Proteolysis and orientation in *Dictyostelium* slugs

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It has been long known that the migrating slugs of the cellular slime moulds are highly sensitive to their environment and orient towards light and in temperature and chemical gradients. There is considerable evidence from past work that these orientations are governed by NH₃ which affects the rate of movement of cells within the slug with such precision that orientation to the external stimuli is achieved. In order to test this hypothesis further, various ways to alter the internal NH₃ concentration were devised. Substances that either increased or decreased proteolysis were applied to one side of the tip of a slug, thereby affecting its orientation. Some of the treatments strongly support the role of internally produced NH₃ in orientation, and all the treatments produce results that are consistent with the hypothesis.

Introduction

In recent years we have been accumulating evidence that orientation in the migrating slugs and rising cell masses of cellular slime moulds is governed by the internal concentration of NH₃ within the mass. First it was shown that the application of minute amounts of external NH₃ gas repels the slugs (Bonner et al., 1986; Feit & Sollitto, 1987; Kosugi & Inouye, 1989) and therefore the volatile orienting substance described much earlier (Bonner & Dodd, 1962) is probably NH₃. It was presumed that it did this by speeding up the cells on the side of the slug, or rising sorogen, which was surrounded by a higher concentration of NH₃, with the result that the cell mass moved away from the NH₃. More recently we have shown by measuring slug speed, and the speed of separate, preaggregation amoebae, that there is an optimal concentration of NH₃ which makes the slugs and the cells move faster, while at higher concentrations the speed is inhibited (Bonner et al., 1989). As Kosugi & Inouye (1989) and Van Duijn & Inouye (1991) have shown, there is evidence that these effects of NH₃ are due to changes in the internal pH of the cells, for NH₃ penetrates cells rapidly and raises their pH. It has been suggested that the striking ability of slime mould cell masses to orient towards light might be explained by the fact that light increases the internal NH₃ production, and because of the ‘lens effect’ the far side will be more illuminated than the near (Bonner et al., 1988). We have also tried to explain orientation in heat gradients in terms of internal NH₃ production, but here the evidence becomes more tenuous because, depending upon the ambient temperature, there may be either a positive or a negative thermotaxis (Whitaker & Poff, 1980) and therefore the hypothesis requires more assumptions. Since it is clear that NH₃ production within the slug might play a role in orientation, I decided to try to find ways of directly influencing the NH₃ production which, in turn, should affect orientation. As will be evident, all of the experiments reported here are consistent with the NH₃ orientation hypothesis, although some provide far more compelling evidence than others.

Methods

The spores of *Dictyostelium discoideum* (strain NC-4) were placed on a mound of *Escherichia coli* B/r made by plunging a loopful of bacteria into 2% (w/v) non-nutrient agar in three spots on a Petri dish, thereby providing plenty of room for the migration of the slugs. All the culturing and experiments were done at room temperature. The experiments were recorded on videotape taken through a 50 mm lens attached through a microscope to a Panasonic video camera (WV-1850) with time lapse (AG-6720A).

Chemicals. The activated charcoal used was Darco G-60 (Fisher Scientific Co.) The papain came in a lyophylized form (Sigma). All the protease inhibitors used were the water-soluble ones in a protease inhibitor kit (Boehringer Mannheim) which was given to me through the generosity of Dr D. Fong (Rutgers Univ., Piscataway, NJ, USA). The hydrolysed polyacrylamide (‘Hypa’) beads were kindly supplied by Drs M. S. Steinberg and J. Drawbridge (Princeton Univ.) following the method of preparation of Zackson & Steinberg (1989). The beads were soaked in the test solutions from 15 min to 1 h.

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Results and Discussion

Activated charcoal

Before considering proteolysis and orientation, I would like to report one experiment suggested to me by Kei Inouye which supports the role of NH₃ in orientation. It has been known for many years that slime mould cell masses will orient towards charcoal (Bonner & Dodd, 1962) presumably because the charcoal absorbs and therefore removes the repellent gas (Fig. 1a). If the charcoal is first placed in an atmosphere of high NH₃ in a desiccator jar (over a mixture of 20 ml 30% (w/v) NH₄Cl and 50 ml 1 M-NaOH in a well) for at least 2 d or more, and then tested by being placed close to a migrating slug, the charcoal will have no effect on the orientation of the slug at all (Fig. 1b). In other words, if the charcoal is saturated with NH₃ it is no longer capable of attracting the slug; it can no longer remove the NH₃ in the atmosphere on one side of the slug.

Another way of testing activated charcoal is to measure the speed of a migrating slug before and after it has been sprinkled lightly with fresh charcoal. In 24 cases the mean speed before treatment with charcoal was 2-10 mm h⁻¹, while after dusting the speed slowed to 0.78 mm h⁻¹. [The difference is significant at a level of $P = 0.0001$ in a paired (one-tailed) t-test.]

Table 1. Effect of different treatments applied to the tip of a slug that cause the slug to either turn away or towards the treated side

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Conc</th>
<th>No. of slugs that:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Turn away</td>
</tr>
<tr>
<td>Papain</td>
<td>5.5 pg ml⁻¹</td>
<td>10</td>
</tr>
<tr>
<td>Wounding</td>
<td></td>
<td>11</td>
</tr>
<tr>
<td>Acid Dowex 50</td>
<td></td>
<td>24</td>
</tr>
<tr>
<td>Antipain</td>
<td>50 pg ml⁻¹</td>
<td>0</td>
</tr>
<tr>
<td>Phosphoramidon</td>
<td>300 pg ml⁻¹</td>
<td>0</td>
</tr>
<tr>
<td>Ethanol</td>
<td>3-30% (v/v)</td>
<td>0</td>
</tr>
</tbody>
</table>

A proteolytic enzyme

In these experiments a hydrolysed polyacrylamide ('Hypa') bead was soaked in a protease solution after which it was placed on one side of the tip of a migrating slug. If papain (5-5 mg ml⁻¹) was used, after 2 to 10 min the slug tip would take a sharp turn away from the side where the bead was placed (Fig. 2a; Table 1). Because it was known from previous work that papain digested the slime sheath (Whitfield, 1964; Takeuchi & Yabuno, 1970) it is reasonable that in the experiment here the papain caused proteolysis in a localized region of the slime sheath. This in turn resulted in a liberation of NH₃ on one side of the slug, causing the cells on that side to move faster, with the result that the slug tip turned away from the bead. If the bead is put in place for as little as 3 min and then removed, the turning effect occurs. Once the NH₃ is made it will diffuse rapidly. It must be remembered that turning can only occur at the tip and the rest of the slug follows, and therefore only the proteolysis that takes place at one side of the tip can have any effect.
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Wounding

If a slug is cut on one side of the tip, or if the cells on that side are disrupted with a fine glass needle, the tip will abruptly turn away from the wound (Fig. 3a; Table 1). One might have imagined that such a localized trauma would cause a decrease in the speed of movement at the site of the wound, but the reverse is true. It is well known that in mammals there is an increase in proteolytic activity at wound sites (review: Rackallio, 1970), and conceivably slime moulds respond the same way. This interpretation is supported by the fact that if charcoal (Darco G-60) is added immediately after the cut is made, the slug is not affected by the cut and wraps itself around the charcoal. This is presumably because the charcoal removes all the NH₃, including the NH₃ generated by the proteases in the wound.

Acid

Another factor which seems to have a profound effect on the rate of cell movement is pH. When slugs crawl over agar of different pH values, the more acid the agar the faster they will move. This was shown in two ways: the slugs were allowed to crawl onto small Nuclepore membranes, and these membranes were first placed over buffered agar of different pH values. Similar results were obtained when groups of slugs crawled directly on buffered agar of different pH values. (If the various results are grouped, the mean speed of slugs at pH 5.5 is 1.47 mm h⁻¹ ± SD 0.19, n = 18; at pH 8.5 the speed is 1.20 mm h⁻¹ ± SD 0.12, n = 8. Using a two-tailed t-test, the difference is significant at the P = 0.001 level.)

The effect can be shown dramatically with the use of ion exchange beads. If an acid Dowex 50 bead is placed on the side, near the tip of a slug, the tip will rapidly move away from the bead: the cells near the point where the bead touches move faster than those on the opposite side (Fig. 3b; Table 1). This is true if the bead is first treated with HCl, NaOH or NH₃Cl. That this effect is due to the acid is supported by various controls, such as basic Dowex 1 beads, or glass chips which do not affect the slug orientation in any way whatsoever when they are similarly placed on the side of a slug tip.

There is the question of why an acid environment would cause cells to move more rapidly. The low pH of the substrate surface is unlikely to affect the pH within the cells as Kay et al. (1986) have shown. One possibility is that there are proteases in the slime sheath among the many different proteins Smith & Williams (1979) found there, and lowering the pH at the contact surface might be more favourable for inducing their proteolytic activity. Furthermore, it is known that slime moulds secrete proteases in large quantities and they are acid proteases, with highest activity at low pH values (North & Harwood, 1979; North, 1982).

Protease inhibitors

To perform the mirror image of the above experiments, various proteolytic enzyme inhibitors were allowed to be soaked up by the polyacrylamide beads and applied to the tips of migrating slugs in the same manner as with papain above. If a mixture of five different water-soluble inhibitors was used the slugs bent around the bead; clearly the inhibitors were slowing the cells on the side to which they were applied (Fig. 2b). When they were tested singly it was found that antipain (50 µg ml⁻¹) and phosphoramidon (300 µg ml⁻¹) were active (Table 1) while leupeptin (0.5 and 1.0 µg ml⁻¹), EDTA-Na₂ (0.5 mg ml⁻¹) and APMSF (4-amidinophenylmethanesulphonyl fluoride; 40 µg ml⁻¹) had no effect. Since the first two inhibitors differ in the kinds of protease they affect (phosphoramidon is a metalloprotease inhibitor and antipain inhibits cysteine- and serine proteases, and since some of the inhibitors that did not work do inhibit the same type of proteases, I suspect the reason for success or failure is due simply to which ones manage to diffuse readily into the cells.

Ethanol

In testing an enzymic method for removing NH₃ [a mixture of 2-oxoglutarate, NADH and glutamate dehydrogenase – first used by Schindler & Sussman (1977) for slime moulds], I found that the slugs curled inwards, towards the bead, as was expected. It was also possible to show that if a drop of the solution was placed on a slug, all migration movement stopped, as Schindler and...
Sussman had discovered. The surprise came upon discovering that the controls with beads containing an NADH solution only also caused slugs to bend around the beads. However, it turned out not to be the NADH but the small amount of ethanol in the NADH preparation. Using beads containing ethanol alone (in concentrations ranging from 3 to 30%, v/v) it was possible to show a reduction of the speed of the cells on the side touching the bead (Table 1), appearing very much like the slug shown in Fig. 2(b).

Because of the enormous medical interest in the effects of ethanol, it has been known since the last century that the small amount of ethanol in the beads. However, it turned out not to be the explanation of why ethanol slows the cells of internal proteolysis and therefore NH$_3$ production on one side of the slug. Obviously, there are other possible reasons, such as the general effect of ethanol in depressing metabolism (review: Wallgren, 1971).

**Conclusion**

All the experiments reported here support the hypothesis that orientation in *Dictyostelium* slugs is propelled by local differences in the NH$_3$ concentration in the slug tip, and that these differences are brought about by variations in the breakdown of proteins in different areas of the slug.

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**References**


