Transsaccadic Representation of Layout: What Is the Time Course of Boundary Extension?

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How rapidly does boundary extension occur? Across experiments, trials included a 3-scene sequence (325 ms/picture), masked interval, and repetition of 1 scene. The repetition was the same view or differed (more close-up or wide angle). Observers rated the repetition as same as, closer than, or more wide angle than the original view on a 5-point scale. Masked intervals were 100, 250, 625, or 1,000 ms in Experiment 1 and 42, 100, or 250 ms in Experiments 2 and 3. Boundary extension occurred in all cases: Identical views were rated as too "close-up," and distractor views elicited the rating asymmetry typical of boundary extension (wider angle distractors were rated as being more similar to the original than were closer up distractors). Most important, boundary extension was evident when only a 42-ms mask separated the original and test views. Experiments 1 and 3 included conditions eliciting a gaze shift prior to the rating test; this did not eliminate boundary extension. Results show that boundary extension is available soon enough and is robust enough to play an on-line role in view integration, perhaps supporting incorporation of views within a larger spatial framework.

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We can never see the surrounding visual world all at once. Instead, we must sample it a part at a time through successive movements of the eyes and head. An interesting aspect of memory for a single view of a scene is that it will often be remembered as having shown more of the scene than was available in the sensory information—observers remember seeing beyond the edges of the view. This is referred to as boundary extension (Intraub & Richardson, 1989). Although boundary extension is an error with respect to the stimulus, it provides a good prediction of the world beyond the view. For this reason, it has been suggested that boundary extension might serve an adaptive function in scene representation by placing each view within its larger spatial framework (Intraub, 2002, 2007; Intraub, Bender, & Mangels, 1992).

This hypothesis has received support from both behavioral and neuroimaging data. Behavioral research has shown that boundary extension is not present in memory for all pictures of objects (e.g., an object on a blank background) but only for those in which the background depicts part of the visual world (i.e., scene layout; Gottesman & Intraub, 2002, 2003; Intraub, Gottesman, & Bills, 1998). A similar distinction between pictures that include scene layout and those that do not is reflected in the heightened neural responses of the parahippocampal place area to pictures of scene layout (Epstein & Kanwisher, 1998). More recently, an fMRI study of brain activation in the presence of boundary extension revealed that indeed the parahippocampal place area was highly activated when boundary extension occurred (Park, Intraub, Yi, Widders, & Chun, 2007). These experiments indicate that boundary extension is part of the representation of scene layout. The purpose of the present research is to determine the early time course of boundary extension. Specifically, at what stage of processing does boundary extension occur?

In the typical experiment, short series of photographs were presented for multisecond durations (e.g., 15 s each), and memory was tested minutes to 48 hr later (e.g., Candel, Merckelbach, & Zandbergen, 2003; Gottesman & Intraub, 2002; Intraub et al., 1992, 1998; Intraub & Richardson, 1989; Mathews & Mackintosh, 2004). The results suggested that boundary extension might be a long-term phenomenon. In fact, Koriat, Goldsmith, and Pansky (2000), in their review of memory errors, grouped boundary extension with memory errors for text in which “...schematic knowledge is used to make inferences and suppositions that go beyond the actual input event” (p. 494). They noted that schema-induced errors such as these tend to increase over time as memory becomes less detailed. If boundary extension requires long retention times to occur, then it could not play a role on-line during visual scanning. It might be the case that at the earliest stages of processing (e.g., between fixations), visual memory for a just-fixated view might be strong enough to maintain fairly veridical boundaries.

However, an experiment described in Previc and Intraub (1997) demonstrated, somewhat surprisingly, that boundary extension did occur rapidly enough to be observed across a series of perception/action cycles during drawing. Observers viewed four photographs for 15 s each and then drew them from memory. Another group
drew them from a projected image on a screen at the front of the room. As expected, boundary extension occurred in the memory group. What was striking was that boundary extension also occurred in drawings made by observers who could see the photographs. Does this mean that boundary extension occurs during perception?

To answer this question it is important to consider the task carefully. While an observer is literally looking at a picture, he or she can see where the picture ends, and can imagine what would likely exist in the world beyond the edges of the view. Both the sensory information and the expected layout beyond the edges are part of the observer’s representation. Unlike well-known perceptual illusions (e.g., the Mueller-Lyer illusion), boundary extension does not occur while the stimulus is in view. However, when drawing, observers did not maintain fixation on the projected image. They shifted their gaze from the projected image to the paper on their desks, thus relying on memory of the view while they drew. They occasionally would look up and sample the image then look down and draw again, alternating between sensory perception and memory. This suggests that boundary extension occurred at least within seconds after the sensory image was gone, while the observer was drawing.

Consistent with this observation, Intraub, Gottesman, Willey, and Zuk (1996) demonstrated that boundary extension occurs for photographs presented in brief rapid serial visual presentation (RSVP) sequences when memory was tested for the last item in the sequence only 1 s later (see also Bertamini, Jones, Spooner, & Hecht, 2005). Given these results, two possibilities are clear. First, boundary extension might occur a second or more following offset of the stimulus, suggesting it is a very short-term memory error—not rapid enough to play a role in transsaccadic memory but available soon after perception. Alternately, boundary extension might occur as soon as the sensory input is gone. Rather than occurring “in memory,” it might instead be part of the unfolding process of scene perception, which involves a rapidly changing cycle of sensory perception and memory.

If boundary extension is available during the time between saccades (i.e., transsaccadic memory; see Irwin, 1991, 1993), then we would expect to see it under the following three conditions. First, it should occur following a brief glimpse of a scene (analogous to the duration of a single fixation). Boundary extension is known to occur following presentations as brief as 250 ms and 333 ms (Bertamini et al., 2005; Intraub et al., 1996; 500 ms: Intraub, Hoffman, Wetherhold, & Stoehs, 2006). Second, it should be evident following a gap in sensory input commensurate with a “typical” saccade (on the order of 30–50 ms; Rayner, 1998). However, retention times briefer than 1 s have not been tested. Third, boundary extension would need to survive a gaze shift caused by a change in the position of the eyes or the head. Recent research shows that boundary extension occurs following a single eye movement when memory is tested 2 s later (Intraub et al., 2006), but it is not known whether it occurs when tested immediately after a gaze shift. It is possible that the process of planning and executing a gaze shift might delay the onset of boundary extension for a couple of seconds, thus preventing its inclusion in transsaccadic memory.

Early Memory Buffers

Might early memory buffers prevent boundary extension? It may be that early visual buffers maintain a fairly veridical representation of layout, essentially protecting the representation from distortions. Iconic memory has been categorized as a high-capacity, brief-duration, veridical buffer that is disrupted by masking by both luminance and pattern. So, for example, if a single fixation on a scene were followed by a fixation on an empty region of space, a veridical representation of the spatial expanse of that view might be maintained in an iconic representation (i.e., as informational persistence) for 100–300 ms (Irwin & Brown, 1987; Irwin & Yeomans, 1986).

What if, however, a single fixation on a view of a scene is followed by another fixation whose contents would disrupt an iconic representation, or alternatively, what if an iconic representation isn’t maintained across a gaze shift? Visual short-term memory (VSTM) has been categorized as a longer lasting (i.e., multisecond) buffer that does not maintain a literal copy of the physical visual stimulus (i.e., it is not a point-by-point copy) and that is capacity limited but not disrupted by visual masking (Irwin, 1991; Phillips, 1974). Numerous studies suggest that VSTM representations are not literal copies of the display (Gordon & Irwin, 1996; Henderson, 1994; Hollingworth, Hyun, & Zhang, 2005; Irwin, 1991, 1992; Irwin & Andrews, 1996; Olson & Jiang, 2004; Phillips, 1974). This does not imply, however, that VSTM representations contain no visual information. Here we ask whether VSTM might maintain a veridical representation for a brief interval following a picture’s offset, thus preventing boundary extension.

Gaze Shifts

Might gaze shifts delay boundary extension? If boundary extension occurs rapidly enough to support view integration on-line, it would not be of use unless it could also survive a shift in gaze. In prior boundary extension research, stimulus and test views have always been presented in the same physical location. Might the demands of attention and/or motor systems engaged during gaze shifts early in processing either delay the onset of boundary extension or cause memory for boundaries simply to be poor—causing errors but not the strong unidirectional error that characterizes boundary extension? Numerous studies have demonstrated that gaze shifts suppress a variety of cognitive processes, whereas others appear to continue across them unimpeded (for a review, see Irwin & Brockmole, 2004). A brief examination of the known properties of transsaccadic memory might shed some light on whether boundary extension might be included in transsaccadic memory.

Most experiments on transsaccadic memory have focused not on scenes as a whole, but on the properties of individual objects that are remembered across a saccade. In general, results suggest that representations of structural descriptions of objects are retained (Carlson-Radvansky, 1999; Carlson-Radvansky & Irwin, 1995; Verfaille & De Graef, 2000; Verfaille, De Troy, & Van Rensbergen, 1994), but there is also evidence for retention of specific visual information (object orientation: Henderson & Sievert, 1999, 2001; but not detailed object contours: Henderson, 1997).

Boundary extension, however, involves an extrapolation of layout. Although the literature on transsaccadic memory has not
focused on scene layout, there have been several studies that have focused on retention of spatial relations. These provide evidence that spatial information is available in transsaccadic memory. It has been shown that information that specifies the structural relations of parts of a single object is included in a transsaccadic representation (Carlson-Radvansky & Irwin, 1995). In addition, information about the configuration, or the spatial relations among different objects, can also be represented (Carlson-Radvansky, 1999; Deubel, 2004; Germey, de Graaf, Panis, van Eccelpoel, & Verfaillie, 2004). The question we ask is whether this transsaccadic representation includes extrapolated layout (boundary extension) or instead maintains a more veridical representation of the view.

The Current Experiments

To explore the early time course of boundary extension and test its resiliency to shifts in gaze, in all three experiments we used Intraub et al.’s (1996) three-picture RSVP method. Three pictures were presented for 325 ms each in a continuous sequence followed by a masked retention interval and subsequent boundary memory test. There were three reasons for choosing this method. First, by embedding a target picture in a rapidly changing series, we could approximate the dynamic nature of visual scanning.1 Second, the design allowed us to test boundary memory following presentations of 0, 1, or 2 intervening items, thus allowing us to determine if factors such as conceptual masking (Intraub, 1984; Potter, 1976) during successive presentation might influence boundary extension. Third, the rapidity of input coupled with observer uncertainty about which picture would be tested would minimize the observer’s ability to develop verbal strategies (e.g., “the man’s head is .5 cm from the top”) for remembering boundary placement over the course of the session.

In all three experiments, on each trial, the observer was required to rate the repeated scene as being the same view or a more close-up or wide-angle view than before on a 5-point scale (Intraub & Richardson, 1989). Across experiments the interval between offset of the last picture and onset of the test picture was always masked and it ranged between 42 ms (comparable to the duration of a saccade) and 1 s (to replicate earlier research). To test the effect of a shift in gaze on boundary extension, in Experiments 1, 3a, and 3b, test pictures were presented either in the same location as the RSVP sequence or to the left or right side of the screen.

During the experiment, when a scene repeated it could be the same, a more close-up view, or a more wide-angle view than one of the pictures in the presentation sequence. In this way, all the patterns of response that are diagnostic of boundary extension could be addressed following each masked interval. These patterns have been replicated in many studies (e.g., Bertamini et al., 2005; Gottesman & Intraub, 2002; Intraub et al., 1992, 1998; Intraub & Richardson, 1989). The three patterns of interest are as follows:

1. When the target picture and test picture are identical close-ups, observers tend to reject the test picture as being the same, reporting instead that it is more close-up than the target picture, thus indicating that the target picture was remembered with extended boundaries. This was tested in Experiments 1–3.

2. Target pictures that are tight close-ups yield more boundary extension than wider angle views of the same scene; in fact, wider angle views tend to yield no directional distortion (see Gottesman & Intraub, 2002; Intraub et al., 1992; Intraub & Berkwits, 1996). This was tested in Experiment 2.

3. When a close-up is the target and a wider view is presented at test, observers rate the pair as being more similar than when the reverse is the case. This is because boundary extension for a closer target causes it to be remembered as looking more like the wider angle test picture. This asymmetry was also observed in neural responses to dissimilar pictures in scene-selective brain regions (Park et al., 2007). This signature pattern was tested in Experiment 2.

Experiment 1

In Experiment 1 boundary extension was tested for a single picture immediately following a three-picture RSVP sequence and a masked interval of 1 s, 625 ms, 250 ms, or 100 ms. We selected the latter three intervals as coarse divisions of the 1-s interval after which boundary extension is known to occur (Bertamini et al., 2005; Intraub et al., 1996). To provide the most sensitive test of boundary extension, on half the trials the same close-up served as both target and test picture. Thus, in the critical case for testing time course (when the target picture is in the final serial position of the RSVP sequence), the same close-up view of a complex scene was interrupted by a 100-ms masked interval. To ensure that observers were focused on the task and were using the scale appropriately, on the remaining trials scenes were divided into two sets: In one a close-up target was tested with a more wide-angle view of the same scene, and in the other, a wider angle view was tested with a close-up version.

Finally, to determine if boundary extension survives a gaze shift between the target picture and the test picture, the test picture appeared equally often in the same location as the target (center screen) or shifted to the right or left side of the screen. If boundary extension does not occur at intervals briefer than 1 s, or if a gaze shift disrupts the extrapolation of expected surrounding space, then boundary extension cannot play an on-line role in view integration during active scanning.

Method

Participants. A total of 144 University of Delaware undergraduates, fulfilling a requirement for an introductory psychology course, participated in the experiment. All reported having normal or corrected-to-normal vision and normal color vision.

Apparatus. All stimuli were presented on a 21” flat-screen CRT monitor in 32-bit color at a resolution of 1024 × 768 pixels

1 Of course, there are several aspects of visual scanning that RSVP does not simulate: for example, the peripheral information made available by the spatial attention shift preceding an eye movement (Deubel & Schneider, 1996; Hoffman & Subramaniam, 1995; Irwin & Zelinsky, 2002; Kowler, Anderson, Dosher, & Blaser, 1995) or the brief period of saccadic suppression that occurs between fixations (see Volkmann, 1986). However, it does allow us to approximate the changes in visual input that result from gaze shifts, as well as the temporal pacing of these changes.
and a refresh rate of 120 Hz that was driven by a video card with 128 MB of video memory. Stimulus presentation was controlled by a Pentium-based PC running Microsoft Windows XP. The software was based on a template program supplied by SR Research (Mississauga, Ontario, Canada) written in C that used Simple DirectMedia Layer (SDL) Version 1.2.9 to access the video hardware. The viewing distance was approximately 80 cm, and on average pictures subtended 9.2° × 10.2° of visual angle (widths ranged from 5.9° to 13.7°; heights ranged from 8.9° to 10.4°).

Stimuli. Stimuli were 96 color photographs that depicted people engaging in various activities, for example, a football player kicking a football, a man tossing a pizza, and a couple dancing. Some of the images were copied, with permission, from the Big Box of Art 615,000 Images database of royalty-free images (Hemera Technologies); others were downloaded from the Internet. All stimuli were presented on a gray background. Of the 96 pictures, 32 served as targets (i.e., to-be-tested pictures; 2 in the practice trials and 30 in the experimental trials). The other 64 pictures served as fillers in the presentation sequence (i.e., the two nontarget pictures in the RSVP trial). A given scene (close or wide version) was always presented with the same filler items. Each target scene could be presented in its close-up version or its more wide-angle version, as is illustrated in Figure 1. Close-ups were created by enlarging the wide-angle version by 8–21% in area and then cropping the enlarged version to be the same size as the original. Thus, both versions were the same size, but the wide-angle version showed a larger amount of background surrounding the main object or objects.

Design and procedure. A depiction of a trial sequence is shown in Figure 2. Each self-initiated trial began with a central fixation point that remained on screen for 500 ms. The RSVP sequence followed (325 ms/picture) in the center of the screen. The fixation point for the next trial then appeared and the observer responded. Observers were asked to rate whether the test picture showed the same view, a more close-up view, or a more wide-angle view than before using a 5-point Likert scale. The alternatives (and their corresponding numerical values) were “much closer up (−2),” “a little closer up (−1),” “the same (0),” “a little farther away (1),” and “much farther away (2).” The test picture was visible until the observer clicked one of these choices with the mouse. Observers then indicated how confident they were about their response by clicking “sure (3),” “pretty sure (2),” “not sure (1),” or “don’t remember that picture (0).” The fixation point for the next trial then appeared and the observer initiated the trial by clicking the mouse.

There were a total of 30 trials. On 15 trials the target and test picture were identical close-ups (referred to as CC trials). On 14 trials the target and test picture were different views of the same scene; half the time a close-up was the target and the test picture was a more wide-angle view (CW trials) and half the time the reverse was true (WC trials). Each target picture was tested in only one of these conditions across observers. The 30th trial was always a “dummy” trial in which the target picture was tested with the same view. This trial had to be added to allow us to show each target at each serial position equally often across observers, while at the same time having an equal number of CW and WC trials. Responses made on this trial were not included in the analyses.

Results and Discussion

Observers were rather confident of their ratings; on average, 21%, 57%, and 20% of their responses were rated as “sure,” “pretty sure,” and “not sure,” respectively. They reported not recognizing the test picture on only 2% of the trials, and these
were excluded from all analyses. A $4 \times 2$ (Masked interval $\times$ Side of display) mixed-design analysis of variance (ANOVA) comparing observers’ mean boundary ratings for test pictures presented on the left versus right side of the screen revealed no main effect of location, $F(1, 140) = 1.73$, ns, and no interaction with the masked interval, $F < 1$. Observers’ mean boundary ratings in these two conditions were subsequently collapsed across this factor.

Critical CC trials: Targets and test pictures are identical close-ups. Figure 3 (left panel) shows the mean boundary rating at each serial position (collapsed over the spatial position of the test picture). The 95% confidence intervals revealed that boundary extension occurred at each serial position for each masked interval. Thus, boundary extension occurred even at the briefest interval tested—when the final picture in a sequence was repeated only 100 ms later. To determine if the size of the boundary extension effect was influenced by the duration of the masked interval, a $3 \times 4$ (Serial position $\times$ Retention interval) mixed-design ANOVA was conducted. It revealed no main effect of the masked interval’s duration, $F(3, 140) = 1.16$, ns; no effect of serial position, $F < 1$; and no interaction, $F < 1$. The lack of a serial position effect shows that the onset of new meaningful pictures during RSVP did not disrupt incorporation of the extrapolated region into the spatial representation of the scenes—that is, there was no effect of conceptual masking (Intraub, 1984; Potter, 1976).

Figure 3 (right panel) shows the mean boundary rating as a function of the spatial location of the test picture for each masked interval (collapsed over serial position). As shown by the 95% confidence intervals in the figure, boundary extension occurred whether or not a shift in gaze intervened between presentation and test. Observers were never forewarned about the location of the test picture, yet when it shifted away from center screen, the concomitant gaze shift had no inhibitory effect on boundary extension. The expanded representation of layout clearly survived the shift in attention and subsequent gaze shift, suggesting that boundary extension is available during the time course of transsaccadic memory.

CW and WC Trials. The mean boundary ratings for CW and WC trials (collapsed over serial position and spatial position) for each masked interval are shown in Figure 4 (left and right panels, respectively). Observers were able to recognize the presence of distractors and were clearly using the scale appropriately. Consistent with the occurrence of boundary extension, the right panel of Figure 4 shows that observers were quite good at recognizing when the test picture was more close-up than the target, whereas the left panel shows that more wide-angle test pictures were sometimes mistaken as being the same as the target.

Experiment 2

Experiment 1 showed that a 100-ms interruption was sufficient to elicit boundary extension. In Experiment 2 we decreased the briefest interval further to 42 ms (commensurate with a saccade);
intervals tested were 250 ms, 100 ms, and 42 ms. To enhance the observer’s ability to retain a veridical representation, the RSVP sequence and test picture were always in the same location (center screen). In addition, we sought to obtain converging evidence for boundary extension through implementation of a design used in many prior boundary extension studies (e.g., Intraub et al., 1998; Intraub & Richardson, 1989), in which targets were either close-up or wide-angle views of a scene, and the test picture was either the same view as the target or its complement. This yielded four different test conditions: (a) close-up view tested with the same close-up view (CC trials), (b) wide-angle view tested with the same wide-angle view (WW trials), (c) close-up view tested with the wide-angle version of the scene (CW trials), and (d) wide-angle view tested with the close-up version of the scene (WC trials). Scenes were counterbalanced across these four conditions. In this way we could determine if all three patterns that are diagnostic of boundary extension (i.e., boundary extension for CC trials, little or none for WW trials, and a CW–WC asymmetry, as discussed in the introduction) would occur.

Method

Participants. A total of 108 University of Delaware undergraduate students, fulfilling a research participation requirement for an introductory psychology course, participated in the experiment. All reported having normal or corrected-to-normal vision and normal color vision, and none participated in the previous experiment.
Apparatus and stimuli. The apparatus and stimuli were the same as in the previous experiment with the exception of the following differences. In the current experiment there were a total of 114 pictures of real-world scenes used, with 38 serving as targets and the remaining 76 serving as fillers for the presentation sequences. Here we added 6 new sequences to the 32 sequences used in the previous experiment.

Procedure and design. The procedure was the same as in Experiment 1 with the exception that all stimuli and test pictures were presented in the center of the screen. There were a total of 36 trials. On half the trials, the target was a close-up view; on the other half, it was a wide-angle view. Within each of these conditions, on half of the trials the test picture was the same view as the target picture (CC, WW); on the other half, it was a different (i.e., the other) view (CW, WC). This yielded 9 trials for each target view–test view combination. Within each of these four target–test combinations, the target picture appeared in Serial Positions 1, 2, and 3 of the initial presentation equally often, yielding 3 trials per cell of the design. Across observers, every target picture appeared in each of these conditions equally often. Each observer was tested with only one of the three masked intervals (42, 100, and 250 ms), with 36 observers being tested with each masked interval.

Results and Discussion

As in Experiment 1, observers were rather confident, rating their responses as “sure,” “pretty sure,” and “not sure” on 27.5%, 53%, and 17%, of the trials respectively. They failed to recognize test pictures on only 2.5% of the trials, and these were excluded from all analyses.

Boundary extension ratings for the final picture in each sequence. To examine the early time course of boundary extension, here we focus on those trials on which the target picture appeared in Serial Position 3. All three patterns consistent with boundary extension were obtained. First, as in Experiment 1, CC trials yielded boundary extension at all masked intervals—even the 42-ms masked interval. Figure 5 shows the mean boundary rating and 95% confidence interval for each serial position at each masked interval. The upper left panel shows that CC trials yielded boundary extension at each masked interval. Thus, the results show that on average, a 42-ms interruption in the sensory input was sufficient to prevent observers from recognizing the repetition of a close-up view—instead, they remembered the first presentation as having extended boundaries.

Figure 5. Observers’ mean boundary ratings for CC trials are shown on the upper left, mean boundary ratings for WW trials are shown on the upper right, mean boundary ratings for CW trials are shown on the lower left, and mean boundary ratings for WC trials are shown on the lower right, for each serial position of the target at each retention interval (Experiment 2). Error bars show the 95% confidence interval of the mean. For CC and WW trials, means that are significantly less than zero reflect boundary extension; means that are significantly greater than zero reflect boundary restriction.
Second, wide-angle views yielded less boundary extension than their close-up counterparts. The mean boundary ratings for WW trials for each masked interval are shown in Figure 5 (upper right panel). A 3 × 2 (Masked interval × Test type) mixed-design ANOVA on observers’ mean boundary ratings revealed that CC trials yielded more boundary extension than WW trials, \( F(1, 105) = 12.36, p < .05, \) partial \( \eta^2 = .11, \) and the lack of interaction with masked interval duration \( (F < 1) \) indicates that this same pattern occurred for each masked interval.

Third, a robust asymmetry in responses to the CW and WC trials, as is shown in Figure 5 (lower left and right panels), provides strong additional support that boundary extension was occurring at these intervals. The same pair of pictures was rated differently (with respect to how different they were from one another) depending on their order (close-up first or wide angle first). A 3 × 2 (Masked interval × Test type) mixed ANOVA on the absolute value of the mean boundary scores on CW and WC trials revealed a main effect of test type, \( F(1, 105) = 3.31, p < .05, \) partial \( \eta^2 = .06, \) and planned comparisons revealed that mean ratings for the 250-ms interval were larger than mean ratings for the 42-ms interval \((–.47 vs. –.19), (t(70) = 2.69, p < .05)\). This small difference could have been the result of a fading visual representation that was attenuating boundary extension at the shortest interval. The results of a recent experiment, however, suggest that this difference may be related to the memory load resulting from the three-picture stimulus sequences (Intraub & Dickinson, 2006). When we presented observers with only a single stimulus for 250 ms and then tested their memories for the views (using the same procedure as in the current experiment), we found no difference in the amount of boundary extension that occurred following 42-ms and 250-ms masked intervals.

**Boundary extension ratings for pictures in Serial Positions 1 and 2.** Analysis of responses to pictures in Serial Positions 1 and 2 showed that boundary extension was not eliminated by the onset of new pictures (i.e., conceptual masking; Potter, 1976). The same three critical data patterns diagnostic of boundary extension occurred on these trials. First, as is shown in the upper left panel of Figure 5, CC trials yielded boundary extension at all masked intervals. Second, a 3 × 2 × 2 (Masked interval × Test type × Serial position) mixed ANOVA comparing mean boundary ratings on CC trials and on WW trials revealed that less boundary extension occurred for WW trials than for CC trials, \( F(1, 105) = 27.39, p < .05, \) partial \( \eta^2 = .21, \) There was no significant interaction of test type with either masked interval, \( F(2, 105) = 1.03, ns, \) or serial position, \( F(1, 105) = 2.80, ns, \) indicating that this pattern occurred at each serial position and each masked interval. Third, a 3 × 2 × 2 (Masked interval × Test type × Serial position) mixed ANOVA on the absolute value of the mean boundary scores on CW and WC trials revealed a main effect of test type, \( F(1, 105) = 59.74, p < .05, \) partial \( \eta^2 = .36, \) Further, this main effect of test type did not interact with either masked interval, \( F < 1, \) or with serial position, \( F < 1, \) indicating that this asymmetry occurred at each serial position and each masked interval.

**Experiments 3a and 3b.**

Experiment 2 demonstrated that boundary extension occurs rapidly enough to play a role in view integration during visual scanning. In Experiments 3a and 3b, we therefore returned to the question of whether or not a shift in gaze would influence boundary extension. In Experiment 3a, as in Experiment 2, the masked intervals tested were 250 ms, 100 ms, and 42 ms. Test pictures were presented in either the same spatial location as the RSVP sequence or in a different (nonoverlapping) location. In Experiment 3b, we recorded observers’ eye movements to determine the extent to which they were maintaining central fixation during stimulus presentation and to determine the duration of the saccade-defined interval between the end of the stimulus sequence and the beginning of observers’ first fixations on the test items in shift-gaze trials.

All trials were CC trials (close-ups tested with close-ups). Unlike Experiment 1, position of the test picture was blocked so that observers always knew where the test picture would appear. We reasoned that by eliminating this uncertainty we would maximize our ability to detect any subtle influences of a gaze shift on memory for the expanse of a view. To determine if the time between the offset of the last picture and the beginning of the saccade would influence boundary extension, we included the same three masked intervals as in Experiment 3a. The end of the interval signaled the start of a saccade.

**Method.**

**Participants.** In Experiment 3a, 108 University of Delaware undergraduate students, fulfilling a research participation requirement for an introductory psychology course, participated. In Experiment 3b, 36 individuals (5 college graduates who were paid $8/hour and 31 students) participated. All reported having normal or corrected-to-normal vision and normal color vision, and none participated in either of the previous experiments.

**Apparatus and stimuli.** The apparatus was the same as in the previous experiment. The stimuli included only the close-up versions of targets and test pictures from the previous experiment. In Experiment 3b, we used an EyeLink II video-based eye tracking system to collect eye movement data (SR Research; Mississauga, Ontario, Canada). Eye position was sampled at 500 Hz, the system’s spatial resolution was estimated to be less than 0.4º, and head position and viewing distance were fixed with a chinrest.

**Design and procedure.** There were a total of 36 trials. Stimuli and test pictures were shown on either the left side or right side of the screen; no items were presented in the screen’s center, and there was no spatial overlap between the stimulus sequence and the test picture when they appeared on different sides of the screen. When the two were presented in different locations, the average center-to-center distance was 11.8º (ranging from 8.4º to 14.2º).
On half of the trials, the test picture was shown on the same side of the screen as the stimulus sequence (maintain-fixation trials); on the other half, it was presented on the other side of the screen (shift-gaze trials). Within each of these conditions, on half of the trials the stimulus sequence was presented on the left side of the screen; on the other half, it was presented on the right side. This yielded 9 trials for each stimulus side–test side combination. Otherwise, the procedure was the same as in the previous experiment.

These conditions were blocked, and observers were informed of the stimulus and test locations at the start of each block. The sequence of blocks was counterbalanced across observers. Within each of these four stimulus–test combinations (test was on the same side: LL, RR; test was on the other side: LR, RL), the target picture appeared in Serial Positions 1, 2, and 3 of the initial presentation equally often, yielding three trials per cell of the design. Across observers, every picture appeared in each of these conditions equally often. Each observer in Experiment 3a was tested with only one of the three masked intervals (42, 100, and 250 ms), with 36 observers being tested with each masked interval. In Experiment 3b, only the 42-ms condition was tested, and maintaining fixation during stimulus presentation was defined as keeping gaze within a $4.4^\circ \times 4.4^\circ$ square surrounding the initial fixation location until the test picture’s onset.

Results and Discussion

Again, observers tended to be rather confident of their responses. On average, the percentages of “sure,” “pretty sure,” and “not sure” responses were 26%, 56%, and 16%, respectively. Observers failed to recognize pictures on only 2% of the trials, and these were excluded from all analyses. Separate ANOVAs conducted on observers’ mean boundary ratings on maintain-fixation and shift-gaze trials revealed no main effect of side of stimulus presentation and no interaction with serial position in either condition in either experiment (all $F$s < 1.66, ns). Mean boundary ratings in each of these conditions were subsequently collapsed across the factor of side of stimulus presentation in both experiments.

Experiment 3a: Serial Position 3. Again, the critical condition for observing time course is when the target is in the final position in the sequence. Replicating Experiment 2, when fixation was maintained, boundary extension occurred for the final picture at each masked interval. The mean boundary ratings and 95% confidence intervals are shown in Figure 6 (left panel). This provides additional evidence to suggest that boundary extension occurred rapidly enough following stimulus offset to be present in transsaccadic memory. BE for the final picture in the sequence also survived a gaze shift for all masked intervals tested. The mean boundary ratings and 95% confidence intervals for shift-gaze trials are shown in Figure 6 (right panel). This shows that boundary extension survived a gaze shift early in processing.

In the maintain-fixation condition, planned comparisons among all three masked intervals showed that when fixation was maintained, the mean rating for Serial Position 3 targets at the 42-ms masked interval was significantly smaller than both the mean rating for Serial Position 3 targets at the 100-ms interval (−.08 vs. −.25), $t(70) = 2.73$, $p < .05$, and the mean rating for those at the 250-ms interval (−.08 vs. −.36), $t(70) = 4.22$, $p < .05$.

Experiment 3a: Serial Positions 1 and 2. As in Experiments 1 and 2, boundary extension was not eliminated by conceptual masking. As can be seen from the 95% confidence intervals for maintain-fixation trials (Figure 6, left panel) and shift-gaze trials (Figure 6, right panel), targets that appeared in Serial Positions 1 and 2 yielded significant boundary extension. In the context of visual scanning, this is important because the contents of each new fixation would serve as a conceptual mask for the contents of the previous one.

**Figure 6.** Observers’ mean boundary ratings for maintain-fixation trials are shown on the left, and mean boundary ratings for shift-gaze trials are shown on the right, for each serial position of the target at each retention interval (Experiment 3a). Error bars show the 95% confidence interval of the mean. Means that are significantly less than zero reflect boundary extension; means that are significantly greater than zero reflect boundary restriction.
Experiment 3b: Oculomotor data. Boundary extension occurred for targets in all three serial positions in both conditions, as is shown in Figure 7. The eye tracking data revealed that observers maintained fixation during stimulus presentation on 84% of maintain-fixation trials and 79% of shift-gaze trials. An ANOVA comparing observers’ mean boundary ratings for all trials with mean ratings for only trials on which fixation was maintained revealed no changes in data patterns (all $F$s < 1). Thus, observers were following instructions, rather than adopting a strategy of fixing near an edge instead of at the center of the stimuli.

The eye tracking data also revealed that the average retention interval on shift-gaze trials, defined as the time between the offset of the final scene in the RSVP sequence and the beginning of the first fixation on the test picture, was 370 ms. This interval can be divided into three components: the masked interval (during which fixation was maintained; 42 ms) the saccade latency (the time required to initiate the saccade following test picture onset; 264 ms), and the saccade duration (64 ms). Leftward and rightward saccade latencies did not differ significantly (255 ms vs. 273 ms), $t(35) = 1.29, ns$, nor did the corresponding retention intervals for left-side or right-side test items (360 ms vs. 381 ms), $t(35) = 1.44, ns$. Thus the time required for observers to shift gaze to the test picture on shift-gaze trials was only slightly longer than typical saccade latencies, which are considered to be in the area of 150–200 ms (Carpenter, 1988; Rayner, 1998). This suggests that observers were neither delaying saccades to test pictures nor initiating them prior to the test picture’s onset.

In conclusion, the results of Experiments 3a and 3b show that boundary extension would, at the least, be available to play a role in the integration of successive views and that it is clearly robust enough to survive a shift in gaze following a dynamic series of briefly presented scenes.

General Discussion

Boundary extension is an error of commission. One usually thinks of such errors as emerging over relatively long periods of time or under conditions of high memory load. In the present experiments we demonstrated that an interruption of a view for as little as 42 ms was sufficient to elicit boundary extension. In these cases observers misidentified the identical view as looking more close-up than the view seen before the interruption. Across experiments, we tested memory for target pictures embedded in rapidly presented streams of three pictures followed by a masked interval of 1 s, 625 ms, 250 ms, 100 ms, or 42 ms. Boundary extension occurred following all of these intervals and did so irrespective of the target’s serial position.

At the outset we had expected that an early visual memory buffer, perhaps iconic memory (which can sometimes persist in spite of a mask; Smithson & Mollon, 2006) or VSTM, might prevent the involvement of boundary extension in the earliest phases of visual scanning, but clearly, at least in the presence of a disruptive mask, boundary extension was evident following an interval commensurate with a saccade. In addition, in all three experiments, boundary extension occurred for pictures that appeared in Serial Positions 1 and 2, suggesting that conceptual masking, caused by the immediate onset of new, meaningful visual information, does not eliminate boundary extension (Intraub, 1984, 1999; Potter, 1976, 1999).

Experiments 1, 3a, and 3b demonstrated that this early, potentially fragile memory of an extended view was robust enough to survive a shift in gaze. This is important because it shows that neither the attention shift nor motor planning involved in this act disrupted what was, for the most part, a rather confidently held assessment that the original view had been more expansive. This occurred both when observers knew exactly where the test picture would appear (Experiments 3a and 3b) and when they had no advance warning and the test picture’s spatial position shifted randomly across trials (Experiment 1). These experiments exploited one of the most striking characteristics of boundary extension—the failure to recognize an identical close-up view.

Experiment 2 provided converging evidence for the occurrence of boundary extension by testing for additional patterns of results.
that are diagnostic of boundary extension under conditions in which observers maintained fixation. Observers were presented with close-up or more wide-angle views, and test pictures were either identical to the target or showed the alternate view (e.g., Gottesman & Intraub, 2002; Intraub et al., 1992, 1998; Intraub & Richardson, 1989). As in the other experiments, repetition of the same close-up at presentation and test was not recognized, as observers tended to think the repetition was more “close-up.” However, it also showed that repetition of the same wider angle views resulted in less boundary extension than had been obtained for the repeated close-ups and that a robust asymmetry occurred for trials on which a different view was shown at test. These patterns occurred not only for the last position in the sequence but across all serial positions, again demonstrating that boundary extension was not eliminated by conceptual masking (Intraub, 1984, 1999; Potter, 1976, 1999).

To test the early time course of boundary extension it was necessary to follow a target picture with the test picture, so of course our test conditions did not provide a simulation of what usually happens during visual scanning in which the image changes with each saccade. It is important to note, however, that our test provides an even stronger test of boundary extension for that very reason. Given that boundary extension occurred both when the same picture repeated and when the observer could control an eye movement with full knowledge that spatial memory would be tested, it is reasonable to conclude that boundary extension is, at the very least, available early enough in processing to play a role in view integration during active visual scanning.

Implications for Transsaccadic Memory and Scene Representation

Most researchers agree that scene representation is not picture-like but instead is more schematic in nature (e.g., Henderson & Hollingworth, 2003; Hochberg, 1978; Intraub, 1997; Simons & Levin, 1997; Simons & Rensink, 2005). The schematic character of representation has also been posited specifically in the case of transsaccadic memory (Carlson-Radvansky, 1999; Carlson-Radvansky & Irwin, 1995; Irwin, 1991, 1992; Verfaillie & De Graef, 2000; Verfaillie et al., 1994). Research has demonstrated that scene layout is obtained rapidly and that in addition to conveying a rough portrait of scene structure, it supports scene comprehension at a conceptual level (Biederman, 1981; Oliva & Schyns, 1997). The present research demonstrates that not only is some characterization of layout maintained in transsaccadic memory but that this representation includes a projection of the layout beyond the bounds of the original sensory information. The rapidity of this effect is striking.

Although it is assumed that some representation of layout is retained above and beyond the individual details of a view (e.g., Rensink, 2000), the current experiments provide evidence that it is retained within transsaccadic memory as well. This adds to our knowledge of the transsaccadic representation because most research in that area has focused primarily on important issues of object representation. For example, several studies have addressed the number of objects (Irwin, 1991, 1992; Irwin & Andrews, 1996) and the types of object properties (Gordon & Irwin, 1996; Henderson, 1997; Henderson & Siefert, 1999, 2001) that can be maintained across a saccade. There has also been research on spatial relations that are maintained across a saccade, both among parts of the same object (Carlson-Radvansky & Irwin, 1995; Verfaillie & De Graef, 2000) and between multiple objects (Carlson-Radvansky, 1999; Deubel, 2004; Germey et al., 2004). The current research suggests, somewhat surprisingly, that a visual interruption lasting the duration of a saccade is enough time for boundary extension to occur.

The extended region is very constrained, perhaps enough to prime upcoming layout (Sanocki, 2003) but not so much that we confabulate and become confused during visual scanning. This unidirectional boundary error might better be classified as a highly constrained prediction rather than an error. After all, the goal of the visual system is to construct a coherent representation of the surrounding visual world, in spite of the fact that scene perception is built from a succession of samples over time. Successive fixations embellish the representation with more detailed information (Hollingworth, 2004; Hollingworth & Henderson, 2002; Melcher, 2001, 2006; Zelinsky & Loschky, 2005). But the location of view boundaries is not among those details. It is more adaptive for the visual system to “ignore” these spurious boundaries and extrapolate the likely continuity of global layout just beyond them.

Implications for the Locus of Boundary Extension

Having more clearly charted boundary extension’s early time course, we return to our original question: At what stage of processing of a given view does boundary extension occur—perception or memory? Finding evidence for boundary extension in memory following a 42-ms gap in sensory input suggests that boundary extension occurs within the span of a single saccade—perhaps as soon as sensory input is removed. What implications do these findings have for the locus of boundary extension? Is there a “process” of extrapolation that rapidly takes place in memory during the brief gap between fixations? A more likely possibility is that the first step of the creation of a boundary-extended view involves processes taking place during sensory perception (i.e., while the sensory information is present). Here it is important to entertain the role of amodal perception and higher order expectation in scene perception. The layout and gist of a scene are grasped very rapidly—requiring only a brief glimpse (less than the duration of a typical fixation; Biederman, 1981; Intraub, 1981; Irwin, 1991, 1992; Verfaillie & De Graef, 2000; Verfaillie et al., 1994). Research has demonstrated that scene layout is obtained rapidly and that in addition to conveying a rough portrait of scene structure, it supports scene comprehension at a conceptual level (Biederman, 1981; Oliva & Schyns, 1997). The present research demonstrates that not only is some characterization of layout maintained in transsaccadic memory but that this representation includes a projection of the layout beyond the bounds of the original sensory information. The rapidity of this effect is striking.

While the stimulus is in view, the observer can readily note where the periphery of the picture terminates and the amodally generated part of the representation begins. Once the sensory input is gone, however, the distinction between what was sensory and what was amodal is no longer clear. The representation of the highly expected, amodally generated region just beyond the view may be mistaken as having been seen, thus resulting in boundary extension. In a sense then, the boundary-extended region is already a part of the representation before the stimulus is gone (albeit an amodal part). Only without sensory input is it mistaken for having been seen. Thus, we observed boundary extension in Experiments
2, 3a, and 3b under extremely artificial conditions that would be expected to support a far better memory representation than what would be expected under normal visual scanning. When fixation was maintained, boundary extension occurred following a 42-ms break in the sensory input when the identical view appeared in the same spatial location.

In conclusion, this series of experiments demonstrates the likelihood that boundary extension is available in transsaccadic memory early enough to help place each successive view in its larger spatial framework (Intraub, 2002; Intraub et al., 1992) and to play a role in the integration of successive views. It is well accepted that transsaccadic memory does not retain a point-by-point representation of the world. The current research shows that the transsaccadic representation of expanse is not simply a vague representation that fluctuates with random error. Instead, the spatial representation includes a constructive unidirectional error that would serve as a good prediction of the continuity of layout. We suggest that this rapid projection of layout may be one of the means by which the visual system supports perception of a continuous, coherent world that can never be seen all at once but must be sampled a part at a time.

References


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