Two-electrode Voltage-clamp Recordings in *Xenopus laevis* Oocytes: Reconstitution of Abscisic Acid Activation of SLAC1 Anion Channel via PYL9 ABA Receptor
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[Abstract] Two-Electrode Voltage-Clamp (TEVC) recording in *Xenopus laevis* oocytes provides a powerful method to investigate the functions and regulation of ion channel proteins. This approach provides a well-known tool to characterize ion channels or transporters expressed in *Xenopus laevis* oocytes. The plasma membrane of the oocyte is impaled by two microelectrodes, one for voltage sensing and the other one for current injection. Here we list a protocol that allows robust reconstitution of multi-component signaling pathways. This protocol has been used to study plant ion channels, including the SLAC1 channel (SLOW ANION CHANNEL-ASSOCIATED 1), in particular SLAC1 activation by either the protein kinase OST1 (OPEN STOMATA 1), Ca²⁺-dependent protein kinases (CPKs) or the GHR1 (GUARD CELL HYDROGEN PEROXIDE-RESISTANT 1) transmembrane receptor-like protein. Data are presented showing reconstitution of abscisic acid activation of the SLAC1 anion channel by the ‘monomeric’ ABA (abscisic acid) receptor RCAR1/PYL9 (PYRABACTIN RESISTANCE1 [PYR]/PYR1-LIKE [PYL]/REGULATORYCOMPONENTS OF ABA RECEPTORS [RCAR]) by co-expressing four components of the abscisic acid signaling core. This protocol is also suitable for studying other ion channel functions and regulation mechanisms, as well as transporter proteins.

**Keywords:** Ion channel, Voltage-clamp, Oocytes, SLAC1, ABA receptor, Slow-type Anion Channel

[Background] Ion channels expressed in *Xenopus laevis* oocytes can be studied using two-electrode voltage-clamping. This protocol provides a method to measure ion channel or transporter currents expressed in oocytes, including plant ion channels. In this protocol, we not only summarize how to prepare cRNA, isolate oocytes, inject cRNA and record currents, but also provide information on how to succeed in completing experiments upon co-expressing a signal transduction cascade from receptor to ion channel.

**Materials and Reagents**

1. Borosilicate glass capillaries (World Precision Instruments, catalog number: 1B100F-4)
2. Parafilm (Sigma-Aldrich, catalog number: P7793-1EA)
3. *Xenopus laevis* oocytes (Ecocyte Bioscience, catalog number: 0-100-2)
4. Vector: pNB1 oocyte expression vector harboring the cDNA of interest using the USER method (Nour-Eldin *et al.*, 2006), or other oocytes expression vector like
5. mMESSAGE mMACHINE® T7 Kit (Thermo Fisher Scientific, Ambion™, catalog number: AM1344)
6. Collagenase D (Roche Diagnostics, catalog number: 11088882001)
7. Mineral oil (Sigma-Aldrich, catalog number: M5904)
8. MES hydrate (Sigma-Aldrich, catalog number: M2933)
10. Calcium chloride (CaCl₂) (Sigma-Aldrich, catalog number: C5670)
11. Magnesium chloride (MgCl₂) (Sigma-Aldrich, catalog number: M8266)
13. Potassium chloride (KCl) (Sigma-Aldrich, catalog number: P9333)
14. Na-gluconate (Sigma-Aldrich, catalog number: S2054)
15. D-sorbitol (Sigma-Aldrich, catalog number: S1876)
16. Gentamicin solution (Sigma-Aldrich, catalog number: G1272)
17. ND96 buffer (see Recipes)
18. Recording buffer (see Recipes)

**Equipment**

1. Two-electrode voltage clamp amplifier (*e.g.*, Warner Instrument, model: Oocyte Clamp OC-725C)
2. Digidata 1440A low-noise data acquisition system (Molecular Devices, model: Digidata 1440A)
3. P-87 flaming/brown microelectrode micropipette puller (Sutter Instrument, model: P-87)
4. Osmometer (*e.g.*, Wescor, model: Vapor Pressure Osmometer 5500)
5. Microdispenser (Drummond Scientific, catalog number: 3-000-510)
6. Custom glass tubing (Drummond Scientific, catalog number: 3-000-210-G8)

**Software**

1. pCLAMP 10 Electrophysiology Data Acquisition and Analysis Software (Molecular Devices).

**Procedure**

A. Prepare cRNA

1. All constructs were cloned into the pNB1 oocyte expression vector using the USER method (Nour-Eldin *et al.*, 2006).
2. cRNAs were synthesized from 0.5-1 μg of linearized plasmid DNA template using the
mMESSAGE mMACHINE® T7 Kit from Thermo Fisher Scientific.

B. Isolation of oocytes
1. Individual *Xenopus laevis* oocytes can be ordered from Ecocyte Bioscience and arrives the next day on dry ice. Alternatively, ovary lobes can be surgically extracted as described in previous reports (Miledi, 1982; Gundersen *et al.*, 1983; Stühmer and Parekh, 1995).
2. To isolate oocytes from ovary lobes, wash ovary lobes with ND96 buffer 3 times, incubate it in ND96 buffer at 16 °C overnight.
3. Place 2-3 ovary lobes in ND96 buffer containing 30 mg/ml collagenase D, shake 2-3 h to remove follicle cell layer at room temperature (~23 °C) (Figure 1A).
4. Wash oocytes with ND96 buffer 5 times, sort stage IV and V oocytes (approximate diameter 1 mm, Wasserman *et al.*, 1984) in ND96 buffer. Incubate sorted oocytes in ND96 buffer at 16 °C overnight before injecting cRNA (Figure 1B).

C. Injection of cRNA
1. Use stage IV and V oocytes, for which the follicle cell layer has been removed (Wasserman *et al.*, 1984). Inject more than 30 oocytes with each mRNA combination that is to be investigated.
2. Pull injection glass needles on P-87 flaming/brown microelectrode micropipette puller using borosilicate glass capillaries to produce injection needles (Figure 1C).
3. Break off the needle tip with forceps so that it is easier to inject cRNA into oocytes (Figure 1D).
4. Fill pipette tip with mineral oil to about one-third to two-thirds.
5. Mount pipette to the microdispenser.
6. Deposit cRNA sample onto Parafilm, to a total volume of cRNA > 1 μl.
7. Fill the pipette with RNA solution with microdispenser by applying negative pressure to the pipette.
8. Inject cRNA into selected oocytes. The volume of cRNA injected into oocytes is about 50 nl (oocyte volume ~500 nl). The concentration of injected cRNA is better more than 2 ng/μl (Figure 1E).
9. Incubate oocytes in ND96 buffer at 16 °C for 2-3 days before recording currents.
Figure 1. Oocytes and glass needle at different steps. A. Oocyte ovary lobes; B. Oocytes after isolation; C. Injection glass needle before the tip is 'broken'; D. 'Broken' needle; E. An oocyte being injected with cRNA. Note that injection pipette is out of focus. F. An oocyte impaled with two electrodes for voltage clamping.

D. Recording
1. Pull recording glass electrodes on P-87 flaming/brown microelectrode micropipette puller using custom glass tubing.
2. In some case, slightly break the needle tip so that it is easier to insert the micropipette into oocytes since needle tips can be too soft for multiple sequential injections into oocytes. Alternatively, press the electrode against the oocyte and tap the recording table. Vibrations aid in impaling the oocyte.
3. Fill the micropipette with 3 M KCl. Place one electrode into each of the two holders, making sure that Ag/AgCl electrode wire contacts the KCl solution in the micropipette. The resistances of the filled electrodes were 0.5-1.5 M (Figure 1F).
4. For anion channel recordings, steady state currents are recorded starting from a holding potential of 0 mV and ranging from +40 to -160 mV in -20 mV decrements, followed by a -120 mV voltage 'tail' pulse (Figure 2).

Note: The time-dependent properties of SLAC1 channel currents in Xenopus laevis oocytes vary among individual oocytes. This may depend on posttranslational modification of the channel protein. This property is also known from guard cell recordings of the SLAC1-encoding S-type anion channels (Schmidt and Schroeder, 1994).
Figure 2. **OST1 activates SLAC1 anion channel currents in *Xenopus laevis* oocytes.**

A. The voltage protocol used for recording SLAC1 anion channel currents. B and C. Example of whole-cell current traces recorded from oocytes injected with cRNA of (B) SLAC1yc only and (C) SLAC1yc+OST1yn (in SLAC1yc, SLAC1 is tagged with the C-terminal ‘half’ of YFP and in OST1yn, OST1 is tagged with N-terminal ‘half’ of YFP. Constructs for similar experiments see Geiger *et al.*, 2009; Lee *et al.*, 2009; Hua *et al.*, 2012; Brandt *et al.*, 2015; Wang *et al.*, 2016).

5. To assess whether ‘monomeric’ abscisic acid ABA receptor PYL9 (Dupeux *et al.*, 2011; Hao *et al.*, 2011) activation of SLAC1 can be reconstituted, we co-injected cRNAs of the SLAC1yc channel, ABI1 (ABA-INSENSITIVE1) protein phosphatase, the PYL9 receptor, and the OST1yn protein kinase into oocytes (Geiger *et al.*, 2009; Brandt *et al.*, 2012). Without ABA, PYL9 did not activate the anion channel currents in oocytes expressing the ABI1 with SLAC1yc and OST1yn. When ABA was injected into oocytes 30 min before measuring of oocyte currents, SLAC1 anion channels activity was dramatically enhanced in oocytes expressing SLAC1yc, OST1yn, PYL9 and ABI1 (Figure 3). Thus, the ‘monomeric’ ABA receptor PYL9 enables ABA signaling reconstitution in oocytes.
Figure 3. Reconstitution of ABA activation of SLAC1 anion channels by PYL9 ABA receptor. Average current of SLAC1 anion channels recorded at -140 mV. In the absence of ABA, the ABA receptor PYL9 is unable to enhance SLAC1 anion channels currents. However, in the presence of injected ABA, SLAC1 anion channels currents are greatly increased. Data are mean ± SEM (SLACyc, n = 9; SLAC1yc+OST1yn, n = 13; SLAC1yc+OST1yn+ABI1, n = 10; SLAC1yc+OST1yn+ABI1+PYL9, n = 12; SLAC1yc+OST1yn+ABI1+PYL9+ABA, n = 15).

Recipes

1. ND96 buffer
   10 mM MES/Tris (pH 7.5)
   1 mM CaCl₂
   1 mM MgCl₂
   96 mM NaCl
   Osmolality is adjusted to 220 mM using D-sorbitol (Geiger et al., 2009; Wang et al., 2016)

2. Recording buffer
   10 mM MES/Tris (pH 7.5)
   1 mM MgCl₂
   1 mM CaCl₂
   2 mM KCl
   24 mM NaCl
   70 mM Na-gluconate
   Osmolality is adjusted to 220 mM using D-sorbitol (Geiger et al., 2009; Wang et al., 2016)

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References


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