# Implied Motion Activation in Cortical Area MT Can Be Explained by Visual Low-level Features

Jeannette A. M. Lorteije<sup>1,2</sup>, Nick E. Barraclough<sup>3,4</sup>, Tjeerd Jellema<sup>3</sup>, Mathijs Raemaekers<sup>1</sup>, Jacob Duijnhouwer<sup>1</sup>, Dengke Xiao<sup>4</sup>, Mike W. Oram<sup>4</sup>, Martin J. M. Lankheet<sup>1,5</sup>, David I. Perrett<sup>4</sup>, and Richard J. A. van Wezel<sup>1,6</sup>

#### Abstract

■ To investigate form-related activity in motion-sensitive cortical areas, we recorded cell responses to animate implied motion in macaque middle temporal (MT) and medial superior temporal (MST) cortex and investigated these areas using fMRI in humans. In the single-cell studies, we compared responses with static images of human or monkey figures walking or running left or right with responses to the same human and monkey figures standing or sitting still. We also investigated whether the view of the animate figure (facing left or right) that elicited the highest response was correlated with the preferred direction for moving random dot patterns. First, figures were presented inside the cell's receptive field. Subsequently, figures were presented at the fovea while a dynamic noise pattern was presented at the cell's receptive field location. The results show that MT neurons

# INTRODUCTION

A photograph of an object or a person in motion can evoke a perception of motion on the basis of the content of the static image in typically developed human observers (Jellema et al., 2009). Human fMRI studies have shown that static photographs depicting an object or a person in motion evoke a higher response in human middle temporal (MT) cortex and its satellites (MT+) than photographs of the same objects or persons without this implied motion (Kourtzi & Kanwisher, 2000; Senior et al., 2000). These studies suggest integration of object and motion information at a relatively early level of visual processing, which makes implied motion an interesting case for studying feedback projections of high-level multimodality cortical areas to low-level single modality areas (Kaas & Collins, 2004).

We tested whether neurons in macaque MT and medial superior temporal (MST) cortex are sensitive to implied motion as expressed by animate agents in static images. did not discriminate between figures on the basis of the implied motion content. Instead, response preferences for implied motion correlated with preferences for low-level visual features such as orientation and size. No correlation was found between the preferred view of figures implying motion and the preferred direction for moving random dot patterns. Similar findings were obtained in a smaller population of MST cortical neurons. Testing human MT+ responses with fMRI further corroborated the notion that low-level stimulus features might explain implied motion activation in human MT+. Together, these results suggest that prior human imaging studies demonstrating animate implied motion processing in area MT+ can be best explained by sensitivity for low-level features rather than sensitivity for the motion implied by animate figures. ■

These recordings allowed us to compare temporal response properties with implied and real motion from the same cell. There is evidence from human EEG and magnetoencephalography studies that implied motion responses in dorsal motion processing areas are delayed compared with responses to real motion (Fawcett, Hillebrand, & Singh, 2007; Lorteije et al., 2006, 2007), potentially resulting from high-level feedback. Cells in the anterior regions of the STS (STSa) are specialized for the perception of bodily actions and postures in macaque monkeys (Barraclough, Xiao, Oram, & Perrett, 2006; Jellema & Perrett, 2003a, 2003b; Jellema, Baker, Wicker, & Perrett, 2000). These cells typically have response latencies of about 80 to 110 msec (Barraclough et al., 2006). Neurons in areas MT have response latencies for real motion stimuli of about 50 to 80 msec (Perge, Borghuis, Bours, Lankheet, & van Wezel, 2005b; Raiguel, Xiao, Marcar, & Orban, 1999; Schmolesky et al., 1998; Lagae, Maes, Raiguel, Xiao, & Orban, 1994). If single neurons in MT show implied motion-related activity as a consequence of feedback from form-sensitive cells in STSa, then MT cells' implied motion responses may appear delayed relative to pure motion responses. In addition, because STSa neurons typically have much larger receptive fields than MT neurons (Jellema & Perrett, 2006), such a

<sup>&</sup>lt;sup>1</sup>Utrecht University, The Netherlands, <sup>2</sup>Erasmus MC, Rotterdam, The Netherlands, <sup>3</sup>University of Hull, United Kingdom, <sup>4</sup>University of St. Andrews, United Kingdom, <sup>5</sup>Wageningen University and Research Centre, The Netherlands, <sup>6</sup>University of Twente, Enschede, The Netherlands

feedback projection from STSa neurons could also induce modulation of MT cell motion responses regardless of the position of an animate agent with respect to the MT neurons' receptive field. Finally, real motion can adapt visually evoked responses to implied motion in humans in a direction specific manner (Lorteije et al., 2007). Because MT neurons are directionally selective for real motion, the question arises whether they show similar direction preference for implied motion and whether modulation of responses to one motion type by the other occurs at the level of MT.

To address the above questions, we performed extracellular single and multiunit recordings in area MT of awake macaque monkeys while presenting the animal with pictures with and without implied motion as well as control stimuli for low-level visual features. In addition, we recorded from single units in the MST cortex a motion-sensitive area that receives direct input from MT but generally has larger receptive fields and is thought to process more complex motion patterns such as optic flow and heading perception (Orban, 2008). In a second experiment, MT responses to dynamic noise in the cells' receptive field were recorded while pictures were presented at the fixation spot. This presentation allowed the animals a clear view of the images, simultaneously removing the differences in low-level features inside the receptive field between conditions. Dynamic noise typically consists of motion energy at a broad range of speeds and in all directions and will evoke activity above spontaneous level in almost all MT cells (Van Wezel & Britten, 2002; Britten & Newsome, 1998; Britten, Shadlen, Newsome, & Movshon, 1993). Thus, modulation of real motion responses by implied motion, via a putative feedback mechanism, can be tested. Finally, a human fMRI experiment was conducted to correlate the single-unit data with previous fMRI studies.

# **METHODS**

# MT/MST Recordings: Surgical and Recording Procedures

The recordings in MT and MST (Experiments 1 and 2) were conducted at the Department of Functional Neurobiology, Utrecht University, The Netherlands. Housing, surgical procedures, recording, handling, and all other procedures used in Experiments 1 and 2 were approved by the Animal Use Committee (DEC) of Utrecht University, and procedures followed national and international guidelines.

Two male rhesus macaques aged 6 and 7 years were implanted surgically with a head-holding device, a scleral search coil for measuring eye movements using the double induction technique (Malpeli, 1998; Reulen & Bakker, 1982), and a stainless-steel recording cylinder placed over a craniotomy above the left occipital lobe. For one animal, a second cylinder was placed dorsally over a craniotomy above the parietal/occipital region. All surgical procedures were performed under  $N_2O/O_2$  anesthesia supplemented with isoflurane. After recovery, the monkeys were trained to fixate a rectangular red spot  $(0.4^{\circ} \times 0.4^{\circ})$ . During experiments, each monkey sat in a primate chair 57 cm from a 19-in. monitor (refresh rate: 120 Hz in Experiment 1, 100 Hz in Experiment 2). When correctly fixating  $(\pm 1^{\circ})$ , sample rate 500 Hz), the monkey was rewarded with water or juice every second during the reverse correlation recordings and after every stimulus presentation during the implied motion recordings.

Extracellular single and multiunit recordings were carried out using standard methods as described previously (Perge, Borghuis, Bours, Lankheet, & van Wezel, 2005a). In short, a parylene-insulated tungsten microelectrode (0.5–2 M $\Omega$  impedance) was inserted manually through a guide tube and then manipulated by a micropositioning controller. Cortical areas MT and MST were identified by the recording position and depth, the transition between gray matter, white matter, and sulci along the electrode track, and by functional properties. For MT, these were the prevalence of direction-selective units, the similarity in direction tuning for nearby single-unit recordings, the receptive field size according to eccentricity, and the change of direction tuning along the electrode penetration. For MST, the functional properties were large receptive fields overlapping the fixation point and extending to the ipsilateral side and selectivity for complex motion patterns. In one monkey, the anatomical positions were confirmed with structural MRI scans of the brain containing a reference electrode inserted at the location of the recordings. Action potentials from single and multiunits were isolated with a window discriminator (BAK Electronics Inc., Mt. Airy, MD) for recordings in Monkey 1 and with online spike sorting software (ASD, Alpha Omega, Nazareth, Israel) for Monkey 2. Spike times were registered at a 0.5-msec resolution for online analysis and data storage using a Macintosh G4 computer with a National Instruments PCI 1200 data acquisition board.

# MT/MST Recordings: Search Procedure

As a search stimulus, we used either the moving random dot patterns (RDPs) or the experimental images described below. Position and size of the receptive fields were mapped by projecting a light bar on a dark monitor with a hand-held projector while the monkey was fixating. Direction tuning for real motion was established using a motion reverse correlation paradigm (Perge et al., 2005a, 2005b; Borghuis et al., 2003). A translating RDP (subtending  $8.5^{\circ} \times 11.5^{\circ}$  in Experiment 1, same size as receptive field in Experiment 2) was presented centered on the cell receptive field. The pattern switched every frame between eight possible randomly chosen directions. The delay between successive steps was 10 or 8.3 msec, with a step size of  $0.12^{\circ}$  corresponding to velocities of 12°/sec and 14.4°/sec, respectively. The preferred direction determined with this reverse correlation paradigm is strongly correlated with the preferred direction measured conventionally with hand mapping or by presenting RDPs moving in different directions for long durations (>1 sec; Borghuis et al., 2003).

# **Experiment 1 Procedure**

In Experiment 1, stimuli were color human or monkey figures implying motion facing left or right (Figure 1A and D) and standing or sitting facing forward (Figure 1C and F). In addition, images of bars that did not convey implied motion (Figure 1I and J), and scrambled images (Figure 1G and H) were presented as controls for low-level effects: orientation, position, and size. All figures subtended  $8.5^{\circ} \times$ 11.5°. Pictures were presented in the receptive field of the cells, at an average eccentricity of  $9^{\circ}$  (SD =  $2^{\circ}$ ). In human psychophysical studies, subjects are well able to discern shapes of similar size at these eccentricities (Näsänen & O'Leary, 1998). In the human pictures series, three different human agents were used (the three different persons are depicted in Figure 1A-C). The monkey pictures only showed one animal per condition (Figure 1D-F). Bars included only one type of tilted bar in two directions (Figure 1I) and one type of vertical bar (Figure 1J). All MT and MST units were tested with the same set of human figures, and 80% of the recordings also included the monkey figures and bars. The whole set of stimuli was presented in blocks, and within each block, the order was randomized. Blocks were repeated 10 to 20 times.

In Experiment 1, stimuli were presented at the center of the receptive field against a black background (Figure 2A) for 500 msec with an interstimulus interval of 250 msec (occasionally interstimulus interval up to 1000 msec). To increase the baseline firing rate of MT or MST cells during the presentation of the stimulus, thereby increasing the likelihood of finding modulating effects, we presented 18 cells in Monkey 1 with human figures flickering at 10 or 20 Hz for 500 msec. Responses of these cells to flickering and continuous stimulation were similar, and therefore these data were included in all analyses (except peristimulus time histograms) and are indicated separately in figures.

#### **Experiment 2 Procedure**

In Experiment 2, we presented foveally images of gray scale human and monkey figures implying motion (Figure 1A and D) or standing/sitting (Figure 1B and E), irrespective of the position of each cell's receptive field. Images subtended  $8.5^{\circ} \times 11.5^{\circ}$  in most recordings; however, to prevent overlap of the images with the receptive field for eight units with near foveal receptive fields, images were reduced to  $4.25^{\circ} \times 5.75^{\circ}$ . Concurrently, a dynamic RDP (500 white dots subtending  $0.2^{\circ} \times 0.2^{\circ}$ , with a limited lifetime of 10 msec, black background) was presented optimally over the receptive field (Figure 2B). Both images and RDPs were presented simultaneously for 500 msec with 250-msec intervals.

Figure 1. Humans and monkeys implying motion and sitting/standing were used in all experiments. The figures implying motion were identical in all experiments (A, D, profiled toward left or right), but the standing/sitting images could be viewed from the front, left, or right. In Experiment 1, monkey figures and human figures faced forward (C, F). In Experiment 2, both monkeys and humans sitting/standing were viewed from the left or right (B, E). To control for low-level stimulus features, we added control stimuli in Experiment 1. Bar images could be vertical to match standing images (J) or tilted left or right to match implied motion images (I). Furthermore, scrambled versions of the figures were used as control stimuli in Experiment 1 (G, H).

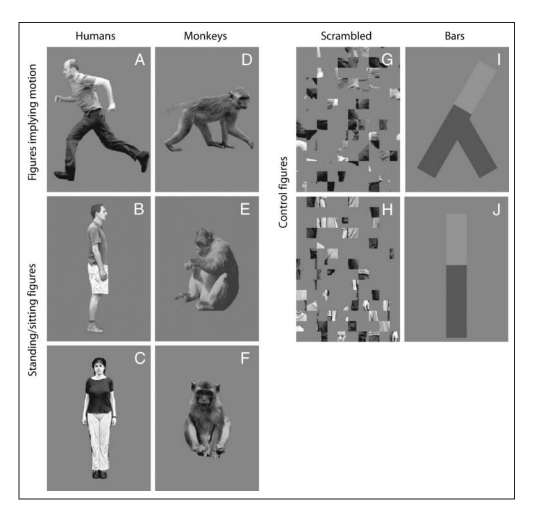
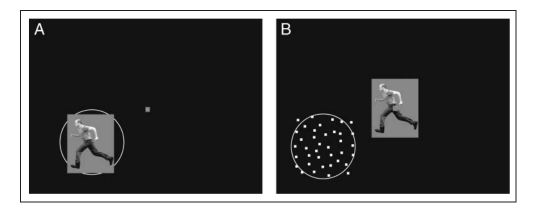


Figure 2. Schematic representation of the stimulus presentation for Experiments 1 and 2. (A) Experiment 1: Monkeys fixated a red dot (illustrated here in gray) on a black background in the middle of the monitor screen in a darkened room. The circle illustrates the receptive field of an MT cell: stimuli were presented in the middle of the receptive field. (B) Experiment 2: White dynamic dots were presented over the receptive field and static images at the fixation point simultaneously.



# MT/MST Recordings: Data Analysis

Data from the reverse correlation recordings were analyzed as described previously (Borghuis et al., 2003). In short, motion direction tuning functions were computed by reverse correlating the response to the rapid sequence of displacements of the RDP. The preferred direction was defined as the direction with maximum correlation and the nonpreferred (null) direction as the direction opposite to the preferred direction. Furthermore, for each cell, we calculated the direction index (DI):

DI = 1-(relative probability in null direction/ relative probability in preferred direction)

Response latency for the moving RDPs was established at the center of a 5-msec response window showing the highest direction-selectivity index. Cells were divided into four groups according to their direction tuning: (1) no direction selectivity, (2) tuned to upward or downward directions, (3) tuned to leftward direction ( $\pm 45^{\circ}$ ), and (4) tuned to rightward directions ( $\pm 45^{\circ}$ ). The responses of cells tuned for leftward or rightward motion were correlated with the responses to static images of figures implying motion facing left and right and bars that were tilted toward left and right.

Responses to static images in Experiments 1 and 2 were analyzed identically. For each cell, the response latency was calculated from the average response to all image conditions. Spontaneous activity and its standard deviation were calculated within a 100-msec window before stimulus onset. Response latencies to each image were calculated by determining the mean firing rate within a sliding 25-msec window (in 1-msec steps). The image response latency was established as the middle of the first window with a mean response of three standard deviations above the spontaneous activity. For some cells with smaller responses (10 MT and 12 MST units in Experiment 1 and 2 MT units Experiment 2), latency was calculated when the mean response was one or two standard deviations above spontaneous activity. For 4 MT units and 7 MST units in Experiment 1 and 1 MT unit in Experiment 2, with very small responses, latencies were set at the respective average MT or MST latency. Mean responses to each image were calculated within a 500-msec period after the respective response latency.

Responses of each cell to each stimulus were analyzed by applying a receiver operating characteristic (ROC) analysis (Britten, Newsome, Shadlen, Celebrini, & Movshon, 1996; Green & Swets, 1966). This analysis does not make any assumptions about the response distributions, and the ROC value expresses the amount of overlap of two response distributions for two conditions (where an ROC value = 0.5 means complete overlap and ROC value = 0 or 1.0means no overlap of the response distributions). In our experiments, by definition ROC values larger than 0.5 signified stronger responses to implied motion stimuli. Furthermore, the ROC analysis allowed us to assess whether an ROC value is significantly different from 0.5 for each individual unit by performing a permutation test (n = 1000; Britten et al., 1996; Elfron & Tibshirani, 1993). Wilcoxon signed rank tests were used to test the differences in responses for different conditions across the population of units.

# Human fMRI: Subjects, Scanning Protocol, and Task Design

Eight healthy subjects (five men and three women, average age = 24.8 years, SD = 6.1 years) who were recruited from the staff and students of Utrecht University participated in the experiment. All subjects gave informed consent for participation (approved by the Human Ethics Committee of the University Medical Center Utrecht).

All images were obtained with a Philips Achieva 3-T MRI scanner (Philips Medical Systems, Best, the Netherlands) with a Quasar Dual gradient set. Functional scans were acquired in sagittal orientation using a navigated 3D-PRESTO pulse sequence (Ramsey et al., 1998; van Gelderen et al., 1995) with the following parameters: repetition time =

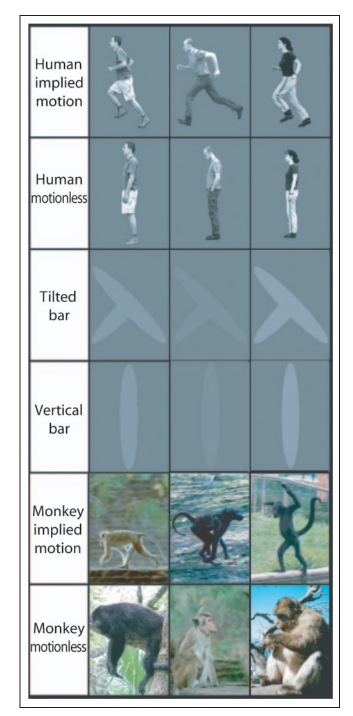
21.75 msec (time between two subsequent RF pulses), effective echo time = 32.4 msec, field of view (anterior-posterior, inferior-superior, and right-left) =  $224 \times 256 \times 128$  mm, flip angle =  $10^{\circ}$ , matrix =  $56 \times 64 \times 32$  slices, voxel size = 4 mm isotropic, eight channel head coil, SENSE factors = 2.0 (left-right) and 1.8 (anterior-posterior). The total acquisition time per volume was 500.3 msec. A T1-weighted structural image was acquired after functional scanning. A PC, a rear projection screen, and a video projector were used for task presentation. All stimuli were projected on a gray background. All events were time locked to the fMRI scans.

Motion-sensitive areas were mapped by presenting RDPs intermittently for 1000 msec, with a 300-msec interval. RDPs alternated between moving left or right (10°/sec) and remaining stationary. Each block lasted 26 sec, and there were 18 blocks in total. The three main image types were human pictures, bars, and monkey pictures. Images of humans implying motion (running) and without implied motion (standing) were generated for three different actors when facing both left and right (Figure 3, top panels). All actors were seen on a gray background with a height of 350 pixels, subtending 13.2° visual angle. Images of bars consisted of three different vertical bars, subtending 13.2° that resembled a standing human actor, and three different combinations of two tilted bars that resembled either a human running left or right (Figure 3, middle panels). The relative luminance of all images (humans and bars) was matched. Images of monkeys implying motion (n = 40) and not implying motion (n = 40) consisted of digitized color photographs (250  $\times$  350 pixels, 9.4°  $\times$  13.2° visual angle) of monkeys in their natural surroundings (Figure 3, bottom panels).

The experimental design that we used was similar to the study of Kourtzi and Kanwisher (2000), except that we used stimuli similar to those used during the electrophysiological recordings and bar stimuli controlling for low-level features, as described above. The fMRI experiment consisted of blocks of seven image types, including monkey and human figures implying and not implying motion and tilted and vertical oval bars. One subject was not presented with the tilted and vertical bars. A new picture was presented every 1000 msec (two scans) for 300 msec (total 20 images/block). In addition, there was a rest condition in which only the red fixation dot was displayed on the screen for 20 sec. There were three scanning sessions containing 21 blocks (3 blocks per condition). During the experiment, subjects were required to fixate a central red dot throughout all stimulus presentations. After the experiment, participants reported that they recognized the presence or the absence of implied motion in the human and monkey pictures, although they were not informed regarding the exact aim of the study nor received explanation about implied motion beforehand.

# Human fMRI: Analysis

The fMRI time series data were preprocessed using SPM2 (http://www.fil.ion.ucl.ac.uk/spm/spm2.html). Preprocess-



**Figure 3.** Pictures containing human figures with and without implied motion, tilted and vertical bars, and pictures of monkeys with and without implied motion used in the fMRI experiment. Human pictures and tilted bars were viewed from both left and right. Three examples of the monkey pictures with and without implied motion are illustrated.

ing steps included realignment, coregistration, normalization, and smoothing (8 mm FWHM). Statistical analysis of fMRI scans was performed with custom-written programs in IDL (Research Systems Inc., Boulder, CO). Data for each subject were submitted to a linear multiple regression analysis. The design matrix for the implied motion experiment contained six factors for stimulus-related changes in BOLD signal during the six conditions in which pictures were shown (the rest condition was used as a reference). All events in the design matrix were convolved with a predefined hemodynamic response function (Friston, Frith, Turner, & Frackowiak, 1995).

MT+ was defined in each subject by the contrast in activation between the presentation of static and moving random patterns in the ascending limb of the inferior temporal sulcus (p < .05, Bonferroni corrected). Subsequently, we calculated the average regression coefficients for the six conditions of the implied motion experiment within area MT+. Differences between the implied and motionless stimuli were tested using a paired samples *t* test.

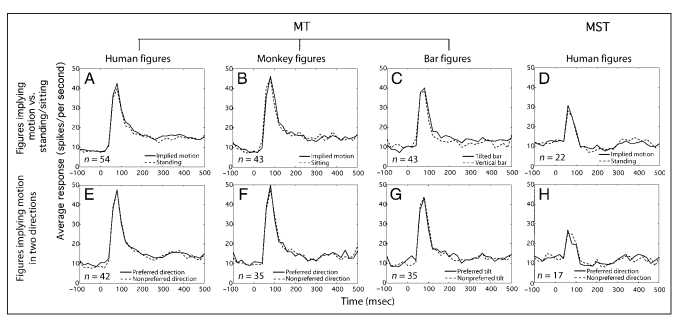
# RESULTS

# **Direction Selectivity**

In total, 68 MT and 26 MST single and multiunits were recorded from two monkeys in Experiment 1 and 69 MT cells from one monkey in Experiment 2. Mean MT unit receptive field eccentricity was 9.6° ( $SD = 4.6^{\circ}$ ) with an average diameter of 8.2° ( $SD = 2.9^{\circ}$ ). MST cells had much larger receptive fields that were often hard to measure precisely, often extending to ipsilateral locations. Direction selectivity for real motion was obtained with the motion reverse correlation paradigm (Borghuis et al., 2003) using eight motion directions, and a DI was calculated. The mean DI of all recorded MT units was 0.4 (SD = 0.3), with an average peak response latency of 59.1 msec (SD = 18.2 msec); the mean MST DI was 0.7 (SD = 0.2), with an average peak response latency of 71.9 msec (SD = 13.4 msec). The DI for MT cells seems low compared with those obtained in previous studies (Snowden, Treue, & Andersen, 1992; Mikami, Newsome, & Wurtz, 1986). These studies varied the speed of their moving stimuli. A huge portion of MT neurons is speed selective, and DI values can be lower at nonoptimal speed (Mikami et al., 1986). Because of time restraints, we tested cells with only one speed  $(12^\circ/\text{sec or } 14.4^\circ/\text{sec})$  to quickly obtain the preferred direction and to continue with the implied motion experiments. These nonoptimized presentation speed would cause a lower DI, but they are in the range reported earlier for MT (Perge et al., 2005a, 2005b; Borghuis et al., 2003).

# Experiment 1: MT/MST Cell Responses to Implied Motion within the Receptive Field

In Experiment 1, we recorded 40 MT single units and 14 MT multiunits as well as 13 MST single units and 9 MST multiunits, with a nonflickering stimulus presentation. Other cells were tested with flickering stimuli (6 MT single units, 8 MT multiunits, and 4 MST single units). Stimuli were presented at the center of the receptive field where the visual responses and motion selectivity were strongest. The average eccentricity of the stimulus was  $9^{\circ}$  ( $SD = 2^{\circ}$ ).



**Figure 4.** Peristimulus time histograms for combined MT and MST responses in Experiment 1 for the nonflickering condition. The response of each neuron was segmented into 20-msec bins. Responses were not normalized to the average firing rate, emphasizing responses from units with high firing rates. Population peristimulus time histograms after normalization to the average response showed similar results (data not shown). The upper panels indicate the responses to figures (A, D, humans; B monkeys; C bars), where figures imply motion (solid lines) or do not imply motion (dashed lines). The lower panels indicate the response to figures (E, H, humans; F monkeys; G bars), where implied motion is in the preferred direction (solid lines) or in the nonpreferred direction (dashed line). Stimulus onset was at 0 msec, and duration for all cells was at least 500 msec.

**Table 1.** MT and MST Cell Numbers in Experiment 1 and Their Responses to Figures Implying and Not Implying Motion or Bar Tilt Are Shown by the Average Firing Rates and the Average ROC Values for the Whole Population and for Cells with a Significant Selectivity for Conditions as Established in the ROC Analysis

Area	Figures	No. Cells	Average Firing Rate Implied Motion/ Standing–Sitting	Average ROC Values	% Significant Cells for Implied vs. Standing–Sitting	Average ROC Value for Significant Cells	Average Firing Rate Implied/ Standing–Sitting for Significant Cells
МТ	Humans	54	$1.29 \pm 1.61$	$0.53 \pm 0.09$	25.9	$0.59 \pm 0.12$	$1.28 \pm 0.3$
MST	Humans flicker	14	$0.95 \pm 0.24$	$0.52 \pm 0.12$	14.3	$0.54 \pm 0.3$	$0.99 \pm 0.16$
	Monkeys	43	$1.44 \pm 2.61$	$0.48 \pm 0.15$	25.3	$0.50 \pm 0.28$	$1.13 \pm 0.65$
	Bars	43	$1.23 \pm 0.57$	$0.6\pm0.14$	23.3	$0.7 \pm 0.13$	$1.68 \pm 0.74$
	Humans	22	$1.00 \pm 0.2$	$0.51 \pm 0.01$	9.1	$0.5 \pm 0.19$	$0.96 \pm 0.41$
	Humans flicker	4	$1.18 \pm 0.11$	$0.56 \pm 0.04$	0		_
	Monkeys	12	$1.11 \pm 0.32$	$0.48\pm0.1$	25.0	$0.46 \pm 0.22$	$1.25 \pm 0.62$
	Bars	12	$1.06 \pm 0.37$	$0.5\pm0.1$	8.3	$0.67 \ (n = 1)$	2.03 $(n = 1)$

Variance is indicated with standard deviation.

The results illustrated in Figure 4 show that there are no clear differences between the responses to figures implying motion and the responses to figures not implying motion in either MT (Figure 4A–C) or MST (Figure 4D) cells (see also Table 1). For units with a DI of at least 0.1 and a preferred direction along the horizontal axis ( $\pm 45^{\circ}$ ) in the motion reverse correlation test, direction selectivity for implied motion was tested. No difference was found when comparing responses with figures implying motion (or tilted bars) when seen from views congruent or incongruent with the preferred motion direction (Figure 4E–H, see also Table 2).

The peristimulus time histograms in Figure 4 illustrate a transient response with an average peak response latency

of 65.0 msec (SD = 16.6 msec) for MT units (n = 54) and 64.6 msec (SD = 18.9 msec) for MST units (n = 22) when presented with nonflickering stimuli. After this peak, average MT responses were sustained and higher than their spontaneous activity until the end of the stimulus presentation. MST responses dropped to or below spontaneous activity. The mean spontaneous firing rate of MT neurons was 7.9 spikes per second (SD = 7.1 spikes per second), and the mean sustained (200–400 msec after stimulus onset) firing rate was 15.1 spikes per second (SD = 16.8 spikes per second). For MST neurons, mean spontaneous activity was 11.0 spikes per second (SD = 6.5 spikes per second), whereas the mean sustained

Table 2. For MT and MST Cells in Experiment 1 with a Horizontal Direction Preference to Real Motion, the Number of Involved
Cells and Their Responses to Figures Implying Motion or Bar Tilt Are Shown by the Average Firing Rates and the Average ROC Values
for the Whole Population and for Cells with a Significant Selectivity for Particular Conditions as Established in the ROC analysis

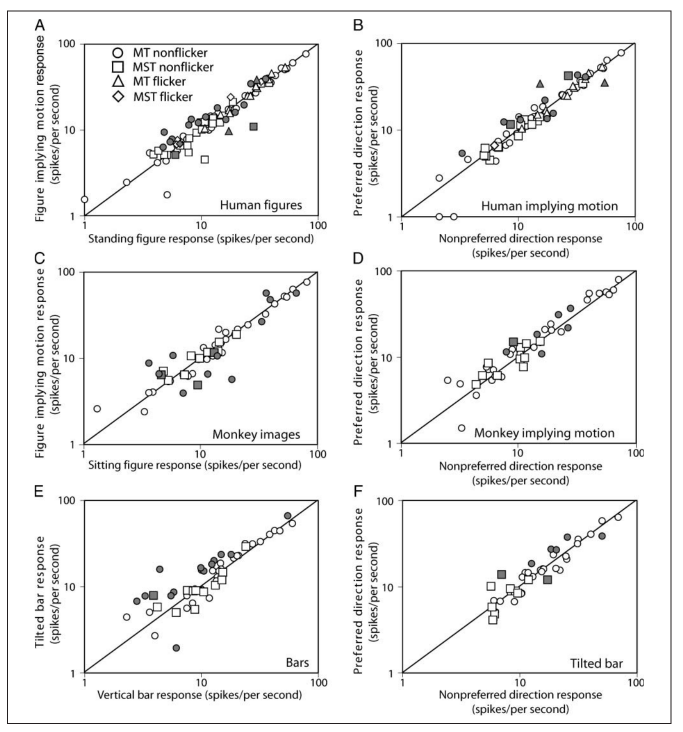
Area	Figures	No. Cells	Average Firing Rate Preferred/ Nonpreferred	Average ROC Values	% Significant Cells for Preferred vs. Nonpreferred	Average ROC Value for Significant Cells	Average Firing Rate Preferred/ Nonpreferred for Significant Cells
MT	Humans	42	$1.08 \pm 0.35$	$0.51 \pm 0.09$	14	$0.57 \pm 0.18$	$1.23 \pm 0.34$
	Humans flicker	12	$1.15 \pm 0.46$	$0.49 \pm 0.15$	17	$0.56 \pm 0.44$	$1.06 \pm 0.37$
	Monkeys	35	$1.09 \pm 0.33$	$0.56 \pm 0.13$	17	$0.60 \pm 0.24$	$1.14 \pm 0.31$
	Bars	35	$1.46 \pm 2.89$	$0.51 \pm 0.13$	14	$0.65 \pm 0.23$	$1.30 \pm 0.31$
MST	Humans	17	$0.98 \pm 0.32$	$0.53 \pm 0.11$	17	$0.73 \pm 0.07$	$0.98 \pm 0.85$
	Humans flicker	1	$1.03 \ (n = 1)$	$0.54 \ (n = 1)$	0	_	
	Monkeys	11	$1.11 \pm 0.29$	$0.51 \pm 0.12$	9	0.77 (m = 1)	1.63 (n = 1)
	Bars	11	$1.09 \pm 0.42$	$0.47 \pm 0.15$	29	$0.56 \pm 0.36$	$1.33 \pm 0.9$

Variance is indicated with standard deviation.

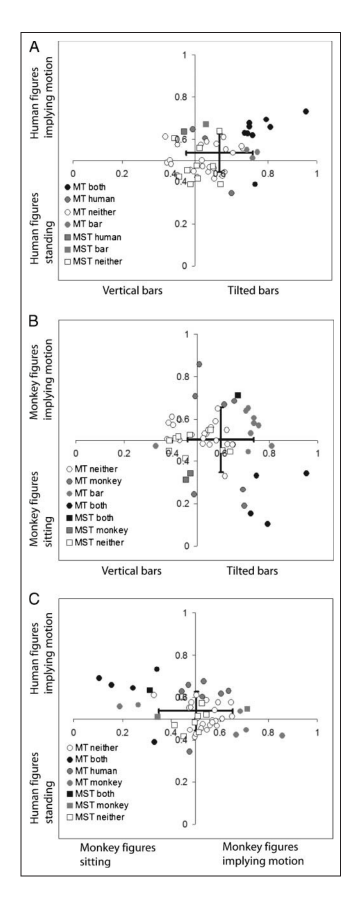
response was 10.0 spikes per second (SD = 7.8 spikes per second). Spontaneous and sustained firing rates were compared using nonparametric tests and differed significantly for MT neurons (Wilcoxon signed rank test, p < .001) but not for MST neurons (Wilcoxon signed rank test, p > .05).

# **Experiment 1: Cell-by-Cell Analysis**

Responses to every condition were averaged for all cells and plotted against each other (Figure 5, see Supplemental Figure 1 for histograms of response ratios). Response preferences of the MT and MST neuronal population were



**Figure 5.** Comparison of mean MT and MST unit responses in Experiment 1. MT responses to nonflickering images are indicated by dots and flickering stimuli by triangles. Diamonds and squares indicate MST responses. The diagonal line is the line of unity when the responses to both types of images are equal. Note that cells with a lower spike rate are more susceptible to noise (e.g., spike burst in 1 trial), and ratios between pictures for cells in the lower left corner are therefore more variable. For each cell, the differences in responses were tested for significance using an ROC analysis with a permutation test. Cells with significantly different responses are indicated by filled symbols.

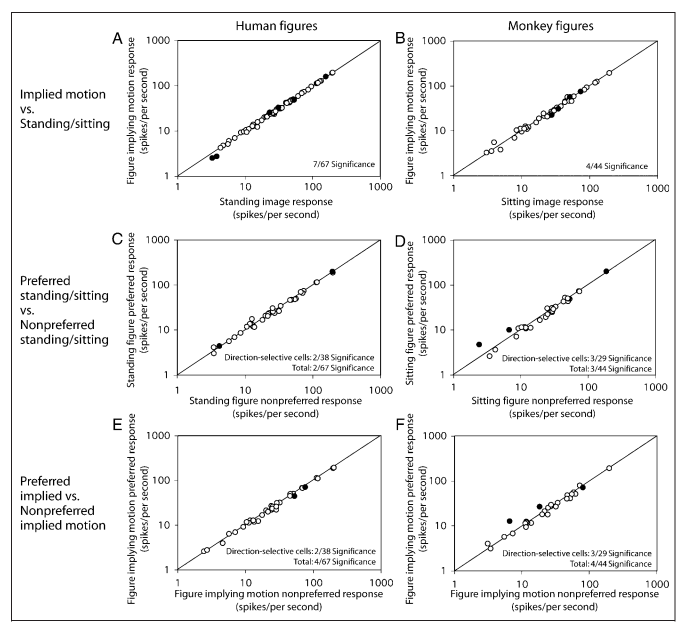


tested with a nonparametric paired test (Wilcoxon signed rank test, p < .05 is considered significant). No significant difference was found between responses to figures implying motion and sitting/standing figures, neither for the human figures (continuously presented or flickering; Figure 5A) nor for the monkey figures (Figure 5C). No difference was found between responses to tilted and vertical bars (Figure 5E). In addition, no significant differences were found in responses to implied motion figures facing the preferred motion direction versus facing the nonpreferred motion direction for both human (Figure 5B) and monkey (Figure 5D) figures or tilted bars (Figure 5F). Comparing response magnitudes of MST neurons in the nonflickering condition (indicated in Figure 5 with square symbols) between figures implying motion and sitting/standing, tilted and vertical bars as well as preferred versus nonpreferred direction did not reveal any significant differences. Average responses to scrambled images versus all unscrambled images for 49 MT and 26 MST units also did not differ significantly (data not shown).

Although we did not find evidence for implied motion processing in areas MT and MST in the population data, ROC analysis of units separately is necessary to exclude the possibility that a subset of neurons is modulated by motion implied by the figures (Table 1). Neurons with a significant ROC value are marked with filled symbols in Figures 5 and 6. Under our definitions, an ROC value > 0.5means a higher response to figures implying motion versus standing figures. Because the average ROC values for the human figures were above 0.5 for MT cells (but equal to 0.5 in MST cells), one might conclude that this small subset of MT neurons was significantly more responsive in the expected direction. However, as will be looked at later, we think this is due to low-level effects, as the response to bar figures shows a similar direction vet larger amplitude. Furthermore, the preference for implied motion stimuli was not present for the monkey images in MT cells and was even opposite in MST cells.

We compared eccentricities of the receptive field for MT cells with significant ROC values versus MT cells with insignificant ROC values in independent *t* tests for the human, monkey, and bar conditions separately. Eccentricity was not significantly different between the two groups (*p* values > .05).

**Figure 6.** A comparison of ROC values for responses to figures of humans and monkeys and tilted bars in Experiment 1. (A) As selectivity for human figures implying motion versus figures of standing humans increased, so did selectivity for tilted bars versus vertical bars. Cells showing significantly different responses for either the human figures or the bar figures are indicated with filled symbols. (B) No clear trend is apparent for ROC values for response preferences to monkey figures and bars. (C) Implied motion preference for human figures and monkey figures was not consistent; five cells with a significant preference for human implied motion even had a significant preference for figures of sitting monkeys. The mean ROC values with standard deviations of all MT cells are shown as black bars.



**Figure 7.** Comparison of MT unit activity induced by dynamic noise onset in the receptive field and static images at the fovea during Experiment 2. Each data point represents average responses from one MT unit. (A) Responses to a combination of dynamic noise with human figures either running or standing. (B) Same for monkey figures either walking or sitting. (C) Responses to the combined presentation of dynamic noise with figures of standing or sitting humans and (D) monkeys facing preferred and nonpreferred direction. (E) Responses to simultaneous presentation of dynamic noise with figures of humans and (F) monkeys running or walking in preferred and nonpreferred direction. Significant differences were detected using an ROC analysis with a permutation test (filled cells). The numbers in the right lower corner of each graph indicate the number of significant cells out of the units that were horizontally directionally selective for real motion (not for A and B) and the number of significant units out of the number of total units that were tested for each condition regardless of real motion direction preference.

In our total MT population, 26% (14/54) of the units had significantly different responses for implied motion versus standing human images in the nonflickering condition. For this subgroup of cells, the ratio of the average response to humans implying motion versus standing human images was, on average, 1.3, corresponding to a difference of 2.2 spikes per second. Response ratios and average response differences for figures with monkeys implying motion versus standing or skewed versus vertical bars for these significant cells were 1.13 with 0.19 spikes per second and 1.68 with 5.55 spikes per second, respectively. This indicated that although a quarter of the MT cells responded significantly more to either figures implying motion or figures standing or sitting, the magnitude of this difference was very small. In MST, the percentages of significant cells were lower than in MT, except for the monkey condition; again, the absolute differences in average firing rate were very small.

We tested whether cells with a selectivity for random dot motion in either left or right direction had a preferences for

**Table 3.** MT Cell Numbers in Experiment 2 and the Modulation of Their Response to Figures Implying and Not Implying Motion or Bar Tilt Are Shown by the Average Firing Rates and the Average ROC Values for the Whole Population and for Cells with a Significant Selectivity for Particular Conditions as Established in the ROC Analysis

Figures	Figures	No. Cells	Ratio of Average Firing Rate Implied Motion/ Standing–Sitting or Preferred/ Nonpreferred	Average ROC Values	% Significant Cells for Preferred vs. Nonpreferred	Average ROC Value for Significant Cells	Ratio of Average Firing Rate Implied/ Standing–Sitting or Preferred/ Nonpreferred for Significant Cells
Implied motion vs.	Humans	67	$0.99 \pm 0.07$	$0.49 \pm 0.05$	10	$0.46 \pm 0.12$	$0.93 \pm 0.15$
standing/sitting	Monkeys	44	$1.01 \pm 0.12$	$0.49 \pm 0.05$	9	$0.40 \pm 0.16$	$0.97 \pm 0.13$
Standing/sitting	Humans	38	$1.02 \pm 0.11$	$0.51 \pm 0.08$	5	$0.70 \pm 0.03$	$1.05 \pm 0.02$
(face direction)	Monkeys	29	$1.01 \pm 0.23$	$0.51 \pm 0.12$	10	$0.58 \pm 0.30$	$1.54 \pm 0.46$
Implied motion	Humans	38	$1.00 \pm 0.08$	$0.48 \pm 0.06$	5	$0.36 \pm 0.01$	$0.90 \pm 0.06$
	Monkeys	29	$1.03 \pm 0.24$	$0.51 \pm 0.12$	10	$0.59 \pm 0.27$	$1.45 \pm 0.54$

Variance is indicated with standard deviation.

implied motion (humans and monkeys) and tilt direction (bars) in the same direction (Table 2 and Figure 5B, D, and F). A small number of cells exhibited a significant preference for implied motion but also tilt in the same direction as their preferred random dot motion direction, although again with only a small increase in spike rate. In addition, 11% (10/94) of all cells recorded in Experiment 1 were selective for upward or downward motion but were also significantly selective for implied motion human figures facing left or right (4 MT cells), in flickering human figures (1 MT cell and 1 MST cell), in monkey figures (2 MT cells), or for tilt direction of bars figures (2 MT cells). These cells had a significant preference for implied motion in a specific direction, which could not be explained by the preferred direction for real motion.

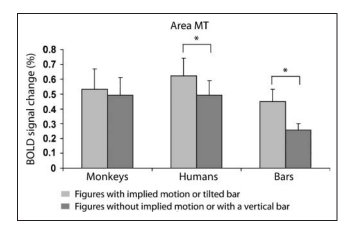
We further examined whether MT neurons that were selective for tilted bars versus vertical bars preferred implied motion versus standing/sitting images by comparing the ROC values (Figure 6). Units with a preference for implied motion often preferred tilted bars, as is indicated by the relatively high number of cells (23) in the upper right quadrant, compared with the lower right quadrant (8) in Figure 6A. No trend was visible for monkeys with implied motion versus bar figures (Figure 6B). For human versus monkey figures implying motion, units with a significant preference for human figures implying motion tended to have a significant preference for monkey figures without implied motion, although units with a significant preference for either monkey or human figures were much more scattered (see Figure 6C). Indeed, regression analysis for these data revealed a significant linear correlation between ROC values for human figures and bar stimuli (R = .64, ANOVA p value = .000), but not for human and monkeys or bars and monkeys. Together, these results indicate that preferences for human figures

implying motion versus standing might be caused by the same low-level stimulus features that were responsible for the tilted bar preference but that may be lacking in the monkey figures.

# Experiment 2: Can MT Responses to Dynamic Noise Patterns Be Modulated by Implied Motion?

We recorded the responses from 69 MT units (43 single units and 26 multiunits) to dynamic noise in the receptive field when static figures were presented at the fovea. We plotted the responses to figures implying motion and figures standing/sitting (Figure 7). We found no significant differences between average responses (Wilcoxon *p* values > .05) to dynamic noise when foveal figures implied motion or foveal figures were standing/sitting (Figure 7A and B) or figures implied motion while facing the preferred motion direction versus facing the nonpreferred motion direction (Figure 7E and F).

ROC values were calculated for all units for figures implying motion versus standing/sitting figures (top rows in Table 3, filled symbols in Figure 7). Roughly 10% of MT cells showed significant ROC values, which could be favorable for either figure type, but with an average preference for the standing/sitting figures (i.e., mean ROC value below 0.5). To quantify what these ROC values mean in terms of firing rate, we calculated the ratio of average response rates to figures with versus without motion. For human figures, this ratio was 0.95, which corresponded to a difference in spike rate of 0.17 spikes per second in favor of the sitting/standing figures. For monkey pictures, the ratio was 0.97, which corresponded to a difference of 0.03 spikes per second in favor of the figures implying motion.



**Figure 8.** Average signal in MT+ during the six conditions relative to passive fixation. Bars indicate standard errors, and asterisks show significant differences (p < .05).

Cells showing preferential responses to horizontal motion were also tested for the difference in response to implied motion figures facing in the preferred and nonpreferred motion direction (bottom rows in Table 3, Figure 7E and F). Few cells had a significant preference for implied motion, and the preferred direction was not consistent over cells. The same result was obtained for comparison of standing/sitting figures facing in the preferred or nonpreferred direction for real motion (middle rows in Table 3, Figure 7C and D).

# Human fMRI Experiment

Robust MT+ activation was detected in seven subjects during the motion-mapping task. For human as well as abstract pictures, there was more activation in MT+ when viewing implied motion pictures than when viewing pictures without implied motion (see Figure 8): human pictures, t(6) = 4.65, p < .005; abstract pictures, t(5) = 4.34, p < .01. The comparison of activation in MT+ between pictures of monkeys implying and not implying motion revealed no differences, t(6) = 1.218, p > .05.

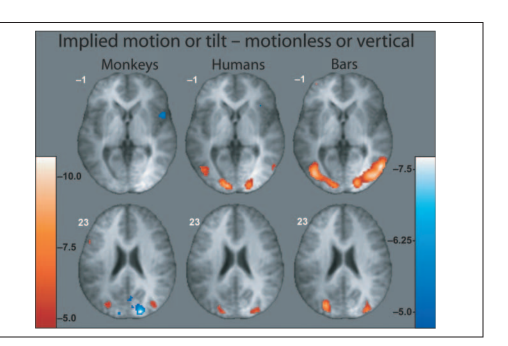
A voxel-based comparison between images of humans and abstract figures implying motion and stationary figures revealed increased activation during observation of implied motion stimuli in several areas, including Brodmann areas 17 and 18 in early visual cortex, MT+ in the MT/occipital region, and parts of the inferior and superior parietal lobe. Differences in activation between images of monkeys implying motion and stationary monkeys were nearly entirely restricted to areas in the parietal lobe (Figure 9; Talairach x, y, and z coordinates: 42, -79, 22 for the left parietal lobe and -30, -79, 91 for the right parietal lobe). The difference in signals to images of humans implying motion and stationary humans in this area was not significant, t(7) = 0.81, p > .05 (Figure 10); however, the difference in signals to tilted and vertical bars was significant, t(6) =2.82, p < .05.

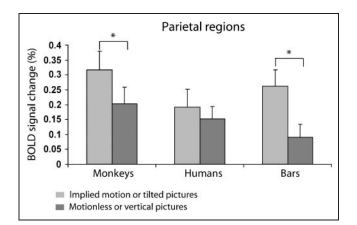
# DISCUSSION

In this study, we measured implied motion processing in cortical areas MT and MST of macaque monkeys and in human MT+. Modulatory input about the form of a moving (in)animate object could, in principle, be integrated in neurons processing real motion. This type of integration of different modalities could be formed by association (Schlack & Albright, 2007), as brief glimpses of moving subjects are often accompanied by actual motion.

There are several reasons why modulatory effects in areas MT and MST can be expected. First, previous human fMRI

**Figure 9.** Results of the contrast between implied and motionless stimuli for monkey, human, and abstract pictures. Significant voxels were thresholded at p < .05 (corrected) and superimposed on the averaged anatomical scan. MNI *z*-coordinates are displayed in white on the top left of each slice.





**Figure 10.** Average signal during the six conditions relative to passive fixation in parietal areas that were significantly more active during implied motion pictures of monkeys than pictures of motionless monkeys. Conventions as for Figure 8.

experiments have shown significantly different BOLD activations in the MT+ when comparing responses to images of humans, animals, or objects moving versus images of humans, animals, or objects standing still (Kourtzi & Kanwisher, 2000; Senior et al., 2000). Second, single-cell recordings in monkeys have shown neurons in the STS respond selectively to different degrees of implied motion (Barraclough et al., 2006; Jellema & Perrett, 2006) and could provide input to MT. Third, responses evoked by implied motion in a human EEG study could be modulated by preadaptation with real motion, which indicates the presence of neurons that are (directionally) sensitive to both types of motion (Lorteije et al., 2007). Neurons in MT and MST are adaptable to real motion and typically directionally sensitive and thus may be responsible for this adaptation effect. Although TMS over MT/MST fails to impair judgments of implied motion within 150 msec of stimulus presentation (Alford, van Donkelaar, Dassonville, & Marrocco, 2007), in an earlier TMS study, MT/MST has shown to be functionally necessary for representational momentum processing (Senior, Ward, & David, 2002). Fourth, modulatory input to MT and MST has been shown in many macaque singlecell studies, for example, modulation by working memory (Schlack & Albright, 2007; Zaksas & Pasternak, 2005; Bisley, Zaksas, Droll, & Pasternak, 2004) and attention (Maunsell & Treue, 2006; Cook & Maunsell, 2002, 2004; Martínez-Trujillo & Treue, 2002; Recanzone & Wurtz, 2000; Treue & Martínez-Trujillo, 1999; Treue & Maunsell, 1996, 1999), and thus any implied motion signal might also modulate MT neuronal activity. In addition, motion implied by Glass patterns has been shown to evoke activity in both human and monkey MT and MST (Krekelberg, Vatakis, & Kourtzi, 2005; Krekelberg, Dannenberg, Hoffmann, Bremmer, & Ross, 2003).

In our study, however, we did not find evidence for implied motion processing in macaque areas MT and MST. At the population level, cells did not respond differently to human or monkey figures with and without implied

motion. A quarter of the cells in the MT population (less in MST, but with a smaller sample size) showed a small but significantly different response to images of humans running and standing. Surprisingly, the cells that were significantly selective for humans implying motion also tended to have larger differences in responses to the vertical and tilted bar figures, suggesting that the effects that were found may be attributable to low-level orientation and size differences in the images. In addition, preferences for human and monkey implied motion were not consistent: Cells selective for humans implying motion appeared to prefer images of stationary monkeys. These results contrast human fMRI results (Kourtzi & Kanwisher, 2000) that show activation within area MT+ by implied motion expressed by both human and animal agents, and this suggests that the response to implied motion is not a response to the implied motion content of the images but could reflect responses to low-level visual features. Our fMRI results support this hypothesis.

For a long time, we have known that MT neurons can respond selectively to static bars (e.g., Albright, 1984). Our results show that in addition to a transient response to stimulus onset, MT neurons have highly significant sustained responses above spontaneous activity for the static images. As MST neurons did not show this sustained response, small eye movements causing motion on the retina probably do not explain this phenomenon. In an fMRI study in monkeys, it has been shown that area MT (but not MST) has a differential response to static images compared with fixation only (Nelissen, Vanduffel, & Orban, 2006). It is thus surprising that we do not find large differences in responses to the different images. The small differences that we find can probably be attributed to low-level visual attributes like orientation and position.

In the second experiment, we found a small percentage (about 10%) of cells whose dynamic dot pattern response could be selectively modulated by static test images at the fovea. The change in firing rate, however, was so small that we question whether an implied motion signal is fed back onto area MT from position and size-invariant STSa neurons coding implied motion. These weak activations and low cell numbers can probably not explain the BOLD activity in MT+ found with human fMRI studies (Kourtzi & Kanwisher, 2000; Senior et al., 2000). Furthermore, the few cells showing differentially sensitive responses had ROC values below 0.5, indicating a preference for images without implied motion rather than images implying motion.

Our human fMRI experiment also strongly indicated that low-level differences in the images (partly) causes differences in responses to images with versus without implied motion because the bar stimuli gave an even stronger response difference than the human stimuli. More importantly, there was no difference in MT+ activity when observing the large set of images of monkeys implying motion and not implying motion. These images all differed considerably in low-level visual features that, on average, there is no consistent visual difference between pictures with and without implied motion, except the presence of implied motion itself.

A direct comparison between human fMRI experiments and single-unit recordings in the monkey is difficult because it has been shown that BOLD signal correlates more with local field potentials (incoming signals) than direct neural activation (Logothetis & Pfeuffer, 2004). Although human homologues of macaque areas MT and MST have been proposed (Goossens, Dukelow, Menon, Vilis, & van den Berg, 2006; Huk, Dougherty, & Heeger, 2002; Dukelow et al., 2001; Peuskens, Sunaert, Dupont, Van Hecke, & Orban, 2001; Morrone et al., 2000), there are also several clear differences in the function and location of motion and objectsensitive cortical areas in humans and monkeys (Nelissen et al., 2006; Orban et al., 2003; Vanduffel et al., 2001). Evidence from fMRI in macaque monkeys suggests that combining object and motion information may mainly occur in areas fundus of the superior temporal region (FST) and a newly defined area lower superior temporal region (LST) (Nelissen et al., 2006). FST and LST would be strong candidate areas for processing implied motion. Because in the monkey these areas are located near area MT, in the human they could be part of MT and its satellites, where implied motion activation was demonstrated (Kourtzi & Kanwisher, 2000; Senior et al., 2000).

STSa contains neurons sensitive to animate implied motion (Barraclough et al., 2006; Jellema & Perrett, 2006). Many cells in this area will also respond to moving random dots (Barraclough et al., unpublished observations). The properties of STSa cells, however, are very diverse and complex; about 60% of those cells that respond to static images of human figures are sensitive to the degree of articulation shown by the human. Of STSa cells sensitive to images of human figures, half prefer implied motion, whereas the other half prefer standing/sitting images (Barraclough et al., 2006). On the basis of these results, one would not necessarily expect a differential response in the human homologue of STSa for implied motion versus standing/sitting images in an fMRI experiment. Furthermore, STSa cell responses can depend on the view of the human or monkey figure. Although cells selective for static images of articulated bodies were more likely to respond to movies of bodies walking forward (Barraclough et al., 2006), typically there is no correlation between responses to implied motion and moving RDPs (Barraclough et al., unpublished observations).

Several arguments could be given as to why we were not able to show evidence for implied motion activity in areas MT and MST. First, although our search stimulus consisted of both moving RDPs and the static images that we used in this study, we cannot completely exclude the possibility that our sample was biased because of search strategies. Also, we may not have sampled neurons from every subregion of MT and MST. However, we searched extensively for implied motion-responsive neurons because we varied recording position and depth to cover different locations within MT and MST, and we recorded in two different animals and for one animal in both hemispheres. Second, it could be argued that our images did not convey a strong signal for implied motion that monkeys were not able to recognize or interpret these images or that the monkeys were not paying attention to the images because their task was only to fixate the fixation dot. These suggestions are unlikely as similar images during the same fixation task can evoke implied motion-selective responses within the STSa of macaques (Barraclough et al., 2006). Furthermore, the same images have been shown to elicit differential responses in human-evoked potential recordings (Lorteije et al., 2006) arising from direction-selective motion-sensitive neurons in dorsal cortex (Lorteije et al., 2007), and the images are very similar to the type of images that were used in the human fMRI study of Kourtzi and Kanwisher (2000). Furthermore, there is a plethora of studies showing that Rhesus macaque monkeys can interpret the content of images depicted on a computer screen as observed from their behavior, and many studies have shown differential neural activation in object recognition areas for different images in monkey fMRI or single-unit recording studies (Kourtzi, Krekelberg, & van Wezel, 2008).

Our results show that MT cells may respond selectively to low-level stimulus features within images with and without implied motion. However, although STSa cells have been shown to respond strongly to the implied motion content within an image, MT cells (and we have shown evidence indicating that this may hold for MST cells too) are insensitive to motion implied by biological forms.

# Acknowledgments

The authors thank Bert van den Berg and Wim van de Grind for their contributions to this project. The project was supported by grants awarded to R. v. W.: Innovational Research Incentives Scheme (VIDI) of the Netherlands Organization for Scientific Research (NWO), the Interuniversity Attraction Poles Programme (IUAP) of the Belgian Science Policy and a High Potential grant of the Utrecht University (UU). This work was further supported by grants from the European Union and the Wellcome Trust awarded to D. P. and M. O.

Reprint requests should be sent to Richard J. A. van Wezel, Division of Pharmacology, Utrecht University, Sorbonnelaan 16, 3584 CL Utrecht, The Netherlands, or via e-mail: r.j.a.vanwezel@uu.nl.

# REFERENCES

- Albright, T. D. (1984). Direction and orientation selectivity of neurons in visual area MT of the macaque. *Journal of Neurophysiology*, 52, 1106–1130.
- Alford, J. L., van Donkelaar, P., Dassonville, P., & Marrocco, R. T. (2007). Transcranial magnetic stimulation over MT/MST fails to impair judgments of implied motion. *Cognitive, Affective & Behavioral Neuroscience, 7,* 225–232.
- Barraclough, N. E., Xiao, D., Oram, M. W., & Perrett, D. I. (2006). The sensitivity of primate STS neurons to walking sequences and to the degree of articulation in static images. *Progress in Brain Research*, 154, 135–148.
- Bisley, J. W., Zaksas, D., Droll, J. A., & Pasternak, T. (2004). Activity of neurons in cortical area MT during a memory for motion task. *Journal of Neurophysiology*, *91*, 286–300.

Borghuis, B. G., Perge, J. A., Vajda, I., van Wezel, R. J., van de Grind, W. A., & Lankheet, M. J. (2003). The motion reverse correlation (MRC) method: A linear systems approach in the motion domain. *Journal of Neuroscience Methods*, *123*, 153–166.

Britten, K. H., & Newsome, W. T. (1998). Tuning bandwidths for near-threshold stimuli in area MT. *Journal of Neurophysiology*, 80, 762–770.

Britten, K. H., Newsome, W. T., Shadlen, M. N., Celebrini, S., & Movshon, J. A. (1996). A relationship between behavioral choice and the visual responses of neurons in macaque MT. *Visual Neuroscience*, *13*, 87–100.

Britten, K. H., Shadlen, M. N., Newsome, W. T., & Movshon, J. A. (1993). Responses of neurons in macaque MT to stochastic motion signals. *Visual Neuroscience*, 10, 1157–1169.

Cook, E. P., & Maunsell, J. H. (2002). Attentional modulation of behavioral performance and neuronal responses in middle temporal and ventral intraparietal areas of macaque monkey. *Journal of Neuroscience, 22*, 1994–2004.

Cook, E. P., & Maunsell, J. H. (2004). Attentional modulation of motion integration of individual neurons in the middle temporal visual area. *Journal of Neuroscience*, 24, 7964–7977.

Dukelow, S. P., DeSouza, J. F., Culham, J. C., van den Berg, A. V., Menon, R. S., & Vilis, T. (2001). Distinguishing subregions of the human MT+ complex using visual fields and pursuit eye movements. *Journal of Neurophysiology*, 86, 1991–2000.

Elfron, B., & Tibshirani, R. (1993). *An introduction to the boot-strap*. New York: Chapman & Hall.

Fawcett, I. P., Hillebrand, A., & Singh, K. D. (2007). The temporal sequence of evoked and induced cortical responses to implied-motion processing in human motion area V5/MT+. *European Journal of Neuroscience, 26*, 775–783.

Friston, K. J., Frith, C. D., Turner, R., & Frackowiak, R. S. (1995). Characterizing evoked hemodynamics with fMRI. *Neuroimage*, *2*, 157–165.

Goossens, J., Dukelow, S. P., Menon, R. S., Vilis, T., & van den Berg, A. V. (2006). Representation of head-centric flow in the human motion complex. *Journal of Neuroscience*, 26, 5616–5627.

Green, D. M., & Swets, J. A. (1966). *Signal detection theory and psychophysics*. New York: John Wiley and Sons, Inc.

Huk, A. C., Dougherty, R. F., & Heeger, D. J. (2002). Retinotopy and functional subdivision of human areas MT and MST. *Journal of Neuroscience*, 22, 7195–7205.

Jellema, T., Baker, C. I., Wicker, B., & Perrett, D. I. (2000). Neural representation for the perception of the intentionality of actions. *Brain and Cognition*, 44, 280–302.

Jellema, T., Lorteije, J., van Rijn, S., van 't Wout, M., De Haan, E. H. F., Van Engeland, H., et al. (2009). Involuntary processing of social cues is compromised in autism spectrum disorders. *Autism Research*, 2, 192–204.

Jellema, T., & Perrett, D. I. (2003a). Perceptual history influences neural responses to face and body postures. *Journal of Cognitive Neuroscience, 15,* 961–971.

Jellema, T., & Perrett, D. I. (2003b). Cells in monkey STS responsive to articulated body motions and consequent static posture: A case of implied motion? *Neuropsychologia*, *41*, 1728–1737.

Jellema, T., & Perrett, D. I. (2006). Neural representations of perceived bodily actions using a categorical frame of reference. *Neuropsychologia*, *44*, 1535–1546.

Kaas, J. H., & Collins, C. E. (2004). The resurrection of multisensory cortex in primates: Connection patterns that integrate modalities. In G. A. Calvert, C. Spence, & B. E. Stein (Eds.), *The bandbook of multisensory processes* (pp. 295–310). Cambridge, MA: MIT Press. Kourtzi, Z., & Kanwisher, N. (2000). Activation in human MT/MST by static images with implied motion. *Journal* of Cognitive Neuroscience, 12, 48–55.

Kourtzi, Z., Krekelberg, B., & van Wezel, R. J. A. (2008). Linking form and motion in the primate brain. *Trends in Cognitive Neurosciences*, 12, 230–236.

Krekelberg, B., Dannenberg, S., Hoffmann, K. P., Bremmer, F., & Ross, J. (2003). Neural correlates of implied motion. *Nature*, 424, 674–677.

Krekelberg, B., Vatakis, A., & Kourtzi, Z. (2005). Implied motion from form in the human visual cortex. *Journal of Neurophysiology*, 94, 4373–4386.

Lagae, L., Maes, H., Raiguel, S., Xiao, D. K., & Orban, G. A. (1994). Responses of macaque STS neurons to optic flow components: A comparison of areas MT and MST. *Journal* of *Neurophysiology*, 71, 1597–1626.

Logothetis, N. K., & Pfeuffer, J. (2004). On the nature of the BOLD fMRI contrast mechanism. *Magnetic Resonance Imaging, 22,* 1517–1531.

Lorteije, J. A., Kenemans, J. L., Jellema, T., van der Lubbe, R. H., de Heer, F., & van Wezel, R. J. (2006). Delayed response to animate implied motion in human motion processing areas. *Journal of Cognitive Neuroscience, 18*, 158–168.

Lorteije, J. A., Kenemans, J. L., Jellema, T., van der Lubbe, R. H., Lommers, M. W., & van Wezel, R. J. (2007). Adaptation to real motion reveals direction selective interactions between real and implied motion processing. *Journal of Cognitive Neuroscience, 19*, 1231–1240.

Malpeli, J. G. (1998). Measuring eye position with the double magnetic induction method. *Journal of Neuroscience Methods*, 86, 55–61.

Martínez-Trujillo, J., & Treue, S. (2002). Attentional modulation strength in cortical area MT depends on stimulus contrast. *Neuron*, 35, 365–370.

Maunsell, J. H., & Treue, S. (2006). Feature-based attention in visual cortex. *Trends in Neurosciences*, *29*, 317–322.

Mikami, A., Newsome, W. T., & Wurtz, R. H. (1986). Motion selectivity in macaque visual cortex: I. Mechanisms of direction and speed selectivity in extrastriate area MT. *Journal of Neurophysiology*, 55, 1308–1327.

Morrone, M. C., Tosetti, M., Montanaro, D., Fiorentini, A., Cioni, G., & Burr, D. C. (2000). A cortical area that responds specifically to optic flow, revealed by fMRI. *Nature Neuroscience*, *3*, 1322–1328.

Näsänen, R., & O'Leary, C. (1998). Recognition of band-pass filtered hand-written numerals in foveal and peripheral vision. *Vision Research*, *38*, 3691–3701.

Nelissen, K., Vanduffel, W., & Orban, G. A. (2006). Charting the lower superior temporal region, a new motion-sensitive region in monkey superior temporal sulcus. *Journal of Neuroscience*, 26, 5929–5947.

Orban, G. A. (2008). Higher order visual processing in macaque extrastriate cortex. *Physiological Reviews*, 88, 59–89.

Orban, G. A., Fize, D., Peuskens, H., Denys, K., Nelissen, K., Sunaert, S., et al. (2003). Similarities and differences in motion processing between the human and macaque brain: Evidence from fMRI. *Neuropsychologia*, *41*, 1757–1768.

Perge, J. A., Borghuis, B. G., Bours, R. J., Lankheet, M. J., & van Wezel, R. J. (2005a). Temporal dynamics of direction tuning in motion-sensitive macaque area MT. *Journal of Neurophysiology*, *93*, 2104–2116.

Perge, J. A., Borghuis, B. G., Bours, R. J., Lankheet, M. J., & van Wezel, R. J. (2005b). Dynamics of directional selectivity in MT receptive field center and surround. *European Journal of Neuroscience*, 22, 2049–2058.

Peuskens, H., Sunaert, S., Dupont, P., Van Hecke, P., & Orban, G. A. (2001). Human brain regions involved in heading estimation. *Journal of Neuroscience*, *21*, 2451–2461. Raiguel, S. E., Xiao, D. K., Marcar, V. L., & Orban, G. A. (1999). Response latency of macaque area MT/V5 neurons and its relationship to stimulus parameters. *Journal of Neurophysiology, 82,* 1944–1956.

Ramsey, N. F., van den Brink, J. S., van Muiswinkel, A. M., Folkers, P. J., Moonen, C. T., & Jansma, J. M. (1998).
Phase navigator correction in 3D fMRI improves detection of brain activation: Quantitative assessment with a graded motor activation procedure. *Neuroimage*, *8*, 240–248.

Recanzone, G. H., & Wurtz, R. H. (2000). Effects of attention on MT and MST neuronal activity during pursuit initiation. *Journal of Neurophysiology*, *83*, 777–790.

Reulen, J. P., & Bakker, L. (1982). The measurement of eye movement using double magnetic induction. *IEEE Transactions on Biomedical Engineering*, 29, 740–744.

Schlack, A., & Albright, T. D. (2007). Remembering visual motion: Neural correlates of associative plasticity and motion recall in cortical area MT. *Neuron*, 53, 881–890.

Schmolesky, M. T., Wang, Y., Hanes, D. P., Thompson, K. G., Leutgeb, S., Schall, J. D., et al. (1998). Signal timing across the macaque visual system. *Journal of Neurophysiology*, 79, 3272–3278.

Senior, C., Barnes, J., Giampietro, V., Simmons, A., Bullmore, E. T., Brammer, M., et al. (2000). The functional neuroanatomy of implicit-motion perception or representational momentum. *Current Biology*, 10, 16–22.

Senior, C., Ward, J., & David, A. S. (2002). Representational momentum and the brain: An investigation into the functional necessity of V5/MT. *Visual Cognition*, 9, 81–92. Snowden, R. J., Treue, S., & Andersen, R. A. (1992). The response of neurons in areas V1 and MT of the alert rhesus monkey to moving random patterns. *Experimental Brain Research, 88,* 389–400.

Treue, S., & Martínez-Trujillo, J. (1999). Feature-based attention influences motion processing gain in macaque visual cortex. *Nature*, 399, 575–579.

Treue, S., & Maunsell, J. H. (1996). Attentional modulation of visual motion processing in cortical areas MT and MST. *Nature*, 382, 539–541.

Treue, S., & Maunsell, J. H. (1999). Effects of attention on the processing of motion in macaque middle temporal and medial superior temporal visual cortical areas. *Journal of Neuroscience, 19,* 7591–7602.

van Gelderen, P., Ramsey, N. F., Liu, G., Duyn, J. H., Frank, J. A., & Weinberger, D. R. (1995). Three-dimensional functional magnetic resonance imaging of human brain on a clinical 1.5-T scanner. *Proceedings of the National Academy of Sciences, U.S.A., 92,* 6906–6910.

Van Wezel, R. J., & Britten, K. H. (2002). Motion adaptation in area MT. *Journal of Neurophysiology*, 88, 3469–3476.

Vanduffel, W., Fize, D., Mandeville, J. B., Nelissen, K., Van Hecke, P., Rosen, B. R., et al. (2001). Visual motion processing investigated using contrast agentenhanced fMRI in awake behaving monkeys. *Neuron*, *32*, 565–577.

Zaksas, D., & Pasternak, T. (2005). Area MT neurons respond to visual motion distant from their receptive fields. *Journal of Neurophysiology*, 94, 4156–4167.