Assessment of movement patterns in Folsomia candida (Hexapoda: Collembola) in the presence of food
Apolline Auclerc, Paul-Antoine Libourel, Sandrine Salmon, Vincent Bels, Jean-François Ponge

To cite this version:

HAL Id: hal-00493987
https://hal.archives-ouvertes.fr/hal-00493987
Submitted on 21 Jun 2010

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Type of contribution: Short communication

Date of preparation: 18 December 2009

Title: Assessment of movement patterns in *Folsomia candida* (Hexapoda: Collembola) in the presence of food

Names of authors: A. Auclerc, P.A. Libourel, S. Salmon, V. Bels, J.F. Ponge

Complete postal addresses or affiliations:

A. Auclerc, S. Salmon, J.F. Ponge: Muséum National d'Histoire Naturelle, CNRS UMR 7179, 4 avenue du Petit-Chateau, 91800 Brunoy, France

P.A. Libourel, V. Bels: Muséum National d'Histoire Naturelle, CNRS UMR 7179, 55 rue Buffon, Case Postale 55, 75231 Paris Cedex 5, France

Full telephone, Fax number and E-mail address of the corresponding author:

Tel. +33 6 78930133
Fax +33 1 60465719
E-mail: ponge@mnhn.fr

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*Complete correspondence address to which the proofs should be sent: Jean-François Ponge, Muséum National d'Histoire Naturelle, CNRS UMR 7179, 4 avenue du Petit-Château, 91800 Brunoy, France*
Abstract

We showed that *Folsomia candida* (a blind soil-dwelling Collembola) was able to shift from non-directional (random or search strategy) to directional (target-oriented) movements at short distance of food. We measured departure from linearity and access (or not) to food by the springtail according to distance to the target position. Video-records and image analysis were used to obtain numerical data at 0.2 sec interval. The probability of food capture within 10 min (maximum duration of the experiment) was negatively related to distance. Two patterns can be observed along successful trajectories in our experimental conditions (22°C, ambient light, still air), non-directional movement being followed by directional movement when the animals approach food at 25 mm.

Keywords: Soil invertebrates; directional movement; food capture

Text

More has to be learned about the way soil invertebrates reach a target food or other favoured place without spending too much energy. While directional movements can be expected to waste less energy and time than random movements they need sophisticated sensory and nervous equipment (Applebaum and Heifetz, 1999). Between random and directional movements a third category consists of efficient ‘search strategy’, using looping behaviour (Bell, 1991). Bengtsson et al. (2004) and Wiktorsson et al. (2004) studied the small-scale movement of springtails in environments without food as attractant. Westerberg et al. (2008) showed that the presence of food decreased the time spent to move and modified looping behaviour. It was demonstrated that chemical cues are used by Collembola for attraction or avoidance, but experiments were made either by transporting odours via air currents in olfactometers (Hedlund et al., 1995) or using odour-conditioned places (Sadaka-Laulan et al., 1998; Nilsson and Bengtsson, 2004). It can be expected that (i) blind or eye-reduced soil-dwelling species will use rather odours as clues, contrary to aboveground species which use visual cues to move over long distances (Hågvar, 1995), and that (ii) this is possible only at short distance if
only brownian motion (diffusion) transports olfactory molecules. Salmon and Ponge (2001) showed that the odour of earthworm excreta attracted the soil-dwelling *Heteromurus nitidus* at 1 cm distance, which is far below the range of higher insects (Laubertie et al., 2006). By studying the movement of animals deposited at varied distances from a food source we expect to find (i) a negative relationship between distance and success of food capture, and (ii) a shift from non-oriented (random or ‘search strategy’) to directional movements when food is perceived.

*Folsomia candida* Willem 1902 was used because of its insensitivity to light, which was verified beforehand: intolerance or attraction to light might interfere with the direction of movement (Salmon and Ponge, 1998). The trajectory was studied by video-tracking naive animals in short-time experiments (<10 min). Experiments were performed without soil in order to avoid any heterogeneity of the substrate and to follow individual animals continuously through an optical system. For that purpose the arena was made of a fine black cotton cloth which was thoroughly rinsed under tap water before each experimental run, thus avoiding pheromone deposition (Verhoef et al., 1977). The absence of air turbulence was provided by placing a glass cylinder 14 cm diameter and 30 cm height above the cotton cloth, which delimited the arena. The temperature was 22±1°C and the light was ambient light, to which *F. candida* was insensitive, as ascertained by preliminary experiments.

The animals were reared for several years on fine quartz sand moistened with tap water and were fed ad libitum with dried powdered cowdung. They also ate their excrements, which formed a thick layer covering the sand. This layer, made of an intimate mixture of cowdung debris and faeces, was used as attractant food in the experiments, *Folsomia candida* being pheromone-conditioned (Leonard and Bradbury, 1984). Animals (adults and sub-adults) were fed *ad libitum* until experiment. Food (~1 mg) was placed as a mound at the centre of the arena, 3-5 minutes before each experimental run, and was covered with a short piece of white cloth of similar area (~0.25 cm²) for the need of image analysis. For each video-recorded run, one naive animal was introduced with a syringe at an uncontrolled distance from the side of the white cloth, varying from 1 to 50 mm (Fig. 1), and its movement was recorded with a Sony® DSR-PD100AP camera mounted on an adjustable support and
connected to a personal computer, starting from the time the animal was deposited in the arena. On each run, which ended when food was encountered and movement ceased (the animal was no longer visible under the white cloth) or at 10 min, white animals were followed on the black background by image analysis using Simulink® software. At each 0.2 sec interval visual data were transformed into X and Y values for further calculations by Excel® software. For the need of calculation, the path followed by the animal within each 0.2 sec interval was assimilated to a straight line. By avoiding vibrations and shocks care was taken that no avoidance jump using the furcula occurred during the experiment. At each time of recording, the length of the trajectory followed by an animal from start was calculated by summing up 0.2 sec segments. This allowed us to compare the distance remaining to travel to the distance in straight line to food. An efficiency index of movement (E.I.) was calculated by dividing the length of the trajectory by the straight line joining start and end of movement record.

The examination of the whole set of records (118) showed that about one third of animals (35) reached the food in the course of their wandering. Systematic search using looping behaviour, as described by Bengtsson et al. (2004), was not observed, although paths followed by animals could be crossed several times in the course of a single run. Successful food capture was negatively influenced by the distance at which the animals were placed at the start of the experiment, the rate of food capture decreasing from 65% at less than 1 cm to zero beyond 4 cm (Fig. 1). Logistic regression was performed with Addinsoft XLSTAT®, using food capture (0 = failure, 1 = success) as dependent variable and distance to food at run start as independent variable. It showed that the probability of food capture (within 10 min) was negatively influenced by the initial distance between the animal and the target food (P<0.0001). The efficiency of movement, as expressed by E.F., was 12.1 (± 2.4 S.E.) in case of failure vs 4.5 (± 0.8 S.E.) in case of success of food capture (Kolmogorov-Smirnov test, P<0.0001).

The trajectory followed by an animal was studied when it reached food within the maximum duration (10 min) of the experiment (35 runs). The latter distance decreases in the course of the experiment, while the former increases or decreases according to the kind of behaviour of the animal.
As an example, Figure 2 displays the characteristic patterns exhibited by an animal that reached food in the course of the experiment. Two patterns can be distinguished, a first pattern corresponding to wandering around the original position of the animal (random movement or ‘search strategy’), and a second pattern where and when the animal approaches food continuously. Although the second pattern is represented by a straight line on Figure 2, the actual 2D-movement may be curvilinear (data not shown). The straight distance to food below which the linear part was observed along the curve measures the distance at which movement was directional. The average value was calculated to be 25.2 mm (± 1.4 mm S.E.). Given that the passage from the first to the second pattern of movement was not exhibited at the same time for the whole set of animals tested, and because the rate of locomotion strongly differed from an animal to another and within the same individual record, it was judged impossible to incorporate time in a predictive model.

Within the limits of our study we confirm that olfactory signals are detected by soil invertebrates over only a few centimetres, as already observed by Salmon and Ponge (2001) in H. nitidus when attracted to earthworm odour. Here our method allows us to estimate it being ~2.5 cm in F. candida, i.e. 1.5 cm longer than in earthworm-attracted H. nitidus. The examination of the whole set of successful trajectories (35) showed that the movement of an animal placed in the vicinity of a food source starts with non-directional movements (as exemplified by the zigzag part of the curve on Fig. 2) followed by directional movements (as exemplified by the final ‘straight line’ on Fig. 2). F. candida being insensitive to light, visual inspection of the environment can be discarded as an explanatory factor, but we cannot rule out that other factors than odour could contribute to reveal the presence of food, such as temperature gradients issuing from microbial respiratory metabolism.

References


**Figure captions**

**Figure 1.** The influence of distance to food at run start (abscissa) on the probability of food capture within experimental time (ordinate). Distances were classified in five groups (n = number of runs).

**Figure 2.** Graphical representation of the distance remaining to travel along a given trajectory according to straight distance to food (an individual run is shown here as example). The zigzag part of the curve (non-directional movement) is followed by a linear part (directional movement).
Fig. 1
Fig. 2