Cellular/Molecular

Branch-Specific Ca\(^{2+}\) Influx from Na\(^{+}\)-Dependent Dendritic Spikes in Olfactory Granule Cells

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Two-photon laser scanning microscopy was used to correlate electrical events detected with whole-cell somatic recordings to Ca\(^{2+}\) transients in dendrites of olfactory bulb granule cells. A subset of spontaneous subthreshold depolarizing events recorded at the soma were shown to correspond to suprathreshold dendritic, Na-dependent action potentials [APs; dendritic spikes (D-spikes)]. These potentials were blocked by intracellular QX-314 (lidocaine N-ethyl bromide), hyperpolarizing current injection at the soma, and by partial inhibition of AMPA/kainate receptors with 0.75 \(\mu\)M DNQX. They were affected only slightly by 100 \(\mu\)M NiCl\(_2\). The majority of D-spikes recorded at the soma had a time to peak of <4 ms, comparable with somatic APs, a nonexponential decay, and amplitudes between 3 and 21 mV. Soma-locally recorded APs produced Ca\(^{2+}\) transients that were observed in spines and dendrites in all parts of the cell. Ca\(^{2+}\) transients from D-spikes were restricted to subsets of distal dendrites and their associated spines but were absent from the soma and dendrite within \(\sim 50–80 \mu\)m of the soma. Ca\(^{2+}\) transients in different branches could be correlated with different-sized D-spikes. D-spike and backpropagating AP-induced Ca\(^{2+}\) transients summed in dendrites, provided the interval between them was >5–6 ms. Generation of a D-spike in a particular dendrite <5–6 ms before a somatic AP blocked backpropagation of the somatic AP into that dendrite. The temporally specific interplay between D-spikes and backpropagating APs may play a role in regulating feedback and feedforward inhibition of groups of mitral cells synapsing on different granule cell dendrites.

Key words: dendrite; dendritic action potential; backpropagating action potential; calcium imaging; olfactory bulb; granule cell

Introduction

Many olfactory neurons have dual-functional dendrites that receive synaptic inputs and release transmitter. Determining the extent to which these cells can compartmentalize their input–output functions is essential to understanding their computational functions.

Active electrical responses in dendrites are proving to be the norm rather than the exception (Segev and London, 2000). The extent to which Na\(^{+}\)-dependent action potentials (APs) backpropagate into or are initiated in dendrites (Bischofberger and Jonas, 1997; Chen et al., 1997; Saraga et al., 2003) varies across neuronal types and often depends on factors such as resting membrane potential, spiking pattern (Hoffman et al., 1997), synaptic activity (Margrie et al., 2001; Lowe, 2002), or neuromodulation (Tsubokawa and Ross, 1997; Hoffman and Johnston, 1999; Davison et al., 2004).

Several roles are evident for APs in dendrites (Hauser and Mel, 2003). In some instances, notably pyramidal cells in mammalian hippocampus, backpropagating APs appear to play a pivotal role in activity-dependent modification of synaptic strength by correlating postsynaptic electrical activity with synaptic input within a narrow temporal window (Migliore et al., 1999; Sjoström and Nelson, 2002; Watanabe et al., 2002; Kampa et al., 2004). In other cells, they can serve to efficiently transmit or enhance the transfer of synaptic input from distal dendrites to the soma/axon compartment (Chen et al., 1997; Hanson et al., 2004).

In neurons such as olfactory granule cells that release transmitter from their dendrites, they obviously can play a major role in initiating transmitter release from synapses distributed throughout the arbor (Isaacson and Strowbridge, 1998). Compartemental modeling has suggested that there are several potential levels of organization of output synapses on granule cell dendrites (Woolf et al., 1991), and it seems probable that whether or under what conditions the granule cell can function in local versus global output modes will have important bearing on the computational role it performs in the olfactory bulb circuit (Shepherd et al., 2004).

We used fluorescence imaging of dendritic Ca\(^{2+}\) transients combined with whole-cell intracellular recordings from somata to study spatial properties of AP propagation in granule cell dendrites of frog olfactory bulb. We focused on Ca\(^{2+}\) transients in dendrites resulting from spontaneous activity and from APs evoked by somatically injected depolarizing current or electrically stimulated synaptic inputs. Somatic APs reliably backpropagate into all dendritic branches and spines to produce robust...
Ca\(^{2+}\) transients, in agreement with previous findings from mammalian slice preparation (Egger et al., 2003). Additionally, we find that primarily Na\(^{-}\)-dependent, dendritic spikes (D-spikes) produce Ca\(^{2+}\) transients that can be restricted to subsets of dendritic branches. This adds to growing evidence for dendritically generated active events in granule cells of turtle and rat reported recently by Pinato and Midggaard (2005) and Egger et al. (2005). In frog, the soma and proximal dendritic segment appear to be a barrier to their propagation, resulting in restriction of D-spike-evoked Ca\(^{2+}\) influx to sets of spines on peripheral branches. Ca\(^{2+}\) transients evoked by backpropagating spikes can add to those from dendritic spikes in those branches that initiate dendritic spikes. However, because dendritically initiated spikes are active events, they block backpropagation of somatic APs for intervals <5–6 ms. These time- and space-dependent interactions between D-spikes and somatic spikes could have branch-specific consequences for transmitter release or synaptic plasticity.

Materials and Methods

Electrophysiology. Procedures were performed in accordance with Canadian Council for Animal Care Guidelines under supervision of the institutional Animal Care and Use Committee. Recordings were made from neurons in hemisected but otherwise intact forebrain of adult Rana pipiens. After deep anesthesia by immersion in MS222 (Sigma, St. Louis, MO), the brain was removed and divided by a razor blade cut along the midline through the thin anterior interbulbar adhesion and callosal fibers of the telencephalon. The olfactory bulb and telencephalon were pinned to the bottom of a Sylgard-lined recording chamber with the cut medial surface facing upwards. Experiments were performed at room temperature, which was regulated to 21 ± 1°C. A bicompartiment-buffered frog Ringer’s solution (in mM: 72 NaCl, 2 KCl, 26 NaHCO\(_3\), 2.5 NaH\(_2\)PO\(_4\), 0.5 NaHPO\(_4\), 1.5 MgSO\(_4\), 2 CaCl\(_2\), 10 glucose, pH 7.4) bubbled with 95%/5% O\(_2\)/CO\(_2\) was superfused at a rate of 1–2 ml/min (Delaney and Hall, 1996). Granule cells were visualized using near-infrared illumination (625–750 nm) with an oblique illumination condenser (Olympus, Melville, NY). Somata were selected for patching between 100 and 170 μm below the surface (the majority were ~120 μm), and most cells were located in the caudal (i.e., interior) portion of the granule cell layers. Average resting membrane potential and input impedance were estimated to be ~59.5 ± 3.4 mV (n = 53) and 231 ± 81 MΩ (mean ± SD; r = 0.94), respectively.

Whole-cell recordings were performed using pipettes pulled from nonfilament thin-wall glass capillaries with resistances of 6–8 MO \(\Omega\) containing a K-glucuronate-based intracellular solution [in mM: 10 HEPES, 5 glucose, 2 MgCl\(_2\), 2 ATPNa\(_2\), 0.3 GTP-Na, 91 K-glucuronate, 195 mM NaCl (measured), pH 7.4 adjusted with KOH]. Seal resistances of at least 3 GΩ were obtained in cells chosen for study, and access resistance of ~30–70 MΩ (55 ± 26; n = 94) was typical after breakthrough. Voltages reported here are not corrected for junction potentials, and series resistance compensation was not performed. Most physiological measurements were performed within 2–2.5 h of breakthrough, during which the electrophysiological properties of the cells, including the occurrence of dendritic spikes, were typically stable. A twisted pair of 125-μm-diameter enamel-coated Ag wires was used to stimulate olfactory nerve (ON) or lateral olfactory tract (LOT). For local stimulation of mitral cells in the vicinity of imaged granule cells, a monopolar glass pipette with a 10-μm-diameter tip filled with 0.5 M NaCl was positioned in the superficial portion of the external plexiform layer using a stage mounted hydraulic manipulator under bright-field visual control.

To test for possible T-type Ca\(^{2+}\) channel contribution to spontaneous D-spike generation, we superfused the bulbs with Ringer’s containing 100 μM NiCl\(_2\). The time to peak, amplitude, and frequency were obtained for all D-spikes in each of six cells during a 5 min control period comprising 20 min after breakthrough and then for 5 min after superfusion with NiCl\(_2\), for 20–25 min. Average values were obtained for these parameters by measuring from between 24 and 123 D-spikes during control and NiCl\(_2\) application. D-spikes were defined as events with \(dV/dt > 0.6\) V/s, and we confirmed in each case that these events were eliminated by hyperpolarizing current sufficient to bring the somatic membrane to ~80 mV. We also confirmed that access resistance remained constant during the course of the experiment and that there was no change in amplitude, rate of rise, or frequency of D-spikes during the 20 min control period in artificial CSF before application of NiCl\(_2\).

Imaging. Ca-Green-1 K+ salt (150 μm; Invitrogen, San Diego, CA) was loaded into neurons by diffusion from the recording pipette. A minimum of 30 min elapsed before commencing recording of fluorescent transients from dendrites. Fluorescence was excited using 810 nm pulsed laser light generated by a Nd:YAG pumped Ti:sapphire laser system (Verdi 10W; MIRA Optima 900F; Coherent Laser, Santa Clara, CA). An x–y–galvonometer mirror assembly (Cambridge Instruments, Deerfield, IL) scanned the beam across the sample through a 40×, 0.8 numerical aperture water dipping lens (Olympus), and fluorescence signals were detected using a GaAsP photomultiplier (H7742P–40 MOD; Hamamatsu, Bridgewater, NJ) using a custom-built microscope modelled after that described by Tsai et al. (2002). A Macintosh (Apple Computers, Cupertino, CA) 867 MHz dual processor computer equipped with an NI PCI6110 analog-to-digital (A/D) board was used to acquire fluorescence signals (600 KHz to 1.25 MHz sampling rate) and to output scan-sweep signals simultaneously with acquisition of electrophysiological signals that were obtained with a National Instruments (Austin, TX) PCI6035E A/D board (5 KHz sampling rate). Electrophysiology and imaging were synchronized through real-time signal integration lines on the two digitizing boards. Hardware control, image acquisition, and analysis software were written in-house using Igor Pro (WaveMetrics, Portland, OR) as a platform. External routines written in C++ were linked to the Igor interface to accelerate some display and image analysis functions. Most imaging was performed by defining scan areas comprising 10–20 horizontal scan lines that encompassed a length of dendrite including spines. Dendrites were aligned along the horizontal scan axis using a hardware-based scan rotation device (Tsai et al., 2002) with bidirectional scanning in the horizontal axis to increase scanning rates. Frame rates of 30–150 frames/s were used in this study.

Polygonal regions of interest were defined around dendrites and spines. Fluorescence changes (ΔF/F) were normalized by dividing by the average baseline fluorescence during a period of inactivity (usually obtained during the first 1–2 s at the beginning of a trace). Photomultiplier offset was adjusted to bring background intensities in unfilled areas to near zero. A smoothing polygonal fit (Savitzyk-Golay; 9 or 11 point window) was used to temporally smooth the fluorescence data presented. At the end of the experiment, cell morphology was reconstructed from sets of optical sections obtained with vertical steps of 1, 2, or 3 μm to determine where optical measurements had been made.

Results

Subthreshold spontaneous events recorded at the soma
Whole-cell current-clamp or voltage-clamp recordings from granule cells obtained with standard intracellular pipette solutions are characterized by numerous spontaneous subthreshold depolarizing events recorded at the soma that occur at an estimated rate of 10–30 Hz (Fig. 1A, B). The majority of these events range in size from 0.3 to 5 mV with times to peak between 5 and 22 ms. This class of events was blocked by DNQX, not affected by intracellular lidocaine N-ethyl bromide (QX-314; 1 mM), and not diminished in size or frequency by hyperpolarizing current injection, consistent with their being primarily AMPA receptor (AMPA)-dependent synaptic potentials.

Another class of subthreshold events was observed with times to peak of <4 ms (typically 2–3.5 ms), amplitudes of 3–21 mV, and a faster initial decay phase (Fig. 1C, D) compared with the majority of subthreshold events. They occurred spontaneously in 71 of 94 cells at normal resting membrane potential of around −60 mV. Because the frequency of these fast potentials varied between different granule cells and depended strongly on resting...
membrane potential, we examined the last 36 cells that we recorded during our study that maintained membrane potential between −57 and −63 mV with either no injected current (18 cells) or between −1 and −13 pA (18 cells). During recording periods of at least 5 min duration obtained within 10 min of initial breakthrough, the frequency of these fast potentials ranged from 0.017 to 1.574 Hz with an average rate 0.30 ± 0.36 Hz (n = 36). In any given preparation, although the overall average rate of these fast subthreshold events was usually <1 Hz, short bursts of two to five events at rates as high as 150 Hz were occasionally seen.

When present in a cell, these events persisted for the duration of recordings without evidence of washout for up to 2 or 3 h after breakthrough with and without high-affinity Ca\(^{2+}\) indicator in the pipettes. Despite the small overlap in size between the two classes of subthreshold potentials, the different time course of the rising and initial recovery phases clearly separates the events into distinct populations (Fig. 1E). For example, for the cells shown in Figure 1E, the average times to peak of the fastest class of events was 2.53 ± 0.30 ms (n = 11) (Fig. 1Ei), 3.65 ± 0.77 ms (n = 22) (Fig. 1Eii), and 2.96 ± 0.24 ms (n = 16) (Fig. 1Eiii) compared with the EPSPs, which had average times to peak of 9.66 ± 3.06 ms (n = 126) (Fig. 1Ei), 8.61 ± 2.60 ms (n = 51) (Fig. 1Eii), and 10.0 ± 3.38 ms (n = 76) (Fig. 1Eiii). The average amplitudes of the fastest class of events were 12.30 ± 5.1 mV (Fig. 1Ei), 6.24 ± 1.6 mV (Fig. 1Eii), and 11.8 ± 5.2 mV (Fig. 1Eiii), whereas the corresponding average EPSP amplitudes were 1.16 ± 0.58 mV (Fig. 1Ei), 1.43 ± 0.89 mV (Fig. 1Eii), and 1.54 ± 0.91 mV (n values as indicated for time to peak data). The faster events were thus readily isolated from the majority of subthreshold potentials by plotting dV/dt versus time for current-clamp recordings (Fig. 1B, F, G). D-spikes had peak rates of rise of >0.5 V/s, typically >2 V/s, whereas derivatized EPSPs were usually not distinguishable above the noise level (<0.2 V/s).

Hyperpolarizing current injection readily blocked the fast class of spontaneous events. Calcium imaging experiments described below led to the conclusion that the fast class of events corresponded to action potentials initiating in the dendrites, that is D-spikes. The combination of a rising phase with maximum dV/dt > 0.6 V/s and susceptibility to hyperpolarization that was always seen for these events served as a defining criterion for D-spikes. We routinely injected hyperpolarizing current as a means of confirming that fast subthreshold events were D-spikes and not large EPSPs, because EPSPs increased in size with hyperpolarization, whereas D-spikes were blocked. Typically, hyperpolarization by 10–20 mV below average resting membrane potential was sufficient to block all spontaneous fast events in all cells tested in this manner (n = 62) (Fig. 2A, C), whereas further hyperpolarization blocked those evoked by strong local synaptic stimulation (n = 3) (Fig. 2A, B).

Full-sized spontaneous APs (SAPs) that were recorded from the soma were often evoked by D-spikes, particularly if they summed with smaller slower potentials (Fig. 1F). We examined 317 SAPs in derivatized voltage recordings in 33 cells and detected that the SAP was triggered from a distinct fast rising prepotential in 171 cases. Across the population of cells, we could detect a distinct prespike preceding 50 ± 35% of SAPs; many cells had no detectable prespikes, whereas many others showed prespikes for all SAPs (Fig. 1D, F, G). Although there are high rates of spontaneous synaptic potentials, the threshold for firing at the soma is 10–15 mV depolarized from resting V\text{m} (Hall and Delaney, 2002; Cang and Isaacson, 2003), thus it is not surprising that a large proportion of spontaneous APs are triggered from large ampli-
tude D-spikes. This average possibly underestimates the frequency of prespikes, because they cannot be detected if they exceed the derivatized record for clarity). The insets are expanded time base segments from portions of the continuous record, as indicated by the arrows. Inset Ai shows evoked D-spikes and an aborted action potential; inset Aii shows spontaneous D-spikes between stimuli. With hyperpolarization to about $-79$ mV, only three D-spikes are evoked and no spontaneous D-spikes occur. At about $-60$ mV, many spontaneous D-spikes occur and synaptic potentials evoke action potentials with several D-spikes. Intermediate hyperpolarization to about $-72$ mV produced one spontaneous D-spike and several evoked D-spikes. B, C, Segments of data extracted from another 2 min of recording from the same cell during which the membrane potential was adjusted similar to the records shown in A. Records in which no D-spikes were seen are presented in gray; those with D-spikes are black. B, Stimulation evokes only synaptic potentials at $-79$ mV (Bi); D-spikes and sometimes action potentials are evoked at $-65$ mV (Bii) or $-71$ mV (Biii) average resting membrane potential. C, Spontaneous D-spikes do not occur at $-79$ mV (C); D-spikes occur as indicated by dashed line, and are seen at a lower frequency for $-71$ mV (Ci) versus $-65$ mV (Cii). D, A cumulative count of all D-spikes shows that 1 min QX-314 added to the pipette gradually blocks the occurrence of spontaneous D-spikes (results from 2 cells shown in left panel). Without QX-314, D-spikes are produced at a constant rate as illustrated by data from three other cells (right). The vertical axes are in millivolts or volts per second for A–C.

We examined the effects of DNQX at varying low concentrations (0.5 and 2 mM, 1 and 1.5 mM, 0.75 mM) on the generation of D-spikes in three preparations. DNQX at 1–2 mM reduced EPSP amplitudes by $\sim 75\%$ and blocked all D-spikes within 5 min. We could not restore D-spikes with depolarization under these conditions. Partial blockade of AMPA/kainate receptors with 0.75 mM DNQX reduced synaptic potentials to about one-half of their normal size (Fig. 3C) and blocked spontaneous production of D-spikes (beginning 1300 s after the start of the experiment) (Fig. 3A,C). After abolition of all D-spikes by 0.75 mM DNQX, depolarization of the cell (Fig. 3B) could restore their occurrence at a low rate but usually not before initiating numerous full-sized somatic APs (Fig. 3C, right). The fast events that could be evoked

Figure 2. Voltage dependence and sensitivity to QX-314 of dendritic spikes. A, The top trace shows a continuous current-clamp recording with and without hyperpolarizing current injection with the corresponding derivatized record underneath. A stimulating pipette inserted into the superficial external plexiform layer stimulated synaptic inputs at 5 s intervals (stimulus artifacts can be seen in voltage recording but were removed from the derivatized record for clarity). The insets are expanded time base segments from portions of the continuous record, as indicated by the arrows. Inset Ai shows evoked D-spikes and an aborted action potential; inset Aii shows spontaneous D-spikes between stimuli. With hyperpolarization to about $-79$ mV, only three D-spikes are evoked and no spontaneous D-spikes occur. At about $-60$ mV, many spontaneous D-spikes occur and synaptic potentials evoke action potentials with several D-spikes.

Figure 3. Partial block of AMPA receptors inhibits spontaneous D-spikes. A, Cumulative count of D-spikes before and after addition of DNQX. B, Record of current injected through the pipette during the experiment. C, Voltage recording. D, Derivitized voltage with segments between 300 and 350 s and 2500–2550 s expanded to show the loss of D-spikes but not somatic APs after partial AMPAR block. E, All D-spikes detected during the periods indicated by the white bars over the data in C.
with depolarization in the presence of AMPAR block appeared to be of similar size and time course as some of the events seen before block (Fig. 3E). These observations are consistent with the spontaneous generation of D-spikes by synaptic depolarization at spike-initiating zones distant from the soma. In the absence of sufficiently large distal synaptic potentials, produced by partial AMPAR block, current injected into the soma preferentially activates a proximal spike generating region in or near the soma rather than a distal dendrite (Fig. 3D).

Evidence for T-type Ca\(^{2+}\) channel dependent active responses in granule cells has been reported by two groups, Egger et al. (2003, 2005) in rat and Pinato and Midggaard (2003, 2005) in turtle. Perfusion with 100 \(\mu\)M NiCl\(_2\) for 20 min had a small (17\%) slowing effect on time to peak of spontaneous D-spikes in all cells tested (see Materials and Methods; mean difference, 0.76 \(\pm\) 0.42 ms; \(n = 6\) cells; paired sample \(t\) test, 0.005 < \(p < 0.01\)). There was also a reliable decrease in amplitude in all cells of 3–28\%, with a mean of 18\% (mean difference, \(-1.54 \pm 0.81\) mV; \(n = 6\); paired sample \(t\) test, 0.005 < \(p < 0.01\)). These small changes were accompanied by a tendency for instability in resting membrane potential and intermittent short bursts of two or three APs, unless a few picoamperes of negative current, sufficient to hyperpolarize the membrane by 5–10 mV, were applied. Consequently, if no additional hyperpolarizing current was applied during NiCl\(_2\) application, D-spike frequency would increase by \(\sim 50\%\) along with SAP generation. We therefore applied enough hyperpolarizing current to try to maintain the SAP rate in NiCl\(_2\) similar to the control rate. Under these conditions, the rate of spontaneous D-spike generation was slightly (115 \(+\) 27\%) but not significantly increased. The increased frequency and depolarization may be the result of reduced inhibition of mitral cells by granule cells, leading to more excitatory input. It is also possible that T-channels may be contributing Ca\(^{2+}\) to activate Ca\(^{2+}\) dependent K\(^+\) channels that control excitability of granule cell dendrites. We have not yet performed imaging experiments in combination with T-channel blockers. Nor have we explored the effect of Ni\(^{2+}\) on T-spikes evoked by localized synaptic stimulation. The time to peak (\(\sim 4\) ms) and initial rapid rate of recovery of our D-spikes (Fig. 1C) are comparable with somatic APs and almost an order of magnitude faster than the stimulus-evoked, low-threshold spikes described in mammalian granule cells [Egger et al. (2003)], their Fig. 6; Egger et al. (2005), their Fig. 8A], suggesting a smaller contribution of T-channels to D-spikes in frog granule cells. Furthermore, unlike rodent granule cells [Egger et al. (2003), their Figs. 4A and 5], we have not yet seen facilitation of Ca\(^{2+}\) transients evoked by backpropagating APs by hyperpolarizing presteps, nor do we observe a voltage-dependent after-spike depolarization (data not shown) (Hall and Delaney, 2002). Thus, although spontaneous production of D-spikes is not blocked by NiCl\(_2\), there is some evidence to suggest a contribution of T-type currents to dendritic excitability in frog granule cells.

The pharmacological and electrophysiological profile of these fast events that we have defined as D-spikes suggests they are active, predominantly sodium-dependent spikes, generated in the periphery by spontaneous synaptic input that occasionally is sufficiently large enough to locally generate a dendritic AP. Activation of excitatory synaptic inputs by stimulation of mitral cells causes a mixed response in granule cells comprised of synaptic potentials, D-spikes, and sometimes SAPs. Similar dendritically generated spikes with a significant Na\(^+\) dependence have been reported recently in response to evoked synaptic excitation in isolated turtle olfactory bulb (Pinato and Midggaard, 2005).

Ca\(^{2+}\) imaging experiments are described in following sections to directly support the conclusion that, although they are subthreshold in the soma, these events correspond to large amplitude suprathreshold action potentials in the dendrites.

**Ca\(^{2+}\) transients in dendrites**

High-speed imaging of dendrites was performed in 73 cells to characterize dendritic Ca\(^{2+}\) transients to backpropagating APs [SAPs or current-evoked APs (EAPs)] and to determine whether D-spikes that were subthreshold at the soma were correlated with Ca\(^{2+}\) transients in the dendrites. Given the relatively low rate of occurrence of spontaneous D-spikes, we adopted a strategy of measuring fluorescence and electrophysiology continuously over periods of 15–22 s (Fig. 4A). Spontaneous APs and EAPs produced Ca\(^{2+}\) transients in proximal and distal dendrites of all 73 cells. We did not attempt to detect differences in the amplitude of fluorescence transients in proximal versus distal dendrites as reported for mammalian granule cells in brain slice (Egger et al., 2003). Rather than focus on differences in backpropagating somatic AP-evoked Ca\(^{2+}\) in different parts of the neuron, we used the reliable Ca\(^{2+}\) transient resulting from a backpropagating EAP as a standard against which SAP and D-spike Ca\(^{2+}\) transients could be normalized (Fig. 4F). Near the beginning of each recording period, we typically stimulated two EAPs at an interval of 1 s by injecting 45 ms depolarizing current pulses through the electrode. Ca\(^{2+}\) transients generated by SAP or by D-spikes were then compared with the transients produced by action potentials evoked through somatic current injection (Fig. 4A, D, F).

When a Ca\(^{2+}\) transient was observed in the dendrite, in response to an EAP or a SAP, spines that were attached to that dendrite showed a corresponding transient. Examples from two cells are illustrated in Figure 5, A and B. We examined 102 spines in 22 granule cells for which we had Ca\(^{2+}\) and electrophysiological recordings of EAPs and SAPs and could discern at least one spine and a portion of the dendrite from which it originated. For every EAP or SAP, there was evidence for a comparably sized transient in both the spine and the dendrite. For 8 of these 22 cells, we were able to obtain good signal to noise in optical traces in spines (one to three per cell) and a piece of adjacent dendrite for which we had responses to between 5 and 14 APs (EAP or SAP). The average fractional change in fluorescence in the spine was 1.05 \(\pm\) 0.29 of that in the adjacent dendrite (\(n = 101\) events).

Action potential-evoked Ca\(^{2+}\) transients were seen in all dendrites of the 73 cells that were imaged. For these 73 cells, we examined one primary branch in 35 cells, two different main branches in 33 cells, and three or more branches in five cells. Of the 73 cells we imaged, 60 had spontaneously occurring electrophysiologically defined D-spikes. Our success in finding a spontaneous D-spike-associated Ca\(^{2+}\) transient was probably reduced by the fact that we did not image all branches. For the 60 cells with D-spikes, we imaged one main branch in 23 cells, two branches in 32 cells, and three branches in five cells. We identified D-spike-associated Ca\(^{2+}\) transients in at least one dendrite in 26 (43\%) of these cells after imaging one main branch in 10 cells, two branches in 14 cells, and three branches in two cells.

If a D-spike produced a Ca\(^{2+}\) transient, it was comparable in size to that of an EAP or an SAP (see Figs. 4D, F, 5B, 7A). Measuring at different points along a dendrite showed that when a D-spike evoked Ca\(^{2+}\), it did so over a large length (see Figs. 4D, 6, 9E–G).

Localized spine Ca\(^{2+}\) transients were described by Egger et al. (2005) in response to minimal synaptic stimulation. In our study, Ca\(^{2+}\) transients in the absence of either an AP or a D-spike that
were restricted to individual spines and absent from the adjacent
dendrite were observed spontaneously in a small number (five) of
instances. Examples of such responses are shown in Figure 5A
threshold by the summation of small synaptic potentials distrib-
uted across a wide area. Using a small-diameter pipette posi-
tioned in the superficial external plexiform layer adjacent to a

Because D-spikes occurred spontaneously at rates on the order of 0.5 per sec-
ond for the cell as a whole and less than once every 10 or more seconds in any given branch,
the opportunities for averaging several responses were limited. Nonethe-
less, when we were able to obtain sufficient signal to noise from single spines, a
D-spike-associated Ca²⁺ transient re-
sponse was always observed coincident with a transient in the adjacent parent dend-
rite (Fig. 5B). In six cells, each with one or
two spines visible (eight spines total) for
which we had at least two recordings of a
D-spike Ca²⁺ transient with good signal to
noise, the fractional fluorescence change in
the spine was 0.98 ± 0.23 of that in the adjacent dendrite (n = 33 events).

Observing that not all D-spikes evoked
Ca²⁺ transients in the segment of dendrite we
were imaging, we hypothesized that
some D-spikes originated in other den-
drite(s) and did not invade the one we
were imaging. Although we could rou-
tinely image across 100 μm of visual field,
simultaneously imaging from segments of
more than one widely separated dendrite
was difficult because of the small chance of
finding two segments from different den-
drites in the same 2- to 3-μm-thick optical
section. We therefore compared the elec-
rophysiological waveform of the D-spikes
that were associated with Ca²⁺ transients
to those that were not to test for dendrite
specific Ca²⁺ transients. We reliably ob-
served that some of the classes of D-spikes
that could be distinguished based on am-
plitude and rise time were never correlated
with Ca²⁺ transients in the dendrite we
were imaging. On the other hand, a subset
of D-spikes defined by their similar size
and shape were correlated with a Ca²⁺
transient (Fig. 6) (see also Figs. 7, 9 B, G, F).
Furthermore, the Ca²⁺ transients ob-
served sequentially in two different main
dendritic branches could be correlated with
the occurrence of two different classes
of D-spikes (Fig. 6).

Using ON or LOT stimulation to evoke
synaptic potentials, we were not able to ac-
tivate a D-spike without also inducing a
full-blown somatic potential. We presume
this is because ON and LOT stimuli acti-
uate synapses distributed across all den-
drites, so the soma can be brought to

Figure 4. Fluorescence (Ca²⁺) transients associated with somatic action potentials and fast subthreshold events (D-spikes). A, Continuous recording (19.8 s) of somatic membrane potential and fluorescence measured from an 80-μm-long segment of dendrite 250 μm distal to the soma (illustrated in D). EAP indicates action potentials evoked by injection of depolarizing current into the soma. Several D-spikes are indicated by asterisks above the recording, one of which is correlated with a fluorescence transient in the dendrite as indicated by the dashed vertical line. The dots in the ∆F/Φ trace show measurements from each frame, whereas the solid line is a smoothing polynomial fit to the data. B, C, Expanded views of segments of recording in A, as indicated by the arrows that show the variation in Ca²⁺ transients produced by spontaneous somatic APs depending on whether they are preceded by dendritic spikes. D, Both dendritic and EAPs produce Ca²⁺ influx along the length of distal dendrites, and transients produced by spontaneous dendritic spikes are comparable with those produced by EAPs. The black solid line measured from entire length and spines as shown in the accompanying image, the gray solid line, and the black dotted line are measurements made from —25% of the distal and proximal ends of the dendrite, as indicated by the lines above the image. E, Fifty D-spike segments extracted from 78 s of recording from this dendrite location. The black traces show eight D-spikes that were associated with either a fluorescence transient alone or a larger than normal SAP-evoked transient (e.g., A). The gray traces show all other occurrences of dendritic spikes. F, Distribution of fluorescence transient amplitudes measured from 20 cells. Amplitudes were normalized to the average of the two EAPs stimulated near the start of each recording period (e.g., A). The top distribution shows responses to
D-spikes. The bottom distribution shows transients from SAPs. Class 1 transients are defined as those that were <150% and class 2 as those that were >150% of the size of the average SAP transient.
filled granule cell dendrite, we were able to generate synaptic potentials from small numbers of mitral dendrite spikes. Dendritic spikes in Olfactory Granule Cells

Figure 5. Ca²⁺ transients in single spines compared with adjacent dendrite. A, Responses recorded simultaneously in four spines and dendrite to EAPs and SAPs. Also indicated by arrows are two rare events showing Ca²⁺ transients that were not associated with either a D-spike or an AP, presumably synaptic in origin, that were restricted to a single spine. B, Two D-spike (*) with Ca²⁺ transients (arrows) observed simultaneously in a spine head and adjacent dendrite. The gray traces show Ca²⁺ responses in the spine head; the black traces show responses in the dendrite.

In two cells, we were able to simultaneously measure from two widely separated dendrites that fortuitously passed through the same optical section. In this case, we were able to observe D-spike-evoked Ca²⁺ transients in one dendrite that did not appear in the other and vice versa (Fig. 8A–C).

In order for D-spike to be restricted to one or a subset of dendrites, they must “fail” at some point. Because they are recorded as subthreshold events in the soma, they obviously fail in this compartment. Frog granule cells in the posterior region of the bulb where we recorded are often multipolar, and in these cases, the soma suffices to isolate one branch from another. In 26 cells, we confirmed the presence of a D-spike-associated Ca²⁺ transient in distal dendrites. In eight of these cells, we also made measurements within 10–80 μm of the soma and did not observe any Ca²⁺ transients in this region in response to a total of >100 D-spike traces. Although there were no Ca²⁺ transients associated with D-spike in dendrite proximal to the soma, in these eight cells, we confirmed that EAPs and SAPs always produced clearly detectable transients, which suggests the electrophysiological failure of D-spike occurs in proximal dendrite near to the soma. Therefore, major branches could possibly be isolated from each other with respect to D-spike if they arose from a branch-point near the soma. Direct evidence for isolation of D-spike to separate distal branches with the same parent dendrite was obtained in the cell with this kind of morphology that we sampled in this way (Fig. 8D,E). Additional specific studies are needed to determine the extent to which this observation is the norm and whether neuromodulation controls the spread of D-spike.

Interactions between D-spikes and somatic APs

Spontaneous APs were generated at irregular intervals. They were often preceded by one or more D-spikes (Figs. 1D, 4B), and this was particularly evident by examining differentiated voltage recordings (Fig. 1F, G). Ca²⁺ transients associated with spontaneous APs were either comparable to those from current-evoked APs or approximately twice the amplitude of D-spike- or current-evoked Ca²⁺ transients (Fig. 4). In every instance (n = 25) in which the Ca²⁺ transient from a SAP was >1.5 times larger than the current-evoked control AP, a D-spike could be detected.
APs are active events, because a brief refractory period would exist after firing of a D-spike, preventing backpropagation of the somatic AP.

**Discussion**

Spontaneously generated active electrical events were detected by a combination of electrophysiological somatic recordings and dendritic Ca\(^{2+}\) imaging. These active events appeared as 3–21 mV depolarizing spikelets, which rose to peak and initially decayed at rates significantly faster than spontaneous synaptic potentials, comparable with full-sized somatically recorded APs. Their sensitivity to low concentration of internally perfused Na\(^+\) channel blocker, their block by hyperpolarizing current, increased frequency with sustained depolarizing current, their rapid time course, and the minimal effects of NiCl\(_2\), all argue against their being primarily Ca\(^{2+}\) spikes supported by T-type channels. Both somatically recorded dendritic spikes (D-spikes) and their Ca\(^{2+}\) transients were abolished by partial AMPA-receptor blockade, indicating they were normally initiated by large amplitude (individual or summed) spontaneous postsynaptic potentials. In support of this conclusion, localized stimulation of synaptic inputs from mitral cells with small bore pipettes positioned close to a dendrite could initiate these D-spikes. Given the low rate of spontaneous D-spikes relative to the frequency of spontaneous synaptic inputs, it seems reasonable to hypothesize that these might result from coincident synaptic inputs to a pair or a few adjacent spines on the same branch. It remains to be determined over what distances synaptic inputs can summate locally to be effective in raising the dendrite above threshold.

Additional evidence that the somatically recorded D-spike spikelets resulted from large amplitude, dendritically generated APs was obtained by imaging Ca\(^{2+}\) transients in dendrites that were correlated with D-spikes. Ca\(^{2+}\) transients associated with D-spikes were restricted to dendrites (and spines), more than ~50–80 \(\mu\)m distal to the soma. The failure to detect Ca\(^{2+}\) transients in the soma or the dendritic proximal to the soma in response to D-spikes does not reflect a lack of voltage-gated Ca\(^{2+}\) channels, because APs that were full sized in somatic recordings produced clear transients in these parts of the cell. Thus, it would appear that D-spikes fail at some point in the proximal dendritic region. Analysis of the waveforms of D-spikes recorded at the soma shows that there are typically several size classes of D-spikes. In most instances, Ca\(^{2+}\) transients in different dendrites were correlated with different D-spike classes, whereas Ca\(^{2+}\) transients measured at different locations along a particular dendrite correlated with only one or a small subset of the electrophysiologically defined D-spikes. This leads to the conclusion that D-spikes are initiated independently in major dendritic branches and are restricted to subsets of dendrites.

Some of our data also show that when a primary dendrite branches a few tens of micrometers from the soma, orthodromically conducting D-spikes from one daughter do not necessarily propagate antidromically out the other. Thus, compartmentalization in the form of branch-specific D-spikes can occur even for granule cells with a single primary dendrite originating from the soma. The different sizes of D-spikes recorded at the soma presumably correspond to properties of the different dendrites such as their diameter or the geometry of their joining the proximal dendritic segment or the soma. Morphological, electrophysiological, and modeling studies will be required to test whether the different D-spike classes correspond to different electrophysiological compartments that are defined by the major branching pattern. Whatever the source of the difference in size, the fact that...
D-spikes from one dendrite can be much larger than others (Figs. 6, 8) means that some branches are inherently more effective for generating the somatic APs that spread throughout the cell. Inhibitory inputs to proximal dendrite or possibly basal dendrites could potentially enhance the segregation of D-spikes or modulate the efficacy of different branches for the initiation of somatic APs.

Action potentials initiated at or near the soma reliably spread throughout all dendrites to cause a Ca\(^{2+}\) influx in all their spines in mammals (Egger et al., 2003) and frog. Thus, inputs that are sufficient to initiate a somatic AP, whether caused by concentrated input to one region or diffuse input to spines on several dendrites, will produce a potential output signal from all release sites. At the other end of the spectrum, the close proximity of NMDA receptors to presynaptic release apparatus in the spine head (Sassoe-Pognetto and Ottersen, 2000; Sassoe-Pognetto et al., 2003) opens the possibility that release can be driven by Ca\(^{2+}\) entering through these receptors. Voltage-clamp experiments have shown that Ca\(^{2+}\) influx through NMDA receptors alone can sustain release under favorable conditions (Halabisky et al., 2000), and so probably at the very least, this influx contributes to AP-evoked release in activated spines. In mammalian slice, recent studies have shown that voltage-dependent channels within the spine compartment can amplify stimulus evoked synaptic potentials, yet under some conditions, the voltage-gated Ca\(^{2+}\) influx is still localized to a spine, so that even voltage-dependent influx can be highly compartmentalized (Egger et al., 2005). Spontaneous Ca\(^{2+}\) transients restricted to single spines were observed too infrequently in the course of our study to characterize their source (NMDA or voltage-gated Ca\(^{2+}\) channels), but their occurrence (Fig. 5A) supports the hypothesis that individual spine input–output functions can occur in amphibian granule cells as well.

Modeling studies that assumed purely passive dendrites (Woolf et al., 1991) suggested that subsets of spines on a single branchlet or local segment of a dendrite could act as a single compartment in response to large synaptic potentials spreading passively. However, at least for frog granule cells, our data suggest that the local excitability of dendrites is generally such that localized synaptic potentials can produce D-spikes that propagate actively along the length of a dendritic branch to all spines Figure 10C. These may not spread to other dendrites, particularly if they originate independently from the soma or from a bifurcation point near the soma.

In many cell types, the soma separates groups of dendrites, for example, apical from basal or basal from basal in pyramidal cells or different primary dendrites of multipolar interneurons. In granule cells, basal dendrites are probably the site of some or all granule–granule inhibitory synapses (Price and Powell, 1970; Wells and Kauer, 1994), whereas apical dendrites are the site for mitral–granule reciprocal synapses, so our data provide evidence for possible separation of output to different classes of targets by D-spikes in these cells.

In addition to synchronizing Ca\(^{2+}\) influx into groups of spines located on the same branch, D-spikes also offer the opportunity to increase the total Ca\(^{2+}\) influx and presumably transmitter release when a somatic AP is initiated within >5 ms after the occurrence of a D-spike. Although APs generated by current injected into the soma have relatively long interspike intervals (>30 ms), our Ca\(^{2+}\) imaging indicates that somatic APs can back-propagate into a dendrite that generated a D-spike at intervals as short as 5–6 ms (150–200 Hz). We have not investigated whether it is possible to locally generate trains of D-spikes in the same dendrite at these high rates with focal synaptic stimulation, but
nonetheless, the invasion of a dendrite by a somatic spike shortly after the occurrence of a D-spike produces an approximate doubling of the total Ca\(^{2+}\) influx into the spines in the dendrite(s) that fired the D-spike(s). This potentially increases the synaptic output from the branch(es) initiating D-spike(s) compared with other branches or may provide a signal for modification of synapses differentially across branches.

So far, our data indicate that for intervals <5–6 ms, backpropagation of a somatically generated AP is blocked or attenuated into a dendrite that has initiated a D-spike (Fig. 10A, B). A possible consequence of this AP collision may be that during periods of synaptic input, when several dendrites are generating D-spikes and releasing transmitter, spines on dendrites that generate D-spikes within <5 ms of a backpropagating AP (i.e., coincident with the SAP) will experience relatively less Ca\(^{2+}\) influx and presumably release comparatively less transmitter (Fig. 10C). Thus, inhibition onto mitral cells synapsing on this dendrite would be expected to be less compared with others. This selective reduction of inhibition to subsets of mitral cells has a parallel in the reduced feedback inhibition of active mitral cells through activation of metabotropic glutamate receptors on granule cell spines, as described by Hayashi et al. (1993). The narrow temporal window for the block of backpropagation might favor firing mitral cells that are active at the phase of the odor-evoked oscillation immediately before the onset of granule cell AP generation. Another consideration though is that the subset of spines that receive synaptic input that is coincident with and/or responsible for initiating a D-spike or a backpropagating AP would have larger Ca\(^{2+}\) transients, as a result of the combination of an
NMII receptor (synaptic) as well as a voltage-gated Ca$^{2+}$ channel (D-spike) influx (note that this is not indicated in Fig. 10C).

Thus, for Ca$^{2+}$ influx into granule cell spines, which will correlate to the efficacy of transmitter release, there are several levels of compartmentalization demonstrated for olfactory granule cells in this and other (Egger et al., 2003, 2005) studies. Ca$^{2+}$ influx restricted to single spines, groups of spines on separate dendritic branches, or all spines on all branches in response to excitatory synaptic input have been observed, demonstrating that there is the scope for these neurons to control their input–output functions across many spatial scales.

References


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