Selective Inactivation of an Extra-cytoplasmic Acid Phosphatase of Yeast-like Cells of *Sporothrix schenckii* by Sodium Fluoride

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Suspensions of intact, yeast-like cells of *Sporothrix schenckii* exhibited an acid phosphatase (EC 3.1.3.2) activity against p-nitrophenyl phosphate of about 5 IU (g dry wt)\(^{-1}\), without recourse to membrane perturbation. This extra-cytoplasmic acid phosphatase was reversibly and competitively inhibited by orthophosphate (\(K_i = 2\) mM at pH 5) but unaffected by L(+) tartrate (in contradistinction to some of the cytoplasmic acid phosphatases of the same organism). Inactivation by NaF of the extra-cytoplasmic isoenzyme was irreversible and followed first order kinetics; sensitivity to NaF was decreased by the presence of citrate, phosphate or substrate. Neither \(K_m (0.3\) mM at pH 5) nor \(V_{max}\) for this enzyme in acetate buffer was greatly affected by pH in the range 3–5 but the first order rate constant for inactivation by NaF was strongly dependent on pH (maximum at pH 3.5). Crude cell-free extracts of yeast cells had nine electrophoretically distinct acid phosphatase activity bands and, on the basis of the pattern of inhibitors, the extra-cytoplasmic activity was identified as Y-I, an isoenzyme that barely penetrates standard polyacrylamide gel electropherograms. Additional evidence for the assignment came from selective inactivation of this isoenzyme by short treatments of intact cells with NaF under conditions that did not allow penetration of the plasma membrane by the inhibitor and did not kill the cells.

**INTRODUCTION**

Mycelial and yeast cells of the dimorphic fungus *Sporothrix schenckii* both exhibit a multiplicity of cytoplasmic and extracytoplasmic sites of acid phosphatase (EC 3.1.3.2) activity (Garrison & Arnold, 1983). The acid phosphatases are ubiquitous among yeast and yeast-like organisms (Arnold, 1981), but they are of special interest with *S. schenckii* because of differences in the isoenzyme patterns among cells of the saprophytic and pathogenic forms (Arnold et al., 1986).

*S. schenckii* has world-wide distribution in soil and on plants as saprophytic mycelia adorned with conidia. Subcutaneous introduction of mycelia or conidia into animals can result in sporotrichosis, due to proliferation in the host tissues of yeast-like cells that arise by transformation at the higher temperature and multiply by budding. Several reports on ultrastructure and immunochemistry (for a review see Travassos & Lloyd, 1980) have appeared, but there has been a paucity of investigations on the enzymology of *S. schenckii*. Our recent findings on acid phosphatase isoenzymes are the first to demonstrate enzymic differences associated with dimorphism in this fungus. Different cell types of *S. schenckii* can be produced simply by regulating the growth temperature and this, together with the medical relevance, commend this organism for further studies of mycotic dimorphism.

We have reported (Arnold et al., 1986) that the nine isoenzymes of acid phosphatase in yeast cells of *S. schenckii* are inhibited to varying degrees by fluoride, L(+) tartrate and orthophosphate. In cytochemical studies with glutaraldehyde-fixed (and concomitantly...
permeabilized) cells, the addition of NaF to control incubations with a modified Gomori (1950) reagent obliterated all the sites of acid phosphatase activity (Garrison & Arnold, 1983). The generality of the inhibition did not help to discriminate the isoenzymes or the sub-cellular sites. However, we now report that the extra-cytoplasmic fraction of acid phosphatase (in intact yeast cells) can be irreversibly and selectively inactivated by low concentrations of NaF. This observation, together with the results of other inhibition studies on intact and broken cells, and correlative electrophoretic analyses, has provided sufficient evidence to identify the isoenzyme in question.

**METHODS**

*Organism.* *Sporothrix schenckii* (strain 29, Institut Pasteur) was grown in liquid YNG medium, which contained yeast extract (0-3%, w/v), neopeptone (0-5%, w/v) and glucose (1%, w/v), at 35 °C, and with shaking (175 orbits min⁻¹). Flasks received a small inoculum of mycelia plus conidia (from a 3 d culture on YNG at 23 °C) and after 3 d at 35 °C (and two orders of magnitude cell mass increase) the cultures were composed almost exclusively of yeast cells (confirmed by light and electron microscopy). Cells were harvested by centrifugation, washed once in 0-25 M-sodium acetate/acetid acid ('acetate') buffer (pH 6-5) and resuspended in the same buffer to 20% (wt/wt/vol).

**Enzyme assays.** One international unit (IU) of acid phosphatase is the amount of enzyme that catalyzes the hydrolysis of 1 μmol p-nitrophenyl phosphate min⁻¹ at 30 °C. The standard conditions were 20 mM-substrate in the presence of 0-2 M-acetate buffer (pH 5-0). Incubations were terminated by addition of NaOH (final concentration 0-4 M), clarified by centrifugation and assayed for nitrophenolate ion (absorbance at 425 nm).

Assays that involved NaF were done in polyethylene tubes. Specific activities are expressed on a dry wt basis; yeast concentrations were monitored turbidimetrically.

**Acid phosphatase isoenzymes.** Cells were broken by three passages of suspension through a French pressure cell operated at 138 MPa and 4 °C. The disrupted material was cleared by centrifugation (20 min, 20200 g, 4 °C), and a sample of the supernatant fraction was transferred to 20 mM-glycine/Tris, pH 7-5, by gel permeation chromatography. Samples were then subjected to electrophoresis in a continous buffer system (80 mM-glycine/5 mM-Tris, pH 8-0), through polyacrylamide gel (7.7%, w/v, total acrylamide; 2.6% crosslinker) cylinders at 4 °C. Acid phosphatase activity bands were revealed by equilibrating individual gels with 0-2 M-acetate buffer (pH 5-0) and then incubating them with a reagent (Davis & Ornstein, 1959) composed of α-naphthyl phosphate and freshly diazotized pararosaniline, also at pH 5-0. Gels were subsequently washed in cold buffer followed by 7% (v/v) acetic acid and stored in the cold. All of the above methods were described in detail by Arnold et al. (1986).

**Materials.** Media components were from Difco. Monomers and catalysts for casting gels were from Bio-Rad; AR grade chemicals were used throughout; NaF was further purified by recrystallization from distilled water and propan-2-ol (Ivett & DeVries, 1941) in plastic-ware.

**RESULTS AND DISCUSSION**

**Extra-cytoplasmic fraction of acid phosphatase**

The amounts of enzyme found in spent media or cellular washes were less than 4% of the total so secretion by live cells was not significant. Freshly harvested and washed cells exhibited 4-6 IU g⁻¹. In intact baker's yeast cells it is known that p-nitrophenyl phosphate and other phosphate esters penetrate the cell wall but are excluded from the cytoplasm by the plasma membrane (Arnold & Lacy, 1977). Assays on untreated cell suspensions give an accurate estimate of extra-cytoplasmic activity, i.e. of enzymes on the plasma membrane (with active sites disposed toward the medium) plus enzymes external to the plasma membrane (Arnold, 1981). In the case of *S. schenckii* yeast cells the extra-cytoplasmic activity is reasonably equated with two of the Golzori-positive sites in micrographs (Garrison & Arnold, 1983), namely the periplasm and the microfibrillar zone.

Mechanical disruption of *S. schenckii* yeast cells results in an increase in acid phosphatase activity of 50-75% (Arnold et al., 1986; this paper); the increment is due to the exposure of the cytoplasmic enzymes to substrate and can be equated with vacuolar and internal membrane sites (Garrison & Arnold, 1983).

**Kinetic parameters for intact cells**

The hydrolysis of p-nitrophenyl phosphate by cell suspensions followed Michaelis kinetics. The *Kₘ* value for this substrate in the presence of 0-2 M-acetate buffer (pH 5-0) was determined
NaF inactivation of S. schenckii phosphatase

Fig. 1. Effect of pH. The Michaelis constant ($K_m$) and the maximum velocity ($V_{max}$) were determined over a range of pH values with p-nitrophenyl phosphate as substrate. $pK_m = -\log K_m$.

Fig. 2. Inhibition by NaF and protection by citrate. Yeast cells were assayed for acid phosphatase at 30 °C and pH 5.0 in the presence of a range of NaF concentrations, in the presence (●) and absence (○) of 80 mM-citrate.

Table 1. Reversible inhibition by phosphate

Cell suspensions were incubated for 30 min at 30 °C in 0.2 M-acetate buffer (pH 5.0) and 20 mM-sodium phosphate as indicated. Acid phosphatase activities were then determined with p-nitrophenyl phosphate (1.6 KIM). The washing and resuspension medium was 0.2 M-acetate buffer (pH 5.0).

<table>
<thead>
<tr>
<th>Addition</th>
<th>Post-treatment</th>
<th>Specific activity [IU (g dry wt)⁻¹]</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>None</td>
<td>3.79</td>
</tr>
<tr>
<td>Phosphate</td>
<td>None</td>
<td>1.13</td>
</tr>
<tr>
<td>Phosphate</td>
<td>Washed once</td>
<td>3.68</td>
</tr>
<tr>
<td>Phosphate</td>
<td>Washed twice</td>
<td>3.86</td>
</tr>
</tbody>
</table>

on several batches of yeast: the mean value was 0.29 ± 0.02 mM ($n = 7$). The effects of pH on $K_m$ and $V_{max}$ are summarized in Fig. 1 where the pH values refer to complete assay mixtures, the buffer in this case being 0.2 M sodium acetate/HCl. There was less than 12% change in $V_{max}$ between pH 3.0 and 5.5. The $K_m$ value was insensitive to pH between pH 3.0 and 5.0 but increased outside that range. Dixon (1953) plots for the data of Fig. 1 were not easily interpreted because less than unit slopes were observed in the regions of change. There was some indication for the participation of two ionizable groups in the enzyme, one with a pK value of <3, and the other with a pK between 5 and 5.5, and for a group in the enzyme–substrate complex with a pK between 5 and 6 (cf. the pK₂ value for p-nitrophenyl phosphate which is approximately 5.7). The closeness of the projected pK values in the region 5 to 6 may explain the non-ideal curves.

Inhibition by phosphate

Sodium orthophosphate was a mild inhibitor of the acid phosphatase activity expressed by intact cells. Phosphate functions as a competitive inhibitor of p-nitrophenyl phosphate hydrolysis; the $K_i$, based on velocity measurements at pH 5.0 and substrate concentrations between 0.5 and 5.0 mM, with and without 10 mM-sodium phosphate, was approximately 2 mM. This inhibition was completely reversed by washing the cells (Table 1).

L(+)-Tartaric acid is a stereospecific inhibitor of some of the isoenzymes; however, at 40 mM it did not inhibit significantly the acid phosphatase of intact yeast cells. This concentration
Fig. 3. Effect of pH on the first order rate constant \( (K) \) for inactivation by NaF. Values of \( K \) were determined from plots of the natural logarithm of initial enzyme activity/residual activity after time \( t \), against \( t \) (not shown).

Fig. 4. Effect of pH on the concentration of fluorine species in aqueous NaF (0.1 mM) at 30 °C. ---, \( F^- \); ---, HF; ---, HF\(_2\) (note that the scale for HF\(_2\) is \( \times 1000 \)).

Table 2. Inhibition by fluoride

<table>
<thead>
<tr>
<th>Time of NaF addition relative to substrate</th>
<th>Acid phosphatase activity (normalized)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Not added</td>
<td>1.00</td>
</tr>
<tr>
<td>Zero time</td>
<td>1.02</td>
</tr>
<tr>
<td>1 min before</td>
<td>0.97</td>
</tr>
<tr>
<td>10 min before</td>
<td>0.62</td>
</tr>
<tr>
<td>20 min before</td>
<td>0.35</td>
</tr>
</tbody>
</table>

elicited only 3% lowering of the rate of \( p \)-nitrophenyl phosphate hydrolysis at pH 5.0, whereas sodium phosphate (40 mM) and NaF (2 mM) caused 66 and 95% decrease respectively under the same conditions.

Inhibition by fluoride

Preliminary trials revealed that phosphate, citrate and substrate can protect against inhibition by NaF. The latter effect is demonstrated by the results of Table 2; inactivation followed first order kinetics. The protective effect by citrate on NaF inactivation is clearly demonstrated by the data of Fig. 2: 0.84 mM-NaF was required for 50% inactivation in the presence of 80 mM-citrate compared with only 0.05 mM-NaF for the same response in the absence of citrate. Inactivation by NaF was essentially irreversible: cells that had suffered 96–98% inactivation of extra-cytoplasmic acid phosphatase activity had an apparent recovery of only 4–9% with up to three successive washes in acetate or citrate buffers respectively.

We also studied the effect of pH: in every case the inactivation by 0.1 mM-NaF followed first order kinetics. Rate constants were plotted against pH (Fig. 3) and showed a steady increase from pH 6 to pH 3.5, and then a precipitous decrease (i.e. the enzyme was less susceptible to
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inactivation). Controls indicated that the enzyme was not significantly inactivated by buffer alone over the pH range discussed.

An aqueous solution of NaF contains HF, F\(^-\), HF\(_2\) and Na\(^+\) in concentrations that are governed by pH and the total concentration of added salt. The following relationships (Broene & DeVries, 1947) apply to our conditions.

\[
\begin{align*}
HF &= H^+ + F^- \\
HF + F^- &= HF_2 \\
[HF] + [F^-] + [HF_2] &= 100 \mu M
\end{align*}
\]

The concentrations of all species were calculated for any pH value by using molar concentrations throughout and were used to construct the curves of Fig. 4. It is apparent that the concentration of HF\(_2\) (note the different scale) is always low. At the maximum, corresponding to the pK of HF, the concentration of the dimer is 10 nM compared with 50 \(\mu M\) each of HF and F\(^-\). Consequently other investigators, under similar conditions but for other enzymes (Dunford & Alberty, 1967; Segal *et al.*, 1968) have ignored HF\(_2\). Reiner *et al.* (1955) claimed that HF\(_2\) is the most important species participating in the reversible inactivation of prostatic acid phosphatase. That contention is not supported here, the only similarity being the biphasic nature for inactivation constant versus pH (Fig. 3). If a single fluorine-containing species were responsible the ratio of the inactivation rate constant to the concentration of that species would be invariant with pH. This was not found for any of the three species. The simplest working hypothesis at present is that both HF and F\(^-\) participate in the mechanism, and that the abrupt lowering in sensitivity below pH 3.5 may be explained by protonation of a critical amino acid side chain followed by a conformational change in the protein that affords protection from inactivation. Protonation alone is probably insufficient explanation because of the steepness of the curve below pH 3.5.

Selective inactivation with fluoride

It has been shown for higher plants (Peters *et al.*, 1965) and baker's yeast (Rothman & Cabib, 1966) that NaF is not toxic at pH values above 6-2 because F\(^-\) (more than 99-9% of the total at these pH values) does not penetrate the plasma membrane (Simon & Beevers, 1952). From the work of Arnold & Lacy (1977) it is clear that anions in general are not freely admitted by yeast plasma membranes in the absence of a metabolizable substrate. These properties, taken together with the irreversibility of inactivation by fluoride in the present case, suggested the following locational studies.

A 20% (w/v) suspension of yeast cells on 0-25 M-acetate buffer (pH 6.5) was divided into two samples. NaF (final concentration 1.125 \(\mu M\)) was added to one sample only and both were then kept at 30 °C for 30 min before washing in the same buffer. Yeast cells were not killed by these pretreatments as indicated by methylene blue exclusion and viability counts on agar plates. Samples were assayed for acid phosphatase activity and the remainder in each case was disrupted in a French pressure cell. Further assays on broken cells and derivative fractions, with and without inhibitors, completed the study summarized in Table 3. A completely independent experiment with another batch of yeast yielded similar results (Table 3). The irreversibility of the NaF treatment permitted washing away of excess inhibitor before cell disruption so that the newly exposed enzyme could be assayed in the absence or presence of NaF and other inhibitors.

The NaF pretreatment of intact cells irreversibly inactivated 90-94% of the extracytoplasmic acid phosphatase activity yet the same cells retained active cytoplasmic acid phosphatase (Table 3). The cytoplasmic activities (increments after cell breakage) were about 80% of those observed for untreated cells and they were also susceptible to inhibition by fluoride. These data indicate a selective effect of fluoride on intact cells, under the specified conditions.

A small but consistent fraction (3-5% of the total) in both sediments and supernatants resisted inhibition by 2 \(\mu M\)-NaF and, for the supernatants at least, may correspond to isoenzymes Y-IId, Y-IIe and Y-V (Arnold *et al.*, 1986) which are known to be resistant. It is worth noting that the
cytoplasmic fraction is to some extent susceptible to inactivation by phosphate and L(+)-
tartrate (Table 3) (Arnold et al., 1986). A significant fraction of the total activity in broken cells
was sedimented by centrifugation at 20200 g for 20 min; this fraction has been equated with
broken cell walls and various cytoplasmic membranes and particles (Arnold et al., 1986). Present
results together with those from unpublished trials indicate that the activity of debris is
consistently about 85% of that for intact cells. Cytochemical findings (Garrison & Arnold, 1983)
indicated that much of this insoluble activity may be associated with microfibrillar material. On
the other hand, a cytoplasmic contribution is apparent from the analyses of untreated cells and is
further supported by analysis of the NaF-treated cells (Table 3), wherein a 2.5-fold increase in
the activity of the sediment from broken cells over that for intact cells was noted.

Cell-free extracts were examined by gel electrophoresis and enzyme staining. The isoenzyme
most affected by NaF pretreatment of intact cells was Y-I, which is evidently a constituent of the
extra-cytoplasmic fraction. This is in keeping with the results of many previous analyses on crude extracts in that Y-I was strongly inhibited by NaF, mildly inhibited by phosphate, and not
inhibited by L(+)-tartrate, i.e. the same pattern of inhibition reported above for the extra-
cytoplasmic fraction of intact yeast cells. Only 6–10% of the extra-cytoplasmic fractions
survived the NaF pretreatment (Table 3), but scanning densitometer analysis of gels indicated
about 40% remaining in the treated extract compared with the control. A possible explanation is
that there is also some Y-I isoenzyme in the cytoplasm.

 Isoenzyme Y-I barely enters the gel in the present system of electrophoresis. We do not
consider that this activity is associated with particles (Arnold et al., 1986); however, it could be
due to a large native molecular mass to self-aggregation, or to association with other polymers in
the crude extract. A limited number of trials with detergents and higher porosity gels have not
yet afforded increased electrophoretic mobility but further attempts are being made.

Interest in isoenzyme Y-I is sharpened by the possibility that it is associated with the
microfibrillar zone of yeast cells, which would certainly allow direct interaction of this enzyme
with the pathogenic cells with macrophages and other host cells. The interaction of S. schenckii
yeast cells with human neutrophils is under current investigation. In this general context,
Remaley et al. (1984) claim a direct interaction of an acid phosphatase on the surface of the
protozoan Leishmania donovani with host macrophages.

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REFERENCES


NaF inactivation of S. schenckii phosphatase


