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Stream/bounce Perception and the Effect of Depth Cues in Chimpanzees (*Pan troglodytes*)

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40 pages (including 8 figures)

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Abstract

The stream/bounce display represents an ambiguous motion event in which two identical visual objects move toward one another and the objects overlap completely before they pass each another. In our perception, they can be interpreted as either streaming past one another or bouncing off each other. Previous studies have shown that the streaming percept of the display is generic for humans, suggesting the inertial nature of the motion integration process. In this study, chimpanzees took part in behavioral experiments using an object-tracking task to reveal the characteristics of their stream/bounce perception. Chimpanzees did not show a tendency towards a dominant "stream" perception of the stream/bounce stimulus. However, depth cues, such as X-junctions and local motion coherence, promoted the stream percept in chimpanzees. These results suggest both similarities and differences between chimpanzees and humans with respect to motion integration and object individuation processes.
Stream/Bounce Perception and the Effect of Depth Cues in Chimpanzees (Pan troglodytes)

In our dynamically changing visual environment, an important task of the visual system is to pursue and identify moving objects and to maintain their inner representations across time and space. Recent studies have focused a great deal of attention on such perceptual aspects as the identification of moving objects. The investigations of the perceptual characteristics of stream/bounce displays are prominent examples of such studies.

In a stream/bounce display, two discs approach each other, overlap at the center of the display, and then separate again. The discs in this ambiguous display can be interpreted either as streaming past or as bouncing off each other. This stream/bounce display enables examination of how visual systems interpret events involving object movements, as well as the type of information selected and integrated to identify and represent moving objects in time and space. Thus, the visual interpretation of this bistable stream/bounce display has been tested using variable intramodal (Bertenthal, Banton, & Bradbury, 1993; Kanizsa, 1979; Metzger, 1934; A. B. Sekuler & Sekuler, 1999) and intermodal stimulus manipulations (Kawachi & Gyoba, 2006; R. Sekuler, Sekuler, & Lau, 1997; K. Watanabe & Shimojo, 2001b). In general, the human visual system interprets the solo presentation of the ambiguous stream/bounce display as streaming, but this can be altered to a bouncing interpretation, depending on intramodal and crossmodal perturbations. These perceptual tendencies reflect the inertial properties of our visual system (Anstis & Ramachandran, 1987), which bias the recruitment of local motion signals to a straight motion path rather than a returning path, and the vulnerability of the maintenance processes of continuous motion to some perturbations.
In addition to the motion integration processes, depth is an important feature in the perception of stream/bounce displays. The ambiguity of the stream/bounce perception is due to the two-dimensional nature of the display. In the natural world, the two objects stream past or bounce off each other depending on their three-dimensional spatial relationship. Therefore, perception of the stream/bounce display might involve visual interpretation of the depth dimension. Bertenthal, Banton and Bradbury (1993) tested this by adding depth information defined by binocular disparity to the stream/bounce stimuli, and demonstrated that depth cues play a crucial role in the resultant percepts.

Although the characteristics of object identification in this kind of bistable motion perception have been well analyzed in humans, few behavioral studies have directly addressed these issues in other animals. For example, in a field experiment on free-ranging rhesus macaques (Macaca mulatta), Flombaum et al. (2004) demonstrated tunnel effects, which also concern the individuation process of moving objects. They showed real objects (a lemon and a kiwifruit) in motion to monkeys in order to attract their attention. When the first object (a lemon) moved behind an occluder and the second object (a kiwifruit) appeared from the other side of the occluder at the appropriate time, the monkeys generally failed to search for the first object as if they had noticed only one continuously moving object. When continuity of the motion was disrupted, the tunnel effect disappeared, as is the case in human studies. These results suggest that properties of the motion integration process that maintain object identity may be shared by humans and monkeys. However, direct comparisons of the spatiotemporal characteristics of such motion integration processes between humans and nonhuman primates are lacking, and therefore further comparative experiments using controlled stimuli are helpful to explore the phylogenetic background of our visual recognition processes.
In the present study, we examined the characteristics of stream/bounce perception in chimpanzees, the closest evolutionary relative of humans. We conducted five experiments using an object-tracking task to reveal how chimpanzees perceive the moving object in stream/bounce displays and how perception is influenced by depth cues. In the first experiment, we compared the characteristics of stream/bounce perception in chimpanzees and humans. In the second to fifth experiments, we examined the effects of two kinds of depth information, X-junctions (Experiment 2) and motion transparency cues (Experiments 3 and 4), on chimpanzees' stream/bounce perception, contrasting with that of humans (Experiment 5).

General Methods

Since the experiments reported here used similar methods, we first describe procedures common to all the experiments.

Participants

Six chimpanzees, Ai (28 years old, female), Ayumu (4.5 years old, male), Chloe (24 years old, female), Cleo (4.5 years old, male), Pal (4.5 years old, female), and Pendesa (29 years old, female), participated in Experiments 1-4 in this order. Ai did not participate in Experiments 2, 3 and 4 and Pal did not participated in Experiment 2, due to non-availability during the experimental schedule. The participants were experienced in various perceptual-cognitive tasks, such as matching-to-sample (Matsuno, Kawai, & Matsuzawa, 2004; Matsuno & Tomonaga, 2007, 2008) and visual search (Matsuno & Tomonaga, 2006; Tomonaga, 2001), and accustomed to the experimental settings used in this study. The performances of the three young participants were not qualitatively different from those of the three adults; therefore the analyses were conducted on the pooled data for all participants.
The participants lived with 8 other chimpanzees in an environmentally enriched outdoor compound and attached indoor residences (Ochiai & Matsuzawa, 1997). They were not deprived of food at any time during the study. Care and use of the chimpanzees adhered to the 2002 version of the Guide for the Care and Use of Laboratory Primates of the Primate Research Institute, Kyoto University. The research design was approved by the Animal Welfare and Animal Care Committee of the Institute.

Apparatus

Chimpanzees were tested in an experimental booth (approximately 1.8 x 1.8 x 2.0 m) with acrylic panels as walls on all four sides. Stimuli were generated on a Pentium-based computer and displayed on 21-in and 22-in CRT monitors (Totoku CV-213PJ for Ayumu, Cleo and Pal and Mitsubishi TSD-221S for the others) equipped with capacitive and surface acoustic wave touch screens. This system served to present the stimuli, and to accurately record responses (touch locations). Monitor resolution was 1024 x 768 pixels with 8-bit color mode. The refresh rate was 75 Hz and the display was synchronized with the vertical retrace of the monitor. Participants observed the monitor at a viewing distance of about 45 cm without head restraints. The viewing distance was restricted by a transparent acrylic panel, which was attached between the monitor and participants to prevent the damage to the monitor by the chimpanzees. A universal feeder (Biomedica, BUF-310) delivered small pieces of a food reward (apples or peanuts) into a food tray below the monitor.

Object tracking task

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Insert Figure 1 about here
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To assess chimpanzees' perception of ambiguous stream/bounce displays in the absence of verbal reports of their subjective experience, we adopted an object-tracking task (e.g. Pylyshyn & Storm, 1988; Figure 1). In the stream/bounce display, two discs appeared, one each at the right and left side of the display, with one of the discs cued by flickering at the beginning of each trial. Then, the two discs started to approach each other, completely overlapped at the center of the display, and separated again. Participants were required to visually track the initially cued disc through the movement phase and to point to the disc after the two discs stopped moving. Pointing to the disc at the side where the cued disc was initially located indicated that the participant perceived the discs as bouncing, whereas pointing to the disc at the opposite side indicated that the participant perceived streaming.

Experiment 1: Stream/bounce perception in chimpanzees and humans.

In the test sessions, two stimulus conditions were tested by varying the movement speed of the discs. In the 100% overlap condition (stream/bounce stimuli), the two discs completely overlapped at the center of the display. In the 50% overlap condition (intermediate partial-overlap stimuli) used as a control, the two discs stopped overlapping when the edge of one disc reached the center of the other disc. We expected that the participants would more frequently perceive the partial-overlap stimuli as bouncing than the completely overlapping stream/bounce event.

Methods

Participants

Six chimpanzees, and 5 adult humans (females) ranging in age from 18 to 25 (mean = 21.2), participated in the experiment. All the human observers had normal or corrected-to-normal visual acuity.
Stimuli

Displays consisted of the two gray discs, identical in shape and color, subtending about 23 x 23 mm (2.9° x 2.9° of visual angle at a viewing distance of 45 cm) on black background (Figures 2 and 3A, B). The two discs were initially horizontally separated at a center-to-center distance of 218 mm (27.8°). In the training phase with moving stimuli and the test phase, the two discs moved horizontally (with slight vertical displacement in some training conditions: See below and Figure 2C, D). The movement continued until the discs reached the approximate horizontal positions where the two discs were initially located. The separation of terminal positions of discs varied from 193 to 218 mm across trials, so that both the final positions of discs and event durations could not be significant cues for discrimination between bouncing and streaming events. The laboratory was dimly illuminated to prevent reflections on the computer screen.

First training phase (static condition)

Prior to the test sessions, the chimpanzees were trained to track the cued disc. In the first phase of the training, chimpanzees were trained under a static condition in which they were required to simply detect a cued disc at its position.

Each trial was initiated by presenting a warning stimulus (an empty gray square subtending 31 x 31 mm) located at the bottom of the screen. After the participant touched the warning stimulus, it disappeared and two discs appeared. One of the discs started to flash at 18.75 Hz for 600 ms and then turned to the same gray color as the other disc. Then two discs remained stationary during delay period (randomly varied from 680 to 760 ms
across trials: 720 ms on average). After the delay period, a gray square (38 x 38 mm) appeared around each disc to signal the start of the response phase. Chimpanzees were required to indicate the cued disc. A correct response was followed by a chime sound and delivery of a food reward. An incorrect response was followed by a buzzer sound and a 4-s timeout. The interval between the end of the trial and the presentation of the warning stimulus for the next trial was 2-s.

A training session under the static condition consisted of 64 trials. The left-right position of the cued disc was counterbalanced in a session. The training phase was continued until the participant reached the criterion for learning, which was set as >90%

Second training phase (movement condition)

In the second phase of training, chimpanzees were trained to track the movement of the target disc with two types of movement path, unambiguous streaming and unambiguous bouncing. Four kinds of trials, two for the streaming movement and two for the bouncing movement, were prepared (Figure 2). In the training trials, the vertical locations of the two discs were initially set to be differentiated at the length of radius of the disc. Under one condition, the discs moved horizontally to the opposite side of the display (Figure 2A). The discs partially overlapped at the center of the display, but their identity was not ambiguous due to the vertical misalignment; human observers perceived unambiguous streaming of the discs. Under another condition (Figure 2B), the discs moved horizontally and reversed their direction at the point where they touched an imaginary vertical center line of the display. Therefore, the discs did not partially overlap, and human observers perceived unambiguous bouncing. Under these two conditions, the vertical relationship of the two discs was maintained in each trial, so the chimpanzees could detect
a cued disc by attending to the vertical position without tracking the disc. Therefore, in the other two types of trials, the relative vertical positions were reversed between pre- and post- oblique movements (Figures 2C and D). Under these conditions, two discs moved with slight vertical displacements so that their relative vertical positions were reversed at the center of the display. The two discs moved on (streaming) or returned at the point where the discs touched the imaginary vertical center line of the display (bouncing). Initial vertical positions (upper or lower of left or right stimuli) were random across trials.

At the movement phase, the discs were horizontally displaced at 144, 287, 431, or 574 mm (18.1°, 35.4°, 51.1°, or 65.1°, respectively) per second. Each stimulus frame lasted 13.3 ms, and the displacement at a frame was less than the size of the disc radius even at the highest speed.

Each trial proceeded as in the static condition except that the disc movement phase described above was inserted just after the cueing flash of one disc ended. Chimpanzees were required to indicate the position of the target disc when the discs stopped moving and a gray squares appeared around each disc to signal the start of the response phase.

A training session with disc movement consisted of 128 trials (eight trials for each movement speed and movement path condition). Each training phase was continued until the participant reached the criterion for learning, which was set as >90% accuracy in three consecutive sessions.

Test phase

In the test phase, we tested how the stream/bounce stimuli and partial-overlap stimuli were perceived. In test probe trials, two discs were initially horizontally aligned and then horizontally moved toward one another. In the stream/bounce display (Figure 3A), the
discs completely overlapped and moved on to the sides of the display. In the partial-overlap display (Figure 3B), the discs turn their movement directions when the edge of one disc reached at the center of the other disc. The discs moved at 144, 287, 431, or 574 mm (18.1°, 35.4°, 51.1°, or 65.1°, respectively) per second.

Probe trials were intermixed with baseline trials, which were the same as those in the training session with moving discs. For chimpanzees a test session consisted eight probe trials, during which each combination of movement speed and overlap condition was tested, and 128 baseline trials. The eight probe trials appeared randomly in a session. The left-right position of the cued disc was counterbalanced in a session. Each chimpanzee participated in 20 test sessions. Feedback in baseline trials was the same as in the second training phase. In probe trials, no feedback was given, and the next trial started after 2-s interval.

Test in humans

For humans, a test session consisted of 80 probe trials. Ten trials under each combination of overlap and speed condition were randomly intermixed in the session. Each human participated in one test session. Prior to the test session, each human observer received 16 baseline trials. They were instructed to track an initially cued disc and touch it after the discs had stopped.

Results

Training phase

In the first training phase, in which static stimuli were used, chimpanzees required 32 sessions on average to reach the learning criterion (53, 10, 5, 24, 80, and 21 sessions for Ai, Ayumu, Chloe, Cleo, Pal, and Pendesa). Individual differences did not reflect age, but probably on their motivation on the new task.
Performance in the second training phase with moving stimuli (Figure 2) was significantly better than chance level (50%) even in the first session (71.1% correct on average, \( t(5) = 4.1, p < .01 \)) both in the streaming condition (73.9%) and the bouncing condition (68.3%). Chimpanzees required 23 sessions on average to reach criterion (33, 4, 8, 18, 59, and 17 sessions for Ai, Ayumu, Chloe, Cleo, Pal, and Pendesa).

The generalization of performance from the first training phase (static discs) to the second training phase (moving discs) means that chimpanzees spontaneously tracked the cued disc when the disc moved position. This also guaranteed that chimpanzees did not solve the task by learning one-to-one stimulus-response associations (associations of 2 cue positions, 4 movement patterns and left/right responses). This was also supported by another study (Matsuno & Tomonaga, in preparation), conducted after this one, in which four of the six chimpanzees were tested on tracking of a target disc among 4 discs that rotationally moved on a shared path. They successfully tracked the target disc (74.5% correct on average) in the first session, irrespective of the novelty of movement patterns. These results confirm that results in the following test probe trials reflected object tracking abilities rather than the other processes associated with the specific cues in the training displays.

Test phase

The performance of chimpanzees and humans in the probe trials is shown in Figure 4. Response tendencies differed between the two groups. Humans predominantly perceived the ambiguous stream/bounce event (100% overlap condition) as streaming, even
in the partial-overlap condition at high speed. However, chimpanzees predominantly perceived the two discs as bouncing, irrespective of the conditions.

A two-way analysis of variance (ANOVA) of overlap and speed conditions in humans revealed a significant main effect of the overlap condition, $F(1, 4) = 63.4, p < .01$, and an interaction, $F(3, 12) = 10.0, p < .01$. Post-hoc simple main effect analyses revealed that performance under the partial-overlap condition varied with movement speed, $F(3, 12) = 5.08, p < .05$. The simple main effect of overlap condition was significant at the slowest speed, $F(1, 5) = 13.8, p < .05$, but not at other speeds.

In chimpanzees, the main effects of moving speed, $F(3, 15) = 0.8, p > .10$, and degree of overlap, $F(1, 5) = 6.1, p > .05$, were not significant but their interaction was, $F(3, 15) = 4.5, p < .05$. Analysis of the simple main effect revealed that performance varied with movement speed under the partial-overlap condition, $F(3, 15) = 4.2, p < .05$, but not under the 100% overlap stream/bounce condition, $F(3, 15) = 1.7, p > .10$. The simple main effect of overlap condition was significant only at the slowest speed, $F(1, 5) = 13.8, p < .05$.

The average streaming responses of chimpanzees in the 100% overlap stream/bounce display (29.2%) were significantly below chance, $t(5) = 3.0, p < .05$. In contrast, those of humans (81.0%) were significantly above chance, $t(4) = 3.9, p < .01$.

**Discussion**

Humans perceived the stream/bounce stimuli as streaming, consistent with previous studies (Bertenthal, Banton, & Bradbury, 1993; A. B. Sekuler & Sekuler, 1999). They also perceived the partial-overlap stimuli as streaming more frequently when the speed of movement was higher. The effects of speed could be due to the difference in the size of stimulus displacement per frame. At higher speeds, the frame-by-frame displacement was larger relative to the size of the stimuli, and the motion correspondence
between frames was more ambiguous. In contrast to humans', the performance of the chimpanzees unexpectedly showed a tendency to perceive both the stream/bounce stimuli and the partial-overlap stimuli as bouncing. Baseline trial performances were very accurate (92% correct on average), indicating the idea that the chimpanzees tracked the target correctly, regardless of the type of movement. In addition, their performance varied depending on the combination of overlap and speed conditions, similar to humans. Therefore, it is difficult to explain the results as a simple response bias such as neglecting the tracking task and merely selecting the disc on the side where the cued disc initially appeared.

What do these results indicate about differences in visual interpretation between chimpanzees and humans? A difference was seen in the overall frequency of percepts of streaming. On the other hand, the similar speed effects in the partial-overlap condition and the shared direction of the effect of stimulus overlap suggest common perceptual mechanisms for resolving the ambiguous motion event. Therefore, the observed species difference might reflect differences in the degree, but not kind, of perceptual functions needed to track and identify a moving object.

As noted previously, humans sometimes predominantly perceive bouncing (Bertenthal, Banton, & Bradbury, 1993; A. B. Sekuler & Sekuler, 1999; R. Sekuler, Sekuler, & Lau, 1997; K. Watanabe & Shimojo, 1998; K. Watanabe & Shimojo, 2001b). The human visual system has a default tendency to assume the continuous movement of objects to individuate them and maintain object identity. Thus, streaming percepts are dominant in solo presentations of a stream/bounce display. However, when an external perturbation interrupts the continuous motion-integration process, such as the abrupt onset of a click sound and sudden cessation of movement, the percept transforms to bouncing. Thus, one
possible reason for the observed species difference is that the motion integration process to one direction is more easily perturbed in chimpanzees than in humans. Although we did not include explicit perturbations to the display, the stimulus configuration of the stream/bounce display itself may have contained perturbing factors to which chimpanzees might have been more sensitive. This might also apply to the small number of trials in which humans perceived the 100% overlap stream/bounce displays as bouncing. Conceivably, a change in stimulus configuration to be less disruptive to and to facilitate the individuation of each object at their coincidence would promote dominant stream percepts in chimpanzees. To investigate this issue further, the stream/bounce perception of chimpanzees was tested in the following experiments.

**Experiment 2: Stream/bounce perception in chimpanzees using ring stimuli**

In the second experiment, we confirmed that the streaming percept is also the default state in chimpanzees when no external perturbation exists and the two discs are more readily individuated. Because no distractor stimulus was used in Experiment 1, the factors that disturb the chimpanzees' motion integration for one direction could have been the overlap or fusion of the two discs themselves. When the two filled stimuli touched, part of the stimulus edge started to fuse and disappeared. Consequently, the local directional signal of the target disc decreased, and the two stimuli became difficult to individuate. This could impair motion integration. Thus, in this experiment, we used open ring stimuli (Figure 3C), that provided an explicit depth cue (X-junctions) when two objects crossed. When open ring stimuli partially overlapped, their edges were still salient and the junctions of the two stimuli could be an explicit signal for the cross-over of the two objects. In addition, the local motion signal was more salient with the ring stimuli than with the filled disc. This might promote continuous motion integration to one direction.
Methods

Four chimpanzees participated in the experiment. Stimuli and procedures were the same as in Experiment 1 except as described here. Displays consisted of the two gray discs used in Experiment 1 or two gray open ring stimuli that matched the contours of the discs (Figure 3C). At the movement phase, the objects were horizontally displaced at 144 mm (18.1°) per second.

In each test session, the configuration of stimuli (filled disc or open ring) was fixed, and the two stimulus conditions were presented in alternate sessions. A test session consisted of 128 baseline trials, which were the same as in Experiment 1 except for the stimuli (open ring in half of the sessions), and eight probe trials, during which four trials in each overlap condition (stream/bounce condition with 100% overlap or partial-overlap stimuli with 50% overlap) were presented. All responses in probe trials were positively reinforced, as were correct responses in baseline trials. Each chimpanzee participated in 10 sessions (five sessions under each stimulus condition). No additional training sessions were conducted.

Results

Chimpanzee perception of the ring stimuli stream/bounce display became predominantly streaming, whereas bouncing was perceived with the uniformly filled gray discs, as in Experiment 1 (Figure 5).

A two-way ANOVA revealed significant main effects of stimulus, $F(1, 3) = 26.7, p < .05$, and overlap conditions, $F(1, 3) = 154.7, p < .01$, and their interaction, $F(1,3)$
Analysis of the simple main effect confirmed that the percentages of perceived streaming differed significantly between stimulus conditions (filled discs or open rings) in the stream/bounce condition with 100% overlap of stimuli, \(F(1, 3) = 243.0, p < .01\), but not in the partial-overlap condition, \(F(1, 3) = 0.3, p > .10\). The simple effects of the overlap condition were significant under both filled disc and open ring conditions, \(F(1, 3) = 33.0\) and \(211.0, p < .01\).

The average streaming response to the 100% overlap ring stimuli (78.8%) was significantly above chance, \(t(3) = 4.0, p < .05\). The chimpanzees performed very accurately in baseline trials with open rings (93% correct responses on average), similar to their performance with filled discs (94%), \(F(1, 3) = 1.2, p > .10\).

**Discussion**

Chimpanzees predominantly perceived the 100% overlap stream/bounce display as streaming when open ring stimuli were used. The filled disc display was less frequently perceived as streaming, though the percentage increased with increasing overlap, as in Experiment 1. The alternation of chimpanzees' perception with stimulus manipulation further confirmed that chimpanzees did not blindly select the disc on the side where the target initially appeared in probe trials. Furthermore, these results suggest that the difference between chimpanzee and human perception shown in Experiment 1 reflects differences in the degree, but not the kind, of the perceptual function needed to track and identify a moving object.

When the two stimuli in stream/bounce displays overlap, the visual system tends to integrate the local motion signals along the same trajectory and to interpret the event as continuous smooth motion. Such default streaming percepts by chimpanzees, however, may be more easily perturbed. The open ring stimuli provided an explicit cue of a crossover
of the two objects in the depth dimension (X-junctions of the contours), which made
individuation of the two stimuli easier, and unambiguous local motion signals. This
enhanced saliency may compensate for the vulnerability of the motion integration process
in chimpanzees.

Experiment 3: Stream/bounce perception in chimpanzees using random dot stimuli

In Experiment 3, we examined the effect of another depth cue, motion
transparency, on the perception of stream/bounce displays in chimpanzees. Coherently
moving dots produce the percept of motion transparency in humans (Braddick, Wishart, &
Curran, 2002; Edwards & Greenwood, 2005). Using random dot stimuli that coherently
moved in opposite directions from each other, we expected that the two objects would be
perceived at different depth dimensions and would be easily discriminated when they were
crossing. We tested chimpanzee perception of the stream/bounce display with random dot
stimuli compared to a display with uniformly filled stimuli.

Methods

Five chimpanzees participated in the experiment. Stimuli and procedures were
the same as in Experiment 2 except as described here. Displays consisted of the two
identical gray squares or two identical random dot squares of 10% density (Figure 3D).
Each stimulus was subtended about 23 x 23 mm (2.9° x 2.9° of visual angle at a viewing
distance of 45 cm). In the movement phase the squares were horizontally displaced at 144
mm (18.1°) per second.

In each test session, the configuration of stimuli (filled squares or random dot
squares) was fixed, and sessions of each stimulus condition alternated. A test session
consisted of 128 baseline trials, the same as in Experiment 2 except for the stimuli (filled
squares or random dot squares), and eight probe trials, in which four trials of each overlap
condition were presented. All responses in the probe trials were positively reinforced, as were correct responses in baseline trials. Each chimpanzee participated in 10 sessions (five sessions in each stimulus condition), with no additional training sessions.

Results

Chimpanzees tended to perceive the stream/bounce display as streaming when the stimuli were random dot squares, whereas bouncing was perceived with uniformly filled gray squares (Figure 6). A two-way ANOVA revealed significant main effects of the stimulus, $F(1, 4) = 42.0, p < .01$, and overlap conditions, $F(1, 4) = 115.1, p < .01$, and their interaction, $F(1,4) = 15.8, p < .05$. Analysis of the simple main effect confirmed that in the stream/bounce condition with 100% overlap of stimuli the percentages of perceived streaming differed significantly between stimulus conditions, $F(1, 4) = 42.5, p < .01$, but not in the partial-overlap condition, $F(1, 4) = 1.1, p > .10$. The simple main effects of degree of overlaps were significant under both filled square and random dot stimulus conditions, $F(1, 4) = 13.4$ and 76.9, $p < .05$.

The average streaming responses of chimpanzees when perceiving the 100% overlap random dot stimuli (64%) was above chance, at a marginally significant level with a two-tailed $t$-test, $t(4) = 2.5, p = .07$. The performance in baseline trials with random dot squares (93% correct responses on average) was highly accurate, and not significantly different from that with filled squares (94%), $F(1, 4) = 1.1, p > .10$.

Discussion

Chimpanzees tended to perceive the stream/bounce display as streaming when
random dot stimuli were used, whereas when filled gray objects was used streaming was
perceived much less frequently. These results further support the view that chimpanzees'
visual system interprets the stream/bounce bistable motion as streaming, as do human
visual systems, when the two stimuli were more easily individuated.

These results also suggest that chimpanzees have some sensitivity to the depth
cues that induce subjective experiences of motion transparency perception in humans. To
further confirm that the variable perceptual interpretation of the stream/bounce stimuli
depended on motion coherency, in Experiment 4 we used incoherently moving random dot
stimuli.

Experiment 4: Stream/bounce perception in chimpanzees using incoherently moving
random dot stimuli

In addition to the motion coherency, the random dot stimuli used in Experiment
3 had some other features that differed from the filled square stimuli. For example, the
luminance intensity of a random dot square was much less than that of a filled square. In
addition, in the random dot condition the stimulus intensity (dot density) was doubled at the
point where the two stimuli overlapped. Such cues, rather than motion coherency, might
induce the observed alteration in stream/bounce perception. To test the effect of coherent
dot motion on the perception of the overlapping stimuli, the experiment was replicated
using incoherently moving random dot stimuli.

Methods

The same five chimpanzees as in Experiment 3 participated. Under the random
dot condition, the spatial arrangement of dots in the stimulus square was randomly
refreshed at each displacement of stimuli. Thus, the density of random dots was duplicated
at the point where the two squares overlapped, but observers were unable to detect coherent
motion by temporally integrating the proximal dot positions. The performance of the
chimpanzees in the incoherent random dot condition was compared to that in the filled
square condition, which was identical to that in Experiment 3. Other procedures were the
same as in Experiment 3. No additional training sessions were given.

Results

The perception of the stream/bounce display was not at all biased toward
streaming when the random dots were not coherently updated (Figure 7). Performance in
the random dot condition did not differ much from that in the filled square condition, even
when the stimuli overlapped 100%.

A two-way ANOVA revealed a significant main effect of overlap, $F(1, 4) = 17.0, \ p < .05$, but neither the main effect of stimuli, $F(1, 4) = 4.1, \ p > .10$, nor the interaction,
$F(1,4) = 2.5, \ p > .10$, was significant. The streaming responses to the 100% overlap
stream/bounce displays with random dot stimuli (37% on average) was not significantly
different from chance, $t(4) = 2.0, \ p > .10$. Performances in baseline trials with both random
dot squares (93% correct responses) and filled squares (91% correct responses) were
similarly accurate, $F(1, 4) = 1.8, \ p > .10$.

Direct comparison between performance in the coherently moving random dot
condition in Experiment 3 and that in the incoherently moving random dot condition in this
experiment revealed that streaming was perceived significantly more frequently in the
former condition. A two-way ANOVA revealed a significant main effect of overlap
condition, $F(1, 4) = 63.7, \ p < .01$, and the interaction between overlap and coherency.
Analysis of the simple main effect confirmed that the percentages of perceived streaming differed significantly between the experiments in the 100% overlap condition, $F(1, 4) = 15.7, p < .05$, but not in the partial-overlap condition, $F(1, 4) = 0.0, p > .10$.

Discussion

Although the stimuli used were almost the same as in Experiment 3, except for the temporal coherence of each dot, they strongly influenced the chimpanzees' perception. In this experiment, chimpanzees tended to perceive the stream/bounce display as bouncing, irrespective of stimulus type. The streaming responses diminished significantly with incoherently moving random dot stimuli compared to coherently moving random dot stimuli. These results suggest that what promoted the perception of streaming in Experiment 3 was neither the low luminance intensity nor the doubling of the dot density at the overlap of the stimuli. Instead, coherent local motion and, conceivably, the perceived motion transparency arising from the coherent motion likely helped the chimpanzees to maintain their continuous tracking of the target.

Experiment 5: Stream/bounce perception in humans using ring and random dot stimuli

In Experiment 5, human participants were tested with the filled disc, filled square, open ring, and coherent and incoherent random dot stimuli used in Experiments 2 to 4, in order to reevaluate the chimpanzees' performances in Experiments 2 to 4 in comparison with those of humans.

Methods

Six humans (1 male and 5 female) ranging in age from 22 to 29 years (mean = 24.4) were tested. Stimuli were the same as those used in Experiments 2 (filled discs and open rings), 3 and 4 (filled squares and coherently and incoherently moving random dot stimuli). In the movement phase the objects were horizontally displaced at 144 mm (18.1°)
per second as in Experiments 2 to 4.

A test session consisted of 100 probe trials. Ten trials of each combination of 2
overlap (stream/bounce condition with 100% overlap or intermediate partial-overlap
stimuli with 50% overlap) and 5 stimulus configuration conditions (filled discs, open rings,
filled squares, and coherently or coherently moving random dots) were randomly
intermixed in the session. Each participant received a single test session, preceded by 20
baseline trials as used in the tests with chimpanzees. They were instructed to track an
initially cued disc and touch the disc after the discs had stopped.

Results

The results of Experiment 5 are shown in Figure 8. The data for circular (filled
disc and open ring) and rectangular stimuli (filled square and coherent and incoherent
random dot) were analyzed separately, as each corresponds to the analysis in Experiment 2
or Experiments 3-4.

With circular stimuli, humans predominantly perceived streaming of the
stream/bounce stimuli, irrespective of the stimulus configuration. In addition, with open
ring stimuli streaming was perceived more frequently than with filled discs.

A two-way ANOVA of overlap and stimulus configuration conditions revealed
significant main effects of overlap, $F(1,5) = 136.0, p < .01$, and stimulus configuration, $F(1,
5) = 15.1, p < .05$. The interaction was not significant, $F(1, 5) = 1.6, p > .10$. The streaming
responses to 100% overlap with open ring stimuli (98.3% on average) was significantly
above chance, $t(5) = 29.0, p < .01$. 
The participants also predominantly perceived filled square and coherently and incoherently moving random dot stimuli to be streaming when the two stimuli completely overlapped. Partial-overlap events were perceived as streaming less frequently than stream/bounce events, and the frequency varied among the conditions of stimulus configuration. Participants perceived partial-overlap of two random dot conditions as streaming more frequently than partial-overlap of filled squares.

A two-way ANOVA revealed significant main effects of overlap condition, $F(1,5) = 53.9, p < .01$, and stimulus configuration, $F(2, 10) = 7.6, p < .01$, and a significant interaction, $F(2, 10) = 10.9, p < .01$. Post-hoc simple main effect analyses revealed that performance under the partial-overlap condition varied with stimulus configuration, $F(2, 10) = 10.8, p < .05$, but the performance under the stream/bounce condition did not, $F(2, 10) = 0.7, p > .10$. The multiple comparison (paired $t$ test with Holm's correction) of performances with 3 stimulus configurations in the partial-overlap condition revealed that stream responses to filled disc stimuli were significantly less frequent than to coherent and incoherent random dot stimuli, $t(5) = 3.6$ and 3.4, $p < .05$. The two random dot conditions were not significantly different from each other, $t(5) = 2.0, p > .10$. The streaming responses to the 100% overlap stream/bounce displays with coherently and incoherently moving random dot stimuli (98.3 and 96.7 % on average) were significantly above chance, $t(5) = 29.0$ and 22.1, $p < .01$.

**Discussion**

Human participants predominantly perceived the 100% overlap stream/bounce event with open rings and coherently moving random dot stimuli as streaming. In addition, these stimuli were perceived as streaming more than the filled stimuli. These results recall those of chimpanzees in Experiments 2 to 4, though the overall frequency of streaming
percepts by humans was much higher, as also shown in Experiment 1.

One notable difference between chimpanzees and humans is that the latter did not change their responses as a function of the coherence of random dot stimuli. Unlike chimpanzees, human participants showed more streaming responses to the incoherently moving random dot stimuli than to the filled square stimuli (in the partial-overlap condition).

More frequent streaming percepts with incoherent random dot stimuli compared to filled stimuli may be explained by the increased salience of the target identity when the two stimuli crossed over. As mentioned above, the filled target fused with the other filled stimulus and target identity became ambiguous at cross-over. Though incoherent random dot stimuli did not have coherent local motion signals, the borders of two crossed-over stimuli were still detected by virtue of the difference of the dot density between the areas where the two random dot stimuli were and were not superimposed. Such cues help human participants to keep attentively track the object motion longer and bias the human percepts toward more frequent streaming. Chimpanzees may be insensitive to such cues, or the effect may be too small to be statistically significant.

General Discussion

This study investigated stream/bounce perception in chimpanzees and compared it with that in humans. We first showed that the stream/bounce perception of chimpanzees with filled discs differed from that of humans. Whereas humans predominantly perceived the stimuli to be streaming, chimpanzees exhibited far more bounce responses. Further experiments using ring and random dot stream/bounce stimuli revealed that chimpanzees also predominantly perceived the stimuli as streaming when the two stimuli were more salient and discriminable from each other due to additional depth cues.
These results reveal both differences and similarities between the perceptual processes of chimpanzees and humans. Chimpanzees, like humans, perceived ambiguous stream/bounce events as streaming when the two stimuli were readily discriminable from each other due to the addition of depth cues, thus indicating that the tendency to keep track of a unidirectional and continuous movement, and its resultant default streaming percepts, is shared by chimpanzees and humans. However, this tendency appears to be more readily perturbed in chimpanzees. Previous studies have identified multiple factors that can alternate stream/bounce perception (Bertenthal, Banton, & Bradbury, 1993; Grassi & Casco, 2009; Kawabe & Miura, 2006; A. B. Sekuler & Sekuler, 1999; K. Watanabe & Shimojo, 2001a, 2001b), and determining a single mechanism that would fully explain the vulnerability of the streaming perception in chimpanzees is difficult.

One possible explanation is that the spatiotemporal integration process of local motion signals may differ between the species. Previous studies in humans have proposed that the dominant streaming perception can be explained by the intrinsic directional bias involving the temporal integration arising from the cooperative interaction between local motion detectors (e.g. Bertenthal, Banton, & Bradbury, 1993). In our study, one prominent difference between stimuli to which chimpanzees did and did not perceive more streaming (open ring and coherent random dot vs. filled and incoherent random dot stimuli) was concerned continuity of the local motion at the brief interval when target and nontarget stimuli crossed over. This implies that the temporal integration window may be smaller in chimpanzees than in humans and that chimpanzees' perception may rely more on the local motion mechanism. Although the temporal integration process has not been well studied in chimpanzees, several comparative studies between chimpanzees and humans revealed species differences in spatial integration processes (e.g. Fagot & Tomonaga, 1999; Fagot &
Tomonaga, 2001). In these studies, chimpanzees were less sensitive to the global configuration of visual stimuli. The relative local bias in chimpanzees' visual processing may be common between temporal and spatial domains. Another possible explanation related to the motion integration process is a difference in the quality of sustained attention between chimpanzees and humans. Visual attention has a critical role in motion perception, selecting and integrating visual information across time and space and keeping track of and identifying moving objects (e.g. Cavanagh, 1992; Choi & Scholl, 2004; Pylyshyn & Storm, 1988). Research on the effects of the state of visual attention on stream/bounce perception has revealed that poorer attentional resources directed to the moving object cause more frequent bouncing percepts (K. Watanabe & Shimojo, 1998), suggesting that sufficient quality of attention is required for constant motion-integration processing and thus the perception of streaming. Thus, the more frequent bouncing responses of chimpanzees might indicate that chimpanzees' attention is more readily disrupted than that of humans.

Species differences in the quality of pursuit of moving objects may also explain the different perception of stream/bounce displays. In our experiments, both chimpanzee and human participants were allowed to observe the display freely, without fixation. Pursuit eye movements can modulate perceived object motion (Baker & Graf, 2010; Kerzel, 2000) and promote streaming percepts in our display. Therefore, the difference in pursuit eye movements between chimpanzees and humans could influence how each species perceives stream/bounce displays. However, previous studies have shown that humans perceive streaming of the stream/bounce stimuli even when the eyes were fixated (e.g. Bertenthal, Banton, Bradbury, 1993; Sekuler & Sekuler, 1999), and the frequency of streaming percepts in those study (approximately 80 to 95%) was similar to that in this study.
Therefore the higher rate of streaming perception by humans in our study can not be
explained simply by effects of pursuit eye movement. Comparative data on smooth pursuit
of moving objects between chimpanzees and humans are lacking; such data are needed to
examine this issue in more detail.

The effect of visual experience during training sessions should also be
considered. Prior to test sessions, our chimpanzees experienced an equal number of
unambiguous streaming and bouncing displays in training trials. However, in the natural
world, a plausible assumption is that an object moving in one direction continues to move
in that direction (Hall-Haro, Johnson, Price, Vance, & Kiorpes, 2008; Spelke, 1994); and
any bouncing event is accidental (K. Watanabe & Shimojo, 2001a). This is consistent with
the tendency of our visual system to interpret bistable ambiguous visual information, such
as a stream/bounce display, as unidirectional movement (Anstis & Ramachandran, 1987;
Bertenthal, Banton, & Bradbury, 1993). Thus, equalized experiences of streaming and
bouncing percepts are unrealistic. This abnormally increased experience of bouncing
percepts may have distorted the prior stochastic expectancy of the event perception by the
chimpanzees, which may have biased their responses to include more bouncing percepts.

However, our data do not seem to support this idea. First, the number of training
sessions experienced by each chimpanzee (ranging from 4 to 59 sessions) and percentages
of perceived bouncing were not positively correlated (Pearson's correlation; \( r = -.14; p
>.10 \)). Second, in additional tests, human participants, who received prolonged experiences
of baseline trials, maintained their predominant streaming percepts (Supplementary
Materials: Experiment S1). The comparative test in humans, however, assessed the effect of
limited number of baseline trials (256 trials) and was not fully equivalent to the tests in
chimpanzees. The effect of longer-term visual experience on the stream/bounce perception
An alternative interpretation of the results focuses on differences in the perception of depth when two-dimensional discs are presented on a flat monitor surface. Our interpretation of a computerized stream/bounce display may relate to structural constraints and the physical laws of the three-dimensional natural world, in which two solid objects on the same depth plane collide and those on different planes pass through (Scholl & Nakayama, 2002; A. B. Sekuler & Sekuler, 1999; K. Watanabe & Shimojo, 2001a). When humans perceive the stream/bounce stimuli as streaming, the two objects are perceived to be on different surfaces, not only in the open ring and random dot conditions but also in the filled disc condition. Chimpanzees may not perceive such depth dimensions on the planar surface of a CRT monitor when explicit depth cues are not given to the objects, and their perceptual processing may interpret objects on the same surface as never passing through, due to physical laws. Although evidence suggests that chimpanzees are capable of correlating projected movies to the real world (e.g. Hirata, 2007; Leighty, Menzel, & Fragaszy, 2008; Menzel, Savage-Rumbaugh, & Lawson, 1985) and perceiving depth from some 2D pictorial cues (Imura & Tomonaga, 2003, 2009; Imura, Tomonaga, & Yagi, 2008), we cannot be sure that they employed these abilities in viewing our stimulus displays without such cues. Given that 2D iconic expressions of the 3D world are a human-specific innovation and that human and chimpanzee participants differed enormously in previous exposure to such media, species differences in responding to such computerized graphical images would not be surprising.

According to this explanation, streaming perception with open ring and coherent random dot stimuli may reflect chimpanzees' sensitivity to two kinds of explicit depth cues, X-junctions and motion transparency. As noted above, the ambiguity of stream/bounce
displays derives from a conflict between the perception of two moving objects with depth deviation (streaming) and without depth deviation (bouncing). The increase in streaming percepts supports the validity of stimulus manipulations as depth cues for chimpanzees' perception.

X-junction is known to be a strong cue for the detection of transparency or overlapping of objects in human vision (e.g., Dresp, Durand, & Grossberg, 2002; Kanizsa, 1979; T. Watanabe & Cavanagh, 1993). Several studies in nonhuman primates have also investigated the role of junctions (T, L, or X) in the perception of object overlap, revealing that monkeys and apes perceive the occlusion or transparency of two-dimensionally displayed objects using the clues of the junctions (e.g. Fujita & Giersch, 2005; Nagasaka, Nakata, & Osada, 2009; Sato, Kanazawa, & Fujita, 1997; Sugita, 1999). Our results, which demonstrate that the addition of X-junctions promoted the streaming percepts, are consistent with the results of these studies.

Aggregations of coherently moving dots moving in opposite directions also strongly induce depth percepts of two different planes in humans (Braddick, Wishart, & Curran, 2002; Edwards & Greenwood, 2005). On the other hand, the perception of motion transparency has not received much attention in comparative perception studies, partly because it is difficult to find behavioral indices to assess such subjective percepts in nonverbal organisms. The method used here to measure the effects of motion transparency cues on stream/bounce perception may be valuable for assessing such visual sensitivity in other species.

In conclusion, our study is the first to demonstrate the perception of stream/bounce displays by chimpanzees. The visual interpretation of chimpanzees differed from that of humans, suggesting species differences in the process of identification of
moving objects. Our study also provides evidence of the chimpanzees' sensitivity to two kinds of depth cues, X-junctions and motion transparency, which could be used to perceptually resolve the interpretations of ambiguous motion events. Further studies comparing other animals would be valuable for tracking the evolutionary origins of the perceptual mechanisms that underlie our representation of the dynamic visual world.
References


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Figure captions

Figure 1. Schematic diagram illustrating a trial in the object-tracking task.

Figure 2. Depictions of stimulus displays used in the training sessions and the baseline trials. Arrows indicate motion. A: unambiguous streaming display without vertical displacement. B: unambiguous bouncing display without vertical displacement. C: unambiguous streaming display with vertical displacement. D: unambiguous bouncing display with vertical displacement.


Figure 4. Collected response data from probe trials in Experiment 1. The vertical axis represents the percentage of stream responses; 0% indicates a bounce response from all subjects in all of the trials, and 100% indicates complete stream responses. The left four bars represent the 100% overlap condition, and the others represent the 50% overlap condition. Each bar indicates a different movement speed condition. Error bars are 1 SE.

Figure 5. Percentages of streaming responses in Experiment 2. The left two bars represent the 100% overlap condition, and the others represent the 50% overlap condition. Each bar indicates a different stimulus condition. Error bars are 1 SE.
Figure 6. Percentages of streaming responses in Experiment 3. The left two bars represent the 100% overlap condition, and the others represent the 50% overlap condition. Each bar indicates a different stimulus condition. Error bars are 1 SE.

Figure 7. Percentages of streaming responses in Experiment 4. The left two bars represent the 100% overlap condition, and the others represent the 50% overlap condition. Each bar indicates a different stimulus condition. Error bars are 1 SE.

Figure 8. Percentages of streaming responses in Experiment 5. The left graph represents the conditions using circular stimuli as in Experiment 2. The right graph represents the conditions using rectangular stimuli as in Experiments 3 and 4.
Figure 1.
Figure 2.

A. Streaming without vertical displacement
B. Bouncing without vertical displacement
C. Streaming with vertical displacement
D. Bouncing with vertical displacement
Figure 3

A. Stream/bounce (disc)

B. Partial overlap (disc)

C. Stream/bounce (ring)

D. Stream/bounce (dots)
Figure 5

% perceived streaming

Filled disc
Open ring

100% overlap
50% overlap
Figure 6
Figure 7

- Filled square
- Incoherent random dot

% perceived streaming

100% overlap  50% overlap

0  20  40  60  80
Figure 8