Primates need to detect and recognize camouflaged animals in natural environments. Camouflage-breaking movements are often the only visual cue available to accomplish this. Specifically, sudden movements are often detected before full recognition of the camouflaged animal is made, suggesting that initial processing of motion precedes the recognition of motion-defined contours or shapes. What are the neuronal mechanisms underlying this initial processing of camouflaged motion in the primate visual brain? We investigated this question using intrinsic-signal optical imaging of macaque V1, V2 and V4, along with computer simulations of the neural population responses. We found that camouflaged motion at low speed was processed as a direction signal by both direction- and orientation-selective neurons, whereas at high-speed camouflaged motion was encoded as a motion-streak signal primarily by orientation-selective neurons. No population responses were found to be invariant to the camouflage contours. These results suggest that the initial processing of camouflaged motion at low and high speeds is encoded as direction and motion-streak signals in primate early visual cortices. These processes are consistent with a spatio-temporal filter mechanism that provides for fast processing of motion signals, prior to full recognition of camouflage-breaking animals.

1. Introduction
Camouflage is a critical evolutionary development for animal survival, as it prevents detection and recognition of both prey and predators in their natural environments (figure 1a). For instance, a camouflaged animal such as a jungle lizard or an ocean cuttlefish alters its appearance to closely match its surroundings and in doing so conceals itself from both prey and/or predators [1,2]. Yet as soon as it moves, the camouflage breaks and the animal becomes visible [3,4]. Because of the critical importance of motion to breaking camouflage, many animals have developed complex movement strategies to continuously minimize their motion signals [5,6]. Even when the camouflage breaks, full recognition of a moving animal within its environment remains challenging, i.e. motion aids detection but not necessarily identification [7]. This is most likely owing to the weak contrast or contour disruptions between the moving animal and its adapted-to environment [8]. Therefore, in a natural setting, the motion awareness of a camouflaged animal often comes before the recognition of the animal itself. Camouflaged predators typically approach their prey stealthily before attacking with a burst of high speed. Thus, motion is the first potent segmentation cue for the detection of a camouflaged animal, whose visibility is often correlated with its speed of motion. Furthermore, fast-moving objects generate a spatial orientation code, commonly known as motion streak owing to the temporal integration of the visual system [9–14]. We hypothesize that a camouflage-breaking motion engages motion-streak processing in the primate early visual system when an object moves above a certain speed [14–16].
Human psychophysical studies, inspired by the motion of camouflaged animals in natural scenes, have investigated the perceptual mechanisms underlying second-order or non-luminance defined motion perception \cite{17,18}. These studies posit the existence of a secondary motion processing stage distinct from the primary motion stage for the detection of luminance displacements \cite{19,20}. Physiological studies of neuronal responses from the visual cortices of old-world macaques have focused on the orientation-cue invariance for the processing of various motion-defined boundaries.

Eighty-nine per cent (143 out of 160) of units recorded in macaque V1 was found to be activated by luminance and temporal texture bars. However, among all these activated units, only around 21.7\% (31 out of 143) of units retained the same orientation preference for both stimuli \cite{21}. As direction-selective cells locate mainly in layer 4B and 6 in macaque V1 \cite{22–24}, it is not surprising that less than 10\% of V1 cells were direction selective. Interestingly, recent findings from the common marmoset, a New World primate, found approximately 42\% (34 out of 81) of V1 cells recorded from displayed invariant responses to stimulus orientation or direction of motion when probed with luminance bars or random-noise bars moving on a flickering background \cite{25}.

Neurons in macaque middle temporal (MT) area have much larger receptive fields (RFs) than in V1 and V2 (figure 1a) \cite{26,27}, and most neurons are selective for dynamic temporal textured bars similar to a subset of V1 neurons \cite{21,28}. Unexpectedly, neurons in macaque MT are not selective to the orientation of static boundaries defined by coherent local motion but instead are thought to encode local motion vectors \cite{29}. The overall percentage of neurons in macaque V4 responding to motion-defined gratings or shapes has also been reported to be quite low (in the range of 10–20\%) \cite{30,31}. Single-unit recordings in macaques appear consistent with findings from human brain imaging where no specific visual area has been found to exclusively encode motion-defined boundaries \cite{32–34}, although areas V3 and V3A respond to oriented shapes generated by motion-defined stimuli \cite{34}. Further along the visual hierarchy, motion-defined shape is assumed to be fully represented in a cue-invariant manner in the inferior temporal (IT) area \cite{35–37}. This might explain why the recognition of a camouflage-breaking animal occurs slowly compared with the initial processing of the motion that gives the animal away. Surprisingly, no studies have directly compared the change of orientation-selective responses to stimuli mimicking camouflaged movements with low and high speeds in macaque V1 and V2, yet this information is critically important for understanding the breadth of camouflage-breaking motions that occur in nature \cite{6}.

In this study, we compared the cortical population responses to moving oriented contours created from sine-wave luminance gratings (LG) to random-noise inducers moving coherently against a static random-noise-textured background. Both types of stimuli had the same orientation and motion direction (figure 1b). Such drifting noise gratings, which disappear immediately when they stop moving, were chosen to more accurately mimic the motion of a camouflaged animal against a static background than static motion-defined contours \cite{38} or dynamic contours moving against a dynamic noise background \cite{25}. We found that direction and motion-streak signals within macaque V1 and V2 underlies the processing of camouflage-breaking stimuli at low and high speeds, respectively.

2. Material and methods

(a) Animal preparation and maintenance

Nine adult rhesus macaques (five male and four female) (Macaca mulatta) weighing 3.0–5.0 kg were prepared and maintained (see the electronic supplementary material) for optical imaging as previously described \cite{15,39,40}.

(b) Visual stimuli

A gamma-corrected monitor (Sony G520, 1280 × 960 pixels, 100 Hz, luminance ranged from 0.2 to 82 cd m\(^{-2}\)) was placed 57 cm in front of the animal eyes covering 40 × 30 degrees. Full-screen visual stimuli were generated using PSYCHTOOLBOX-3 and MatLab 2010a run under WINDOWS XP and were presented binocularly. We used eight orientations of LG stimuli (figure 1b) with spatial frequency (SF) of 1–2 cycles degree\(^{-1}\) and temporal frequency of 5 cycles s\(^{-1}\). For random-dot stimuli, dot diameter was 0.2°, density 5 dots degree\(^{-2}\), and a dot speed of 1–2 s\(^{-1}\) (degree per second, with no accumulation). We generated camouflage-breaking stimuli (figure 1b), which were moving oriented camouflage contours (CC) created from local random-noise inducers moving coherently against a static random-noise-texture background. The two-dimensional noise texture was composed of randomly positioned noise elements with each element spanning approximately 3.6 arcmin and had a mean luminance of 41 cd m\(^{-2}\). The CC SF was 0.375 cycles degree\(^{-1}\) and the speeds of CC inducers were 1° s\(^{-1}\) or 7° s\(^{-1}\).
in this experiment. The equations for generating CC stimuli together with flash movie simulations of the stimuli are presented in the electronic supplementary material.

(c) Optical imaging and data analysis
Intrinsic-signal optical imaging was used as described previously [15,39,40] to measure the cortical response of V1, V2 and V4. Orientation and direction preference maps were classically constructed using a vector summation algorithm [41]. Angular differences between different pairs of orientation and direction preference maps were calculated and histograms were constructed to display the results. Response profile analysis was performed to extract the orientations best represented by the differential orientation maps [15,42,43] (see the electronic supplementary material for details). For the quantification of the response amplitude, the max $\Delta R/R$ values (changes of reflected light) in each responsive patch of a differential map were averaged.

(d) Model simulation
To simulate the neural responses of V1 and V2 populations, we used a spatio-temporal energy model [15,39,44]. For the simulation responses to camouflage-breaking stimuli, a total of 32 trials were averaged. The responses to two orthogonal stimuli were subtracted to match our optical imaging results. The equations and descriptions of the energy model are detailed in the electronic supplementary material.

3. Results
Most neurons in V1 and V2 of macaques are orientation selective with small spatio-temporal RFs and precise retinotopic coordinates [45,46]. These orientation-selective neurons respond invariantly to bidirectional moving gratings or bars and are clustered into columns/domains, from which the population activities (mainly within cortical layers 2/3) can be recorded topographically using intrinsic-signal optical imaging [47,48].

(a) Camouflage-breaking stimuli activated orientation domains
We generated differential maps of orientation preference by subtracting intrinsic signals evoked by alternating full-field stimuli with orthogonal orientations. We did this for both LG and camouflage-breaking stimuli, moving bidirectionally at $7 \text{s}^{-1}$ (a speed that generates robust cortical responses). Figure 2r depicts examples of the $0^\circ$ and $90^\circ$ stimuli. In the differential response maps, dark regions prefer the first stimulus condition and bright regions prefer the second (figure 2b). Both LG and camouflage-breaking stimuli activated orientation domains within the same region of interest (ROI) of V1 and V2. However, when measuring the relative change in the amount of reflected light ($\Delta R/R$), response magnitudes elicited by camouflage-breaking stimuli were only about 50% of those activated by LG stimuli (figure 2b, see scale bars). These weaker population responses are consistent with the idea that camouflaged movements are harder to detect. We constructed orientation preference maps for the LG and camouflage-breaking stimuli (figure 2c,d) and found that the angular differences of preferred orientations for camouflage-breaking and LG stimuli peaked around $\pm 90^\circ$ using pixel-by-pixel subtraction (figure 2c).

(b) Orientation preferences to the camouflage-breaking stimuli dependent on speed
Without exception population responses of V1 and V2 to camouflage-breaking stimuli moving at $7 \text{s}^{-1}$ displayed robust tuning with $90^\circ$ shifts in orientation preferences compared with LG stimuli (figure 2c). A recent study [15] investigating the motion-streak [11,14,49] (where faster texture/dot motion can activate orientation-selective cells with preferences parallel to the motion direction) has shown that full-field noise patterns moving at $7 \text{s}^{-1}$ activated orientation domains with orthogonal preference to those activated at $1 \text{s}^{-1}$ or by moving LGs. We wondered whether the cortical population responses were dependent on the speed of the camouflage-breaking stimuli.

We tested this hypothesis in macaque V1, V2 and V4 by comparing the cortical responses to camouflage-breaking stimuli moving at speeds of 1 and $7 \text{s}^{-1}$ (figure 3). Within the response regions of V1 and V2 (figure 3a), the differential orientation responses to camouflage-breaking stimuli moving forwards and backwards (figure 3b) at speeds 1 and $7 \text{s}^{-1}$ were superimposed with coloured iso-orientation contours derived from the orientation preference map generated using LG stimuli (figure 3c,d). With orientation response profile analysis [42], we found that the orientation preference responses in both V1 and V2 elicited by camouflage-breaking stimuli moving at $1 \text{s}^{-1}$ were in close register to those evoked by LG stimuli, while those activated by camouflage-breaking stimuli moving at $7 \text{s}^{-1}$ were orthogonal, with a roughly $90^\circ$ shift in orientation preference (tuning curves in figure 3c,d). Similar results were also observed in V4 (see an example in the electronic supplementary material, figure S1).

(c) Unidirectional camouflage-breaking stimuli activated direction domains in V2
The majority of neurons in macaque V1, V2 and V4 are orientation selective and respond equivalently to a LG stimulus moving in either of two opposing directions. We thus inferred that at $1 \text{s}^{-1}$, it was the direction of camouflage-breaking stimuli moving forwards and backwards that was responsible for activating the orientation columns as it was with the LG stimuli. In order to confirm this, we asked whether the unidirectional motion of the camouflage-breaking stimuli at $1 \text{s}^{-1}$ could activate direction-selective domains in macaque V2. No direction-selective domains can be mapped in macaque V1 by using intrinsic optical imaging [15,50]. We used full-field random dots moving in four opposite directions ($0^\circ$, $180^\circ$, $90^\circ$ and $-270^\circ$) at a speed of $1 \text{s}^{-1}$ as our control stimuli (electronic supplementary material, figure S2a,b). Camouflage-breaking stimuli moving at a speed of $1 \text{s}^{-1}$, activated direction domains with similar response patterns in the same locations of V2 as the random dots (electronic supplementary material, figure S2c). Pixel-by-pixel subtraction of the angles of preferred directions for random dots and camouflage-breaking stimuli showed histograms peaking around $0^\circ$ for both ROIs (electronic supplementary material, figure S2d). Similar results were observed in all macaques tested. These findings demonstrate that unidirectional signals in camouflage-breaking stimuli moving at $1 \text{s}^{-1}$ generate responses from direction columns in segregated locations of V2.
(d) Local motion cues responsible for the population responses

As shown above, the population responses to camouflage-breaking stimuli are dependent on motion speed. We reasoned that it must be the motion of the local-noise inducers, not the global camouflage contours, that activates the orientation domains in V1 and V2. This idea is outlined in the electronic supplementary material, figure S3. Motion-defined contours can be generated by noise moving in any direction [19, 20]. So we designed different variants (CC1–5) of the camouflage-breaking stimuli to test whether it was the local motion cues that dominated the cortical responses. We analysed the responses of orientation domains from the same locations in V1 and V2 when the camouflage contours were induced by noise moving either parallel or perpendicular to the contour orientations (figure 4a).

In our standard camouflage-breaking stimuli, the direction of coherent noise motion was perpendicular to the orientation of the camouflage contours. Consistent with the results in figures 2 and 3, the preferences of orientation domains activated by the LG and camouflage-breaking stimuli moving at 7° s⁻¹ were orthogonal in both V1 and V2 (figure 4b,c; LG versus CC). For the CC1 variant, the global camouflage contour moved identically to the camouflage-breaking stimuli, but the direction of the local noise inducers was parallel rather than perpendicular to the orientation of the camouflage contours. The pattern of orientation domains activated by CC1 was closely in register with that elicited by LG in both V1 and V2 (figure 4d; LG versus CC1). CC2 contained exactly the same horizontal and vertical global camouflage contours as CC and CC1; however, in this case, the local noise inducers of the two stimulus conditions moved along an identical axis of motion (perpendicular to the horizontal camouflage contours and parallel to the vertical camouflage contours). This configuration contained the same local motion cues in both cases (that move in the same direction) and failed to produce any responsive domains in either V1 or V2, resulting in an almost flat tuning curve (figure 4e; CC2) despite the fact that global camouflage contours with orthogonal orientations were used. In our last configuration, CC3, population responses were activated by global camouflage contours of the same orientation, but with local noise motion set at right angles...
to each other (i.e. the same orthogonal axes of motion as in CC). Once again, reversed orientation preference maps emerged from responses in both V1 and V2 (figure 4b,f; LG versus CC3). Electronic supplementary material, figure S4, shows the results from another macaque, where the noise moved at an acute angle to the camouflage contours (CC4 and CC5). Similar findings were observed in all animals studied. Together, these findings demonstrate that it was the motion of the local-noise inducers, rather than the orientation of the global camouflage contours, that was responsible for the activation of orientation domains in V1 and V2.

(e) A spatio-temporal energy filter model simulated all population responses

Recently, V1 population responses across species [15,44,51,52] have been successfully reproduced by a spatio-temporal linear filter model. We implemented this generic energy model to systematically simulate the population responses in macaque V1 and V2 to camouflage-breaking stimuli at high and low speeds. We used the camouflage-breaking stimuli of 0° and 90° orientations moving in eight opposing directions at 7° s⁻¹ to simulate V1 and V2 population responses (electronic supplementary material, figure S5a). The orientation response

Figure 3. Orientation domains in V1 and V2 activated by CC stimuli at speeds of 1 and 7° s⁻¹. (a) The surface vasculature of the left hemisphere of macaque 080921022 with ROI of V1 and V2. (b) The LG and CC stimulus pairs. (c) Differential maps for 0° and 90° orientation pairs from V1 (upper panel) and V2 (lower panel) as areas boxed in (a) superimposed with coloured iso-orientation outlines derived from the orientation preference map generated with LG stimuli. The response profiles depict the orientation preferences of CC stimuli moving at speeds of 1 and 7° s⁻¹. (d) The results for CC stimuli of 45° and 135° orientation pairs moving at speeds of 1 and 7° s⁻¹. The responses represent the average pixel values within orientation bins defined by using the LG orientation preference map.
profiles derived from the model simulation closely matched those in our experiments. Specifically, the resulting response profiles were reversed for CC and CC1 stimuli (as in figure 4c,d) and also for CC4 and CC5 stimuli (as in electronic supplementary material, figure S4). We further simulated V1 and V2 population responses to camouflage-breaking stimuli at 18 s⁻¹ and the resulting simulated orientation response profiles were reversed to those derived from the camouflage-breaking stimuli moving at 78 s⁻¹ (electronic supplementary material, figure S5b), consistent with the observations revealed by optical imaging.

In both V1 and V2, the simulated response strength for LG stimuli was the highest, while the simulated response strength for camouflage-breaking stimuli moving at 18 s⁻¹ was the lowest and that for camouflage-breaking stimuli moving at 78 s⁻¹ was in the middle (electronic supplementary material, figure S5c). These simulations closely matched the trend of the average response strengths directly measured in the optical imaging experiments (electronic supplementary material, figure S5d). We found that the average population responses to LG stimuli were significantly larger than those to camouflage-breaking stimuli moving at speed 78 s⁻¹ in both V1 and V2 (p < 0.05, n = 12, paired t-test). In addition, the responses to camouflage-breaking moving at speed 18 s⁻¹ were significantly weaker than those to camouflage-breaking stimuli moving at speed 78 s⁻¹ (p < 0.05, n = 12, paired t-test). Together, these results suggest that the population responses of V1 and V2 to camouflage-breaking stimuli can be accounted for by a simple linear spatio-temporal filter mechanism.

4. Discussion

Animal camouflage is a striking example of a natural evolutionary structural and behavioural adaptation that has
long fascinated many branches of scientific investigation [8]. How the visual brain detects camouflaged animal movements and subsequently recognizes the camouflaged animal remains unclear. This study explores the first part of this question by examining population responses across visual areas in macaques, the best animal model for human vision. We found that orientation-selective domains within macaque V1, V2 and V4 engaged in the processing of different components of camouflage-breaking motion at low and high speeds.

(a) The population responses of macaque V1, V2 and V4 to camouflage-breaking stimuli

In macaques, most direction-selective neurons in V1 are simple or complex cells residing in layers 4B and 6 [22,23,46]. The neurons in these layers are too deep for their activity to be recorded by intrinsic-signal optical imaging. As the majority of neurons in V1 layers 2/3 are orientation-selective and respond equally well to motion in opposite directions, it was the noise inducers moving bidirectionally at 1°s⁻¹ that were responsible for the activation of orientation domains in V1 and thereafter in V2 and V4. However, as previously reported, when the speed of motion was increased to 7°s⁻¹, the movement of the random-noise texture generated a spatial orientation signal (commonly referred to as the motion-streak) parallel to the motion direction of the moving noise stimuli [15]. Thus, the orientation-selective neurons with preferences parallel to the motion streak were activated and their responses were subsequently recorded by intrinsic optical imaging as orientation domains in V1, V2 and V4. The temporal integration of visual inputs along the axis of motion was also precisely accounted for by the spatio-temporal energy model, consistent with the notion that motion can be linearly processed in primate early visual cortices [15,53,54]. In comparison to LG stimuli, the weaker population responses to camouflage-breaking stimuli may relate to the poor visibility of concealed animals even when moving, supporting the notion of motion-streak using random dots and noise patterns [15,16]. Perhaps the presence of a twinkling background, in the Bourne et al. [25] study, made it impossible for the motion-streak signal to develop and so interfered with the orientation selectivity.

Another study reported even a lower percentage of cells (less than 10%) in macaque V1 and V2 showing cue-invariant responses to the orientation of motion-defined but static contours [8], although this study also did not look at the effect of motion streak. As intrinsic signals are derived from pooled neuronal activities, it is unlikely that intrinsic optical imaging would reveal this small percentage of cells in macaque V1 and V2; and even in V4 where the number of neurons responding to motion-defined contours and shapes is reported to be higher (up to around 20%) [30,31]. Hence, the population response in V1, V2 and V4 mainly encodes the local motion signals in camouflage-breaking stimuli as revealed here both experimentally and mathematically. A very recent study in awake macaques has however claimed that these small number of neurons in macaque V2 but not V1 can be recorded by intrinsic optical imaging, and that motion-defined boundaries are thus mapped in orientation domains of V2 [59]. These authors used static boundaries defined by dot motion fields moving at slow speeds (1 to 2°s⁻¹) and the motion axis of the moving dots was always at a 45° angle with respect to the motion defined strip borders. Unfortunately, they did not test their stimuli at higher speeds and did not vary the local motion direction of the random dots relative to the global motion-defined boundary orientation. Hence, it is uncertain if their motion boundary results are invariant or not to speed as well as to local inducer motion direction. In both V1 and V2 of anesthetized macaques as we have demonstrated, it is the movement of the local-noise inducers within our camouflage-breaking stimuli that dominates the neural responses we recorded. However, neurons encoding motion-defined contours were prevalent in cat visual area 18 [60], suggesting carnivores may have developed more efficient ways for detecting camouflaged movements.

It is expected that neurons in MT, the motion centre of primates which receives direction signal inputs from V1 layers 4B/6, V2 thick strips, and directly from the thalamus [22–24,61], will also encode motion direction/axis signals within camouflage motion. Furthermore, as a higher percentage of cells in both the V3 and IT area have been found to respond to simple geometric shapes defined by motion-defined contours [31,32,34–37], it is more likely that robust responses to motion-defined contours as well as to camouflage contours will be found in these higher visual areas. Thus, the recognition of a camouflaged animal on the move must engage multiple visual areas working together.

5. Concluding remarks

A thought-provoking implication of our findings relates to the wide prevalence of conspicuous oriented stripes and zig-zags found across many animal species (for example zebras) and used by humans in camouflage designed for moving objects (famously warships in World War I and II). These oriented-stripe patterns, though more salient when static, are hypothesized to generate a ‘motion dazzle’ that makes it difficult for either predator or prey to judge speed and/or direction of motion [8]. In addition, the interference
by oriented motion dazzle has been observed in human psychophysical tests, where oriented patterns can impede motion task performance [62–64], and/or cause misperceptions of speed [65,66]. Thus, a more detailed understanding of the neural population responses to direction and motion-streak signals across speeds can usefully constrain and inform future behavioural studies of motion crypsis. In summary, low-speed camouflage-breaking movements are encoded by both direction- and orientation-selective cells corresponding to the direction of motion of the concealed animal, while high-speed camouflage-breaking movements generate a motion-streak signal that is encoded by a complementary set of orientation-selective cells within V1, V2 and V4 of the macaque ventral visual stream.

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Ethics statement. All experimental procedures for primate research including animal euthanasia were approved by the Institute of Neuroscience Institutional Animal Care and Use Committee and by the local ethical review committee of the Shanghai Institutes for Biological Sciences.

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