Minimum-Risk Path Finding by an Adaptive Amoebal Network

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When two food sources are presented to the slime mold Physarum in the dark, a thick tube for absorbing nutrients is formed that connects the food sources through the shortest route. When the light-avoiding organism is partially illuminated, however, the tube connecting the food sources follows a different route. Defining risk as the experimentally measurable rate of light-avoiding movement, the minimum-risk path is exhibited by the organism, determined by integrating along the path. A model for an adaptive-tube network is presented that is in good agreement with the experimental observations.

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Introduction.—The plasmodium of Physarum polycephalum is an amoebalike organism with a body made up of a tubular network through which nutrients, signals, and body mass are transported. Studies of this organism have shown that it is able to determine the shortest path through a maze as well as “solve” other geometric puzzles [1–3]. In a maze, a starved organism forms a tube that connects food sources (FS) placed at the two exits of the maze via the shortest path, while nearly the entire protoplasm of the amoeba gathers over the two FS. The organism meets its physiological requirements in adopting this shape by absorbing nutrients from the FS as rapidly as possible while maintaining sufficient connectivity to permit intracellular communication. Such behavior in a primitive organism of this kind may offer insights into the evolutionary origins of biological information processing.

Here we give the plasmodium a new type of task involving optimization behavior. Two separate FS are presented to the organism, which is illuminated by an inhomogeneous light field. Because the plasmodium is photophobic, tubes connecting the FS do not follow the simple shortest paths but form according to the illumination inhomogeneity. We report on the behavior of the organism under these conditions and discuss its physiological significance. We also propose a mathematical model for the cell dynamics and present a computational algorithm for its problem solving.

Organism and methods.—The plasmodium of Physarum polycephalum, which regenerated from the sclerotia in ca. one-half day in the dark (25 °C), was used in the experiments. A plastic film was placed onto a 1% agar gel, leaving a rectangular area (1 × 2 cm²) of the gel uncovered. A few pieces (0.5 × 1 cm²) of the regenerated plasmodium were placed in the rectangular area, and the preparation was placed in the dark for a few hours. The organism readily spread over the rectangular area. Food sources, consisting of agar blocks containing powdered oat flakes (oat wt/agar solution = 0.1 g/ml), were presented at two sites, the upper left-hand and the lower right-hand corners. A part of the rectangular region was then illuminated with cold white light (Luminar, Hayashi Co., Japan). The illumination gives rise to the intracellular production of reactive oxygen, which is avoided by the organism [4]. As a control, the plasmodium was either uniformly illuminated (bright control) or not illuminated (dark control).

We next estimated the level of risk resulting from the illumination field. A plasmodium was placed at the end of the narrow channel of 1% agar gel (0.5 × 5 cm²), as shown in Fig. 1(a) (t = t₀). After a few hours, the plasmodium had extended to ca. 3 cm along the channel (t = t₁). We measured the rate of body mass transport from the rear (left-hand side) to the front (right-hand side) of the plasmodium by video image analysis [5], in which the organism was viewed from below by infrared imaging. The light intensity at each pixel of the image reflects the cell thickness. The posterior region of the organism (20%–30%) was illuminated with white light, and the infrared light intensity of this region was measured before and during the illumination as a function of time. An increase in intensity was found during white light illumination such that the ratio α₁ = (α_light)/(α_dark) > 1, where α_light and α_dark are the rates of intensity increase in the presence and absence, respectively, of white light illumination. This intensity increase is an indication of the risk experienced by the region of the organism when illuminated by white light, and we therefore refer to the ratio α₁ as the relative risk. Hence, the level of risk of the illuminated field was relative to that of the dark field, which was set at 1. A second estimation of risk was carried out in which we measured the tip position x of the extending plasmodium and cal-
calculated its extension velocity over a period of 20 min before and during illumination, defined as $V_0 = (x_2 - x_1)/(t_2 - t_1)$ and $V = (x_4 - x_3)/(t_4 - t_3)$, respectively [see Fig. 1(a)]. The velocity is also related to the risk of the organism that extends linearly. The organism (gray) before and during illumination, defined as calculated its extension velocity over a period of 20 min of the organism that extends linearly. The organism (gray) relative levels of risk in photoavoidance behavior. (a) Top view

FIG. 1. Schematic illustration of the procedure for measuring
relative levels of risk in photoavoidance behavior. (a) Top view
of the organism that extends linearly. The organism (gray)
extends from left to right in the time sequences from $t_0$ to $t_4$.
PL: plasmodium, $X_i$: position of the plasmodium tip. (b) Side
view of the experimental setup. VC (video camera for video
image analysis), FL (filter to eliminate visible light), WL (cold
white light to stimulate the posterior of the plasmodium), IR
(infrared light to observe the plasmodium).

Connection path with minimum risk.—At the beginning of the experiment the organism initially had a sheetlike morphology and a rectangular shape. Two FS were placed diagonally opposite one another, and an area of the rectangular organism was then illuminated with cold white light, indicated by the dashed outline in Fig. 2(a). The organism moved toward the FS and also formed several thick tubes that connected the FS. Eventually, only one main tube remained, as shown in Figs. 2(d)–2(f). Figure 2(d) shows a typical path, where a shift is exhibited at the boundary between the dark and illuminated fields. The path length in the illuminated field was shorter and the path length in the dark field was longer than the corresponding path in the homogeneous field, shown in Figs. 2(b) and 2(c). The connecting path varied from experiment to experiment; however, a statistical trace of the paths averaged over several experiments showed the path to consist of two straight lines, meeting at a point on the boundary between the dark and illuminated fields, as shown in Fig. 3. The point of intersection shifted to the left along the boundary as the illumination intensity was increased. The path length in the illuminated field therefore decreased progressively as the toxic effects of the light increased. This implies that the risk of damage to the organism was reduced at the expense of an increase in the total length of the path. The implications of this are considered next.

In order to consider the physiological significance of the path, the minimum-risk path between the FS, in which the risk is a minimum when integrated along the path, was calculated using the two relative levels of risk, $\alpha_1$ and $\alpha_2$, as shown by the dotted lines and the dashed lines in Fig. 3. Measured values of $\alpha_1$ and $\alpha_2$ were used for these comparisons. Since the connecting path in the dark control (uniformly dark conditions) was not significantly different from that in the bright control (uniformly bright conditions), the key parameter is not the absolute value of $\alpha$ but the ratio of $\alpha$ in the dark and bright fields. The minimum-risk paths are in good agreement with the measured path in strong light conditions, Figs. 3(d)–3(f), while there are larger but acceptable deviations in weak light conditions, Figs. 3(b) and 3(c). We conclude that Physarum is able to locate the minimum-risk path in an inhomogeneous risk field. Note that we assume the value of $\alpha$ is uniform in each region of uniform light intensity, and we neglect possible secondary influences on $\alpha$ such as boundary effects. From a mathematical point of view, minimizing the functional defining the minimum-risk path is a variational problem, which is typically solved by using Euler’s equation for the necessary conditions for the extremum of the functional. This method is based on a process of picking the best candidate from all possible paths. The plasmodium, however, does not use an algorithm of this kind. Since the mechanism by which the plasmodium solves the problem is an interesting example of a biological computation, we propose a mathematical model for this process.

FIG. 2. Photographs of the connecting paths between two food sources (FS). (a) The rectangular sheetlike morphology of the organism immediately before the presentation of two FS and illumination of the region indicated by the dashed white lines. (b),(c) Examples of connecting paths in the control experiment in which the field was uniformly illuminated. A thick tube was formed in a straight line (with some deviations) between the FS. (d)–(f) Typical connecting paths in a nonuniformly illuminated field (95 Klx). Path length was reduced in the illuminated field, although the total path length increased. Note that fluctuations in the path are exhibited from experiment to experiment.
which edge $M_{ij}$ plasmodium to be a randomly meshed lattice of tubes, in mutually antagonistic processes. The tube structure changes according to a balance of these accelerated in the illuminated part of the organism. Thus, organism moves toward the FS. This thickness decrease is diameter of the plasmodial tube) decreases with time as the thickness therefore adapts to flow rate. The converse pro-

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Mathematical model for tube selection.—Let us begin with the morphogenesis of the tubular structure of Physarum plasmodium: the mechanism for the tube appearance and disappearance. Cytological experiments in-
dicate that protoplasmic shuttle streaming plays a key role in tube formation [6]. Tubes become thicker and thinner as the flow rate increases and decreases, respectively. Tube thickness therefore adapts to flow rate. The converse process of tube collapse is also involved: cell thickness (or diameter of the plasmodial tube) decreases with time as the organism moves toward the FS. This thickness decrease is accelerated in the illuminated part of the organism. Thus, the tube structure changes according to a balance of these mutually antagonistic processes.

Let us suppose the initial sheetlike structure of the plasmodium to be a randomly meshed lattice of tubes, in which edge $M_{ij}$ connects nodes $N_i$ and $N_j$, as shown in Fig. 4(a). Two special nodes, $N_1$ and $N_2$, correspond to the food sources. One of the food source nodes ($N_1$) always acts as a flux source and the other ($N_2$) acts as a flux sink.

The variable $Q_{ij}$ is the flux through $M_{ij}$ from $N_i$ to $N_j$. Assuming approximate Poiseuille flow, the flux $Q_{ij}$ is given by $Q_{ij} = (D_{ij}/L_{ij})(p_i - p_j)$ and $D_{ij} = (\pi r_{ij}^4)/8\xi$,

where $p_i$ is the pressure at node $N_i$, $L_{ij}$ is the length of the edge $M_{ij}$, $D_{ij}$ is its conductivity, and $\xi$ and $r_{ij}$ are the viscosity coefficient and radius of the tube, respectively.

By considering the conservation law of flux at each node, we have $\sum_j Q_{ij} = 0$, where $j \neq 1, 2$. For the source node $N_1$ and the sink node $N_2$, $\sum_i Q_{ij} + I_0 = 0$ and $\sum_i Q_{ij} - I_0 = 0$, where $I_0$ is the flux flowing into the source node and out of the sink node. It should be noted that $I_0$ is a constant in our model.

The network Poisson equation for the pressure, derived from the above equations, is

$$\sum_i D_{ij} (p_i - p_j) = \begin{cases} -I_0 & \text{for } j = 1, \\ I_0 & \text{for } j = 2, \\ 0 & \text{otherwise}. \end{cases}$$

By setting $p_2 = 0$ as the basic pressure level, all of the $p_i$ can be determined by solving Eq. (1), in which each $Q_{ij} = (D_{ij}/L_{ij})(p_i - p_j)$ is also obtained.

In order to describe the dynamics of tube thickness, it is assumed that the conductivity $D_{ij}$ changes in time according to the flux $|Q_{ij}|$,

$$\frac{d}{dt} D_{ij} = |Q_{ij}| - aD_{ij}. \quad (2)$$

FIG. 3. Comparison between the measured connection path and the two routes based on minimum-risk estimates. Mean paths, averaged over ca. 10 experiments, are shown by the filled circles (with error bars) at light intensities of 0 lx (a), 15.000 lx (b), 35.000 lx (c), 55.000 lx (d), 75.000 lx (e), and 95.000 lx (f). Each point indicates the tube position on the vertical axis at the corresponding horizontal position. The dotted lines and the dashed lines indicate the minimum-risk paths calculated according to the relative rate of decrease in thickness $\alpha_1$ and the relative migration velocity $\alpha_2$, respectively. The $\alpha$ values were $\alpha_1 = (1.14, 1.03), (1.61, 1.04), (1.50, 1.40), (2.07, 1.49)$, and $(2.14, 1.62)$ at 15, 35, 55, 75, and 95 Klx, respectively. (The fluctuations in the $\alpha$ values are likely due to the difficulty of preparing each organism in exactly the same state.)

FIG. 4 (color online). Simulation of path formation. (a) Schematic illustration of the mathematical model. The tube network of Physarum is represented by a random mesh lattice. Between junctions $i$ and $j$ the tube has fixed length $L_{ij}$, variable radius $r_{ij}$, and the conductance $D_{ij} = r_{ij}^4$. All tubes initially have a similar small value of $r_{ij}$ with random fluctuations around the mean value. (b)–(e) Results of simulation. A constant current flows into the source in the upper left-hand corner and out of the sink on the lower right hand. Some paths including the minimum-risk path appear at an intermediate stage (c),(d) and finally only the minimum-risk path remains (e). (f) The minimum-risk path obtained by a conventional path finding algorithm, Dijkstra’s algorithm [10], where the length of a lattice edge was scaled by the relative level of risk. Above the dashed line $a = 2$ (corresponding to the illuminated region) and below the dashed line $a = 1$. The simulations utilized 20 000 nodes in order to generate a sufficiently fine meshwork. No change in the qualitative behavior was found when the number of nodes was increased.

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This relation implies that the conductivity tends to vanish exponentially according to the second term \(-a D_{ij}\), while it is enhanced by the flux along an edge according to the first term \([Q_{ij}]\). The tube lengths \(L_{ij}\) remain constant throughout the adaptation process, and changes in the conductance therefore result in changes in tube radius. Note that \(a\) is a kinetic constant in the process of tube thinning.

It should be noted that the variable \(D_{ij}\) evolves following the adaptation Eq. (2), while the variables such as \(p_i, Q_{ij}\) are determined by solving the network Poisson Eq. (1) characterized by the values of \(D_{ij}\) at each moment. (See [7,8] for mathematical details of the model.) Here we focus on the effects of parameter \(a\) on the model behavior in order to develop an understanding of the effects of homogeneity in the experimental system. We next consider the relationship of the process of tube thinning [second term in Eq. (2)] to the experimentally measured sensitivity to the exponent (4, 3, or 2) and only affects the time to the final path. [Note that the total flux \(I_0\), plays an essential role in the global optimization in the model.

**Discussion.**—We note that the minimum-risk path exhibited by the *Physarum* plasmodium and in our adaptive-tube network model has geometric features much like the path of light in two materials with differing refractive indices. While the path of light according to Snell’s law depends on the speed of light in different materials, the minimum-risk path results from an optimization process in which tube thickness depends on competing processes that are influenced by the photoavoidance response of the plasmodium.

The global evolution of the tube network arises from local tube dynamics. The simplicity of such dynamics offers insights into the optimization behavior of the plasmodium. We also note that the evolution of the tube network shares features in common with Hebbian learning found in neuronal networks [9], as the tubes grow or shrink and disappear based on their level of activity.

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Each tube evolves in parallel over time according to the local dynamics, which may be regarded as a form of parallel computation. The model equations lead to the optimal solution much like the experimentally observed optimization in the plasmodium. The conserved quantity, the total flux \(I_0\), plays an essential role in the global optimization in the model.

For the sake of simplicity we assumed a constant flux \(I_0\); however, the behavior is much the same if the flux is replaced by \(I_0 \sin 2\pi \omega t\). This is consistent with the experimental observation that protoplasmic sol flows periodically back and forth between the FS. An important requirement for path finding is sufficient flow of sol through the tubes. Since the total amount of sol in the organism is conserved, shuttle streaming of sol is necessary for achieving flow. Protoplasmic shuttle streaming thus plays an essential role in path finding.

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