Mechanisms of Face Perception

Doris Y. Tsao¹ and Margaret S. Livingstone²

¹Centers for Advanced Imaging and Cognitive Sciences, Bremen University, D-28334 Bremen, Germany; email: doris@nmr.mgh.harvard.edu
²Department of Neurobiology, Harvard Medical School, Boston, Massachusetts 02115; email: mlivingstone@hms.harvard.edu

Key Words
face processing, face cells, holistic processing, face recognition, face detection, temporal lobe

Abstract
Faces are among the most informative stimuli we ever perceive: Even a split-second glimpse of a person’s face tells us his identity, sex, mood, age, race, and direction of attention. The specialness of face processing is acknowledged in the artificial vision community, where contests for face-recognition algorithms abound. Neurological evidence strongly implicates a dedicated machinery for face processing in the human brain to explain the double dissociability of face- and object-recognition deficits. Furthermore, recent evidence shows that macaques too have specialized neural machinery for processing faces. Here we propose a unifying hypothesis, deduced from computational, neurological, fMRI, and single-unit experiments: that what makes face processing special is that it is gated by an obligatory detection process. We clarify this idea in concrete algorithmic terms and show how it can explain a variety of phenomena associated with face processing.
INTRODUCTION

The central challenge of visual recognition is the same for both faces and objects: We must distinguish among often similar visual forms despite substantial changes in the image arising from changes in position, illumination, occlusion, etc. Although face identification is often singled out as demanding particular sensitivity to differences between objects sharing a common basic configuration, in fact such differences must be represented in the brain for both faces and nonface objects. Most humans can easily identify hundreds of faces (Diamond & Carey 1986), but even if one cannot recognize a hundred different bottles by name, one can certainly distinguish them in pairwise discrimination tasks. Furthermore, most of us can recognize tens of thousands of words at a glance, not letter by letter, a feat requiring expert detection of configural patterns of nonface stimuli. Thus, face perception is in many ways a microcosm of object recognition, and the solution to the particular problem of understanding face recognition will undoubtedly yield insights into the general problem of object recognition.

The system of face-selective regions in the human and macaque brain can be defined precisely using fMRI, so we can now approach this system hierarchically and physiologically to ask mechanistic questions about face processing at a level of detail previously unimaginable. Here we review what is known about face processing at each of Marr's levels: computational theory, algorithm, and neural implementation.

Computer vision algorithms for face perception divide the process into three distinct steps. First, the presence of a face in a scene must be detected. Then the face must be measured to identify its distinguishing characteristics. Finally, these measurements must be used to categorize the face in terms of identity, gender, age, race, and expression.

Detection

The most basic aspect of face perception is simply detecting the presence of a face, which requires the extraction of features that it has in common with other faces. The effectiveness and ubiquity of the simple T-shaped schematic face (eye, eye, nose, mouth) suggest that face detection may be accomplished by a simple template-like process. Face detection and identification have opposing demands: The identification of individuals requires a fine-grained analysis to extract the ways in which each face differs from the others despite the fact that all
faces share the same basic T-shaped configuration, whereas detection requires extracting what is common to all faces. A good detector should be poor at individual recognition and vice versa.

Another reason why detection and identification should be separate processes is that detection can act as a domain-specific filter, ensuring that precious resources for face recognition [e.g., privileged access to eye movement centers (Johnson et al. 1991)] are used only if the stimulus passes the threshold of being a face. Such domain-specific gating may be one reason for the anatomical segregation of face processing in primates (it is easier to gate cells that are grouped together). A further important benefit of preceding identification by detection is that detection automatically accomplishes face segmentation; i.e., it isolates the face from background clutter and can aid in aligning the face to a standard template. Many face-recognition algorithms require prior segmentation and alignment and will fail with nonuniform backgrounds or varying face sizes.

Measurement and Categorization

After a face has been detected, it must be measured in a way that allows for accurate, efficient identification. The measurement process must not be so coarse as to miss the subtle features that distinguish one face from another. On the other hand, it must output a set of values that can be efficiently compared with stored templates for identification. There is a zero-sum game between measurement and categorization: The more efficient the measurement, the easier the classification; conversely, less efficient measurement (e.g., a brute force tabulation of pixel gray values) makes the classification process more laborious.

COMPUTER VISION ALGORITHMS

A comprehensive review of computer algorithms for face recognition can be found in Zhao et al. (2003) and Shakhnarovich & Moghaddam (2004). Our goal here is to discuss algorithms that offer special insights into possible biological mechanisms.

Detection

How can a system determine if there is a face in an image, regardless of whose it is? An obvious approach is to perform template matching (e.g., search for a region containing two eyes, a mouth, and a nose, all inside an oval). In many artificial face-detection systems a template is swept across the image at multiple scales, and any part of the image that matches the template is scored as a face. This approach works, but it is slow.

To overcome this limitation, Viola & Jones (2004) introduced the use of a cascade of increasingly complex filters or feature detectors. Their reasoning was that the presence of a face can be ruled out most of the time with a very simple filter, thus avoiding the computational effort of doing fine-scale filtering on uninformative parts of the image. The first stage in their cascade consists of only two simple filters, each composed of a few rectangular light or dark regions (Figure 1a). Subsequent stages of filtering are performed only on regions scoring positive at any preceding stage. This cascade approach proved just as accurate, but 10 times faster, than single-step face-detector algorithms.

Sinha’s face-detection algorithm (Sinha 2002a) is based on the observation that qualitative contrast relationships between different parts of a face are highly conserved, even under different lighting conditions (Figure 1b). Even though any single contrast relationship between two facial regions would be inadequate to detect a face, a set of such relationships could be adequate (because probabilities multiply). A subset of Sinha’s directed contrasts ([r2, r3] and [r4, r5]) are equivalent to the first stage of the Viola-Jones face detector.

Effective primitives for face detection can also be computed using an information theory approach by identifying fragments (subwindows) of face images that are maximally
informative about the presence or absence of a face (Ullman et al. 2002). The resulting fragments consist of medium-resolution face parts, e.g., an eye, rather than the whole face, so in this algorithm, face detection is triggered by detection of a threshold number of such fragments.

All three algorithms discussed above use basic feature detectors much simpler than a whole face (rectangle features in the Viola-Jones algorithm, qualitative contrast ratios between pairs of face regions in the Sinha algorithm, and face parts in the Ullman algorithm). Yet, all three algorithms perform holistic detection, that is, they obligatorily detect faces as correctly arranged wholes. This is because all three algorithms detect overlapping constellations of elemental features that cover the whole face. The feature overlaps implicitly enforce the correct overall arrangement of features.

Measurement

Once a face has been detected, it may need to be identified or classified. Algorithms for the identification of individual faces are generally either feature-based or holistic. In feature-based methods, fiducial points (e.g., eyes, mouth, nose) are identified and used to compute various geometric ratios. As long as the features can be detected, this approach is robust to position and scale variations. In holistic methods, the entire face is matched to memory templates without isolating specific features or parts. One advantage of holistic methods is that all parts of the face are used, and no information is discarded.

The simplest holistic recognition algorithm is to correlate a presented image directly to a bank of stored templates, but having templates for every face is expensive in time and memory space. Turk & Pentland (1991) developed the eigenface algorithm to overcome these limitations. The eigenface algorithm exploits the fact that all faces share a common basic structure (round, smooth, symmetric, two eyes, a nose, and a mouth). Thus the pixel arrays defining various faces are highly correlated, and the distinguishing characteristics of a face can be expressed more efficiently if these correlations are removed using principal components analysis (PCA). When PCA is performed on a large set of faces, the eigenvectors with largest eigenvalues all look like faces, and hence are called “eigenfaces” (Figure 2a). An arbitrary face can be projected onto a set of eigenfaces to yield a highly compressed representation; good face reconstructions can typically be obtained with just 50 eigenfaces and passable ones with just 25. In other words, something as ineffable as an identity can be reduced to 25 numbers (Figure 2b).

PCA on sets of faces varying in both expression and identity generates some principal components that are useful for only expression or only identity discrimination and others that are useful for both (Calder et al. 2001). This partial independence of PCs can successfully model the independent perception of expression and identity (Cottrell et al. 2002).

The eigenface algorithm does not perform well if the sample face is not accurately aligned in scale and position to the template eigenfaces. Human face perception, however, is tolerant to changes in both scale and position. Moreover, if a face is transformed further along the morph line representing the deviation of that face from the average face, the transformed face is easily recognized as the same individual; this is the basis of caricature (Leopold et al. 2001). The process of morphing one individual into another (Wolberg 1996) involves both an intensity transform (which eigenfaces model very well) and a simultaneous geometric transform (Figure 3a). Because eigenfaces represent axes

---

**Figure 1**

(a) The two most diagnostic features defining a face comprise the first level of the detection cascade in the Viola-Jones algorithm for face detection. From Viola & Jones 2004. (b) The Sinha algorithm for face detection, showing the ratio-templates defining a face. From Sinha 2002a.
Figure 2

The eigenface algorithm for face recognition. (a) The first 25 eigenvectors computed from the Yale face database (a collection of 165 face images). (b) Eigenface reconstructions of 5 different images, using the 25 eigenfaces shown in panel a. Note that nonface images can have nontrivial projections onto eigenfaces. Courtesy of C. DeCoro.
A computational approach that can represent both spatial and intensity variations. (a) The computer graphics technique of morphing, in which the identity of one individual can be continuously transformed into that of another, provides insights about the nature of the face template. In the middle row, the individual outlined in red is continuously morphed into the individual outlined in green, which requires both a geometric transform and an intensity transform. The top and bottom rows show pure geometric transforms (morphing of the mesh) of the same 2 faces (the top rows show the geometric distortion of the red face into the shape of the green face, and the bottom row shows the distortion of the green face into the shape of the red face). The middle row shows a weighted intensity average of the aligned meshes from the top and bottom rows. From Wolberg 1998. (b) Bags of Pixels variant on the eigenface algorithm. The \((x,y)\) coordinate of each pixel is elevated to the same status as the intensity value. (c) Adding or subtracting traditional eigenfaces to an average face produces only intensity variations at each pixel. Adding or subtracting eigenfaces computed using Bags of Pixels, however, can produce geometric variations in addition to intensity variations. From Jebara 2003.
of intensity values on a fixed spatial basis, the eigenface approach does not interpret caricature transformations as the same individual.

Jebara (2003) proposed a clever way to get around the spatial rigidity of the original eigenface approach: Instead of performing PCA on the intensity values, the size of the representation is tripled, so each pixel conveys not only the image intensity value but also the intensity value’s \((x, y)\) location. Then PCA can be done on the triple-sized image containing a concatenation of \((x, y, I)\) values (Figure 3b,c). The power of this approach is that spatial coordinates are treated just like intensity coordinates, and thus the resultant eigenfaces represent both geometric and intensity variations. The fact that this bags of pixels approach performs three orders of magnitude better than standard eigenface analysis on face sets with changes in pose, illumination, and expression is computational proof of the importance of representing geometric variations in addition to intensity variations.

**Categorization**

Turk and Pentland used a simple Euclidean distance metric on face eigen-coordinates to perform recognition. More powerful classifiers that have been applied to the problem of face recognition include Fisher linear discriminants (Belhumeur et al. 1997), Bayesian estimation (Moghaddam et al. 2000), and support vector machines (Shakhnarovich & Moghaddam 2004). These classification techniques can be regarded as second-tier add-ons to the basic eigenface measurement system. Measurement yields analog descriptions, whereas classification is nonlinear and yields discrete boundaries between descriptions.

Separating the process of measurement from the process of classification gives a computational system maximum flexibility because different categorizations (e.g., emotion, identity, gender) can all operate on the same set of basic eigenvector projections. Gender determination can be based on large eigenvalue eigenvectors, whereas identification of individuals relies on lower-value eigenvectors (O’Toole et al. 1993). Furthermore, because classifications are necessarily nonlinear, the independence of classification mechanisms from measurement mechanisms would be very exciting from an experimental point of view because the templates for measurement could thus be linear, and therefore their detailed structure could be mapped. We will return to the idea of linear measurement mechanisms when we discuss tuning properties of face cells.

**Invariance**

Developing position and scale invariant recognition is a huge challenge for artificial face-recognition systems. Initial attempts to compute a meaningful set of eigen-coordinates for a face required that the face be accurately aligned in scale, position, and rotation angle to the template eigenfaces. However, if, as we propose, face detection precedes measurement, the detector can determine the location, size, and rotation angle of the eyes and face outline and then use these to normalize the input to face-measurement units.

**Summary**

The main lesson we can extract from artificial systems for face processing is that detection and recognition are distinct processes, with distinct goals, primitives (coarse contrast relationships vs. detailed holistic templates), and computational architectures (filter cascade vs. parallel measurements). By preceding recognition, detection can act as a domain-specific filter to gate subsequent processing and can include alignment and segmentation, preparing the face representations for subsequent measurement. The effectiveness of the eigenface algorithm for face recognition shows that faces can be represented by their deviation from the average in a compressed subspace. To characterize faces most effectively, this subspace needs to include spatial variations as well as intensity variations.

Some machine vision models of recognition use common meta-algorithms to learn the primitives for both detection and recognition
of faces (Riesenhuber & Poggio 2000, Ullman 2007). Thus the two processes may share core computational principles. Whether biological systems use discrete steps of detection, measurement, and classification to recognize faces is a question that can only be resolved empirically.

**HUMAN BEHAVIOR AND FUNCTIONAL IMAGING**

The extensive behavioral literature on face perception provides a rich source of clues about the nature of the computations performed in processing faces (Figure 4). One of the hallmarks of face processing is that recognition performance drops substantially when faces are presented upside down (Figure 4a) or in negative contrast, and both effects are much smaller for objects (Kemp et al. 1990, Yin 1969). We propose that both these properties can be explained if only upright, positive-contrast faces gain access to the face-processing system, i.e., if an upright, positive-contrast template is used for face detection. This template may be innate in humans, as evidenced by the tendency for newborns to track normal schematic faces longer.

![Figure 4](https://www.annualreviews.org/doi/abs/10.1146/annurev.neuro.31.060607.164509?journalCode=arns)

**Figure 4**

Behavioral observations on the nature of human face processing. (a) Flip the page upside down. The Thatcher Illusion shows that faces are obligatorily processed as wholes (an identical pair of features such as the upright and inverted mouth can appear similar or dramatically different depending on the surrounding context). From Thompson 1980. (b) Robustness of face identification to caricature. (c) Adaptation: Run your eyes along the 5 red dots for a minute, and then shift your gaze to the single red dot. From Afraz & Cavanagh 2008. (d) Robustness to compression. From Sinha et al. 2006. (e) The importance of external features. From Sinha & Poggio 1996. (f) Robustness to low resolution. From Sinha 2002b.
Prosopagnosia: highly specific inability to recognize faces, due to either congenital brain miswiring ("developmental prosopagnosia") or focal brain lesions ("acquired prosopagnosia")

Norm-Based Coding

Caricatures are remarkably powerful in evoking recognition (Figure 4b). Caricatured faces are often more identifiable than veridical photographs (Lee et al. 2000). This finding has led to the proposal that faces are represented in terms of their deviation from the norm, or average, face (Leopold et al. 2001, Rhodes et al. 1987). Furthermore, the existence of face aftereffects (Figure 4c) shows that the face norm is adaptable (Webster & MacLin 1999). Because such face aftereffects transfer across retinal positions (Leopold et al. 2001) and image sizes (Jeffery et al. 2006), they apparently do not reflect adaptation to specific low-level image features, but instead indicate adaptation of higher-level representations. This face identity aftereffect was interpreted as indicating that adaptation to a given face shifts the norm or average face in the direction of the adapting face, making faces on the opposite side of the norm more distinctive (i.e., more different from the norm). To explain these results Rhodes & Jeffrey (2006) propose that face identity is coded by pairs of neural populations that are adaptively tuned to above-average and below-average values along each dimension of face space.

Opposite adaptation can occur simultaneously for upright and inverted faces, consistent with the idea that distinct neural pathways underlie the coding (and adaptation to) upright versus inverted faces (Rhodes & Jeffery 2006). Finally, although norm-based coding can work only for classes of stimuli that have similar enough first-order shape that a norm can be defined, this situation may not be unique to faces. Rhodes & McLean (1990) showed evidence for norm-based coding for images of birds, and adaptation effects can also be observed for simple shapes such as taper and overall curvature (Suzuki & Cavanagh 1998). Thus adaptive norm-based coding may be a general feature of high-level form-coding processes.

Detection

As argued in the modeling section, it is computationally efficient to separate detection and recognition and to have detection precede recognition because detection can act as a domain-specific filter to make the recognition process more efficient (by focusing recognition on regions actually containing faces). That there are also separate detection and recognition stages in human face processing fits with one of the most striking findings from the neuropsychology literature: Patient CK, who was severely impaired at object recognition, including many basic midlevel visual processes, was nonetheless 100% normal at face recognition (Moscovitch et al. 1997). His pattern of deficits indicated that face processing is not simply a final stage tacked onto the end of the nonface object recognition pathway but rather a completely different pathway that branches away from object recognition early in the visual hierarchy, and it is this branching off that we propose to equate with the detection process. CK’s dissociation is illustrated by his perception of the painting of a face made up of vegetables by Arcimbaldo—CK sees the face but not the constituent vegetables.

CK’s ability to recognize famous or familiar faces was at least as good as normal controls, until the faces were shown upside down, and then his performance became much worse than that of controls. Conversely, patients with prosopagnosia perform better than controls in recognizing inverted faces (Farah et al. 1995). This double dissociation of the inversion effect is consistent with the existence of a face-specific processing system that can be accessed only by upright faces, present in CK and absent in prosopagnosics. Presumably, CK can process objects using only the face-specific system, prosopagnosics have a general object-recognition system but not the face-specific system, and normal subjects have both systems. The general nonface object system is not as good at processing faces as the face-specific system (hence the inversion effect in normal subjects), is missing in CK (hence his
Holistic Processing of Faces

Face processing is said to be distinct from nonface object processing in that it is more holistic; that is, faces are represented as nondecomposed wholes rather than as a combination of independently represented component parts (eyes, nose, mouth) and the relations between them (Farah et al. 1998). Evidence for holistic processing of faces comes from a number of behavioral paradigms, of which the two most cited are the part-whole effect (Tanaka & Farah 1993) and the composite effect (Young et al. 1987). In the part-whole effect, subjects are better at distinguishing two face parts in the context of a whole face than in isolation. In the composite effect, subjects are slower to identify half of a chimeric face aligned with an inconsistent other half-face than if the two half-faces are misaligned (Young et al. 1987). As with the part-whole effect, the composite effect indicates that even when subjects attempt to process only part of the face, they suffer interference from the other parts of the face, suggesting a lack of access to parts of the face and mandatory processing of the whole face.

One interpretation of the uniqueness of face processing is that it uses special neural machinery not shared by other kinds of objects, an idea that is consistent with functional imaging studies, as described below. Another interpretation is that holistic processing is characteristic of any kind of object that must be distinguished on a subordinate level, especially objects with which the subject is highly trained or familiar (Diamond & Carey 1986). It is not yet clear what the perceptual phenomenology of holistic processing implies either mechanistically or computationally. We suggest that holistic face processing can be explained by an obligatory detection stage that uses a coarse upright template to detect whole faces (Figure 5). This model explains the composite effect because an aligned chimera would be detected as a whole face and therefore would be processed as a unit by subsequent measurement and classification stages.

However, we cannot rule out alternatives, such as one-stage models in which both face detection and identification are carried out by the same set of face-selective cells. In this case, to explain holistic properties of face processing, we would have to postulate that individual face cells, unlike nonface cells, are selective not just for local features but for whole faces or that the readout of face information must comprise all or most of the population code. Either or both of these models would produce the behavioral holistic effects, even without an antecedent detection gate. The key evidence favoring our early detection gating hypothesis over a single-stage system comes from the identification of a series of face-selective areas in the macaque (Pinsk et al. 2005, Tsao et al. 2003) and the finding that an area early in this hierarchy already consists entirely of face-selective cells (Tsao et al. 2006); both these results are discussed more extensively below.

Although faces are unique in the degree to which they are processed holistically, other nonface objects can also show holistic effects, especially well-learned categories; for review see Gauthier & Tarr (2002). Words may approach faces in the degree to which they are processed holistically: Coltheart et al. (1993) found that some acquired dyslexics can read whole words and understand their meanings but cannot distinguish individual letters making up the words. And Anstis (2005) showed that word recognition can show the composite effect, in that observers cannot tell whether two words have same or different top halves.

HUMAN FUNCTIONAL IMAGING

Positron emission tomography studies initially showed activation of the fusiform gyrus in a variety of face-perception tasks (Haxby et al. 1991, Sergent et al. 1992), and fMRI subsequently
### a) Stimulus Detection and Winner-Take-All

<table>
<thead>
<tr>
<th>Stimulus</th>
<th>After detection</th>
<th>Bill template</th>
<th>Jesse template</th>
<th>Winner take all</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image1" alt="Bill" /></td>
<td><img src="image2" alt="Bill" /></td>
<td><img src="image3" alt="Bill" /></td>
<td><img src="image4" alt="Bill" /></td>
<td>Bill</td>
</tr>
<tr>
<td><img src="image5" alt="Jesse" /></td>
<td><img src="image6" alt="Jesse" /></td>
<td><img src="image7" alt="Jesse" /></td>
<td><img src="image8" alt="Jesse" /></td>
<td>Jesse</td>
</tr>
<tr>
<td><img src="image9" alt="Winner" /></td>
<td><img src="image10" alt="Winner" /></td>
<td><img src="image11" alt="Winner" /></td>
<td><img src="image12" alt="Winner" /></td>
<td>Winner take all</td>
</tr>
</tbody>
</table>

### b) Car Detection and Winner-Take-All

<table>
<thead>
<tr>
<th>Stimulus</th>
<th>After detection</th>
<th>Beetle template</th>
<th>Isuzu template</th>
<th>Winner take all</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image13" alt="Beetle" /></td>
<td><img src="image14" alt="Beetle" /></td>
<td><img src="image15" alt="Beetle" /></td>
<td><img src="image16" alt="Beetle" /></td>
<td>Beetle</td>
</tr>
<tr>
<td><img src="image17" alt="Isuzu" /></td>
<td><img src="image18" alt="Isuzu" /></td>
<td><img src="image19" alt="Isuzu" /></td>
<td><img src="image20" alt="Isuzu" /></td>
<td>Isuzu</td>
</tr>
<tr>
<td><img src="image21" alt="Winner" /></td>
<td><img src="image22" alt="Winner" /></td>
<td><img src="image23" alt="Winner" /></td>
<td><img src="image24" alt="Winner" /></td>
<td>Winner take all</td>
</tr>
</tbody>
</table>

![Red Box](image25)
revealed more specificity in these cortical regions for faces with demonstrations of fusiform regions that responded more strongly to faces than to letter strings and textures (Puce et al. 1996), flowers (McCarthy et al. 1997), everyday objects, houses, and hands (Kanwisher et al. 1997). Although face-specific fMRI activation can also be seen in the superior temporal sulcus (fSTS) and in part of the occipital lobe [the occipital face area (OFA)], the most robust face-selective activation is consistently found on the lateral side of the right mid-fusiform gyrus, the fusiform face area (FFA) (Kanwisher et al. 1997) (Figure 6). The fact that this part of the brain is activated selectively in response to faces indicates that activity in this region must arise at or subsequent to a detection stage.

Many studies support the idea that the FFA is activated specifically by faces and not by the low-level stimulus features usually present in faces, that is, activity in the FFA indicates that stimuli have been detected as faces: The FFA shows increased blood flow in response to a wide variety of face stimuli: front and profile photographs of faces (Tong et al. 2000), line drawings of faces (Spiridon & Kanwisher 2002), and animal faces (Tong et al. 2000). Furthermore, the FFA BOLD signal to upright Mooney faces (low-information two-tone pictures of faces; Mooney 1957) is almost twice as strong as to inverted Mooney stimuli (which have similar low-level features but do not look like faces) (Kanwisher et al. 1998). Finally, for bistable stimuli such as the illusory face-vease, or for binocularly rivalrous stimuli in which a face is presented to one eye and a nonface is presented to the other eye, the FFA responds more strongly when subjects perceive a face than when they do not, even though the retinal stimulation is unchanged (Andrews et al. 2002, Hasson et al. 2001).

Although the FFA shows the strongest increase in blood flow in response to faces, it does also respond to nonface objects. Therefore, two alternative hypotheses have been proposed to the idea that activity in the FFA represents face-specific processing. First is the expertise hypothesis. According to this idea, the FFA is engaged not in processing faces per se, but rather in processing any sets of stimuli that share a common shape and for which the subject has gained substantial expertise (Tarr & Gauthier 2000). Second is the distributed coding hypothesis: In an important challenge to a more modular view of face and object processing, Haxby et al. (2001) argued that objects and faces are coded via the distributed profile of neuronal activity across much of the ventral visual pathway. Central to this view is the suggestion that nonpreferred responses, for example, to objects in the FFA, may form an important part of the neural code for those objects. The functional significance of the smaller but still significant response of the FFA to nonface objects will hopefully be unraveled by the combined assaults of higher-resolution imaging in humans and single-unit recordings in nonhuman primates.

Measurement and Categorization

Does the human brain use separate systems for face measurement and face classification? Some fMRI evidence suggests that it does. For

---

**Figure 5**

We propose that holistic (composite) effects of face processing can be explained by a detection stage that obligatorily segments faces as a whole. Subjects are asked to identify the top (faces) or left (car) part of each chimera (third and fourth rows) or simply to identify the object (first and second rows). Four face (a) and car (b) stimuli are detected, projected onto holistic templates, and then identified through a winner-take-all mechanism. The numbers in the third and fourth columns indicate the result of projecting each stimulus, after detection, onto the respective templates. Aligned faces are obligatorily detected as a whole, but misaligned faces and cars are not, and therefore their attended parts can be processed independently. According to this hypothesis, the essential difference between face (a) and nonface (b) processing occurs at the detection stage (red boxes). Subsequent measurement and classification could use similar mechanisms.
Face-selective regions in one representative subject. Face-selective regions (yellow) were defined as regions that respond more strongly to faces than to houses, cars, and novel objects ($p < 10^{-4}$). From Grill-Spector et al. 2004.

example, in a study of morphing between Marilyn Monroe and Margaret Thatcher, adaptation strength in the OFA followed the amount of physical similarity along the morph line, while in the FFA it followed the perceived identity (Rotshtein et al. 2005), suggesting that the OFA performs measurement and the FFA performs classification. However, another study indicates that release from adaptation occurs in the FFA when physical differences are unaccompanied by changes in perceived identity (Yue et al. 2006).

According to Bruce & Young (1986), the processing of facial expression (one form of categorization) and facial identity (another form of categorization) takes separate routes. Haxby and colleagues (2000) proposed a neural basis for this model. According to this idea, the inferior occipital gyri are involved in early perception of facial features (i.e., measurement). The pathway then diverges, with one branch going to the superior temporal sulcus, which is proposed to be responsible for processing changeable aspects of faces including direction of eye gaze, view angle, emotional expression, and lip movement. The other projection is to the lateral fusiform gyrus, which is responsible for processing identity. A recent review has challenged the Bruce and Young model, arguing that changeable aspects and invariant identity may instead be processed together and rely on partially overlapping visual representations (Calder & Young 2005).

Invariance

Several studies have used fMRI adaptation for face identity in the FFA and found invariance to image size (Andrews & Ewbank 2004) and spatial scale (Eger et al. 2004). Thus representations in the FFA are not tied to low-level image properties, but instead show at least some invariance to simple image transformations, though not to viewpoint (Pourtois et al. 2005).

Summary

Behavioral studies complement computational approaches by indicating that specialized...
Machinery may be used to process faces and that a face-detection stage gates the flow of information into this domain-specific module. The filters, or templates, used by this detection stage require an upright, positive contrast face, with the usual arrangement of features, and images that do not fit the template are analyzed only by the general object-recognition system. Even images that pass into the face-specific module are probably also processed in parallel by the general system, but the face module appears to process images differently from the general object system: Face processing is holistic in the sense that we cannot process individual face parts without being influenced by the whole face. We suggest that this difference arises early in the face processing pathway. The face-detection stage may, in addition to gating access, obligatorily segment faces as a whole for further processing by the face module. Finally, substantial recent evidence suggests that face identity is coded in an adaptive norm-based fashion.

Human imaging studies converge on the conclusion that faces are processed in specific locations in the temporal lobe, but the degree of specialization for faces within these locations is debated. The modular interpretation is consistent with neurological findings and, as described below, with single-unit recordings in macaques. The role of experience in determining both the localization of face processing and its holistic characteristics is also debated. And the relationship, if any, between modular organization and holistic processing is completely unexplored. Only a few visual object categories show functional localization in fMRI: faces, body parts, places, and words (for review see Cohen & Dehaene 2004, Grill-Spector & Malach 2004). Faces, bodies, and places are all biologically significant, and their neural machinery could conceivably be genetically programmed, but the use of writing arose too recently in human history for word processing to be genetically determined. Therefore, at least one kind of anatomical compartmentalization must be due to extensive experience. We have suggested that the existence of discrete brain regions dedicated to face processing implies an obligatory detection stage and that an obligatory detection stage results in holistic processing. What we know about word processing suggests that it too displays holistic properties, and it is localized, interestingly, in the left hemisphere in an almost mirror symmetric location to the position of the FFA in the right hemisphere (Cohen & Dehaene 2004, Hasson et al. 2002).

MONKEY fMRI AND SINGLE-UNIT PHYSIOLOGY

Detection

The seminal finding by Gross and his colleagues (1969, 1972) that there exist cells in inferotemporal cortex (IT) that are driven optimally by complex biologically relevant stimuli, such as hands or faces, was novel and initially not well accepted, despite the fact that Konorski (1967) had predicted the existence of face-selective cells, or gnostic units, and that they would be found in IT. Although IT cells do not generally appear to be detectors for complex objects, there are consistently observed populations of cells selectively responsive to faces, bodies, and hands, suggesting that faces, bodies, and hands are treated differently from other types of complex patterns, consistent with their also being among the only object categories, aside from words and numbers, that show localization in human fMRI. But the strong possibility remained that these cells were not really tuned to biologically relevant objects, but rather to some more abstract basis set, in which all possible shapes are represented by different cells and some cells were tuned to particular parameters that happened to fit the face or hand stimuli better than any of the other objects tested. Foldiak et al. (2004) recently provided evidence that face selectivity is not just an incidental property of cells tuned to an exhaustive set of image features: They presented 600–1200 stimuli, randomly chosen from several image archives, to cells recorded from both the upper and the lower bank of the STS and found...
that the distribution of tuning to these images showed bimodality, i.e., cells were either predominantly face selective or not face selective. It is not unprecedented to have specialized neural systems for socially important functions: Birds have evolved specialized structures for the perception and generation of song, and in humans there are specialized parts of the auditory and motor systems devoted specifically to language. Direct evidence that some face cells are used for face detection comes from a microstimulation study by Afraz et al. (2006). Monkeys were trained to discriminate between noisy pictures of faces and nonface objects. Through systematic sampling, Afraz et al. identified cortical locations where clusters of face-selective cells could be reliably recorded. When they stimulated these regions and observed the monkeys’ perceptual choices, they found a shift in the psychometric curve favoring detection of a face.

**Holistic Processing of Faces**

In general, face cells require an intact face and are not selective just for individual features (Bruce et al. 1981; Desimone et al. 1984; Kobatake & Tanaka 1994; Leonard et al. 1985; Oram & Perrett 1992; Perrett et al. 1982, 1984; Scalaidhe et al. 1999; Tsao et al. 2006). Figure 7 shows nonlinear combinatorial response properties of a face-selective cell recorded in IT by Kobatake & Tanaka (1994). Out of a large number of three-dimensional objects, this cell responded best to the face of a toy monkey (panel a), and by testing various simplified two-dimensional paper stimuli, they determined that the cell would also respond to a configuration of two black dots over a horizontal line within a disk (panel b) but not in the absence of either the spots or the line (panels c and d) or the circular outline (panel e). The contrast between the inside and the outside of the circle was not critical (panel g), but the spots and the bar had to be darker than the disk (panel b). Thus the cell responded only when the stimulus looked like a face, no matter how simplified.

The response selectivity of face cells indicates that they must not only combine features nonlinearly but also require them to be in a particular spatial configuration. However, such spatial-configuration selective responses and nonlinear combination of features are not restricted to face cells as such behavior has been reported for other kinds of complex object-selective cells in the temporal lobe (Baker et al. 2002, Kobatake & Tanaka 1994, Tanaka et al. 1991). Even earlier in the temporal pathway, nonlinear combinatorial shape selectivity can be seen (Brincat & Connor 2004).

**Anatomical Specialization of Face Cells**

Most studies on face cells reported face-selective cells scattered throughout the temporal lobe, though they tended to be found in clusters (Perrett et al. 1984). Because other kinds of shape selectivities also tend to be clustered (Desimone et al. 1984, Fujita et al. 1992, Tanaka et al. 1991, Wang et al. 1996), it was assumed that within the temporal lobe there was a columnar organization for shape, in which face columns represented just one of many shape-specific types of columns. However, this view was inconsistent with emerging evidence from human neurology and functional imaging that human face processing was localized to specific, reproducible regions of the temporal lobe. The apparent discrepancy was resolved by two recent studies by Tsao et al. (2003, 2006), who found that in monkeys, as in humans, face processing, as revealed by functional imaging, is localized to discrete regions of the temporal lobe, and they further showed that even at the single-unit level, face processing is highly localized (Figure 8; note also Figure 7, top).

Tsao et al. used functional imaging to localize regions in the macaque temporal lobe that were selectively activated by faces, compared with nonface objects, and then they recorded almost 500 single units within the largest of these face-selective regions in two monkeys. They found a remarkable degree of face selectivity within this region; 97% of the cells were face selective, on average showing almost 20-fold larger responses to faces than to nonface.
objects. The region where they recorded was quite posterior in the temporal lobe (6 mm anterior to the interaural canal, corresponding to posterior TE/anterior TEO). The fact that an area consisting almost entirely of face-selective cells exists so early in the ventral stream provides strong support for the hypothesis that the face pathway is gated by an obligatory detection stage.

In light of the clear large-scale organization of face processing in macaques revealed by Tsao et al. and recently by Pinsk et al. (2005),

Figure 7
Holistic face detection. (Top) recording site and receptive-field location of a face cell. (a–h) Response selectivity. From Kobatake & Tanaka (1994).
we reexamined all previous physiological studies that mapped out locations of face-selective cells, and by remapping their face-cell localizations onto a common map, we found that, taken en masse, these studies do show a concentration of face selectivity in two major regions of the temporal lobe, regions that correspond to the middle and anterior face patches described by Tsao and colleagues using functional imaging (Figure 8d).

**The Functional Significance of the Anatomical Localization of Face Processing**

The cerebral cortex is functionally parcellated: Neurons concerned with similar things are
organized into areas and columns, each having extensive interconnections and common inputs and outputs (Mountcastle 1997). It is not surprising that face processing, being an important, identifiable and discrete form of object recognition, is also organized into anatomically discrete processing centers. Individual neurons connect with only a small fraction of the rest of the neurons in the brain, usually to nearby cells, because longer axons delay neural transmission, are energetically expensive, and take up space. Barlow (1986) has noted that facilitatory interactions within a functional area or column could underlie Gestalt linking processes—clustering cells concerned with color or motion might facilitate interactions between parts of the visual field having common color or motion. However, enriched local inhibitory interactions and sharpening of tuning might be an even more important function of colocalization because inhibitory neurons are always local, and long-range intracortical connections are invariably excitatory (Somogyi et al. 1998). Wang et al. (2000) recorded responses in anterior IT to a set of complex stimuli before, during, and after applying the GABA antagonist bicuculline near the recording electrode. In many cases, for both face-selective and nonface-selective cells, blocking local inhibition revealed responses to previously nonactivating stimuli, which were often activating stimuli for neighboring cells. This suggests that neighboring cells refine each other’s response selectivity by mutual inhibition.

**Time Course of Feature-Combination Responses**

Although a large fraction of the information about which face stimulus was shown is carried by the earliest 50 ms of the response of face-selective cells (Tovee et al. 1993), several studies have shown that the information carried by the early part of the response is different from the information carried by later spikes. In particular, the earliest spikes in a response are sufficient for distinguishing faces from other object categories, but information about individual facial identity does not develop until ~50 ms later (Sugase et al. 1999, Tsao et al. 2006).

Similarly, responses in IT to nonface stimuli also become more selective, or sparser, over time (Tanaka et al. 1991, Tamura & Tanaka 2001). Similar temporal dynamics indicative of early detection activity followed by later individual identification activity have been observed for face-selective MEG responses in human occipitotemporal cortex (Liu et al. 2002). The observations that global information precedes finer information are consistent with a role for local inhibition in sharpening tuning within a local cluster of cells having similar response properties. Such response dynamics suggest a feedback or competitive process, whereby cells that respond best to a given stimulus inhibit nearby cells, resulting in a winner-take-all situation.

**Norm-Based Coding**

Recently an idea has emerged for both face processing and general object coding in the temporal lobe—that firing rate represents the magnitude of deviation from a template or norm for that property. Cells in V4 can be tuned to curvature, but the optimal values for curvature are most often found at either extreme or zero curvature, with few cells tuned to intermediate curvature (Pasupathy & Connor 2001). Kayaert and colleagues (2005a) found norm-based tuning for shapes in IT; neurons tuned to different shapes tended to show monotonic tuning, with maximum responses to extreme values of those shapes. Lastly, Leopold et al. (2006) recorded from face-responsive cells in anterior IT and found that most cells were tuned around an identity-ambiguous average human face, showing maximum firing to faces farthest from an average face (i.e., tuning was V-shaped around the average). Freiwald et al. (2005), on the other hand, reported that many cells in the macaque middle face patch showed monotonic tuning curves to different feature dimensions in a large cartoon face space, with the maximum response at one extreme and the minimum response at
the opposite extreme (Figure 9). This ramp-shaped tuning is consistent with the model proposed by Rhodes et al. (2004) for explaining the face-adaptation effect (Figure 5b)—that each face feature axis is coded by two opponent cell populations; thus the face norm would be implicitly represented as the virtual point of intersection between face cell populations with opponent ramp-shaped feature tuning curves. For both faces and nonface objects, many cells show tuning to several feature dimensions, and the tuning is separable, or independent, for the different tuning axes (Freiwald et al. 2005, Kayaert et al. 2005b).

**Invariance**

Face-selective cells in the temporal lobe are usually position and scale invariant in their ability to detect and distinguish faces, but they are seldom view and angle invariant (Desimone et al. 1984; Perrett et al. 1984, 1985, 1989, 1991; Rolls & Baylis 1986; Tanaka et al. 1991; Tovee et al. 1994; Tsao et al. 2006). The marked view selectivity of some IT cells may reflect a role in interpreting social gestures (who is looking at whom) (Argyle & Cook 1976, Bertrand 1969). De Souza et al. (2005) recently found a striking pattern of view selectivity in rostral versus
caudal anterior STS. In caudal anterior STS, they found mirror-symmetric view-tuned cells, but in rostral anterior STS, view tuning was not mirror symmetric. Furthermore, view angle and gaze direction interacted, with neurons selective for a particular combination of face view and direction of gaze and often were strongly modulated by eye contact.

Recordings from the medial temporal lobe of human epilepsy patients have revealed the existence of cells that respond to familiar individuals in a highly invariant manner (Quiroga et al. 2005), as expected of a grandmother cell. For example, some cells responded to multiple pictures of a well-known individual as well as to a letter string of the person’s name but were unresponsive to all other images. Such individual-specific cells have not been found in the lateral inferior temporal lobe, where most face cells in monkeys have been recorded, although as a population, cells in the anterior inferior temporal gyrus of the macaque can represent view-invariant identity (Eifuku et al. 2004).

**Summary**

The correlation between fMRI localization of face processing in macaques and the strong clustering of physiologically identified face-selective cells supports the idea of domain specificity, suggested by neurological findings and fMRI studies in humans. The strength and predominance of face selectivity within the middle face patch are not consistent with either the expertise hypothesis or the distributed coding model. The existence of neurons located at an early stage of form processing in the macaque brain that respond selectively to faces supports the idea that face processing begins with a detection stage, and the response properties of face cells indicate that this stage is highly nonlinear.

However, face cells seem to measure different face variables independently and linearly, so how does this reconcile with evidence that face perception in humans is holistic; i.e., how can we explain the composite effect and the part-whole effect neurally? We suggest that both these apparently nonlinear perceptual effects are consistent with a linear neural measuring stage if the preceding detection stage is holistic and nonlinear. One surprising result from physiological studies on face processing is the preponderance of view-selective units, but what role they play in face processing is still unclear.

**FUTURE DIRECTIONS**

1. Is face processing unique? We do not yet understand the details of how either faces or nonface objects are represented in the brain—perceptual studies have shown major differences in the ways that faces and objects are recognized, but there are nevertheless similarities in the response properties between face-selective cells and object-selective cells in IT. Both face- and object-selective cells in IT show tuning characteristics of a norm-based code. A variety of evidence suggest that our perception of faces is holistic, but processing of some nonface objects, like words, also shows important context effects. One fact is clear: The basic computational challenges to face processing are common to all object recognition (namely, detection, measurement, and classification). What is a face template in computational and neural terms, and how does it differ from a chair template? A truly satisfying answer to the question of whether face processing is unique will come only when we understand the precise neural mechanism underlying both face and nonface object recognition.
2. Is face processing modular? Perhaps the most striking result to come from the neurobiological research on face perception in the past decade is that specialized machinery is used for processing faces. Evidence reveals a fundamental specialization both at the gross anatomical level and at the level of single cells. It will be exciting to move forward along this pathway to understand how these face cells are used for different high-level percepts and behaviors; e.g., conveying invariant identity, expression, direction of attention, social dominance. But we believe that equally important new insights will come from looking back, asking how these cells acquire their face selectivity—undertaking a systematic study of the face-detection process.

3. What makes face processing special? We have proposed that what is special about face processing is that it is gated by an obligatory detection process. Such a design would be computationally elegant (by allowing for fast domain-specific filtering, segmentation, and alignment prior to fine-grained identification) and could explain the existence of face cells, face areas, prosopagnosia, and holistic processing. This detection-gating hypothesis naturally leads to the idea that there are two distinct classes of face cells: face-recognition cells, which encode different kinds of face templates, and face-detector cells, which (contrary to their name) could perform the triple function of detection, segmentation, and alignment. However, it is also possible that detection and discrimination are carried out by the same cells (either simultaneously or sequentially). Either way, we should at least be able to find out the answer. Because we know that face-selective cells are coding faces, we can distinguish detection-related activity from discrimination-related activity, which is impossible when one is studying a cell whose form specialization is unknown. Perhaps what is truly special about face processing is that it is now amenable to being understood. We have a beautiful hierarchy, a gift from nature, and we should exploit it in both directions.

DISCLOSURE STATEMENT

The authors are not aware of any biases that might be perceived as affecting the objectivity of this review.

LITERATURE CITED

Barlow HB. 1986. Why have multiple cortical areas? Vision Res. 26:81–90


Jebara T. 2003. *Images as bags of pixels.* Presented at IEEE Int. Conf. Comp. Vis. (ICCV’03), 9th, Nice, France


Rolls ET, Baylis GC. 1986. Size and contrast have only small effects on the responses to faces of neurons in the cortex of the superior temporal sulcus of the monkey. *Exp. Brain Res.* 65:38–48


Contents

Cerebellum-Like Structures and Their Implications for Cerebellar Function
Curtis C. Bell, Victor Han, and Nathaniel B. Sawtell ........................................... 1

Spike Timing–Dependent Plasticity: A Hebbian Learning Rule
Natalia Caporale and Yang Dan .................................................................................. 25

Balancing Structure and Function at Hippocampal Dendritic Spines
Jennifer N. Bourne and Kristen M. Harris ................................................................. 47

Place Cells, Grid Cells, and the Brain’s Spatial Representation System
Edward I. Moser, Emilio Kropff, and May-Britt Moser ........................................... 69

Mitochondrial Disorders in the Nervous System
Salvatore DiMauro and Eric A. Schon ........................................................................ 91

Vestibular System: The Many Facets of a Multimodal Sense
Dora E. Angelaki and Kathleen E. Cullen ................................................................. 125

Role of Axonal Transport in Neurodegenerative Diseases
Kurt J. De Vos, Andrew J. Grierson, Steven Ackerley, and Christopher C.J. Miller ... 151

Active and Passive Immunotherapy for Neurodegenerative Disorders
David L. Brody and David M. Holtzman ................................................................. 175

Descending Pathways in Motor Control
Roger N. Lemon ....................................................................................................... 195

Task Set and Prefrontal Cortex
Katsuyuki Sakai ....................................................................................................... 219

Multiple Sclerosis: An Immune or Neurodegenerative Disorder?
Bruce D. Trapp and Klaus-Armin Nave ................................................................. 247

Multifunctional Pattern-Generating Circuits
K.L. Briggman and W.B. Kristan, Jr. ........................................................................ 271

Retinal Axon Growth at the Optic Chiasm: To Cross or Not to Cross
Timothy J. Petros, Alexandra Rebsam, and Carol A. Mason ................................. 295
Brain Circuits for the Internal Monitoring of Movements
Marc A. Sommer and Robert H. Wurtz .......................................................... 317

Wnt Signaling in Neural Circuit Assembly
Patricia C. Salinas and Yimin Zou ................................................................. 339

Habits, Rituals, and the Evaluative Brain
Ann M. Graybiel .............................................................................................. 359

Mechanisms of Self-Motion Perception
Kenneth H. Britten ......................................................................................... 389

Mechanisms of Face Perception
Doris Y. Tsao and Margaret S. Livingstone .................................................. 411

The Prion’s Elusive Reason for Being
Adriano Aguzzi, Frank Baumann, and Juliane Bremer .................................. 439

Mechanisms Underlying Development of Visual Maps and
Receptive Fields
Andrew D. Huberman, Marla B. Feller, and Barbara Chapman ................. 479

Neural Substrates of Language Acquisition
Patricia K. Kuhl and Maritza Rivera-Gaxiola ............................................... 511

Axon-Glial Signaling and the Glial Support of Axon Function
Klaus-Armin Nave and Bruce D. Trapp ...................................................... 535

Signaling Mechanisms Linking Neuronal Activity to Gene Expression
and Plasticity of the Nervous System
Steven W. Flavell and Michael E. Greenberg ............................................. 563

Indexes

Cumulative Index of Contributing Authors, Volumes 22–31 ...................... 591
Cumulative Index of Chapter Titles, Volumes 22–31 .................................... 595

Errata

An online log of corrections to *Annual Review of Neuroscience* articles may be found at http://neuro.annualreviews.org/