Combining calcium imaging with other optical techniques.
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ABSTRACT
Ca\textsuperscript{2+} imaging is a commonly used approach to measure Ca\textsuperscript{2+} signals at high spatial resolution. The method is often combined with electrode recordings to correlate electrical and chemical signals or to investigate Ca\textsuperscript{2+} signals following an electrical stimulation. To obtain information on electrical activity at the same spatial resolution, Ca\textsuperscript{2+} imaging must be combined with membrane potential imaging. Similarly, stimulation of subcellular compartments requires photostimulation. Thus, combining Ca\textsuperscript{2+} imaging with an additional optical technique facilitates the study of a number of physiological questions. The aim of this chapter is to introduce some basic principles on how to combine Ca\textsuperscript{2+} imaging with other optical techniques. We will focus on: 1. the design of the optics; 2. the design of experimental protocols; 3. the optical characteristics of the Ca\textsuperscript{2+} indicators and of the additional optical probe used in combination with the Ca\textsuperscript{2+} indicator; 4 the affinity of the Ca\textsuperscript{2+} indicator in relation to the type of measurement. This chapter will enable the reader to devise an optimal strategy for combined optical experiments.
INTRODUCTION

Ca^{2+} imaging is a very powerful approach allowing measurement of the intracellular free Ca^{2+} concentration change (Δ[Ca^{2+}]) or of the Ca^{2+} influx (I_{ca}) simultaneously from multiple sites, including different cells in a network or different compartments within a cell. Using confocal microscopy or two-photon imaging, the improved spatial resolution permits recordings from sub-micron structures such as axonal terminals or dendritic spines. Whereas the principles of Ca^{2+} imaging and of related microscopy techniques are described in other chapters, here we address the topic of combining Ca^{2+} imaging with additional optical techniques, i.e. how to image simultaneously another important physiological variable or how to measure a Ca^{2+} signal following an optical stimulation.

Correlating a Ca^{2+} signal with another variable of biological interest at the same spatial resolution is often necessary to unambiguously answer a specific question. The example that we address here is the combination of Ca^{2+} imaging using calcium indicators with membrane potential (V_m) imaging using voltage-sensitive dyes. Several types of Ca^{2+} signals are directly correlated with V_m changes and the simultaneous recording of the two signals from same locations may facilitate the interpretation of results. I_{ca} initiated by depolarization may also contribute to the further change in V_m when it originates from Ca^{2+} channels in the plasma membrane, such as voltage-gated Ca^{2+} channels (VGCCs), glutamate receptors (in particular NMDA receptors) and other Ca^{2+} permeable pores. In this case, the influx of Ca^{2+} cations represents a depolarizing current. In contrast to I_{ca} through calcium channels in the plasma membrane, Ca^{2+} signals are supposedly not directly correlated with V_m when they are due exclusively to Ca^{2+} flux through internal membranes, such as those of the endoplasmic reticulum. In this case, Ca^{2+} flux does not contribute to the current underlying ΔV_m. Sometimes the source of Ca^{2+} signal can be determined by using appropriate pharmacological tools. More often, however, Ca^{2+} signals originate from multiple sources and the combined Ca^{2+} and V_m imaging is the ideal experimental tool to determine the composition of Ca^{2+} signals. In the subheading dedicated to this technique, we will give some examples illustrating this concept.

Ca^{2+} imaging can also be combined with photostimulation. Photostimulation may be necessary in experiments where physiological stimulation does not allow the separation and the study of different components of Ca^{2+}. An example that we will describe below is the investigation of Ca^{2+} signals associated with activation of glutamate receptors using L-glutamate photorelease from caged compounds. In this type of study, the different Ca^{2+} components associated with synaptic signals (I_{ca} from AMPA or NMDA receptors and Ca^{2+} signals associated with activation of metabotropic glutamate receptors) cannot be easily separated from Ca^{2+} signals from VGCCs using synaptic stimulation, because pharmacological manipulation of VGCCs also affects presynaptic function. In a separate
section, we will discuss how combining Ca\(^{2+}\) imaging with L-glutamate uncaging can be used to investigate Ca\(^{2+}\) signaling associated with activation of metabotropic receptors.

**MICROSCOPY PRINCIPLES OF COMBINING TWO OPTICAL TECHNIQUES**

Combining optical techniques for functional studies requires particular attention to the design of the experiment in order to make the measurement possible and to facilitate the interpretation of the results. The design of the experiment starts from the optical properties of the two molecules that are utilized. In fluorescence measurements, the relevant properties of optical probes include the absorption and the emission spectra of the indicators and the absorption (or excitation) spectrum of the caged compound. For a “dual” imaging experiment, the two indicators can be excited at the same wavelength if the emission spectra of the two dyes are different permitting separation and independent acquisition of signals with two detectors. This arrangement, however, has the disadvantage that both indicators must always be excited at same intensity and for the same duration, a condition that is rarely ideal. Another possibility is to alternate two excitation wavelengths and to use only one detector utilizing dyes that have different absorption spectra but largely overlapping emission. Using this protocol, however, a strictly simultaneous imaging cannot be achieved. The ideal condition is met by using two indicators having separate absorption and emission spectra. In this case, it is possible to design the experiment to permit simultaneous illumination at two separate wavelengths and independent acquisition of the two fluorescence signals (Fig. 1A). The simultaneous \(V_m\) and Ca\(^{2+}\) imaging can be achieved by using Fura Ca\(^{2+}\) indicators, that can be excited by UV light and that emit in the blue/green region, together with commercial styryl voltage-sensitive dyes (for example di-4-ANEPPS, di-8-ANEPPS and JPW-1114) that are excited with green light and that emit in the red/IR region (Canepari et al. 2007; Canepari et al. 2008; Canepari and Vogt 2008; Vogt et al. 2011). The same principle of dual illumination also applies to combine Ca\(^{2+}\) imaging with photostimulation. In uncaging experiments, UV light is normally used for photorelease while blue light is typically inert to caged compounds. Thus, blue-excitable Ca\(^{2+}\) indicators such as Fluo, Oregon Green or Calcium Green dyes are ideal for this application. For dual illumination, two light sources can be directed to the epifluorescence port of the microscope using a dichroic mirror (Fig. 1B) and reflected to the preparation. This configuration is not ideal to obtain whole-field illumination with lasers (Canepari et al. 2010). In this case, illumination can be done directly from above the objective (Fig. 1C) and a multiple-band dichroic mirror should be chosen if the transmission of two separate fluorescence bands is necessary (Vogt et al. 2011). Details on the specific configuration for simultaneous \(V_m\) and Ca\(^{2+}\) imaging are given in Protocol 1.
APPLICATION 1: COMBINING $\text{C}^{2+}$ AND $V_m$ OPTICAL MEASUREMENTS

A minimal overlap in the absorption and emission, meeting the ideal condition for combining $\text{C}^{2+}$ and $V_m$ optical measurements, is obtained with styryl voltage-sensitive dyes (e.g. di-8-ANEPPS or JPW1114) and with Fura based $\text{Ca}^{2+}$ indicators (Fig. 1D). It is only recently that it became possible to image $\text{Ca}^{2+}$ and $V_m$ simultaneously (Vogt et al. 2011, see also Protocol 1). Thus, sequential $\text{Ca}^{2+}$ and $V_m$ imaging was utilized in several studies where signals were proven to be consistent in different trials (Canepari et al. 2007; Milojkovic et al. 2007; Berger et al. 2007; Canepari and Vogt 2008). In this type of experiment, the choice of the $\text{Ca}^{2+}$ indicator is fundamental. Fura indicators are available with different dissociation constants ($K_d$). The $K_d$ and the concentration of the indicator determine whether a transient $\text{Ca}^{2+}$ in the cytosol binds mostly to the dye or to endogenous buffer of the cell (Canepari et al. 2008). In the first case, which is in most cases met with high-affinity $\text{Ca}^{2+}$ indicators ($K_d < 1 \mu\text{M}$) such as Fura-2 or
Bis-Fura-2 at concentrations >300 µM, the Ca\(^{2+}\) bound to the dye is nearly the total Ca\(^{2+}\). Thus, it is possible to correlate $\Delta V_m$ with the cumulative total cytosolic Ca\(^{2+}\) concentration which is proportional to the fractional change of fluorescence (i.e. the ratio between the change of fluorescence and the initial fluorescence). However, because the free Ca\(^{2+}\) is tightly buffered by the indicator, physiologically important downstream signaling and Ca-dependent processes that regulate Ca influx, release and uptake are prevented from occurring. In the second case, which requires the use of low-affinity Ca\(^{2+}\) indicators (Kd > 5 µM) such as Fura-FF or Mag-Fura-2, the Ca\(^{2+}\) bound to the dye has minimal impact on the endogenous cell reactions. Thus, it is possible to correlate $\Delta V_m$ with the physiological $\Delta [\text{Ca}^{2+}]_i$, proportional to the change of Ca\(^{2+}\) indicator fluorescence normalized to the indicator fluorescence at resting [Ca\(^{2+}\)]. Fura indicators allow ratiometric measurements in which fluorescence increases or decreases with Ca\(^{2+}\) depending on whether the indicator is excited below or above its isosbestic wavelength (~360 nm). Imaging above the isosbestic wavelength has several advantages. First, the resting fluorescence, which corresponds to nominally zero Ca\(^{2+}\), is the maximal fluorescence. Second, the contrast of healthy cells is high because they have low resting Ca\(^{2+}\) and the fluorescence of the extracellular dye is quenched. Third, since the dynamic range is ~1, after subtraction of fluorescence at saturating Ca\(^{2+}\), these measurements allow better quantitative estimates of Ca\(^{2+}\) signals (Konishi et al., 1991; Ogden et al., 1995). The comparison of the ratios between bound Ca\(^{2+}\) and free Ca\(^{2+}\) (buffering capacity), a parameter that can be estimated both for the indicator ($k_{\text{dye}}$) and for the cell ($k_{\text{cell}}$), is the key factor to determine in which regime a Ca\(^{2+}\) imaging experiment is performed. For the indicator, $k_{\text{dye}}$ is given by the ratio between its concentration and its K\(_d\). For the cell, $k_{\text{cell}}$ can be measured using specific imaging protocols (see for example Neher and Augustine 1992; Zhou and Neher 1993; Lips and Keller 1998; Palecek et al. 1999).

The choice of $k_{\text{dye}}$>>$k_{\text{cell}}$ is advisable when the aim of the study is to correlate $\Delta V_m$ with $I_{\text{Ca}}$ via channels located in the plasma membrane. An example of this type of experiment is the measurement of Ca\(^{2+}\) signals associated with induction of long-term potentiation of excitatory synapses CA1 hippocampal pyramidal neurons (Canepari et al. 2007). At these synapses, back-propagating action potentials (bAPs) activate VGCCs whereas excitatory postsynaptic potentials (EPSPs) produce $I_{\text{Ca}}$ via Ca\(^{2+}\) permeable ionotropic glutamate receptors, mainly NMDA receptors. When EPSPs are paired with bAPs, depolarization from bAPs unblocks NMDA receptors from Mg\(^{2+}\) increasing $I_{\text{Ca}}$ through these channels, whereas depolarization from EPSPs may inactivate A-type K\(^+\) channels boosting the depolarization (Magee and Johnston 1997). The result is a potentiated $I_{\text{Ca}}$ correlated with an increased or more prolonged depolarization, exclusively at the site where EPSPs originate. Here we use the experiment to investigate this phenomenon as an example of recordings with $k_{\text{dye}}$>>$k_{\text{cell}}$. In Fig. 2A-C, a CA1 hippocampal pyramidal neuron (Fig. 2A) was loaded with the voltage-sensitive dye JPW-3028 and
300 µM of the Ca\(^{2+}\) indicator Bis-Fura-2 (\(K_d \sim 500\) nM). In this type of neuron \(k_{cell}\) has been estimated to be \(\sim 100\) in the dendrites and \(\sim 20\) in the spines (Sabatini et al. 2002). Thus, the condition \(k_{dye} >> k_{cell}\) is valid both in the dendrite and in the spines. The analysis of this experiment concentrates on \(\Delta V_m\) and Ca\(^{2+}\) signals from two sample dendritic regions (a and b). One region (a) is recognized as the site of synaptic inputs since Ca\(^{2+}\) signals are observed with EPSPs evoked by adjacent extracellular stimulation (data not shown). The other region (b), far from the origin of EPSPs, is used as control site. The action potential initiated in the soma propagates back in the dendrites (Fig. 2B, top) producing a small I\(_{Ca}\) in both a and b. The amplitude of the action potential was boosted by pairing with a burst of 5 EPSPs in a, where also the associated Ca\(^{2+}\) signal was strongly enhanced (Fig. 2B, bottom). The same phenomenon was not observed in the control region b. Thus, as expected, a larger I\(_{Ca}\) was correlated with a larger depolarization (Fig. 2C).

The relationship \(k_{cell} >> k_{dye}\) must be satisfied when the aim of the experiment is to correlate \(\Delta V_m\) with \(\Delta [Ca^{2+}]_i\) under physiological calcium homeostasis. Indeed, if an investigated phenomenon depends on the intact dynamics of Ca\(^{2+}\) which is naturally regulated by Ca\(^{2+}\)-binding proteins, a significant buffering by the indicator will produce artificial results. As example of this scenario we use another interesting phenomenon; also related to synaptic plasticity, occurring at cerebellar parallel fibre (PF) – Purkinje neuron (PN) synapses. PF-PN EPSPs can evoke local Ca\(^{2+}\) spikes and high-frequency (50-200 Hz) of EPSPs can evoke bursts of local Ca\(^{2+}\) spikes (Canepari and Vogt 2008). \(\Delta [Ca^{2+}]_i\) signals associated with Ca\(^{2+}\) spikes, however, progressively increase during the burst producing a phenomenon of non-linear Ca\(^{2+}\) summation. The mechanism underlying this increase is the local and transient saturation of the endogenous buffer which is not correlated with size of the spikes. In Fig. 2D-E, a PN was loaded with the voltage sensitive dye JPW-1114 and with 1 mM of the Ca\(^{2+}\) indicator Fura-FF (\(K_d \sim 10\) µM). Because dendritic \(k_{cell}\) in the PN is estimated to be \(\sim 2000\) (Fierro and Llano 1996), the condition of \(k_{cell} >> k_{dye}\) is valid in this experiment. Optical signals related to membrane potential transients were also calibrated on an absolute scale (in mV) using a prolonged non-attenuating hyperpolarization (Roth and Häusser 2001) as described in Canepari at al. (2008). In the recording of Fig. 2D, a burst of seven PF-EPSPs at 100 Hz produced Ca\(^{2+}\) spikes of the same size, but \(\Delta [Ca^{2+}]_i\) components sequentially increasing (Fig. 2E). Under conditions of \(k_{cell} << k_{dye}\), obtained by using millimolar concentrations of Bis-Fura-2, the additional buffering capacity of the indicator prevents the transient saturation resulting in equal Ca\(^{2+}\) components associated Ca\(^{2+}\) spikes. The need for \(k_{cell} >> k_{dye}\) is therefore obvious for this type of measurement.

**APPLICATION 2: COMBINING CA\(^{2+}\) IMAGING WITH NEUROTRANSMITTER PHOTORELEASE**
The analysis of Ca$^{2+}$ signaling associated with excitatory synaptic transmission is complicated by the fact that extracellular application of pharmacological tools, necessary to separate various Ca$^{2+}$ components, often causes presynaptic effects. A way to investigate the consequence of glutamate receptor activation, without excitation of presynaptic terminals, is to use photorelease of L-glutamate from caged compounds. This approach fulfills the ideal requirement of mimicking synaptic transmission on a small spatial scale and a fast time scale. A widely used caged compound is the 4-methoxy-7-nitroindolinyl L-glutamate (MNI-glutamate), caged L-glutamate which has sub-microsecond kinetics (Morrison et al 2002) and thermally stable and efficient in photorelease (Papageorgiou et al. 1999; Papageorgiou and Corrie 2000). It has been in use for more than 10 years, and the only negative side – action found is a low affinity block of GABA$\text{A}$ receptors by the cage itself (IC(50) block = 0.6 mM, Fino et al. 2009). Nevertheless it is essentially biologically inert in its caged precursor form at 1 mM and in the photolytic intermediates and by-products (Canepari et al. 2001a).

Good spatial resolution in experiments combining imaging with photolysis can be achieved in two ways. First, excitation of photolysis at a spatial resolution of 1 micron can be achieved with laser photolysis with one or two photon excitation. With extracellular cages like MNI-L-glutamate the main difficulty is the absorption of photolysis light by the cage in the bath – an inner filtering- resulting in poor penetration to the focus at the peak absorption wavelengths near 340 nm. This effect can be overcome by using wavelengths away from the absorption peak. For MNI-caged ligands a 405 nm laser is effective, producing a 1 $\mu$m spot, and efficient enough to produce full photolysis conversion without phototoxicity (Trigo et al, 2009). Two-photon photolysis (Matsuzaki et al., 2001; Carter and Sabatini 2004; Noguchi et al. 2005; Nikolenko et al. 2007) also avoids the inner filtering effect, however two-photon excitation of MNI-glutamate, and all other cages currently available, has very poor efficiency relative to one photon excitation. Because of the small volume in laser spot photolysis it is necessary to use brief stimulations, less than 300 $\mu$s duration, to maintain spatial resolution without smearing due to diffusion. In the second approach spatial resolution can be achieved by uniform wide field photolysis combined with spatially resolved imaging to monitor the distribution and time course of fluorescence. In this case photolysis pulses of 1 ms can be used with a flashlamp (Canepari et al. 2001a, 2001b, 2004; Canepari and Ogden 2003, 2006), or longer exposures with LED (Bernardinelli et al. 2005) or shuttered arc lamp illumination.

An important requirement in combined Ca$^{2+}$ imaging and photolysis is that the excitation wavelength of the Ca$^{2+}$ indicator does not produce photolysis. For MNI-glutamate, wavelengths $\geq$ 420 nm satisfy this requirement. Imaging with Fura indicators can still be done at 420 nm (Ogden et al. 1995; Canepari et al. 2004), but blue-excitable Ca$^{2+}$ indicators (such as Fluo, Oregon-Green or Calcium Green dyes) which show a fluorescence increase may be preferable for this application. Fig. 3 shows an example of
combined Ca\textsuperscript{2+} imaging with L-glutamate uncaging (from Canepari and Ogden 2006). A rat PN was filled with the Oregon Green BAPTA-5N indicator (K\textsubscript{d} ~ 35 µM) at 0.9 mM concentration. L-glutamate was photoreleased over the whole dendritic tree. Fig. 3A shows a dendritic region from which Ca\textsuperscript{2+} fluorescence was averaged. To study Ca\textsuperscript{2+} signals mediated by type-1 metabotropic glutamate receptors, voltage-gated Na\textsuperscript{+} and Ca\textsuperscript{2+} channels, AMPA receptors, NMDA receptors and GABA\textsubscript{A} receptors were blocked by tetrodotoxin (1 µM), agatoxin IVA (250 nM), NBQX (100 µM), AP5 (50 µM) and bicuculline (20 µM). Activation of mGluR1 triggered two independent intracellular Ca\textsuperscript{2+} pathways. At -77 mV holding potential (HP), mGluR1 activated a slow non-selective cation conductance (Canepari et al. 2001b) permeable to Ca\textsuperscript{2+} (Canepari et al. 2004) (Fig. 3B left). At -12 mV HP, mGluR1 also triggered Ca\textsuperscript{2+} release from stores via InsP\textsubscript{3} receptors, leading to an outward K\textsuperscript{+} current from BK channels (Canepari and Ogden 2006) (Fig. 3B right), thus distinguishing two mGluR pathways based on kinetics and prior Ca\textsuperscript{2+} influx.

**FIGURE 2.** Combining \(V_m\) and Ca\textsuperscript{2+} imaging (A) Area of optical recording of a CA1 hippocampal pyramidal neuron filled with the voltage sensitive dye JPW1114 and with the Ca\textsuperscript{2+} indicator Bis-Fura-2 (300 µM); two regions \(a\) and \(b\) and the position of the stimulating electrode indicated. (B) Somatic \(\Delta V_m\) (black traces) and Ca\textsuperscript{2+} signals from \(a\) and \(b\) (grey traces) associated with a bAP (top) and with the pairing of 5 EPSPs and 1 bAP. (C) \(\Delta V_m\) optical signals from \(a\) and \(b\) associated with the bAP (white traces) and with the pairing protocol (black traces). (D) A PN filled with the voltage sensitive dye JPW1114 and with the Ca\textsuperscript{2+} indicator Fura-FF (1 mM); a region and the position of the stimulating electrode indicated. (E) \(\Delta V_m\) (left) and Ca\textsuperscript{2+} (right) signals from the region in D associated with 7 PF- EPSPs at 100 Hz; calibration described in Canepari et al. 2008.
**FIGURE 3.** Ca^{2+} signals mediated by mGluR1 activation and elicited by L-glutamate photorelease in cerebellar Purkinje neurons. (A) Fluorescence image of a sub-region of a PN filled with Oregon Green BAPTA-5N. (B) Somatic membrane current (black traces) and Ca^{2+} signals (gray traces) from the region in A and mediated by mGluR1 activation and elicited by L-glutamate photorelease; left: signals at -77 mV holding potential (HP); right: signals at -12 mV holding potential.
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