| 1  | Finding decodable information that can be read out in behaviour                                      |
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#### Abstract

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Multivariate decoding methods applied to neuroimaging data have become the standard in 22 23 cognitive neuroscience for unravelling statistical dependencies between brain activation patterns 24 and experimental conditions. The current challenge is to demonstrate that decodable information 25 is in fact used by the brain itself to guide behaviour. Here we demonstrate a promising approach 26 to do so in the context of neural activation during object perception and categorisation behaviour. 27 We first localised decodable information about visual objects in the human brain using a multivariate decoding analysis and a spatially-unbiased searchlight approach. We then related 28 29 brain activation patterns to behaviour by testing whether the classifier used for decoding can be 30 used to predict behaviour. We show that while there is decodable information about visual 31 category throughout the visual brain, only a subset of those representations predicted 32 categorisation behaviour, which were strongest in anterior ventral temporal cortex. Our results have important implications for the interpretation of neuroimaging studies, highlight the 33 importance of relating decoding results to behaviour, and suggest a suitable methodology towards 34 35 this aim.

# 37 **1 Introduction**

38 Multivariate pattern analysis (MVPA), also called brain decoding, is a powerful tool to establish 39 statistical dependencies between experimental conditions and brain activation patterns (Carlson, 40 Schrater, & He, 2003; Cox & Savoy, 2003; Haxby et al., 2001; Haynes, 2015; Kamitani & Tong, 41 2005; Kriegeskorte, Goebel, & Bandettini, 2006). In these analyses, an implicit assumption often made by experimenters is that if information can be decoded, then this information is used by the 42 43 brain in behaviour (de-Wit, Alexander, Ekroll, & Wagemans, 2016; Ritchie, Kaplan, & Klein, 2017). However, the decoded information could be different (e.g., epiphenomenal) from the signal that is 44 relevant for the brain to use in behaviour (de-Wit et al., 2016; Williams, Dang, & Kanwisher, 2007), 45 highlighting the need to relate decoded information to behaviour. Importantly, this implicit 46 47 assumption of decoding models leads to testable predictions about task performance (Naselaris, 48 Kay, Nishimoto, & Gallant, 2011). Previous work has for example correlated decoding performances to behavioural accuracies (Bouton et al., 2018; Freud, Culham, Plaut, & Behrmann, 49 2017; Raizada, Tsao, Liu, & Kuhl, 2010; van Bergen, Ji Ma, Pratte, & Jehee, 2015; Walther, 50 51 Caddigan, Fei-Fei, & Beck, 2009; Williams et al., 2007). However, this does not model how individual experimental conditions relate to behaviour. Another approach has been to compare 52 53 neural and behavioural similarity structures (Bracci & Op de Beeck, 2016; Cichy, Kriegeskorte, 54 Jozwik, Bosch, & Charest, 2017; Cohen, Dennett, & Kanwisher, 2016; Grootswagers, Kennedy, 55 Most, & Carlson, 2017; Haushofer, Livingstone, & Kanwisher, 2008; Mur et al., 2013; Proklova, 56 Kaiser, & Peelen, 2016; Wardle, Kriegeskorte, Grootswagers, Khaligh-Razavi, & Carlson, 2016). 57 While this approach allows to link behaviour and brain patterns at the level of single experimental conditions, it is unclear how this link carries over to decision making behaviour such as 58 59 categorisation (but see Cichy et al., (2017) for recent developments).

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61 Recently, a novel methodological approach, called the distance-to-bound approach (Ritchie & 62 Carlson, 2016), has been proposed to connect brain activity directly to perceptual decision-making 63 behaviour at the level of individual experimental conditions. The rationale behind this approach 64 (Bouton et al., 2018; Carlson, Ritchie, Kriegeskorte, Durvasula, & Ma, 2014; Kiani, Cueva, Reppas, 65 & Newsome, 2014; Philiastides & Saida, 2006; Ritchie & Carlson, 2016) is that for decision-making 66 tasks, the brain applies a decision boundary to a neural activation space (DiCarlo & Cox, 2007). 67 Similarly, MVPA classifiers fit multi-dimensional hyperplanes to separate a neural activation space. In classic signal-detection theory (Green & Swets, 1966) and evidence-accumulation models of 68 choice behaviour (Brown & Heathcote, 2008; Gold & Shadlen, 2007; Ratcliff & Rouder, 1998; Smith 69 & Ratcliff, 2004), the distance of the input to a decision boundary reflects the ambiguity of the 70 71 evidence for the decision (Green & Swets, 1966). Decision evidence, in turn, predicts choice 72 behaviour (e.g., Ashby, 2000; Ashby & Maddox, 1994; Britten, Newsome, Shadlen, Celebrini, & Movshon, 1996; Gold & Shadlen, 2007; Shadlen & Kiani, 2013) which also has clear neural 73 correlates (e.g., Britten et al., 1996; Ratcliff, Philiastides, & Sajda, 2009; Roitman & Shadlen, 2002). 74 If for a decision task (e.g., categorisation), the brain uses the same information as the MVPA 75 76 classifier, then the classifier's hyperplane reflects the brain's decision boundary. This in turn 77 predicts that distance to the classifier's hyperplane negatively correlates with reaction times for 78 the decision task. In the distance-to-bound approach, finding such a negative distance-RT-79 correlation shows that the information is then suitably formatted to guide behaviour. "Suitably 80 formatted to guide behaviour" here means that the information is structured in such a way that 81 the brain can apply a linear read out process to this representation to make a decision (importantly, this does not imply a causal link with behaviour). Carlson et al. (2014) demonstrated 82 83 the promise of the distance-to-bound approach in a region of interest based analysis using fMRI. Here we go beyond this work by using the distance-to-bound method and a spatially unbiased 84

- 85 fMRI-searchlight approach to create maps of where in the brain information can be used to guide86 behaviour.
- 87

#### 88 2 Materials and Methods

In this study, we separately localised information that is decodable, and information that is 89 90 suitably formatted to guide behaviour in the context of decodable information about visual 91 objects and object categorisation behaviour. To ensure robustness and generality of our results, 92 we analysed in parallel two independent fMRI datasets (Cichy et al., 2014, 2016), with different 93 stimulus sets, and in relation to partly overlapping categorisation behaviours. Overall, this allowed 94 us to investigate the relationship between decodable information from brain activity and 95 categorisation behaviour for seven different distinctions: animate versus inanimate, faces versus 96 bodies, human versus animal, natural versus artificial, tools versus not tools, food versus not food, 97 and transport versus not transport. Note that the negative 'not-X' category was defined as all 98 stimuli that did fall into one of the aforementioned classes. Categorisation reaction times for those 99 stimuli were collected on Amazon's Mechanical Turk. In this section, we describe the two-step 100 searchlight procedure used to create decoding and correlation maps of areas involved in visual 101 object categorisation.

102

#### 103 2.1 Experimental design

104 Stimuli

Stimuli for experiment 1 consisted of 92 visual objects, segmented on a white background (Figure 106 1A). Stimuli consisted of animate and inanimate objects. The animate objects could be further 107 divided into faces, bodies, humans and animals. Inanimate objects consisted of natural (e.g., 108 plants or fruits) and man-made items (e.g., tools or houses). The stimulus set for experiment 2 109 consisted of 118 visual objects on natural backgrounds (Figure 1C). A small proportion of the objects (27) were animate. The inanimate objects included subcategories such as tools, or food items. In both experiments, participants were presented with the visual object stimuli while performing an orthogonal task at fixation. Stimuli were displayed at 2.9° (Experiment 1) and 4.0° (Experiment 2) visual angle with 500 ms duration. Images were displayed (overlaid with a grey fixation cross) for 500 ms in random order.

115

#### 116 fMRI recordings

The first experiment (Cichy et al., 2014) had high resolution fMRI coverage of the ventral visual stream (Figure 1B) from 15 participants with a 2 mm isotropic voxel resolution. The second experiment (Cichy et al., 2016) had whole brain from 15 participants with a 3 mm isotropic voxel resolution. In both experiments, at the start of a session, structural images were obtained using a standard T<sub>1</sub>-weighted sequence. fMRI data were aligned and coregistered to the T1 structural image, and then normalized to a standard MNI template. General linear models were used to compute t-values for each stimulus (92 and 118, respectively) against baseline.

124

#### 125 Reaction time data

We obtained reaction times for the stimuli in multiple different categorisation contrasts (Figure 126 127 1A&B). For experiment 1, these were animate versus inanimate, face versus body, human versus animal, and natural versus artificial. For experiment 2, we tested animate versus inanimate, tool 128 129 versus not tool, food versus not food, and transport versus not transport. The RTs were collected using Amazons Mechanical Turk (MTurk). For each of the categorisation contrasts, 50 unique 130 131 participants performed a categorisation task using the same stimuli as were used in collecting the fMRI data. Participants were instructed to "Categorise the images as fast and accurate as possible 132 133 using the following keys: (z for X, m for Y)", where X and Y would be replaced with the relevant categories (e.g., animate and inanimate) for the contrast. On each trial, an image was presented 134

for 500ms, followed by a black screen until the participant's response (Figure 1C). The presentation order of the stimuli was randomized and stimuli did not repeat. This resulted in 50 reaction time values per exemplar (one for each participant). Each participant's reaction times were z-scored. Next, we computed the median reaction time (across participants) for each exemplar. his resulted in one reaction time value per exemplar, which were used in the rest of the study.

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# 142 2.2 Statistical Analysis

#### 143 Searchlight procedure

For each categorisation contrast and subject, we used a searchlight approach (Haynes et al., 2007; Kriegeskorte et al., 2006) to create maps of decoding accuracy and of correlations between distance to the classifier boundary and categorisation reaction time. In contrast to pre-defined ROI's, which are used to test a-priori hypotheses about the spatial origin of information in the brain, the searchlight results in a spatially unbiased map of decodable information. An overview of the approach is presented in Figure 1D.

150

151 To create the decoding accuracy maps, we used a standard searchlight decoding approach 152 (Grootswagers, Wardle, & Carlson, 2017; Haynes, 2015; Kriegeskorte et al., 2006; Pereira, Mitchell, 153 & Botvinick, 2009), as implemented in the CoSMoMVPA decoding toolbox (Oosterhof, Connolly, & Haxby, 2016). In detail, at each spatial location (voxel) in an fMRI image, a support vector machine 154 (SVM) was used to classify visual object category based on local brain patterns, resulting in a map 155 156 of classification accuracies. We then determined the subset of the locations at which brain 157 patterns were suitably formatted for read-out by the brain using the distance-to-bound approach (Ritchie & Carlson, 2016) in a second searchlight analysis. Analogous to the decoding analysis, at 158 159 each voxel, an SVM was trained to classify visual objects. Diverging at this point from the decoding

160 approach we did not test the classifier, but rather obtained the distance for each exemplar to the hyperplane set by the SVM. We then correlated those distances to reaction times acquired in 161 separate categorisation tasks. The contribution of each category was assessed individually, by 162 163 performing the correlations separately for the two sides of the categorisation (e.g., one correlation for animate and one for inanimate exemplars). For each categorisation task this 164 resulted in two correlation maps per subject. The maps of decoding accuracy and correlations 165 166 were assessed for significance at the group level using sign-rank tests for random-effects 167 inference. The results were thresholded at p < 0.05, using the false discovery rate (FDR; (Benjamini & Hochberg, 1995)) to correct for multiple comparisons at the voxel level. 168

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# 170 Relating the results to topographical locations of the visual system

171 For the animacy categorisation contrasts, we identified the locations of the significant voxels with 172 respect to ROIs of the visual system. The significant voxels in the decoding maps and correlation maps were compared to probabilistic topographic maps of visual processing areas (Wang, 173 Mruczek, Arcaro, & Kastner, 2015), which represent for each voxel the visual area with the highest 174 probability. A percentage score for each ROI was then computed, reflecting the percentage of 175 176 voxels in this ROI that were significant at the group level. We obtained a bootstrapped distribution 177 of percentage scores for each ROI by repeating this procedure 10,000 times, while randomly sampling the subjects with replacement and recomputing the group level statistics. We report the 178 5<sup>th</sup>, 50<sup>th</sup> and 95<sup>th</sup> percentiles of this distribution. This approach allows quantifying the difference 179 180 between the number of decoding voxels and correlation voxels per visual ROI.



181

**Figure 1. General experimental rationale.** Stimuli (**A**,**C**) used to map fMRI brain responses and brain coverage (**B**,**C**) for fMRI study 1 and 2 respectively. **E**. Acquisition of reaction times on object categorisation tasks. Reaction times for categorisation contrasts were collected in a different pool of participants than the ones participating in the fMRI experiment. On each trial, a stimulus was displayed for 250ms, and participants categorised it into two categories (exemplarily here: animate vs inanimate) by pressing one of two keys. **F**. The two-partite approach to separately

188 localize decodable information and information that is suitable for read out in behaviour. For both 189 parts, a local cluster of neighbouring voxels (i.e., searchlight) was used to train a linear support 190 vector machine (SVM) on an image category classification task (e.g., animacy). To localize decodable information, the classifier was tested on left-out data, storing the classification 191 192 accuracy at the centre voxel of the searchlight. To localise information that was suitably formatted 193 for read-out in a categorisation task, the distances of objects to the classifier hyperplane were 194 correlated with the reaction times for the same object images on the same classification task. 195 Repeated for every voxel, this resulted for each subject in one map of decoding accuracies and 196 one of correlations. For visualisation, significant correlation voxels were superimposed on 197 significant decoding accuracy voxels, each showing group average values in significant voxels.

198

#### 199 **3 Results**

200 We examined the relationship between decodable information and information that is suitably formatted for read-out by the brain in the context of decodable information about visual objects 201 202 and object categorisation behaviour. We determined the relationship between decodable 203 information and behaviour separately. First, we determined where information about objects is 204 present in brain patterns using decoding in a standard fMRI searchlight decoding analysis (Haynes 205 et al., 2007; Kriegeskorte et al., 2006). We then determined the subset of the locations at which 206 brain patterns were suitably formatted for read-out by the brain using the distance-to-bound 207 approach (Ritchie & Carlson, 2016) in a second searchlight analysis. The subject-specific 208 searchlight results were subjected to inference statistics at the group level using one-sided sign 209 rank tests and thresholded at p < 0.05 (fdr-corrected for multiple comparisons across voxels).

210

# 3.1 A subset of locations that have decodable information about animacy also had information suitably formatted for animacy categorisation behaviour

Animacy is a pervasive and basic object property according to which any object can be classified as
animate or inanimate (Caramazza & Shelton, 1998). Previous studies have shown that the division

215 of animate versus inanimate objects is reflected in the large-scale architecture of high-level visual 216 areas such as the ventral temporal cortex (VTC) (Caramazza & Shelton, 1998; Grill-Spector & Weiner, 2014; Kriegeskorte et al., 2008), However, it has also been shown that animacy can be 217 218 decoded not only from VTC, but from the whole ventral visual stream (Cichy et al., 2016; Grill-219 Spector & Weiner, 2014; Long, Yu, & Konkle, 2017). Furthermore, categorical object responses have also been found in the dorsal visual stream (Bracci, Daniels, & op de Beeck, 2017; Freedman 220 221 & Assad, 2006; Konen & Kastner, 2008) and in frontal areas (Freedman, Riesenhuber, Poggio, & Miller, 2001, 2003). This prompts the question of where in the visual system object 222 representations are suitably formatted for read-out by the brain for animacy decisions. 223

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225 Corroborating previous studies, we found decodable information about animacy in the entire 226 ventral visual stream from the occipital pole to anterior ventral temporal cortex (Figure 2AB, Table 227 1AE, N = 15, one-sided sign-rank test, p < 0.05 fdr-corrected). In addition, we found decodable information in dorsal and prefrontal cortex (Figure 2B) in experiment 2 which had full brain 228 coverage. Localising the brain representations suitable to guide animacy categorisation behaviour 229 230 (using the distance-to-bound approach) revealed convergent evidence across experiments that 231 only a subset of voxels containing decodable information fulfilled this criterion. In detail, distance-232 RT-correlations for animate objects were strongest in the high-level regions of the ventral and the 233 dorsal stream. For inanimate objects, we found no voxels with significant distance-RT-correlations 234 (Carlson et al., 2014; Grootswagers, Ritchie, Wardle, Heathcote, & Carlson, 2017).



Figure 2. Relationship between decodable information and categorisation behaviour for animacy. Decodable information is shown in hot colours and distance-RT-correlations in cool colours. Colour intensities reflect the mean across subjects. Only significant voxels (N=15, sign-

240 rank test, p<0.05 fdr-corrected) are shown. Data are projected onto axial slices of a standard  $T_1$ 241 image in MNI space. A. In experiment 1, decodable animacy information (hot colours) was found 242 throughout the ventral stream. A correlation between distance to the classifier boundary and 243 reaction time for animate stimuli (cool colours) was found in a subset of these areas. The colour 244 intensities depict the mean across subjects. B. The results of the analysis for experiment 2 245 corroborated these findings, and showed decodable information in prefrontal areas and in the 246 dorsal visual stream. Correlations between distance and reaction time were also present in the 247 dorsal stream.

| $\mathbf{r}$ | л | n |
|--------------|---|---|
| Z            | 4 | 3 |
|              |   |   |

| Contrast                                     | #significant | Max/min | х   | Y   | Z   |
|--|--------------|---------|-----|-----|-----|
|  | voxels       |         |     |     |     |
| A) decoding 'animate' vs 'inanimate' (exp 1) | 11745        | 0.80    | 36  | -52 | -15 |
| Distance-RT-correlation 'animate'            | 6410         | -0.38   | 38  | -58 | -19 |
| Distance-RT-correlation 'inanimate'          | 0            | -0.16   | -48 | -58 | 5   |
| B) decoding 'human' vs 'animal'              | 4863         | 0.69    | 22  | -90 | -13 |
| Distance-RT-correlation 'human'              | 0            | -0.29   | 30  | -58 | -15 |
| Distance-RT-correlation 'animal'             | 0            | -0.17   | 48  | -46 | -4  |
| C) decoding 'face' vs 'body'                 | 10661        | 0.84    | 44  | -78 | -10 |
| Distance-RT-correlation 'face'               | 226          | -0.32   | 40  | -76 | -15 |
| Distance-RT-correlation 'body'               | 0            | -0.20   | -54 | -68 | 16  |
| D) decoding 'natural' vs 'artificial'        | 0            | 0.63    | 30  | -52 | -17 |
| Distance-RT-correlation 'natural'            | 0            | -0.27   | 42  | -72 | -4  |
| Distance-RT-correlation 'artificial'         | 0            | -0.24   | 26  | -88 | 3   |
| E) decoding 'animate' vs 'inanimate' (exp 2) | 8824         | 0.80    | 36  | -55 | -11 |
| Distance-RT-correlation 'animate'            | 2015         | -0.34   | 51  | -73 | -2  |
| Distance-RT-correlation 'inanimate'          | 0            | -0.12   | -21 | -43 | -2  |
| F) decoding 'tool' vs 'not tool'             | 0            | 0.58    | -30 | -94 | 7   |
| Distance-RT-correlation 'tool'               | 0            | -0.25   | -33 | -13 | 19  |
| Distance-RT-correlation 'not tool'           | 0            | -0.24   | -33 | -52 | -17 |
| G) decoding 'transport' vs 'not transport'   | 0            | 0.59    | 33  | -94 | 1   |
| Distance-RT-correlation 'transport'          | 0            | -0.32   | 15  | 50  | 4   |
| Distance-RT-correlation 'non-transport'      | 0            | -0.18   | -33 | -55 | -14 |
| H) decoding 'food' vs 'not food'             | 1092         | 0.62    | 36  | -55 | -14 |
| Distance-RT-correlation 'food'               | 0            | -0.16   | -18 | 26  | -5  |
| Distance-RT-correlation 'not food'           | 154          | -0.13   | 27  | -40 | -14 |

Table 1. Results for all categorisation contrasts. For all categorisation contrasts, we report the number of significant voxels (after correction for multiple comparisons), its peak value (maximum for decoding or minimum for distance-RT-correlation), and the peak's location in MNI-XYZ coordinates.

#### **3.2 The proportion of region-specific representations suitably formatted for behaviour increases**

# along the ventral stream and decreases along the dorsal stream

257 We next explicitly determined the degree to which representations in single brain regions within 258 the ventral and dorsal streams are suitably formatted for behaviour. For this we parcellated the 259 cortex (Figure 3A) using a probabilistic topographic map of visual processing areas (Wang et al., 260 2015). For each region, we calculated the ratio between the number of significant voxels in the 261 decoding analysis and the total number of voxels, so that a high ratio indicates that a large part of a region contains object representations with categorical information. Similarly, we calculated the 262 263 ratio between the number of significant voxels in the distance-to-bound analysis and the total 264 number of voxels. Here, a high ratio indicates that a large part of a region contains object 265 representations that are suitably formatted for read out in a categorisation task.

266

267 In the ventral stream, our results suggest that these ratios increase with processing stage, from early visual areas to high-level visual areas, with highest ratios in ventral occipital (VO) and 268 269 parahippocampal (PHC) cortex (Figure 3 B&C). In contrast, in the dorsal stream we observed a 270 decrease of the correlation ratio with processing stage. In addition, significant animacy decoding 271 information was found in similar proportions in the ventral-temporal areas as in lateral-occipital 272 areas, however, the proportion of voxels with information suitable for categorisation was lower in 273 lateral-occipital areas. This is consistent with the notion that while both these regions contain 274 object representations, the VTC contains location-invariant representations which are essential for 275 object categorisation (Cichy et al., 2013; Haushofer et al., 2008; Schwarzlose, Swisher, Dang, & 276 Kanwisher, 2008; Williams et al., 2007). The results were similar between experiments, with the 277 exception for area TO, which had a smaller proportion of voxels with RT-correlations in experiment 1. It is possible that this difference was caused by the differences between the stimuli 278 279 (e.g., segmented objects versus objects in scenes) used in the experiments. Alternatively, this

280 difference could be attributed to the size of the searchlight sphere, which was larger in
281 experiment 2 than in experiment 1 due to their different voxel sizes.

282

In sum, these results show that representations along the ventral stream are suitably formatted for read-out of categorical information (Cichy et al., 2013; Grill-Spector & Weiner, 2014). In contrast, representations in the dorsal stream might be shaped for the read-out in different tasks (Bracci et al., 2017; Freud et al., 2017). These results also suggest that intermediate stages along the ventral and dorsal streams may be similar or partly shared, as suggested by the similar ratios of information suitable for read-out.



291 Figure 3. Quantifying the decodable information in visual areas and their contribution to 292 categorisation behaviour. A. Locations of topographical ROIs of the visual system (Wang et al., 293 2015), containing early visual cortex (EVC) areas V1 and V2, mid-level areas V3 and hV4, high level ventral occipital (VO) and parahippocampal cortex (PHC), temporal occipital (TO) and lateral 294 295 occipital (LO) areas, areas in the intra-parietal sulcus (IPS), the superior parietal lobule (SPL), and the frontal eye fields (FEF). B-C. The ratio between significant voxels in an ROI and the size of the 296 297 ROI. Orange points show the ratio of voxels within the ROI that had significant animacy decoding 298 performance. Blue points show the ratio of voxels with a significant correlation between distance to the hyperplane and RT for 'animate'. The lower, middle and upper points on these lines indicate 299 5<sup>th</sup>, 50<sup>th</sup>, and 95<sup>th</sup> percentiles (bootstrapping of participants 10,000 times). These results quantify 300 301 the increasing contribution of early to late areas in the ventral visual stream to animacy 302 categorisation behaviour.

# 303 **3.3 Decodable information about subordinate categorisation tasks is also suitably formatted for**

# 304 categorisation behaviour

305 While animacy categorisation may be based on large-scale representational differences in the 306 visual brain (Carlson, Tovar, Alink, & Kriegeskorte, 2013; Downing, Chan, Peelen, Dodds, & 307 Kanwisher, 2006; Grill-Spector & Weiner, 2014; Kriegeskorte et al., 2008), subordinate 308 categorisation tasks (e.g., faces, bodies, tools) may depend more on fine grained patterns in focal 309 brain regions (Downing, Jiang, Shuman, & Kanwisher, 2001; Downing & Peelen, 2016; Kanwisher, 310 McDermott, & Chun, 1997). Here, we tested whether decodable information about subordinate category membership is also suitably formatted for read out in respective categorisation tasks. We 311 tested two subordinate contrasts for the animate exemplars in experiment 1: face versus body, 312 313 and human versus animal using the same general procedure as for animacy. We found that both 314 contrasts were decodable (Table 1B-C). We found a significant correlation between distance to the 315 classifier hyperplane and reaction times for faces in the face versus body task (Figure 4A). We found no significantly decodable information or significant correlations for the natural versus 316 artificial objects (Table 1D). Of the subordinate categorisation contrasts in experiment 2 (food, 317 318 transport or tool versus everything else), transport and tool versus everything else were not 319 significantly decodable information nor had they significant correlations (Table 1F-G). Food versus not food resulted in significant decodable information, and significant distance-RT correlations 320 were present for this contrast in the 'not food' category (Figure 4B, Table 1H). Taken together, for 321 322 some subordinate categorisation contrasts that were decodable, we were successful in localising brain patterns suitably formatted for read-out in behaviour. 323



Figure 4. Relationship between decodable information and behaviour for subordinate categorisation tasks. Decodable information is shown in hot colours and distance-RT-correlations in cool colours. Colour intensities reflect the mean across subjects. Only significant voxels (N=15,

328 sign-rank test, p<0.05 fdr-corrected) are shown. Data are projected onto axial slices of a standard 329 T<sub>1</sub> image in MNI space. **A.** In experiment 1, decodable face versus body information (hot colours) 330 was found in the entire ventral stream. A distance-RT-correlation for the face stimuli (cool colours) 331 was found in a subset of these areas. **B.** In experiment 2, food versus not food was decodable in 332 some areas in the ventral visual stream. A distance-RT-correlation for the 'not food' stimuli was 333 found in a subset of these areas.

334

335 4 Discussion

#### **4.1 Dissociating between decodable information and information that is used in behaviour**

337 The aim of this study was to examine where in the brain decodable information is suitably 338 formatted for read-out by the brain in behaviour. We found that only a subset of information that 339 is decodable could be related to behaviour using the distance-to-bound approach, which argues for a partial dissociation between decodable information and information that is relevant for 340 341 behaviour. This speaks to a current challenge in neuroimaging, which is to show that information 342 visible to the experimenter is in fact used by the brain (de-Wit et al., 2016; Ritchie et al., 2017). To 343 illustrate, consider the question about what regions are used by the brain to perform an object 344 animacy categorisation task (DiCarlo, Zoccolan, & Rust, 2012; Grill-Spector & Weiner, 2014). On its 345 own, the result of the animacy decoding searchlight might be interpreted as the brain using 346 animacy information from anywhere in the ventral stream. However, when investigating this interpretation directly using the distance-RT-correlation results, it becomes clear that object 347 348 animacy information is suitably represented for read-out in mid- and high-level visual areas only.

349

350 It is important to note that not finding a correlation between distance to the classifier hyperplane 351 and RT does not imply that the information revealed using the decoding approach is irrelevant or 352 epiphenomenal. The distance-to-bound approach taken here makes specific assumptions about 353 the brain's read-out process, such as distance in representational space as the measure for

354 evidence, and a monotonic relationship between distance and reaction time (Ritchie & Carlson, 2016). Note that this model of readout follows from the assumptions behind the decoding 355 methods (Ritchie & Carlson, 2016; Ritchie et al., 2017). While the model may not be perfect, our 356 357 results stress the importance of explicitly testing models of readout when decoding information from the brain. Finding the correct model of readout would significantly increase the capacity of 358 359 cognitive neuroscience to infer brain-behaviour relationships. Other assumptions follow from 360 those imposed by the decoding approach, such as the binary classification, the size of the searchlight radius, the choice of classifier. For example, it could be that the representations are 361 362 relevant in a different task (Grootswagers, Ritchie, et al., 2017; Ritchie & Carlson, 2016), or that 363 read-out involves pooling over larger spatial scales or multiple brain areas. Therefore, the current approach only allows the positive inference on the level of suitability of decoded information for 364 365 behaviour in the context of the current task and decoding parameters. On the other hand, a 366 correlation with behaviour still does not prove that the information is used by the brain, but it shows that the information is at least formatted in a way that is suitable to be used by the brain 367 368 for decisions. Future work can use causal measures (e.g., TMS) targeting the areas highlighted in the current results. 369

370

# **4.2** The contribution of ventral and dorsal visual regions to categorisation behaviour

We found that neural representations suitably formatted for behaviour in categorisation were most prominently located in the anterior regions of the VTC. This corroborates previous studies (Afraz, Kiani, & Esteky, 2006; Carlson et al., 2014; Hong, Yamins, Majaj, & DiCarlo, 2016; Hung, Kreiman, Poggio, & DiCarlo, 2005), and reinforces the tight link between VTC and visual categorisation behaviour. In these areas, our results provide converging evidence for the (implicit) assumption made in neuroimaging studies, which is that information that is available to the experimenter is also available for read out by the brain in behaviour (cf. de-Wit et al., 2016).

379

However, we found that correlations between distance to boundary and RT were not restricted to 380 anterior regions of the VTC, but were also prominent in V3 and hV4. This is consistent with the 381 382 view that lower level visual features encoded in mid-level visual regions could aid faster read-out 383 of category information. V4 is thought of as an intermediate stage of visual processing that 384 aggregates lower level visual features into invariant representations (Riesenhuber & Poggio, 385 1999). It has been proposed that direct pathways from V4 to decision areas allow the brain to exploit visual feature cues for fast responses to ecologically important stimuli (Hong et al., 2016; 386 387 Kirchner & Thorpe, 2006; Thorpe, Fize, & Marlot, 1996), such as identifying faces (Crouzet, 388 Kirchner, & Thorpe, 2010; Honey, Kirchner, & VanRullen, 2008). An alternative possibility is that read out is not happening directly from V4, but its representational structure is shaped by the low-389 390 level feature differences in animacy. This structure is then largely preserved when it is 391 communicated to more anterior areas, leading to similar distance-RT-correlations. Both of these 392 accounts are also consistent with recent findings that show differential responses for object 393 categories in mid-level visual areas (Long et al., 2017; Proklova et al., 2016). The extent to which 394 visual features contribute to the read-out process could be further investigated by using the 395 approach from this study with different stimulus sets that control for these features (Kaiser, 396 Azzalini, & Peelen, 2016; Long et al., 2017; Proklova et al., 2016).

397

We found that distance-RT-correlations were also present in early parietal areas. The classical view is that the ventral and dorsal visual streams are recruited for different function (Ungerleider & Mishkin, 1982). However, areas in the ventral and dorsal streams have been found to exhibit similar object-selective responses (Freud et al., 2017; Konen & Kastner, 2008; Sereno & Maunsell, 1998; Silver & Kastner, 2009). Consistent with this, we found similar RT-distance-correlations in mid-level areas in the ventral and dorsal streams. However, our results also showed that the

404 proportion of correlations decreased along the dorsal stream, while they increased along the
405 ventral stream. This suggests that representations in the ventral and dorsal streams undergo
406 similar transformations at first, and then diverge for different goals.

407

# 408 **4.3** Without a task, neural object representations in the VTC are formatted for read-out in 409 categorisation decisions

410 Here, the fMRI participants performed an orthogonal task, and were not actively categorising. Despite this, categorisation reaction times could still be predicted from representations in the 411 412 visual stream. This highlights that, without a categorisation task, information in the visual system 413 is represented in a way that is suitable for read out in behaviour (Carlson et al., 2014; Ritchie, 414 Tovar, & Carlson, 2015). This representation possibly reflects a more general property of the object that aids its categorisation, such as how typical it is for that category (Grootswagers, 415 416 Ritchie, et al., 2017; Iordan, Greene, Beck, & Fei-Fei, 2016), or how frequently we encounter the 417 object in our lives. In addition, the orthogonal task in the scanner has the advantage that it avoids 418 RT- and difficulty confounds (see e.g., Hebart & Baker, 2017; Woolgar, Golland, & Bode, 2014). 419 Future studies might use the distance-to-bound approach with participants actively performing 420 the same task in the scanner, where we predict that areas involved in the decision making and 421 execution processes would contain information that correlates with reaction times. For example, 422 some areas preferentially represent task-relevant information, such as areas in the prefrontal 423 cortex (Duncan, 2001; Jackson, Rich, Williams, & Woolgar, 2016; Woolgar, Jackson, & Duncan, 2016), and in the parietal stream (Bracci et al., 2017; Freedman & Assad, 2016; Jeong & Xu, 2016). 424 425 In the absence of an animacy categorisation task, one would predict that animacy information 426 would not be strongly represented in these areas. Yet, our results showed that animacy information can be decoded from prefrontal and parietal areas when participants perform an 427 428 orthogonal task. However, our results did not provide evidence that the animacy information in

429 these areas was suitably formatted for readout. This again argues for a dissociation between information that can be decoded, and information that is suitable for read out in behaviour. A 430 prediction that follows from this is that performing an active object categorisation task in the 431 432 scanner would change the representations in these task-relevant areas so that they become predictive of reaction times (Bugatus, Weiner, & Grill-Spector, 2017; McKee, Riesenhuber, Miller, 433 434 & Freedman, 2014). Similarly, representations can change when participants perform different 435 tasks on the same stimuli, such as categorising a specific feature (e.g., colour), for which suitably formatted information would be predicted in other areas. 436

437

# 438 **4.4 Asymmetric distance-RT-Correlations in binary categorisation tasks**

439 In both experiments, we found correlations between distance and reaction times for animate stimuli, but none for the inanimate stimuli. This is consistent with previous work (Carlson et al., 440 441 2014; Grootswagers, Ritchie, et al., 2017; Ritchie et al., 2015), which argued that this discrepancy might be caused by inanimate being a negatively defined category (i.e., "not animate"). Under this 442 hypothesis the animacy categorisation task can be performed by collecting evidence for animate 443 444 stimuli and responding inanimate only when not enough evidence was accumulated after a certain 445 amount of time. Here, we tested a prediction of this hypothesis by contrasting two positively 446 defined categories, face versus body, and found that there was a distance-RT-correlation only for faces. This goes against the notion of the negative definition of inanimate as the main reason for a 447 448 lack of correlation. However, it still is possible that observers still treated these tasks as 'A' or 'NOT A', with 'A' being the category that is easiest to detect (Grootswagers, Ritchie, et al., 2017). For 449 450 example, perceptual evidence for a face would be easier to obtain than evidence for a body-part, 451 as faces share low level visual features (Crouzet & Thorpe, 2011; Honey et al., 2008; Wu, Crouzet, Thorpe, & Fabre-Thorpe, 2015). Thus, while not explicitly specified as a negative category, it could 452 453 have been treated as such.

454

This suggests that the binary categorisation might be an unnatural way of approaching human 455 categorisation behaviour in the real world. Other operationalisations such as picture naming or 456 457 visual search may be better suited to capture the relevant behaviours (cf. Krakauer, Ghazanfar, 458 Gomez-Marin, Maclver, & Poeppel, 2017). Still, it is important to note that the binary task matches 459 the brain decoding task performed by the classifier. The above-chance decoding accuracy in the 460 brain decoding task is commonly interpreted as a similar dichotomy in the brain's representation that the brain can use in a decision. However, when only the information in one of the categories 461 462 (i.e., animals or faces) can be used to predict decision behaviour, as shown here, then this 463 interpretation needs to be revisited.

464

#### 465 **4.5 Limitations of the approach**

466 Our results highlight the importance of relating decoding to behaviour and demonstrated one possible methodology to address this issue. However, the approach taken here is subject to a set 467 468 of limitations which may preclude its application in other settings. Firstly, here we studied a binary 469 visual object categorisation task. It is not possible to describe all behaviours as binary tasks, and 470 reaction times are not always a meaningful measure for behaviour. This can restrict the 471 generalisability of the current approach to other domains. Secondly, finding an RT-correlation 472 does not reveal the source of the variance in evidence for a decision. As the method remains correlational, it is important to stress that it can only go as far to show that information is suitably 473 formatted to be used by the brain for decisions, and that the critical test of this relationship will 474 475 require causal measures. In the animacy task, one possible source of variance is typicality, which 476 modulates animacy categorisation (Posner & Keele, 1968; E. H. Rosch, 1973; E. Rosch & Mervis, 1975) and decoding performance (lordan et al., 2016), and typicality ratings have been shown to 477 478 correlate with distance to the classifier decision boundary (Grootswagers, Ritchie, et al., 2017).

479 Yet, there is always the possibility of other covariates. For example, larger or more colourful objects may be easier to identify and therefore have a faster reaction times and larger distances, 480 and it is not always feasible to control for all possible covarying features. Note that this is also a 481 482 strength of the approach; if colourful objects are indeed easier to identify (i.e., the brain is using the feature), then areas where this feature is represented would have stronger correlations 483 between distance to boundary and reaction times. If a stimulus property is thought to 484 485 (unintentionally) drive decoding and behaviour in the same way, then this property should be 486 controlled for in the stimulus set.

487

## 488 **4.6 Conclusion**

In this study, we combined the distance-to-bound approach (Ritchie & Carlson, 2016) with a searchlight decoding analysis to find brain areas with decodable information that is suitable for read-out in behaviour. Our results showed that decodable information is not always equally suitable for read-out by the brain in behaviour. This speaks to the current debate in neuroimaging research about whether the information that we can decode is the same information that is used by the brain in behaviour (de-Wit et al., 2016).

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