Optimal Random Avoidance Strategy in Prey-Predator Interactions

Masato S. Abe\textsuperscript{1,*} and Minoru Kasada\textsuperscript{2,3}

1. RIKEN Center for Advanced Intelligence Project, 1-chome Mitsui Building, 15th floor, 1-4-1 Nihonbashi, Chuo-ku, Tokyo 103-0027, Japan (Tel: +81-048-467-3627)
2. Graduate School of Life Sciences, Tohoku University 6-3, Aoba, Sendai 980-8578 Japan
3. Department of Experimental Limnology, Leibniz-Institute of Freshwater Ecology and Inland Fisheries, Alte Fischerhuette 2, 16775, Stechlin, Germany

* Author for correspondence (MSA: masatoabee@gmail.com)
Abstract

It has recently been reported that individual animals, ranging from insects to birds and mammals, exhibit a special class of random walks, known as Lévy walks, which can lead to higher search efficiency than normal random walks. However, the role of randomness or unpredictability in animal movements is not very well understood. In the present study, we used a theoretical framework to explore the advantage of Lévy walks in terms of avoidance behaviour in prey-predator interactions and analysed the conditions for maximising the prey’s survival rate. We showed that there is a trade-off relationship between the predictability of the prey’s movement and the length of time of exposure to predation risk, suggesting that it is difficult for prey to decrease both parameters in order to survive. Then, we demonstrated that the optimal degree of randomness in avoidance behaviour could change depending on the predator’s ability. In particular, Lévy walks resulted in higher survival rates than normal random walks and straight movements when the physical ability of the predators was high. This indicates that the advantage of Lévy walks may also be present in random avoidance behaviour and provides new insights into why Lévy walks can evolve in terms of randomness.

Keywords: Lévy walks, avoidance behaviour, prey-predator interactions, movement ecology
1. Introduction

To an observer, animal behaviour often seems random and unpredictable. Even when individual animals are subjected to the same experimental apparatus or external stimuli, behavioural variability is often observed [1–5]. It has been suggested that, to some extent, such variability stems from the nervous system that can initiate spontaneous actions [1,3,5–8]. However, how is variability or randomness in animal behaviour associated with evolutionary advantages? One of the candidate hypotheses is that such variability may improve a prey’s efficiency in avoiding a predator’s attacks in prey-predator interactions [6,9,10]. If the prey moves in the same manner every time, the predator can easily predict its movement and predation will be successful. Apart from top predators in a food web, most organisms are exposed to predation risk; therefore, the adaptive evolution of strategies that cannot be easily predicted must occur in many taxa. Consequently, in order to gain a full understanding of the role of variability or randomness in animal movements, it is crucial to explore the kind of movements exhibited by prey under predation risk and what the optimal avoidance strategy is, based on the cost and benefit involved [11,12].

Studies on the escape responses of prey to predators’ attacks have examined this topic from the perspectives of behavioural ecology and neurophysiology [9,10,13–15]. In particular, animal escapology has focused on how animals escape from threatening stimuli in an extremely short time-scale (e.g. 5-10 ms) [10,16–18]. Empirical evidence has shown that the variability in the direction of the prey’s movement may be an adaptive trait as a response to the predator that approaches it [2]. In contrast, another important consideration in prey-predator interactions is that when the predator is far from a prey individual, it can predict its movement by employing a kind of interception strategy and then it attacks it for a longer time-scale. For instance, a bird flying in the sky predicts the movement of an insect walking on the ground and then tries to attack it (figure 1a). Another example is an insect, such as a mosquito or a cockroach, which escapes from a human who is trying to catch it. In these situations, the predator aims for the prey target and tries to catch it (figure 1a). Although the speed with which the predator attacks is much faster than the speed with which the prey escapes, there would be a gap between an actual prey position and a predicted position before the attack. Besides, the prey would seek refuge if this would help it elude some predator attacks. In such cases, unpredictable movements such as random walks (figure 1b) would benefit the prey, although such movements would pose the risk of delayed arrival to the refuge. Therefore, it may be crucial for prey to exhibit movements with randomness rather than escape responses, such as reactive responses characterised by an extremely short time scale.

It has recently been reported that various animals ranging from insects to fishes, birds, and mammals, including humans, exhibit a special class of random walks, called Lévy walks [19–
The step length \( l \) (i.e. the distance between consecutive reorientations) of Lévy walks follows a power law distribution, \( P(l) \sim l^{-\mu} \) where \( \mu \in (1, 3] \) is a power law exponent characterising the movements. Lévy walks are characterised by rare ballistic movements among short steps (figure 1b), resulting in anomalous diffusions. Theoretical studies reported that Lévy walks have higher search efficiency than Brownian walks where the targets are distributed sparsely and patchily [26,27]. Therefore, it has been suggested that the prevalence of Lévy walks in animal movements is a consequence of natural selection that facilitates search efficiency. However, recent findings suggest that the various mechanisms that can produce Lévy walk patterns depend on several conditions [28,29]. Thus, it is imperative to explore why and when animals exhibit Lévy walks depending on various ecological conditions. Nevertheless, the impact of Lévy walks, other than search efficiency, on the fitness of individual animals remains poorly understood.

In the present study, we hypothesised that individual animals performing Lévy walks would benefit in terms of avoiding a predator’s attacks. The questions examined included what movement strategy is the most efficient in surviving in prey-predator interactions and how is the optimal behaviour affected by the ecological conditions and the predator’s cognitive and physical abilities [30]. To answer these questions, we constructed a general framework of the predator avoidance behaviour based on a random walk paradigm (figure 1a, b) and analysed the theoretical model to determine the efficiency of avoidance.

2. Methods
To explore the most efficient avoidance behaviour, we constructed a framework in which a prey individual avoided a predator that overlooked an open arena (figure 1a, b) and calculated the survival rate of the prey by using simulations and analytical solutions.

First, we considered a prey individual to be at the centre of the 2D circular field with radius \( R \) and a predator that overlooked the entire field from above (figure 1a, b). Then, the prey started moving and performed a class of random walks (i.e. Lévy walks with a specified \( \mu \)) at a constant velocity (one space unit per unit time) without stopping. The step lengths \( l \) were drawn from a truncated power law distribution which is often best fitted to empirical data [19,20,23,24] and the turning angles between consecutive steps were drawn from a uniform distribution (see electronic supplementary material for details). In our analysis, \( \mu \) ranged from 1.1 to 3.0 by 0.1. Lévy walks with \( \mu = 1.1 \) and 3.0 correspond approximately to straight-line movements and Brownian movements, respectively.

We assumed that predators were rational and had relatively high cognitive ability. The predator could perceive the present position of the prey, denoted by \( X_t = (x_t, y_t) \), and could also memorise its past positions. Then, the predator would have to predict the future position \( X_{t+\tau} = (x_{t+\tau}, y_{t+\tau}) \) of the prey after a \( \tau \) time step, based only on the information it had about the present (\( t \))
and past positions of the prey. The predicted position was denoted by $\hat{X}_{t+\tau} = (\hat{x}_{t+\tau}, \hat{y}_{t+\tau})$. While the time lag $\tau$ could be considered as a predator attack characteristic, it could also be a characteristic of the prey individual as its position after $\tau$ could be scaled by its velocity. Therefore, if the prey’s velocity was fast, $\tau$ could be relatively large. Moreover, if the predator attacked slowly after making a prediction, $\tau$ could also become large. Intuitively, the larger the $\tau$, the higher the prey’s advantage in escaping the attacks.

To model how the predator would predict the prey’s position in a simple manner, we assumed that the probability distributions of step lengths and turning angles of the prey did not depend on its spatial positions. Therefore, we could straightforwardly derive the optimal prediction rule adopted by predators by using a straight line (figure 1c). This is similar to interception strategies which have been observed in birds, bats, fishes, and insects [31–35]. The predicted position can be described as follows:

$$\left( \begin{array}{c} \hat{x}_{t+\tau} \\ \hat{y}_{t+\tau} \end{array} \right) = \frac{\tau}{\sqrt{(x_t - x_{t-1})^2 + (y_t - y_{t-1})^2}} \left( \begin{array}{c} x_t - x_{t-1} \\ y_t - y_{t-1} \end{array} \right) + \left( \begin{array}{c} x_t \\ y_t \end{array} \right)$$

(1)

where $(x_{t-1}, y_{t-1})$ is the past position of the prey used to obtain the direction of the prediction (figure 1c). When the distance between the predicted position and the actual position was less than or equal to an attack radius $r$, the predator could successfully catch the prey. If the prediction failed, the predator would predict the new position again after the waiting time ($W$) had elapsed. $W$ represents the time interval from the present attack to the next attack and is indicative of a characteristic of the predator that cannot attempt the next attack immediately. In contrast, once the prey reached the safety area (i.e. the edge of the circular field) (figure 1) without having been predicted by the predator, it would never leave it and would be regarded as having survived. To obtain the survival rate of the prey, this trial (from the beginning to the point of being caught or having survived) was independently iterated $10^6$ times for a specified parameter set in our simulation.

Although, based on our assumption, the prey could not obtain any information about when and where the predator would attack, we considered two cases: the prey was either aware (precaution case) or not aware (unwariness case) of the presence of the predator. The difference of these cases would result in a different step length at an initial condition ($t=0$). In the precaution case, the prey could choose a new step length from the probability distribution at the initial condition and then it could start moving. On the other hand, in the unwariness case, the prey was on the move with step length $l$. Therefore, the initial step length is likely to be longer than that of the precaution case. Under these assumptions, we conducted simulations to estimate the survival rate of the prey.
To obtain an analytical solution of the survival rate based on the aforementioned assumptions, we split it into two components; the prediction rate and the time length to reach the safety area. In this case, the survival rate $\phi$ of a prey can be described as follows:

$$\phi = (1 - Q)^{T/W}$$  \hspace{1cm} (2)

where $Q$ is the predictability of the prey’s movements by the predators and $T/W$ is the average number of attacks until the prey reaches the safety area. $Q$ and $T$ are functions of the parameter associated with the movements, namely $\mu$, but the explicit derived expression can be obtained only for $Q$ with $r \to 0$ (see electronic supplementary material for details). Therefore, we used the simulations for $T$.

In our framework, the most unpredictable strategy included movements with many reorientations. In contrast, the straight-line movement was the easiest to predict. However, movements with reorientations could result in a delay in reaching the safety area, while straight-like movements could lead to short-term exposure to predation (figure 1c); the Lévy walk model could be an intermediate strategy between the two.

3. Results
First, we demonstrated the relationship between the predators’ prediction rate and the prey’s time length to reach the safety area for specified movement patterns, which were obtained from simulations and analytical solutions (see figure 2 for the precaution case and figure S1 for the unwariness case). The smaller the $\mu$ exponents, the shorter the time it would take prey to reach the safety area. This roughly corresponds to the mean squared displacement, which characterises the diffusion property and increases as the exponents decrease [19,26]. As for the prediction rate, when the exponents $\mu$ were small, it was easier for the predators to predict the prey’s position. This is because Lévy walks with large $\mu$ are characterised by frequent reorientations. Note that the prediction rate of Lévy walks with very small $\mu$ was close to one. This result indicates that there is a trade-off relationship between the predictability and the time to reach the safety area for any $\tau$ and $R$, that is, the shorter the time it takes to reach the safety area, the higher the predictability and vice versa. An intuitive explanation for these results is shown in figure 1b. The trade-off relationship also implies that the prey cannot decrease both parameters simultaneously, although both the lower predictability and the shorter time length to reach the safety area can lead to high survival rates for the prey. Therefore, we expect that Lévy walks with intermediate $\mu$ will represent a better strategy for increasing the survival rate compared with straight movements and Brownian walks (figure 2).

Next, we analysed the relationship between the survival rate and the movement pattern ($\mu$) and examined the dependency of the other parameters ($\tau$, $R$, and $W$) on this relationship. Figure
3 shows the results for the precaution case, suggesting that Lévy walks with intermediate $\mu$ can lead to a higher survival rate for $W = 50$ and $100$, $R = 50$ and $100$ and a small prediction time lag $\tau$. In contrast, when $\tau$ was large, Brownian walks (i.e. Lévy walks with $\mu = 3$) resulted in higher survival rates than Lévy walks with $\mu < 3$ did in any scenario. When $\tau$ was small, the difference in the prediction rates for large and small $\mu$ was relatively small (figure 2). Therefore, the advantage of Brownian walks was negligible in terms of decreasing the prediction rate.

The results also showed that Brownian walks were favoured when the radius $R$ of the area was small ($R = 10$). This could be because of the short time length to reach the safety area (e.g. $T = 40$), even for Brownian walks, compared with the time length to reach the area for $R = 50$ and $100$ (figure 2). Therefore, for small $R$, Brownian walks resulted in few attacks with a low prediction rate. In contrast, for large $R$, Brownian walks were more likely to be attacked, owing to the longer time it took to reach the safety area.

The waiting time $W$ was also a key factor in determining the best avoidance strategy, for $R = 50$ and $100$ (figure 3) in particular. $W$ could affect the number of possible attacks because the average number of attacks was represented by the $T/W$ ratio. When $R$ and $\mu$ were large, $T$ could also be larger. In this case, Brownian walks caused more predator attacks, thereby resulting in a low survival rate in spite of the low prediction rate.

These tendencies were also observed in the unwariness case and the semi-analytical results for $r \to 0$ (figures S2, S3, and S4). Our results suggest a general tendency of Lévy walks to result in a higher survival rate for small $r$; for large $r$, the lower prediction rate strategy (i.e. $\mu = 3$) was more efficient.

4. Discussion

In the present study, we created a theoretical framework to investigate random avoidance behaviour in terms of Lévy walks, which have been observed widely in biological movements [19–25]. We constructed a framework for prey-predator interactions based on the prey’s movements and the predator’s prediction (figure 1) and calculated the predator’s prediction rate, the time length to reach the safety area and the prey’s survival rate. We found that there is a trade-off relationship between the predator’s prediction rate and the prey’s time length to reach the safety area (figure 2). We can interpret the latter as the degree of exposure to predation risk. Thus, the trade-off corresponds to the relationship between the predictability and the degree of exposure to predation risk, suggesting that prey individuals cannot decrease both of these factors, although they need to. Consequently, they must identify a point of compromise in order to achieve higher survival rates.

Then, we showed that Lévy walks could outperform Brownian walks in terms of the survival rate when the time lag $\tau$ between the prediction and the attack was short and the area was
not too small (figure 3). This implies that if the predator attacks the prey fast after predicting its
position, Lévy walks can be the most efficient avoidance strategy. The attack characteristics
associated with the time lag $\tau$ can be classified into two categories; the predator’s physical and
cognitive abilities. The short time lag (small $\tau$) is equivalent to the high physical ability that results
in fast attacks or in highly flexible changes of attack directions. Likewise, it can be suggested that
if making a prediction is a, somewhat, time-consuming task, the predator’s high cognitive ability
results in a shorter time lag; however, in our model we did not make explicit assumptions about
time regarding the predator’s predictions. Moreover, as the prey’s slow movement corresponded
to a small time lag $\tau$, prey individuals with low physical ability may experience higher survival
rates by employing Lévy walks. As the small size of the area may correspond to a situation where
obstacles such as grass where a prey can immediately hide under are present, Lévy walks can be
favoured in relatively large open areas. Thus, these results show that there is a crucial connection
between the prey’s fitness and the ecological and physical conditions that result from the prey’s
movements and the predator’s predictions.

When the time lag was large, Brownian walks represented a better escape strategy
(figure 3). If the movements were sufficiently random, they decreased the chances of the prey’s
position being predicted by the predator. Furthermore, when $W$ was small, the predator had a
higher chance of attacking; in this case, the random movements represented an advantage. In both
cases, eluding the attack was relatively more important for the prey’s survival than reaching the
refuge rapidly was. This result suggests that Brownian walks still represent an effective strategy
of escaping from a predator, which aims for and catches prey.

So far, the main advantage of Lévy walks has been the high encounter rate with targets
(e.g. food, mates and habitats), which are distributed scarcely and patchily [26,27]. However,
recent findings suggest that the origins of Lévy walks may not be unique, but may depend on
ecological and physical conditions [28,29]. Therefore, it is crucial to explore when and why
animal individuals would exhibit Lévy walks and which ecological traits, physical constraints or
environmental factors are responsible for this phenomenon. In this study, we focused on the prey-predator interactions once they encounter each other. This scenario occurs in relatively shorter
time and smaller spatial scales than the scenario in which a predator moves in search for prey
moving in the environment [36,37]. Our results suggest a novel advantage of Lévy walks in the
context of avoiding a predator’s attacks. Moreover, although the essential role of variability or
randomness in Lévy walks is not very well understood, our results indicate that the extent of
randomness plays a key role in determining the optimal avoidance strategy. Additionally, based
on our framework, Lévy walks should be expressed as spontaneous behaviour, which is generated
from the internal state of the prey, as the prey determines the reorientation and step length
independent of external factors such as environmental cues. Therefore, these results point to one
of the evolutionary origins of spontaneously produced Lévy walks [38].

A hunting strategy by predators with high cognitive abilities would not just be a reflective response to their prey’s movements. It has been shown that predators, including vertebrates and invertebrates, can predict the future position of a prey by using internal models [31,39,40]. In our model, we assumed that predators simply predicted the prey’s future position based on the prey’s direction, which reflects the prediction that is based on internal models. The evolution of the neural circuits associated with this prediction may be driven by the movements of prey. Further studies could reveal the co-evolution of prey’s movements and predators’ cognitive abilities.

In our framework, the environmental conditions were limited (e.g. circle). Indeed, in natural conditions, complex obstacles may be present, however the trade-off between the prediction rate and the time of exposure to the predation risk would still exist. It is important that future research addresses how environmental heterogeneity and the strategy’s dependency on environmental influences, affect prey-predator interactions [41]. Furthermore, considering situations in which a predator and a prey predict one another’s movements, is important in helping us gain a full understanding of prey-predator interactions.
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Authors’ contribution

M.S.A designed the study. M.S.A and M.K. conducted the analysis. M.S.A. wrote the first draft of the manuscript. All authors contributed to the final version of the manuscript, gave final approval for publication, and agree to be held accountable for the work performed therein.

Competing interests

We declare we have no competing interests
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Figure 1. Random avoidance in prey-predator interactions.

(a) We considered a scenario in which a predator observes from above a prey moving in an open area. (b) A predator’s view and the example of movement trajectories of a prey for $\mu = 1.1, 2.0$ and 3.0. Note that $\mu = 1.1$ and 3.0 roughly correspond to straight movement and Brownian walks, respectively. The green area is a safety zone. (c) How to predict the prey’s position. At time $t$ the predator must predict the future (after time step $\tau$) position of the prey. A red circle is an attack area with radius $r$ and its centre is the predicted position $\hat{X}_{t+\tau} = (\hat{x}_{t+\tau}, \hat{y}_{t+\tau})$. If the prey moves within the red circle after time step $\tau$, the predator can be considered successful in having predicted the prey’s position and in attacking it. When the prey is outside the red circle after time step $\tau$, the predator’s attack is regarded as a failure. Then, the predator tries to predict the next future position (after waiting time $W$ has elapsed) and attack the prey. The solid arrows represent the actual trajectory of the prey’s movement and the dashed arrow represents the predator’s prediction.
Figure 2. The relationship between the prediction rate and time length to reach the safety area.
The horizontal axis is a prediction rate $Q$ for $r \to 0$ obtained by an analytical solution and the vertical axis is the average time length to reach the safety area for the precaution case, which was obtained by simulations with different area sizes (a) $R = 10$, (b) $R = 50$ and (c) $R = 100$. The colour represents the prey’s strategy parameter $\mu$. The cut-off values of step length that we used were $l_{\text{min}} = 1$ and $l_{\text{max}} = 1,000$. The number of iterations for our simulation was $10^6$. 
Figure 3. The simulated survival rate for the precaution case depended on $\mu$. The horizontal axis is a prey’s strategy parameter $\mu$ and the vertical axis is the survival rate of the prey. The colour indicates the prediction time lag $\tau$. The points represent the maximum survival rate for a specified parameter set ($R$, $W$, and $\tau$). The cut-off values of step length that we used were $l_{\text{min}} = 1$ and $l_{\text{max}} = 1,000$. The attack radius $r$ was set as 0.01. The number of iterations for our simulation was $10^6$. 