 0.012] \) and the OSIVQ object score \( [t_{(62)} = 2.64, p = 0.010] \) were significant quantitative predictors. Furthermore, letter was a significant categorical predictor. Specifically, letter ‘T’ \( [t_{(62)} = 5.58, p < 0.001] \) presented with significantly improved correlation values over the reference letter ‘H’, whereas letters ‘S’ \( [t_{(62)} = -3.88, p = 0.0003] \) and ‘C’ \( [t_{(62)} = -2.25, p = 0.028] \) presented with significantly decreased correlation values with respect to the reference. Neither the OSIVQ verbal score \( [t_{(62)} = 0.0278, p = 0.978] \) nor the ROI were significant predictors of reconstruction quality.

Next, we examined the second-level correlation metric of reconstruction quality. Correlations between physical and reconstruction pairwise first-level correlation vectors were 0.60 (95% CI [0.28, 0.80], \( p = 0.103 \)) for V1, 0.65

## Reconstruction

### Raw imagery data

We reconstructed the visual field from average imagery voxel patterns in response to each letter (see Figs. 4, 5). Mean correlations between reconstructed imagery and physical letters are presented in Table 3 (for comparison, Table 4 shows correlations between reconstructed perception and physical letters). As can be appreciated from these results as well as the figures, first-level reconstruction quality varies across ROIs as well as across subjects. Differences between subjects might be due to differences in their ability to imagine shapes accurately and vividly as measured by the VVIQ and OSIVQ questionnaires. Differences between ROIs might be due to differences with respect to their retinotopy (most likely receptive field sizes) or due to different numbers of voxels included for analysis of each ROI. Only the former would be a true ROI effect. We investigate which factors account for observed correlations (transformed to Fisher z-scores for analyses) by performing a mixed-model regression with questionnaire scores, ROI (using dummy coding, V1 = reference), letter (dummy coding, ‘H’ = reference), and number of selected voxels as predictors. A number of voxels were grouped by ROI. Furthermore, the regression model included the VVIQ and the OSIVQ spatial and object scores. However, the VVIQ score was not included since it correlated highly with the OSIVQ verbal score (leading to collinearity). To further prevent collinearity, we also only included single-area ROIs in this analysis and not the combined ROI. A number of voxels \( [t_{(62)} = 2.59, p = 0.012] \) and the OSIVQ object score \( [t_{(62)} = 2.64, p = 0.010] \) were significant quantitative predictors. Furthermore, letter was a significant categorical predictor. Specifically, letter ‘T’ \( [t_{(62)} = 5.58, p < 0.001] \) presented with significantly improved correlation values over the reference letter ‘H’, whereas letters ‘S’ \( [t_{(62)} = -3.88, p = 0.0003] \) and ‘C’ \( [t_{(62)} = -2.25, p = 0.028] \) presented with significantly decreased correlation values with respect to the reference. Neither the OSIVQ verbal score \( [t_{(62)} = 0.0278, p = 0.978] \) nor the ROI were significant predictors of reconstruction quality.

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### Table 1 Probing accuracy (averages over participants and time)

|       | T1                   | T2                   | T3                   |
|-------|----------------------|----------------------|----------------------|
| Visible| 60.42 (95% CI [48.2, 72.64]) | 75.39 (95% CI [66.70, 84.08]) | 77.73 (95% CI [69.36, 86.10]) |
| Invisible| 62.02 (95% CI [44.57, 79.45]) | 73.18 (95% CI [65.98, 80.38]) | 81.57 (95% CI [75.47, 86.10]) |

### Table 2 Contrast analysis comparing the physical letter corresponding to the imagery with all remaining letters

|       | H | T | S | C |
|-------|---|---|---|---|
| V1 t(21) | 32.11, p = 0.0004 | t(21) = 48.00, p = 0.0002 | t(21) = 14.10, p = 0.0025 | t(21) = 29.84, p = 0.0006 |
| V2 t(21) | 25.21, p = 0.0008 | t(21) = 67.63, p = 0.0001 | t(21) = 19.64, p = 0.0013 | t(21) = 47.48, p = 0.0002 |
| V3 t(21) | 47.90, p = 0.0006 | t(21) = 27.60, p = 0.0007 | t(21) = 11.48, p = 0.0038 | t(21) = 32.83, p = 0.0005 |

Each of the 12 letter–ROI combinations was significant after applying Bonferroni correction \( (\alpha_c = 0.0042) \).
(95% CI [0.34, 0.83], \( p = 0.082 \)) for V2, 0.48 (95% CI [0.15, 0.71], \( p = 0.167 \)) for V3, and 0.64 (95% CI [0.34, 0.83], \( p = 0.084 \)) for V1V2V3, respectively. Finally, we performed a mixed regression to assess which factors account for the observed correlations (again transformed to Fisher z-scores). We included OSIVQ verbal, spatial, and object scores, ROI (dummy coding, V1 = reference), and number of selected voxels (grouped by ROI) as predictors. The OSIVQ object score \( t(11) = 3.26, p = 0.0076 \) and number of voxels \( t(11) = 3.94, p = 0.0023 \) significantly predicted second-level correlations, while the spatial \( t(11) = 0.71, p = 0.492 \) and verbal scores \( t(11) = -0.81, p = 0.436 \) did not. Furthermore, there was significant effect of ROI, since neither V2 \( t(11) = -1.56, p = 0.148 \) nor V3 \( t(11) = 0.40, p = 0.697 \) significantly differed from V1.

**Processed imagery data**

Our results confirm that visual mental imagery preserves perceptual topographic organization. This can be leveraged to obtain improved reconstructions of mental imagery. Specifically, an autoencoder trained to retrieve perceptual voxel patterns from their noise-corrupted version can be utilized to enhance imagery data. Figure 6 shows how the autoencoder affects first-level reconstruction quality on a single-trial basis for V1. As shown in the figure, reconstruction quality was best for ‘T’, followed by ‘H’, ‘C’, and ‘S’. A subject effect is also clearly apparent with participants three and five generally displaying the best results. Finally, imagery reconstruction quality was generally inferior to perception prior to using the autoencoder. However, using the autoencoder pushed imagery reconstruction quality towards perception levels. Indeed, the autoencoder maps imagery voxel patterns onto the corresponding perception voxel patterns it has learned previously. This explains two important observations. First, for some trials, using the autoencoder decreased resemblance to the physical letter. This is especially apparent for participants four and six, whose reconstructions were generally not particularly good. Such decrements in reconstruction quality result from imagery voxel patterns in response to one letter falling within the attraction domain of another letter (resembling the activation pattern of that letter slightly more) and hence get mapped onto the wrong pattern.
Second, even the few imagery trials, whose reconstructions match the physical letter better than the perceptual data were mapped onto the perceptual pattern. A notable example is two trials for the letter ‘S’ by participant two. This implies that reconstruction quality of the perceptual data used to train the autoencoder constitutes an upper limit for imagery when using the autoencoder.

As a general effect, the autoencoder maps imagery voxel patterns onto their perceptual counterpart for most individual trials. Hence, reconstructions of average imagery voxel patterns as well as of individual trials more strongly resemble the corresponding physical letter. Figure 7 shows reconstructions from average imagery voxel patterns after feeding the data through the autoencoder.

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**Fig. 4** Reconstructed visual field images (participants 1–3). Reconstructed average visual field images are visualized for each ROI of participants one, two, and three. Reconstructions of the remaining three subjects are shown in Fig. 5. Perceptual as well as imagery voxel patterns were obtained from raw BOLD time-series.
Fig. 5 Reconstructed visual field images (participants 4–6). Reconstructed average visual field images are visualized for each ROI of participants four, five, and six. Reconstructions of the remaining three subjects are shown in Fig. 4. Perceptual as well as imagery voxel patterns were obtained from raw BOLD time-series.

Table 3  First-order correlations between reconstructed imagined letters and physical stimuli (averages over participants)

|       | H       | T       | S       | C       |
|-------|---------|---------|---------|---------|
| V1    | 0.24 (95% CI [0.09, 0.40]) | 0.49 (95% CI [0.38, 0.58]) | 0.08 (95% CI [0.03, 0.19]) | 0.14 (95% CI [0.07, 0.21]) |
| V2    | 0.21 (95% CI [0.13, 0.30]) | 0.46 (95% CI [0.36, 0.55]) | 0.10 (95% CI [0.04, 0.17]) | 0.12 (95% CI [0.07, 0.18]) |
| V3    | 0.20 (95% CI [0.10, 0.30]) | 0.28 (95% CI [0.14, 0.46]) | 0.04 (95% CI [0.03, 0.12]) | 0.15 (95% CI [0.01, 0.29]) |
| V1V2V3| 0.27 (95% CI [0.16, 0.37]) | 0.51 (95% CI [0.45, 0.56]) | 0.12 (95% CI [0.02, 0.21]) | 0.14 (95% CI [0.08, 0.20]) |
Figures 8 and 9 show reconstructions of individual trials in a single run of participants three and five, respectively. Obviously, these participants are not representative of the population at large but provide an indication of what is possible for people with a strong ability to imagine visual shapes. Table 5 shows the mean correlation values across trials of participant three and five when the data of these participants were fed through the autoencoder and without using the autoencoder.

### Classification

Having established support for the hypothesis that activity in early visual cortex in response to imagery exhibits a similar topographical profile as perception, we proceeded to test whether it is possible to pretrain latent representations for an imagery classifier using purely perceptual data. The classifier consists of three layers with the output layer being a softmax classifier stacked onto the hidden

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**Table 4** First-order correlations between reconstructed perceived letters and physical stimuli (averages over participants)

|     | H        | T        | S        | C        |
|-----|----------|----------|----------|----------|
| V1  | 0.40 (95% CI [0.35, 0.44]) | 0.65 (95% CI [0.60, 0.69]) | 0.27 (95% CI [0.15, 0.38]) | 0.32 (95% CI [0.22, 0.40]) |
| V2  | 0.37 (95% CI [0.31, 0.42]) | 0.58 (95% CI [0.50, 0.64]) | 0.19 (95% CI [0.09, 0.29]) | 0.31 (95% CI [0.23, 0.38]) |
| V3  | 0.25 (95% CI [0.19, 0.31]) | 0.41 (95% CI [0.30, 0.51]) | 0.06 (95% CI [−0.06, 0.18]) | 0.27 (95% CI [0.25, 0.30]) |
| V1V2V3 | 0.41 (95% CI [0.36, 0.46]) | 0.63 (95% CI [0.56, 0.68]) | 0.22 (95% CI [0.12, 0.32]) | 0.31 (95% CI [0.24, 0.38]) |
layer of an autoencoder pretrained to denoise perceptual voxel patterns (see “Methods” for details). We trained the classifier on imagery data using a leave-one-run-out procedure; that is, we trained the classifier on three of the four imagery runs and tested classification accuracy on the left-out run. Figure 10 shows average classification accuracies per subject and ROI (including the combined ROI ‘V1V2V3’). For five of the six participants, average classification accuracies exceeded theoretical chance levels (25% correct) as well as the 95th percentile.
Fig. 8: Reconstructed visual field images from denoised single trials in a single run of participant 3. Each run comprised of 8 trials (columns) per letter (rows). Recognizable reconstructions can be obtained for a number (though not all) individual trials.

Fig. 9: Reconstructed visual field images from denoised single trials in a single run of participant five. Each run comprised of eight trials (columns) per letter (rows). Recognizable reconstructions can be obtained for a number (though not all) individual trials.

Table 5: Effect of the autoencoder on mean correlation values across trials for two participants.

|       | H          | T          | S          | C          |
|-------|------------|------------|------------|------------|
| P03   |            |            |            |            |
| Autoencoder | 0.39 (95% CI [0.32, 0.45]) | 0.55 (95% CI [0.46, 0.62]) | 0.10 (95% CI [0.04, 0.16]) | 0.09 (95% CI [0.06, 0.12]) |
| Raw   | 0.19 (95% CI [0.15, 0.21]) | 0.33 (95% CI [0.28, 0.38]) | −0.02 (95% CI [−0.06, 0.02]) | 0.02 (95% CI [−0.02, 0.06]) |
| P05   |            |            |            |            |
| Autoencoder | 0.28 (95% CI [0.20, 0.35]) | 0.53 (95% CI [0.43, 0.61]) | 0.08 (95% CI [0.00, 0.17]) | 0.21 (95% CI [0.12, 0.31]) |
| Raw   | 0.12 (95% CI [0.08, 0.15]) | 0.32 (95% CI [0.25, 0.37]) | 0.02 (95% CI [−0.01, 0.06]) | 0.07 (95% CI [0.03, 0.10]) |
of 1000 permutation runs (randomly scrambled labels) in all ROIs. For participant six, theoretical chance levels as well as the 95th percentile were (barely) exceeded for V2 only.

We performed a mixed-model regression with the OSIVQ object, spatial and verbal scores, ROI (using dummy coding, V1 = reference), and number of selected voxels (again grouped by ROI) as predictors to assess which factors account for the observed accuracies. Number of voxels \( t_{(11)} = 4.80, p = 0.0006 \), the object sub-score of OSIVQ \( t_{(11)} = 4.83, p < 0.0005 \), and the spatial sub-score of OSIVQ \( t_{(11)} = 3.45, p = 0.006 \) were significant predictors of accuracy, whereas the verbal sub-score of OSIVQ \( t_{(11)} = 0.656, p = 0.525 \) was not. Furthermore, neither V2 \( t_{(11)} = -1.72, p = 0.113 \) nor V3 \( t_{(11)} = 1.85, p = 0.092 \) differed significantly from reference (V1).

**Discussion**

The aim of the present study was to investigate whether visual imagery exhibits sufficient topographic organization to preserve the geometry of internally visualized objects. To that end, we trained participants to maintain a vivid mental image of four letter shapes. Subsequently, we obtained sub-millimeter resolution 7T fMRI measurements from early visual cortex, while participants viewed or imagined the same letter shapes. Finally, we conducted a series of encoding, reconstruction, and decoding analyses to establish the degree of similarity between imagined and perceived shapes. Our results reveal that an object’s geometry is preserved during visual mental imagery.
Over training sessions, all participants reached a high probing accuracy for both imagery and perception trials, showing that they could reliably indicate the location of the invisible letter shape in visual space. The ability to imagine the borders of the letter in the absence of visual stimulation suggests that participants were able to generate a precise internal representation of the instructed letter. While providing explicit instructions to participants prohibited them from engaging in a more ecologically valid form of imagery, it is unlikely to fundamentally alter the neural processes underlying imagery. Instead, it allowed us to have a reasonable degree of confidence in the ground truth of imagined shapes. Next, we showed that patterns of voxel activations predicted by a pRF-encoding model and a physical (binary) letter stimulus can account for observed activation patterns in response to mental imagery of the letter corresponding to the physical stimulus. Given that pRF mapping has been shown to accurately predict fMRI responses to visual stimuli (Wandell and Winawer 2015), our results suggest that intrinsic geometric organization of visual experiences is also maintained during visual mental imagery. Our encoding analysis is somewhat reminiscent of that employed by Naselaris et al. (Naselaris et al. 2015) who used a more computational complex encoding model to identify an imagined artwork from a set of candidates. Given that our encoding model is limited to retinotopy, our approach is more restricted in its applications than that detailed in (Naselaris et al. 2015). However, a more restricted approach has the advantage of affording tighter experimental control providing a stronger basis for drawing conclusions. By focusing on a single feature (retinotopic organization), using stimuli differing solely with respect to their geometric properties, and directly comparing the predictions based on each stimulus regarding the activation profiles in early visual cortex, allowed us to draw specific conclusions regarding the topographic organization of mental imagery.

With respect to reconstructions, we found significant overlap between reconstructed imagery and the physical stimulus in terms of object geometry. While we anticipated this given findings that visual mental imagery exhibits retinotopic organization in early visual cortex (Slotnick et al. 2005; Albers et al. 2013; Pearson et al. 2015), these results were, nonetheless, exciting, because the previous reconstructions of mental imagery based on retinotopy did not preserve object geometry (Thirion et al. 2006). Indeed, to the best of our knowledge, we present the first visually recognizable reconstructions of mental imagery, even at the single-trial level. Our first-level correlation metric of reconstruction quality revealed that reconstruction quality of letter ‘S’ was significantly reduced, while that of letter ‘T’ was significantly improved with respect to that of letter ‘H’. This fits with the notion that stimuli exhibiting finer (coarser) spatial layouts would be harder (easier) to reconstruct. Furthermore, the OSIVQ object score was a significant predictor of first-level reconstruction quality, whereas the OSIVQ verbal score was not. This indicates that participants relying on an object-based imagery strategy were generally more successful at imagery of the letter shapes than participants relying on verbal strategies. Our findings are further in line with recent observations that neural overlap between imagery and perception in the visual system depends on experienced imagery vividness (Dijkstra et al. 2017). Interestingly, while inspection of Figs. 4 and 5 would suggest that reconstruction quality is ROI-specific, ROIs do not constitute a significant predictor of first-level reconstruction quality. Rather, the number of voxels included for any given ROI determined the quality. However, this does not imply that uncritically adding more voxels will definitely lead to higher classification accuracies. We included only those voxels for which pRF mapping yielded a high fit. It is likely that reconstructions benefit from a large number of voxels, whose pRF can be estimated to a high degree of precision (i.e., which show a strong spatially selective visual response, especially with high-resolution 7T fMRI) rather than a large number of voxels per se. The OSIVQ object score and number of voxels were also significant predictors of second-level reconstruction quality for reasons similar to those just mentioned.

Both our encoding and reconstruction results show that it is possible to extract similar information from perceived and imagined shapes. This possibility has previously been suggested to be strongly indicative of the pictorial nature of vividly experienced mental images (Brogaard and Gatzia 2017). In conjunction with the observation that the object but not the verbal score of the OSIVQ significantly predicts reconstruction quality, these results support the view that mental imagery is represented pictorially, at least within early visual cortex. At later stages of (visual) processing, mental imagery may become increasingly symbolic. As such, we do not wish to imply that our results settle the imagery debate, as they do not rule out the existence of (additional) symbolic representations.

We further show that the tight topographic correspondence between imagery and perception in early visual cortex allows for improved reconstruction and opens new avenues for classification. Specifically, our results indicate that training a denoising autoencoder on perceptual data creates an attractor landscape with one attractor per perceived letter. Importantly, the resemblance of imagery activation profiles in early visual cortex is sufficiently similar to its perceptual pendant to ensure that activation patterns of a large proportion of imagery trials fall within the attraction domain of the correct letter. The autoencoder then projects these imagery activation patterns onto the corresponding perceptual activation patterns. Though this is not the case for every trial with some being projected onto the wrong perceptual activation pattern pointing to
intra-individual fluctuations regarding successful imagery (Dijkstra et al. 2017). Nevertheless, these projections allow for perception-level reconstruction quality even for individual imagery trials for those participants with good imagery ability. It may be an interesting avenue for future research to study the attractor landscape formed through training the autoencoder and investigate under which conditions imagery trials fall inside or outside the attraction domain of each letter. Our current observations regarding the autoencoder imply that perception provides an upper limit on the achievable reconstruction quality. That is, any improvements of perceptual reconstructions, for instance, obtaining a more accurate encoding model by correcting for eye movements during pRF mapping (Hummer et al. 2016) should improve imagery reconstructions as well.

Furthermore, the autoencoder can be utilized to pretrain a classifier purely based on perceptual data before fine-tuning it on imagery data. We showed the feasibility of this approach using it for classifying imagined letters with a high degree of accuracy from at least one region of interest (between 50 and 70% correct) in five out of six participants. Statistical analyses revealed that both the OSIVQ object and OSIVQ spatial scores are significant predictors of classification accuracy. The finding that the OSIVQ spatial score constitutes a significant predictor here indicates that for classification, a cruder retinotopic organization of mental imagery might already be sufficient. Indeed, classification may rather benefit from an increased signal-to-noise ratio (SNR) which could be achieved by lowering the spatial resolution. Here, we opted for high spatial resolution to obtain precise receptive field estimates (additionally trading temporal resolution for SNR). In any case, successful classification may not be sufficient to draw conclusions regarding the precise geometry of imagined objects. As before, the number of voxels also constitutes a significant predictor of classification accuracy.

The autoencoder enables leveraging perceptual data to improve reconstructions of imagined letters and pretrain classifiers. This may eventually be utilized for the development of content-based BCI letter-speller systems. So far, fMRI-based BCI communication systems have mostly focused on coding schemes arbitrarily mapping brain activity in response to diverse mental imagery tasks (e.g., mental spatial navigation, mental calculation, mental drawing, or inner speech), and hence originating from distinct neural substrates, onto letters of the alphabet (Birbaumer et al. 1999; Sorger et al. 2012). As such, current BCI speller systems do not offer a meaningful connection between the intended letter and the specific content of mental imagery. This is demanding for users, as it requires them to memorize the mapping in addition to performing imagery tasks unrelated to intended letters and words. Our results constitute a proof-of-concept that it may be possible to achieve a more natural, content-based, BCI speller system immediately decoding internally visualized letters from their associated brain activity.

In conclusion, our letter encoding, reconstruction, and classification results indicate that the topographic organization of mental imagery closely resembles that of perception. This lends support to the idea that mental imagery is quasi-perceptual not only in terms of its subjective experience but also in terms of its neural representation and constitutes an important first step towards the development of content-based letter-speller systems.

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Compliance with ethical standards

Conflict of interest The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Studies involving human participants All procedures were conducted with approval from the local Ethical Committee of the Faculty of Psychology and Neuroscience at Maastricht University.

Informed consent All participants gave written informed consent and were paid for participation in this study.

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