Turning the other cheek: the viewpoint dependence of facial expression after-effects

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How do we visually encode facial expressions? Is this done by viewpoint-dependent mechanisms representing facial expressions as two-dimensional templates or do we build more complex viewpoint independent three-dimensional representations? Recent facial adaptation techniques offer a powerful way to address these questions. Prolonged viewing of a stimulus (adaptation) changes the perception of subsequently viewed stimuli (an after-effect). Adaptation to a particular attribute is believed to target those neural mechanisms encoding that attribute. We gathered images of facial expressions taken simultaneously from five different viewpoints evenly spread from the three-quarter leftward to the three-quarter rightward facing view. We measured the strength of expression after-effects as a function of the difference between adaptation and test viewpoints. Our data show that, although there is a decrease in after-effect over test viewpoint, there remains a substantial after-effect when adapt and test are at differing three-quarter views. We take these results to indicate that neural systems encoding facial expressions contain a mixture of viewpoint-dependent and viewpoint-independent elements. This accords with evidence from single cell recording studies in macaque and is consonant with a view in which viewpoint-independent expression encoding arises from a combination of view-dependent expression-sensitive responses.

Keywords: facial expressions; adaptation; after-effects; viewpoint dependence; psychophysics

1. INTRODUCTION

In this paper, we investigate viewpoint dependence in our encoding of facial expression. Investigation of viewpoint dependence runs through the literature on human object perception as it addresses the nature of the underlying models. The basic issue is whether we encode objects as series of two-dimensional templates or whether we construct three-dimensional viewpoint invariant representations (Biederman 1987; Bülbö® & Edelman 1992). In relation to facial identity, these questions have traditionally been studied using paradigms in which subjects are trained on an initially unfamiliar identity at one viewpoint and then tested at another (Troje & Bülbö® 1996; Hill et al. 1997; Newell et al. 1999; Watson et al. 2005); an approach that would clearly be difficult to extend to facial expressions. Recent adaptation-based paradigms offer a novel and powerful method for addressing this issue.

Adaptation paradigms examine how prolonged view of a stimulus can modify the perception of subsequent-related stimuli. For example, you might examine a line tilted leftwards of vertical for 30 s or so. When you are then briefly presented with a vertical test line, this will appear to be tilted rightwards of vertical (Gibson & Radner 1937). The phenomenon appears ubiquitous within the visual system and is believed to represent recalibration of the neural systems encoding the adapted attribute

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three-quarter views), a substantial after-effect is still obtained (Benton et al. 2006).

In comparison to facial identity, little is known about viewpoint dependence in our encoding of facial expressions. Indeed, there is no particular reason to expect it to follow a similar pattern to that found with identity. Facial expression and identity are largely held to be processed by different mechanisms (Bruce & Young 1986) and to be encoded in different brain regions. Permanent aspects of faces (such as identity and gender) are thought to be encoded by face-sensitive neurons in the fusiform gyrus while changeable aspects (such as facial expression, gaze direction and viewpoint) are thought to be encoded in the superior temporal sulcus (Perrett et al. 1987; Hasselmo et al. 1989a; Haxby et al. 2000; Rolls 2000).

This leads us to differ in this expectation of viewpoint dependence in adaptation to identity and adaptation to facial expression. The identity encoding system is conceived as not encoding viewpoint, or at least, not as its final outcome. However, viewpoint independence might reasonably be achieved through summing the outputs of viewpoint-dependent neurons (Rolls 2000). Such a model is supported by the mixture of viewpoint dependence and independence found in adaptation to facial identity (Benton et al. 2006). In contrast, facial expression is held to be encoded by a neural substrate that also encodes viewpoint. On this basis, we might reasonably expect expression adaptation to show strong viewpoint dependence. This expectation is supported by studies showing that gaze direction and viewpoint modulate our processing of facial expressions (Kappas et al. 1994; Lyons et al. 2000; Adams & Kleck 2003).

In the present study, we use a recently developed adaptation-based methodology (Benton et al. 2006) to examine viewpoint dependence in our encoding of facial expressions. We wished our stimuli to be as naturalistic as possible; we therefore used pictures of actors producing facial expressions taken simultaneously from a variety of views. In contrast to our initial expectation, we find that our encoding of facial expression displays substantial viewpoint invariance.

2. METHODS AND RESULTS

We created morphs between different facial expressions to produce sequences of images that changed gradually from one expression to another. The expressions at either end of the sequence are readily identified, but somewhere in the middle of the sequence, each subject will have a balance point, the estimated point along the morph sequence where a subject is equally likely to judge an image as either of the two original expressions. For example, with a happy to sad morph sequence, when subjects adapt to the happy image, the balance point shifts closer to the happy end of the sequence. Conversely, when subjects adapt to sad, the balance point shifts towards the sad end of the sequence.

In the following experiments, we measured after-effects by measuring adaptation-induced shifts in balance points. We used a classic adaptation/top-up paradigm in which subjects were presented with sequences of pairs of images. Each pair contained an adaptation image followed by a brief test image (to which subjects make responses). Within a sequence, the adaptation image in the initial pair was presented for 30 s. Each subsequent adaptation image served to top up the initial adaptation and was presented for 5 s.

(a) Materials

We recruited actors (11 females and 8 males) from the Drama Department at the University of Bristol and elicited facial expressions (produced upon demand) while filming simultaneously from a variety of angles using a high-definition (HD) multi-camera rig. The five Dalsa DS-25-02M30 colour cameras were placed at $-45^\circ$, $-22.5^\circ$, $0^\circ$, $22.5^\circ$ and $45^\circ$ equidistant from a stool on which the actors sat. The angles refer to rotations parallel to the ground plane (seen from above) with $0^\circ$ referring to a front view (i.e. full face) of the actor. An angle of $45^\circ$ means that the face is seen in a three-quarter rightward facing view. The cameras captured HD frames $(1920 \times 1080$ pixels) at $25$ Hz. We gathered $2$ s of sequence for each actor for each of five facial expressions (anger, disgust, fear, happiness and sadness). Filming took place under controlled lighting conditions in a dedicated studio; we used a mixture of direction and non-directional lighting placed overhead and at knee level to light our actors.

From our database of actors and expressions, we generated three expression morph sequences by morphing between images of full-blown expressions (Tiddeman et al. 2001). We used happy to sad for actor 18 (male) and happy to sad and anger to disgust for actor 5 (female). These are shown in figure 1 and were chosen on the basis of the quality of the expression (judged by the experimenters) and the ability to produce high-quality morphs between the neutral and the full-blown expressions. Each morph sequence contained 101 images (i.e. morph increments of $1\%$). Images were cropped and the edges of the images were blurred to display mean luminance (using Gaussian blur of standard deviation $10$ pixels) so that no hard image edges would be present when the images were displayed on the mean luminance background. Examples from our morph sequences can be found in the electronic supplementary material.

Our images were linearized and then displayed on an Iiyama Vision Master Pro 513 (MA203DT) monitor set to a resolution of $1600 \times 1200$ pixels and a frame rate of $75$ Hz. Display mean luminance was $43.2$ cd m$^{-2}$. The experiments were controlled by a PC; the images were rendered using the Cogent Graphics MATLAB extension developed by John Romaya at the LON at the Wellcome Department of Imaging Neuroscience. Images were presented in the centre of the screen with the remainder set to display mean luminance. Subjects viewed images from $1$ m. At this distance, the interocular distance for the faces presented during the experiments was $1.33^\circ$ of visual angle for actor $18^\circ$ and $1.40^\circ$ for actor 5 (figures given for full face images). The screen itself subtended $22.1^\circ \times 16.7^\circ$.

(b) General procedures

We measured balance points by using an adaptive method of constants procedure (Watt & Andrews 1981) in which subjects viewed images from the morph sequences and were asked to classify these images as one of the two facial expressions from which the images are drawn. We use the responses from $64$ such image presentations (or trials) to
estimate each balance point by fitting a cumulative Gaussian to the resultant data (Wichmann & Hill 2001a). We refer to the group of 64 trials used to measure a balance point as a run.

Subjects completed two types of run, those without adaptation and those with adaptation. We used the unadapted runs primarily for training. In those without adaptation, subjects viewed each image for 1000 ms and indicated their judgement of expression using the arrow keys on a keyboard. There was a minimum gap of 500 ms between stimuli. In adaptation runs, subjects viewed the adaptation image for 30 s before being presented (after a 500 ms gap) with the first test image. Thereafter, each test image was presented after viewing the adaptation image for 5 s (to top up the initial adaptation).

In all, but our final experiment, we used the following procedure to control for low-level retinotopic adaptation (Fang & He 2005; Benton et al. 2006). Subjects were required to fix a small spot presented in the centre of the screen. All adaptation and test images were moved in a circular trajectory around the fixation spot with an angular velocity of 1 revolution every 5 s. The radius of the circle was 0.5°. For each image, the starting point and initial direction of travel were random. For each morph sequence at each angle, we had to decide on a centre point—i.e. the part of the image that actually moves in a circle with the fixation spot at its centre. Vertically these were chosen to lie halfway between the eyes and the mouth. Horizontally they are different for each angle. In the case of the 0° images, they fall on the faces’ vertical midlines; in the 45° images, they lie directly under the centremost eye (Benton et al. 2006); and in the 22.5° images, they lie midway between these points.

Subjects completed a number of runs within a session. A session refers to a number of balance points in a single sitting. Within each session only one adapter angle was used. There was a minimum of a 12 h gap between sessions to minimize any carry over of adaptation. Within sessions the runs were not interleaved—in other words, we gathered one balance point at a time. For each combination of adaptation angle, test angle and morph sequence that we tested, we gathered both unadapted balance points and balance points under adaptation to the two ends of the morph sequence. Within sessions containing unadapted and adapted runs, we always gathered the unadapted balance points first.

When we measure the strength of adaptation, we look at the shift in balance point between the two adaptation conditions. The two adaptation conditions refer to adaptation to the two ends of a morph sequence (i.e. the original expressions). For example, in experiment 1, we measured adaptation to actor 18 using a happy to sad morph sequence. Adaptation to happy makes subsequent images appear less happy so that the balance point shifts towards the happy end of the morph sequence. Under adaptation to sad, the balance points shift in the opposite direction. We take the difference between adapters (rather than between adapted and unadapted balance points) as our metric of strength of
adaptation. We do this because (i) the difference between adapted balance points provides the better signal-to-noise ratio, (ii) adaptation runs occur after training and are therefore likely to be more stable, and (iii) adaptation runs differ only in their adapter whereas adaptation and non-adaptation runs differ in other respects (Benton et al. 2006).

(c) Experiment 1
Three subjects (an experimenter S1 and two naives, S2 and S3) adapted to happy and sad expressions from actor 18. In this experiment, the adaptation viewpoint was always 45°. Test viewpoint was varied over all five available angles. We initially gathered four unadapted balance points for each angle for each participant. We then gathered the adapted balance points—again four for each subject for each combination of adapter type, adapter viewpoint and test viewpoint.

To assess statistical variability, we used parametric bootstrapping to generate 10,000 bootstrap estimates for each balance point (Wichmann & Hill 2001b). We then propagated the bootstrap populations through the relevant averaging and differencing calculations (Benton et al. 2006) to generate 95% confidence limits. These are calculated using the percentile method (Efron & Tibshirani 1993).

Note that all hypotheses testing in the current study is achieved through the use of 95% confidence limits—standard practice in statistical bootstrapping. In terms of the data presented, if the x-axis (difference equals zero) line lies outside the confidence limits associated with a data point, then that point can be considered significantly different from zero with \( p<0.05 \) (two-tailed). When comparing two means, these are significantly different with \( p<0.05 \) when there is less than about 50% overlap of error bars. When there is no overlap, they are significantly different with \( p<0.01 \) (Cumming & Finch 2005).

Results are depicted in figure 2a which essentially shows the strength of the after-effect (coded as the difference between balance points) as a function of test viewpoint. We carried out linear regressions on the bootstrap populations underlying these data to assess the strength of the after-effect as a function of test viewpoint. Our first experiment showed viewpoint dependence in facial expressions at 90° change in test viewpoint. Our first experiment showed viewpoint dependence with substantial after-effects at an angular difference of 90° between adapter and test viewpoints. The results from experiments 2 to 5 demonstrate that this result is not due to the position of the test viewpoint itself but is due to the angular difference between test and adaptation viewpoints. The experiments also demonstrate that the effect extends across different naive subjects and to different actors and different facial expressions. The final experiment shows that the findings generalize to natural viewing conditions.

A comparison of results across experiments 1 and 2 for the two common subjects (S1 and S3) shows similar levels of adaptation within the congruent and incongruent (90° difference) conditions. This indicates little effect of training such as that found with identity adaptation where increased familiarity with a particular identity leads to increased transfer across viewpoint (Jiang et al. 2007). In the current study, any familiarity effect would most probably not be evident owing to the comparatively extensive initial training undergone by our subjects.

3. DISCUSSION
This study uses a novel multi-view face database to investigate viewpoint dependence in facial expressions over changes in viewpoint commonly seen in our visual worlds. We studied the effects of expression adaptation at one viewpoint on the perception of facial expressions at other viewpoints. We found high-level non-retinotopic adaptation that generalized across differences in adapter and test viewpoint. However, we also observed a substantial decrease in the strength of that adaptation as the angle between test and adaptation viewpoints increased. Over a change of 90°, between three-quarter leftward and three-quarter rightward facing views,
there was approximately a 40% decrease in the strength of adaptation.

Recent single cell recordings in macaque superior temporal sulcus show viewpoint-dependent tunings that would predict upwards of a 70% decrease in cell response over a 90° viewpoint change (Földiák et al. 2003). Perrett et al. (1991) looking at view-dependent neurons in macaque superior temporal sulcus, found a number of bimodal neurons which showed responses to both left and right three-quarter views. However, out of a total of 110 neurons responsive to perspective views, only 3 showed this characteristic. Based on these data, it is unlikely that the relatively small size of the viewpoint-dependent decrease found in our study can be accounted for by wide tuning or bimodality of viewpoint-dependent neurons. Instead, our findings are commensurate with a view in which facial expression is encoded by a mixture of viewpoint-dependent and viewpoint-independent mechanisms.

On the surface, our results would seem to argue against a model of expression encoding in which a viewpoint-independent representation is the final outcome. However, adaptation should target those neural systems encoding faces, whether or not they encode viewpoint as well. The endpoint of expression analysis may well be a viewpoint-independent representation; however, if this is built from view-dependent responses which are themselves adapted, this could readily give rise to viewpoint-dependent behaviour. Our adaptation-based findings therefore show simply that expression encoding occurs at viewpoint-dependent and viewpoint-independent levels.

Nevertheless, the fact that we have a substantial viewpoint-independent component implies the existence of viewpoint-independent mechanisms in addition to viewpoint-dependent mechanisms. Given that the input to the visual system is necessarily a two-dimensional image smeared over our retinas, it is clear that three-dimensional representations (whether explicit or not) must be built from two-dimensional information. The question then is, if three-dimensional representations of expressions do exist, are they built from intermediate two-dimensional expression representations or are they built directly,
using something akin to Biederman’s recognition by components model (Biederman 1987)? Transfer of after-effect across viewpoint cannot in itself distinguish between these possibilities. However, the fact that we see viewpoint-dependent behaviour, in addition to viewpoint independence, supports a view in which a viewpoint-independent representation of facial expression is constructed from viewpoint-dependent mechanisms that encode facial expression.

This notion of a mixed mechanism is supported by the neurophysiological literature dealing with viewpoint-dependent responses to faces. Single cell recording studies in macaque have described face-sensitive neurons in the superior temporal sulcus (Perrett et al. 1982, 1985; Hasselmo et al. 1989b), an area believed to underlie the processing of changeable aspects of faces, such as expressions (Haxby et al. 2000). While many of these cells show view-dependent responses, a number show viewpoint-independent responses. Based on these findings, it has been proposed that viewpoint-independent face mechanisms may be created from the summation of responses of viewpoint-dependent neurons tuned to a variety of different viewpoints (Rolls 2000).

The encoding of changeable and fixed aspects of faces is thought to occur in different neural substrates; with the latter in the fusiform gyrus and the former in the superior temporal sulcus (Haxby et al. 2000). This anatomical distinction accords with a prevalent model of face processing in which identity and facial expression are processed through different routes (Bruce & Young 1986). Note that Fox & Barton (2007) showed a decrease in expression after-effects when adaptation and test identities differed. This finding cannot however be taken as evidence for dependence of expression on identity because the effect may not be based on perceived identity per se but may be based on some factor that clearly covaries with identity (such as facial structure).

Adaptation studies of viewpoint dependence in facial identity, when taken together, show a mixture of viewpoint dependence and viewpoint independence (Benton et al. 2006; Jefferay et al. 2006; Jiang et al. 2006) similar to that found with facial expressions in the current study. This mixture seems therefore to be a general property of the encoding of both the changeable and the fixed aspects of faces. The similarity in viewpoint-dependent response between expression and identity seems to occur in spite of the fact that facial identity and facial expression are rather different qualities. Most obviously, the number of facial identities with which we are faced is far larger than the number of discrete facial expressions. Based on these differences, one might reasonably expect to find what we do not—evidence of substantial processing differences between the two. The concordance between our expression and identity findings hints at a common substrate where identity and expression form part of a common distributed representation (Calder et al. 2001; Calder & Young 2005).

In conclusion, our results show viewpoint-dependent expression after-effects over changes in angular difference between adaptation and test viewpoints. However, even over large changes in viewpoint, the effect of adaptation is still substantial. The viewpoint-dependent adaptation that we describe is non-retinotopic and can be observed in different identities across different facial expressions and is preserved in natural viewing conditions. Our findings demonstrate that the human encoding of facial expression occurs through a mixture of viewpoint-dependent and viewpoint-independent mechanisms. This is similar to results obtained in single cell recording studies of macaque and may well indicate a hierarchical organization in which the responses of viewpoint-independent expression-encoding neurons are summed to produce viewpoint invariant expression-dependent responses.

REFERENCES


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