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3 A neural correlate of image memorability

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Some images are easy to remember while others are easily forgotten. While variation in 11 image memorability is consistent across individuals, we lack a full account of its neural 12 correlates. By analyzing data collected from inferotemporal cortex (IT) as monkeys 13 performed a visual memory task, we demonstrate that a simple property of the visual 14 encoding of an image, its population response magnitude, is strongly correlated with its 15 memorability. These results establish a novel behavioral role for the magnitude of the IT 16 response, which lies largely orthogonal to the coding scheme that IT uses to represent 17 object identity. To investigate the origin of IT memorability modulation, we also probed 18 convolutional neural network models trained to categorize objects. We found brain-19 20 analogous correlates of memorability that grew in strength across the hierarchy of these networks, suggesting that this memorability correlate is likely to arise from the 21 optimizations required for visual as opposed to mnemonic processing. 22 23

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26 We have a remarkable ability to remember the images that we have seen, even after a single viewing [1, 2]. Although this capacity appears general and may serve a wide variety of functions, 27 we remember some images better than others [3]. Image memorability is consistent across 28 individuals [3, 4], however, a full account of the sources of image memorability has remained 29 elusive. For example, while some types of natural image content are known to impact 30 memorability – such as images with people, which tend to be more memorable than scenes [3]. 31 and abnormal objects, such as chair shaped like a hand, which tend to be more memorable 32 33 than typical objects [4] - we lack a complete account of how image content determines image 34 memorability.

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36 What neural processes determine memorability? The sources of memorability could range from 37 variation in the perceptual organization of images in visual cortex to the processes that support memory formation and/or memory recall. The neural correlates of memorability are likely to 38 reside at higher stages of the visual form processing pathway, where image memorability can 39 be decoded from human fMRI activity patterns [5, 6], and more memorable images evoke larger 40 fMRI BOLD responses [5]. However, we lack a deeper understanding of how the 41 representations of memorable and non-memorable images differ. Similarly, some insight into 42 the neural correlates of memorability can be gained from convolutional neural network (CNN) 43 models trained for object classification, which have been demonstrated to mimic other (i.e. 44 object identity) representations in the form-processing pathway (reviewed by [7]). Image 45 memorability can be reasonably decoded from the higher layers of at least one of these 46 47 networks [8], but we do not understand how memorability is reflected in this CNN nor whether this CNN reflects memorability like the brain. 48 49

50 The fact that image memorability is linearly decodable in higher visual brain areas such as

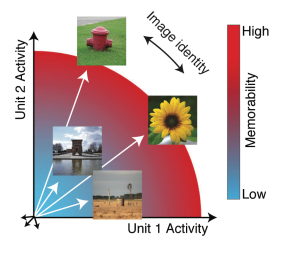
inferotemporal cortex (IT) [5, 6] could imply that information about image memorability is

52 represented in the same fashion as information about object identity in these areas. Within IT, representations of image and object identity are generally thought to be encoded as different 53 54 patterns of spikes across the IT population, consistent with neurons that are individually "tuned" for distinct image and object properties. In a population representational space, these distinct 55 spike patterns translate into population response vectors that point in different directions, and 56 information about object identity is formatted such that it can be accessed from IT neural 57 responses via a weighted linear decoder (Fig. 1a; reviewed by [9]). Similarly, image 58 memorability could be represented by population vector direction in IT. However, under this 59 60 proposal, it is not clear how our experience of image identity and image memorability would be represented as by the same population of neurons, i.e. the fact that one image of a person can 61 be more memorable than another image of that same person, and at the same time, identity 62 63 information, such as the class of an object, explains only a limited amount of how memorable an image will be [3]. 64

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Here we present an alternative proposal, hinted at by the fact that more memorable images 66 evoke larger fMRI responses [5]: we propose that memorability variation is determined 67 principally by the magnitude of the IT population response, or similarly, the total number of 68 spikes across the IT population (Fig. 1). This scenario is consistent with general accounts in 69 which visual perceptual processing precedes memory storage and images that evoke larger 70 71 numbers of spikes, and consequently have more robust visual representations, are remembered best. This scenario incorporates a representational scheme for memorability that is orthogonal 72 to the scheme IT uses to support object identity, and it is thus attractive from the perspective 73 74 that it would provide a straightforward account of how IT multiplexes visual information about image content (as the population vector direction) as well as memorability (as population vector 75 magnitude). The plausibility of this scenario rests on whether there is sufficient variation in 76 population response magnitude across the class of natural images to account for memorability, 77 given the host of homeostatic and normalization mechanisms that act to maintain constant 78 grand mean firing rates across a cortical population [10]. 79

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Figure 1. The hypothesis: the magnitude of the IT population response encodes image memorability. In

geometric depictions of how IT represents image identity, the population response to an image is

depicted as a vector in an N-dimensional space, where N indicates the number of neurons in the

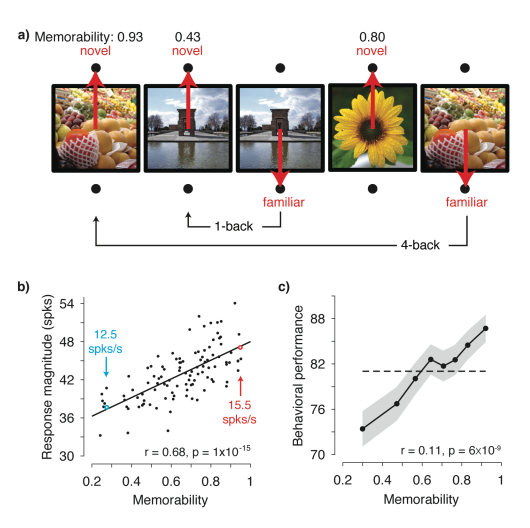
population, and identity is encoded by the direction of the population vector. Here we test the hypothesis

that image memorability is encoded by the magnitude (or equivalently length) of the IT population vector, where images that produce larger population responses are more memorable

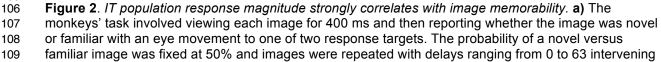
- where images that produce larger population responses are more memorable.
- 91 92

To test the hypothesis presented in Fig. 1, we obtained image memorability scores by passing 93 images through a model designed to predict image memorability for humans ([4]; Supp. Fig. 1). 94 The neural data, also reported in [11], were recorded from IT as two rhesus monkeys performed 95 a single-exposure visual memory task in which they reported whether images were novel (never 96 before seen) or were familiar (seen once previously; Fig. 2a). In each experimental session, 97 neural populations with an average size of 26 units were recorded, across 27 sessions in total. 98 99 After screening for responsive units, data were concatenated across sessions into a larger pseudopopulation in a manner that aligned images with similar memorability scores (see 100 Methods and Supp. Fig. 1). The resulting pseudopopulation contained the responses of 707 IT 101 102 units to 107 images, averaged across novel and familiar presentations. 103







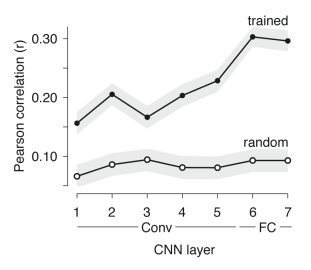


trials (4.5 s to 4.8 min). Shown are 5 example trials with image memorability scores labeled. The 110 111 memorability of each image was scored from 0-1, where the score reflects the predicted chance-112 corrected hit rate for detecting a familiar image (i.e., 0 maps to chance and 1 maps to ceiling, [4]). b) The relationship between image memorability scores and IT population response magnitudes. Each point 113 corresponds to a different image (N=107 images). Population response magnitudes were computed as 114 the L2 norm $\left(\sqrt{\sum_{i=1}^{N} r_i^2}\right)$, where r_i is the spike count response of the ith unit, across a pseudopopulation of 115 707 units. Spikes were counted in an 80 ms window positioned 180 to 260 ms following stimulus onset 116 (see Supp. Fig 2c for different window positions). The Pearson correlation and its p-value are labeled. 117 The solid line depicts the linear regression fit to the data. For reference, the mean firing rates for two 118 119 example images are also labeled (see also Supp. Fig 3b). c) Mean and standard error (across 120 experimental sessions) of monkey behavioral performance on the memory task as a function of humanbased image memorability scores. Performance was binned across images with neighboring memorability 121 scores and pooled across monkeys (see Supp. Fig 4 for plots by individual). The dashed line corresponds 122 to the grand average performance, and if there were no correlation, all points should fall near this line. 123 The point-biserial correlation and its p-value, computed for the raw data (i.e. continuous memorability 124 125 scores and binary performance values for each image in each session) are labeled. 126 127 Fig. 2b shows the correlation between image memorability and IT population response 128 magnitudes, which was strong and highly significant (Pearson correlation: r = 0.68; $p = 1x10^{-15}$). 129 This correlation remained strong when parsed by the data collected from each monkey 130 individually (Supp. Fig. 2a-b) and, after accounting for the time required for signals to reach IT. 131 132 across the entire 400 ms viewing period (Supp. Fig. 3a). The correlation also remained strong when computed for a quantity closely related to response magnitude, grand mean firing rate 133 (Supp. Fig. 3b), as well as when the highest firing units were excluded from the analysis (Supp. 134 Fig. 3c). 135 136 The strength of the correlation between memorability and IT response magnitude is notable

The strength of the correlation between memorability and IT response magnitude is notable
 given the species difference, as the memorability scores were derived from a model designed to
 predict what humans find memorable whereas the neural data were collected from rhesus
 monkeys. Likewise, we found that estimates of human memorability scores were predictive of
 the images that the monkeys found most memorable during the single-exposure visual memory
 task (Fig. 2c).

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As described above, image memorability can be reasonably decoded from at least one CNN 144 145 trained to categorize objects and scenes, but not explicitly to score memorability [8]. This hints at the fact that the neural correlate of memorability variation may be a consequence of the 146 optimizations required for visual (as opposed to mnemonic) processing, however, before making 147 this conclusion, one would want to establish that this CNN reflects memorability in a manner 148 analogous to the brain. We found that this was the case: the correlation between image 149 memorability scores and their corresponding population response magnitudes was significantly 150 higher in the trained as compared to a randomly initialized version of the network in all layers, 151 and the strength of this correlation generally increased across the hierarchy (Fig. 3). These 152 results were also replicated in other CNNs trained for object classification, where correlation 153 strength also systematically increased across the hierarchy throughout much of the network 154 (Supp. Fig. 4), suggesting that this signature is not unique to this particular architecture or 155 training procedure. These results suggest that variation in population response magnitude 156 across images is likely to be a natural consequence of visual systems that classify objects, and 157 that this variation is directly related to variation in image memorability. 158 159



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Figure 3. *Correlations between memorability and population response increase in strength across layers* of a *CNN trained to classify objects and scenes*. Mean and 95% CIs of the Pearson correlations between image memorability and population response magnitude for each hierarchical layer of the CNN described in [8], up to the last hidden layer. "Conv": convolutional layer; "FC": fully connected layer. p-values for a one-sided comparison that correlation strength was larger for the trained than the randomly connected network: p < 0.0001 for all layers.

169 170

171 Discussion

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173 Here we have demonstrated that variation in the ability of humans and monkeys to remember images is strongly correlated with the magnitude of the population response in IT cortex. These 174 results indicate that memorability is reflected in IT via a representational scheme that lies largely 175 176 orthogonal to the one IT uses for encoding object identity (Fig. 1). For example, investigations of how monkey IT and its human analogs represent objects using 'representational similarity 177 analysis' typically begin by normalizing population response vector magnitude to be the same 178 for all images such that all that is left is the direction of the population response pattern, under 179 the assumption that population vector magnitude is irrelevant for encoding object or image 180 181 identity [12]. Before our study, data from human fMRI had pinpointed the locus of memorability to the human analog of IT, but we did not understand "how" the representations of memorable 182 and non-memorable images differed. Our results point to a simple and coherent account of how 183 IT multiplexes representations of visual and memorability information using two complementary 184 representational schemes (Fig. 1). 185

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How might variation in IT population response magnitudes lead to variation in how visual 187 memories are stored? These results are consistent with general accounts of memory in which 188 189 visual processing precedes memory storage and images with more robust visual representations are those that are best remembered. Our results demonstrate that despite the 190 host of homeostatic mechanisms that contribute to maintaining constant global firing rates 191 across a cortical population [10], changes in image content can result in IT population response 192 magnitudes that differ by up to 19% (Fig. 2b; Supp. Fig. 3b). Of course one naturally expects 193 that classes of images that are known to be more robustly represented in IT should be better 194 remembered – for example, natural images should be better remembered than their scrambled 195 counterparts [e.g. 13]. The significance of our result follows from the unexpected finding that 196 there is variation in the robustness of visual representations within the class of natural images 197

that correlates with our understanding of the content that makes images more or less
memorable. For example, unusual objects, such as a chair shaped like a hand, are known to be
more memorable than typical objects, but the fact that unusual objects have more robust visual
representations has not been previously established. As such, our results give insight not only
into visual memorability, but also vision itself.

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Our neural data were recorded from the brains of monkeys that could both see and remember 204 what they had seen. To tease apart whether the origin of memorability could be attributed to 205 206 optimizations for visual as opposed to mnemonic processing, we investigated CNNs optimized to categorize objects but not explicitly trained to predict the memorability of images. Prior to our 207 study, memorability was demonstrated to be linearly decodable from higher layers of one of 208 these CNNs, but it was unclear how memorability was reflected in this CNN and how that 209 compared to the brain. Additionally, while this class of models has been demonstrated to mimic 210 many aspects of how IT represents visual object identity (reviewed by [7]), image memorability 211 has a distinct representational scheme from identity (Fig. 1), and in the context of the many 212 illustrations that CNNs solve the same problems as brains using different strategies (e.g. [14]), it 213 need not have been the case that CNNs reflected memorability in the same way as the brain. 214 The fact that CNNs trained for object recognition mimic the neural representation of a distinct 215 behavior – visual memorability – is compelling evidence that this strategy of multiplexing visual 216 identity and memorability results from the computational requirements of optimizing for robust 217 object representations. These modeling results also offer insight into the nature of the 218 mechanism underlying memorability. The brain perceives and remembers using both 219 220 feedforward and feedback processing, and this processing is modulated by top-down and bottom-up attention. Because of this, it is difficult to pinpoint the locus of an effect like the one 221 we describe to any single mechanism using neural data alone. The fact that variations in 222 223 response magnitudes that correlate with memorability emerge from static, feed-forward, and fixed networks suggests that memorability variation is unlikely to follow primarily from the types 224 of attentional mechanisms that require top-down processing, recurrent processing, or plasticity 225 beyond that required for wiring up a system to identify objects. 226

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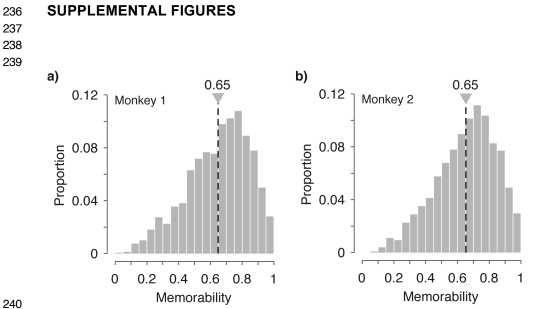
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234 CONFLICTS OF INTEREST

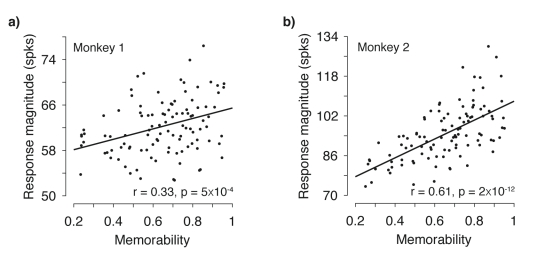
None.





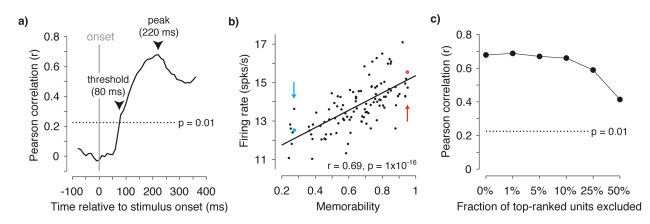
Supplemental Figure 1. Distributions of memorability scores for the images used in these experiments.
 Memorability scores range from 0-1, where the score reflects the predicted chance-corrected hit rate for
 detecting a familiar image and 0 maps to chance (see Methods and [4]).







Supplemental Figure 2. The correlation of memorability and population response magnitude, for each 249 monkey individually. a-b) Fig 2b replotted for each monkey individually (monkey 1: 353 units; monkey 2 250 354 units). To compensate for parsing the data, the spike count window was increased to 250 ms in these 251 plots (positioned 150 ms - 400 ms) relative to the 80 ms window depicted in Fig. 2. The Pearson 252 correlation and its p-value are labeled. The following two points were included in computing the 253 correlations but fall outside the boundaries of the plot or are obscured by text: Monkey 1 (panel a): 254 memorability = 0.86, response magnitude = 83.6; Monkey 2 (panel b): memorability = 0.57, response 255 256 magnitude = 71.5. Solid lines depict the linear regression fits to the data. 257

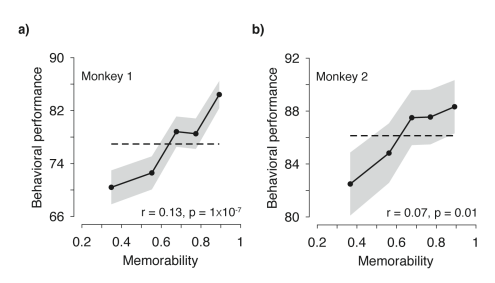


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Supplemental Figure 3. The correlation of memorability and the IT population response, applied to 260 different time windows, assessed with firing rate, and determined with top-ranked firing units removed. a) 261 The same analysis described for Fig. 2b, but applied to 80 ms windows shifted at different positions 262 relative to stimulus onset, where the correlations are plotted against the center of each time bin. Fig. 2b is 263 shown at the peak of this plot (220 ms). Also shown (dotted line) is the critical correlation threshold for p < 264 0.01, which the population reached at 80 ms following stimulus onset. b) Correlations between 265 266 memorability and grand mean firing rate across the 707 units (in contrast to the plots of response 267 magnitude in Fig. 2b). The two example images from 2b are indicated. Solid line depicts the linear regression fit to the data. c) The analysis in Fig. 2b with N% top-ranked firing rate units excluded from the 268 pseudopopulation for different N. The dotted line indicates the critical correlation for the significance level 269 p = 0.01. 270 271



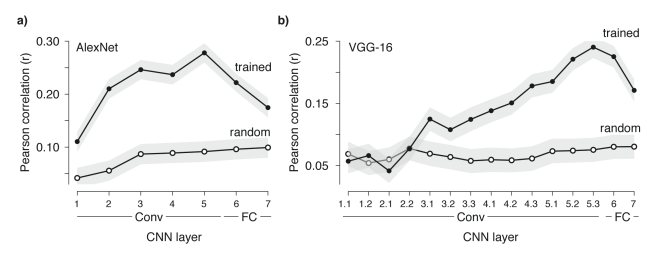




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Supplemental Figure 4. Human-based memorability scores predict what monkeys find memorable. The 275 analysis presented in Fig. 2c, applied to each monkey individually. To compensate for parsing the data, 276 the data is parsed into 5 bins as opposed to the 7 bins in Fig. 2c. The dashed lines correspond to the 277 grand average performance, and if there were no correlation, all points should fall near this line. The 278 point-biserial correlation and its p-value, computed for the raw data, are labeled. 279

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Supplemental Figure 5. Correlations between memorability and population response magnitude are 284 also reflected in two other CNNs. Mean and 95% CIs of the Pearson correlations between image 285 286 memorability and population response magnitude for each hierarchical layer for two CNNs, including a) AlexNet [15], b) VGG-16 [16], up to the last hidden layer. "Conv": convolutional layer; "FC": fully 287 connected layer, p-values for a one-sided comparison that correlation strength was larger for the trained 288 than the randomly connected network, AlexNet: p < 0.0001 for all layers; VGG-16: p = 0.8, 0.2, 0.9, and 289 0.5 for Conv 1.1, 1.2, 2.1, and 2.2, respectively, p = 0.0008 for Conv 3.2, and p < 0.0001 for all other 290 291 layers.

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1 **METHODS:**

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As an overview, three types of data are included in this paper: 1) Behavioral and neural data 3 collected from two rhesus monkeys that were performing a single-exposure visual memory task; 4 5 2) Human-based memorability scores for the images used in the monkey experiments, and 3) The responses of units at different layers of three convolutional neural network models trained 6 7 to classify objects and scenes. The Methods associated with each type of data are described below. 8 9

- 10 Behavioral and neural data collected from two rhesus monkeys that were performing a single-exposure visual memory task 11
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Experiments were performed on two adult male rhesus macaque monkeys (Macaca mulatta) 13 with implanted head posts and recording chambers. All procedures were performed in 14 15 accordance with the guidelines of the University of Pennsylvania Institutional Animal Care and Use Committee. Monkey behavioral and neural data were also included in an earlier report that 16 examined the relationship between behavioral reports of familiarity as a function of the time 17 between novel and familiar presentations (e.g., "rates of forgetting") and neural responses in IT 18 cortex [1]. The results presented here cannot be inferred from that report. 19 20 The single-exposure visual memory task: 21 22

All behavioral training and testing were performed using standard operant conditioning (juice 23 reward), head stabilization, and high-accuracy, infrared video eye tracking. Stimuli were 24 25 presented on an LCD monitor with an 85 Hz refresh rate using customized software (http://mworks-project.org). 26

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Each trial of the monkeys' task involved viewing one image for at least 400 ms and indicating 28 whether it was novel (had never been seen before) or familiar (had been seen exactly once) 29 30 with an eye movement to one of two response targets. Images were never presented more than twice (once as novel and then as familiar) during the entire training and testing period of the 31 experiment. Trials were initiated by the monkey fixating on a red square (0.25°) on the center of 32 a gray screen, within an invisible square window of $\pm 1.5^{\circ}$, followed by a 200 ms delay before a 33 4° stimulus appeared. The monkeys had to maintain fixation of the stimulus for 400 ms, at which 34 time the red square turned green (go cue) and the monkey made a saccade to the target 35 indicating that the stimulus was novel or familiar. In monkey 1, response targets appeared at 36 37 stimulus onset; in monkey 2, response targets appeared at the time of the go cue. In both cases, targets were positioned 8° above or below the stimulus. The association between the 38 target (up vs. down) and the report (novel vs. familiar) was swapped between the two animals. 39 The image remained on the screen until a fixation break was detected. The first image 40 presented in each session was always a novel image. The probability of a trial containing a 41 novel vs. familiar image quickly converged to 50% for each class. Delays between novel and 42 familiar presentations were pseudorandomly selected from a uniform distribution, in powers of 43 two (n-back = 1, 2, 4, 8, 16, 32 and 64 trials corresponding to mean delays of 4.5s, 9s, 18s, 36s, 44 1.2 min, 2.4 min, and 4.8 min, respectively). 45 46

The images used in these experiments were collected via an automated procedure that 47

- downloaded images from the Internet. Images smaller than 96*96 pixels were not considered 48
- and eligible images were cropped to be square and resized to 256*256 pixels. An algorithm 49

50 removed duplicate images. Within the training and testing history for each monkey, images were 51 not repeated.

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The activity of neurons in IT was recorded via a single recording chamber in each monkey. 53 Chamber placement was guided by anatomical magnetic resonance images in both monkeys. 54 The region of IT recorded was located on the ventral surface of the brain, over an area that 55 56 spanned 5 mm lateral to the anterior middle temporal sulcus and 14-17 mm anterior to the ear canals. Recording sessions began after the monkeys were fully trained on the task and after the 57 depth and extent of IT was mapped within the recording chamber. Combined recording and 58 behavioral training sessions happened 4-5 times per week across a span of 5 weeks (monkey 59 1) and 4 weeks (monkey 2). Neural activity was recorded with 24-channel U-probes (Plexon, 60 61 Inc) with linearly arranged recording sites spaced with 100 µm intervals. Continuous, wideband neural signals were amplified, digitized at 40 kHz and stored using the Grapevine Data 62 Acquisition System (Ripple, Inc.). Spike sorting was done manually offline (Plexon Offline 63 Sorter). At least one candidate unit was identified on each recording channel, and 2-3 units were 64 65 occasionally identified on the same channel. Spike sorting was performed blind to any experimental conditions to avoid bias. For quality control, recording sessions were screened 66 based on their neural recording stability across the session, their numbers of visually responsive 67 units, and the numbers of behavioral trials completed. A multi-channel recording session was 68 included in the analysis if: (1) the recording session was stable, guantified as the grand mean 69 firing rate across channels changing less than 2-fold across the session; (2) over 50% of 70 neurons were visually responsive (a loose criterion based on our previous experience in IT). 71 assessed by a visual inspection of rasters; and (3) the number of successfully completed 72 novel/familiar pairs of trials exceeded 100. In monkey 1, 21 sessions were recorded and 6 were 73 removed (2 from each of the 3 criterion). In monkey 2, 16 sessions were recorded and 4 were 74 removed (1, 2 and 1 due to criterion 1, 2 and 3, respectively). The resulting data set included 15 75 76 sessions for monkey 1 (n = 403 candidate units), and 12 sessions for monkey 2 (n = 396candidate units). Both monkeys performed many hundreds of trials during each session (~600-77 1000, corresponding to ~300-500 images each repeated twice). The data reported here 78 correspond to the subset of images for which the monkeys' behavioral reports were recorded for 79 both novel and familiar presentations (e.g. trials in which the monkeys did not prematurely break 80 fixation during either the novel or the familiar presentation of an image). Finally, units were 81 82 screened for stimulus-evoked activity via a comparison of their responses in a 200 ms period before stimulus onset (-200 ms - 0 ms) versus after stimulus onset (80 - 280 ms) with a two-83 84 sided t-test, p < 0.01. This yielded 353 (of 403) units for monkey 1 and 354 (out of 396) units for monkey 2. 85

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To perform our analyses, we concatenated units across sessions to create a larger 87 pseudopopulation. In the case of the pooled data, this included 27 sessions in total (15 sessions 88 from monkey 1 and 12 from monkey 2). When creating this pseudopopulation, we aligned data 89 across sessions in a manner that preserved whether the trials were presented as novel or 90 familiar, their n-back separation, and image memorability scores (obtained using methods 91 described below). More specifically, the responses for each unit always contained sets of 92 93 novel/familiar pairings of the same images, and pseudopopulation responses across units were always aligned for novel/familiar pairs that contained the same n-back separation and images 94 95 with similar memorability scores. When the number of images in a session exceeded the number required to construct the pseudopopulation, a subset of images were selected 96 separately for each n-back by ranking images within that n-back by their memorability scores, 97

98 preserving the lowest-ranked and highest-ranked images within that session, and selecting the

number of additional images required as those with memorability scores that were evenly

spaced between the two extreme memorability scores for that session. The resulting

101 pseudopopulation consisted of the responses to 107 images presented as both novel and

102 familiar (i.e. 15, 15, 16, 17, 17, 15 and 12 trials at 1, 2, 4, 8, 16, 32 and 64-back, respectively).

To perform the neural analyses (Fig 2b, Supp Figs 2, 3), a memorability score for each of the

104 107 pseudopopulation images was computed as the mean of the memorability scores across all 105 the actual images that were aligned to produce that pseudopopulation response. The average

standard deviation across the set of memorability scores used to produce each

pseudopopulation response was 0.05, where memorability ranges 0-1. To perform behavioral
 analyses (Fig 2c, Supp Fig 4), the memorability score as well as binary performance values
 (correct/wrong at reporting that a familiar image was familiar) were retained for each of the 107

- images, across each of the 27 sessions.
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112 **Human-based memorability scores for the images used in the monkey experiments**

We obtained memorability scores for the images used in the monkey experiments using 114 MemNet [2] estimates. MemNet is a convolutional neural network (CNN) trained to estimate 115 image memorability on a large-scale dataset of natural images (LaMem [2], publicly available at 116 memorability.csail.mit.edu). LaMem consists of 60K images drawn from a diverse range of 117 sources (See [2] for more detail). Each image in this dataset is associated with a memorability 118 score based on human performances in an online memory game on Amazon's Mechanical Turk. 119 Behavioral performances were corrected for the delay interval between first and second 120 121 presentation to produce a single memorability score for each image. After training, MemNet estimates visual memorability of natural images near the upper bound imposed by human 122 performance: MemNet estimates reach 0.64 rank correlation with mean human-estimated 123 memorability, while the upper bound of consistency between human scores has a rank 124 125 correlation of 0.68. Here we treat MemNet memorability estimates as a proxy for human 126 memorability scores. 127 The memorability scores were obtained using the network weights reported in [2] and publicly 128 129 available at http://memorability.csail.mit.edu/download.html. This network was originally trained

using the Caffe framework [3], and we ported the trained network to Pytorch [4] using the caffe-

131 to-torch-to-pytorch package at <u>https://github.com/fanq15/caffe_to_torch_to_pytorch</u>. Before

passing images into MemNet, we preprocessed them as described in [5]: we resized images to
 256 × 256 pixels (with bilinear interpolation), subtracted the mean RGB image intensity

(computed over the dataset used for pretraining, as described in [5]), and then produced 10
crops of size 227 × 227 pixels. The 10 crops were obtained by cropping the full image at the
center and at each of the four corners and by flipping each of these 5 cropped images about the
vertical axis. All 10 crops were passed through MemNet. The average of these 10 scores was
used as the mean prediction of the model for the input image. This mean prediction was then
linearly transformed to obtain the estimated memorability score:

- 140
- 141 142

Memorability_score = min (max ((output - mean_pred)*2 + additive_mean, 0), 1)

where following [2], we set mean_pred = 0.7626 and additive_mean = 0.65.

The responses of units at different layers of CNN models trained to classify objects and 145 146 scenes.

147

We evaluated the correlation between response magnitude and image memorability on images 148 from the LaMem dataset [2] using three commonly used convolutional neural networks (CNNs). 149 All reported models were evaluated on the full test set of split 1 of LaMem, which contains 150 10,000 images. We chose to use LaMem images, as each image in this dataset is labeled with 151 a memorability score computed directly from human behavioral performance (i.e. not estimated 152 with a model; see above and [2] for details of data collection and memorability score 153 computation). All networks were run in TensorFlow 1.10 ([6], software available 154 from tensorflow.org), using custom Python evaluation code. 155

156

The results presented in Fig 3 were obtained by running images from this dataset through 157

- HybridCNN [5]. HybridCNN is a network with an identical architecture to AlexNet [7]. HybridCNN 158 was first trained to classify natural images of objects and scenes using data from the ImageNet
- 159 Large Scale Visual Recognition Challenge (ILSVRC) 2012, a 1000-way object classification
- 160
- dataset [8], as well as the Places 183-way scene classification dataset [5], for a combined 1183-161
- 162 way classification task. For details of training, see [5]. Results were obtained using the network
- weights reported in [5] and publicly available at http://places.csail.mit.edu/downloadCNN.html. 163 This network was originally trained using the Caffe framework [3], and we ported the trained 164
- network to TensorFlow using the caffe-tensorflow package https://github.com/ethereon/caffe-165

166 tensorflow. Random initialization baselines were obtained using the same architecture, but

- randomly sampling the weights using the initialization algorithm described in [9]. 167
- 168

Before passing images into each network, we preprocessed them as described in [5] and 169 above: we resized images to 256 x 256 pixels (with bilinear interpolation), subtracted the mean 170 171 RGB image intensity (computed over the training dataset), and then cropped the central 227 x 227 and passed it into the network. The response magnitude (L2 norm) of each layer was 172 computed over the full output vector of each hidden layer. In all cases, we show the magnitude 173 of hidden layer output after applying the nonlinear operation. Results for the two networks 174 presented in the supplement (Supp Fig 5) were obtained in an identical manner, except for the 175 176 image preprocessing step. For each network, images were preprocessed as described in the original papers (AlexNet: [7], VGG-16: [10]). 177

178

For all three networks (HybridCNN, AlexNet, and VGG-16), we computed correlations for all 179 convolutional and fully-connected hidden layers. The Pearson correlation coefficient was used 180 181 to measure correlation. All correlations were computed over the full set of 10,000 images described above. 95% confidence intervals for the correlation coefficient of each layer were 182 183 obtained by bootstrapping over the set of 10,000 per-image layer magnitudes and memorability scores. 95% confidence intervals were estimated empirically as the upper and lower 97.5%-184 centiles of the bootstrapped correlation coefficients for each layer and condition. Bootstrapped 185 resampling was performed independently for each layer and each condition (trained or randomly 186 187 connected). In all cases, bootstrap estimates were performed using 10,000 samples (with replacement) of the full dataset of 10,000 images. The bootstrapping procedure was also used 188 to conduct one-tailed tests to determine whether the correlations between memorability and 189 response magnitude were stronger in the trained as compared to the randomly initialized 190 191 network at each layer separately. p-values were estimated by taking pairs of correlation coefficients computed on the bootstrapped data for each condition and measuring the rate at 192 which the correlation for the random layer exceeded the correlation for the trained layer. 193

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