Analysis of convolu	tional	neural netwo	rks reveals the computational properties
essentia	l for su	ubcortical pr	ocessing of facial expression
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37 Abstract

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39 Perception of facial expression is crucial in the social life of primates. This visual information 40 is processed along the ventral cortical pathway and the subcortical pathway. Processing of face 41 information in the subcortical pathway is inaccurate, but the architectural and physiological properties that are responsible remain unclear. We analyzed the performance of convolutional 42 43 neural networks incorporating three prominent properties of this pathway: a shallow layer 44 architecture, concentric receptive fields at the first processing stage, and a greater degree of 45 spatial pooling. The neural networks designed in this way could be trained to classify seven 46 facial expressions with a correct rate of 51% (chance level, 14%). This modest performance was 47 gradually improved by replacing the three properties, one-by-one, two at a time, or all three 48 simultaneously, with the corresponding features in the cortical pathway. Some processing units 49 in the final layer were sensitive to spatial frequencies (SFs) in the retina-based coordinate, 50 whereas others were sensitive to object-based SFs, similar to neurons in the amygdala. Replacement of any one of these properties affected the SF coordinate of units. All three 51 52 properties constrain the accuracy of facial expression information in the subcortical pathway, 53 and are essential for determining the coordinate of SF representation.

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56 Keywords

57 Subcortical visual pathway; ventral pathway, convolutional neural network; amygdala; face 58 recognition

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60 Introduction

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62 Perceiving the facial expressions of other individuals plays a critical role in the social life of primates, including humans. Two neural pathways, the ventral cortical pathway and the 63 64 subcortical pathway, contribute to this perceptual ability (Fig. 1A; Pessoa and Adolphs, 2010; Tamietto and de Gelder, 2010; Petray and Bickford, 2019). The ventral cortical pathway consists 65 of a network of areas in the occipito-temporal region of the cerebral cortex, and processes a 66 variety of visual features of objects, people, and environments, including shape, color, texture, 67 68 material properties, and binocular disparity (Ungerleider and Mishkin, 1982; Connor et al., 69 2007; Conway et al., 2010; Roe et al., 2012; Kravitz et al., 2013; Vaziri et al., 2014; Verhoef et 70 al., 2016; Komatsu and Goda, 2018). Neurons that preferentially respond to images of faces or 71 facial features are found in several clusters along this pathway (Desimone et al., 1984; Perrett et al., 1987; Fujita et al., 1992; Haxby et al., 2000; Tsao and Livingstone, 2008; Duchaine and 72 73 Yovel, 2015; Freiwald et al., 2016). They constitute the neural system that analyzes facial details 74 such as expression, identity, and direction of attention. The subcortical pathway consists of a 75 few processing stages in phylogenetically ancient regions: the superior colliculus in the 76 midbrain, the pulvinar nucleus in the posterior thalamus, and the amygdala in the medial limbic 77 system. The subcortical pathway is suggested to mediate rapid behavioral and physiological 78 (autonomic) responses to sensory signals related to possible dangers such as fearful faces 79 (Tamietto and de Gelder, 2010; Nakano et al., 2013; for a critical review, see Pessoa and Adolphs, 2010). The ventral cortical pathway and the subcortical pathway intersect at the 80 amygdala. 81

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Psychological and brain imaging studies suggest that the subcortical pathway subserves the 83 84 ability of some patients with lesions in the primary visual cortex (V1) to discriminate facial expressions despite lacking visual awareness ("affective blindsight"; deGelder et al., 1999; 85 Pegna et al., 2005; Striemer et al., 2019). These patients also reflexively exhibit specific facial 86 87 expressions and pupillary reactions when exposed to fearful or happy faces (Tamietto et al., 88 2009). Studies have also shown that the subcortical pathway supports unconscious face perception in neurologically healthy subjects (Morris et al., 1999, 2001). Furthermore, 89 90 orientation bias toward faces or face-like patterns by newborn babies is suggested to be mediated 91 by the subcortical pathway (Cassia et al., 2001; Johnson, 2005; but see Buiatti et al., 2019). 92 Importantly, these perceptual abilities are not perfectly accurate, instead resulting in modest 93 performance at above-chance levels. These findings suggest that information on faces conveyed 94 by the subcortical pathway is less accurate than that carried by the ventral cortical pathway.

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Electrophysiological studies have demonstrated that processing of facial expression in the 96 97 subcortical pathway is indeed fast and not very accurate. Méndez-Bértolo and colleagues (2016) 98 showed that intracranial local field potentials in the human amygdala respond differentially to 99 fearful faces versus other faces within 74 ms after stimulus onset. A recent single-neuron 100 recording study in the monkey revealed that a population of amygdala neurons responded to 101 threatening faces even within 50 ms (Inagaki et al., 2022a). This early response, when combined 102 across an ensemble of neurons, carries information that allows linear classifiers to discriminate threatening faces from neutral and affiliative faces. The rate of correctly discriminating the three 103 104 expressions is around 50%; this is well above chance (33%), but significantly worse than perfect.

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Some neurons in the superior colliculus and pulvinar of the monkey also respond to faces and face-like patterns with an even shorter latency of 30–50 ms (Nguyen et al., 2013, 2014).

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108 What architectural and physiological properties of the subcortical pathway are responsible for 109 its fast, crude processing? The fast processing most likely arises from the small number, or 110 "shallowness," of processing stages in the subcortical pathway, given that the ventral cortical pathway and its upstream area (the lateral geniculate nucleus) consist of a larger number of 111 regions (at least six before reaching the amygdala) than the subcortical pathway (only two), and 112 113 that every transition from one cortical region to the next takes at least 10 ms (Schmolesky et al., 114 1998). It is unclear whether the shallow processing similarly explains the low accuracy of the 115 information transmitted by the subcortical pathway to the amygdala. This uncertainty arises 116 from the fact that in addition to the difference in the number of processing stages, visual 117 response properties differ markedly between the two pathways. Neurons in the superior 118 colliculus at the first stage of the subcortical pathway show circular receptive fields with center-119 surround organization, which can be modeled using the difference-of-Gaussian (DoG) function 120 (Cynader and Berman 1972; Updyke 1974; Marino et al. 2008; Churan et al., 2012). By contrast, 121 simple cells of V1 at the first stage of the cortical pathway have elongated receptive fields with 122 side-by-side ON and OFF subregions, which can be modeled by two-dimensional Gabor 123 functions (Jones and Palmer, 1987). Furthermore, the receptive field is typically larger in the 124 superior colliculus (for the foveal field, 1.5–10° in superficial layers, 10–20° in deep layers; Goldberg and Wurtz, 1972; Wallace et al., 1997) than in V1 (1.18° in simple cells, 1.3° in 125 complex cells; Van den Bergh et al., 2010) and the extrastriate areas V2 and V4 (Freeman and 126 127 Simoncelli, 2011). Thus, spatial pooling across ascending stages occurs over a wider visual field area in the subcortical pathway than in the ventral cortical pathway. 128

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130 In the present study, we addressed how these properties of the subcortical pathway, i.e., the 131 shallowness of processing stages, DoG-type receptive fields at the initial stage, and spatial 132 pooling over a wider visual field, influence facial expression processing. To this aim, we 133 constructed convolutional neural networks (CNNs) and analyzed their performance in facial 134 expression discrimination. CNNs are one type of multilayer perceptron, and can be optimized 135 ("learn") to classify inputs by varying connection weights between processing units through 136 supervised learning algorithms (LeCun et al., 2015). Typical CNNs have several to tens of layers (deep neural networks, DNNs). DNNs developed for classifying visual objects share 137 architectural and representational features similar to those of the ventral cortical pathway 138 139 (Yamins et al., 2014; Güçlü and van Gerven, 2015; Yamins and DiCarlo, 2016; Hassabis et al., 140 2017). We designed our CNNs to imitate the subcortical pathway by reducing the number of 141 processing stages and by implementing DoG-type receptive fields and a wider extent of pooling. 142 These CNNs, hereafter referred to as shallow neural networks (SNNs), learned to discriminate 143 facial expressions with modest correct rates. Replacing the three properties, one-by-one, two at 144 a time, or all three simultaneously, with the corresponding properties in the ventral cortical 145 pathway gradually improved discrimination performance, suggesting that all three features are 146 responsible for limiting the performance of the SNNs. We further showed that like some neurons 147 in the amygdala, a major group of units in the final processing layer of the SNNs were sensitive 148 to spatial frequency (SF) in the retina-based reference frame as initially detected in the first

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149 processing layer, and that the three subcortical properties contribute to preserving the retina-

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153 Materials and Methods

based SF sensitivity.

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155 Architecture of SNNs.

We constructed SNNs incorporating the distinct properties of the primate subcortical pathway 156 157 (Fig. 1B–D; Table 1). Unlike typical DNNs, the SNNs consisted of only two sets of convolution 158 and pooling layers followed by two fully connected layers (FC1, FC2), approximating the small 159 number of processing stages of the subcortical pathway. The first convolution layer incorporated 160 32 DoG-type filters (Fig. 1C, top) with a spatial resolution of 11×11 pixels, whereas weights in the second convolution layer were initially random, i.e., the filters had no structure, and 161 162 gradually changed through training. A rectified linear unit (ReLU) was used as the activation function of a unit in the convolution layers and FC1; the ReLU forwards the processing results 163 164 directly to the next stage if they are positive, otherwise it outputs zero. A max pooling operation 165 was performed over 5×5 sliding regions with a stride of 4 for the outputs of convolution layers (Fig. 1D, top). Max pooling selected the largest value among the responses of units within a 166 167 sliding window over the preceding convolution laver, and forwarded the value to the next laver. 168 A local response normalization process was added after the pooling layers to aid generalization (Krizhevsky et al., 2012; we used slightly different parameters from theirs; k = 1, n = 5, $\alpha = 2 \times$ 169 10^{-5} , $\beta = 0.75$). Every unit in FC1 and FC2 received inputs from all units in the immediately 170 171 preceding layer, i.e., each was fully connected. FC1 is the final processing layer, and FC2 172 outputs the results of entire processing by the SNNs. These features were implemented to 173 capture the architectural and computational properties of the subcortical pathway, i.e., fewer 174 processing stages compared to the ventral cortical pathway (Fig. 1A), DoG-type receptive fields 175 in the superficial layer of the superior colliculus (Churan et al., 2012), and large receptive fields of deeper superior colliculus neurons (Wallace et al., 1997). The first three processing layers 176 177 were intended to represent the superior colliculus, pulvinar, and amygdala, respectively. The 178 processing types of these layers, i.e., convolution and pooling in the first two layers and full 179 connection in FC1, were chosen to match the retinotopic organization of the three brain regions. 180 The convolution and pooling processes in the first two layers exploit retinotopy, as the superior colliculus and pulvinar contain retinotopic maps (Bender, 1981; Chen et al., 2019). The FC1 181 182 layer loses retinotopic information because of the fully convergent connection from the earlier 183 stage, as the amygdala does not have a retinotopic map (Morawetz et al., 2010).

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The SNNs were trained to discriminate images of facial expressions representing seven basic emotions: angry, disgusted, fearful, happy, sad, surprised, and neutral (Fig. 1E; see below for details). For each input image, the seven units in FC2 yielded scores ranging from 0 to 1 for the seven expression categories, representing the probabilities of classified expressions. The expression with the highest score was taken as the output of the model.

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193 Figure 1. Shallow neural networks (SNNs) and modifications. (A) Cortical and subcortical 194 visual pathways for processing facial expressions in the primate brain. AMYG: amygdala; AIT, 195 PIT: anterior and posterior parts of inferior temporal cortex; LGN: lateral geniculate nucleus; 196 PU: pulvinar nucleus; SC: superior colliculus; V1, V2, V4: visual areas 1, 2, 4. (B) A schematic 197 illustration of the SNNs and full-replacement models. In the full-replacement models, 198 processing layers were added, the filters in the initial layer were changed with Gabor filters, and 199 the range of pooling was narrowed. (C) DoG filters for the SNNs and DoG models (upper) and 200 Gabor filters for the Gabor models (lower). (D) The pooling range for the SNNs (5×5) and the 201 narrow-pooling models (3×3) . (E) Examples of presented face images with seven expressions 202 (angry, disgusted, fearful, happy, sad, surprised, neutral) of two individuals (upper: female, 203 AF06; lower: male, AM27). The original images were obtained from the Karolinska Directed 204 Emotional Faces database (Lundqvist et al., 1998).

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206	<i>Table 1.</i> Architecture of the Shallow Neural Network (SNN). Each row describes a layer <i>i</i> with
207	calculation operator Fi, output resolution $H_i \times W_i$, and the number of output channels C_i . Conv
208	denotes convolution layer, and Pool denotes max pooling layer.
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Input &	Operator	Resolution	#Channels
Layer	F_i	$H_i \times W_i$	C_i
i			
Input		227×227	3
Layer 1	11×11 Conv (stride 4) & 5×5 Pool (stride 4)	55 × 55	32
Layer 2	5×5 Conv (stride 1) & 5×5 Pool (stride 4)	4×4	64
Layer 3	Fully connected	1×1	32
Layer 4	Fully connected	1×1	7

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213 The DoG-type filters of the first convolution layer were built using the following formula:

- 214 215 $DoG(r) = \pm A_1 exp\left(-\frac{r^2}{2\sigma_1^2}\right) \mp A_2 exp\left(-\frac{r^2}{2\sigma_2^2}\right),$
- 216

where *r* is the polar radius from the filter center, A_1 and A_2 are the amplitudes of exponentials of two Gaussian functions, and σ_1 and σ_2 are the standard deviations. Values of A_1 , A_2 , σ_1 , and σ_2 were chosen empirically so that DoG curves took the shapes of Mexican hats. A_1 values were 0.4, 0.67, 0.8, and 1.0. A_2 values were determined based on $A_1 - A_2 = 0.4$. When A_1 was 0.4 (i.e., A_2 is 0, and σ_2 cannot be defined), we set the σ_1 value at $1/2\sqrt{2}$, $1/4\sqrt{2}$, $1/8\sqrt{2}$, or $1/16\sqrt{2}$. Otherwise, the σ_1 value was $1/2\sqrt{2}$ or $1/4\sqrt{2}$. The σ_2 value was based on $\sigma_1/\sigma_2 = 0.5$ or 0.25. The same number of filters was generated for each A_1 value.

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225 We also constructed modified models in which the three properties of the SNNs were replaced 226 one-by-one, two at a time, or all three simultaneously with the corresponding properties in the 227 ventral cortical pathway. First, we added additional convolution layers with filters of 3×3 pixels 228 after each of the first two convolution layers to increase the number of processing stages (add-229 layer model). In adding the convolution layers, the stride of sliding filters was reduced to 1 to 230 keep the output resolutions unchanged before and after adding the new layers. Also, to keep the 231 number of the output channels unchanged, the new layers contained the same number of filters 232 as the preceding layers.

233

Second, we replaced the DoG-type filters with Gabor-type filters (Gabor model). Gabor-type
filtering occurs in simple cells of V1, and emerges in the first layer of DNNs after they are
trained to classify object images (Krizhevsky et al., 2012; Rai and Rivas, 2022). We constructed
the Gabor-type filters with the following formula:

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$$g(x, y; f, \theta) = Aexp(-\frac{x^2 + y^2}{2\sigma^2})exp(2\pi i f(x\cos\theta + y\sin\theta)),$$
240 (2)

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241 where *A* is the amplitude of the Gaussian envelope, *i* is $\sqrt{-1}$, *f* is the carrier frequency of a Gabor 242 filter, and θ is the orientation (Movellan, 2002). *A* was fixed at 0.4 to match the amplitude of 243 the DoG filters. σ was fixed at 0.125 so that the half-amplitude width was half of the filter width.

244 f values were 2 or 4 cycles/image. Orientation θ was 0, 22.5, 45, 67.5, 90, 112.5, 135, or 157.5.

245 We built even- and odd-symmetric filters for every combination of variables. In total, we

- obtained 32 Gabor-type filters (Fig. 1C, bottom).
- 247

Finally, we made the pooling window (convergence field) in the max pooling layers smaller (3 × 3; Fig. 1D, bottom) than that of the SNNs (5 × 5; Fig. 1D, top), enabling better spatial resolution of processing (narrow-pooling model) to mimic the smaller receptive fields in the visual cortices compared to the superior colliculus and pulvinar (Wallace et al., 1997; Vand den Bergh et al., 2010; Freeman and Simoncelli, 2011). The pooling range of 3×3 is often used in DNNs (e.g., AlexNet of Krizhevsky et al., 2012; ResNet of He et al., 2016).

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255 Face images and training of the SNNs.

256 Face images were obtained from Karolinska Directed Emotional Faces (KDEF; Lundqvist et al., 257 1998) and Radboud Faces Database (RaFD; Langer et al., 2010). Images of the seven 258 expressions of 40 individuals (half females, half males) were chosen from each database (the 259 total number of images was $560 = 7 \times 40 \times 2$). We converted the images from color into 260 grayscale, and extracted the face region by removing hair, neck, and ears with the face-detection function of a computer vision library, OpenCV (Open Source Computer Vision Library; Bradski, 261 2000). The isolated faces were pasted on a gray background (198×198 pixels; RGB values = 262 263 128; Fig. 1E). We augmented the number of face images by changing size and position, and by flipping horizontally; seven sizes $(28 \times 28, 56 \times 56, 85 \times 85, 113 \times 113, 141 \times 141, 170 \times 170,$ 264 265 and 198×198 pixels), five positions (center, left-top, right-top, left-bottom, and right-bottom; directional displacements = 10 pixels), and two horizontally flipped images. The augmentation 266 267 increased the number of images by 70 times to 39,200. At each training session, we randomly 268 split this augmented set of face images into a training set (29,400 images), a validation set (2,450 269 images), and a test set (7,350 images). The number of images per facial expression was identical 270 within each of these stimulus sets. To avoid the inadvertently biased assignment of face images 271 of a particular size, position, or horizontal flip state into a given set, all images from the same 272 individual were assigned into the same set.

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The training was performed through supervised learning, and was conducted individually 20 274 275 times with randomized initial weights except for the built-in weights of the fist convolution layer, 276 i.e., 20 SNNs with different initial states were built. In training, the weights other than the first 277 convolution layer were optimized for classification of face images into the seven categories. 278 Stochastic gradient descent was used for weight optimization. For each iteration, 32 samples 279 were randomly selected from the training set as a mini-batch. The averaged cross-entropy 280 (Goodfellow et al., 2016) across the 32 images in a mini-batch was calculated as an estimate of 281 loss value, which is a measure of the difference between a model output and a supervised signal, 282 and is used for quantifying the training effect. The number of iterations (i.e., weight-updating 283 processes with single mini-batches) was set at 240,000. Initial weight parameters followed a 284 normal distribution with a mean of 0 and a standard deviation of $\sqrt{(2/N)}$ (N is the total number of weights; He et al., 2015). Weights were updated at each iteration with a constant learning rate 285

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of 0.001. This learning rate was determined empirically; a preliminary analysis based on 10 286 constructed SNNs (different from the 20 SNNs in the main analysis) revealed no decrease in 287 288 loss values (i.e., no learning) with a learning rate of 0.01, which has frequently been used for 289 DNNs in the literature (e.g., Simonyan and Zisserman, 2014). A dropout process was added 290 before FC2 to facilitate learning across all units. The proportion of units dropped out of each 291 weight update was set to 0.5. The training was conducted in a Python environment (Chainer 3.0.0; Tokui et al., 2015) on a graphics processing unit (GPU) machine (Intel[®] CoreTM i7-5820K 292 Processor. Intel, Santa Clara, CA, USA; The GeForce® GTX 1080 Ti, NVIDIA, Santa Clara, 293 CA, USA). While the SNNs were being trained with the training set, the correct rate and loss 294 295 value for the validation set were periodically checked to monitor signs of overfitting. After 296 training was completed, the performance of the models was evaluated using the test dataset that 297 had not been used for training. This was done to ensure that the models acquired a genuine 298 ability to classify the facial expressions, as opposed to simply sorting the training images into 299 the seven facial expression categories according to the instruction signals.

300

301 Test for reference frames of SF tuning of model units.

302 A difference in visual responses between the two pathways is the reference frame of neuronal 303 tuning to SFs (Inagaki and Fujita, 2011). Neurons in the inferior temporal cortex, the final stage 304 of the ventral cortical pathway, are tuned to object-based SFs (cycles/object) and represent face 305 patterns in a size-invariant, hence distance-invariant, manner (Fig. 2A, right). Thus, the ventral 306 cortical pathway converts the representation of SFs in the retina-based coordinate 307 (cycles/degree) to that of object-based SFs. By contrast, many amygdala neurons preserve 308 sensitivity to retina-based SFs. When the stimulus size is changed, these neurons change their 309 preferred object-based SFs; for large stimuli, they respond to higher object-based SFs, which 310 correspond to the same retina-based SFs (Fig. 2A, left). We analyzed the reference frame of 311 FC1 SF tuning to evaluate how well our models captured this characteristic of subcortical 312 processing.

313

314 Bandpass-filtered face images were used to examine the SF tunings of FC1 units (Fig. 2B). 315 These images were created by multiplying Gaussian functions with the original face images on 316 the polar Fourier domain. Gaussian functions had 61 different center frequencies between 1 317 cycle/object and 64 cycles/object. The center frequencies had discrete values at steps of 0.1 318 cycles/object on a log scale. Gaussian functions shared the same variance at 2.4 octaves, 319 regardless of their center frequencies. The filtered images had amplitude spectra that were 320 determined solely by the Gaussian function because their spectra were set to be flat before the 321 multiplication. To balance the total luminance contrast among the filtered face images, the peak 322 amplitude of the Gaussian function was set inversely proportional to the center image-based SF 323 (Inagaki and Fujita, 2011). These bandpass-filtered images were created for the seven facial expressions at two different sizes (99 \times 99 and 198 \times 198 pixels). 324

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328 Test determining whether units are tuned for object- or retina-based spatial Figure 2. 329 frequencies (SFs). (A) Hypothetical tuning curves for object-based SFs of units ideally tuned to 330 retina-based SFs (left: peak shift = 1) or object-based SFs (right: peak shift = 0). (B) The models 331 were fed face images with two different sizes (198×198 and 99×99 pixels) and 64 different bandpass filtering. These images were created by applying two-dimensional bandpass filters 332 333 that shared the same center object-based SF across different sizes. For each unit in FC1, we 334 obtained responses to images of different center SFs to create tuning curves for object-based 335 SFs.

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337 To characterize the reference frame of SF tuning of each unit, the peak SFs for the two stimulus sizes were estimated, defined by the SFs at which filtered face stimuli activated a unit most 338 339 strongly. For a given unit, 14 peak SFs were determined (two sizes \times seven expressions). The 340 degree to which unit responses to SFs depended on the stimulus size was quantified by 341 calculating differences in peak SFs on a log scale between the two face sizes. A peak shift of 0 342 indicates that a unit respond to the same cycles/object regardless of the image size, and is perfectly tuned to object-based SFs (Fig. 2A, right). A peak shift of 1 means that an SF tuning 343 344 curve shifts by the amount corresponding to the change in the stimulus size, indicating that a 345 unit is perfectly tuned to retina-based SFs (Fig. 2A, left). This analysis excluded cases in which 346 units did not respond to face images or were not sensitive to SFs, and cases in which peak SFs 347 were at either end of the tested range of SFs and the peak positions could not be determined.

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349 Analysis of the effects of the max pooling operation on SF selectivity.

350 The max pooling operation collapses positional information of edges, which is detected by 351 convolution filters and is critical for encoding the SFs of facial images. We therefore examined 352 the effects of the max pooling on the SF selectivity of units. We were particularly interested in 353 the role of the max pooling in converting sensitivity to retina-based SFs to sensitivity to object-354 based SFs. Bandpass-filtered images of the two stimulus sizes (198×198 and 99×99 pixels) 355 were fed to the models, and it was determined how different stimulus sizes affected the 356 responses of units to SFs in the first convolution layer (before pooling) and the first max pooling 357 layer (after pooling). Changes in response patterns across the units associated with different 358 stimulus sizes were quantified by calculating the dissimilarity index. The dissimilarity index D 359 (x, y) for responses x to the large stimuli and responses y to the small stimuli was defined by the Euclidean distance between *x* and *y* as follows: 360

- 362 D(x, y) = ||x y||/(NM)
- 363

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364 where $\|\cdot\|$ is the Euclidean distance, and N is the number of elements of x and y. M is the 365 maximum value among the 6,405 Euclidean distances calculated for 61 center SFs and the seven 366 facial expressions of 15 individuals. To probe the roles of the max pooling, the ratio of the 367 dissimilarity index before pooling in the first convolution layer and after pooling in the first max 368 pooling layer was then calculated. This analysis was applied to the case of wide pooling (5×5) and narrow pooling (3×3) , as well as to the case of the SNNs and the Gabor models. By dividing 369 370 $\|\mathbf{x} - \mathbf{y}\|$ by M, the dissimilarity index was normalized across the layers (convolution vs. max 371 pooling) and the models (SNNs vs. Gabor models), taking values from 0 to 1.

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- 374 Results
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376 *Performance of SNNs in facial expression classification.*

The SNNs were trained to classify each face image in the training set into one of the seven facial expressions. The training improved the classification performance rapidly over the initial iterations and then slowly thereafter. The correct rate across the seven facial expressions rose from the chance level (0.14), surpassed 0.6 around 50,000 iterations, further improved to around

381 0.8 over 150,000 iterations, and reached an asymptote (Fig. 3A for an example SNN; 3B for the

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average of the 20 constructed SNNs; orange lines). The correct rate for the validation set 382 383 saturated at around 0.5, which was substantially lower than for the training set, indicating 384 insufficient generalization to "unseen" images. However, the validation correct rate reached a 385 plateau in a similar way to the training correct rate. This indicates that the low correct rate was 386 not the result of inadequate training, but represents the limited learning ability of the SNNs. It 387 also indicates that no overfitting occurred. The loss value also quickly decreased over the initial 388 50,000 iterations, and became gradually stable (Fig. 3A, B; cvan lines). The results indicate that 389 within the range of adopted iterations (240,000), the SNNs were trained to classify facial 390 expressions without overfitting.

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392 The correct rates for the test set were higher than the chance level (1 / 7 = 0.14) for all facial 393 expressions, but were modest; the average correct rate across the seven expressions was 0.51394 for the example SNN shown in Fig. 3A. The average correct rate across the 20 constructed SNNs 395 was $0.51 (\pm 0.03, \text{ s.d.})$. This was not different from the average correct rate across 20 additional 396 SNNs that were trained with 3,000,000 iterations (0.50 ± 0.03 ; p = 0.289, *t*-test). The training 397 performance thus did not improve even when the SNNs underwent overly excessive training, 398 assuring that the modest correct rate was not due to insufficient training but instead reflected the 399 limited ability of the SNNs. Confusion matrices showed that the correct rates varied among the 400 facial expressions (Fig. 3A, B, right panels). Based on the averaged performance, the 401 classification performance of the SNNs was best for happy (0.74) and surprised faces (0.72), 402 followed by angry (0.50), disgusted (0.47), and fearful (0.44) faces, and was worst for sad (0.37)403 and neutral (0.34) faces. Sad faces were often confused with neutral, angry, and fearful faces. 404 Neutral faces were often confused with sad and angry faces. This expression-dependent 405 performance was consistent across the 20 constructed SNNs (Fig. 3C; p < 0.001 for expressions, 406 p = 0.562 for models, two-way ANOVA).

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410 Figure 3. Learning curves and confusion matrices of an example SNN (A) and the average of 20 SNNs with different initial weights (B). Left panels show changes in correct rates that 411 412 occurred during training in the training set (orange) and the validation set (red), and loss values 413 (cyan). Confusion matrices on the right indicate the rate of classification of each facial 414 expression (true label) as one of the seven expressions (prediction label). (C) The correct rates 415 for the seven expressions. Black line indicates the mean of the 20 SNNs, and lines with other 416 colors indicate the individual performance of the 20 SNNs. The order of facial expression is 417 based on the mean correct rate.

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419 *Effects of modification of SNNs on classification performance.*

420 Replacement of one or more of the three subcortical properties, namely the shallowness of 421 processing stages, the DoG-type receptive fields at the initial stage, and spatial pooling over a 422 wider visual field, with the corresponding cortical properties improved the classification 423 accuracy (Fig. 4A; p < 0.001, ANOVA). The correct rates averaged across the seven expressions 424 and the 20 constructed models of each modification were increased from 0.51 in the SNNs to 425 0.55 in the narrow-pooling models, 0.64 in the Gabor models, and 0.69 in the add-layer models 426 $(p < 0.01 / 28 = {}_{8}C_{2}; t-\text{test}$ with Bonferroni correction). Among these models with one replaced 427 property, the add-layer models exhibited the best performance. The results indicate that all three 428 properties had effects on classification performance, and the layer structure was the most 429 influential.

430

When two properties were replaced together, the narrow-pooling + Gabor models and the 431 432 narrow-pooling + add-layer models performed better than the narrow-pooling models and the 433 Gabor models ($p < 0.01 / 28 = {}_{8}C_{2}$; t-test with Bonferroni correction), but comparably to the add-434 layer models (correct rate, 0.69, p = 0.778 for the narrow-pooling + Gabor models; 0.69, p =435 0.564 for the narrow-pooling + add-layer models). The Gabor + add-layer models performed 436 better than all one-property-replacement models (correct rate, 0.75; $p < 0.01 / 28 = {}_{8}C_{2}$). When 437 all three properties were replaced together (full-replacement models), the correct rate was 0.77, 438 which was better than all other models ($p < 0.01 / 28 = {}_{8}C_{2}$) except for the Gabor + add-layer 439 models ($p = 0.00846 > 0.01 / 28 = {}_{8}C_{2}$). The performance was improved for all facial expressions 440 (Fig. 4B; mean correct rates: happy = 0.92; surprised = 0.85; disgusted = 0.81; neutral = 0.78; 441 angry = 0.75; fearful = 0.66; sad = 0.65). As in the SNNs, it was highest for happy and surprised 442 faces, and lowest for fearful and sad faces. The performance was improved most for neutral 443 faces (SNNs, 0.37; full-replacement models, 0.78). The variance of the correct rates was affected 444 both by facial expressions and models (Fig. 4C; p < 0.001 for facial expressions, p = 0.00285445 for models, two-way ANOVA). The improved performance of the two-property-replacement 446 and full-replacement models indicate that the effects of the three features on classification 447 performance were partially additive, suggesting that the three features exerted their effects 448 partially independently.

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452 Figure 4. Effects on model performance of replacing subcortical properties with corresponding 453 cortical properties. (A) Discrimination performances of the SNNs and the modified models. The 454 discrimination performance differed across the models (ANOVA, p < 0.001). The pairs of 455 models with statistically significant differences in the performances are linked with horizontal 456 lines in the upper part (*t*-test, Bonferroni correction, $p < 0.01/28 = {}_{8}C_{2}$). (B) Confusion matrix 457 for the full-replacement models (average of the 20 constructed models). (C) The correct rate for 458 the seven expressions across the 20 full-replacement models. The black line indicates the mean, 459 and lines with other colors indicate the data for individual full-replacement models. 460

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461 Spatial frequency representation in the FC1 layer.

462 As shown above, the SNNs exhibited modest performance in facial expression classification, 463 and this performance was improved by changing SNN subcortical properties to corresponding 464 cortical properties. These findings suggest that the SNNs captured aspects of processing in the 465 subcortical pathway to the extent that they explained the suboptimal perceptual performance of V1-lesioned patients. We next looked into individual computational units to gain insights about 466 the processing in the models. We examined SF sensitivities of FC1 units using two different 467 sizes of input images (198×198 and 99×99 pixels). This procedure allowed us to determine 468 469 whether units were sensitive to retina- or object-based SFs (Fig. 2; see Materials and Methods).

470

471 FC1 units of the SNNs exhibited a variety of dependencies of SF tunings on stimulus size (Fig. 472 5A). Some units responded to the same range of object-based SFs for both large and small 473 stimuli, and the peak positions of the SF tuning curves remained unchanged (Fig. 5Aa, Ab). 474 Other units exhibited different preferred SFs for large and small stimuli, and in these cases the 475 peak position shifted horizontally along the abscissa (Fig. 5Ac-f). We quantified these shifts by 476 measuring the difference between preferred SFs on a log scale for the two stimulus sizes. A peak 477 shift of 0 means that the unit encoded SFs in the object-based coordinate, whereas a peak shift 478 of 1 means that the unit encoded SFs in the retina-based coordinate. The peak shifts of the 479 example units shown in Fig. 5A were 0.0 (a), 0.1 (b), 0.1 (c), 0.9 (d), 1.0 (e), and 1.3 (f).

480

We plotted the peak positions of 2,401 FC1 units of the 20 SNNs in a two-dimensional space 481 482 defined by the peak SF for the large stimuli on the abscissa, and the peak SF for the small stimuli 483 on the ordinate (Fig. 5B, left). Note that 46% of FC units were excluded from this analysis, 484 either because they were not sensitive to SFs (23%) or because the largest responses were found 485 at the end of the examined range of SFs and the peak SFs could not be determined (23%). The 486 diagonal solid line in Fig. 5B represents the responses of a peak shift of 0, and the dashed line 487 next to it represents the responses of a peak shift of 1. FC1 units of the SNNs were clustered in 488 multiple groups in this scatter plot. One conspicuous group was selective to low SFs and was 489 centered on the diagonal, i.e., peak shift values around 0. Another group was selective to higher 490 SFs, and was clustered on the dashed line indicative of peak shift values around 1. The 491 multimodality of the distribution can also be seen in the histogram (Fig. 5C, left). We applied 492 an excess mass test for multimodality (Ameijeiras-Alonso et al., 2019, 2021) to this distribution. 493 This test statistically determines the number of peaks in the distribution, with the null hypothesis 494 that the true number of peaks is N(N = 1, 2, 3, ...). The true number of peaks is estimated as the 495 smallest N under which the null hypothesis is not rejected. The excess mass test also estimates 496 the locations and heights of peaks from Gaussian kernel density estimation. The test revealed 497 that there were three peaks in the distribution of the SNNs (first *p*-value < 0.001, second *p*-value 498 < 0.001, third *p*-value = 0.096). Based on the probability density function derived from the 499 histogram (Ameijeiras-Alonso et al., 2021), the peaks were estimated to be located at -4.85, 500 0.144, and 0.909 (open and solid arrowheads in Fig. 5C, left). Units sensitive to low SFs below 501 2 cycles/object were most frequent around a peak shift of 1 (gray columns). Comparing this 502 result and the density map, the peak around 0 was mostly from the low spatial frequency group 503 and the peak around 1 was from the high spatial frequency group. Although the third peak at the 504 far periphery (at -4.85, open arrowhead) was statistically detected, it was much smaller in height 505 than the other two peaks (1.1%) of the peaks near 0 and 1). The results indicate that FC1 506 contained two major groups of units, those sensitive to low SFs, encoding SFs in the object-507 based coordinate, and those sensitive to high SFs, encoding SFs in the retina-based coordinate.

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511 Figure 5. SF tuning reference frames of FC1 units of the SNNs and full-replacement models. 512 Responses of FC1 units to SF-filtered face images were examined at two different sizes (198 \times 513 198, 99 \times 99 pixels). (A) Six example FC1 units of SNNs with a different peak shift (PS). (B) 514 Two-dimensional histograms of peak SFs at large images versus small images for the SNNs 515 (left) and the full-replacement models (right). Solid lines indicate peak shifts of 0, and dashed 516 lines indicate peak shifts of 1. (C) Distribution of peak shifts of units in the 20 SNNs (left) and the 20 full-replacement models (right). Arrowheads indicate the estimated locations of multiple 517 518 peaks in the distribution (solid: major peaks, open: statistically detected but less obvious peaks). 519 Gray columns indicate units with a response peak at SFs below 2 cycles/object for large and/or 520 small stimuli.

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521 The distribution of peak shift values was drastically altered in the full-replacement models. In 522 the two-dimensional plot shown in Fig. 5B (right), most data points were diffusely distributed 523 in an elongated area between the diagonal and dashed lines, indicating that the SF reference 524 frame of most units was intermediate between retina-based and object-based. An excess mass 525 test again detected three peaks located at -4.84, 0.436, and 4.98 (Fig. 5C right, solid and open arrowheads; first p-value < 0.001, second p-value < 0.001, third p-value = 0.096). The second 526 527 and third peaks at -4.84 and 4.98 (open arrowheads) were smaller than the primary peak at 528 0.436 (3.1% and 2.6% of the primary peak, respectively), making the distribution nearly 529 unimodal.

530

531 Given the change of the peak shift distribution in the full-replacement models, we next analyzed 532 one- and two-property replacement models to determine which subcortical properties were 533 essential for the multimodal distribution of the SNNs. All these modified models exhibited 534 unimodal distributions of the major peak (solid arrowheads) at different peak positions (Fig. 6). 535 An excess mass test for multimodality detected two other less obvious peaks (open arrowheads) 536 in each model as in the cases of the SNNs and the full-replacement models (Fig. 5C). The heights 537 of these smaller peaks were 2.6–26 % of those of the major peaks, and were located at the 538 periphery of the distribution.

539

540 Each of the three one-property-replacement models showed a characteristic distribution of the 541 peak shift values. The narrow pooling models contained units with peak shift values between 0 542 and 1 in addition to units with peak shift values around either 0 or 1. The distribution became 543 unimodal and broad, and was estimated to be centered at 0.85. In the Gabor models, units with peak shift values intermediate between 0 and 1 were the most abundant with a smaller number 544 545 of units of peak shift values around 0 and 1. The distribution peak was estimated at 0.51. In the 546 add-layer models, units with peak shift values around 0 were predominant, and exhibited a sharp 547 distribution peak at 0.32. As to the two-property-replacement models, the narrow-pooling + 548 Gabor models and the Gabor + add-layer models showed a broad distribution straddling the 549 peak values from 0 to 1 (peak for the former, 0.56; peak for the latter, 0.52), whereas the narrow-550 pooling + add-layer models showed a sharp distribution peak at 0.20. As in the SNNs and the 551 full-replacement models, units sensitive to low SFs (below 2 cycles/object) were most 552 frequently found around the peak shift of 0 in all of the one- and two-property replacement 553 models (gray columns). The results indicate that all of the three computational properties were 554 responsible for the multimodal distribution of peak shift values observed in the SNNs. In 555 particular, the smaller number of units with peak shift values around 1 in the Gabor models and 556 the add-layer models suggests that the shallowness and the DoG-type filters were critical for 557 preserving the unit sensitivities to retina-based SFs. The broad distribution observed for the 558 narrow-pooling models and the narrow-pooling + Gabor models suggests that the wide pooling employed in the SNNs contributed to the two peaks at 0 and 1, by reducing units with peak shift 559 560 values intermediate between 0 and 1.

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564 Figure 6. Distributions of peak shifts of FC1 units of the SNNs and one-property or two-565 property-replacement models. (A) Data from narrow-pooling model, add-layer model, and 566 Gabor model. (B) Data from models with two modifications: narrow-pooling + Gabor, Gabor + 567 add-layer, narrow-pooling + add-layer. Gray columns indicate units with a response peak at SFs below two cycles/object for large or small stimuli. Arrowheads indicate the estimated locations 568 569 of multiple peaks in the distribution (solid: major peaks, open: statistically detected but small 570 peaks). Gray columns indicate units with a response peak at SFs below 2 cycles/object for large 571 and/or small stimuli.

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573 *Effects of max pooling on SF tuning.*

We showed above that FC1 units of the SNNs were roughly grouped into two populations in terms of the reference frame of SF encoding. Because the max pooling yields the same output from a population of convolution layer units in response to slightly different spatial arrangement of local features, the max pooling operation is likely to affect the encoding of global configuration of face components. This information of global configuration will be reflected in a low range of SFs. Therefore, we next compared the effect of max pooling on the representation of SFs across different SF ranges.

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582 We first analyzed the responses of the 96,800 units (32 filters \times 55 \times 55 resolution) in the first 583 convolution layer. We obtained the response patterns across these units by feeding bandpass-584 filtered faces of two sizes (198×198 and 99×99 pixels; Fig. 2B) to the models, and quantified 585 the difference between the SF tunings obtained for the two stimulus sizes by calculating the 586 dissimilarity index (see Materials and Methods). In the SNNs with DoG filters, the dissimilarity index was high (around 0.6) for a low SF range up to approximately four cycles/object, but 587 gradually decreased over a higher range of SFs (Fig. 7A, black curve). In the pooling layer, the 588 589 dissimilarity index of the 6,272 units (32 filters \times 14 \times 14 resolution) became lower for a low 590 SF range of less than four cycles/object than that of the convolution layer. For a high SF range 591 of greater than four cycles/object, by contrast, it became higher than that of the convolution 592 layer (Fig. 7A, compare the orange and cyan curves with the black curve). Thus, max pooling 593 resulted in the SF tuning becoming similar between the two stimulus sizes for a low SF range, 594 consistent with the results of peak shift analysis (Fig. 5C). Although these changes were 595 observed both for wide pooling (5×5) and narrow pooling (3×3) , the effects were larger for 596 the former than for the latter (Fig. 7A, compare the orange curve with the cyan curve). This was 597 more evident when we plotted the ratio of dissimilarity indices between before and after pooling 598 (Fig. 7B). Furthermore, the ratio curve for wide pooling had smaller standard deviations than 599 that for narrow pooling (shown as shades in Fig. 7B), indicating that wide pooling exerted its 600 effects more consistently across the 15 individual faces and the seven facial expressions than 601 narrow pooling.

602

603 By contrast, the convolution-layer units of the Gabor models exhibited a constantly high 604 dissimilarity index over most of the SF range (Fig. 7C, black). However, when we applied max 605 pooling with windows of either 5×5 or 3×3 in size, the dissimilarity index became small over 606 almost the entire SF range, with the largest decrease for 1–16 cycles/object (Fig. 7C; compare 607 the orange and blue curves with the black curve). As in the case of the SNNs, the effect was 608 stronger for wide pooling than for narrow pooling (Fig. 7D). These results demonstrated that 609 max pooling rendered the SF tuning more invariant to stimulus size for units sensitive to low SFs, enabling them to represent SFs in the object-based coordinate. Regardless of the filter type 610 611 in the first convolution layer (i.e., DoG vs. Gabor), wide pooling was more effective than narrow 612 pooling in creating this response property.





616 Figure 7. Effects of max pooling on the size-invariant responses to SFs. (A) Dissimilarity index 617 curves of responses of units in the first convolution layer of the SNNs before max pooling 618 operation (black), after wide pooling (blue), and after narrow pooling (orange). Dissimilarity 619 indices, defined by the Euclidean distances of unit responses between different stimulus sizes 620 (see Method and Methods), are plotted against the center SFs of input images. Solid lines 621 indicate the means of dissimilarity indices across the seven facial expressions. Shades indicate standard deviations. Each dissimilarity index was normalized by the number of units and the 622 623 maximum values. (B) Dissimilarity ratios of inputs and outputs of the max pooling operation 624 (upper, after wide pooling; lower, after narrow pooling). (C, D) Data from units in the first 625 convolution layer of the Gabor models. The conventions are the same as in A and B. 626

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627 Effects of alternation of sliding strides on SF tuning.

Finally, we examined the effect of another free parameter of our models, the stride size, on the SF sensitivity of FC1 units. We changed the stride of the two max pooling layers of the SNNs from 4 to 2. The stride size of 2 was also employed in the narrow-pooling model. This modified model with a smaller stride of 2 achieved a mean correct rate of 0.54, which was better than the SNNs (0.51) but similar to the narrow pooling models (0.55) (vs. SNN, p = 0.0020; vs. the narrow pooling, p = 0.23; *t*-test with Bonferroni correction).

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635 An analysis of FC1 responses to two stimulus sizes $(198 \times 198 \text{ and } 99 \times 99 \text{ pixels})$ revealed that 636 the distribution of the peak shift had three peaks, at -4.17, -0.0107, and 0.976 (first *p*-value < 637 0.001, second *p*-value < 0.001, third *p*-value = 0.098; excess mass test for multimodality; solid 638 and open arrowheads in Fig. 8). Two of them were conspicuous and located near 0 or 1 (solid 639 arrowheads), and the third one at the periphery of -4.17 (open arrowhead) was small (3.2% and 640 3.3% of the two major peaks). Comparisons with Fig. 5B, C show that the small-stride model 641 exhibited a similar SF representation as in the SNNs, in that there were two main groups of units, 642 one sensitive to low SFs, representing SFs in the object-based coordinate (peak shift around 0), 643 and the other sensitive to high SFs, representing SFs in the retina-based coordinate (peak shift 644 around 1). The change of the stride from 4 to 2 had little effects on the reference frame of SF 645 sensitivity of FC1 units. 646

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Figure 8. Effects of alternation of sliding strides of pooling windows on SF tuning reference frames of FC1 units of the SNNs. Responses of FC1 units were obtained in the same way as in Fig. 5. (A) A two-dimensional histogram of peak SFs obtained with large (198×198) versus small (99×99) stimulus images. (B) Distribution of peak shifts of units. Arrowheads indicate the estimated locations of multiple peaks in the distribution.

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658 Discussion

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660 We analyzed the ability of the SNNs and modified models to classify facial expressions, with the goal of determining what architectural or physiological properties underlie the modest 661 662 performance of facial expression discrimination supported by the primate subcortical pathway. 663 The SNNs were implemented with the three prominent subcortical properties, i.e., shallow processing, DoG-type filters at the first processing stage, and spatial pooling over wide areas 664 665 (Fig. 1). The SNNs classified the seven basic facial expressions with modest performance (Fig. 666 3). Replacement of any one of these properties with the corresponding cortical properties 667 resulted in higher performances (Fig. 4). Replacement of a combination of two or three 668 properties further improved classification performances in a partially additive manner (Fig. 4). 669 These results suggest that all three subcortical properties of the SNNs underlie the modest 670 performance. A major group of units in the final processing layer (FC1) of the SNNs was 671 sensitive to SFs defined in the retina-based coordinate, whereas another group responding to low SFs encoded SFs in the object-based coordinate (Fig. 5). The number of retina-based units 672 673 was reduced in most of the modified models, suggesting that the three features are also important 674 for preserving retina-based SF information (Figs. 5, 6). Max pooling over the wide window 675 employed in the SNNs contributed to object-based SF representation of units sensitive to low 676 SFs (Fig. 7). These findings advance our understanding of the computational processes utilized 677 by the subcortical pathway in facial expression recognition.

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Modest performance of the SNNs and neural computations of the subcortical pathway.

Based on psychological assessment and brain imaging in V1-lesioned patients, it has been 681 682 proposed that affective blindsight is mediated by components of the subcortical pathway spared 683 by the lesions, including the superior colliculus, pulvinar, and amygdala (de Gelder et al., 1999; 684 Pegna et al., 2005; Striemer et al., 2019). One view assumes that the shortest route directly 685 connecting the three subcortical structures conveys facial expression information from the 686 superior colliculus via the pulvinar to the amygdala (Tamietto and de Gelder, 2010). A different 687 view proposes that information from the pulvinar reaches the amygdala through the facial 688 processing system in the temporal cortex, under an assumption that "the direct connections of 689 the pulvinar with the amygdala are likely insufficient in themselves for recognizing emotional 690 expressions" (Gerbella et al., 2019). The present study demonstrates that the SNNs with only 691 three processing stages and the subcortical physiological properties can successfully acquire an 692 ability to discriminate facial expressions.

693

694 The average correct rate of classifying the seven facial expressions in the present study was 0.51. 695 This rate was well above chance (1/7 = 0.14), but was far from perfect. The modest correct rate 696 is in line with the performance of patients with affective blindsight. Pegna et al. (2005) reported 697 that a patient with bilateral lesions in V1 discriminated happy faces from either angry, sad, or 698 horrified faces at correct rates of 0.58-0.62, marginally above the chance level of 0.5. Another 699 patient with bilateral lesions in V1 exhibited correct rates of 0.64–0.67 for happy vs. fearful or 700 angry faces (chance level = 0.5; Striemer et al. 2019). The residual ability of facial expression 701 classification in these patients was only moderate compared to the nearly perfect performance

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in healthy people. This raises the question of why subcortical processing supports vision morepoorly than visual functions mediated by the cortical pathway.

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705 A traditional explanation is that neurons in the subcortical pathway respond to low SFs and are 706 less sensitive to high SFs than the cortical pathway (e.g., Vuilleumier et al., 2003; Méndez-Bértolo et al., 2016; Burra et al., 2019). This will limit the ability of the subcortical pathway to 707 708 analyze the fine details of visual images, and can itself result in the inaccurate processing of face 709 images. However, the dependence of the subcortical response on low SFs has been disputed by 710 other researchers (De Cesarei and Codispoti, 2013; McFadyen et al., 2017). Our results suggest 711 that low SF sensitivity, if important, was not the only cause, because the DoG filter models 712 combined with narrow-pooling or add-layer modifications exhibited improved performances, 713 despite the fact that our DoG filters were tuned to low SFs, and had full width at half maximum 714 of 0.067–1.0 cycles/degree. Note that we estimated this value on an assumption of the image 715 size of 30.5° based on our DoG parameters, the filter resolution, and the RF size of superior 716 colliculus neurons representing the foveal region. The range of DoG-filter width corresponds to 717 that applied in models of the superior colliculus in a recent simulation study (Méndez et al., 718 2022).

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720 Another explanation is that the small number of processing stages in the cortical pathway 721 hampers detailed analysis of visual inputs. However, a previous study (Dailey et al., 2002) 722 showed that CNNs that had only two processing layers, with Gabor filters at the first stage, 723 performed highly accurate discrimination of facial expressions (the mean correct rate for 724 classifying six facial expressions was 0.90). The performance of our SNNs incorporating the 725 three subcortical properties was not this high. This was not due to inadequate training, because 726 the performance reached a plateau and stayed stable over a large number of iterations in the 727 training sessions (Fig. 3). This was further verified by showing that the correct performance for 728 the test set remained unchanged even after overly excessive training with 3,000,000 iterations. 729 Furthermore, replacing not only the small number of processing layers but also the filter type at the first processing layer and the width of the pooling window with the corresponding cortical 730 731 properties improved the performance of the SNNs (Fig. 4). The three properties at least partially 732 underlie the less accurate processing of facial images in the subcortical pathway, and hence, 733 may be responsible for the low performance in affective blindsight.

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Confusions of facial expressions in the SNNs, DNNs and patients.

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737 The classification accuracy of the SNNs varied across facial expressions (Fig. 3A, B). The 738 classification performance of the SNNs was best for happy and surprised faces and worst for 739 sad and neutral faces. The rank order of performance on the seven facial expressions was largely 740 consistent across the 20 SNNs trained independently from random states (Fig. 3C). It also 741 corresponds to the classification performance by previously developed AlexNet-based DNNs 742 (Inagaki et al., 2022b). These DNNs were trained to discriminate between the seven expressions 743 derived either from the KDEF database or the Kokoro Research Center (KRC) facial expression 744 database (Ueda et al., 2019). Like the SNNs, the DNNs exhibited the best performance for happy 745 and surprised faces (KDEF: 0.93 for happy, 0.84 for surprised; KRC: 0.91 for happy, 0.84 for 746 surprised; chance level, 0.14). This coincidence may simply suggest that within each database,

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747 facial features are consistent across faces with happy or surprised expressions, but are more diverse across faces with sad or neutral expressions. However, the variations across examples 748 749 of facial expressions within a database are not the sole reason for the difference in the 750 performance across facial expressions, because neutral faces were classified poorly by the SNNs 751 (Fig. 3B; correct rate = 0.34), but the DNNs of Inagaki et al. (2022b) classified them with high 752 correct rates (0.85 for KDEF, 0.82 for KRC). An alternative, yet-to-be-tested explanation is that the ease (or difficulty) of classification may vary across the facial expressions owing to 753 754 differences in the conspicuousness of component facial actions underlying various expressions. 755 Similarities between neural networks regarding expression-specific performance may vary 756 according to these differences.

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758 The relatively poor ability to distinguish between sad and neutral faces was also observed in 759 another CNN with the first layer of DoG filtering and average pooling (Méndez et al., 2022). 760 This CNN was constructed to simulate facial processing in the superior colliculus, and was 761 trained to discriminate three facial expressions: happy, sad, and neutral. The CNN showed the 762 best performance for happy faces and moderate performance for sad faces, but classified neutral 763 faces into neutral faces with a classification rate of 0.49 and into sad faces with a rate of 0.39. 764 The fact that this CNN and the SNNs in the present study demonstrated this confusion, whereas 765 AlexNet-based DNNs and our add-layer models (0.52 for sad, 0.67 for neutral) did not, suggests 766 that the convolution processes after the initial DoG filtering (in the case of add-layer models) or 767 the convolution by the Gabor filters (in the case of AlexNet-based DNNs) may be critical for 768 classification of sad and neutral faces.

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Finally, we point out that the expression-dependent performance of the SNNs also had both similarities and dissimilarities to that observed in a V1-lesioned patient. The patient reported by de Gelder et al. (1999) classified happy and sad faces with a higher correct rate than angry and fearful faces; our SNNs and this patient classified happy faces well, whereas the performance for sad faces was poor in the SNNs but good in the patient.

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Reference frame of coding SF information and invariance of visual responses.

FC1 units of the SNNs consisted of two major groups, each with different properties regarding 778 779 SF processing (Fig. 5). One group of units responded best to the same object-based SFs (cycles/object) regardless of the stimulus size (peak shift around 0). This size-invariant response 780 781 indicates that these units represent SFs in the object-based coordinate. Most of these units were 782 tuned to a low SF range (around one to two cycles/object). In the other group of units, the 783 optimal object-based SFs shifted when testing was performed with different stimulus sizes. The 784 direction of the shift was consistent with the interpretation that the units were tuned to retinabased SFs (cycles/degree) (peak shift around 1). That is, for larger stimuli, the units responded 785 786 to higher object-based SFs that corresponded to the same retina-based SFs. The DoG filters at 787 the initial stage and the shallow architecture appear to be critical for preserving the SF 788 representation based on the retina-based coordinate, because FC1 units with retina-based SF 789 sensitivity were reduced in number when the first convolution layer were changed to Gabor 790 filters or when the number of processing layers was increased (Fig. 6A, middle, bottom).

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792 A major group of the object-based SF units in the SNNs were tuned to low SFs (Fig. 5B). This 793 curious bias of the object-based units towards low SF sensitivity likely resulted from the wide 794 max pooling process. Lowpass-filtered facial images contain only coarse structure such as solid 795 blobs at eye or mouth positions. Positional information of these blobs is initially detected by 796 DoG filters, and is encoded as response patterns across units in the convolution layer. These 797 blobs appear in different positions and scales for images of different sizes, and thus the response 798 patterns vary between different sizes. After the max pooling operation, however, response 799 patterns would become more similar between different sizes, because this operation renders 800 units in the pooling layer insensitive to slight changes in spatial arrangement of local features. 801 Indeed, the dissimilarity index for lowpass-filtered images decreased after max pooling in our 802 data (Fig. 7). This effect might result in object-based SF tuning (i.e., preferential responses 803 invariant of image size to a particular range of object-based SFs) for lowpass-filtered images. 804 Wider pooling window would enhance this effect at the expense of losing fine details of inputs. 805 When the pooling window is narrower, this effect would be incomplete, and units with 806 intermediate peak shift values would increase, as we found in the narrow-pooling models (Fig. 807 6A, top). 808

809 One may wonder why FC1 units of the SNNs maintained sensitivity to retina-based SFs, i.e., 810 size-dependent representation of SFs, despite the demand that we imposed on the SNNs to 811 classify facial expressions regardless of the seven different face image sizes. One possibility is 812 that the architecture of our SNNs cannot achieve sufficient object-based representation, and 813 remains suboptimal for the required task even after the excessive training sessions. This may be 814 a reason for the modest classification performance of the SNNs. Indeed, replacement of the 815 subcortical processing properties with the cortical properties resulted in the representation 816 becoming more object-based (Fig. 5B, C, Fig. 6) and improved the classification performance 817 (Fig. 4). However, if object-based SF encoding was the only requirement for optimal 818 performance under our training conditions, the models that showed object-based SF encoding 819 should have had the highest correct rate, but this was not the case. The add-layer models and the 820 narrow-pooling + add-layer models exhibited the best object-based encoding of SFs (Fig. 6A 821 bottom, 6B bottom), while they performed worse than the full-replacement model (Fig. 4A). 822 The representation acquired for the classification depended not only on the task demand of size-823 invariant classification of facial expressions, but also on other, yet unspecified, constraints 824 deriving probably from the architecture of the models.

825

In the primate amygdala, the responses of many neurons are affected by retina-based SFs, and only a minority of neurons have perfect object-based SF sensitivity (Inagaki and Fujita, 2011). By contrast, many FC1 units tuned to low SFs of the SNNs exhibited object-based SF sensitivity. The paucity of evidence for units with object-based SF sensitivity in the amygdala may be related to the fact that the previous electrophysiological study (Inagaki and Fujita, 2011) did not present face images with very low SFs, and may have overlooked the neurons with object-based SF sensitivity in this range of SFs.

833

Some inferior temporal cortex neurons exhibit invariant responses to changes in shape sizes (Rolls and Baylis, 1986; Ito et al., 1995). The max pooling operation may help achieve these invariant responses. To some degree, max pooling ignores positional changes of inputs in each region of interest. Because size changes involve alternations in edge positions without

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modifications in topologies, if the changes are small enough to be covered by each region of interest, stimuli before and after the changes would yield similar responses. The effects of wide pooling shown in Figure 7 suggest that some aspects of the invariant responses of inferior temporal cortex neurons can simply be achieved by bypassing early cortical areas with high spatial resolutions such as V1. Such shortcut routes indeed exist, including the projection from the pulvinar to V2 and then to the posterior inferior temporal cortex and the projection from the pulvinar to V4 and then to the anterior inferior temporal cortex (Pessoa and Adolphs, 2010).

845

846 The size invariance in low SFs is important in newborns. They have blurred visions that relies 847 on low SFs (Atkinson et al., 1974; Dobson & Teller, 1978), but respond to faces or face-like 848 patterns irrespective of the stimulus size or the viewing distance (Cassia et al., 2001; De Heering 849 et al., 2008). These findings indicate that the ability of size-invariant face recognition based on 850 low SFs is innately implemented in our visual system. Convergence of inputs from the 851 superficial layer of the superior colliculus to the deep layer, which is already present in newborns 852 (Wallace et al., 1997), may be part of the neural substrate supporting this aspect of size-invariant 853 face recognition.

854

855 Concluding remarks

856

857 The present study provides the first computational model for facial expression processing along 858 the subcortical pathway (see Méndez et al., 2022 for a model of face processing in the superior 859 colliculus). Despite the celebrated success of DNNs in modeling visual processing in the ventral 860 cortical pathway, it has remained unclear whether and how the CNN architecture can be adapted 861 to processing in the subcortical pathway. We demonstrated that the SNNs implemented with the 862 three computational properties of the subcortical pathway, i.e., a shallow layer architecture, 863 concentric receptive fields at the first processing stage, and a greater degree of spatial pooling, 864 were successfully trained to discriminate facial expressions with a modest correct rate. The three 865 properties were all essential for reproducing the modest performance seen in V1-lesioned 866 patients, as well as the representation of SFs in the retina-based coordinate observed in a 867 population of amygdala neurons. Research interest in the role of subcortical structures in 868 cognitive functions has recently surged, but physiological data are still much sparser for 869 subcortical structures than for the cerebral cortex (Janacsek et al., 2022). Computational 870 approaches such as the one we present here are expected to partially compensate for this data 871 scarcity and to guide future research.

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1099 Author contributions

- 1100 CL, MI, TS, and IF designed the research; CL, MI, and TS performed the research; CL, MI, TS,
- and IF wrote the paper. All authors approved the submitted version.

1102 Competing interests

1103 The authors declare no conflicts of interest.

1104 Data availability

1105 All data and analysis codes are available from the corresponding author upon request.

1106 **Ethics statement**

1107 Written informed consent was obtained for the publication of any identifiable images included 1108 in this article.