Multidimensional internal dynamics underlying the perception of motion

Mark Wexler

When ambiguous visual stimuli are presented continuously, they often lead to oscillations between usually two perceptions. Because of these oscillations, it has been thought that the underlying neural dynamics also arises from a binary or two-state system. Contradicting the binary assumption, it has been shown recently that the perception of some ambiguous stimuli is governed by continuously varying internal states, measured as biases that differ considerably from one observer to the next and that can also evolve over time (Wexler, Duyck, & Mamassian, 2015). Here I study bias patterns in the motion quartet, an ambiguous apparent motion stimulus, as the quartet’s orientation is varied. The bias patterns are robustly idiosyncratic, and are even more complex than those that have been described previously. There are two qualitatively different bias types: Some observers prefer a translation axis, while others show preference for a rotation direction. Each type also varies parametrically: the orientation of the preferred axis, and the direction of preferred rotation. There are also clear cases of combination of the two bias types. When measured repeatedly over 9 hr, the bias patterns usually remain stable, but also sometimes evolve both parametrically (e.g., change of preferred axis), as well as across bias type (change from axial to rotational bias). Control experiments revealed that the variety of bias patterns observed across subjects, and their changes over time, are not due to voluntary decisions. Overall, these results exhibit the multidimensional complexity of internal states underlying the perception of even simple stimuli.

Introduction

Reversible or bistable stimuli have long attracted attention because, in the absence of external cues that favor a particular interpretation, the dynamics of perceptual choice provides a uniquely sensitive probe of the neural mechanisms of perception (Necker, 1832; Wheatstone, 1838; Rubin, 1921; Wallach, 1935; Attneave, 1971; Leopold & Logothetis, 1999; Blake & Logothetis, 2002; Long & Toppino, 2004; Sterzer, Kleinschmidt, & Rees, 2009). Such ambiguous stimuli have been used to study the mechanisms of adaptation, priming, context, history, memory, cue integration, and cortical dynamics, to cite some examples. It is often assumed that neural dynamics is isomorphic to overt perceptual dynamics. For example, if the perception of stimulus switches between two interpretations, then the underlying neural representations must also be binary. Here I show that this assumption underestimates the complexity of the underlying brain circuits. By measuring the population distribution and temporal evolution of the individual observers’ biases in the perception of a well-known twofold-ambiguous stimulus—the motion quartet—I show that the biases reflect complex and multidimensional internal states.

The motion quartet (also called the “bistable quartet” or “stroboscopic alternative motion”) is a two-frame ambiguous motion stimulus, illustrated in Figure 1a (Schiller, 1933; Gengerelli, 1948; Ramachandran & Anstis, 1983b). In the first frame of the usual version of the quartet, two identical objects appear at opposite corners of an imaginary square; in the second frame, the same two objects appear in the other two corners. If the visual system matches each of the objects in the second frame to a single and unique object in the first frame, two apparent motions can be perceived, illustrated in Figure 1b and c. Because the distances in the two solutions are the same, neither the slow-motion prior nor any other perceptual cue can disambiguate the two solutions. The motion quartet has been used to probe contextual and priming effects in motion perception (Ramachandran & Anstis, 1983a, 1983b; Anstis & Ramachandran, 1987), hysteresis and sequence effects (Hock, Kelso, & Schöner, 1993; Maloney, Martello, Sahm, & Spillmann, 2005), and analogies between visual and tactile motion perception (Carter, Konkle, Wang, Hayward, & Moore, 2008; Liai, Bach, Tebartz van Elst, Heinrich, & Kornmeier, 2016; Haladjian, Anstis, Seizova-Cajic, Wexler, & Cavanagh, 2018), as well as to study the neural correlates of visual consciousness (Sterzer & Kleinschmidt, 2007; Zhang et al., 2012) and the effects of...
neuroanatomy on motion perception (Genç et al., 2011; Shimono, Mano, & Niki, 2012).

The subject of interindividual differences in vision has always elicited a certain amount of interest, going back to philosophers’ questions about the subjective appearance of colors (Locke, 1690) and the “personal equation” in astronomy (Bessel, 1823), as well as more recently (Coren & Porac, 1987; Wilmer, 2008; Grzeczkowski, Clarke, Francis, Mast, & Herzog, 2017), including in the context of multistable stimuli (Kleinschmidt, Sterzer, & Rees, 2012). Nevertheless, most visual psychophysics is carried out with the tacit assumption that the appearance of suprathreshold stimuli is invariant across observers, or that any variations are small and can be ignored. This assumption is probably the reason why the large individual differences in the appearance of The Dress—a photograph that spread on the Internet in 2015—impressed not only naive observers but psychophysicists as well (Witzel, O’Regan, & Hansmann-Roth, 2017).

In the current study I measure individual biases in the perception of the motion quartet. These biases turn out to have novel and interesting properties. In the first of the two experiments described here, I measured the bias patterns for the motion quartet in a relatively large sample (106 subjects). Conventional studies of ambiguous stimuli like the motion quartet tend to concentrate on their bistable aspect and present the exact same
stimulus either continuously or repeatedly with intervening blanks. In the current studies, on the other hand, the two-frame sequence was presented only once on each trial, and its overall orientation varied randomly from trial to trial. Not having witnessed spontaneous oscillations from repeated exposure to the same stimulus, many subjects did not realize that the stimuli are ambiguous, judging from their qualitative reports. In the second experiment, I measured the time evolution of the individual bias patterns by repeatedly measuring them (45 times) over a period of 9 hr.

To get a good feel for the stimulus, I urge the reader to try an online demonstration at http://lab-perception.org/demo/q. The demonstration is a briefer version of Experiment 1. Additionally, after the 24 trials (which take about a minute or two to complete), the reader will be shown his or her own bias pattern, which can be compared to the data in this article.

Experiment 1: Population sample of bias patterns

This experiment measured bias patterns for the motion quartet in a fairly large sample of subjects ($N = 106$), at one point in time. The quartet was presented once (two frames) on every trial, with its orientation varying randomly from trial to trial.

Methods

In order to obtain a large sample, this experiment was performed by subjects inside an Internet browser on their own computers outside the laboratory. I used methods developed in an earlier study (Wexler et al., 2015) in order to control stimuli and experimental procedures as much as possible in this unconventional setting (see below for details). Similar results were obtained in a more traditional laboratory setting, including pilot studies for this experiment, as well as in the second experiment presented later in this article.

Stimuli

Stimuli were displayed and responses collected in an Internet browser, with the experiment programmed in Javascript and HTML5. Tablets and telephones were not allowed, in order to control monitor orientation. It was possible to measure video frame rate, which was above 30 Hz for all subjects. Subjects were encouraged to make the browser window full screen or as large as possible, to minimize extraneous stimuli; the browser window filled at least 60% of the monitor’s area in all cases but one (where it filled 47%). Subjects were instructed to sit upright at a comfortable distance from the monitor. For further details, see Wexler et al. (2015).

Stimulus geometry will be given in pixels. It is impossible to translate from pixels to degrees of visual angle, because both pixel size and the distance between subject and monitor are unknown. Additionally, some systems with high-resolution monitors scale pixel units in order to make small text and images easier to see, so that a logical pixel may correspond to more than one physical pixel. However, at the subjects’ median resolution of 1,366 × 768, assuming a common 13-in. 16/9 diagonal monitor viewed at a typical distance of 50 cm, and no pixel scaling, 150 pixels (the distance from the center of the imaginary square to its corners) is about 3.3° of visual angle.

The main stimulus consisted of two frames, each containing two disks lying on the opposite corners of an imaginary square (labeled 1 and 2 in Figure 1a). The disks had radii of 20 pixels, and were drawn in black on a medium gray background. They were drawn so that their centers were at a distance of 150 pixels from the center of the window. The entire figure (the imaginary square) was oriented at one of 24 angles: 3.75°, 11.25°, 18.75°, . . . , 176.25°. (The orientations were between 0° and 180° because adding 180° to the stimulus orientation leaves the stimulus unchanged.) In addition to the disks, a white cross (30-pixels wide) was displayed in the center of the window, which the subject was instructed to fixate. The fixation cross was displayed alone for 750 ms, followed by the first frame with the fixation cross for 500 ms, immediately followed by the second frame and cross for 500 ms. Two icons showing the clockwise (CW) and counterclockwise (CCW) percepts (similar to Figure 1b and c) were then displayed on the right and left sides of the window (chosen randomly), each at a distance of 200 pixels from the window’s horizontal center. The subject selected and clicked on icon corresponding to his or her perceived motion, using the computer mouse.

Procedure

The main experimental block consisted of 48 trials, composed of two subblocks of 24 trials, in which the stimuli were presented in each of the 24 angles, in random order (different random order for every subject). The order of the stimuli was identical in the two subblocks, in order to measure subjects’ consistency. Prior to the main experimental block, subjects were given written instructions and performed 20 practice trials, similar to the trials in the main block but excluded from data analysis. Trials followed one another without pause, with breaks every 16 trials. The main experimental block had a median duration of 2.6 min.
Subjects

One hundred and six subjects took part in the experiment (40 men and 66 women, ages 18–66, median 27 years). Prior to taking part in the current experiment, these subjects had participated in a separate Internet-based longitudinal study on the evolution of individual biases in the perception of ambiguous visual stimuli (experiment 3 in Wexler et al., 2015). The stimuli used in the prior longitudinal study were different from those used here. The subjects had been originally recruited through an announcement to a mailing list of several thousand people willing to participate in cognitive science experiments (organized by the Relais d’information sur les sciences de la cognition [RISC], Paris, France). The current experiment was carried out as the final session of the prior longitudinal study. Subjects were paid 1€ for their participation in this session. Informed consent was obtained prior to the experiment in accordance with the Declaration of Helsinki. In France, the legal ethics committees do not examine noninvasive behavioral studies.

Results

The results of 12 representative subjects are shown in Figure 2, using the graphical representation explained in Figure 1d and e. For clarity, only the data from the first subblock are shown. The full data of all 106 subjects and both subblocks are shown in Supplementary Figure S1.

A simple hypothesis is that the perceptual decision on each trial is a stochastic choice, with 50% chance of responding CW and 50% CCW, independent of all other trials. If so, the response patterns should resemble random, isotropic mixtures of red and green arrows (because stimulus orientations were presented in random order). This is visibly not the case for a majority of subjects (as can be seen in Figure 2 and Supplementary Figure S1), whose data show large-scale spatial patterns.

The randomness of the response patterns was tested using a procedure based on the number of boundaries between CW and CCW responses. If one considers $n = 48$ independent binary variables on a circle (with equal probability of CW and CCW values), the number of boundaries $b$ between CW and CCW regions follows a binomial distribution $P(b, n) = n!/(2^{n-1} b!(n-b)!)$, which was taken as the null hypothesis. The large-scale spatial structure of many of the response patterns, with large homogeneous regions, results in an improbably small number of boundaries, as in the Wald-Wolfowitz runs test (Wald & Wolfowitz, 1940). The Benjamini-Hochberg correction (Benjamini & Hochberg, 1995) was applied to take into account multiple tests, with false-discovery ratio set to 0.05. Using this procedure, I found that 71% of the subjects had response patterns that differed significantly from random and independent choices of CW or CCW at each orientation.

In fact, as can be seen in Figure 2 and Supplementary Figure S1, most subjects’ response patterns seem to cluster around two distinct types. In one pattern, many subjects’ responses show four alternating CW/CCW regions of about equal size (i.e., 90°). Subjects presenting this pattern are shown at the beginning of Figure 2 and Supplementary Figure S1. This pattern is indicative of an axial bias, or a preference for translation, in either direction, closest to a particular axis. This is illustrated in Figure 3. The overall
orientation of the four-region pattern determines the preferred axis, as shown in the top line of Figure 3.

A second, qualitatively different pattern can be seen in subjects at the end of Figure 2 and Supplementary Figure S1: all or nearly all responses being either CW or CCW. This kind of pattern will be called a rotational bias, a preference for motion compatible with either CW or CCW rotation.

When individual data was tested for the two bias patterns, sizable fractions of the subject pool showed statistically significant biases of one or the other type: 41% of the subjects had a significant axial bias, while 45% had a significant rotational bias (with corrections for multiple tests using a Benjamini-Hochberg procedure with a false-discovery rate of 0.05). The axial bias was tested as follows: For each trial, I took one of the two reported translation directions, and multiplied the angle by 2 (the standard way of handling axial data), and applied the Rayleigh test (Mardia & Jupp, 1999). The orientation of the preferred axis (the black arrows in Figure 2 and Supplementary Figure S1 was calculated by multiplying the angle of the perceived axis vector by 2, taking a circular mean, and diving the resulting angle by 2). Rotational bias was tested using a sign test on the raw response data.

Individual biases are represented in Figure 4 as points in a 3-D space. The first two dimensions are the Cartesian components of the preferred axis vector, whose direction is the orientation of the preferred translation axis, and whose length is the strength of the axial bias. (Since axial bias is bidirectional, each subject is represented by two opposite points in the first two dimensions.) The third dimension is the rotational bias, calculated by taking the mean of each subject’s raw responses, with +1 corresponding to CW and -1 to CCW. The subjects with a significant axial bias are colored blue in Figure 4a, and those having a significant rotational bias colored yellow.

Interestingly, there were three subjects who had statistically significant biases of both the axial and the rotational types (colored red in Figure 4). This result is important because it shows that the two bias types are not mutually exclusive. Rather, individual biases are properly seen as multidimensional combinations of the two underlying bias types. Although only a small fraction of subjects had significant axial and rotational biases, many more appear to have combinations of the two bias types (Supplementary Figure S1), and would have likely been significant in an experiment with a larger volume of data (more than 48 trials).

In the subjects with significantly axial patterns, the distribution of axial directions was highly nonuniform (see Figure 4b), with peaks in the cardinal directions (88% of the subjects had axis directions closer to the cardinals than to the main diagonals, significantly above 50% by bootstrap), and with a higher peak in the vertical than in the horizontal direction (63% of the axes closer to vertical than horizontal, significantly above 50% by bootstrap). Of the subjects with a significant rotational pattern of responses, a majority (73%, significantly above 50% by bootstrap) had a bias in the CW direction (Figure 4c).

Discussion

The results of this experiment have shown that when the motion quartet is presented at different orientations on successive trials, perceptual decisions about the direction of motion depend on large-scale bias patterns. The bias patterns seem to be multidimensional combinations of two extreme types. The first type is axial bias, favoring translation, in either direction, along a given axis (which can vary between 0° and 180°). The second type of bias pattern is rotational, favoring motion that is globally CW or CCW. Quite a few subjects have bias patterns that are pure or almost pure axial, or rotational. Other subjects have patterns that seem like combinations of axial and rotational, with
several subjects having statistically significant biases of both types.

If a subject has an axial bias with a vertical axis, or an axial bias with a horizontal axis, or a CW rotational bias, or a CCW one, or some combination of axial and rotational biases—is this a permanent condition? In our previous work on individual perceptual biases, it has been shown that such biases are on the whole stable, but also sometimes change, either continuously or discontinuously (Wexler et al., 2015). In our previous study, we probed the temporal evolution of the individual biases with daily measurements over a period of several months. In the second experiment in the current study, I measured changes over time in the bias pattern for the motion quartet, testing each subject 45 times over a period of 9 hr.

**Experiment 2: Temporal evolution of individual biases**

In this experiment I directly tested the stability of the individual bias patterns, by measuring them every 12 min over a period of 9 hr, for a total of 45 measurements in every subject. One possibility is that the bias patterns are stable, at least over this duration. A second possibility is that the bias type—axial or rotational—remains stable, but with the axis or direction of rotation changing over time. Finally, it is possible that biases vary over their parameter space (such as Figure 4a), which would additionally allow changes from axial to rotational biases, or vice versa.

There is a possible criticism of the conclusion from Experiment 1 that the individual differences in the bias patterns are somehow intrinsic to the subject: The variety of individual response patterns (e.g., Figure 2) could have been due to the different order of trials for every subject. For example, perhaps the first few orientations seen by each subject somehow fixed the subject’s subsequent response pattern. If so, one should find much less interindividual variation if the trials were in the same order for all subjects. This was tested in Experiment 2, in which trials were always in one and the same (but random) order for all subjects, and for all blocks for every subject.

**Methods**

In contrast to Experiment 1, this experiment was performed in the laboratory.
The stimulus was very similar to that in Experiment 1. On every block a single repetition of the same 24 orientations as in Experiment 1 was tested. The orientations were presented in random order, but in the same random order on every block, for every subject. (This was done in order to test whether the variety of individual response patterns found in Experiment 1 could have been due to the different order of trials for every subject.)

Stimulus disks were presented with their centers at 2° of visual angle from the center of the monitor. Each disk had radius of 0.25° of visual angle, and was white (112 cd/m²) on a black background (0.85 cd/m²). Subjects were instructed to fixate a red disk drawn at the center of the monitor (radius 0.1° of visual angle). On each trial, the fixation disk was presented alone for 500 ms, followed each of the two frames (with the fixation disk) presented for 400 ms, followed by the response icons, as in Experiment 1. Stimuli were presented on a Sony GDM F520 monitor (resolution 1,024 ×3768, refresh rate 120 Hz, size approximately 38°×29° dva) at an approximate distance of 57.3 cm from the subject, whose head was restrained with a chinrest during the experimental blocks. The experiment was performed in a room with normal indoor lighting.

Blocks began every 12 min. The median duration of one block was about 54 s, so subjects had a break of about 11 min between blocks. During these breaks they were free to do what they wished, for example to read, study, listen to music, eat, and so on; they could also leave the experimental room, for example, to use the restroom or smoke. Thirty seconds before the start of each block, the computer emitted a series of beeps and flashes to warn the subject to put his or her head into the chinrest, and that the block was about to begin. Before the start of the series of 45 blocks, subjects performed a practice block, identical to the experimental blocks. Six subjects took part in the experiment (ages 23–37, median 28 years, four women and two men), and were paid 10€/hr.

Results

The complete time series of the bias patterns of all six subjects are shown in Supplementary Figure S2. The first thing to note about these data is that, clearly, the samples are not independent of one another; in most cases, the shape of the bias pattern is retained from one measurement to the next. To check this, I performed a test based on the fact that the number of differences between two independent binary samples follows the binomial distribution. The number of differences in successive samples was calculated, and the Benjamini-Hochberg procedure (false discovery rate 0.05) was used to correct for multiple tests. In 79% of the cases, responses in successive blocks differed by significantly fewer trials than predicted by the null (no-memory) hypothesis. Thus, one can conclude that bias patterns are generally stable from one measurement to the next, 12 min later.

Nevertheless, there are changes in the bias patterns over time. Three examples of change, excerpted from the time series of three separate subjects (12 blocks, about 2.2 hr), are shown in Figure 5. In the first series in Figure 5 a horizontal axial bias becomes roughly vertical, while in the second series a rotational bias cycles between CW and CCW several times. In the third series a strong axial bias undergoes a transition to an almost completely rotational bias, while seemingly retaining a small “memory” of its previous axial direction. (This memory could be the result of taking the shortest path between the two biases in the 3-D space of Figure 4a.) Thus, the bias patterns seem to be able to change within type—change of preferred axis in axial bias, change of preferred rotation direction in rotational bias—as well as between types—change from axial to (mostly) rotational bias.

The behavior of the time series was analyzed in a different way, by calculating mean response (with individual responses coded as +1 = CW, −1 = CCW) for each session, and analyzing the time series of the means using the autoregressive integrated moving-average
(ARIMA) framework (Box & Jenkins, 2008). The analysis, using the forecast package in R (Hyndman & Khandakar, 2008), automatically identified the best model by searching through ARIMA$(p, d, q)$ models with $p, d, q \in \{0, 1\}$. The search was performed by applying the KPSS unit-root test (Kwiatkowski, Phillips, Schmidt, & Shin, 1992) to determine the differencing order $(d)$, and then searching the $p, q$ subspace for the model that best fits the series, subject to penalties from the Akaike information criterion (Hyndman & Khandakar, 2008). The same analysis was carried out on the 3-month time series by Wexler, Duyck, and Mamassian (2015).

The most common time series model found was ARIMA$(0, 1, 1)$, identified by the automatic method for three subjects as the best model. This model is equivalent to an underlying random-walk or Brownian-noise variable, observed through independent measurement noise (Box & Jenkins, 2008; Wexler et al., 2015). The time series of two other subjects were fitted by ARIMA$(1, 1, 0)$ and ARIMA$(1, 0, 0)$ models, while the final subject—the same one who had no significant autocorrelations—was fitted with a white-noise ARIMA$(0, 0, 0)$ model.

Discussion

By repeatedly measuring bias patterns to obtain time series of significant length, the results of this experiment showed that the bias patterns at successive samples (separated by 12 min) were not independent, but generally remain stable over time. However, there were significant changes, both within bias types (changes in preferred axis for axial biases, and in preferred rotation direction for rotational biases), and between bias types (an axial bias that became a nearly complete rotational bias).

A time-series analysis of mean responses, averaged over stimulus orientation, revealed that the time series of a plurality of our subjects were best described by one particular model, a random walk measured through noise. The same result was obtained by Wexler et al. (2015) on different measures coming from biases pertaining to different stimuli (biases for surface orientation in structure-from-motion stimuli, and biases for motion direction in transparency-from-motion stimuli). This model, as well as the other time-series results, fit in with a picture that the individual biases are reflections (through noisy measurements) of persistent brain states. These brain states evolve, either through exposure to external stimuli or through internal neural dynamics, and their evolution can at least partly be described as a random walk.

It should be noted that, although in this experiment trials were always in the same order for all subjects and all blocks (in contrast to Experiment 1), there was wide variability, both between and within subjects (Supplementary Figure S2). Thus, variation in response patterns is due to variable and evolving states of the visual system, rather than different order of stimuli in Experiment 1. Nor could have the between-subjects variability in Experiment 1 been (all) due to differences in experimental conditions in that internet-based experiment, as the data in Experiment 2—performed in controlled laboratory conditions—also shows a great deal of between-subjects variability.

Experiment 3: Involuntary perception or voluntary control?

So far we have implicitly interpreted the differences in bias patterns—both between subjects (Experiments 1 and 2) and over time within the same subject (Experiment 2)—as involuntary perceptual effects due to internal state changes in the visual system. However, some observers can sometimes voluntarily control their percepts. Although subjects certainly weren’t instructed to voluntarily control their perceptions, and had no obvious reason to do so (except possibly boredom in Experiment 2), so far we cannot exclude that bias differences in Experiments 1 and 2 were at least partly due to voluntary, decisional factors rather than involuntary, perceptual ones.

Voluntary control of the direction of motion in the motion quartet was noticed by Ramachandran and Anstis (1983b, 1985), who remarked that their continuously cycling quartet display was controllable voluntarily for temporal frequencies below roughly 3 Hz. In our single-shot (two-frame) version of the quartet, we have noticed that the critical factor is, in fact, the duration of the first frame. We have found that when this duration is above roughly 200–400 ms, some observers can reliably voluntarily control the perceived direction of motion. When the duration of the first frame is below about 200 ms, we have found no observer who can control the direction of motion. These observations are at least in qualitative agreement with those of Ramachandran and Anstis (1983b, 1985), who remarked that their continuous cycling quartet display was controllable voluntarily for temporal frequencies below roughly 3 Hz. In our single-shot (two-frame) version of the quartet, we have noticed that the critical factor is, in fact, the duration of the first frame. We have found that when this duration is above roughly 200–400 ms, some observers can reliably voluntarily control the perceived direction of motion. When the duration of the first frame is below about 200 ms, we have found no observer who can control the direction of motion. These observations are at least in qualitative agreement with those of Ramachandran and Anstis (1983b, 1985).

The goal of the present experiment was to repeat the time series measurements of idiosyncratic biases of Experiment 2, but in conditions in which voluntary control would have been impossible. In practice, I searched for a duration threshold for the first frame below which subjects could not voluntarily control the perceived direction of motion. To measure this threshold, I performed an auxiliary experiment, de-
scribed the Supplementary Materials. In this experiment subjects were instructed, for first-frame durations from 200 to 1,000 ms, to try to force themselves to perceive either vertical, horizontal, CW, or CCW motion—but were nevertheless told to truthfully report which of the usual two motions they perceived on any given trial. Of course subjects could have simply reported having seen what they were asked to force themselves to see. However, this would have only exaggerated the measured effectiveness of voluntary control, or in other words, decreased the temporal threshold of voluntary control, which is not a problem for the present purpose. It was found that the degree of voluntary control increased significantly as a function of first-frame duration. There was no measurable control at 200 ms (see Supplementary Figure S4). For the reason stated above, this represents a lower bound for the control threshold. The reader is referred to the Supplementary Materials for further details.

Because the durations of the first frame in Experiments 1 and 2 were 500 and 400 ms, respectively, it cannot be excluded that subjects had some voluntary control of the perceived motion direction. I therefore performed another experiment, similar to Experiment 2, but with the duration of the first frame set to 100 ms, well below the lower threshold for voluntary control. In all other respects, the conditions were identical to those of Experiment 2. Six naive subjects, none of whom had participated in any other experiment in this study, took part in this experiment (ages 21–31, median 22 years, three women).

Results

The time series for all subjects are shown in Supplementary Figure S3. The bias patterns are perhaps a bit noisier than in Experiment 2 (Supplementary Figure S3), but the general features found in time series of Experiment 2 are present. The memory effect is still found: Patterns on successive measurements tend to be alike. Quantitatively, calculating the number of different responses in successive samples, in 69% of the cases, responses in successive blocks differed by significantly fewer trials than predicted by the null, no-memory hypothesis.

Significantly, there were cases of bias change over time. Namely, one subject with a vertical bias pattern several times either lost a clear bias (second subject in Supplementary Figure S3), or developed a CCW bias, each time for several blocks and each time eventually regaining vertical bias (third subject in Supplementary Figure S3). Another subject, part of whose time series is shown in Figure 6, who initially had a very strong axial bias, with an approximately vertical axis, gradually underwent a change of bias to a very strong CCW bias.

It is worth noting two things about the time series of bias patterns in Figure 6. First, during the axial phase the preferred axis is not exactly vertical, but slightly CCW of vertical. What is interesting is that this deviation from vertical seems to be systematic: It actually holds for 17 blocks (over 3 hr) before the axial bias disappears (as can be seen at the bottom of Supplementary Figure S3). Thus, this CCW deviation is most likely \( p = 2^{-(17-1)} \approx 1.5 \times 10^{-5} \) part of the actual internal state of the visual system, rather than noise. Second, the change from axial to rotational bias is gradual (over about 30 min) and monotonic, an orderly path in parameter space (Figure 4).

Discussion

This experiment was a replication of Experiment 2, but in conditions where no voluntary control of the perceived direction of motion should be possible. The requisite conditions—a first frame with duration 100 ms—were determined through an auxiliary experiment, described in the Supplementary Materials. The results replicated the essential features of Experiment 2, namely idiosyncratic bias conditions that are usually stable over time, but that occasionally undergo changes that are sometimes large. This shows that these effects are not due, or at least not solely due, to conscious, volitional effects, but are truly perceptual in nature.

General discussion

The results of this study show that complex structures underlie the perception of the motion quartet. When a two-frame quartet is shown at orientations that vary from trial to trial, responses as a function of orientation are organized around two distinctive types of pattern in most subjects. One of the
patterns consists of four CW/CCW lobes of about 90°, which corresponds to a bias for translation (in either direction) along a preferred axis, whose orientation depends on the orientation of the pattern. In subjects who have this pattern the distribution of preferred axes has peaks in the cardinal directions, with the vertical direction more common than the horizontal. The other pattern is a prevalence of uniformly CW or CCW responses, a bias for rotation in one of these directions. In subjects with rotational bias, CW preference is more common than CCW.

The two bias patterns are actually extremes in a continuum. Although many subjects have the extreme patterns, others have combinations of axial and rotational biases. Thus, individual biases patterns are actually points in a multidimensional space, with several subjects having biases that have significant contributions of both bias types.

A second experiment, using repeated measurements over 9 hr, showed that biases are usually stable over time, but also sometimes undergo significant change. I have found cases where an axial bias changes preferred axis, and a rotational bias changes preferred direction. In at least one case, an axial bias became almost completely rotational.

Finally, two control experiments showed that the variety of bias patterns found across subjects, and the patterns’ changes over time within subjects, are not due to voluntary or decisional processes. The first control experiment revealed that voluntary control of the motion quartet is possible only when the duration of the first frame is above a threshold of 200–400 ms. Since the duration of the first frame in Experiments 1 and 2 was slightly above this threshold, I performed another control experiment, similar to the time series measurement of Experiment 2, but with the duration of the first frame set to 100 ms, well below the threshold for voluntary control. Results similar to those of Experiment 2 show that between- and within-subject variability cannot be due to voluntary or decisional effects.

Certain findings presented here agree with earlier observations on the motion quartet, but place these earlier observations in a broader context. It has long been known that, when the quartet is in its usual upright orientation, perception of vertical motion is more common than that of horizontal motion (Generelli, 1948), with this difference attributed to the difference between intra- and interhemispheric connections (Genç, Bergmann, Singer, & Kohler, 2011; Shimono, Mano, & Niki, 2012). This discrete fact can now be seen as part of a larger picture: Observers who have an axial bias have a preferred axis with a population distribution estimated by Figure 4b. This distribution has a larger peak centered on vertical than horizontal orientations, but, interestingly, oblique preferred axes also occur.

However, it is important to note that the temporal changes in preferred axis observed in Experiment 2, as well as changes from axial to rotational bias, invalidate a strong version of the hypothesis that brain anatomy alone determines individual differences in bias (Genç et al., 2011; Shimono et al., 2012). However, brain anatomy may play a role in the vertical-horizontal asymmetry in the population distribution of preferred axes.

Another observation that enlarges previously known facts is that, when the quartet is shown over several cycles, some observers perceive a rotation in a consistent direction (CW or CCW), rather that back-and-forth translation along a particular axis (Ramachandran & Anstis, 1983b). Once again, this can be seen in the context of the current findings, as likely to arise in subjects having rotational (rather than axial) bias. Interestingly, the consistent rotation percept is reported as disappearing above a certain temporal frequency (Ramachandran & Anstis, 1983b), which may indicate that individual bias patterns depend on temporal frequency—a possibility not tested here.

Adding to our previous results on individual biases (Wexler et al., 2015), the current study demonstrates an additional family of stimuli governed by robust-but-evolving biases with large individual differences. However, the biases governing the motion quartet are more complex than the structure- and transparency-from-motion biases reported by Wexler et al. (2015), which were shown to exist in a two-dimensional parameter space (i.e., a large part of between- and within-subject variability can be described by two parameters). Here I have shown that the motion quartet biases exist in a 3-D space, but their actual dimensionality may be even higher. For example, if it is found the bias patterns depend idiosyncratically on temporal frequency (as is likely—see previous paragraph), position in the visual field, stimulus size, or other stimulus parameters, the dimensionality of the biases will be shown to be higher than three. Other studies of idiosyncratic biases may have also demonstrated high-dimensional biases (Carter & Cavanagh, 2007; Afraz et al., 2010; Houltsby et al., 2013; Kosovicheva & Whitney, 2017), but precise estimation of dimension is difficult. The question is important because the biases are actually brain states that can be measured psychophysically, that govern perception of families of stimuli, and that evolve in response to stimuli or autonomously. Understanding the spaces in which they evolve and the laws governing this evolution is crucial to complete description of perception.

Keywords: individual differences, motion perception, apparent motion, motion quartet, internal states
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Corresponding author: Mark Wexler.
Email: mark.wexler@parisdescartes.fr.
Address: Laboratoire Psychologie de la Perception, CNRS and Université Paris Descartes, Paris, France.

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