Thigmotropism and stretch-activated channels in the pathogenic fungus *Candida albicans*


Author for correspondence: N. A. R. Gow. Tel: +44 1224 273179. Fax: +44 1224 273144.
e-mail: n.gow@abdn.ac.uk

The direction of growth of hyphae of the pathogenic fungus *Candida albicans* responds thigmotropically to surface contours by following scratches, ridges and grooves and by penetrating pores. Here it is shown that the thigmotropic response to ridges is attenuated by GdCl₃ and verapamil [blockers of stretch-activated (SA) ion channels and L-type calcium channels, respectively]. At low concentrations, both compounds reduced the percentage of hyphae reorienting on contact with a ridge without markedly affecting hyphal extension rate, suggesting a possible role for SA or other calcium channels in the transduction of the thigmotropic response. In addition, patch-clamp recordings demonstrated SA channel activity in the plasma membrane of both yeast and hyphal cells of *C. albicans*. Two distinct SA channels with conductances of 54 pS and 20-25 pS in 200 mM KCl were observed in protoplasts from yeast cells and one channel of 51 pS was found in protoplasts from hyphal cells.

Keywords: *Candida albicans*, thigmotropism, stretch-activated channels

INTRODUCTION

*Candida albicans* is an opportunistic pathogen of humans, occurring as a normal commensal of up to 50% of asymptomatic individuals (Odds, 1988). It is a common agent of superficial mycoses of mucosal epithelia and an increasingly serious life-threatening pathogen in immunocompromised patients, being the fourth most common agent of septicemia (Beck-Sagué & Jarvis, 1993). The budding yeast form of this dimorphic fungus adheres strongly to mucosal surfaces, forming cavitations in epithelial surfaces (Ray & Payne, 1988). Yeast cells can also give rise to tip-growing hyphae. The direction of hyphal growth is guided by grooves, ridges and pores in the substratum and it has been suggested that this contact guidance (thigmotropism) may facilitate hyphal invasion of host epithelia through membrane invaginations, wounds or points of weakened surface integrity (Sherwood et al., 1992; Gow et al., 1994; Gow, 1996).

Thigmotropism is a well-known property of cells that grow on surfaces and within solid substrates. Some plant-pathogenic fungi, including the rust fungi *Uromyces appendiculatus* and *Puccinia hordei* (Hoch et al., 1987; Read et al., 1992) and the rice blast fungus *Magnaporthe grisea* (Xiao et al., 1994), display thigmotropism during germ tube growth on plant surfaces. This behaviour seems to expedite location of stomata through which the fungi invade the plant surface. In addition, topographical features of stomatal guard cells serve as signals for induction of appressorium formation. Fabricated topographies on inert substrata, thought to mimic inductive topographies associated with guard cells, induced in vitro thigmodifferentiation of rust fungi germ tube apices into appressoria (Hoch et al., 1987; Read et al., 1997). The plasma membrane from germ tubes of *U. appendiculatus* also contains stretch-activated (SA) ion channels, which are permeable to K⁺ and Ca²⁺ ions and inhibited by the SA channel blocker Gd³⁺ (Zhou et al., 1991). This suggests that localized stretching of the membrane by contact with a ridge may provide a sensing mechanism of substrate topological changes, possibly through local regulation of SA channels in regions of the apical membrane that are adpressed to a ridge or other topological discontinuity. Ion-channel-mediated changes in membrane voltage or cytoplasmic ion concentration, especially Ca²⁺, may then initiate signal transduction cascades leading to appressorium development or redirection of the axis of...
hyphal growth (Zhou et al., 1991; Gow, 1995). SA putatively calcium-permeable channels have also been implicated in the regulation of hyphal tip growth of Saprolegnia ferax and Neurospora crassa via the control of calcium influx (Garrill et al., 1993; Gow, 1995; Levina et al., 1995). SA ion channels regulated by membrane tension have also been described in a wide variety of animal, plant and bacterial cells and in other fungi (Sackin, 1995; Ding & Pickard, 1993; Martinac et al., 1987; Gustin et al., 1988; Zhou & Kung, 1992). The proposed roles for these channels include transduction of responses to changes in osmotic pressure, regulation of cell volume and growth and the sensation of touch or sound (Sackin, 1995).

Here, the effects of verapamil (a blocker of L-type Ca2+ channels) and gadolinium have been tested on the thigmotropic response of C. albicans hyphae, to investigate the possible roles of ion channels in the transduction of its thigmotropic behaviour. In addition, patch-clamp electrophysiology has been applied to C. albicans for the first time to demonstrate the presence of SA channels in the plasma membrane of both the hyphal and yeast forms of the fungus and to assess whether morphology-specific SA channels exist.

**METHODS**

**Culture maintenance.** Candida albicans strain 3153 was obtained originally from the London Mycological Reference Laboratory, Colindale, now transferred to the Mycological Reference Laboratory, Department of Microbiology, University of Leeds, UK. This is a germ-tube-positive strain that is fully virulent in systemic infections in mice. The strain was maintained on YPD (1%, w/v, yeast extract; 2%, w/v, peptone; 2%, w/v, glucose; Sigma) solidified with 2% (w/v) agar, incubated at 30°C.

**Hyphal thigmotropism and extension rates.** Topographically defined surfaces were used as described by Gow et al. (1994). The polystyrene membranes were a kind gift from Harvey Hoch, Cornell University, and had a ridge height of 1.53 μm and pitch (distance from centre of ridge to centre of next ridge) of 200 μm. Polystyrene replica membranes were pressed, contoured surface down, onto serum agar plates containing 20% (w/v) newborn calf serum (Gibco), solidified with 2% (w/v) agar. The membrane was then lifted off, inverted and placed contour side up on top of a fresh serum agar plate. These membranes are impermeable to nutrients but this procedure provided sufficient media on top of the membrane to support fungal growth. The small amount of medium that was lifted from the agar surface did not appear to influence the response to the topographical contours. In hyphal extension rate experiments, C. albicans was grown on serum agar plates without ridged membranes. For inhibition studies, channel blockers were filter-sterilized prior to incorporation into serum agar (stock inhibitor concentration of 1 mM). An inoculum of C. albicans yeast cells was grown for 16 h at 25 °C in 125 ml liquid YPD medium to obtain stationary phase cells. For both thigmotropism and growth rate experiments, these cells were inoculated onto serum agar plates at approximately 2 x 10⁶ cells ml⁻¹ per plate, and incubated at 37°C to promote hypha formation.

Hyphal growth was recorded by time-lapse video microscopy (total magnification x400) in a constant temperature room at 37°C. An Olympus CK2 inverted microscope was used to calculate hyphal extension rates and angles of hyphal approach to ridges by linking to a BBC master computer with image analysis software (Gray & Morris, 1992). Experiments were recorded on a Panasonic AG-6720 time-lapse video cassette recorder. The angle of approach of the hypha at the point at which it encountered a ridge was determined as in Fig. 1. The numbers of hyphae which reoriented and subsequently grew along the groove or ridge upon encountering the ridge were expressed as a percentage of the total, subjected to arcsin transformation (John & Quenouille, 1977) and plotted against the angle of approach. Hyphal extension rates were calculated over a 6 h growth period.

**Preparation of protoplasts.** The method for protoplast preparation and channel recording was adapted from that described for Saccharomyces cerevisiae (Bertl & Slayman, 1992). To prepare protoplasts from yeast cells, C. albicans was grown overnight at 25°C (shaking at 200 r.p.m.) in Difco Yeast Nitrogen Base (YNB; with amino acids) supplemented with 5 g glucose liter⁻¹. A 0.5 ml sample of cells was centrifuged (1500 g, 4°C, 5 min), resuspended in 0.5 ml of a solution containing 50 mM KH₂PO₄, 5 mM EDTA, 50 mM DTT adjusted to pH 7.2 with KOH and incubated with gentle agitation at 30°C for 30 min. An equal volume of a solution containing 50 mM KH₂PO₄, 40 mM 2-mercaptoethanol, 2.4 M sorbitol, 3 μg chitinase ml⁻¹ (10 units g⁻¹; Sigma), 1.8 μg lyticase ml⁻¹ (200–1000 units mg⁻¹; Sigma) and 12 μl β-glucuronidase ml⁻¹ (10000 units ml⁻¹; Sigma), pH 7.2 with KOH, was added. The cells were incubated for a further 45 min at 30°C with gentle agitation. Protoplasts were harvested by centrifugation (1500 g, 4°C, 5 min) and resuspended in 0.5 ml of a solution containing (in mM) 200 KCl, 10 CaCl₂, 10 MgCl₂, 10 glucose, 345 d-sorbitol and 5 HEPES/Tris, pH 7.2. To prepare protoplasts from hyphal cells, an inoculum of yeast cells was first grown overnight as described before. The cells were washed in distilled water, diluted 1/10 into YNB (final volume 25 ml; adjusted to pH 8.0 with 20 mM Tris/MES) and incubated in a shaking water bath (200 r.p.m.) at 37°C for 4 h. Hyphae were harvested and protoplasts were prepared as before. The final protoplast suspension was filtered through a nylon mesh filter (20 μm porosity) to remove non-digested cells and resuspended as for yeast protoplasts.

**Patch-clamp recordings.** Standard patch-clamp techniques as described by Hamill et al. (1981) were used. Pipettes were fabricated from thin-walled borosilicate glass capillaries (Kimble) and coated with Sylgard (Dow Corning). The pipettes had a resistance of 15–20 MΩ when filled with pipette solution. This consisted of 50 mM KCl and 1 mM MgCl₂, buffered to pH 5.6 with 5 mM MES/Tris for yeast protoplasts or to pH 7.2 with 5 mM HEPES/Tris for hyphal protoplasts. The reference Ag/AgCl half cell was filled with the relevant pipette solution and solidified with 1% (w/v) agar. The bath solution consisted of 200 mM KCl, 10 mM CaCl₂ and 10 mM MgCl₂, buffered to pH 7.2 with 5 mM HEPES/Tris. The osmolarity of both pipette and bath solutions was adjusted to 925 mOsm with d-sorbitol (> 98% purity; Sigma). The patch-clamp amplifier was an I/M PC (List Electronics). Voltage pulse protocols, data acquisition and analysis were performed using CED software (Cambridge Electronic Design). Data were low passed filtered at 1 kHz and sampled at 3 kHz. Pressure applied through the pipette was measured with a pressure monitor (World Precision Instruments). The mean number of open channels as a function of applied pressure was determined by dividing the integral of the total single channel current by the channel unitary current and the length of the voltage pulse.
RESULTS

Growth of C. albicans hyphae on ridged surfaces

Previous work demonstrated that the hyphal cells of C. albicans responded to thigmotropic cues, following grooves, ridges and entering pores of membrane substrates (Sherwood et al., 1992; Gow et al., 1994). It was observed that the angle of approach of a hypha to a ridge was related to whether the hypha reoriented upon making contact with that ridge and this was used to quantitate the thigmotropic response. Hyphae were grown on polystyrene membranes with ridges of 1:53 μm height and 20:0 μm pitch; the percentage of hyphae which changed growth direction to follow the ridge was recorded as a function of the angle of approach (Fig. 1). Greater than 82–90% of the hyphae changed their direction of growth on contact with a ridge when approaching at angles of 0–30°. As the angle increased, the number of hyphae reorienting decreased up to approach angles of 71–90° where virtually no reorientation in growth direction was observed (Fig. 1).

In the presence of either 50 μM verapamil or 50 μM GdCl₃, the percentage of hyphae reorienting on contact with a ridge was reduced (Fig. 1). For example, at approach angles of 41–50° only 4–8% of hyphae reoriented in the presence of verapamil or gadolinium compared to 67% of hyphae in control experiments. The percentage response in the presence of either verapamil or Gd³⁺ was significantly different to control values (by Student’s t-test) at the 0:1% confidence limit in each of the 10° intervals up to and including 70°. Control experiments showed that 50 μM GdCl₃ or 50 μM verapamil did not inhibit hyphal extension rates on solid serum-containing agar. In the absence of inhibitors, the hyphal extension rate was 25:2±5:8 μm h⁻¹ compared to 29:5±11:1 μm h⁻¹ and 27:6±5:4 μm h⁻¹ (mean±sd, n = 12–19) in the presence of 50 μM gadolinium and 50 μM verapamil, respectively. Measurable attenuation of the reorientation effects of hyphae to ridges could also be measured in 10 and 20 μM gadolinium and 10 and 20 μM verapamil, although these were less marked than at the higher concentration (data not shown).

Other channel blockers, for example 50 μM Co²⁺ and 50 μM La³⁺, were also tested but found to reduce the hyphal extension rate significantly in addition to affecting the thigmotropic response.

SA channels in the plasma membrane of C. albicans

Enzymic digestion of the cell wall formed spherical protoplasts, typically of 5 μm diameter. Seals of 10 GΩ were obtained in approximately 50% of attempts with yeast cell protoplasts but seal formation on hyphal cell protoplasts proved to be more difficult. The inside-out (IO) excised patch configuration was readily obtained by the addition of strong positive pressure lasting from a few seconds to several minutes. Whole-cell recording was also achieved but was found to be too fragile to withstand the application of pressure. Hence the outside-out excised patch configuration for testing the effects on single channel activity of channel blockers at the external membrane face was not attained. However, in the longer term it may be possible to develop a pipette perfusion system suitable for use with Candida which would allow drug delivery to this site.

Two types of SA channels were observed in IO patches from yeast protoplasts. An example of SA channel activity from a yeast cell protoplast is shown in Fig. 2. The activity of this channel increased appreciably when pressure was applied (Fig. 2a). The channel had a conductance of 51 pS (Fig. 4b), similar
Fig. 2. SA channel activity from a C. albicans yeast cell protoplast. Pipette and bath solutions were as given in Methods. (a) Recordings from an IO excised patch: left panel, single channel activity without applied pressure (P = 0); right panel, the same patch under +60 mm Hg pressure. The holding voltage was +20 mV; applied voltages (V_m) are indicated on the right of each set of traces. The current level with the channel in a closed state is marked '<'. (b) Current–voltage relationship of the SA single channel activity represented in (a). Current flowing above the voltage axis represents positive charge flow from the cytosolic to the extracellular membrane face. The equilibrium voltage for K⁺ is marked as 'E_K'. Channel conductance (G) was estimated as 54 pS from the regression fitted to channel activity; relative permeability was estimated from the extrapolated voltage intersect. (c) Effect of pressure on activity of the 54 pS channel.

DISCUSSION

Fungal hyphae, including those of C. albicans, have been shown to respond to thigmotropic cues. Previous work with the plant-pathogenic fungus U. appendiculatus suggested that SA channels may be involved in the sensing of substrate topography and induction of appressorium formation (Zhou et al., 1991). In this study, the numbers of C. albicans hyphae responding to a thigmotropic cue of ridges on polystyrene replica membranes were reduced in the presence of an SA channel inhibitor, gadolinium, and an L-type calcium channel blocker, verapamil. This suggests that SA and/or other calcium-permeable channels could be involved in the thigmotropic response, although there remains the possibility of non-specific effects of the inhibitors on whole cells. Verapamil can produce direct block of potassium selective channels (Thomine et al., 1994) and gadolinium can affect ostensibly non-SA channels (Allen & Sanders, 1994). However, verapamil and gadolinium did not inhibit hyphal elongation, even at the highest concentration used, suggesting that their effects were specific to the thigmotropic response.
Ion channels and thigmotropism in *Candida albicans*

The presence of SA channels in the plasma membrane of *C. albicans* was demonstrated by patch-clamp electrophysiology. Although a preliminary characterization, it represents the first single channel recordings from a medically important fungus. Small protoplast size has not precluded the formation of high resistance seals from either yeast or hyphal forms of a clinical isolate and artificially enlarged strains, for example those produced by colchicine treatment (Jibiki *et al.*, 1993), did not need to be used.

The smaller of the two SA channels observed in the yeast form (20-25 pS) bears some resemblance to the YORK channel, a 26 pS outward rectifier of the *Sac. cerevisiae* plasma membrane (Lesage *et al.*, 1996). Although the latter appears to be stretch-insensitive when expressed in *Xenopus laevis* oocytes, the smaller *C. albicans* yeast channel could be a functional homologue. As such the physiological role of this low conductance channel in *C. albicans* yeasts cells may lie primarily in facilitating rapid cellular K+ efflux for osmoregulation. If this channel were retained in the 'hyphal form, its Ca2+ permeability (which was not determined in this study) could be a determinant of its ability to play a role in thigmotropic sensing.

Two weakly cation-selective SA channels of similar conductance were observed in the yeast (54 pS) and hyphal (51 pS) forms; however, as differences were observed in their voltage dependence and response to pressure it cannot be concluded at this point that the 54 pS yeast channel is equivalent to the 51 pS hyphal channel. If this yeast channel proved to be yeast-specific, then it may play a role in turgor responses as it was activated by both positive and negative pressures. In *Sac. cerevisiae*, a non-selective mechanosensitive channel has been suggested to play such a role in osmoregulation (Gustin *et al.*, 1988). However, it is also possible that the observed channel responses to pressure were related to the patch-clamp configuration used. For example, an SA channel characterized in *Schizosaccharomyces pombe* plasma membrane was activated by positive pressure in cell-attached mode but could be activated by either pressure or suction in excised patches (Zhou & Kung, 1992).

The hyphal 51 pS channel is perhaps the most obvious candidate for a thigmotropism signal transducer. However, the hyphal channel was activated by negative pressure, whereas membrane deformation of a hyphal tip on making contact with a ridge would be equivalent to application of positive pressure in the patch-clamp experiments. This channel may yet have a role in thigmotropism if it is considered that the hydrostatic pressure created in *vivo* within the hypha and directed towards the tip is functionally equivalent to the negative pressure applied to the I0 patch. It is possible that in *vivo* the channel is maintained preferentially in an open state and it is its closure on contact of the tip with a solid surface that is relevant to thigmotropic sensing. As yet this remains a speculative explanation with no direct supporting evidence from other tip-growing cell types.

Our pharmacological analysis of the thigmotropic response of *Candida* hyphae suggests that the specific
plasma membrane channels involved in this behaviour may not be essential for hyphal extension. In *N. crassa*, 500 μM Gd³⁺ incompletely inhibited stretch channel activity and only inhibited hyphal extension transiently, indicating that such channels were not essential for tip growth (Levina et al., 1995). However, in the oomycete *Sap. ferax Gd³⁺ dissipated the tip high cytosolic calcium gradient and stopped hyphal growth (Garrill et al., 1993). Although a functional role for SA channel activity in *Candida* hyphal thigmotropism has not been established unequivocally in this study, the ability to form stable IO patches from the plasma membrane will enable further investigation of this hitherto unexplored facet of *Candida* biology. Thorough characterization of *Candida* single channel activity will enable assessment of channel interaction with, for example, the signal transduction cascades and changes in cytoskeletal architecture which are thought to be relevant not only to contact sensing but also to osmotolerance, polarized hyphal growth and the dimorphic switch.

ACKNOWLEDGEMENTS

We thank the following for financial support: (to N. A. R. G.) University of Aberdeen Research Committee (with I. R. Booth), the BBSRC (ROPA grant no. 1/CEL 04556), The Royal Society/Leverhulme Senior fellowship scheme, the Wellcome Trust and Glaxo Group Research; (to J. M. D.) BBSRC (grant no. P04954) and The Royal Society of London. The use of patch-clamp equipment at the University of York (Professor D. Sanders) is gratefully acknowledged as well as useful advice from Natasha Levina.

REFERENCES


Ion channels and thigmotropism in *Candida albicans*


Received 6 August 1997; revised 22 October 1997; accepted 31 October 1997.