Sequence-dependent sensitivity explains the accuracy of decisions when cues are separated with a gap

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Abstract
Most decisions require information gathering from a stimulus presented with different gaps. However, the neural mechanism underlying this integration is ambiguous. Recently, it has been claimed that humans can optimally integrate the information of two discrete pulses independent of the temporal gap between them. Interestingly, subjects’ performance on such a task, with two discrete pulses, is superior to what a perfect accumulator can predict. Although numerous neuronal and descriptive models have been proposed to explain the mechanism of perceptual decision-making, none can explain human behavior on this two-pulse task. In order to investigate the mechanism of decision-making on the noted tasks, a set of modified drift-diffusion models based on different hypotheses were used. Model comparisons clarified that, in a sequence of information arriving at different times, the accumulated information of earlier evidence affects the process of information accumulation of later evidence. It was shown that the rate of information extraction depends on whether the pulse is the first or the second one. Moreover, our findings suggest that a drift diffusion model with a dynamic drift rate can also explain the stronger effect of the second pulse on decisions as shown by Kiani et al. (Journal of Neuroscience, 33 (42), 16483-16489, 2013).

Keywords Perceptual decision-making · Discrete pulses · Random-dot motion · Accuracy · Accumulation to bound model

Introduction
Struggling with an extensive range of decisions with various degrees of complexity and type is an inseparable part of life, from voting for a competent president by integrating different types of discrete information over a period of time to simple immediate decisions such as picking up a pen. Recent studies on perceptual decisions in primates have investigated the neurobiological mechanisms underlying simple decisions such as a binary choice between two likely stimuli (Britten et al., 1996; Heekeren et al., 2004; Horwitz et al., 2004; Kiani et al., 2008; Latimer et al., 2015; Newsome et al., 1989; Roitman & Shadlen, 2002; Schall, 2003). Neural responses in the lateral intra-parietal area (LIP) gradually increase to reach a specific level of firing rate when a monkey wants to make a decision about the motion direction in a stimulus with its eye movement (Cook & Maunsell, 2002; Gold & Shadlen, 2001, 2007; Hanks et al., 2014; Heitz & Schall, 2012; Huk & shadlen, 2005; Kiani & Shadlen, 2009; Kiani et al., 2008; Philiaestides & Sajda, 2005). Therefore, it has been hypothesized that a variety of perceptual and mnemonic decisions in primates can be explained by a “bounded evidence accumulation” framework (Kira et al., 2015; Philiaestides & Sajda, 2005; Ratcliff & McKoon, 2008; Ratcliff & Tuerlinekx,
A well-known class of models, drift diffusion model (DDM) and its extensions, has been proposed to explain this process based on the integration of noisy evidence in favor of different alternatives to reach a decision bound (Gold & Shadlen, 2007; Link, 1992; Ratcliff & McKoon, 2008). Different extensions of the DDM can interpret the reaction time, accuracy, confidence, speed–accuracy trade-off, and some other aspects of decision-making (Hanks et al., 2014; Heitz & Schall, 2012; Kiani et al., 2014; Smith & Vickers, 1988; Usher & McClelland, 2001; Wang, 2002).

Indeed, the vast majority of decisions involve a complicated procedure that is the accumulation of evidence from separated cues over time. Along the way, the brain needs to evaluate and integrate various pieces of information arriving at different times (Kiani et al., 2013; Kira et al., 2015; Yang & Shadlen, 2007). Recently, it has been demonstrated that the brain can optimally accumulate evidence from a set of noisy observations and attach more weight to the more decisive evidence (Kira et al., 2015; Yang & Shadlen, 2007). It has also been shown that humans can gather information from two discrete cues separated by a gap of time, independent of the length of gaps. More importantly, Kiani et al. indicated that the most recent evidence shows far more influence on decisions (Kiani et al., 2013). However, it still remains unclear which mechanism can explain the process of decision-making in the presence of two pulses separated by time gaps.

The aim of this study was to investigate the mechanism of decision-making of a two-pulse experiment suggested by Kiani et al. (2013). A behavioral experiment including trials with one (single-pulse) and two (double-pulse) pulses of random-dot motion, separated with different interim periods (similar to Kiani et al., 2013) was implemented. Behavioral results can replicate what Kiani et al. (2013) had previously clarified. First, subjects’ accuracy improved in double-pulse trials more than what had been expected from single-pulse trials, and accuracy was independent of the interim period between pulses. Moreover, the second pulse had a larger effect on subjects’ performance.

Using a well-known decision-making model, DDM, the plausible mechanism underlying this behavior was explored (Gold & Shadlen, 2007; Mazurek et al., 2003; Shadlen et al., 2006). The DDM contains three basic parameters consisting of a starting point, drift rate, and width of the interval between decision thresholds (Bogacz et al., 2006; Ratcliff, 1978; Ratcliff & Rouder, 1998; Ratcliff & Tuerlinckx, 2002; Ratcliff et al., 2016). According to the parameters of the model, three different hypotheses were examined. Results showed that the process of information accumulation differs across double-pulse and single-pulse trials. The accumulated information in the first pulse plays the role of a starting point for the accumulation process in the second pulse. More importantly, the rate of accumulation, not the bound, depends on the sequence of pulse presentation. Interestingly, the proposed model can well explain the sequence-dependent behavior in trials with unequal pulse strengths, even when the parameters are fitted using different sets of trials with equal pulse strength.

Material and methods

Subjects

Four human subjects, two females and two males aged 24–31 years, were tested in this experiment. Two subjects (one female and one male) were naïve to the purpose of the experiment, and all subjects had normal or corrected-to-normal vision. Subjects signed an informed consent form before participating in the experiment. The Ethics Committee of Iran University of Medical Sciences approved the experimental protocol.

Before the main experiment, subjects were extensively trained (Supplemental Material Methods and Fig. S1) to achieve a high level of accuracy and a predefined reaction time (Fig. S2 and Fig. S3). In the main experiment, each subject completed 11–15 blocks containing 300 trials.

Stimulus

Subjects were asked to identify the direction of motion stimuli consisting of three independent sets of dots displayed on consecutive video frames (dot density: 16.7 dots/deg²/s). The size of dots was 2×2 pixels (side: 0.088°) (Roitman & Shadlen, 2002; Shadlen & Newsome, 2001). Each set of white dots was shown on a black background for one frame (13.3 ms) and was updated three frames later (after 40 ms). Therefore, the positions of the dots in frames one, two, and three were respectively updated in frames four, five, and six, and so on. To update, each dot was either moved coherently along the motion direction (right or left) at the velocity of 6 deg/s or relocated randomly. Task difficulty was controlled through the coherence of motion, which is the probability that each dot will be moved toward the motion direction.

Procedure

All experiments were conducted in a dimly lit room. Subjects were seated in front of a 19-in. LG CRT display monitor (screen resolution: 800 × 600, refresh rate: 75 Hz) at a distance of 57 cm. An adjustable chin rest and a forehead bar were used to ensure the stability of subjects’ heads. MATLAB’s Psychtoolbox (Brainard & Vision, 1997; Pelli, 1997) was employed to display the stimuli. The procedure of trials is described below.
1) **Fixation:** A red fixation point with a 0.3° diameter at the center of the screen was on during the trial to maintain subjects’ attention. Each trial started with the appearance of the red fixation point. Then, subjects were asked to gaze at the fixation point and maintain their gaze within the trial. Upon pressing the space key, the second step would start.

2) **Target on:** Two red targets with a 0.5° diameter were presented on the right and left sides of the fixation point at a distance of 10° to cue subjects for possible direction of motion. After a random delay (200–500 ms, truncated exponential), the motion stimuli were presented.

3) **Motion:** The motion stimulus was displayed within a 5° circular aperture at the center of the monitor. As mentioned before, two types of stimulus presentation, single-pulse and double-pulse trials, were utilized as below:

- **Single-pulse:** Motion coherence was randomly selected from these values: 0%, 3.2%, 6.4%, 12.8%, 25.6%, and 51.2%. After 120 ms of stimulus presentation, the next step was followed (Fig. 1A).

- **Double-pulse:** In each pulse, motion coherence was randomly selected from these values: 3.2%, 6.4%, and 12.8%. However, both pulses had the same net direction of motion, and subjects were informed. Two pulses were separated by a gap from these values: 0, 120, 360, or 1,080 ms. In total, there were 9 × 4 types of trials. Each pulse was presented for 120 ms and, after the second pulse, the next step was followed (Fig. 1B).

4) **Delay:** As in the second step, two targets and a fixation point were shown for a random delay (400–1,000 ms, truncated exponential).

5) **Go:** After the random delay, the fixation point extinguished and subjects were instructed to report their decision within 1 s after the Go cue by pressing two specific keys for right and left choices. After that, an auditory feedback with two frequencies (low frequency for correct and high frequency for error) beeped for correct and error responses. More importantly, on trials with 0% motion coherence, the frequency of auditory feedback was randomly selected.

Subjects completed one or two blocks a day, each block including 300 trials. In total, 42 (6 + 36) types of trials were randomly interleaved in each block. Subjects were asked to perform each trial as accurately as possible. The probability correct of each block was displayed at the end of the block as a feedback to the subjects.

**Data analysis**

To estimate the effect of parameters on the binary choices, different logistic regression models were applied. In the following regression models, Logit[P] indicates log (P/1-P) and β denotes fitted coefficients.

To approximate the performance of single-pulse trials, the first regression model was fitted as below:

$$\text{Logit}[P_{\text{correct}}] = \beta_0 + \beta_1 C$$  \hspace{1cm} (1)

where C is the coherency level of single-pulse trials.

To evaluate whether the probability correct is affected by the interim period or not, the following regression analysis was utilized:

$$\text{Logit}[P_{\text{correct}}] = \beta_0 + \beta_1 C_1 + \beta_2 C_2 + \beta_3 T + \beta_4 C_1 T$$  \hspace{1cm} (2)

where $C_1$ and $C_2$ are the motion coherence of the first and second pulse, respectively, and $T$ is the interim period. The null hypothesis is the lack of relationship between the probability correct and the interim period ($H_0: \beta_{3.4} = 0$).

In the next step, a regression model was fitted to compare the expected accuracy of a perfect integrator to the observed performance on discrete trials:

$$\text{Logit}[P_{\text{correct}}] = \text{Logit}[P_e] + \beta$$  \hspace{1cm} (3)

where $P_e$ is the expected probability correct resulting from a perfect accumulator (Eq. 4). A positive value for $\beta$ indicates that the observed accuracy is higher than the expected one.

$$P_e = 1 - \Phi(0, e_1 + e_2, \sqrt{2})$$  \hspace{1cm} (4)

where $\Phi$ is the normal cumulative distribution function, and is calculated as:

$$\Phi(x, \mu, \sigma) = \int_{-\infty}^{x} N(v, \mu, \sigma)dv$$  \hspace{1cm} (5)

where $N(v, \mu, \sigma)$ is the normal probability density function with mean ($\mu$) and SD ($\sigma$). $e_1$ and $e_2$ are the pieces of evidence of two pulses. The distribution of the evidence can be calculated from the probability correct of single-pulse trials:

$$e_i = \Phi^{-1}(P_i, 0, 1), \hspace{0.5cm} i = 1, 2$$  \hspace{1cm} (6)

where $P_i$ is the probability correct of single-pulse trials based on Eq. 1, and $\Phi^{-1}$ is the inverse of the normal cumulative distribution function in Eq. 5.

The following regression model was used to test the effect of pulse sequence on the probability correct:

$$\text{Logit}[P_{\text{correct}}] = \beta_0 + \beta_1 |C_1 + C_2| + \beta_2 |C_1 - C_2|$$  \hspace{1cm} (7)

where $C_1$ and $C_2$ are the motion coherence of the first and second pulse, respectively, and $\beta_2$ represents the change in probability correct from the weak-strong pulse ($C_1 < C_2$) to the strong-weak one ($C_1 > C_2$). The null
hypothesis is that the probability correct does not depend on the pulse sequence (H₀: β₂ = 0).

Modeling

To explain the plausible mechanism for decisions with two discrete pulses, the DDM (Eq. 8) (Ratcliff, 1978; Ratcliff & McKoon, 2008) implemented by Voss et al. was applied in a computationally efficient and accessible program called fast-dm (Lerche & Voss, 2017; Voss & Voss, 2007; Voss et al., 2004, 2013).

\[ dv = V dt + \xi \sqrt{dt} \text{ where } v(0) = z \]  

(8)

This model is described via an accumulation-to-bound mechanism that starts with gathering noisy sensory evidence (explained by drift rate (v)) from a starting point (z) and terminates upon crossing the pre-specified response criterion (a). These three basic parameters existing in early versions of the diffusion model are compatible with both neurophysiological and behavioral data. The refined version of DDM was designated by seven parameters grouped into the following three classes: (I) the basic parameters of the decision process (decision threshold a, mean starting point z, and mean drift rate v), (II) the non-decision process parameter (non-decision time t_{ND}), and (III) the changeability across-trial parameters (changeability in stimulus quality η, changeability in starting point sz, and starting point in non-decision time s_{ND}) (Ratcliff, 1978; Ratcliff & McKoon, 2008; Ratcliff & Tuerlinckx, 2002).

Fast-dm estimated the DDM’s parameters using the partial differential equation (PDE) method (Voss & Voss, 2007), which had swift calculations to compute the cumulative distribution function (CDF) and the Kolmogorov-Smirnov statistic (Voss et al., 2004). The diffusion model can describe the perceptual decision process in the brain and is actually a well-entrenched model in this field (Ratcliff, 1978; Ratcliff & McKoon, 2008).

In this study and based on our hypothesis, three modified models were employed and compared using the Bayesian information criterion (BIC) (Kass & Wasserman, 1995; Liddle, 2007; Smith & Spiegelhalter, 1980).

Results

Four human subjects were asked to make a decision about the net direction of motion in trials in which a 120-ms motion pulse was displayed once or twice. Single-pulse and double-pulse trials were randomly presented. In double-pulse trials, both pulses had the same direction, and subjects were informed about this consistency. The net direction, motion strength, and interim period were random from trial to trial. Here, we first demonstrated that our data can reproduce the results reported by Kiani et al. (2013), and then fitted different DDMs to shed light on the mechanism of decision-making under discrete conditions.

Since 86% of trials were double-pulses, it may be hypothesized that the first pulse would be ignored and the subjects made their decisions merely based on the second pulse. However, the psychometric function of the single-pulse trials (Fig. 2) showed that the subjects accurately used the information of the first pulse. As expected from previous studies, the probability of being correct increased as a function of motion strength (Kiani et al., 2013), supporting the claim that subjects did not ignore the first pulse.

Achieving the same accuracy levels independent of the interim period

Consistent with previous results, accuracy was invariant in the interim period between pulses in double-pulse trials for both trials with equal pulse strength, or so-called same discrete trials (Fig. 3A, Eq. 2, β₃=0.31±0.02, p=0.19; β₄=-0.05±0.03 p=0.15, for each subject; see Table S1) and trials with unequal pulse strength, or so-called different discrete trials (Fig. 3B, Eq. 2, β₃=-0.11±0.29, p=0.70; β₄=0.01±0.02 p=0.57; β₅=-0.002±0.02, p=0.92, for each subject; see Table S1) (see equivalence test result in Table 1).
Accordingly, this finding demonstrated that the information of the first pulse was not lost but maintained throughout the interim period. Furthermore, the probability correct in weak-strong double-pulse trials was enhanced to above the expected performance of a perfect accumulator (Eq. 3, $\beta = 0.10 \pm 0.04$, $p = 0.018$, for each subject; see Table S2), which, at least, provides evidence for the use of the information of both pulses. This better-than-anticipated performance also shows that the process of decision-making in double-pulse trials might be different from simply doubling the information of one pulse.

What mechanism can explain the sequence-dependent performance of subjects in double-pulse trials?

Our data also replicated what Kiani et al. (2013) had demonstrated about the sequence of pulses. As stated above, subjects performed better than expected in weak-strong sequence of double-pulse trials, where the second pulse is stronger than the first pulse and, as illustrated in Fig. 4, accuracy increased in double-pulse trials when the motion strength of the second pulse was stronger than that of the first one (Eq. 7, $\beta_2 = 0.03 \pm 0.005$, $p = 8.60 \times 10^{-8}$, for each subject; see Fig. S4).

In this part of the Results section, we propose a plausible mechanism for the observed sequence-dependent performance that caused the over-expected accuracy in double-pulse trials.

Assuming the accumulation to bound model as a true model of the decision process, we used the DDM to investigate three plausible hypotheses for the mechanism of the decision process in double-pulse trials: (1) Decision-making in double-pulse trials involves the same mechanism as single-pulse trials such that information accumulation in the first pulse will be continued in the second pulse until crossing a bound. This means that the rate of accumulation (drift rate) in the diffusion model depends on the summation of coherency of two pulses. (2) The first pulse affects the rate of information accumulation in the second pulse. (3) The first pulse affects the decision threshold or the bound of information accumulation in the second pulse.

In the following section, the results of examining these three hypotheses are presented.

Model fitting

We used the DDM to test our hypotheses and examine the mechanism of decision-making in double-pulse trials. In so doing, based on our alternative hypotheses, we ran and compared three variations of the DDM to explore how and which parameters were influenced by discretizing information.

In the first model, we assumed that the information accumulation is similar in both single-pulse and double-pulse trials. This means that the interim period is not considerable, and the two pulses are considered as one pulse. To examine this assumption, we made the drift-rate ($v$) a function of the sum of...
the coherency of two pulses in double-pulse trials and called it model$_{sum}$. The second and third models were fitted to explore whether the process of decision-making in the second pulse is influenced by the accumulated information of the first pulse or not. To do so, in the second hypothesis, we assumed that the accumulated information of the first pulse can affect the rate of information accumulation in the second pulse. In order to check this hypothesis, we fitted model$_{starting point}$. This model is based on the assumption that in the second pulse the starting point of the accumulation process is equal to the accumulated evidence from the first pulse, which itself depends on the stimulus strength in that pulse. In other words, the starting point of the accumulation process in the second pulse depends on the drift rate in the first pulse. In model$_{starting point}$, the starting point ($z$) and the drift-rate ($v$) of the DDM are a function in the first pulse and expect a stronger one in the second pulse, respectively. As a result, this model can potentially assign different drift rates for the first and the second pulses.

The third DDM model, model$_{bound}$, was based on the fact that the decision bound determines the speed and accuracy trade-off in which a higher decision bound results in higher accuracy with the cost of higher reaction time and a lower decision bound causes faster but less accurate decisions. Subjects may increase the decision bound when they receive weak information in the first pulse and expect a stronger one in the second pulse.

**Table 1** Equivalence test results

| Gap condition (ms) | 0 and 120 | 0 and 360 | 0 and 1,080 | 120 and 360 | 120 and 1,080 | 360 and 1,080 |
|--------------------|-----------|-----------|-------------|-------------|---------------|---------------|
| Motion strength    |           |           |             |             |               |               |
| 3.2–3.2%          | B= ±0.1029| B= ±0.04  | B= ±0.448   | B= ±0.1176  | B= ±0.1377    | B= ±0.0731    |
|                   | CI=[0.07;0.103] | CI=[-0.027;0.036] | CI=[-0.043;0.012] | CI=[0.048;0.115] | CI=[-0.133;0.095] | CI=[-0.065;0.001] |
|                   | P= 0.048 | P= 0.032  | P= 0.037    | P= 0.039    | P= 0.021      | P= 0.021      |
| 6.4–6.4%          | B= ±0.1016| B= ±0.1068| B= ±0.1036  | B= ±0.0201  | B= ±0.021     | B= ±0.0309    |
|                   | CI=[-0.099;0.073] | CI=[-0.106;0.081] | CI=[-0.098;0.064] | CI=[-0.019;0.002] | CI=[-0.011;0.02] | CI=[-0.003;0.028] |
|                   | P= 0.022 | P= 0.044  | P= 0.016    | P= 0.03     | P= 0.044      | P= 0.027      |
| 12.8–12.8%        | B= ±0.0217| B= ±0.0232| B= ±0.0597  | B= ±0.0145  | B= ±0.0472    | B= ±0.0461    |
|                   | CI=[-0.005;0.021] | CI=[-0.006;0.022] | CI=[0.03;0.056] | CI=[-0.011;0.012] | CI=[0.025;0.045] | CI=[0.023;0.046] |
|                   | P= 0.042 | P= 0.041  | P= 0.016    | P= 0.021    | P= 0.04        | P= 0.044      |
| 3.2–6.4% and      | B= ±0.0151| B= ±0.0252| B= ±0.0273  | B= ±0.0208  | B= ±0.0271    | B= ±0.0359    |
| 6.4–3.2%          | CI=[-0.012;0.01] | CI=[-0.022;0.003] | CI=[0.005;0.025] | CI=[-0.021;0.002] | CI=[0.006;0.026] | CI=[0.02;0.036] |
|                   | P= 0.017 | P= 0.012  | P= 0.026    | P= 0.049    | P= 0.037      | P= 0.048      |
| 3.2–12.8% and     | B= ±0.0187| B= ±0.0327| B= ±0.0111  | B= ±0.0225  | B= ±0.0175    | B= ±0.0345    |
| 12.8–3.2%         | CI=[0.002;0.016] | CI=[0.016;0.032] | CI=[-0.009;0.007] | CI=[0.008;0.021] | CI=[-0.017;0.004] | CI=[-0.032;0.017] |
|                   | P= 0.012 | P= 0.033  | P= 0.022    | P= 0.015    | P= 0.034      | P= 0.018      |
| 6.4–12.8% and     | B= ±0.0138| B= ±0.0501| B= ±0.0157  | B= ±0.015   | B= ±0.0234    | B= ±0.0591    |
| 12.8–6.4%         | CI=[-0.011;0.004] | CI=[-0.049;0.037] | CI=[-0.003;0.014] | CI=[-0.047;0.031] | CI=[-0.001;0.019] | CI=[0.039;0.057] |
|                   | P= 0.015 | P= 0.026  | P= 0.024    | P= 1        | P= 0.011      | P= 0.027      |

B, CI, and P represent equivalence bounds, 90% confidence interval of TOST test, and p-value, respectively.
pulse, and vice versa. Therefore, the third hypothesis is that the first pulse can influence the decision bound of the information accumulation in the second pulse (similar to Olianezhad et al., 2019). In order to test this hypothesis, we made the bound (a) of the DDM dependent on the motion strength of the first pulse and the drift-rate (v) on the motion strength of the second pulse. The fitted parameters of each model are listed in Tables 2, 3 and 4 (standard errors (SEs) were calculated with 100 iterations of bootstrap).

Here, we utilized the BIC (Kass & Wasserman, 1995; Liddle, 2007; Smith & Spiegelhalter, 1980) to compare the models (Table 5). As shown in this table, the BIC of model starting point is smaller than that of the other models, indicating that this model can better explain the observed behavior of subjects in double-pulse trials (see quantile probability function of the best model in Fig. 5).

In the next step, we also investigated the differences among the dependent parameters for the best model (model starting point). As mentioned in Table 3, the winner model produced three starting points and drift rates. Therefore, the significance of the differences among them was tested by the nonparametric bootstrap method (using a t-test with 100 iterations of bootstrap) (Efron, 1992; Efron & Tibshirani, 1994). These differences were highly significant (p_{z,3.2,2.6.4} < 0.05, p_{z,2.2,2.12.8} < 0.05, and p_{v,6.4,2.12.8} < 0.05), indicating that the winner model wins the competition not due to small differences between its unique free parameters. It also shows that the accumulated evidence after the first pulse is significantly dependent on the pulse strength.

### Analyzing the parameters of the best model

Table 3 illustrates the parameters of the best model. Considering that each pulse lasts 120 ms, one can calculate the drift rate of the first pulse using Eq. 9:

\[
\mu = \frac{(z-0.5)}{0.120}
\]  

where \( \mu \) is the drift rate and \( z \) is the starting point for the second pulse or the accumulated information at the end of the first pulse. Moreover, 0.5 is the starting point of the first pulse. As a result, the drift rate of each stimulus at each pulse is presented in Table 6.

### Table 3 Fitted parameters (mean±SE) of the second DDM (model starting point)

| Parameter | Value         |
|-----------|--------------|
| \( z_{3,2} \) | 0.5206±0.0300 |
| \( z_{6,4} \) | 0.5425±0.0294 |
| \( z_{12,8} \) | 0.5772±0.0290 |
| \( a_{3,2} \) | 1.0909±0.0238 |
| \( v_{3,2} \) | 0.7684±0.0238 |
| \( v_{6,4} \) | 1.0882±0.1430 |
| \( v_{12,8} \) | 1.6308±0.1674 |
| \( t_{ND} \) | 0.7181±0.0116 |
| \( s_{ND} \) | 0.3407±0.0150 |

### Table 4 Fitted parameters (mean±SE) of the third DDM (model bound)

| Parameter | Value         |
|-----------|--------------|
| \( z \) | 0.5379±0.0222 |
| \( a_{3,2} \) | 1.0829±0.0260 |
| \( a_{6,4} \) | 0.7773±0.0216 |
| \( a_{12,8} \) | 1.1160±0.0238 |
| \( v_{3,2} \) | 0.8011±0.0116 |
| \( v_{6,4} \) | 1.1178±0.1270 |
| \( v_{12,8} \) | 1.6638±0.1452 |
| \( t_{ND} \) | 0.7131±0.0082 |
| \( s_{ND} \) | 0.3419±0.0146 |

### Table 5 Model performance comparison via BIC and R^2 metrics (mean±SD across subjects)

| Model | Total parameters | R^2 ± SD | BIC ± SD |
|-------|------------------|---------|---------|
| Model sum | 15 | 0.8181±0.0550 | -29.7407±2.8348 |
| Model starting point | 14 | 0.8660±0.0573 | -35.1189±4.2153 |
| Model bound | 14 | 0.7470±0.0884 | -29.0620±3.3791 |

**Fig. 5** Quantile-probability function for average data in the double-pulse trials. The percentage of correct responses is on the right side of the panel. In the left side we have the distributions for the error responses. The continuous lines show the fit of the Model starting point and the xs are the behavioral data. Response times are grouped to produce the five conditions.
Table 6  Drift rate of the first and second pulse

|                  | First pulse | Second pulse |
|------------------|-------------|--------------|
| $v_{3.2}$        | 0.172       | 0.6915       |
| $v_{6.4}$        | 0.3542      | 1.0125       |
| $v_{12.8}$       | 0.6433      | 1.5319       |

**Model starting point can explain the sequence-dependent behavior of subjects**

Figure 6A demonstrates that model starting point can mimic the same trend as behavioral data (Fig. 4). As reported by Kiani et al. (2013) and replicated here, subjects’ accuracy in double-pulse trials is sequence-dependent. They rely more on the second pulse than on the first pulse to make their decisions. Here, we also investigate whether the best model suggested by BIC and fitted on double-pulse trials with equal pulse strength can replicate the sequence-dependent behavior of subjects in double-pulse trials with unequal pulse strength (Fig. 6B). It is noteworthy that, in Fig. 6B, the model is totally blind to the subject data in the double-pulse trials with unequal pulses strength.

**Discussion**

The decision-making process tends to integrate discrete pieces of information in favor of various options (Kiani et al., 2013; Kira et al., 2015; Yang & Shadlen, 2007). Kiani et al. (2013) showed that, in a fixed-duration direction discrimination task, the integration of direction cues is independent of the temporal gap between them. Moreover, they revealed that participants’ accuracy is more affected by the last pieces of information than the primary ones and, more importantly, their accuracy in double-pulse trials is more than what is expected from their performance in single-pulse trials. Here, using a similar psychophysics task, in addition to the confirmation of previous results, we suggested a mechanism to explain participants’ behavior on such a task. Based on the DDM, we demonstrated that among a set of modified models, a sequence-dependent drift rate model can provide a better fit for the behavioral results. The model also accounts for both the sequence dependence and the over-expected accuracy of participants. It is notable that Kiani et al. (2013) concluded that the accuracy of weak-strong trials where the first pulse is weaker than the second one and strong-weak trials where the first pulse is stronger than the second one is significantly higher than that of the perfect integrator fitted to the performance of single-pulse trials. However, this over-expected accuracy is reproduced in our results only in trials with a weak-strong sequence. Our behavioral results, in contrast to Kiani et al. (2013), do not show a significant difference between the expected accuracy and the accuracy in trials with strong-weak sequence trials. As discussed by Kiani et al. (2013), “the better-than-expected performance in double pulse trials was actually explained by underperformance in single-pulse trials.” Thus, performing better in single-pulse trials diminishes the better-than-expected performance effect. The discrepancy between our and Kiani’s

![Image](image_url)
data stems from the fact that our subjects perform better than Kiani’s subjects in single-pulse trials with low motion strength (3.2 % and 6.4%), which is evident in our psychometric functions.

Based on our results, the drift rate is not always a static parameter as assumed in previous studies. Based on Table 6, the rate of information extraction in the first pulse differs from that of the second pulse. This change in sensitivity also occurs in double-pulse trials with equal pulse strength in a way that the model fitted to double-pulse trials with equal pulse strength can truly predict the sequence-dependent performance in double-pulse trials with unequal pulse strength (Fig. 6B).

The change in the sensitivity of the second pulse can be explained by two potential mechanisms: (1) The time variant sensitivity (drift rate), according to which attention increases as the trial is near the end. This time variant decision-making has previously been suggested to explain longer reaction times in error trials (Ditterich, 2006). Ditterich showed that within-trial variability of sensitivity can better explain the distribution of reaction time in a random dot task than the variability in decision threshold. (2) The accumulated evidence (decision variable state) variant sensitivity (drift rate), according to which the sensitivity of the second pulse depends on the state of the accumulated evidence at the end of the first pulse. This hypothesis stems from the idea that when the accumulated evidence in the first pulse is less than what we expect, the brain will try to extract more information in the next pulse to improve his decision. On the other hand, when the accumulated evidence from the first pulse is more than what we expect, the brain does not need to increase the attention for gaining more evidence in the second pulse. As a result, the overall sensitivity improved in the second pulse. However, since both time and accumulated evidence are highly correlated variables, distinguishing between these hypotheses needs a more complex and well-controlled design of experiments.

In summary, our results shed more light on the mechanism of decision-making where pulses of information are separated by temporal gaps. We showed that the drift rate of the accumulation to bound model is modulated not only with the strength of the stimulus but also with the temporal position of pulses. However, our results investigated the process of decision-making when stimulus is discretized only in two pulses. More investigations into the process of decision-making with stimuli containing more than two pulses are needed to better understand the mechanism of decision-making in discretized environment.

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