



Neural construction of 3D medial axis from the binocular fusion of 2D MAs



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ABSTRACT

The perceptual constancy of shape, including view invariance, is an amazing property of the visual system. Cortical representation by the medial axis (MA) is an attractive candidate for maintaining the constancy of a wide range of arbitrary shapes. Recent physiological studies have reported that neurons in the primary visual cortex (V1) show a response to two-dimensional (2D) MAs, and those in the inferior temporal cortex (IT) are selective to three-dimensional (3D) MAs. However, little is known about the neural mechanisms underlying the transformation of 2D to 3D MAs. As a first step toward investigating the cortical mechanism, we have proposed as a hypothesis that a pair of monocular 2D MAs is fused to generate a 3D MA. We examined the computational plausibility of the hypothesis; specifically, whether an energy-based fusion model is capable of generating 3D MAs. We generated blob-like, physiologically plausible 2D MAs, and used a standard energy model to detect the disparity between a pair of 2D MAs. The model successfully generated 3D MAs for a variety of objects that included typical shape characteristics. A reconstruction test showed that the computed 3D MAs captured the essential structure of the objects with reasonable accuracy and view invariance. These results indicate that the fusion of monocular blob-like 2D MAs is capable of generating a reasonable 3D MA within the framework of the energy model.

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1. Introduction

Robust perception of the shape of objects is an amazing property of the visual system. Although the view and the size of an object on a retinal image change dramatically as we see the object from different directions and distances, our visual system perceives a stable, invariant shape for the object. The representation of shape in the visual cortex should play a crucial role in realizing such invariance in shape perception. An object-centered representation that describes shape as a spatial arrangement of parts has been supported widely by psychological and physiological studies [1–3], as it has the ability to establish the perceptual constancy of shape, including view and distance invariance. The medial axis (MA) is considered suitable for a parts-based representation among theorists [4,5]. MA representation encodes each part of the object with a medial line that is derived from the local symmetry of the part. This representation, based on an object-centered coordinate, is independent of view and capable of describing shape efficiently using two types of parameters: the spatial arrangement and relative length of the axes corresponding to the parts [4,6,7]. MA is an attractive

candidate for the cortical representation of shape, as a robust and efficient coding scheme [8].

Recently, Hung et al. showed that a number of neurons in the inferior temporal cortex (IT) encode three-dimensional (3D) MA configurations, supporting the idea that the MA plays a critical role in the representation of shape in the ventral pathway [9]. IT has been reported to encode the 3D structure of shape [10,11], but little was known about the representation scheme for 3D shapes. The selectivity for 3D MA configurations reported recently in IT has provided crucial direct evidence to support MA coding for the cortical representation of shape. A recent fMRI study has also reported the cortical representation of MA structure in the ventral stream [12]. However, the computational processes that constitute the 3D MA along the ventral pathway remain unknown. One of the keys to understanding these processes lies in the lower cortex: cells in the primary visual cortex (V1) show strong responses to the MA of a textured figure [13,14]. Computational studies have shown that the MA response in V1 can be generated by simultaneous arrival of traveling spikes that are initiated by nearby V1 cells [14], or from onset synchronization of border-ownership (BO)-selective cells in V2 [15,16]. These computational studies have also reported that the generated MA encodes arbitrary two-dimensional (2D) shapes. These studies note that the MAs were not like thin skeletons as previous studies have assumed, but rather, the MAs were elongated blobs with spatial extent. This

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blob-like MA is expected to be robust for 3D construction. Because the structure of skeletons is sensitive to the direction of view (binocular difference) and noise in the contours, small changes in view and contour dramatically alter the structure of skeleton-like MAs, leading to erroneous stereo-matching. However, blob-like MAs are expected to be insensitive to such changes [17]. Investigating the fusion of blob-like MAs rather than conventional skeleton-like MAs is essential. The intermediate areas of the ventral visual pathway such as V4 are known to play a crucial role in the binocular fusion of object shapes [18,19]. A certain translation function that takes place along the ventral pathway may contribute to the construction of the 3D MA observed in IT from the 2D MAs observed in V1.

We investigated the cortical mechanisms underlying the construction of 3D-shape representation, by focusing on blob-like 2D MAs and their fusion along the ventral pathway. Fusion of 2D MAs based on their disparity is a plausible candidate mechanism for filling the gap between the 2D MA in the primary cortex and the 3D MA in the higher cortex. It is conceivable that the 2D MAs resulting from the left and right retinal images are fused in an intermediate-level area by a process based on disparities in the 2D MAs, thereby establishing a 3D MA in IT. An alternative mechanism for the construction is that the MA responses in V1 are binocular with absolute disparity, and are thus “3D MA segments”. The 3D MA segments in V1 would then be integrated along the visual pathway to establish a global 3D MA with relative disparities in IT. Although a number of V1 cells are selective to the binocular disparity of contours, it is not at all certain whether cells responding to MAs are selective for the binocular disparity of the local MA. V1 cells could respond to the depth of contours, but not necessarily to that of the MA. Specifically, the depths of both sides of an object as well as its MA are generally different. This concept is illustrated by a cuboid with a different depth for each side of the object; for example, the left side is nearer and the right side is farther (see Fig. 1A). Although the depths of these sides can be determined correctly, the depth of the MA is inherently ambiguous; the MA could be located anywhere between the two sides and there is no way to determine its depth from the depth of the sides. On the other hand, in the former case involving 2D MAs, the local disparities between the 2D MAs could be integrated without ambiguity (see Fig. 1B). This idea appears to be consistent with the tuning of three-dimensional orientation in the macaque V4 [19]. In the present study, we focused on the fusion of monocular 2D MAs that are formed in V1, and are fused along the ventral pathway based on the disparities between the axes, to generate a 3D MA in IT.

Physiological evidence for the generation process of a 3D MA has not been available. As a first step toward investigating our hypothesis, we conducted computational studies to determine whether the fusion of monocular, blob-like (physiologically plausible) 2D MAs is capable of generating a 3D MA, and how accurately this method would work. Specifically, we constructed a fusion model based on a standard energy model [20] that is thought to capture the essential functions of physiological properties in early- to intermediate-level visual areas. We examined whether the model is capable of generating a correct 3D MA, and whether the computed 3D MA captures the essential structure that is sufficient for the reconstruction of a 3D shape. Our simulation results showed that the model was capable of generating 3D MAs for a variety of shapes including those of natural objects. The results also showed that the reconstruction of 3D shapes based on the computed 3D MAs was successful, with similar levels of accuracy for various shapes with different degrees of shape complexity, which is one of the most remarkable features of the visual system. Furthermore, we tested view invariance of the model in terms of the reconstruction error. Similar reconstruction errors were observed for images from different views,

suggesting that the representation of a 3D MA from the fusion of 2D MAs has invariance to rotation. View invariance has been reported in MA-selective cells in IT [9]. Our results indicate that the energy-based fusion of monocular blob-like 2D MAs is capable of generating a 3D MA with robustness in terms of shape complexity and view invariance. Therefore, the generation of a 3D MA from the fusion of 2D MAs is a plausible candidate for the cortical mechanisms underlying the representation of 3D shape.

2. The model

To investigate whether the fusion of physiologically plausible 2D MAs is capable of generating a correct 3D MA, and whether the computed 3D MA captures the structure essential for the reconstruction of 3D shape, we constructed a computational model and conducted simulations. An outline of the model is illustrated in Fig. 1. The model is composed of two stages: (i) the detection of monocular 2D MAs based on the distances from surrounding contours, and (ii) the generation of a 3D MA from the disparities between the two 2D MAs (Fig. 1B). A unit in the first stage computes the distances between the unit and the points on the contours surrounding the unit, and evaluates how much the unit is similarly distant from the surrounding contours by taking pairwise differences between the distances. Units with small differences (similar distances) tend to be located around local symmetry axes, thus their locations are highly likely a part of the 2D MA. The second stage fuses a pair of 2D MAs using a standard energy model to generate a 3D MA. Note that the model includes neither the representation nor the reconstruction of a 3D object. We conducted the reconstruction in Section 3 solely for the evaluation of the computed 3D MA.

2.1. The detection of 2D MA

A computational study by Hatori and Sakai has shown that onset synchronization of BO-selective cells appears to generate V1 activities in response to 2D MAs [16,21]. BO-selective cells on the contour of a figure depolarize if the figure is located on their preferred side [22]. The spikes from BO-selective cells, which are initiated at the same time and travel at the same speed, reach the center of the figure at the same time. Temporal integration of the traveling spikes would result in strong responses of cells located at the center of the figure and along the axes of local symmetry, generating the V1 activity corresponding to the MA. The magnitude of the activity depends on how much the cell is similarly distant from the contours. Taking into account the essence of their idea, the present model computes the possibility of being a 2D MA based on distances from the surrounding contours. Although Hatori's model was capable of processing multiple objects, we limited our model to deal with a single object for the sake of simplicity. We computed an index that describes how much a cell is similarly distant from the contours. If the value of the index exceeds a certain threshold, we consider it as an indication of the MA.

The input to the model was a pair of stereo images with a spatial resolution of 200×200 pixels (considered as $5 \times 5^\circ$ of visual angle). To evaluate the similarity of distances from nearby contours, we measured the distance, $\text{dist}(p, q_i)$, between a point within a figure, p , and every point on the contour, q_i

$$\text{dist}(p, q_i) = \|p - q_i\|, \quad (1)$$

where $\|\cdot\|$ represents the Euclidean distance between the two points. The distances between p and q_i were measured for every 5°

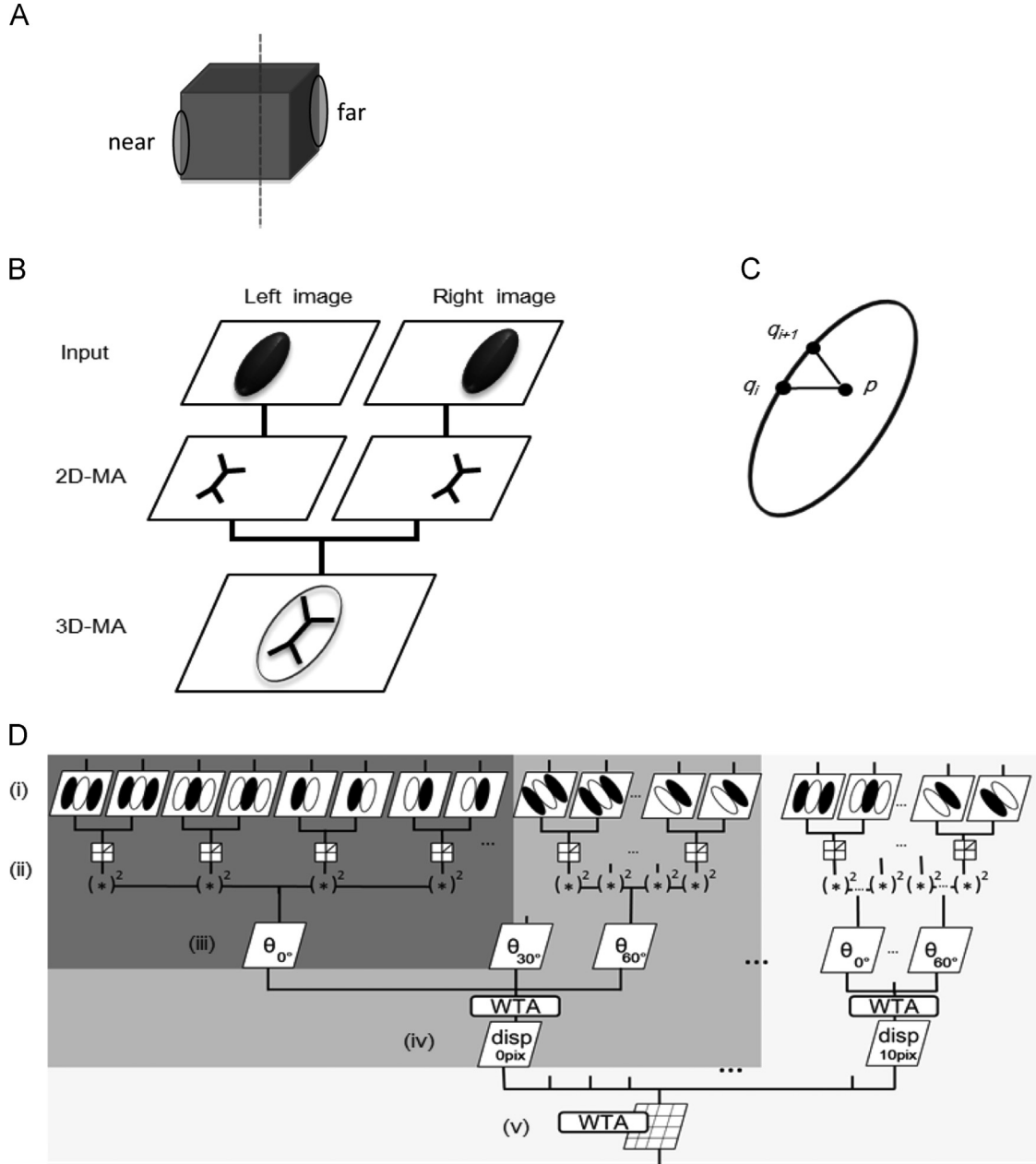


Fig. 1. (A) An illustration showing that, in general, the depths of the sides and the MA of an object could be different. In this example, the right and left sides of the cube are far and near, respectively, with respect to the vertical dotted line. (B) The model comprises two distinct stages: detection of monocular 2D MAs based on the distances from surrounding contours, and generation of a 3D MA from the disparities between the two 2D MAs. (C) A 2D MA is defined as a set of points (e.g., p) equidistant from nearby contours (q_i, q_{i+1}). (D) A detailed illustration of the model. Activities of a pair of simple cells with a certain phase difference (e.g., an in-phase pair for disparity=0) are summed (i), and pass through a half-squaring computation (ii). A model complex cell pools four quadrature pairs of simple cells whose preferred orientation is one of three orientations ($0^\circ, 30^\circ$, or 60°) (iii). The responses of three complex cells with a distinct preferred orientation are integrated by winner-take-all (iv). There are 11 channels with distinct phase differences, corresponding to 11 distinct disparities. The optimal disparity at each spatial position is chosen from the 11 distinct disparities by winner-take-all (v).

(Fig. 1C)

$$q \in \mathbf{Q}, \quad \text{where} \quad \mathbf{Q} = \{q_i | \angle q_i p q_{i+1} = 5^\circ\}. \quad (2)$$

The equidistance index, $E(p)$, is given by a mean of the pairwise differences in the distance between p and q_i

$$E(p) = \frac{1}{|\mathbf{Q}|} \left\{ \sum_{i=1}^{|\mathbf{Q}|} \sum_{j=i+1}^{|\mathbf{Q}|} s(\text{dist}(p, q_i) - \text{dist}(p, q_j)) \right\}, \quad (3)$$

where $|\mathbf{Q}|$ indicates the number of the elements of \mathbf{Q} . To reproduce the nonlinearity of neural responses, we introduced a

sigmoidal function for $s(\cdot)$

$$s(x) = 1 - \frac{1 - e^{(-x/w)}}{1 + e^{(-x/w)}}, \quad (4)$$

where a constant, w , controls the rate of sigmoidal decay. Throughout the simulations, we set w to 6 so that the decay is 10% if the difference in distance is 18 pixels.

We computed the equidistance index for all points within a figure. A unit with a higher index value is likely to be located around the local axes of symmetry. We consider that units with an index value higher than or equal to a threshold, $E_{\text{threshold}}$, correspond to the MA. Therefore, an index to represent how much

a unit is likely to be part of the MA is given by the equidistance index with a threshold, $E_{threshold}$

$$MA_index(p) = \begin{cases} E(p), & \text{if } E(p) \geq E_{threshold}, \\ 0, & \text{otherwise,} \end{cases} \quad (5)$$

We chose empirically $E_{threshold} = 0.26$. This value is crucial for the formation of MA. Although this threshold could be fixed for all stimuli, we chose to fine-tune the value within 15% because details of the formation of 2D MAs are not the focus of our study. To avoid an abrupt distribution of the MA, we introduced Gaussian smoothing to **MA_index**

$$MA(p_x, p_y) = (MA_index * Gauss)(p_x, p_y), \quad (6)$$

where (p_x, p_y) is the spatial position of a point p , and $*$ and *Gauss* represent convolution and a 2D Gaussian with $\sigma_x = \sigma_y = 2$ pixels, respectively. The optimal size of the Gaussian could be different among objects depending on their spatial extent. However, our test showed that the computed MAs were barely sensitive within the range of 3σ . As previously noted, we define a 2D MA as a set of points that are located nearly equidistant from surrounding contours. Therefore, our 2D MA is a fat region with spatial extent, which is distinct from an engineering MA that is defined by skeletons. A 2D MA was computed for each ocular image. A binocular pair of 2D MAs was used to generate a 3D MA as described in the next section.

2.2. The detection of 3D MA

To obtain a 3D MA from a pair of monocular 2D MAs, we computed disparities between the two axes. Fig. 1D shows a diagram of the computation. We used a standard energy model for binocular disparity [23]. We assumed that a fusion mechanism similar to the energy model might take place along the ventral pathway probably in intermediate-level visual areas. The model consists of a cascade of simple- and complex-type cells with half-wave rectification. A pair of 2D MAs was used as input, and the disparities were determined as described below.

A model complex cell consists of a pooling of four quadrature pairs of model simple cells (Fig. 1D(i)) with a particular binocular disparity. The response of a pair of simple cells, $O^1(x, y)$, was computed by the convolution of a monocular image (2D MA) and an oriented Gabor function with a particular orientation, phase, and disparity. We summed up the responses for the right and left images, and passed them through a half-wave rectification step (Fig. 1D(ii))

$$O^1(x, y) = \begin{cases} \text{sum_simple}(x, y), & \text{if } \text{sum_simple}(x, y) \geq 0, \\ 0, & \text{otherwise,} \end{cases} \quad (7)$$

where

$$\text{sum_simple}(x, y) = (nMA_{left} * Gabor_{left})(x, y) + (nMA_{right} * Gabor_{right})(x, y). \quad (8)$$

nMA_{left} and nMA_{right} represent a normalized 2D MA for the left and right images, respectively. $MA(x, y)$ of an image was normalized to its maximum value so that nMA_{left} and nMA_{right} range between 0 and 1. $Gabor_{left}$ and $Gabor_{right}$ represent the oriented receptive field in V1 for the left and right images, respectively. A detailed description of $Gabor_{left}$ and $Gabor_{right}$ is given in Appendix A.

The response of a model complex cell, $O^2(x, y)$, was given by the summation of squared outputs of the four quadrature pairs of the model simple cells, O^{1,ϕ_i} (Fig. 1D(iii)):

$$O^2(x, y) = \sum_{i=1}^4 (O^{1,\phi_i})^2. \quad (9)$$

To establish orientation-invariant selectivity, we pooled three types of complex cells with distinct optimal orientations ($\theta = 0, \pi/6, \pi/3$) by using a winner-take-all mechanism. Although the three channels for orientation appear fewer than those in V1, we chose three for the sake of simplicity. The response of the winner complex cell with disparity, ψ_j , is given by (Fig. 1D(iv))

$$O^{3,\psi_j}(x, y) = \max_k (O_{\theta_k}^{2,\psi_j}(x, y)). \quad (10)$$

The model has 11 distinct disparity channels ($j = 1-11$), resulting in the range of disparity between 0 and 10 pixels. The disparity of a location is given by a winner-take-all mechanism, that is, the preferred disparity of a cell with the strongest response among the 11 disparity channels is chosen as the disparity of the location (Fig. 1D(v)):

$$\text{disp}(x, y) = \underset{\psi_j}{\operatorname{argmax}} \{O^{3,\psi_j}(x, y)\}. \quad (11)$$

We defined horizontal disparity as

$$\text{disparity}(x, y) = \begin{cases} (\text{disp} * Gauss)(x, y), & \text{for } (x, y) | nMA_{right}(x, y) > N_{threshold} \\ -1, & \text{otherwise} \end{cases} \quad (12)$$

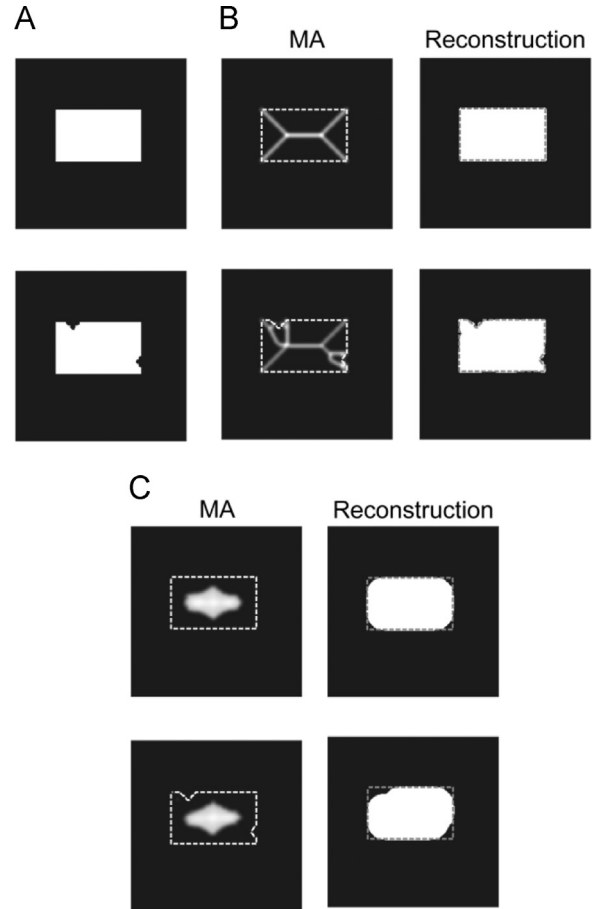


Fig. 2. (A) Stimuli used for the computation of 2D MAs, with and without noise on the contour (bottom and top, respectively). (B) The engineering MAs (left) and their reconstructions (right). Dotted lines indicate the object contour (shown for presentation purposes). The two engineering MAs were different, with a correlation coefficient of 0.77. The reconstructed images were accurate with reconstruction errors of 0.05 for both stimuli. (C) The biological MAs (left) and their reconstructions (right), with (bottom) and without (top) contour noise. The two biological MAs were similar with a high correlation coefficient of 0.99. Although the reconstructions were less accurate (errors of 0.09) than those of engineering MAs, reasonable shapes were achieved.

where $N_{threshold}$ indicates the threshold for eliminating unnecessary smoothing. We set $N_{threshold}$ to 0.1; however, the results were similar when the threshold is less than or equal to 0.3. $Gauss(x, y)$ represents the Gaussian for smoothing with $\sigma_x = \sigma_y = 3$ pixels. The optimal size (σ) of the Gaussian could depend on the size of an object. However, our test showed that the size of the Gaussian was relatively insensitive to the results; an enlargement of 50% did not alter the results. The relation between the disparity of a location, $disparity(x, y)$, and the depth of 3D MA, $depth(x, y)$, is given by

$$\begin{aligned} depth(x, y)^2 - \left(\frac{\delta}{disparity(x, y)} + fix \right) * depth(x, y) + fix^2 &= 0, & \text{for } disparity(x, y) > 0, \\ depth(x, y) &= 0, & \text{for } disparity(x, y) = 0, \end{aligned} \quad (13)$$

where f is the focal length (5 cm; 142 pixels) and δ is the distance between the two eyes (8 cm; 227 pixels). fix is the distance between a fixation point and the frontal plane including the eyes. The nearest point of an object was chosen as the fixation point, and its depth was considered zero. The depth of 3D MA in the model is given by

$$MA_{depth}(x, y) = \frac{depth(x, y)}{r}, \quad \text{for } (x, y | disparity(x, y) > 0), \quad (14)$$

where r represents the ratio between the size of the real object and its projection onto the retina (image).

To evaluate the model, we reconstructed the shape from the computed 3D MA, as described in Section 3. For the reconstruction, we needed the distances between the MA and the surrounding contours as well as the location of the MA. Because the model does not compute the distances, we preserved the distances

between the 2D MA and the contour of the object in the right image for the purpose of evaluation. This procedure assures consistency and objectivity in the determination of the distances, and adequately evaluates the location of the MA.

3. Results

We constructed a computational model that generates a 3D MA from the fusion of physiologically plausible 2D MAs, based on a standard energy model [20] that is thought to capture the essential functions of physiological properties in early- to intermediate-level visual areas. We examined whether the model is capable of generating a correct 3D MA, and whether the computed 3D MA is adequate for the reconstruction of a 3D shape. The model has two novel characteristics: (1) the 2D MA is defined as a set of points that are nearly equidistant from surrounding contours, thus, our 2D MA has a spatial extent, unlike a skeleton as defined in engineering; (2) we detected the binocular disparities between such “fat” 2D MAs using an energy model. We performed the simulations of the model with a variety of 3D objects that included distinct features of shape. Firstly, we present the results of examples with elementary geometric features such as a capsule and a cuboid. Secondly, we present the results for typical geometric features, such as a variation in thickness and a bend, together with other complex features. We also present the results for pairs of real images. For the evaluation of the computed MA, we reconstructed a 3D shape based on the MA, and computed the reconstruction accuracy. To thoroughly test the reconstruction of the 3D shape, we examined the reconstruction error using three criteria: depth from the eyes, 3D shape (relative depth), and the shape of the

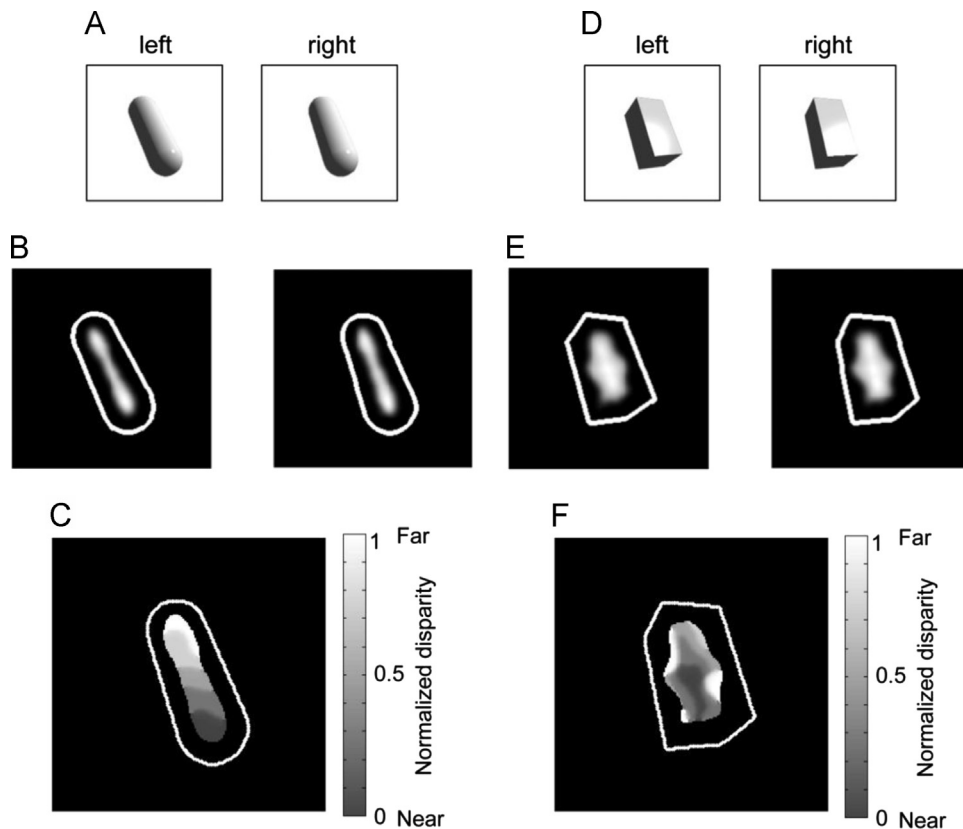


Fig. 3. Computation of 3D-MAs from binocular 2D-MAs ((A)–(C) for a capsule; (D)–(F) for a cuboid). (A) and (D) Two input images for the left and right eyes. (B) and (E) The computed 2D MAs. White lines indicate object contours for presentation purposes, and are not computed by the model. (C) and (F) The 3D MA fused from the binocular 2D-MAs. The disparity computed by the model is plotted in gray. White/dark gray represents a far/near disparity. (A)–(C) The fixation point (depth=0) was set at the bottom end of the major axis, so that the disparity increases toward the top. The computed 3D-MA shows a smooth gradient for the disparity consistent with the ground truth. (D)–(F) The fixation point was set at the nearest corner of the cuboid, so that the disparity increases toward the top. The disparity in the 3D-MA is somewhat complicated because of the sharp corners.

2D projection with respect to the eyes (comparable with the retinal images). We also present the results for testing view invariance of the computed MA.

3.1. The proposed 2D MA

Retinal images of an object can be noisy for various reasons, such that contours of an object might be deformed. However, our visual system is capable of generating a stable percept of the object's shape. The representation of shape in the cortex appears to be robust with respect to noise on the contour. In contrast, the skeletal representation that is used in engineering is sensitive to noise on the contour. An example is given in Fig. 2, which shows MAs and their reconstruction, with and without noise. In Fig. 2A, the top and bottom panels show the rectangles without and with noise, respectively. Here, we introduced two notches as contour noise. The two engineering MAs for the rectangles with and without noise appear very different (correlation=0.77), as shown in Fig. 2B (left). The change in the MA structure that is caused by slight noise often produces considerable differences in MAs between the left and right images, which could be a major reason that binocular fusion of the engineering MA is difficult. On the other hand, the physiological MA appears to be stable with respect to noise, and produces a robust structure in the presence of noise.

To demonstrate the insensitivity of a physiologically plausible MA with respect to contour noise, we computed the MAs for the same two stimuli used above, with and without noise, and compared the two MAs. Fig. 2C (left panels) shows the computed 2D MAs. The MAs for the object with and without noise were very similar (correlation=0.99), indicating that the physiologically plausible MA produces a stable structure insensitive to noise on the contour. To demonstrate the accuracy in the reproduction of the original image from the physiologically plausible 2D MAs, we reconstructed the object shape from the computed MA. The reconstruction was conducted by placing circles for all points on the MA, with the radius of the circles equal to the distance to the nearby contour [16]. Fig. 2C (right) shows the reconstructed shape from the computed physiologically plausible MAs. Although the reconstruction is not as ideal as that from the engineering MAs (the reconstruction errors (see [16] Eq. (10)) for the engineering MAs were 0.05 regardless of noise), the rough shape appears to be reproduced (the errors were around 0.09). The result suggests that the physiologically plausible 2D MA produces a stable structure that is capable of representing an object's shape with robustness.

3.2. The generation of 3D MAs for elementary shapes

The computed 2D and 3D MAs for a capsule, the simplest shape for representation by a MA, are shown in Fig. 3. Input images for the left and right eyes are shown in Fig. 3A. The computed 2D MAs for each eye is shown in Fig. 3B. We observe a rod-like MA elongated along the major axis of the capsule. The 2D MA for the left eye appears slightly tilted compared with that for the right eye, indicating that the top side (in 2D image) of the capsule is farther than the bottom side. We set the fixation point (depth=0) at the bottom end of the major axis, such that the disparity increases toward the top side. The computed 3D MA is shown in Fig. 3C. We observed a rod-like MA with its depth increasing toward the top side, showing agreement with the shape of the capsule.

We computed 2D and 3D MAs for a cuboid, which is another elementary shape with sharp corners (Fig. 3D). The computed 2D MA for the cuboid is shown in Fig. 3E. Similarly to the capsule, the tilt of the 2D MAs (Fig. 3E) is slightly different between the left and right images (the left MA is more tilted). We set the fixation point (depth=0) at the nearest corner of the cuboid, such that the

disparity increases toward the top side. The computed 3D MA is shown in Fig. 3F. We observe a vase-like MA with its depth increasing toward the top side, showing agreement with the shape of the cuboid. These results show that the model computed reasonable 3D MAs for elementary shapes with simple structure.

3.3. Evaluation by the reconstruction of 3D shape for elementary shapes

To evaluate the adequacy of the computed 3D MA, we reconstructed a 3D shape based on the 3D MA, and computed how accurately the computed 3D MA is capable of reproducing a 3D shape in terms of its depth and shape. For the reconstruction, we needed the distances between the MA and the surrounding surface, as well as the location of the MA. The model focuses on the location of the MA, and it does not determine the distances to the surface. For the 3D reconstruction, we used the Euclidean distance between the MA and the nearest contour that is stored separately from the model, as described in Section 2. We reconstructed the 3D shape by placing a number of overlapping spheres along the 3D MA. The centers of the spheres were aligned with the MA, and the radii were set equal to the distance to the nearby contour.

We evaluated quantitatively the accuracy of the reconstruction in terms of depth and shape. The reconstruction error for *depth* was defined as the difference in the depth maps between the original and the reconstruction

$$Error_{depth} = \sqrt{\frac{\sum_{x,y} (D_{original}(x,y) - D_{reconstruct}(x,y))^2}{\sum_{x,y} D_{original}(x,y)^2}},$$

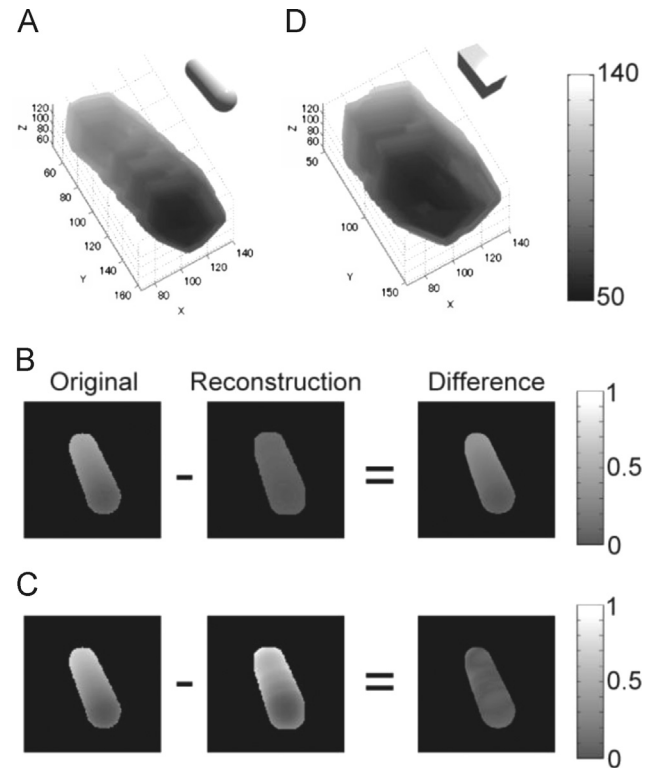


Fig. 4. (A) Reconstructed shape of a capsule. The reconstruction was given by the superposition of overlapping spheres onto the computed 3D MA. For details, see Section 2. The x–y axes represent the plane projected onto a camera. The z-axis and gray represent depth (a larger value indicates farther away). (B) Evaluation of the difference in *depth* between the original and the reconstruction of the capsule. The right panel shows the difference in gray (between 0 and 1). The overall error for depth was 0.78. (C) Evaluation of the difference in *shape* (relative depth) between the original and the reconstruction of the capsule. The error for shape was 0.16. (D) Reconstructed shape of a cuboid.

for $(x, y) \mid D_{\text{original}}(x, y) \cap D_{\text{reconstruct}}(x, y) \neq \emptyset$, (15)

where $D_{\text{original}}(x, y)$ and $D_{\text{reconstruct}}(x, y)$ represent the depth map of the original and reconstruction, respectively. The depth map indicates the distances of all points on the object surface from the eye. This index computes the difference in depth for all points where the original and the reconstruction overlap. To evaluate the reconstruction of *shape*, we introduced an index, $Error_{\text{shape}}$, which was defined by the normalization of $Error_{\text{depth}}$ to the maximum depth within each map. This normalization cancels out the absolute depth so that shape (or relative depth) is evaluated. Note that $Error_{\text{shape}}$ estimates the shape of the front side, not the overall 3D shape, because the model does not estimate the back side of an object. These error indices become zero when the reconstruction is perfect (equal to the original), and one when the reconstruction is twice as large as the original.

Fig. 4 shows the reconstruction of the two elementary shapes, the capsule and cuboid. The columnar shape and the rounded ends of the capsule were reconstructed smoothly (Fig. 4A). Fig. 4B shows the difference in depth from the viewing point. The overall difference, $Error_{\text{depth}}$, was 0.78. As we discuss later, the error appears to be caused by the simplification of the energy model in which only one and three channels are provided for spatial frequency and orientation, respectively. Fig. 4C shows the difference in shape (relative depth). The overall difference, $Error_{\text{shape}}$, was 0.16, indicating that the model successfully reproduced the shape with rounded surfaces. Further evaluation of the errors is discussed in the next section. Because the shape of the capsule is

composed of a set of spheres, it was expected that the reconstruction from overlapping spheres along the 3D MA would reproduce the shape of the capsule with high accuracy. A cuboid with sharp corners was expected to be difficult for the model. Fig. 4D shows the reconstruction of a cuboid. Although the reconstructed shape is somewhat rounded compared with the original cuboid, we can still observe corners that are a crucial feature of a cuboid. $Error_{\text{depth}}$ for the cuboid was 0.79, indicating a level of accuracy similar to the capsule. The reconstruction of the surface was fairly successful with $Error_{\text{shape}}$ of 0.52. These results indicate that the 3D MA computed by the model is fairly capable of representing the shape of objects with elementary shapes.

3.4. Evaluation of 3D MA for shapes with typical features

To investigate the accuracy of the model for the representation of 3D shape, we performed simulations of the model with a variety of objects with distinct features. In this section, we report in detail the results of three examples with typical features: (i) a shape with varying thickness along its major axis, (ii) a shape with a curved axis, and (iii) a combination of multiple features. Overall evaluation of the model for various shapes is discussed in the next section.

Fig. 5A shows the results of a vase whose radius varies along the major axis. The reconstruction from the computed 3D MA shows the depth increasing along the major axis from the center to both ends, indicating successful reproduction of the crucial features of the vase. $Error_{\text{depth}}$ and $Error_{\text{shape}}$ were 0.57 and 0.62,

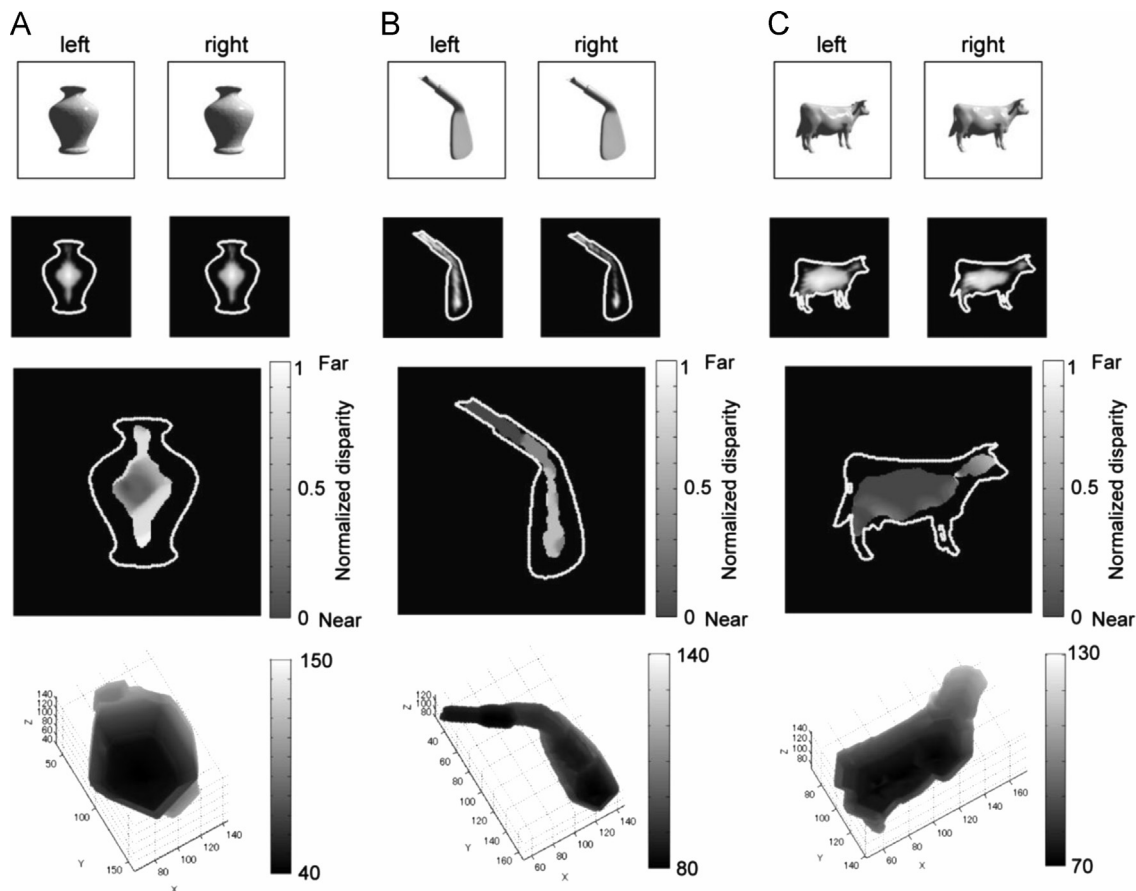


Fig. 5. Simulation results for the stimuli with typical features. The top row shows the binocular stimuli for a vase (A), a golf club (B), and a cow (C). The second row shows the computed 2D-MAs. The third row shows the computed 3D MAs. Conventions are the same as in Fig. 3. The bottom row shows the reconstructed shapes from the 3D MAs. Conventions are the same as in Fig. 4. The errors for depth were 0.57 (A), 0.85 (B), and 0.62 (C), and the errors for shape were 0.62 (A), 0.64 (B), and 0.68 (C).

respectively. The vase was expected to be easy for the model, similarly to the capsule, because both surfaces are rounded. However, the reconstruction errors were still larger than the cuboid that consists of flat surfaces and sharp corners. This large error is attributable to the failure of the reproduction along the top and bottom of the vase where surfaces splay out. This change is barely detected by the disparity in the contours of the vase (the boundary between the vase and background) that extends horizontally at the top and bottom ends. Because the overall surface (except for both the ends) was reproduced successfully, the results indicate that the 3D MA computed from the model is capable of representing a shape with varying thickness along the axis.

Fig. 5B shows the results for a golf club that contains a bent axis and a flat surface. In both the reconstruction and in the computed 3D MA, we can observe a sharp bend at the middle of the head of the club. The depth of the reconstruction increases from the bottom end toward the top end, which is consistent with the structure of the original shape. $Error_{depth}$ and $Error_{shape}$ were 0.85 and 0.64, respectively, similar to the range for other stimuli. The major cause of the error was the flatness of the club head. As discussed with the cuboid, we reconstruct objects using spheres along the axis, so that the reconstruction of a flat surface is difficult. These results indicate that the computed 3D MA is capable of representing an object shape that contains a sharp bend along the major axis.

Fig. 5C shows the results for a cow that has a complex structure. The head and body of the cow appear to be reproduced smoothly and successfully. The values of $Error_{depth}$ and $Error_{shape}$ were 0.62 and 0.68, respectively. The failure of the reconstruction of the legs was a major source of the error. Because the present model has only a single frequency, small parts are disregarded. The results show that the model is capable of representing a complex structure with an error similar to that of simple structures, which is consistent with the characteristics of the human visual system. This result supports the robustness of the model in its representation of shape.

To examine the representation of shape from real images (not created by CG), we conducted a simulation of the model using stereo photographs that may include a variety of noise. A pair of convergent stereo images of a miniature duck was taken, as shown in Fig. 6A, and used as an input stimulus. The fixation point was set at the center of the duck's chest. Fig. 6B–D shows the results of the simulation. The depth of the computed 3D MA, as shown in Fig. 6C, increases as it diverges from the center of the chest. Fig. 6D shows the reconstruction in which the shape of the head and body of the duck appear to be reasonably reproduced. These results suggest that the model is capable of generating a reasonable 3D MA from real images.

3.5. Overall evaluation of the reconstruction error

We evaluated quantitatively the accuracy of the reconstruction in terms of depth and 3D shape. The reconstruction error for depth, $Error_{depth}$, represents how accurately the absolute depth is reproduced by taking the difference between the depth maps of the reconstruction and the original, and the error for shape, $Error_{shape}$, represents how accurately 3D shape is reproduced by canceling out the absolute depth. We reconstructed the 3D shape from the 3D MA for a variety of objects, in addition to those with typical shapes as shown above. The eight input stimuli, the computed 3D MAs, and the reconstructions of shape are shown in Fig. 7. The $Error_{depth}$ and $Error_{shape}$ for all objects (including those shown in the previous sections) are shown in Table 1. The mean and SD of the depth error were 0.69 and 0.13, respectively, indicating that the capability of the model to represent 3D depth is

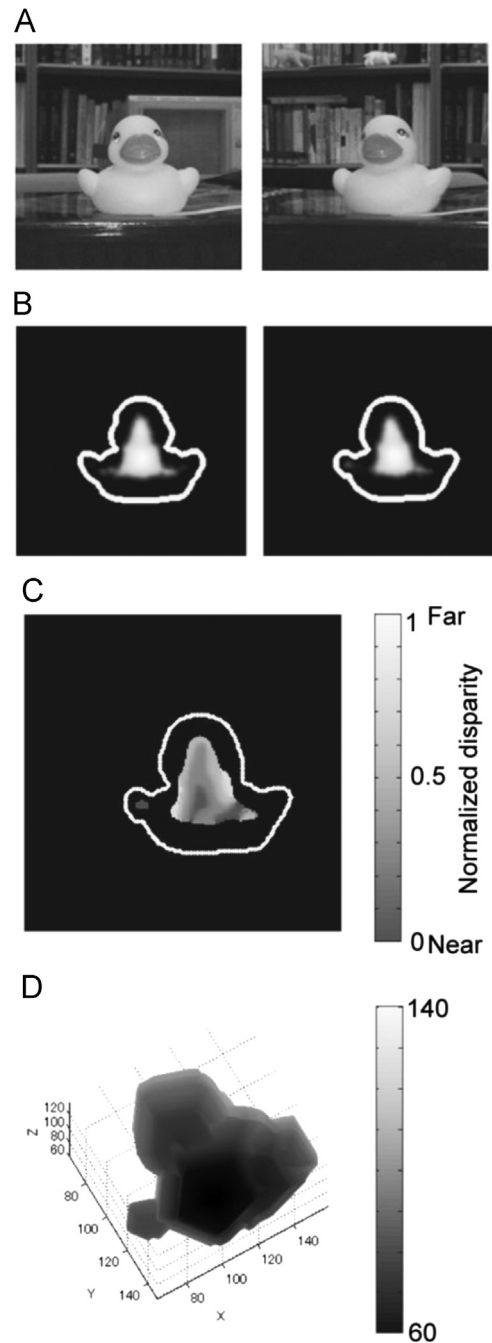


Fig. 6. Simulation results for real stereo images of a duck. Conventions are the same as in Fig. 5. (A) The images of the duck taken using a stereo camera with the fixation point set at the center of the front of the body. (B) The computed 2D MAs. (C) The 3D MA obtained from the binocular 2D-MAs. The computed depth increases as it departs from the center of the front of the body. (D) The reconstructed shape from the 3D MA. The head and body of the duck are visible.

relatively independent of the complexity of the shape and structure of the object. The mean and SD of the shape error were 0.70 and 0.34, respectively. It appears that low errors were observed for the objects whose surface is smoothly rounded or relatively simple when viewed from the designated eye position. The duck showed the worst error among these objects, because the width of its neck differed between the left and right images so that the shape of their 2D MAs was very distinct; this discrepancy caused the failure of binocular fusion leading to an inaccurate representation of depth in the 3D MA. These results indicate that the proposed

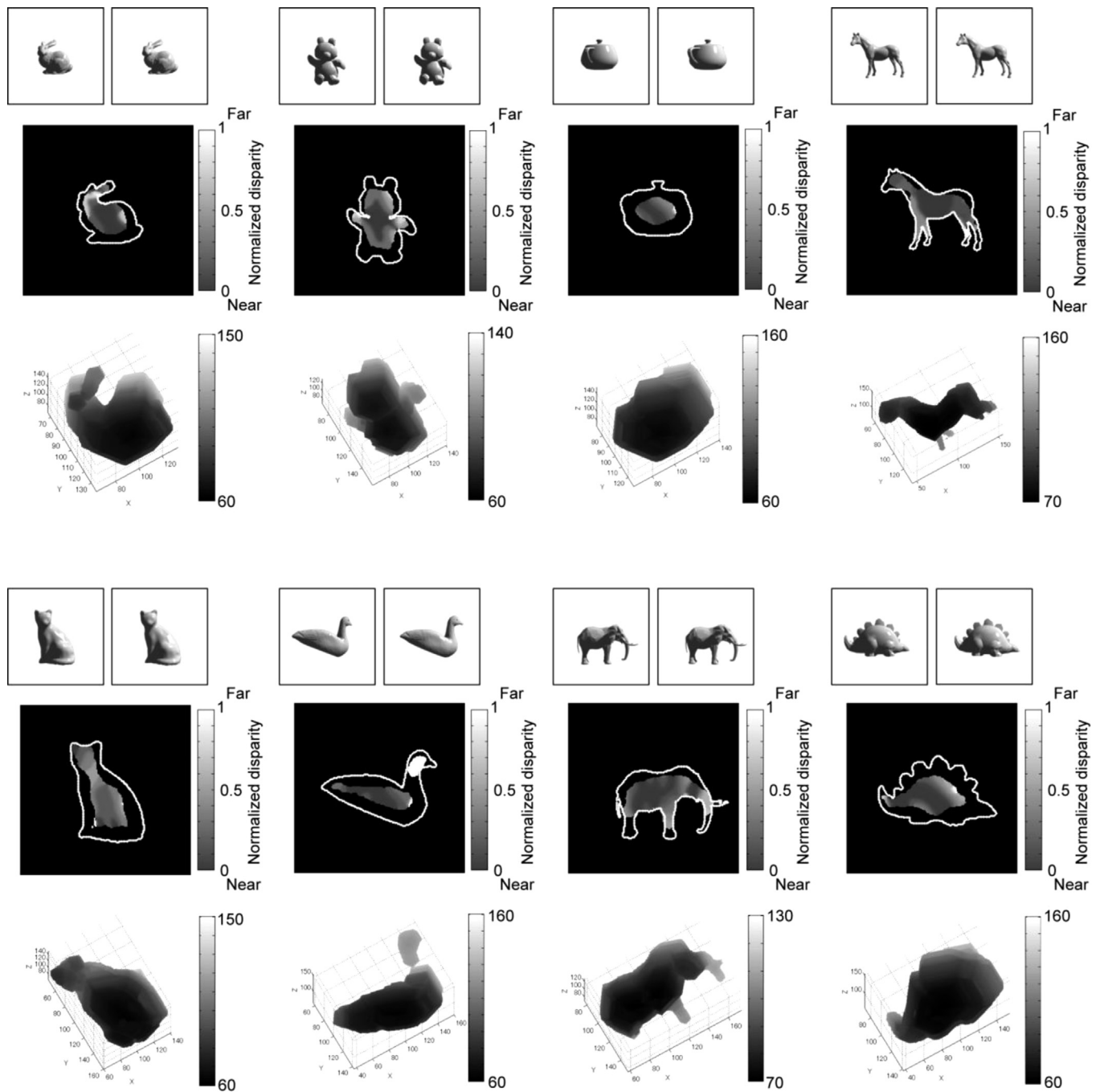


Fig. 7. Simulation results for other stimuli such as a rabbit, a bear, a pot, a horse, a cat, a duck, an elephant, and a stegosaurus. Conventions are the same as in Fig. 5. The errors for reconstruction are summarized in Table 1. All shapes were reasonably reconstructed, including those with complex shapes.

model is capable of representing the rough shape of various 3D objects. Given the limited number of frequency and orientation channels (1 and 3 for frequency and orientation, respectively), the reproduction should be considered surprisingly successful.

3.6. Evaluation by the reconstruction of 2D stimulus (frontal projection)

As an evaluation of the internal representation of the model, we examined the capability of the model to reconstruct the original input stimulus from the computed 3D MA. We defined the error in 2D projection, $Error_{2D}$, as an index to indicate how

accurately the model is capable of reproducing the 2D shape

$$Error_{2D} = \frac{|S_{original} - S_{reconstruct}|}{S_{original}}, \quad (16)$$

where $S_{original}$ and $S_{reconstruct}$ indicate the surface areas that are projected onto an eye (the 2D area seen from a viewing point) of the original and reconstruction, respectively. This index is important in that it shows the capability of the model to reconstruct the original stimulus from the internal representation of the model. The index becomes zero when the reconstruction is perfect, and one when the reconstruction is twice as large as the original. The errors of all objects are shown in Fig. 8. The mean and SD of the error were 0.14 and 0.05, respectively. The error was less than 20%

Table 1
Reconstruction errors in depth and shape for all stimuli.

Stimulus	Depth error	Shape error
Capsule	0.7771	0.1559
Cuboid	0.7879	0.5244
Vase	0.5822	0.6298
Club	0.8469	0.6408
Cow	0.6176	0.6803
Rabbit	0.5861	0.6577
Bear	0.7205	1.0237
Pot	0.7412	0.4924
Horse	0.7412	0.6748
Cat	0.6860	0.7660
Duck	0.6173	1.5153
Elephant	0.9318	1.0348
Stegosaurus	0.4262	0.4011
Mean	0.6891	0.7043
SD	0.1325	0.3368

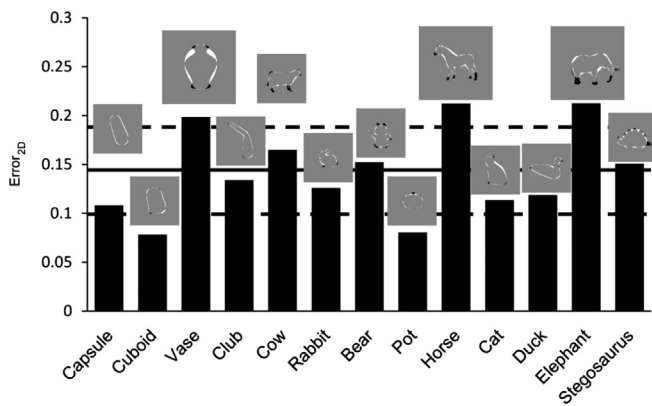


Fig. 8. Evaluation by the reconstruction of 2D images (surface area). The differences in the surface areas between the original and reconstructed shapes ($Error_{2D}$) are plotted. The surface area was determined by projecting a 3D shape onto a camera. The solid line and dotted lines indicate the mean and SD of the errors, respectively. The errors were less than 20% except for the horse and elephant. The over- and under-estimation of the areas are shown in the insets by white and black, respectively. Overestimation is often observed around concave contours, whereas underestimation occurs around small parts.

Table 2
Reconstruction errors in 2D projection for all stimuli.

Stimulus	$Error_{2D}$	Overestimated	Underestimated
Capsule	0.1084	0.1084	0
Cuboid	0.0785	0.0703	0.0083
Vase	0.1986	0.1635	0.0351
Club	0.1341	0.1224	0.0117
Cow	0.1651	0.0700	0.0951
Rabbit	0.1260	0.1015	0.0245
Bear	0.1522	0.0713	0.0809
Pot	0.0806	0.0612	0.0194
Horse	0.2124	0.1347	0.0778
Cat	0.1136	0.0962	0.0174
Duck	0.1188	0.1066	0.0121
Elephant	0.2126	0.1101	0.1025
Stegosaurus	0.1507	0.0913	0.0594
Mean	0.1424	0.1006	0.0419
SD	0.0451	0.0291	0.0362

for most of the objects except the horse and the elephant whose legs were too thin to be reproduced. We also calculated separately the errors for the over- and underestimation of the areas (the

positive and negative parts of the index). The results are shown in white and black in the insets of Fig. 8, and the values are given in Table 2. The model appears to show overestimation where the contour of an object is concave, and underestimation where the part is small. Because the shape is reconstructed by superimposing spheres, concave regions tend to be masked by the spheres (overestimated). Small parts are often missed because the present model consists of a single spatial frequency channel. If multiple frequency channels were provided, the model would be capable of detecting these small parts and avoid underestimation. Multiple frequency channels would also be helpful in reducing the overestimation caused by concave surfaces. These results support the capability of the model to represent object shape.

3.7. View invariance of the reconstruction

IT neurons that are selective for 3D MA showed view-invariant responses [9]. The human visual system also shows view invariance in its perception of object shape, although the reaction time often varies. It is expected that view invariance is an inherent characteristic of the representation by the MA. Here, we evaluated whether our model reproduces view invariance in the reconstruction error. We computed the 3D MA and the reconstruction error for a series of images viewed from distinct directions. Specifically, we used the stimuli of a cow viewed from its side, tail, and an in-between position. The input stimuli are shown in Fig. 9 (generated by rotating the cow shown in Fig. 5C), together with the computed 3D MAs and the reconstructions. The head and body of the cow were reproduced in all views, although mostly its thin legs were not. The error in depth, $Error_{depth}$, for each view was 0.54 (Fig. 9A), 0.40 (Fig. 9B), and 0.62 (Fig. 9C), and the mean of the three was 0.52. The error in shape, $Error_{shape}$, for each view was 0.56 (Fig. 9A), 0.75 (Fig. 9B), and 0.83 (Fig. 9C), and the mean of the three was 0.71. Both errors in depth and surface show small variation: all views show errors that are within 10% of the means. This result shows that the model is capable of reproducing shapes from a variety of viewpoints with similar amounts of error. This view invariance is consistent with the characteristics of IT neurons tuned to a 3D MA configuration, and also human vision.

4. Discussion

Numerous studies have suggested object-centered coordinates for the cortical representation of shape [1–3]. Although theoretical studies have favored the advantages of the MA representation for more than three decades, only a few physiological studies have reported supportive results [13]. Recently, an electrophysiological study has provided direct evidence that neurons in IT show selectivity for the 3D MA configuration [9]. However, the mechanisms by which the 3D MA is constructed through the visual pathway have not been clarified. The present study examined neural processes for the generation of a 3D MA. A physiological study has reported that neurons in V1 respond to the medial region of a textured figure [13]. Such a response in V1 could be produced by the synchronization of BO-selective neurons in V2, and the 2D MA has been reported to be capable of coding object shape [16]. We focused on the transformation of the 2D MA reported in V1 into the 3D MA observed in IT. The latency of V1 cells that respond to the edges of an object range between 40 and 60 ms [13,24], and that to 2D MA range between 90 and 110 ms [13,24]. The onset latency of IT cells is generally more than 90 ms [25] and the latency for 3D MA is considered to be much longer than 90 ms. Given this time constraint, afferent connections appear to play a crucial role in the transformation from 2D MAs to a 3D MA, probably in combination with efferent connections.

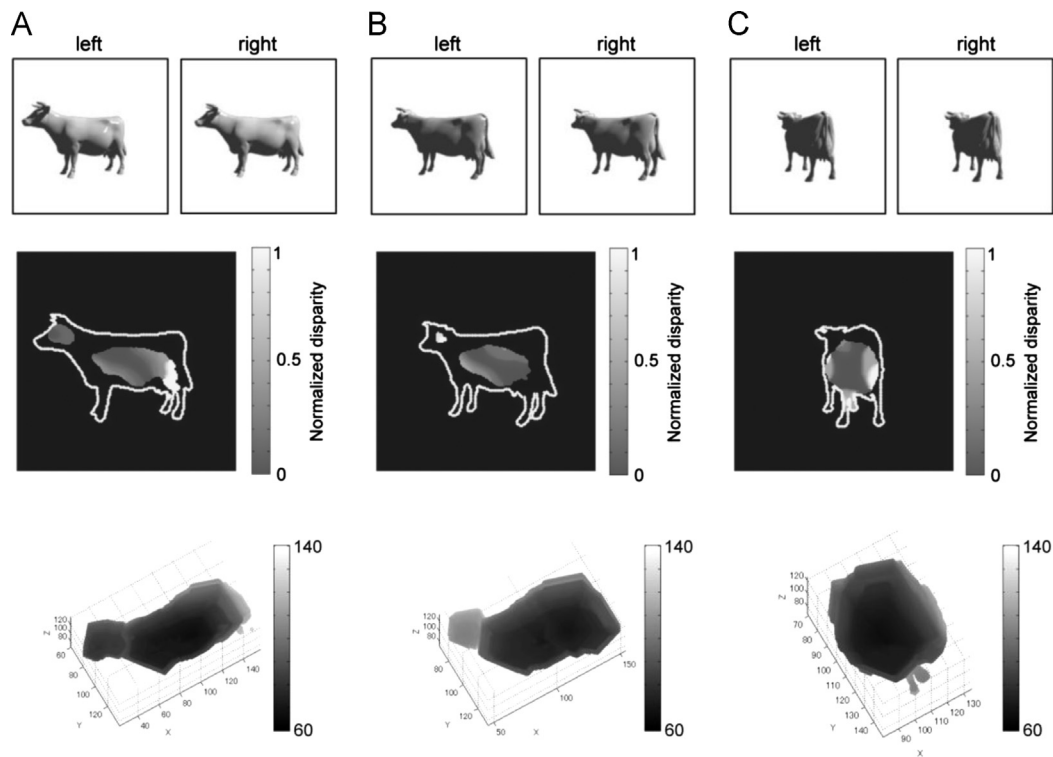


Fig. 9. View invariance of the reconstruction. Conventions are the same as in Fig. 5. (A)–(C) The simulation results for a cow viewed from three different directions. The top row shows the stereo stimuli that were viewed from distinct points. The middle row shows the 3D MAs computed from the binocular 2D MAs. The bottom row shows the reconstructed shapes computed from their 3D MAs. The depth errors were 0.54 (A), 0.40 (B) and 0.62 (C). The shape errors were 0.56 (A), 0.75 (B) and 0.83 (C). Both types of error in the reconstruction show small variation, indicating that the model is capable of reproducing reasonable shapes regardless of viewpoint.

Therefore, the present study investigated the generation of a 3D MA by the binocular fusion of 2D MAs. Note that the present model does not account for these latencies. It is expected to further study the temporal properties of the representation of 3D shape.

In the process of binocular fusion, it is crucial to determine whether the responses of V1 cells to MA are monocular or binocular. If the MA is monocular, a retinal image of an object is transformed into a monocular 2D MA by V1 cells, and then the fusion of a binocular pair of 2D MAs generates a 3D MA based on the disparity between the 2D MAs. On the other hand, if the MA in V1 is binocular, a local segment of the 3D MA is produced from a binocular pair of local contours of the object image, and the integration of the 3D MA segments generates a global 3D MA in IT. Consider the case where a contour of one side of an object is nearer than the fixation point, and that of the other side is farther. In the monocular case, the disparity-selective cells fuse a pair of 2D MAs based on the disparity between the axes to generate a 3D MA that represents correct depth. On the other hand, in the binocular case, the fusion of a binocular pair of local contours would be extremely difficult because the fusion requires V1 cell that is tuned to near on one side and far on the other side, and that signals depth at the middle of the two. An alternative would consider feedback from higher cortical areas to V1. Because disparity-selective cells in V1 detect local disparity and the higher cortical regions are required to produce global depth, a higher region such as V4 and IT would generate 3D contours and send feedback to generate local, binocular 2D MAs in V1. Although feedback may play an important role, an assumption of such complex pathways prevents the construction of a plausible computational model. In the present study, we focused on the monocular case, and proposed the hypothesis that a pair of 2D MAs that encode monocular projections of object shape is fused to generate

a 3D MA, as a first step toward understanding the transformation of MA from V1 to IT.

We defined the physiologically plausible 2D MA to mimic the activities of V1 cells responding to 2D MA. The physiologically plausible 2D MA is capable of representing the outline of an object with around 10% error. A major downside of MA representation in general could be high sensitivity to noise on contours. In the real world, a retinal image of an object often includes noise on contours for a variety of reasons. In fact, an engineering MA that is defined by a set of axes (skeleton) often changes considerably in response to noise, so that even the graph structure that represents the object shape varies. Given that the visual system is able to perceive shape with stability and robustness in the presence of noise, the engineering MA may not be a suitable candidate for cortical representation. In the present study, we propose that the physiologically plausible MA overcomes this disadvantage. To reproduce V1 responses to 2D MA, we defined the physiologically plausible MA as having an equal distance between the point under examination and nearby contours. Specifically, we computed the equality of the distances from the contours, and determined the region of the MA by setting a threshold for equality. Because of this processing, the MA is defined by a set of points whose distances from the contours are similar, but not exactly equal, giving it robustness with respect to noise on contours. Therefore, the physiologically plausible MA is inherently more robust than the engineering MA, at the expense of accuracy. The physiologically plausible MA appears meaningful in terms of its stability and robustness.

The present model uses a standard energy model [23] to determine the binocular disparities of physiologically plausible 2D MAs. The disparity at each location along the axis is detected by using a winner-take-all mechanism among disparity-selective cells, each of which is tuned to a distinct disparity. For the sake

of simplicity, the model has only a single spatial frequency channel and three orientation channels. Therefore, the accuracy for disparity detection is very limited, and much lower than that of the visual system. It should be noted that a model with this simple structure is capable of generating a 3D MA whose disparity varies reasonably according to the original shape, and reproduces the overall form of the original object. These results support the plausibility of binocular fusion of physiologically plausible MAs using the energy model.

We constructed the model for the generation of a 3D MA based on the binocular fusion of physiologically plausible 2D MAs, and examined whether this model is suitable for the representation of 3D shape. We computed a 3D MA from a number of objects, with a variety of shape characteristics, including natural objects. The model was capable of generating a reasonable 3D MA for a wide range of objects. We also reconstructed the 3D shape of the test objects based on the computed 3D MA. The model showed excellent reconstruction accuracy for somewhat rounded objects such as a capsule, and reasonable accuracy for all other objects including those with sharp corners, flat surfaces, and complex structures. Given the limited number of frequency and orientation channels, the reproduction should be considered as surprisingly successful. Furthermore, the simulation results showed view invariance in the reconstruction, which is consistent with the results of physiological experiments [9]. These results show that a model based on the fusion of a binocular pair of physiologically plausible 2D MAs generates a reasonable 3D MA with robustness in representing the 3D structure independent of viewpoint, indicating the plausibility of the model as a candidate for the cortical computation of 3D MA.

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Appendix A

Gabor_{left} and **Gabor_{right}** represent the oriented receptive field of the model simple cells for the left and right images, respectively

$$\text{Gabor}_{\text{left}} \theta, \phi_i(x, y) = \frac{1}{2\pi} \cos \left(2\pi \frac{(x-x_0) \cos(\theta) - (y-y_0) \sin(\theta)}{\lambda} + \phi_i \right) * e^h, \quad (17)$$

$$\text{Gabor}_{\text{right}} \theta, \phi_i, \psi_j(x, y) = \frac{1}{2\pi} \cos \left(2\pi \frac{(x-x_0) \cos(\theta) - (y-y_0) \sin(\theta)}{\lambda} + \phi_i + \psi_j \right) * e^h, \quad (18)$$

where

$$h = - \left[\left\{ \frac{(x-x_0) \cos(\theta) - (y-y_0) \sin(\theta)}{\sigma_x} \right\}^2 + \left\{ \frac{(x-x_0) \sin(\theta) + (y-y_0) \cos(\theta)}{\sigma_y} \right\}^2 \right],$$

where x_0 and y_0 represent the center of the Gabor filters, and θ , λ , σ_x , and σ_y show the orientation, wavelength and SDs of the Gabor filters, respectively. ϕ_i represents the phase of the left receptive field, and ψ_j represents the ocular difference in phase. We set λ , σ_x and σ_y to 20, 8 and 8 pixels, respectively, so as to mimic V1 cells (λ and $\sigma_x(\sigma_y)$ equal to 0.5° and 0.2° in visual angle, respectively).

ϕ_i ($i=1-4$) were set to 0, $\pi/2$, π and $3\pi/2$. ψ_j ($j=1-11$) ranged between 0 and 2π in increments of $\pi/11$. The size (spatial extent) of the Gabor filter was set to 40×40 pixels ($1 \times 1^\circ$) so as to be consistent with that of the receptive field of V1 neurons [25].

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