Identification and Characterization of Stretch-Activated Ion Channels in Pollen Protoplasts

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Pollen tube growth requires a Ca\(^{2+}\) gradient, with elevated levels of cytosolic Ca\(^{2+}\) at the growing tip. This gradient’s magnitude oscillates with growth oscillation but is always maintained. Ca\(^{2+}\) influx into the growing tip is necessary, and its magnitude also oscillates with growth. It has been widely assumed that stretch-activated Ca\(^{2+}\) channels underlie this influx, but such channels have never been reported in either pollen grains or pollen tubes. We have identified and characterized stretch-activated Ca\(^{2+}\) channels from *Lilium longiflorum* pollen grain and tube tip protoplasts. The channels were localized to a small region of the grain protoplasts associated with the site of tube germination. In addition, we find a stretch-activated K\(^+\) channel as well as a spontaneous K\(^+\) channel distributed over the entire grain surface, but neither was present at the germination site or at the tip. Neither stretch-activated channel was detected in the grain protoplasts unless the grains were left in germination medium for at least 1 h before protoplast preparation. The stretch-activated channels were inhibited by a spider venom that is known to block stretch-activated channels in animal cells, but the spontaneous channel was unaffected by the venom. The venom also stopped pollen tube germination and elongation and blocked Ca\(^{2+}\) entry into the growing tip, suggesting that channel function is necessary for growth.

The growth of lily pollen tubes is a dynamic affair, with the tubes extending at the average rate of about 10 μm min\(^{-1}\). Furthermore, the growth rate is not steady but oscillatory, with a period of 40 to 60 s (Feijó et al., 2001; Robinson and Messerli, 2002). Tip-high gradients of cytosolic Ca\(^{2+}\) are an absolute requirement for pollen tube extension (Rathore et al., 1991; Miller et al., 1992), and it is well established that the Ca\(^{2+}\) gradient oscillates at the same frequency as growth (Holdaway-Clarke et al., 1997; Messerli and Robinson, 1997), although not in the same phase (Messerli et al., 2000). The medium in which lily pollen tubes germinate and grow is quite simple and requires only K\(^+\), Ca\(^{2+}\), a trace of boron, and somewhat acidic pH. No inorganic anions are required (Weisenseel and Jaffe, 1976). Ion-specific self-referencing electrodes have detected the oscillatory uptake of three cations at the tip, Ca\(^{2+}\), K\(^+\), and H\(^+\) (Holdaway-Clarke et al., 1997; Feijó et al., 1999; Messerli et al., 1999), as well as an oscillatory efflux of Cl\(^-\) of a magnitude that exceeds the magnitude of the other fluxes by a factor of 10 or more (Zonia et al., 2002).

The role of Ca\(^{2+}\) influx at the tip has been brought into question by the fact that the rise in cytosolic Ca\(^{2+}\) occurs 7 to 12 s before the measured influx of Ca\(^{2+}\) (for review, see Robinson and Messerli, 2002). This result gave rise to the suggestion that the Ca\(^{2+}\) influx did not enter the cytosol directly but entered intracellular stores or was bound up in the wall (Holdaway-Clarke et al., 1997). Nevertheless, all recent models of the organization of the machinery to maintain the focused elongation at the pollen tube tip assume the existence of mechanically sensitive, stretch-activated (SA) calcium channels (Derksen, 1996; Feijó et al., 2001; Holdaway-Clarke and Hepler, 2003). No such channels have been identified in pollen grains or tubes, while a number of potassium currents and channels have been identified in pollen by electrophysiological and molecular techniques (Obermeyer and Kolb, 1993; Obermeyer and Blatt, 1995; Fan et al., 2001, 2003; Mouline et al., 2002). SA channels have been reported in other tip-growing organisms, including oomycete hyphae (Garrill et al., 1992) and algal rhizoids (Taylor et al., 1996).

Using the whole-cell patch clamp configuration, Griessner and Obermeyer (2003) recently detected both inward and outward potassium currents in the plasma membranes of protoplasts made from *Lilium longiflorum* pollen grains and pollen tubes. In contrast to previous studies on lily pollen grains, they found significant activation of inward potassium currents at holding potentials that were in the range of the measured membrane potentials of lily pollen tubes (−102 mV when bathed in 3 mM K\(^+\); Messerli et al., 1999). The measured influx of K\(^+\) at the lily pollen tube tip is strongly pulsatile, varying in each growth cycle from a basal level of less than 25 pmol cm\(^{-2}\) s\(^{-1}\) to average peaks of 688 pmol cm\(^{-2}\) s\(^{-1}\) or more (Messerli et al., 1999). However, significant variations in membrane potential have not been measured despite efforts to detect them (K. R. Robinson, unpublished data), making it unlikely that changes in K\(^+\) flux are regulated by voltage.

Thus, the factor(s) that regulates oscillatory K\(^+\) influx as well as the other ion fluxes is not known. It is

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possible that K\(^+\) channel activity at the tip is influenced by cytosolic Ca\(^{2+}\); however, the maximum influx of K\(^+\) is delay by nearly 7 s from the maximum of cytosolic Ca\(^{2+}\). Another obvious factor to consider is the tension in the membrane. That tension is likely to oscillate as the rate of elongation oscillates, so mechanical sensitivity of the channels is an attractive possibility. SA Ca\(^{2+}\), K\(^+\), and Cl\(^-\) channels, three channels that would seem to be required at the pollen tube tip, have been found in the plasma membranes of Vicia faba guard cells (Cosgrove and Hedrich, 1991). We have carried out patch clamp studies of outside-out patches from lily pollen protoplasts designed to detect SA channels, and we have characterized both K\(^+\) and Ca\(^{2+}\) SA channels in the grain, but found no Cl\(^-\) channel under any of the conditions that were used. In addition, we found a K\(^+\) channel that was not activated by stretch or voltage. In patches pulled from wall-free tips of tubes, only SA Ca\(^{2+}\) channels were detected.

RESULTS

The formation of seals between the patch pipettes and the plasma membrane usually required the application of suction (5–10 kPa) for 1 to 2 min. As a result of the suction, the membrane deformed into the pipette, and occasionally SA channels were noted in the form of currents that varied with suction. Occasionally, seals formed almost immediately after contact and with little or no suction; the properties of channels isolated in these patches did not differ from those from patches that required more prolonged suction. After the formation of high resistance seals (3–6 GΩ), the patches were analyzed in order to characterize channels with regard to their mechanical sensitivity and conductance. Three distinct channels were detected in the grains, two of which were stretch activated. In order to detect SA channels, a period of incubation in the standard solution was required. The pollen grains were suspended in the standard solution for 10 min, 30 min, or 1 h before the enzymatic digestion of cell wall. SA channels were only found when the pollen grains were suspended in the standard solution for at least 1 h. Membrane at the growing pollen tube tip was exposed as described in “Methods and Materials,” below, and only an SA Ca\(^{2+}\) channel was detected with similar, but not identical, properties to the grain SA Ca\(^{2+}\) channel.

Spontaneous K\(^+\) Channels in Grain Protoplasts

The most abundant channel that we observed, and the only channel seen in the absence of mechanical deformation, was a K\(^+\) channel that was detected when the internal solution contained 150 mM K-glucuronate and 30 mM sorbitol, buffered to pH 7.2 with 5 mM Tris/HEPES, and the external solution was 24 mM K-glucuronate or 1 mM K-glucuronate, and 300 mM sorbitol, buffered to pH 5.6 with HEPES/MES. The channel was present in >90% of about 80 patches that were successfully pulled, regardless of the incubation time of the grain before enzymatic removal of the cell wall. Examples of channel activity are shown in Figure 1A, and these data from 75 successful patches are summarized in the I-V curves in Figure 1B. This channel had a conductance of 67 pS when the external solution contained 24 mM K\(^+\), and 72 pS conductance in 1 mM external K\(^+\). The channel open probability changed little with changes in the holding potential (0.18 at −60 mV and 0.21 at +60 mV). The

![Figure 1](image-url)
reversal potential was about $-40$ mV when the external solution contained 24 mM K$^+$ and shifted to $-102$ mV if the external K$^+$ was 1 mM. The calculated equilibrium potential for K$^+$ in the two external K$^+$ concentration solutions was $-40$ mV and $-120$ mV, assuming that the activity coefficients for K$^+$ were 0.75, 0.97, and 0.96 for 150 mM, 24 mM, and 1 mM K$^+$, respectively.

The effect of pressure application up to 10 kPa and the SA channel blocker Gd$^{3+}$ was tested on the spontaneous K$^+$ channel. Neither treatment affected any aspect of the channel’s activity (Fig. 1A). Even 50 μM Gd$^{3+}$ had no effect on this channel’s properties.

**Stretch-Activated K$^+$ Channels in Grain Protoplast**

If 10 kPa of suction was applied to patches that were pulled from protoplasts made from grains that were incubated for at least 1 h or more, a different K$^+$ channel appeared in about 8% of the patches. Representative sweeps are shown in Figure 2A, and the I-V curve for many such channels is shown in Figure 2B. The average conductance of the channel in 24 mM K$^+$ was 33 pS. The SA K$^+$ channel did not show fatigue after 30 min of pressure application; that is, its activity in patches did not decline as is common with mechanically sensitive channels (Gustin et al., 1988; Ding and Pickard, 1993).

As shown in Figure 2A, no channel activity was found in patches lacking the spontaneous K$^+$ channels unless suction was applied. When 10 μM Gd$^{3+}$ was added to the bath, channel activity ceased (Fig. 2A) after about 10 min. If 50 μM Gd$^{3+}$ was added, channel activity ceased immediately. In addition to its lower conductance and sensitivity to Gd$^{3+}$, the SA K$^+$ channel differed from the spontaneous K$^+$ channel in its average open probability, which was 0.45 at $-80$ mV versus 0.20 at $-80$ mV for the spontaneous K$^+$ channel.

**Stretch-Activated Ca$^{2+}$ Channels in Grain Protoplasts**

Under conditions where K$^+$ currents are suppressed (no K$^+$ and 30 mM Ca-gluconate in the bath), no channel activity was detected in the absence of suction applied to the patch. However, if the patch was pulled from protoplasts made from grains that were hydrated for 1 h or more before enzyme treatment, and if suction was applied to the pipette, SA Ca$^{2+}$ channels were occasionally seen (Fig. 3). These channels had an instantaneous conductance of about 15 pS, and the currents were rapidly blocked by 10 μM Gd$^{3+}$. The average open probability of the channels was 0.28 at $-80$ mV and 10 kPa pressure. Unlike the SA K$^+$ channel, this channel exhibited fatigue; that is, the activity of a channel in a patch declined noticeably after 30 min of pressure application. Data from the first 10 min after pressure application were used for analysis.

The overall chance of detecting an SA Ca$^{2+}$ in a patch was about 2%. It was noticed that the SA Ca$^{2+}$ channels appeared to be localized on the protoplasts. Occasionally, it was possible to return to the same place on a protoplast from which a patch containing an SA Ca$^{2+}$ channel was pulled. In those cases, a second patch from the same place inevitably also contained an SA Ca$^{2+}$ channel. Shortly after the protoplasts are released from the pollen wall, a groove can be seen on the protoplast that is left from the groove in the wall from which the pollen tube will later emerge (see Fig. 4). This groove in the protoplasts quickly disappeared. We found that keeping the protoplasts cold extended the time during which the groove could be seen and that SA channels could be detected in about 97% of the successful patches pulled from this region. SA K$^+$ channels were never found in this region.

The internal (pipette) solution contained 150 mM K$^+$ and the external solution contained 30 mM Ca$^{2+}$. These conditions allow the use of a modified form of the Goldman-Hodgkin-Katz equation to evaluate the
permeability of the channel to Ca\textsuperscript{2+} compared to K\textsuperscript{+} using the equation

\[ \frac{P_{\text{Ca}}}{P_{\text{K}}} = \frac{a_{\text{Ca}}}{a_{\text{K}}} \left( \frac{[\text{K}]}{[\text{Ca}]} \right) e^{E_{\text{rev}} F / RT} \]

where \( a_{\text{K}} \) is the activity coefficient of K\textsuperscript{+} (0.75), \( a_{\text{Ca}} \) is the activity coefficient of Ca\textsuperscript{2+} (0.52), and \( E_{\text{rev}} \) is the estimated reversal potential from Figure 3, which we estimate to be at least +50 mV (Fatt and Ginsborg, 1958; Ding and Pickard, 1993). This equation gives a value for \( P_{\text{Ca}}/P_{\text{K}} \) of 98 under these conditions, indicating that the channel has a high preference for Ca\textsuperscript{2+} over K\textsuperscript{+}.

**Stretch-Activated Ca\textsuperscript{2+} Channels in Protoplasts from Growing Tips**

If the osmolarity of the growth medium is raised, extension of pollen tubes can be temporarily halted and the plasma membrane retreats from the cell wall at the tip due to water loss (plasmolysis). Because the cell wall at the tip is composed of pectin, it can be digested with pectinase, leaving intact the more complex cell wall distal to the tip. The pollen tube recovers turgor pressure and extrudes a protoplast in the presence of pectinase that is suitable for patch clamp analysis (Fig. 4C). We exploited this mechanism to characterize the ion channels at the tip of the growing pollen tube. When outside-out patches were drawn from tip protoplasts, only one type of channel could be detected. It was an SA Ca\textsuperscript{2+} channel with similar properties to the Ca\textsuperscript{2+} channel found in grain protoplasts. The density of the channels in the tip protoplasts was greater than in the grain, and multiple channels were often detected in patches pulled with our standard 3-μm tip, as shown in Figure 5. Also shown in Figure 5 is a record made in the cell-attached mode. No currents were detected in the absence of pressure, but complex pattern of inward current was seen when negative pressure of 10 kPa was applied.

Single channels were further analyzed in outside-out patches made using smaller electrode tips. Figure 6A shows the effects of increasing negative pressure on a channel, and the effects of both negative and positive pressure on the channel open probability are shown in Figure 6B. The channels responded symmetrically to negative and positive pressures in these isolated patch preparations.

The tip SA Ca\textsuperscript{2+} channel differed from the one found in the grain in its conductivity. Figure 4B shows the I-V curves for the two channels, and the tip channel’s conductance at −100 mV is about 50% greater than the grain channel.

**Effects of Spider Venom on Channels and Growth**

One problem in the study of SA channels has been the lack of specific pharmacological blockers for these channels. It has been reported that the crude venom of the spider *Grammostola spatulata*, at a dilution of about 1,000, could block SA channels in outside-out patches from pituitary cells (Chen et al., 1996). More recently, a 35-amino acid peptide from the venom has been identified as the active component (Suchyna et al., 2000). We have tested the crude venom (obtained from Spider Pharm, Yarnell, AZ) on the three channel types from pollen protoplasts and also on the growing tubes themselves. We find that the venom, at a final dilution of 3,000, rapidly blocks both the SA K\textsuperscript{+} channel and the SA Ca\textsuperscript{2+} channel but had no effect on the spontaneous K\textsuperscript{+} channel (Figs. 3A and 7).

The venom, at a final dilution of 3,000, also blocked pollen tube growth. When added directly to the medium in which the tubes were grown, growth stopped as soon as the tubes could be examined (1–2 min). In order to observe the effects on single pollen tubes, the pollen grains were germinated in a thin layer of low temperature gelling agarose covered with medium, and the venom was added to the covering medium. After a delay of a few minutes due to diffusion through the agarose (Messerli and Robinson, 1997), growth stopped (Fig. 8, A–F). Following cessation of growth, the clear zone disappeared. Streaming at the tip was halted but continued in regions of the...
Figure 4. Photographs of protoplasts. Shown in A is a recently isolated protoplast with the groove (arrow) still visible. This groove was the locus of the SA Ca\textsuperscript{2+} channels on the grain protoplasts. Shown in B is the same protoplast as the patch electrode makes contact with the groove region. The protoplast was displaced by the electrode from its position shown in A. The patch pulled from this protoplast contained an SA Ca\textsuperscript{2+} channel. C shows the swelling of a pectinase-treated tube (arrow) from which a patch was successfully pulled.

The tube closer to the grain. Many tubes exposed to venom exhibited plasmolysis at the tip (Fig. 8G). The addition of venom at a final dilution of 4,000 did not affect growth within 30 min.

Effects of Spider Venom on Ca\textsuperscript{2+} Currents

We suspect that the SA Ca\textsuperscript{2+} channels that we detect at the tips of growing pollen tubes are the means by which Ca\textsuperscript{2+} enters the tubes. If so, inhibiting the channels with venom should reduce or abolish the known influx of Ca\textsuperscript{2+} into the tips. Using the self-referencing probe, we tested this inference. We found that the venom rapidly stopped the usual oscillatory Ca\textsuperscript{2+} influx (Fig. 9). However, within about 1 min, a period of substantial Ca\textsuperscript{2+} efflux occurred, followed by a large burst of influx. At that point, streaming in the pollen tube stopped and the flux again reversed direction. Variations on this complex pattern were seen in all six tubes that were treated with venom. In all cases, an initial elimination of normal influx occurred, followed by much larger alternations of influx and efflux. Cessation of streaming near the tip always coincided with the onset of large Ca\textsuperscript{2+} influx.

Lack of Cl\textsuperscript{-} Channels

Using protocols that others have used to detect Cl\textsuperscript{-} channels in guard cells (Cosgrove and Hedrich, 1991), we attempted to identify Cl\textsuperscript{-} channels in pollen protoplasts. In no case were such channels detected, either in the presence or absence of suction or at any holding potential. We conclude that such channels must be exceedingly rare or nonexistent.

DISCUSSION

We have identified three, and possibly four, ion channels in the membrane of pollen protoplasts. The channels differ in their frequency of occurrence, conductivity, sensitivity to pressure, selectivity, and distribution. While we have not carried out detailed analysis of the ion specificity of the channels, it should be recognized that lily pollen tubes germinate and grow in a simple medium, with Ca\textsuperscript{2+} and K\textsuperscript{+} the only cations (except for protons) in the extracellular medium. The membrane potential of pollen tubes is more negative than the equilibrium potential of any of the cations, so currents through open channels will be inward, carried by one of these three ions. Thus, we characterize these channels as K\textsuperscript{+} channels or Ca\textsuperscript{2+} channels. We did evaluate the permeability ratio of the Ca\textsuperscript{2+} channel for Ca\textsuperscript{2+} versus K\textsuperscript{+}, and found that the channel was about two orders of magnitude more permeable to Ca\textsuperscript{2+} than K\textsuperscript{+}.
are a common feature of pollen tube extension. The SA Ca\(^{2+}\) channel is not present in functional form in the recently hydrated pollen grain; it only appears after an hour or more of incubation in the germination medium, shortly before germination would occur. It will be interesting to learn if the SA Ca\(^{2+}\) channel is synthesized and inserted into the membrane or modified to make it functional during that 1-h period. It is unclear if the tip SA Ca\(^{2+}\) channel is a slightly modified form of the grain channel or a completely different entity. The only difference that we noted between the channels from the two locations is in somewhat greater conductivity of the tip channel at more negative holding potentials.

The SA Ca\(^{2+}\) channels are highly localized on the protoplasts. The faint indentation that runs along one part of the newly isolated protoplast persists long enough to permit sampling it for the presence of channels. That region proved to be a rich source of SA Ca\(^{2+}\) channels, while the remainder of the protoplast was devoid of the channels. We think that the indentation marks the region from which the pollen tube would emerge, and, thus, it is likely that the SA Ca\(^{2+}\) channels there would also be found in the tube tip. The SA K\(^{+}\) channels were never found in this region.

There is an important difference in the physical situation in which we have characterized the channel, compared to the situation in the intact pollen grain or tube. We have disrupted any putative links to the cytoskeleton, the extracellular matrix (the cell wall), and the endoplasmic reticulum (Reuzé et al., 1997). The tethering of both extracellular and intracellular domains of SA channels has been shown to affect the mechanical gating of some channels, while others are gated by membrane deformation (for review, see Hamill and Martinac, 2001). We do not know how the behavior of the SA Ca\(^{2+}\) channel (or the SA K\(^{+}\) channel) would be modulated in the normal

![Figure 6](image_url)

**Figure 6.** Pressure sensitivity of the SA Ca\(^{2+}\) channels. A, A single channel from a tip protoplast was subjected to positive pressures of different magnitude. All sweeps were at a holding potential of −100 mV. B. The points show the open probability of tip-derived channels in patches subjected to varying positive and negative pressures. The error bars indicate the se of the mean for five different patches.

While the SA Ca\(^{2+}\) channels and the SA K\(^{+}\) channels differed in a number of regards, the most compelling reason for assuming that they are distinct entities rather than different states of the same channel is their distribution. SA K\(^{+}\) channels were never detected in the groove of the pollen grain protoplasts, and SA Ca\(^{2+}\) channels were never detected except in the groove. Likewise, SA K\(^{+}\) channel activity was never detected in the protoplasts made from growing tips, but that region was densely populated with SA Ca\(^{2+}\) channels.

The identification of an SA Ca\(^{2+}\) channel in the grain and at the tip is of considerable significance. Such a channel has long been hypothesized to exist in pollen tube tips to explain the localized, oscillating entry of Ca\(^{2+}\) there. Extracellular Ca\(^{2+}\) and Ca\(^{2+}\) gradients are essential for lily pollen tube growth (Brewbaker and Kwack, 1963; Rathore et al., 1991; Miller et al., 1992), and blocking the entry of Ca\(^{2+}\) abolishes cytoplasmic Ca\(^{2+}\) oscillations (Messerli and Robinson, 1997). Geitmann and Crepaldi (1998) have shown that lanthanides at low concentrations abolish the growth pulses of Petunia pollen tubes, suggesting that SA channels

![Figure 7](image_url)

**Figure 7.** The effect of spider venom at a dilution of 3,000 on pollen protoplast channels. All sweeps at −100 mV. A, C, and E show channel opening of the SA Ca\(^{2+}\) channel, the SA K\(^{+}\) channel, and the spontaneous K\(^{+}\) channel, respectively. B, D, and F show the effects of venom on these channels. The venom abolishes activity of two SA channels (B and D) but has no effect on the spontaneous channel (F). The current scale bar represents 1 pA for A and B, 2 pA for C and D, and 5 pA for E and F.
membrane environment. The whole cell recordings done at the tip do not address this issue, as the cell wall was eliminated and the cytoskeleton within the tip may have been greatly modified from its normal state in an untreated pollen tube.

A surprising feature of our results is the lack of any K\textsuperscript+ channels in the tip protoplasts. It is known that there is a large, oscillatory influx of K\textsuperscript+ into the growing tip (Messerli et al., 1999), and the K\textsuperscript+ flux is an order of magnitude larger than the Ca\textsuperscript{2+} flux. The electrochemical driving force for K\textsuperscript+ entry is vastly smaller than that for Ca\textsuperscript{2+}, so K\textsuperscript+ channels would be expected to be abundant in order to support the large measured influx. K\textsuperscript+ is a required component of the growth medium; however, K\textsuperscript+ must be rigorously removed from the medium in order to stop growth, and K\textsuperscript+ concentrations as low as 10 \(\mu\)M are sufficient to support normal growth rates (Messerli and Robinson, 2005). It has been suggested previously that perhaps the measured K\textsuperscript+ influx pulses involved cotransport with H\textsuperscript+ rather than movement through an ion channel (Messerli et al., 1999). This suggestion was based on the observation that K\textsuperscript+ pulses and H\textsuperscript+ pulses were spatially and temporally coincident and had similar magnitudes. Our present failure to detect K\textsuperscript+ channels at the growing tip lends support to the notion of H\textsuperscript+ / K\textsuperscript+ cotransport.

The spontaneous K\textsuperscript+ channel was ubiquitously distributed on the protoplasts. It had the largest conductance of the three channels that we detected. Its I-V curve was symmetrical around the zero current point, and it was unaffected by membrane tension, Gd\textsuperscript{3+}, or spider venom. The distribution of K\textsuperscript+ influx over the surface of the pollen tube has not been carefully studied; attention has been focused on the tip. If this channel were widely distributed on the pollen tube, it would lead to steady K\textsuperscript+ uptake. The membrane potential of the pollen tube is somewhat more negative than the K\textsuperscript+ equilibrium potential (Messerli et al., 1999) and does not vary with time. It would be surprising, however, if this channel were completely unregulated, so its behavior in the intact membrane may be quite different.

The finding that spider venom blocked both SA channels gives us confidence that the channels are in fact stretch activated and are distinct from the spontaneous K\textsuperscript+ channel. The effect of the venom on germination and growth emphasizes the importance of the SA channels to pollen tube physiology. The same dilution of the venom that blocks the channels rapidly stops elongation of the tubes and disrupts the normal

**Figure 8.** Effect of spider venom at a dilution of 3,000 on pollen tube growth. A to F. Successive photographs of a pollen tube taken at the indicated time with respect to the time of addition of venom, which was just after the photograph in B was taken. Pollen tubes were embedded in a thin layer of agarose, which is covered with medium, so small molecules take a few minutes to diffuse to the tubes after they are added to the medium. Growth slows within 5 min and ceases by 10 min after venom addition, and some swelling of the tip is evident. Scale bar for A to F = 50 \(\mu\)m. G. A different tube at higher magnification, 30 min after venom addition. The plasmolysis at the tip and the complete loss of the clear zone are typical results of treatment with venom.

**Figure 9.** The effect of spider venom on Ca\textsuperscript{2+} fluxes. The Seris Ca\textsuperscript{2+} probe was used to measure Ca\textsuperscript{2+} fluxes at the growing pollen tube tip. Initially, the usual oscillatory influx was detected. Upon adding venom, the flux rapidly declined and then reversed direction, becoming outward briefly. About 3 min after venom addition, massive Ca\textsuperscript{2+} influx occurred and with it, the cessation of streaming near the tip. Note that the pollen tube was not embedded in agarose, as in Figure 8, so the venom reached to tube tip immediately as a result of gentle mixing.
Ca\(^{2+}\) influx. The effects of venom on Ca\(^{2+}\) influx were complicated. As expected from the effect on isolated channels, the immediate effect of venom was to reduce the amplitude of the normal Ca\(^{2+}\) influx oscillations. However, within a few minutes, Ca\(^{2+}\) efflux occurred, followed by massive influx that stopped cytoplasmic streaming. We do not understand the mechanism that underlies these later events, which presumably are a pathological response to the initial disruption of normal Ca\(^{2+}\) influx.

Finally, we failed to detect any evidence of anion channels, stretch activated or otherwise. One possibility that we did not explore is Ca\(^{2+}\) regulation of an anion channel. Zonia et al. (2002) have reported massive oscillatory Cl\(^{-}\) efflux at the pollen tube tip. Their measured fluxes were an order of magnitude larger than any other measured fluxes, thus the channel underlying the fluxes would be expected to be both abundant and high conductance. The Cl\(^{-}\) fluxes reported by Zonia et al. (2002) were disrupted by the injection of Ins(3,4,5,6)P\(_4\), so perhaps our patch preparation lacked some critical aspect of inositol phosphate control. It should be noted that others have found that lily pollen germination and growth are independent of extracellular Cl\(^{-}\) (Weisenseel and Jaffe, 1976).

MATERIALS AND METHODS

Protoplast Isolation

Anthers from *Lilium longiflorum* flowers were dried at room temperature for 2 d, and pollen grains were then separated from the anthers by shaking them in a sieve. The grains were then stored in small vials at \(-20^\circ\text{C}\). Protoplasts were prepared following the method described by Fan et al. (1999). In order to prepare protoplasts, the pollen grains were hydrated in a standard solution containing 1 m\(\text{M}\) KNO\(_3\), 0.2 m\(\text{M}\) KH\(_2\)PO\(_4\), 3 m\(\text{M}\) MgSO\(_4\), 1 \(\mu\)m KI, 0.1 \(\mu\)m CuSO\(_4\), 5 \(\mu\)m CaCl\(_2\), 5 \(\mu\)M MES (pH 5.8 adjusted with Tris), 500 \(\mu\)M Glc, and 500 \(\mu\)M sorbitol (osmolality = 1.5 Osmol kg\(^{-1}\)). The hydrated grains were then filtered through 80-\(\mu\)m nylon mesh and centrifuged at 160g for 5 min. An enzyme solution was prepared by adding to the standard solution 2% (w/v) cellulase, 1% (w/v) macerozyme R-10 (Yokult Honsha, Tokyo), and 0.2% potassium dextran sulfate, and the pelleted pollen grains were suspended in it. The protoplasts were again centrifuged at 160g for 5 min, and the pellet was resuspended with 2 mL of the standard solution. This centrifugation/resuspension cycle was repeated three times, and the washed protoplasts were stored on ice until use.

Preparation of Tip Protoplasts

Pollen grains were incubated in modified Dickenson’s medium (Messerli et al., 2000) and allowed to germinate. Pollen tubes were plasmolyzed by increasing the osmolality of the bathing medium and then were treated with an enzyme solution containing only 2% pectinase (Sigma, St. Louis). As the tubes recovered turgor, a protoplast was extruded from the tip from which patches could be pulled. Before patching, the enzyme solution was replaced with the desired patch clamp solution.

Patch Clamp Methods

Patch pipettes were fabricated from Kimax 51 capillaries 1.5 to 1.8 mm o.d. and 100 mm long using two-step pulling with Narishige vertical puller, and the pipettes were fire polished. The pipettes were backfilled with desired solution, and seals were formed between the pollen plasma membrane and the patch pipettes, and outside-out patches pulled as described by Hamill et al. (1981). The Axopatch-1D amplifier (Axon Instruments, Union City, CA), Digidata 1200, and programs Clampex 8.0 and Clampfit 8.0 were used for recording and analysis of the patch clamp data. After successful patches were pulled, single channel recordings were carried out at membrane potentials from 100 mV to \(-100\) mV (\(-160\) mV in some cases) at the intervals of every 20 mV. The sweep lengths were either 2,000 or 2,500 ms. The records were filtered at 3 kHz.

The experiments were designed to characterize three different channels: Cl\(^{-}\), K\(^{+}\), and Ca\(^{2+}\). The outside-out patches were analyzed with three different sets of solutions. For Cl\(^{-}\) channels, the internal solution contained 150 m\(\text{M}\) KCl, 2 m\(\text{M}\) MgCl\(_2\), 2 m\(\text{M}\) Mg-ATP, 1 m\(\text{M}\) EGTA buffered to pH 7.2 with 10 m\(\text{M}\) Tris/HEPES, and external solution consisted of 40 m\(\text{M}\) CaCl\(_2\), 2 m\(\text{M}\) MgCl\(_2\), and 220 m\(\text{M}\) sorbitol buffered to pH 5.5 with 10 m\(\text{M}\) MES/Tris. For K\(^{+}\) channels, internal solution contained 150 m\(\text{M}\) K-gluconate and 30 m\(\text{M}\) sorbitol, buffered to pH 7.2 with 4 m\(\text{M}\) Tris/HEPES, and external solution was 24 m\(\text{M}\) K-gluconate and 300 m\(\text{M}\) sorbitol buffered to pH 5.6 with 2 m\(\text{M}\) HEPES/MES. For Ca\(^{2+}\) channels, the internal solution contained 150 m\(\text{M}\) K-gluconate, 30 m\(\text{M}\) sorbitol, buffered to pH 7.2 with HEPES/MES, and external solution contained 30 m\(\text{M}\) Ca-gluconate and 300 m\(\text{M}\) sorbitol buffered to pH 5.6 with 2 m\(\text{M}\) HEPES/MES (Cosgrove and Hedrich, 1991). Pressure, both positive and negative, was applied to patches by activating a solenoid valve that connected the patch electrode to a gas-tight syringe equipped with a screw drive attached to the plunger. The pressure generated in the system was measured continuously by a digital manometer and was stable for the duration of each recording.

Solutions containing Gd\(^{3+}\) were prepared shortly before use by dissolving GdCl\(_3\) directly in the appropriate media. Spider venom was reconstituted from lyophilized solid and stored in small aliquots at \(-70^\circ\text{C}\). The venom was used within 2 weeks of reconstitution.

Ca\(^{2+}\) Flux Measurements

Ca\(^{2+}\) fluxes were measured using the self-referencing ion selective (Seris) probe (Smith et al., 1994) as modified for pollen tip measurements (Messerli et al., 1999). Pollen tubes were germinated in modified Dickenson’s medium and then transferred to plastic petri dishes that were treated with poly-L-lys (500 \(\mu\)g/mL) solution for 30 min and then rinsed. This treatment caused the pollen tubes to stick slightly to the dish bottom, which facilitated flux measurements.

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LITERATURE CITED


1405

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