Neural computation of surface border ownership and relative surface depth from ambiguous contrast inputs

Birgitta Dresp, Stephen Grossberg

To cite this version:

Birgitta Dresp, Stephen Grossberg. Neural computation of surface border ownership and relative surface depth from ambiguous contrast inputs. Frontiers in Psychology, 2016, 10.3389/fpsyg.2016.01102. hal-01202930

HAL Id: hal-01202930
https://hal.science/hal-01202930
Submitted on 22 Sep 2015

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Neural computation of surface border ownership
and relative surface depth from ambiguous contrast inputs

Birgitta Dresp-Langley
ICube UMR 7357 CNRS and University of Strasbourg
2, rue Boussingault
67000 Strasbourg
FRANCE

Stephen Grossberg
Center for Adaptive Systems
Graduate Program in Cognitive and Neural Systems
Center for Computational Neuroscience and Neural Technology
Department of Mathematics
Boston University
677 Beacon Street
Boston, MA, 02215
Abstract

The segregation of image parts into foreground and background is an important aspect of the neural computation of 3D scene perception. To achieve such segregation, the brain needs information about border ownership; that is, the belongingness of a contour to a specific surface represented in the image. This article presents psychophysical data derived from 3D percepts of figure and ground that were generated by presenting 2D images composed of spatially disjoint shapes that pointed inward or outward relative to the continuous boundaries that they induced along their collinear edges. The shapes in some images had the same contrast (black or white) with respect to the background gray. Other images included opposite contrasts along each induced continuous boundary. Results show that figure-ground judgment probabilities in response to these ambiguous displays are determined by the orientation of contrasts only, not by their relative contrasts, despite the fact that many border ownership cells in cortical area V2 respond to a preferred relative contrast. The FACADE and 3D LAMINART models are used to explain these data.

Keywords: figure-ground separation; border ownership; perceptual grouping; surface filling-in; bipole cells; surface contours; V2; V4; FACADE theory; 3D LAMINART model
Introduction

The non-ambiguous perceptual organization of planar visual images into figure and ground requires the visual system to be able to generate a three-dimensional (3D) representation from a two-dimensional (2D) stimulus input. During viewing of a natural 3D scene, objects that are closer to the viewer may block or occlude the view of objects that are further away. Boundaries of these occluding objects are perceived as belonging to them, a property called *border ownership*. Because occluding objects occur closer in depth than the objects that they occlude, border ownership in response to a 3D scene typically coexists with a percept of being closer in depth. The importance of surface border ownership to what may seem nearer to us was already noticed by Galileo (see the review by Dresp-Langley, 2014). The borders of occluding surfaces generally occur in the foreground, while the borders of occluded surfaces generally occur in the background.

An important problem in visual perception concerns how border ownership assignment occurs in response to 2D pictures, and what role it may play in determining 3D percepts of such pictures. In response to 2D pictures, there are famous examples where the perceptual assignment of surface borders to 3D percepts of foreground and background may be reversible, leading to totally different interpretations of the objects in each representation. (Figure 1). Such spontaneous changes in figure-ground perception occur only under particular circumstances due to competition between multiple, approximately balanced, 3D interpretations of the 2D image.

Figure 1

Von der Heydt and his colleagues have published important data from their systematic series of neurophysiological experiments about the border ownership properties of neurons in cortical area V2 of monkeys. In particular, Zhou, Friedman, and von der Heydt (2000) reported data from neurons in cortical area V2 that tend to respond to borders with different firing rates depending on whether the border is owned by an occluding or an occluded surface. These neurons are often maximally excited by a preferred combination of direction-of-contrast and border ownership. Zhang and von der Heydt (2010) further studied the contribution of individual edges to border ownership assignment by decomposing figural contours into fragments. Fragments on the preferred side-of-figure produced facilitation, while fragments on the opposite side produced suppression of neural responses. Border-ownership signals also persist for about a second in the brain (O'Herron and von der Heydt, 2009; 2011). Border-ownership signals are generally consistent over multiple variations in shape geometry, configuration, and contrast (Qiu and von der Heydt, 2005; Qiu, Sugihara,
Grossberg (2015) has proposed a unified explanation of these data properties using the FACADE (Form-And-Color-And-DEpth) model of 3D vision and figure-ground perception, and its further development and extension by the 3D LAMINART laminar cortical model, which together have explained and predicted many data about how the brain consciously sees 3D surface percepts in response to 2D pictures and 3D scenes (Cao and Grossberg, 2005, 2012; Fang and Grossberg, 2009; Grossberg, 1994, 1997, 1999; Grossberg and McLoughlin, 1997; Grossberg and Pinna, 2012; Grossberg, Srinivasan, and Yazdanbakhsh, 2011; Grossberg and Swaminathan, 2004; Grossberg and Yazdanbakhsh, 2005; Grossberg et al., 2008; Kelly and Grossberg, 2000; Leveille, Versace, and Grossberg, 2010; McLoughlin and Grossberg, 1998; Yazdanbakhsh and Grossberg, 2004).

As noted above, the von der Heydt et al. data show that various neurons in V2 that are sensitive to border ownership also respond with a preferred contrast polarity. The experiments that are reported in the current article show, however, that figure-ground assignments can often be made that are not sensitive to contrast polarity, indeed that the same figure-ground properties can occur even if contrast polarities are mixed, or are switched from one polarity to the opposite polarity, across the stimulus fragments that induce a percept of 3D surfaces. This difference between the current experimental data and those of von der Heydt et al. emphasize that conscious 3D surface percepts arise from mechanisms that include properties of V2 neurons, but also employ other brain mechanisms as well. After summarizing these data, an explanation of them will be offered using the same FACADE and 3D LAMINART model mechanisms that were used to explain the von der Heydt and various other 3D vision and figure-ground perception data.

**Materials and method**

The psychophysical experiments were conducted in accordance with the Declaration of Helsinki (1964). Experimental sessions were run under laboratory conditions of randomized free trial-by-trial image viewing using a Dell PC computer equipped with a mouse device and a high resolution color monitor (EIZO LCD ‘Color Edge CG275W’). This screen has an in-built calibration device which uses the Color Navigator 5.4.5 interface for Windows. The images were generated in Photoshop using selective combinations of Adobe RGB increments to generate contrast inputs (see also Dresp-Langley, 2015). The luminance levels for each
RGB triple could be retrieved from a look-up table after calibration and the values were also cross-checked on the basis of standard photometry using an external photometer and adequate interface software (Cambridge Research Instruments).

Subjects
Ten unpracticed observers, mostly students in computational engineering who were unaware of the hypotheses of the study, participated in the experiments. All subjects had normal or corrected-to-normal visual acuity.

Figure 2

Stimuli
The stimuli (Figure 2) consisted of six images with different edge contrast inputs. The luminance of the background was 50.5 cd/m$^2$ (148,148,148 RGB) in all eight images. The luminance of the black contrast fragments was 1.5 cd/m$^2$ (0,0,0 RGB) and the luminance of the white contrast fragments was 99.5 cd/m$^2$ (255,255,255 RGB), yielding perfectly balanced Weber contrasts ($L_{\text{feature}}/L_{\text{background}}/L_{\text{background}}$) of -0.97 and 0.97 for negative and positive polarities in the six images with the fragmented edge contrasts. The height of the central surfaces was 10 cm on the screen, whereas the width was 12 cm. In the six images with the ambiguous fragmented edge contours, about 50% of the inner surface contour was void of a contrast, so that 50% of the boundary contour had to be completed perceptually.

Task instructions
A classic psychophysical forced choice procedure with three response alternatives was used to measure perceptual decisions for relative depth (figure-ground). Observers were asked to indicate whether the central surface appeared to "stand in front" of", to "lie behind”, or to be in the "same plane" as the surrounding surface. It was made sure that all observers understood the instructions correctly before an experimental session was initiated.

Procedure
Subjects were seated at a distance of 1 meter from the screen and asked to look at the center of the screen. The experiments were run in a dimmed room (mesopic conditions), with blinds closed on all windows. The six images were presented in random order for about one second each, and each image was presented four times in a session. Inter-stimulation intervals were measured. They typically varied from one to three seconds, depending on the observer, who
initiated the next image presentation by striking a key on the computer keyboard. Each individual session consisted of 32 trials per subject.

Figure 3

Experimental Results
The results from the depth judgment experiment (Figure 3) show that the central surface in the images has a high probability to be perceived as figure ("in front") when the local contrast edges of the fragmented contour configurations are inward directed, and a high probability to be perceived as ground ("behind") when the local edges are outward directed. These perceptual decisions do not depend on the contrast sign of the local edges. Analysis of variance (ANOVA) on means for a 2x2 experimental design, with two levels of the factor "contrast edge direction" and two levels of the factor "contrast sign" returned statistically significant effects of "contrast edge direction" on perceptual decisions for "in front" (F(1,1)=228.30, p<.001) and "behind" (F(1,1)=212.77, p<.001), and, as expected, no effect of contrast sign (see also Dresp, Durand and Grossberg, 2002) for neither type of perceptual decision (F(1,1)=2.58, NS and F(1,1)=0.25, NS respectively).

Theoretical Results

Bipole boundary completion can pool over opposite contrast polarities. A unified mechanistic neural explanation can be given of these various percepts using FACADE and 3D LAMINART model mechanisms (Figures 4 and 5). These mechanisms are summarized here with enough detail to achieve a self-contained exposition.

Figure 4
In response to all of the images in Figure 2, boundaries can be completed inwardly between pairs of adjacent colinear inducers. The completion process uses the oriented long-range horizontal cooperation of bipole grouping cells in layer 2/3 of cortical area V2, balanced by shorter-range disynaptic inhibition (Figures 5 and 6a). Bipole cells can complete boundaries in response to colinear inducers with the same relative contrasts with respect to the background, as in the leftmost two columns of Figure 2, as well as between inducers with opposite relative contrasts with respect to the background. This is true because bipole cells receive their inputs, after several stages of additional processing, from complex cells in layer 2/3 of cortical area V1 (Figures 4 and 5). Complex cells, in turn, pool inputs from simple cells in layer 4 of V1 that have the same preferences for position and orientation, but opposite contrast polarities. As a result, bipole cells can complete boundaries around objects that lie in front of textured backgrounds whose relative contrasts reverse along the perimeter of the
object. In the present cases, bipole cells complete rectangular boundaries that abut all their inducers.

**Figure 5**

**Bipoles are sensitive to T-junctions: End gaps.** The long-range cooperation and short-range competition processes whereby bipoles complete boundaries are sensitive to any T-junctions that lie along the boundaries that they complete (Figure 6a). In the images with incomplete boundaries, there are no explicit T-junctions in the image. However, when a rectangular boundary is completed, T-junctions are created at the corners of the colinear inducing contrasts. The bipole cells that lie along the orientation of a completed boundary (the “head” of the T) get more excitatory input than do the bipole cells that lie near the head of the T, but whose orientational preference is along the perpendicular or oblique orientation of the inducing contrast (the “stem” of the T). This is true because the bipole cells that are activated along the head of the T receive strong excitatory inputs from both sides of their receptive fields, whereas the bipole cells that are activated along the stem of the T receive strong excitatory inputs from just one side of their receptive fields (Figure 6a). The more strongly activated bipole cells inhibit surrounding bipole cells more than conversely through a spatially short-range competitive network. As a result, the bipole cells near the head that are along the stem get inhibited. An *end gap* hereby forms in each boundary near where the stem of a T touches its head (Figure 6a).

**Figure 6**

Because the bipole cells can complete rectangular boundaries in response to spatially disjoint inducers with the same relative contrasts with respect to their surrounding regions, or in response to combinations of inducers with opposite relative contrasts, end gaps at the T-junctions can also form in either case.

As originally explained in Grossberg (1994, 1997), and simulated in such articles as Kelly and Grossberg (2000), Grossberg and Swaminathan (2004), and Grossberg and Yazdanbakhsh (2005), end gaps trigger a process of figure-ground perception and border ownership in which the rectangular boundaries are perceived in front of the regions that they enclose, which are themselves perceived as a ground at a slightly further depth. For example, the percepts of the Necker cube (Figure 5b; Grossberg and Swaminathan, 2004) can be explained in this way, as can the way that shifts in attention can make an attended disk in Figure 6c look both nearer and darker (Grossberg and Yazdanbakhsh, 2005; Tse, 2005). These concepts are reviewed and extended below in order to explain the conscious 3D surface percepts that are generated by the 2D images in Figure 2.
In order to motivate this explanation, it is useful to ask the following question: If it is indeed the case that these figure-ground relationships do not depend on having inducers with the same contrast polarity, then why do so many cortical area V2 cells that are sensitive to border ownership also exhibit a particular contrast preference; e.g., Zhou, Friedman, and von der Heydt (2000). This can be understood by going into more detail about how end gaps trigger figure-ground perception and border ownership.

Figure 7

*Feedback between boundaries and surfaces achieves complementary consistency.* The FACADE and 3D LAMINART models (Figures 4 and 5) detail how the figure-ground perception process utilizes feedback between the boundary completion process in the interblob cortical stream and the blob cortical stream within V1, V2, and V4 of visual cortex. This feedback enables boundaries and surfaces to generate a consistent percept, despite the fact that they obey computationally complementary laws. This property is called *complementary consistency.* As will be noted shortly, the mechanisms that ensure complementary consistency also contribute to 3D figure-ground separation.

In particular, the completed boundaries with their end gaps are projected topographically from the interstripes, or pale stripes, of V2, at which boundaries are completed, to the thin stripes of V2, at which one stage of surface filling-in occurs. When surface filling-in occurs within these boundary inducers, brightness and color can flow out of the end gaps, thereby equalizing the filled-in brightnesses and colors on both sides of the remaining boundaries near these gaps (Figure 7). Only the boundary of the rectangle is closed, so only it can fully contain its surface-filling in. However, in these images, the regions both inside and outside the rectangles are surrounded by closed boundaries, since the frame of the image provides another closed boundary that can contain filling-in between it and the bipole-generated rectangular boundary that lies within it. The significance of this fact will be discussed below.

Figure 8

*Closed boundaries, surface contours, and boundary pruning.* As filling-in occurs, feedback can occur from the surfaces in the thin stripes to the boundaries in the interstripes (Figure 8). These feedback signals take the form of *surface contours* that are generated by contrast-sensitive on-center off-surround networks that act across position and within the depth represented by each filling-in domain. These contrast-sensitive networks sense sufficiently sharp spatial discontinuities in the filled-in brightnesses or colors within their filling-in domain. They hereby generate surface contour output signals only at the surface positions that
are surrounded by closed boundaries. In response to the incomplete inducers in the top row of Figure 2, these regions lie on both sides of the completed boundaries. However, due to the end gaps, surface contour signals are not generated at the boundary positions of the inducers themselves.

The surface contour output signals hereby generate topographic feedback signals to a subset of the boundary representations that induced them (Figure 8). These feedback signals are delivered to the boundary representations via an on-center off-surround network whose inhibitory off-surround signals act within position and across depth (Figure 8). The on-center signals strengthen the boundaries that generated the successfully filled-in surfaces at the same depth, whereas the off-surround signals inhibit spurious boundaries at the same positions but further depths. This inhibitory process is called *boundary pruning*. Surface contour signals hereby strengthen consistent boundaries and prune, or inhibit, redundant boundaries.

Because surface contour signals are generated by the contrasts of a filled-in surface, they are sensitive to a particular contrast, and not to the opposite one. Their feedback to boundaries thus makes the responses of the recipient bipole cells also sensitive to this contrast, even though the bipole cells, in the absence of surface contour feedback signals, respond to both contrast polarities, due to their inputs from V1 complex cells, so that they can complete boundaries of objects in front of textured backgrounds. Thus, after surface contour signals act, their target bipole cells also exhibit sensitivity to a particular contrast polarity, as in the data of Zhou et al. (2000).

In response to 3D scenes, boundary pruning is part of the process of *surface capture* whereby feature contours can selectively fill-in visible surface qualia at depths where binocular fusion of object boundaries can successfully occur, thereby contributing to the formation of closed boundaries that can contain the filling-in process. Surface contour and boundary pruning signals hereby work together to generate 3D percepts based on successfully filled-in surface regions.

For example, the open boundary at Depth 2 in V1 and the V2 pale stripes of Figure 8 can be created due to a monocularly viewed vertical boundary that is seen by only one eye, as occurs during daVinci stereopsis (Cao and Grossberg, 2005; Gillam, Blackburn, and Nakayama, 1999; Nakayama and Shimojo, 2000), and by a pair of horizontal boundaries that do not give rise to strong binocular disparities. Such depth-nonselective boundaries are projected to all depth planes along the line of sight (Cao and Grossberg, 2005; Grossberg and Howe, 2003). The closed boundary at Depth 1 in Figure 8 is due to these boundaries plus a left vertical boundary that is formed at that depth due to binocular disparity matching between
the two eyes. As a result of surface filling-in within V2 thin stripes and the formation of surface contours only at Depth 1, the closed boundary at Depth 1 is strengthened, whereas the spurious open boundary at Depth 2 is inhibited.

**Figure 9**

*From boundary pruning to figure-ground separation.* Remarkably, by eliminating spurious boundaries, the off-surround signals that are activated by surface contours also enable figure-ground separation to proceed. They do so by separating occluding and partially occluded surfaces onto different depth planes, after which partially occluded boundaries and surfaces can be amodally completed behind their occluders. For example, the three rectangles in Figure 9a are perceived as a vertical rectangle in front of a partially occluded horizontal rectangle. Due to the action of surface contours, the redundant copy of the vertical rectangle at a further depth (denoted by D2 in Figure 9a) is inhibited, thereby enabling the horizontal boundaries corresponding to the smaller rectangles to be colinearly completed within depth D2. In response to the picture in Figure 9b, the redundant vertical rectangular boundary is inhibited at depth D2, thereby restoring the boundary fragments at depth D2 that previously were inhibited by the D2 vertical boundaries at end gaps. That is why end gaps are not seen in the final depthful percept.

*How the disparity filter eliminates some spurious boundaries in the near depth.* Although the boundaries that contained end gaps are eliminated by surface contours at the further depth D2 in Figure 9a, they are not eliminated in this way from depth D1. These near depth boundary fragments are eliminated by the disparity filter (Figure 6), an inhibitory circuit in V2 that operates along the line of sight and across depth to help solve the correspondence problem (Cao and Grossberg, 2005; Grossberg and Howe, 2003; Grossberg and McLoughlin, 1997). The D1 near depth end gap boundary is inhibited by the D2 far depth rectangular boundary at corresponding positions by the disparity filter, because the latter boundary, being closed, is strengthened by surface contour signals, whereas the former boundary is not. Hence the D2 boundary can inhibit the D1 boundary more than conversely.

Although the disparity filter can eliminate the near depth end gap boundary in response to the image in Figure 9a, it cannot do so in response to the image in Figure 9b. This is because the D2 far depth boundary is not closed in this case, and thus is not strengthened by surface contour feedback signals. The same kind of situation occurs in response to the fragmented inducers in Figure 2. How, then, are end gap near-depth boundaries eliminated in this case?
From unoccluded and occluded recognition in V2 to unoccluded seeing in V4. In order to explain how these spurious boundaries are also eliminated, it needs to be explained how additional mechanisms generate the modal, or consciously visible, percepts of the unoccluded parts of both occluding and occluded objects in depth. FACADE theory proposes how boundaries and surfaces may be amodally completed in V2 for purposes of recognition, but also that conscious qualia of the unoccluded surfaces of opaque objects are predicted to be represented in V4. These proposed V2 and V4 representations enable the brain to complete the representations of partially occluded objects behind their occluders for purposes of object recognition, without forcing all occluders to appear transparent, which would be the case if the completed boundaries and surfaces that are illustrated in Figure 9a could generate visible surface qualia. How these V2 and V4 mechanisms may cooperate to achieve both effective recognition and seeing were first described in Grossberg (1994, 1997) and then further developed and simulated in many further articles; e.g., Fang and Grossberg (2009) and Kelly and Grossberg (2000). Grossberg and Yazdanbakhsh (2005) additionally explained and simulated how both opaque and transparent percepts can be generated using the same model mechanisms.

Before summarizing these V2-to-V4 mechanisms for conscious seeing, it is worth noting here that surface contour signals also help to control where the eyes look and thereby help to regulate how the brain learns invariant object categories. The first role arises because surface contour signals are strongest at the distinctive features of an attended object, such as at high curvature positions along a boundary. In addition to the (thin stripe)-to-(pale stripe) feedback that enhances some boundaries while pruning others, a parallel pathway, that is predicted to occur through cortical area V3A, clarifies how these enhanced surface contour positions can also determine target positions of eye movements that explore an attended object’s surface. In all, these signals are proposed to determine where the eyes will look next on an attended surface, and thereby enable inferotemporal cortex to learn view-, size-, and positionally-invariant object categories as the eye movements explore this surface. Thus, the 3D LAMINART model is part of a more comprehensive 3D ARTSCAN Search architecture for active vision wherein 3D boundary and surface representations help to control eye movements for attending, seeing, searching, learning, and recognizing invariant object categories (Cao, Grossberg, and Markowitz, 2011; Chang, Grossberg, and Cao, 2014; Fazl, Grossberg, and Mingolla, 2009; Foley, Grossberg, and Mingolla, 2012; Grossberg, 2009; Grossberg, Srinivasan, and Yazdanbakhsh, 2014).
**Boundary enrichment and surface pruning in V4.** To set the stage for these V2-to-V4 processes, keep in mind that the boundary pruning process spares the closest surface representation that successfully fills-in at a given set of positions, while removing redundant copies of the boundaries of occluding objects that would otherwise form at further depths. This type of process is called "the asymmetry between near and far". When the competition from these redundant occluding boundaries is removed, the boundaries of partially occluded objects can be amodally completed behind them on boundary copies that represent further depths. Moreover, when the redundant occluding boundaries collapse, the redundant surfaces that they momentarily supported collapse as well. Occluding surfaces are hereby seen to lie in front of occluded surfaces.

These surface representations in V2 are depth-selective due to their depth-selective capture by binocular boundaries, but they do not combine brightness and color signals from both eyes (Figure 4). They are said to be computed within monocular Filling-In-DOmains, or FIDOs. The computation of binocular surfaces that combine brightness and color signals from both eyes takes place in V4. These networks are called binocular FIDOs. Here monocular surface signals from both eyes are binocularly matched (pathways 8). The successfully matched binocular signals are pruned by inhibitory signals from the monocular FIDOs (pathways 9). These surface pruning inhibitory signals eliminate redundant feature contour signals at at their own positions and further depths. As a result, occluding objects cannot redundantly fill-in surface representations at multiple depths. This surface pruning process is a second example of the "the asymmetry between near and far".

As in the case of the monocular FIDOs, the feature contour signals to the binocular FIDOs can initiate filling-in only where they are spatially coincident and orientationally aligned with binocular boundaries. Boundary pathways 10 in Figure 5 hereby carry out depth-selective surface capture of the binocularly matched feature contour signals that survive surface pruning. In all, the binocular FIDOs fill-in feature contour signals that: (a) survive within-depth binocular feature contour matching (via pathways 8) and across-depth feature contour inhibition (via pathways 10); (b) are spatially coincident and orientationally aligned with the binocular boundaries (pathways 10); and (c) are surrounded by a connected boundary, or fine web of such boundaries.

Figure 10
In addition, at the binocular FIDOs, the binocular boundaries of nearer depths are added topographically to those that represent further depths (Figure 10b). This third instance of the asymmetry between near and far is called boundary enrichment. These enriched boundaries
prevent opaque occluding objects from looking transparent by blocking filling-in of occluded objects behind them.

The total filled-in surface representation across all binocular FIDOs—after all three processes of boundary pruning, surface pruning, and boundary enrichment act—represents the visible surface percept (Figure 10c). It is called a FACADE representation because it combines together, or multiplexes, properties of Form-And-Color-And-DEpth. As to the three asymmetries between near and far, it is possible that they arise during development due to the asymmetric optic flows that are caused by moving forwards much more than backwards.

**Top-down attention from V4 to V2 eliminates end gap boundaries at D1.** Contour-sensitive top-down feedback from the V4 filled-in surfaces to their generative V2 boundaries obeys the ART Matching Rule (e.g., Carpenter and Grossberg, 1987, 1991), which predicts how top-down object attention works. The ART Matching Rule is defined by a modulatory on-center, off-surround network. Its form has been supported by psychological and neurobiological experiments, and there is even convergence about the mathematical form that the rule should take; see Grossberg (2013) for a review. In the present instance, the modulatory on-centers at each depth, D1 and D2, can strengthen the boundaries that generated the corresponding filled-in surface, while inhibiting other boundaries in its broad off-surround. One consequence of this inhibition is elimination of the spurious end gap boundary at depth D1 (Figure 10d).

**Discussion**

The 3D boundary and surface representations that are depicted in Figures 9 and 10 provide an explanation of how the fragmented images in Figure 2, each of which is caricatured by the image in Figure 9b, generate their depthful figure-ground percepts, notably why the relative depths of figure and ground depend on the positions of the T-junctions relative to the completed boundaries, but not on the relative inducer contrasts that caused them. In response to the fragmented images in Figure 2, these boundaries need to be completed by bipole grouping cells before T-junctions can be created at the fragmented inducers. Once that happens, surface-filling in within closed boundaries ensues. Figures 9 and 10 clarify how the boundary and surface representations within V2 can lead to recognition of figure and ground objects in V2, without these representations also leading to visible surface qualia. The filled-in surface representations within V4 are predicted to support conscious percepts of the qualia of the unoccluded parts of opaque surfaces. Both unique and bistable transparent percepts can
also be explained by these FACADE and 3D LAMINART mechanisms, as has been shown by Grossberg and Yazdanbakhsh (2005).
References

Cao, Y., and Grossberg, S. (2005). A laminar cortical model of stereopsis and 3D surface perception: Closure and da Vinci stereopsis. *Spatial Vision*, 18, 515-578.

Cao, Y., and Grossberg, S. (2012). Stereopsis and 3D surface perception by spiking neurons in laminar cortical circuits: A method of converting neural rate models into spiking models. *Neural Networks*, 26, 75-98.

Cao, Y., Grossberg, S., and Markowitz, J. (2011). How does the brain rapidly learn and reorganize view- and positionally-invariant object representations in inferior temporal cortex? *Neural Networks*, 24, 1050–1061.

Carpenter, G.A., and Grossberg, S. (1987). A massively parallel architecture for a self-organizing neural pattern recognition machine. *Computer Vision, Graphics, and Image Processing*, 37, 54-115.

Carpenter, G. A., and Grossberg, S. (1991). *Pattern recognition by self-organizing neural networks*. Cambridge, MA: MIT Press.

Chang, H.-C., Grossberg, S., and Cao, Y. (2014). Where's Waldo? How perceptual cognitive, and emotional brain processes cooperate during learning to categorize and find desired objects in a cluttered scene. *Frontiers in Integrative Neuroscience*, doi: 10.3389/fnint.2014.0043.

B. Dresp and O. K. Langley (2005) Long-range spatial integration across contrast signs: a probabilistic mechanism? *Vision Research*, 45, 275-284.

B. Dresp and K. Langley (2006) Fine structural dependence of ultraviolet reflections in the King Penguin beak horn" *The Anatomical Record A* 288, 213-22.

Dresp, B. (1997). On illusory contours and their functional significance. *Current Psychology of Cognition*, 16(4), 489–518.
Dresp, B., Durand, S., and Grossberg, S. (2002). Depth perception from pairs of overlapping cues in pictorial displays. *Spatial Vision*, 15, 255-276.

Dresp-Langley, B. (2014). On Galileo's visions: Piercing the spheres of the heavens by eye and mind. *Perception*, 43, 1280-1282. doi: 10.1068/p4311rvw.

Dresp-Langley, B. (2015). 2D geometry predicts perceived visual curvature in context-free viewing. *Computational Intelligence and Neuroscience*, Article ID 708759, 1-9.

Fang, F., Boyaci, H., and Kersten, D. (2009) Border ownership selectivity in human early visual cortex and its modulation by attention. *The Journal of Neuroscience* 29, 460-465.

Fang, L., and Grossberg, S. (2009) From stereogram to surface: How the brain sees the world in depth. *Spatial Vision*, 22, 45-82.

Fazl, A., Grossberg, S., and Mingolla, E. (2009). View-invariant object category learning, recognition, and search: How spatial and object attention are coordinated using surface-based attentional shrouds. *Cognitive Psychology*, 58, 1-48.

Foley, N.C., Grossberg, S., and Mingolla, E. (2012). Neural dynamics of object-based multifocal visual spatial attention and priming: Object cueing, useful-field-of-view, and crowding. *Cognitive Psychology*, 65, 77-117.

Gillam, B., Blackburn, S., and Nakayama, K. (1999). Stereopsis based on monocular gaps: Metrical encoding of depth and slant without matching contours. *Vision Research*, 39, 493-502.

Grossberg, S. (1994). 3D vision and figure-ground separation by visual cortex. *Perception and Psychophysics*, 55, 48-120.

Grossberg, S. (1997). Cortical dynamics of three-dimensional figure-ground perception of two dimensional figures. *Psychological Review*, 104, 618-658.
Grossberg, S. (1999). How does the cerebral cortex work? Learning, attention and grouping by the laminar circuits of visual cortex, *Spatial Vision*, 12, 163-186.

Grossberg, S. (2009). Cortical and subcortical predictive dynamics and learning during perception, cognition, emotion, and action. *Philosophical Transactions of the Royal Society of London*, 364, 1223-1234.

Grossberg, S. (2013). Adaptive Resonance Theory: How a brain learns to consciously attend, learn, and recognize a changing world. *Neural Networks*, 37, 1-47.

Grossberg, S. (2015). Cortical dynamics of figure-ground separation in response to 2D pictures and 3D scenes: How V2 combines border ownership, stereoscopic cues, and Gestalt grouping rules. Submitted for publication.

Grossberg, S., and Howe, P.D.L. (2003). A laminar cortical model of stereopsis and three-dimensional surface perception. *Vision Research*, 43, 801-829.

Grossberg, S., and McLoughlin, N.P. (1997). Cortical dynamics of 3-D surface perception: Binocular and half-occluded scenic images. *Neural Networks*, 10, 1583-1605.

Grossberg, S., and Pinna, B. (2012). Neural dynamics of Gestalt principles of perceptual organization: From grouping to shape and meaning. *Gestalt Theory*, 34, 399-482.

Grossberg, S., Srinivasan, K., and Yazdabakhsh, A. (2011). On the road to invariant object recognition: How cortical area V2 transforms absolute to relative disparity during 3D vision. *Neural Networks*, 24, 686-692.

Grossberg, S., Srinivasan, K., and Yazdanbakhsh, A. (2014). Binocular fusion and invariant category learning due to predictive remapping during scanning of a depthful scene with eye movements. *Frontiers in Psychology: Perception Science*, doi: 10.3389/fpsyg.2014.01457.
Grossberg, S., and Swaminathan, G. (2004). A laminar cortical model for 3D perception of slanted and curved surfaces and of 2D images: development, attention and bistability. *Vision Research*, 44, 1147-1187.

Grossberg, S., and Yazdanbakhsh, A. (2005). Laminar cortical dynamics of 3D surface perception: stratification, transparency, and neon color spreading. *Vision Research*, 45, 1725-1743.

Grossberg, S., Yazdanbakhsh, A., Cao, Y., and Swaminathan, G. (2008). How does binocular rivalry emerge from cortical mechanisms of 3-D vision? *Vision Research*, 48, 2232-2250.

Kelly, F. J., and Grossberg, S. (2000). Neural dynamics of 3-D surface perception: Figure-ground separation and lightness perception. *Perception and Psychophysics*, 62, 1596-1619.

Leveille, J., Versace, M., and Grossberg, S. (2010). Running as fast as it can: How spiking dynamics form object groupings in the laminar circuits of visual cortex. *Journal of Computational Neuroscience*, 28, 323-346.

McLoughlin, N.P. and Grossberg, S. (1998). Cortical computation of stereo disparity. *Vision Research*, 38, 91-99.

Nakayama, K., and Shimojo, S. (1990). da Vinci stereopsis: depth and subjective occluding contours from unpaired image points. *Vision Research*, 30, 1811-1825.

O'Herron, P., and von der Heydt, R. (2009). Short-term memory for figure-ground organization in the visual cortex. *Neuron*, 61 (5), 801-809.

O'Herron, P., and von der Heydt, R. (2011). Representation of object continuity in the visual cortex. *Journal of Vision*, 11, 12. doi: 10.1167/11.2.12.
Qiu, F. T., Sugihara, T., and von der Heydt, R. (2007). Figure-ground mechanisms provide structure for selective attention. *Nature Neuroscience* 10 (11), 1492-1499.

Qiu, F. T., and von der Heydt, R. (2005). Figure and ground in the visual cortex: V2 combines stereoscopic cues with Gestalt rules. *Neuron* 47(1), 155-166.

Rubin, E. (1921). *Visuell Wahrgenommene Figuren: Studien in psychologischer Analyse*. Kopenhagen: Gyldendalske.

Tse, P. U. (2005). Voluntary attention modulates the brightness of overlapping transparent surfaces. *Vision Research*, 45, 1095-1098.

Tzvetanov, T., & Dresp, B. (2002). Short- and long-range effects in line contrast detection. *Vision Research*, 42, 2493-2498.

Yazdanbakhsh, A., and Grossberg, S. (2004). Fast synchronization of perceptual grouping in laminar visual cortical circuits. *Neural Networks*, 17, 707-718.

Zhang, N. R., and von der Heydt, R. (2010). Analysis of the context integration mechanisms underlying figure-ground organization in the visual cortex. *The Journal of Neuroscience* 30(19), 6482-6496.

Zhou, H., Friedman, H. S., and von der Heydt, R. (2000). Coding of border ownership in monkey visual cortex. *The Journal of Neuroscience* 20(17), 6594-6611.

**Figure Captions**

**Figure 1.** Two faces or a vase? In these variations on the famous reversible figures of Rubin (1921), with surface contrasts of opposite signs, the perceptual assignment of border ownership to foreground and background may be influenced by both shifts in spatial attention and prior learning of object categories.

**Figure 2.** The visual configurations (see Dresp, 1997) presented in the psychophysical experiment. The spatially discontinuous shapes in these images induce processes of boundary completion and surface filling-in that generate unambiguous figure-ground percepts of continuous surfaces in depth. In the upper row of these images, the outward-directed contrast edges make the central surface more likely to be seen as lying "behind" the surrounding...
surface, whereas in the lower row of images, the inward-directed edges make the central surface more likely to be seen as standing out "in front" of the surround.

**Figure 3.** Probabilities of perceptual decisions for figure ("in front") or ground ("behind") assignment of the surface in the center of the images with fragmented edge contours, plotted as a function of the direction of the local edge contrasts and their contrast sign.

**Figure 4.** FACADE model macrocircuit. the illuminant-discounted inputs from Right and Left Monocular Preprocessing stage, which is composed of center-surround cells, output to the Left and Right Monocular boundaries composed of simple cells via pathways 1. Left and Right Monocular Boundaries are binocularly fused via pathways 3. Pathways 4 and 5 complete these boundaries using bipole grouping at the Binocular Boundaries stage. Depthful binocular boundaries mutually interact with the Monocular Surfaces stage (pathways 6), where the closed boundaries are filled-in by the illuminant-discounted surface input. The attached boundaries to the successfully filled-in surfaces generate surface contour outputs signals. These signals strengthen the boundaries that induced them, and prune the redundant boundaries at the same positions and further depths (pathways 7). The Binocular Surfaces stage binocularly fuse excitatory inputs from the Left and Right Monocular Preprocessing stages (pathways 8) while surface pruning occurs of redundant feature contours at further depths (pathways 9). Boundary enrichment of the Binocular Boundaries occurs at the Binocular Surfaces and regulates surface filling-in there. Boundaries are enriched by adding boundaries at same positions from near depths to far depths. Due to surface pruning, the illuminant-discounted surface inputs that are contained by the enriched boundaries are pruned from the further depths where boundaries are added.

**Figure 5.** 3D LAMINART model circuit diagram. This laminar visual cortical model consists of a boundary stream that includes V1 interblobs, V2 pale stripes (also called interstripes), and part of V4, and computes 3D perceptual groupings in different scales; and a surface stream that includes V1 blobs, V2 thin stripes, and part of V4, and computes 3D surfaces that are infused with lightness in depth. Both the boundary and surface streams receive illuminant-discounted signals from LGN cells with center-surround receptive fields, and both converge in V4, where visible 3D surfaces are consciously seen that are separated from their backgrounds. Model V2 and V4 also output to inferotemporal cortex (not shown) where object recognition takes place. Model V1 interblobs contain both monocular and binocular cells. Binocular simple cells become disparity-sensitive by binocularly matching left and right
scenic contours with the same contrast polarity in layer 3B before pooling opposite polarity responses at complex cells in layer 2/3A. Monocular and binocular boundary cells control filling-in of monocular 3D surfaces within V1 blobs. Closed boundaries can contain the filling-in process, and can send feedback to V1 interblobs that selectively strengthens the closed boundary components. Monocular and binocular V1 boundaries are pooled in V2. V2 pale stripes can complete 3D perceptual groupings while inhibiting false binocular matches using the disparity filter to solve the correspondence problem. These completed boundaries form compartments in the V2 thin stripes within which filling-in of monocular 3D surfaces occurs. Xlosed boundaries can contain the filling-in process and send surface-to-boundary surface contour feedback signals to enhance their generative boundaries, while also suppressing redundant boundaries at the same positions and further depths. These completed boundaries and filled-in surfaces complete the representations of partially occluded objects. They do not generate visible percepts, but can be recognized by activating inferotemporal cortex. Visible surfaces in which figures are separated in depth from their backgrounds are formed in V4. Here, left and right eye feature contour signals from the LGN are binocularly matched, while redundant feature contour signals are pruned at further depths by inhibitory signals from the thin stripes. Then the pruned feature contour signals induce filling-in of a visible surface percept within enriched binocular boundaries. V4 emits output signals that lead to recognition and grasping of unoccluded parts of opaque surfaces. [Reproduced with permission from Fang and Grossberg (2009).]

**Figure 6.** (a) T-Junction Sensitivity: (left panel) T-junction in an image. (middle panel) Bipole cells provide long-range cooperation (+), and work together with inhibitory interneurons that provide short-range competition (-). (right panel) An end gap in the vertical boundary arises because, for cells near where the top and stem of the T come together, the top of the T activates bipole cells along the top of the T more than bipole cells are activated along the T stem. As a result the stem boundary gets inhibited whereas the top boundary does not. [Reprinted with permission from Grossberg (1997).] (b) Necker cube. This 2D picture can be perceived as either of two 3D parallelograms whose shapes flip bistably through time. (c) When attention switches from one circle to another, that circle pops forward as a figure and its brightness changes. See Grossberg and Yazdanbakhsh (2005) for an explanation. [Reprinted with permission from Tse (2005).]
Figure 7. The top row illustrates how, at a prescribed depth, a closed boundary contour abuts an illuminant-discounted feature contour. When this happens, the feature contours can fill-in within the closed boundary. The bottom row (left panel) depicts how filling-in of the feature contours is contained by this closed boundary contour, thereby generating large contrasts in filled-in activity at positions along the boundary contour. Contrast-sensitive surface contour output signals can then be generated in response to these large contrasts. The bottom row (right panel) depicts a boundary contour that has a big hole in it at a different depth. Feature contours can spread through such a hole until the filled-in activities on both sides of the boundary equalize, thereby preventing contrast-sensitive surface contour output signals from forming at such boundary positions. [Reprinted with permission from Grossberg (2015)].

Figure 8. A closed boundary can form at Depth 1 by combining a binocular vertical boundary at the left side of the square with three monocular boundaries that are projected along the line of sight to all depths. Surface contour output signals can thus be generated by the FIDO at Depth 1, but not the FIDO at Depth 2. The Depth 1 surface contours excite, and thereby strengthen, the boundaries at Depth 1 that controlled filling-in at Depth 1. These surface contours also inhibit the redundant boundaries at Depth 2 at the same positions. As a result, the pruned boundaries across all depths, after the surface contour feedback acts, can project to object recognition networks in inferotemporal cortex to facilitate amodal recognition, without being contaminated by spurious boundaries. [Reprinted with permission from Grossberg (2015)].

Figure 9. Initial steps in generating a 3D percept of figures at different depths in response to a 2D picture with particular occlusion. (a) This figure is composed of three abutting rectangles but generates a percept of a vertical rectangle that partially occludes a horizontal rectangle. Due to mechanisms described in the text, the boundary of the vertical rectangle is separated onto a near depth D1 and achieves border ownership of its shared boundaries with the two smaller rectangles. The remaining boundaries are separated onto a slightly further depth D2, where they can use bipole completion to complete the boundary of the partially occluded horizontal rectangle (dotted lines). This picture does not show the boundary fragments at depth D1 in which end gaps have been generated. The text and Figure 10 propose how end gap boundaries are eliminated. (b) This figure is composed of two abutting rectangles. Although there is no completion of the horizontal rectangle behind the vertical
rectangle, a 3D percept can nonetheless be generated using the same mechanisms. [Adapted with permission from Grossberg (1997).]

**Figure 10.** How spurious end gap boundaries are eliminated. This figure illustrates how spurious end gap boundaries are eliminated from the near depth D1 in the 3D percept that is generated by the 2D picture in Figure 9b. In this case, the end gap boundaries at depth D1 in (a) cannot be eliminated, as they can in response to the percept generated by Figure 9a, by the disparity filter in V2 after surface contour feedback strengthens closed boundaries at the pale stripes from thin stripes. This is true because the boundary at depth D2 is not closed; see (a). On the other hand, this boundary is closed by boundary enrichment in V4; see (b). As a result, top-down attention from the filled-in surfaces in V4 (see (c)) can strengthen the boundaries of closed regions in V2 (see thicker lines in (d)). After this happens, the disparity filter in V2 can eliminate the end gap boundary at depth D1.
Figure 3

Figure 4
Figure 5
Figure 6
Before Filling-in

3D Boundary

Illuminant-discounted surface input

After Filling-in

No Gap

Gap

Figure 7
Figure 8
Figure 9
Figure 10