Intrinsic bursting of AII amacrine cells underlies oscillations in the rd1 mouse retina

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Submitted 11 June 2014; accepted in final form 26 June 2014

Intrinsic bursting of AII amacrine cells underlies oscillations in the rd1 mouse retina. J Neurophysiol 112: 1491–1504, 2014. First published July 9, 2014; doi:10.1152/jn.00437.2014.—In many forms of retinal degeneration, photoreceptors die but inner retinal circuits remain intact. In the rd1 mouse, an established model for blinding retinal diseases, spontaneous activity in the coupled network of AII amacrine and ON cone bipolar cells leads to rhythmic bursting of ganglion cells. Since such activity could impair retinal and/or cortical responses to restored photoreceptor function, understanding its nature is important for developing treatments of retinal pathologies. Here we analyzed a compartmental model of the wild-type mouse AII amacrine cell to predict that the cell’s intrinsic membrane properties, specifically, interacting fast Na and slow, M-type K conductances, would allow its membrane potential to oscillate when light-evoked excitatory synaptic inputs were withdrawn following photoreceptor degeneration. We tested and confirmed this hypothesis experimentally by recording from AIIIs in a slice preparation of rd1 retina. Additionally, recordings from ganglion cells in a whole mount preparation of rd1 retina demonstrated that activity in AIIIs was propagated unchanged to elicit bursts of action potentials in ganglion cells. We conclude that oscillations are not an emergent property of a degenerated retinal network. Rather, they arise largely from the intrinsic properties of a single retinal interneuron, the AII amacrine cell.

All amacrine cell; bursting; oscillation; retinal degeneration


Ganglion cells in rd1 retinas are driven to fire bursts of action potentials by phasic synaptic inputs arriving at ~10 Hz (Marc et al. 2007; Margolis et al. 2008; Stasheff 2008). Because oscillatory activity might inhibit normal retinal responses to restored photoreceptor function or might inhibit cortical responses to restored visual input by inducing maladaptive changes in visual circuits, interest in elucidating the circuitry generating it has grown. Speaking to the importance of understanding circuit changes following vision loss, restoration of photoresponsiveness in a blind human patient did not restore visual perception (Fine et al. 2003).

The synaptic drive underlying ganglion cell bursting likely originates in the network of electrically coupled AII amacrine cells and ON cone bipolar (ONCB) cells (Borowska et al. 2011; Trenholm et al. 2012). This notion is supported by the observation that bursting in ON and OFF alpha ganglion cells is anticorrelated (Margolis et al. 2014), just as light-evoked synaptic excitation to these cell types in the normal retina is anticorrelated (Murphy and Rieke 2006); the AII network generates this anticorrelation by providing simultaneously indirect excitation to ON alpha cells and direct inhibition to OFF alpha cells (Demb and Singer 2012).

Here, we demonstrate that oscillations in the rd1 retina arise from intrinsic membrane properties of AIIIs; they are not an emergent property of a network of coupled neurons with heterogeneous intrinsic electrical properties, as has been suggested (Trenholm et al. 2012). Specifically, we tested the hypothesis that AIIIs in the rd1 retina are hyperpolarized relative to their wild-type counterparts and that this hyperpolarization permits an interaction between a fast, depolarizing Na conductance and a slow, hyperpolarizing, M-type K conductance to generate phasic bursting, just as it does in the wild-type retina (Cembrowski et al. 2012).

A model of the AII (Cembrowski et al. 2012) allowed us to predict effects of M-type K current activators and inhibitors on oscillations observed experimentally during whole cell recordings from AIIIs in a retinal slice preparation. Pharmacologically induced alterations in AII activities were propagated unchanged to ganglion cells, as revealed by whole cell recordings from individual cells and by multielectrode array recordings of population activity. We conclude that the intrinsic properties of rd1 and wild-type AIIIs need not differ; rather, hyperpolarization arising from reduced light-dependent synaptic input is sufficient to unmask bursting. Electrical coupling is relevant only because it affects the AII hyperpolarization. Our finding indicates that treatment of retinal degeneration might benefit from preventing hyperpolarization of AIIIs. Inducing light sensitivity in ON bipolar cells (see Lagali et al. 2008) may accomplish this.

MATERIALS AND METHODS

Electrical recordings from in vitro retinal preparations. C3HeJ mice (rd-1/rd-1; RD1), which carry a mutation in the Pde6b gene encoding the β-subunit of cGMP-phosphodiesterase and lose the outer nuclear layer by 4 wk of age, were obtained from the Jackson

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Laboratories (Bar Harbor, ME) and killed at postnatal days 40-50, when their retinas were not responsive to light due to the loss of photoreceptors. Mice were anesthetized by isoflurane inhalation and decapitated before enucleation, as approved by the Animal Care and Use Committee at the University of Maryland. Retinas were isolated into cooled (-15°C) bicarbonate-buffered Ames’ medium (Sigma) equilibrated with 95% O2-5% CO2 (Carbogen). Whole cell recordings were made from retinal amacrine cells in retinal slices and from ganglion cells in retinal whole mount preparations (e.g., Margolis et al. 2014).

For retinal slice preparation, retinas then were embedded in low-melting temperature agarose (Sigma type VIIA, 3% in a HEPES-buffered saline), and slices (300 μm) were cut on a vibrating microtome (Leica). Slices were stored in Carbogen-bubbled Ames’ medium at room temperature until use. For retinal whole mount preparation, a ~2-mm square of retinal tissue was mounted flat, photoreceptor-side down, onto Anodic filter paper (Whatman, Florham Park, NJ). Both retinal slices and whole mounts were placed in a recording chamber mounted below an upright video-microscope (Zeiss) so that cells of interest could be visualized and targeted for whole cell recordings with pipettes containing the following (in mM): 95 K gluconate, 15 KCl, 5 NaCl, 10 HEPES, 0.2 EGTA, 8 Tris-phosphocreatine, 4 MgATP, and 0.4 NaGTP. For ganglion cells, alpha cells were targeted based on soma size (diameter 18–25 μm). Access resistances were <25 MΩ and were not compensated. For extracellular recording of ganglion cell spiking, a ~2-mm square of retinal tissue was mounted flat, ganglion cell-side down, onto a 60-channel multielectrode array (MEA; square array; 30-μm electrodes with 200-μm spacing; Multichannel Systems).

In all cases, during recordings, the tissues were super-fused with Ames’ medium (~34°C) to which drugs were added as noted. Picrotoxin (100 μM), strychnine (0.5 μM), dinitroquinoline-2,3-dione (DNQX; 25 μM), and APV (50 μM) were added to block GABA receptor-, glycine receptor-, AMPA/kainate receptor-, and NMDA receptor-mediated currents, respectively. L- (+)-2-amino-4-phosphonobutyric acid (l-AP4; 2–10 μM) was added to activate mGluR6 receptors on ON bipolar cells. Tetrodotoxin (TTX; 0.5 μM) was added to block voltage-gated Na channel-mediated currents, linopirdine hydrochloride (LP; 30 μM) was added to block M-type K currents, flupiritine maleate (flu; 10 μM) added to block voltage-gated Ca currents, and melaminophenol (MPA; 100 μM) was added to block the long neurite branching (Schmitzer-Torbert et al. 2005) (see Fig. 4Ai and Aii). The morphologically detailed model was built based on an AII filled with a fluorescent tracer (see Fig. 4Aii). Na, A-type K, and slowly activating M-type K conductances were inserted into the distal varicosity; the long neurite and the soma were taken to be passive, and the rest of the cell had a low-density A-type K conductance (Cembrowski et al. 2012). The previous study has shown that, in the morphologically detailed model, voltage responses to a brief current injection at the distal varicosity were very similar when measured in the soma, the lobules at the end of the primary dendrite, and the arboreal dendrite (Cembrowski et al. 2012). Thus, excepting the distal varicosity that was shown to be the spike-initiation site (Cembrowski et al. 2012) and the attached long neurite, when spiking the rest of the AII behaves as if it is electrotonically compact. This justifies treating the soma and arboreal dendrite as a single compartment.

Specifically, our AII model consisted of a large compartment (length = 25 μm, diameter = 25 μm) representing the soma and arboreal dendrite of the AII, a thin cable-like compartment (length = 32 μm, diameter = 0.3 μm) that represents the long neurite branching from the primary dendrite, and the small initiation site (length = 2 μm, diameter = 2 μm; see Fig. 4Aii). For convenience, we will call the large compartment the “soma,” the thin cable compartment the “cable,” and the small varicosity compartment the “initiation site.” Unlike in Cembrowski et al. (2012), here we treated the cable as 1 compartment instead of 11 compartments, after confirming that this change does not make a significant difference. In all three compartments, there was a passive leak conductance, with membrane resistivity of 40,000 Ω cm². The specific membrane capacitance and axial resistivity were 1 μF/cm² and 150 Ω/cm respectively. In addition, the initiation site had a fast Na conductance (gIS,Na = 0.2 S/cm²), an incompletely inactivating A-type K conductance (gIS, A = 0.08 S/cm²), and a slow noninactivating M-type K conductance (gIS,M = 0.03 S/cm²), as this single distal initiation site has been shown to be the site of spike generation as well as of its slow modulation (Cembrowski et al. 2012). Although the soma itself was found to be passive (Tian et al. 2010), since the somatic compartment of the three-compartment model was taken to include arboreal dendrites, an A-type K conductance was included in this compartment (gIS, A = 0.004 S/cm²). These active conductances, with the reversal potentials $E_{Na} = 50$ mV and $E_{K} = -77$ mV, had the following kinetics. The Na current was given by

$$i_{Na} = \frac{g_{IS,Na} h_{IS}(V - E_{Na})}{(1 + \exp\left(\frac{V - V_{1/2, Na}}{k_{Na}}\right))^{-1}}$$

where the activation variable and the inactivation variable were governed by

$$\tau_{m_{Na}} \frac{dm_{Na}}{dt} = m_{Na} - m_{Na},$$

$$\tau_{h_{Na}} \frac{dh_{Na}}{dt} = h_{Na} - h_{Na},$$

with

$$m_{Na}(V) = \left[1 + \exp\left(-\frac{V - V_{1/2, m_{Na}}}{k_{m_{Na}}}\right)\right]^{-1},$$

$$h_{Na}(V) = \left[1 + \exp\left(-\frac{V - V_{1/2, h_{Na}}}{k_{h_{Na}}}\right)\right]^{-1},$$

$$V_{1/2, m_{Na}} = -48 \text{ mV}, k_{m_{Na}} = 5 \text{ mV},$$

$$V_{1/2, h_{Na}} = -49.5 \text{ mV}, k_{h_{Na}} = 2 \text{ mV}.$$
The M-type K current was given by

\[ i_M = g_M m_M(V - E_K) \]

where the activation variable was described as

\[ \tau_{m_M} \frac{dm_M}{dt} = m_{\infty} - m_M \]

with

\[ m_{\infty}(V) = \left[1 + \exp\left(-\frac{(V - V_{1/2,m})}{k_{m_M}}\right)\right]^{-1}, \]

\[ V_{1/2,m} = -40 \text{ mV}, k_{m_M} = 4 \text{ mV}. \]

The M-type K current is noninactivating and activates slowly with a time constant \( \tau_{m_M} = 50 \text{ ms} \).

Finally, the A-type K current was given by

\[ i_A = g_A m_A \left[ch_A^{(1)} + (1 - c) h_A^{(2)}\right](V - E_K), \]

\[ c = \left[1 + \exp\left(-\frac{(V + 45)}{15}\right)\right]^{-1}, \]

where the activation and inactivation variables were governed by

\[ \tau_{m_A} \frac{dm_A}{dt} = m_{\infty} - m_A, \]

\[ \tau_{h_A} \frac{dh_A}{dt} = h_{\infty} - h_A, \]

\[ \tau_{h_A}^{(1)} \frac{dh_A^{(1)}}{dt} = h_{\infty}^{(1)} - h_A^{(1)}, \]

\[ \tau_{h_A}^{(2)} \frac{dh_A^{(2)}}{dt} = h_{\infty}^{(2)} - h_A^{(2)}, \]

with

\[ m_{\infty}(V) = \left[1 + \exp\left(-\frac{(V - V_{1/2,m})}{k_{m_A}}\right)\right]^{-1}, \]

\[ V_{1/2,m} = -10 \text{ mV}, k_{m_A} = 7 \text{ mV}. \]

\[ h_{\infty}(V) = f \left[1 + \exp\left(-\frac{(V - V_{1/2,h})}{k_{h_A}}\right)\right]^{-1} + (1 - f), \]

\[ V_{1/2,h} = -40.5 \text{ mV}, k_{h_A} = 2 \text{ mV}, f=0.83. \]

The A-type K current had a relatively fast activation with \( \tau_{m_A} = 1 \text{ ms} \) along with a fast inactivation \( h_A^{(1)} \) and a slow inactivation \( h_A^{(2)} \). Inactivation was incomplete as captured by \( f \) in the steady-state expression above, with time scales

\[ \tau_{h_A}^{(1)}(V) = 25 - 20 \left[1 + \exp\left(-\frac{(V + 35)}{6}\right)\right]^{-1}, \]

\[ \tau_{h_A}^{(2)}(V) = \min\left\{\frac{(V + 17)^2}{4} + 26, 100\right\}. \]

The parameters for the A-type K conductance were well constrained by voltage-clamp recordings (Tian et al. 2010), while other parameters were fit to reproduce the experimentally observed spikelets and bursts in wild-type AII (Cembrowski et al. 2012). In this three-compartment AII model, we omitted an L-type Ca conductance that is present in AII cells (Habermann et al. 2003) since it was not needed to reproduce realistic AII responses in the regime of interest. For the first set of simulations (see Fig. 4), the model consisted of this three-compartmental AII in isolation, showing that a single AII is capable of generating tonic spiking and bursting. We adjusted the leak reversal potentials of the AII to reproduce the experimentally recorded membrane potentials for the wild-type and rd1 AII. We used the leak reversal potential of −10 mV to put the AII in the tonic spiking state.

Fig. 2. A model based on gap junctions between heterogeneous AIIss does not explain the MFA-induced frequency decrease. The model in Trenholm et al. (2012), which designates the gap-junction coupled network of heterogeneous AIIss as the key to the rd1 oscillations, does not explain the experimental observed frequency decrease under MFA application. The model consists of 2 slightly heterogeneous AII cells with Na, delayed rectifier K, and leak conductances, and an ON cone bipolar (ONCB) cell with a hyperpolarization-activated current and a leak current. All cells have a single compartment and are electrically coupled to each other. As the coupling strength decreases, mimicking the effect of MFA, the oscillation amplitude decreases, but the oscillation frequency increases in contrast to our experiments with MFA (Fig. 1B). All the parameters used here are the same as in Trenholm et al. (2012).
regime, while using −50 mV to put the AII in the bursting regime, capturing the rd1 AII being relatively hyperpolarized compared with the wild-type AII.

For the rest of the simulations (see Figs. 5–8, 13, and 14), an ONCB cell was electrically coupled to the three-compartmental AII cell. The ONCB cell was modeled as a passive single compartmental cell with surface area of 440 μm² and membrane resistivity 12,000 Ω cm². The ONCB model was coupled to the “soma” compartment of the AII model, capturing its coupling to the AII arboreal dendrite, via a gap junction with a conductance of 750 pS. This is in the range of reported coupling strengths, which spans the range from 100 to 1,500 pS for type 5, 6, and 7 ONCB cells (Veruki and Hartveit 2002). The leak reversal potentials for the AII and the ONCB cell in this coupled AII-ONCB model were adjusted to −65 and −35 mV, respectively, to reproduce the experimentally observed membrane potentials in rd1 retinas.

RESULTS

MFA first slows down and then eliminates oscillations in rd1 retinas. It has been reported previously that the gap junction blocker MFA eliminates the oscillations in rd1 retina (Menzler and Zeck 2011; Trenholm et al. 2012). Based on this observation, it has been suggested that oscillations are an emergent property of the electrically coupled network of AII amacrine and ONCB cells. More specifically, in a model network of AIIIs with slightly heterogeneous properties, electrical coupling could cause the AIIIs to enter an oscillatory state even if the individual AIIIs are not intrinsically oscillatory (Trenholm et al. 2012).

To characterize in more detail the effect of MFA on the AII network, we recorded membrane voltages of AII amacrine cells of rd1 mice. Our recordings confirmed spontaneous oscillatory activity in the AIIIs of rd1 mice (Trenholm et al. 2012; Yee et al. 2012; Menzler and Zeck 2011; Stasheff 2008), with frequencies in the range 5–15 Hz in agreement with previous studies (Trenholm et al. 2012; Yee et al. 2012; Menzler and Zeck 2011; Stasheff 2008; Margolis et al. 2014). We then applied MFA and measured the oscillation frequency and the membrane potential of rd1 AIIIs (n = 4). In our experiments, MFA application indeed was followed by elimination of the oscillations (Fig. 1A). During the onset of the MFA effect, oscillations gradually decreased in frequency as the AIIIs hyperpolarized. During MFA washout, oscillation frequency increased slowly until the original frequency was restored, and the mean AII membrane potential increased at the same time. Figure 1B shows that oscillation frequency varied directly with membrane potential.

Next, we investigated whether such behavior during MFA application could be explained by the model put forward in Trenholm et al. (2012). In this network model, which consists of two AIIIs with heterogeneous intrinsic electrical properties and one ONCB cell, it was shown that reducing the gap junction conductances is followed by elimination of the oscillations. Based on this simulation, an emergent property of a network of heterogeneous AIIIs has been suggested as the source of the oscillations. In this model, the two AIIIs are electrically coupled to one another and the ONCB cell is electrically coupled to one of the AIIIs. The ONCB cell has a hyperpolarization-activated cationic current, a delayed rectifying K current, and a leak current. Each AII has a fast Na conductance, a delayed rectifying K conductance, and a leak conductance. Importantly, the AIIIs are mildly heterogeneous in their maximal Na and leak conductances (for details, see Trenholm et al. 2012) (Fig. 2).

![Fig. 3. Bursting in the hyperpolarized wild-type (WT) AII induces rd-like oscillations in the AII-cone bipolar network. A: representative intracellular rd1 and WT AII recordings. Ai: oscillations in rd1 AII without hyperpolarizing current injection. Aii: WT AII that exhibits spontaneous spikelets at rest (up to t = 300 ms) undergoes a transition to bursting when it is hyperpolarized (after t = 300 ms). B: under tetrodotoxin (TTX), the resting membrane potentials of WT AIIIs (n = 4) are more depolarized than those of rd1 AIIIs (n = 6).](http://jn.physiology.org/doi/10.1152/jn.00437.2014)
We decreased the gap junction coupling strength in the model of Trenholm et al. (2012) and observed that the oscillation amplitude decreased and eventually the oscillations stopped. The oscillation frequency, however, increased during this process (Fig. 2), which is inconsistent with the experimentally observed decrease induced by MFA (Fig. 1). The marked inconsistency between the behavior of the model of Trenholm et al. (2012) and the neuronal behaviors that we observed experimentally motivated us to develop an alternative model that would describe experimentally measured parameters of rd1 oscillations.

AII oscillations resemble cable-filtered intrinsic bursting in AII. Previous electrophysiological and computational analyses revealed that the small somatic spikelets recorded in AII are attenuated versions of full action potentials initiated in an electrically distal compartment (Cembrowski et al. 2012). This referenced computational analysis made use of a multicompartment model; single-compartment models do not describe spiking in AII adequately (Apollo et al. 2013).

The multicompartment model in Cembrowski et al. (2012) comprised a soma, a single spiking process, and the remainder of the dendritic tree (represented as many individual compartments) as well as experimentally constrained membrane conductances (Fig. 4Ai). Because the Na conductance was limited to a single, distal spike initiation site, full action potentials at this site induced only small voltage deflections (“spikelets”) in the soma. Experimentally observed bursting in hyperpolarized AII was explained by interactions between the Na conductance and a slow, M-type K conductance that suppressed the spiking: in the bursting regime, fast spiking generated by the Na current slowly activates the M-type K current, which in turn hyperpolarizes the neuron, terminating the spiking, and completing a burst.

Because voltages in the dendrites and in the soma were more negative than those of the wild-type AII (−50.2 ± 2.1 mV, n = 6 vs. −42.3 ± 4.9 mV, n = 4; two-valued t-test, P < 0.015), as shown in Fig. 3B.

AII are hyperpolarized in the rd1 retina. Upon closer inspection of the voltage waveforms recorded from rd1 AII, small spikelets were apparent during the depolarized phase of the oscillation (Fig. 3Ai, inset) (also see Borowska et al. 2011). These oscillations were strikingly similar to those we observed previously following mild hyperpolarization of wild-type AII (Cembrowski et al. 2012) (Fig. 3Aii).

Because hyperpolarization of the membrane potential was necessary to induce bursting in wild-type AII, we postulated that the somatic voltages of AII in rd1 retinas would be hyperpolarized relative to their wild-type counterparts. Because the wild-type AII were spontaneously spiking (within the range of −45 to −40 mV, n = 6, also see Cembrowski et al. 2012) (Fig. 3Aiii, before the hyperpolarizing current injection) and the rd1 AII were spontaneously oscillating (in the range of −65 to −50 mV, n = 6; Fig. 3Ai), it was not possible to compare their resting potentials directly. Therefore, we compared the resting potentials of wild-type and rd1 AII in the presence of TTX: although there was notable variability, the resting potentials of the rd1 AII under TTX were significantly lower than those of wild-type AII (−50.2 ± 1.1 mV, n = 6 vs. −37.1 ± 0.7 mV, n = 4; two-valued t-test, P < 0.015).

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Because voltages in the dendrites and in the soma were predicted to be similar during bursting, we reduced the multicompartment model to a three-compartment model by lumping the soma and the dendrite into a single compartment (Fig. 3Ai). Before the hyperpolarizing current injection and the rd1 AII were spontaneously oscillating (in the range of −65 to −50 mV, n = 6; Fig. 3Ai), it was not possible to compare their resting potentials directly. Therefore, we compared the resting potentials of wild-type and rd1 AII in the presence of TTX: although there was notable variability, the resting potentials of the rd1 AII under TTX were significantly lower than those of wild-type AII (−50.2 ± 1.1 mV, n = 6 vs. −37.1 ± 0.7 mV, n = 4; two-valued t-test, P < 0.015).

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Fig. 4. Simulations showing tonic spiking and bursting of an AII. Ai: confocal image of an AII (left) and the morphologically detailed model (right). Aii: reduced 3-compartmental model of the AII consists of an “initiation site” (IS) and a “cable” (C) corresponding to the thin neurite branching from the primary dendrite of the AII (arrow in Ai), as well as a “soma” (S), which is taken to represent the rest of the AII (soma and dendrites). Bi: tonic spiking in the IS leads to small spikelets in the soma as well as a “soma” (S), which is taken to represent the rest of the AII (soma and dendrites). Bii: tonic spiking in the IS leads to small spikelets in the soma as well as a “soma” (S), which is taken to represent the rest of the AII (soma and dendrites).
oscillations in the soma that included small spikes during the depolarized phase (Fig. 4Bii).

The similarities between the voltage waveforms in the rd1 AII and the hyperpolarized wild-type AII led us to the hypothesis that oscillations in the rd1 retina reflect the intrinsic bursting activity of AIIIs rather than some emergent property of an electrically coupled network. Below, we tested this hypothesis by manipulating gap junctional and K conductances in AIIIs and determining whether the experimental observed effects were consistent with our model of the AII.

The model reproduces the effects of MFA on bursting in the rd1 retina. A first test of this new model is whether it correctly reproduces the slowing down and elimination of bursting due to MFA application. MFA affects the AII-ONCB network beyond blocking the coupling between AIIIs; it also reduces coupling between AIIIs and ONCB cells (Pan et al. 2007). Blocking electrical coupling between AIIIs and ONCB cells in the rd1 retina should hyperpolarize AIIIs because these ONCB cells are depolarized ($V_m$ oscillating around $-50$ mV, Borowska et al. 2011) relative to AIIIs ($V_m$ oscillating around $-65$ to $-50$ mV; Fig. 3Ai). Our model predicts that additional hyperpolarization of bursting AIIIs will suppress bursting (Fig. 5A). Such behavior mimics our experimental results (Fig. 1).

To test this interpretation further, we extended our model to include gap-junction coupling to a passive ONCB cell. We simulated the effect of MFA by reducing the coupling strength between the AII and the ONCB cell. As expected, weakening the coupling strength between the AII and the ONCB cell hyperpolarized the AII, which in turn decreased the oscillation frequency and then eliminated oscillations in AIIIs, in agreement with our experiments (Fig. 5B). A similar result has been reported in Trenholm et al. (2012). If MFA stops oscillations by hyperpolarizing intrinsically oscillating AIIIs, then, it should be possible to restore the oscillations by injecting depolarizing current injection following MFA application restores oscillations in rd1 AII. A: experimental recording from an rd1 AII in which a depolarizing current injection of 15 pA restored oscillations that had been silenced by MFA application. B: simulation of a depolarizing current injection of 5 pA to the model rd1 AII, with its gap junction coupling to the ONCB reduced to 100 pS to mimic the effect of MFA. The oscillations are eliminated by a reduction in the gap junction coupling (Fig. 5B) and are restored by the depolarizing current injection. The restored oscillations have the typical waveform (bottom).
current into the AII following the application of MFA. Indeed, somatic current injections of \( \sim 15-20 \) pA restored bursting in AIIIs in both the experiments (Fig. 6A; \( n = 4 \)) and the model (Fig. 6B).

The model accurately predicts the effects of K channel manipulation on bursting in the rd1 retina. Our model posits a central role for a M-type K conductance in bursting. Therefore, we tested the effects of potentiating and blocking M-type K conductances on bursting in AIIIs of the rd1 retina to determine whether the effects were consistent with the predictions of our model. First, we tested the effect of the M-current activator flupirtine. Flupirtine is known to activate the M-type K current by shifting its activation to more negative voltages (Wu et al. 2012).

During the washin of flupirtine, oscillations first decreased in frequency and then ceased (Fig. 7, Aii and Aiii). This pattern was reversed upon flupirtine washout: silenced AIIIs first began to oscillate at low frequency, and then oscillation frequency rose. Concomitant with the change in frequency, the waveform of the oscillations evolved (Fig. 7Ai). To quantify the change in the waveform, we measured the width of the bursts at a membrane potential that corresponded to 30% of the maximal oscillation amplitude in each sweep (1,667-ms long). As Fig. 7, Aiii and Aiv, shows, the bursts became narrower during the washin of flupirtine (from 79.3 ± 4.5 to 44.9 ± 9.5 ms). The effect due to flupirtine was found to be significant (\( P < 10^{-3} \) by two-way ANOVA). Biophysically, with the left shift of the activation curve of the M current fewer spikes are required to generate an outward current sufficient to shut off the spiking.

We also simulated the effect of flupirtine in our model AII by shifting the half-activation voltage (\( V_{1/2} \)) of the M-type K current to more negative voltages. Consistent with the experiments, this manipulation decreased bursting frequency and then silenced the cell (Fig. 7B). While the frequency generally decreased as \( V_{1/2} \) of the M current became more negative, there were also discrete jumps around \( V_{1/2} \) = -42 mV and \( V_{1/2} = \)

![A](image1.png)  
![B](image2.png)

Fig. 7. Opening M current reduces frequency and burst length and stops oscillations. A: experiment during flupirtine washin on an rd1 AII. Ai: representative traces of the AII membrane potential during flupirtine washin. Aii: spectrogram of the recording in Ai shows that the frequency decreased under flupirtine application, eventually resulting in termination of the oscillations. Aiii: burst length at the end of the flupirtine washin (final) is significantly smaller than at the beginning (initial) (\( n = 3 \); \( P < 10^{-3} \)). Aiv: burst length during the flupirtine washin of a representative recording. The length was measured at 30% of the maximal oscillation amplitude for each 1,667-ms-long sweep (dotted line in Ai). “Initial” and “Final” indicate where the initial and final burst lengths were measured. B: simulation mimicking flupirtine application by moving the half-activation voltage \( V_{1/2} \) of the M conductance leftward. Bi: representative membrane potential traces for the model AII. Along with the frequency decrease, the number of spikes per burst decreases as well, resulting in a burst length decrease. Bii: in the model the burst frequency decreases as the half-activation voltage \( V_{1/2} \) of the M conductance is reduced. The discrete jumps correspond to reduction in the number of spikes per burst. Eventually the AII becomes quiescent.
−45 mV (Fig. 7Bi). These jumps occurred due to a reduction in the number of spikes per burst. In the experiments the individual bursts were too variable to allow the identification of such details (cf. Fig. 3Ai). As in the experiment, however, the burst width decreased (Fig. 7Bi).

To determine how antagonizing the M-type K channel affects the rd1 oscillations, we used LP (see Cembrowski et al. 2012). Blocking the M current reduced the burst frequency (Fig. 8, Ai and Aii; n = 5). The burst duration, however, increased from 87.4 ± 41.1 to 175.0 ± 34.7 ms; this is opposite to the effect of flupirtine on the burst duration.

Despite its marked variation across the recorded cells, the increase was highly significant (P < 10⁻³; Fig. 8, Aiii and Aiv). In agreement with this observation, the computational model also generated lower burst frequencies and longer burst durations when the maximal M conductance was reduced (Fig. 8B).

This was expected, since the M current has been identified as being responsible for terminating spiking (Cembrowski et al. 2012).

In the model, decreasing the M current caused the burst duration to increase without bound and eventually change to tonic spiking. In the experiments, however, LP did not eliminate all oscillations and it did not induce tonic spiking. Instead, when the burst duration increased beyond ~200 ms, a transition to a different type of low-frequency (~1 Hz) oscillation occurred (Figs. 8Ai, bottom, and 9B). Riding on top of these slow oscillations were faster, irregular, 5- to 10-Hz bursts of spikes. These bursts were reminiscent of the characteristic rd1 oscillations that we observed in the absence of LP. This phenomenon was also observed when ONCB cells first were hyperpolarized by application of t-AP4, which on its own does not block oscillations in the rd1 retina (Borowska et al. 2011) (n = 6). Because of this transition to a qualitatively different type of oscillation, the measurement of the increase in the burst duration due to LP depended on the identification of the
transition time. We used the sudden increase in the variability of the duration as a criterion for that transition.

The slow-frequency oscillation resembled both in frequency and shape the slow, Ca channel-mediated oscillations that have been observed in ONCB cells of rat, mouse, and goldfish retina when inhibition was blocked (Ma and Pan 2003; Yee et al. 2012; Zenisek and Matthews 1998). To test whether the slow oscillations were indeed Ca channel mediated, we applied cadmium (Cd^{2+}) and nickel (Ni^{2+}), two nonselective voltage-gated Ca^{2+} channel blockers, after the slow oscillations induced by LP had been established (in the presence of L-AP4; n = 4). The characteristic shape and frequency of the slow oscillations induced by LP were disrupted by Ni^{2+} and Cd^{2+}, supporting our hypothesis that the slow oscillations that appear under LP application require Ca channels and therefore arise from a mechanism that differs from that underlying the ~10-Hz oscillations (Fig. 9).

Thus either blocking or potentiating an M-type K conductance eliminates the characteristic 5- to 10-Hz oscillations in rd1 AII cells (Figs. 7 and 8). The observed changes in oscillation frequency and burst duration are predicted by our model of a single AII. This suggests that the slow M-type K current plays the same key role in generating the rd1 oscillations as it does in generating bursting in wild-type AII.

Note that the model investigated in Trenholm et al. (2012) does not include an M current and does therefore not allow any predictions pertaining to these experiments.

Intrinsirc oscillations in AIIIs are propagated unaltered to ganglion cells of the rd1 retina. In rd1 retina, the 5- to 15-Hz oscillations are also observed in retinal ganglion cells (RGCs). These oscillations reflect rhythmic synaptic input (Margolis et al. 2008; Borowska et al. 2011; Menzler and Zeck 2011; Stasheff 2008; Ye and Goo 2007; Yee et al. 2012), and anticorrelation of activity in ON and OFF alpha RGCs implicates the AII as a major contributor to this synaptic input (Margolis et al. 2014). Here, having found evidence to support the hypothesis that the rd1 oscillations in AIIIs are intrinsically generated by their M-type K conductances, next we investigated whether these AII oscillations are indeed the origin of the RGC oscillations by testing the effects of LP and flupirtine on the oscillations in RGCs in the rd1 retina. The oscillations were measured with intracellular recordings of single alpha ganglion cells in the rd1 whole mount retina and by extracellular recordings of single and multuniits via MEA.

Both flupirtine and LP disrupted oscillatory synaptic currents and membrane potentials recorded in the whole cell configuration in ON and OFF alpha RGCs. Observed changes in frequency and burst duration were consistent with those of AII oscillations; flupirtine eliminated the oscillations following a decrease in both frequency and burst duration, and LP decreased the bursting frequency and increased the burst duration before eliminating the rd1 oscillations. Representative traces of ganglion cell membrane potentials along with their frequency and burst duration changes under flupirtine and LP are shown in Figs. 10, Aii, and Aiv, and Bi, Bii, and Biv, respectively. Figure 10, Biii and Biv, shows the collective data of burst duration changes under flupirtine and LP. In retinas treated with LP (n = 4), the width of ganglion cell bursting increased (from 38.5 ± 27.8 to 305.3 ± 108.5 ms, P < 10^{-3}). In three out of four cells under flupirtine, the burst widths decreased (from 51.7 ± 12.4 to 17.4 ± 8.4 ms, P < 10^{-3}). In one case, application of flupirtine was followed by almost instant elimination of the oscillations, so it was not possible to observe slow changes in the burst duration. In addition, in three out of four cells treated with flupirtine, the bursting frequency became irregular before the cells became quiescent. However, in all cases, the spikes became less frequent after flupirtine application.

In the extracellular electrode recordings, we studied the oscillations present in each single unit as well as in the multiunit activity (MUA). Both single and multiunit activities agreed with the changes in AII oscillations under LP application. A representative single unit activity is shown in Fig. 11A. It is apparent that the oscillations under LP are slower than those in L-AP4 and washout conditions. Similarly, the interspike intervals show a nonzero peak at a much higher value (810 ms) under LP than under L-AP4 or washout. Figure 11B shows the comprehensive bursting statistics of single units. Under L-AP4 (n = 61), the single unit burst lengths had a median value of 49 ms with an interquartile range (IQR) of 45–59 ms. The frequency of these oscillations had a median 6.2 Hz with IQR 4.8–6.7 Hz. When LP was applied, the burst lengths of the single units (n = 18) increased to 64 ms and their IQR ranged from 40 to 120 ms. The frequency of these oscillations decreased to a median of 1.37 Hz with an IQR of 1.08–1.6 Hz, consistent with what we observed from AIIIs under LP. Then, for the washout condition, the oscillation frequencies of the single units (n = 34) increased back to near the values found for the initial L-AP4 condition, and the mean burst length decreased to 51 ms, with IQR = 39–67 ms.

We also inspected the effects of LP on MUA. The spectrograms of the MUA in Fig. 12A show that LP application slowed down the MUA oscillations. Based on the MUA, oscillating channels under the L-AP4 condition (n = 52) had a median oscillation frequency of 7.5 Hz and an IQR of 4.8–9.6 Hz. For the LP condition (n = 70), the oscillating channels had a median frequency of 1.75 Hz and IQR of 1.2–4.8 Hz. Then, for the washout condition (n = 66), we observed that the oscillation frequencies increased back towards their initial values (Fig. 12B).

Fig. 9. Blocking Ca^{2+} channels disrupts the slow-frequency oscillations. A: representative recording of an rd1 AII under LP and Ca^{2+} blockers. The characteristic rd1 oscillations, which were unaffected by L-(+)-2-amino-4-phosphonobutyric acid (L-AP4; A), were disrupted by LP and replaced by slow-frequency oscillations (B). The slow-frequency oscillations were eliminated by Ni^{2+} and Cd^{2+} (C).
The extracellular recordings were also made under flupirtine (2 experiments), and we observed either complete elimination of the oscillations or a decreased bursting frequency. However, we do not report extracellular recording summary statistics here for the bursting activity of RGCs under flupirtine, because we did not isolate a statistically significant number of single units for this condition ($n$). In conclusion, intracellular recordings, single unit extracellular recordings, and multiunit extracellular recordings all show elimination of the oscillatory activities in RGCs under LP and flupirtine. Moreover, the changes in burst frequency and burst duration in RGCs agree with those in AIIs, providing strong evidence that the oscillations in RGCs indeed come from the upstream AII cells in the rd1 retina.

**DISCUSSION**

A combined experimental and computational approach was used to test the hypothesis that oscillations of AIIs in the rd1 retina are intrinsic and the result of interactions between fast, depolarizing Na and slow, hyperpolarizing, M-type K conductances. In this scenario, the oscillations arise as the result of spontaneous bursting in a distal compartment, generated because rd1 AIIs are hyperpolarized relative to wild-type AIIs. We analyzed a compartmental model of the AII (Cembrowski et al. 2012) to predict the expected electrophysiological responses of AIIs to pharmacological manipulation of M-type K currents and performed these experiments. The combined computational and exper-

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**Fig. 10.** The intracellularly recorded rd1 oscillations in retinal ganglion cells (RGCs) change similarly to those in AIIs under M-current activator and blocker. **A:** rd1 oscillations in RGCs were disrupted in the presence of flupirtine. **Ai:** representative RGC membrane potential under flupirtine. Flupirtine decreased oscillation frequency (Aii) and decreased burst duration (Aiv). Aiii: initial and the final burst lengths during flupirtine washin show a significant decrease ($n = 3, P < 10^{-3}$). **B:** rd1 oscillations in RGCs were disrupted in the presence of LP. **Bi:** representative RGC membrane potential under LP. LP induced a decrease in oscillation frequency (Bii) and increase in burst duration (Biv). The burst durations were averaged for each 5-s-long sweep. Biii: initial and the final burst lengths during LP washin show a significant increase ($n = 4, P < 10^{-3}$).
Experimental results are fully consistent with our hypothesis. Additionally, single cell and multielectrode array recordings from ganglion cells demonstrated that altered AII activity was propagated unchanged to ganglion cells. Our conclusion is that the oscillatory output of ganglion cells in the rd1 retina reflects the intrinsic properties of AII amacrine cells.

Mechanisms underlying bursting in neurons in the rd1 retina. Spontaneous oscillatory activity in interneurons of the rd1 retina has been documented in a number of studies (Borowska et al. 2011; Margolis et al. 2014; Toychiev et al. 2013; Trenholm et al. 2012). Because blocking electrical transmission among AII and between AII and ONCB cells inhibits
it, this activity has been characterized as an emergent property of a network of heterogeneous neurons (Trenholm et al. 2012).

Our study, however, supports a different conclusion: oscillations in the rd1 retina are explained largely by the intrinsic properties of AII amacrine cells, which appear to rest at potentials hyperpolarized relative to those observed in wild-type AII (Fig. 3). A number of experimental results, predicted by our compartmental model of the AII, support this assertion. In particular, the burst waveforms of AII in the rd1 retina closely resemble bursts in wild-type AII elicited by hyperpolarization (Fig. 3). Our compartmental model reproduces these bursts, and this allowed us to make predictions about the characteristics of bursting, specifically about the role played by M-type K currents, that then could be tested experimentally.

Our model suggested that potentiating the M current would first reduce burst duration and decrease oscillation frequency and then eliminate the oscillations and that blocking the M current would increase burst duration and decrease oscillation frequency; these predictions were borne out by our experiments (Figs. 7 and 8). The corresponding effects of pharmacological manipulation of M currents were also observed in ganglion cells (Figs. 10–12), indicating that activity in AII is propagated unchanged to ganglion cells.

Bursting by AII is generated intrinsically in our model, which suggests that hyperpolarization of AII in rd1 mice may be the primary cause of oscillations. Because our model was developed in a study of wild-type AII (Cembrowski et al. 2012), we suggest that AII in rd1 retinas are functionally similar to their wild-type counterparts. Consistent with this notion, TTX blocks bursting in rd1 AII, as it does in wild-type AII (Cembrowski et al. 2012; Trenholm et al. 2012; Margolis et al. 2014). The lower concentration of TTX used in Menzler and Zeck (2011), however, did not eliminate the oscillations in the local field potential associated with oscillations in ganglion cells of the rd1 mouse.

Taken together, the rd1 oscillations are cable-filtered bursts generated by the interplay between the fast, spike-generating Na and the slowly activating M-type K conductances. When the AII are relatively depolarized as in the wild-type retinas, the cells exhibit spontaneous tonic spiking, while too much hyperpolarization brings the cells to the quiescent regime. However, in between, there is a range of

Fig. 13. FitzHugh-Nagumo type model of Ca\textsuperscript{2+} mediated slow oscillations in an ONCB coupled to an AII. In this simulation, $g_M$ decreases in a sigmoidal fashion mimicking LP application. The burst duration increases, and when it reaches a critical value, the waveform makes a transition from the characteristic rd1 oscillations to qualitatively different slow oscillations. The number of spikes per burst with decreasing $g_M$ increases as indicated.

Fig. 14. In the model, depolarizing current injection into an ONCB suppresses bursting in the coupled AII. A: passive ONCB is coupled to an AII cell with a coupling strength of 750 pS. B: current injection of 20 pA in the ONCB (Bi) depolarizes the ONCB (Bii) and eliminates bursting in the coupled AII (Biii and Biv).

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membrane potential that induces spontaneous bursting in the AII.

As others have observed (Trenholm et al. 2012), in our experiments an antagonist of gap junctions slowed and then blocked oscillations in the rd1 retina (Fig. 1); our model predicts this as well (Fig. 5). Moreover, our model further predicted, and our experiments confirmed, that bursting in individual AIIIs in the presence of MFA can be restored by injection of depolarizing current (Fig. 6). This supports that the oscillations arise from individual AIIIs, not from a network effect as previously suggested (Trenholm et al. 2012). Based on these results, we suggest that the main effect of MFA is hyperpolarization of AIIIs through elimination of their coupling to more depolarized ONCB cells. Thus our study supports the conclusion that homotypic electrical coupling between AIIIs does not contribute to bursting by AIIIs and that heterotypic electrical coupling between AIIIs and ONCB cells is relevant primarily because it affects the membrane potentials of AIIIs.

Slow Ca channel-dependent oscillations. We found that application of the M-current antagonist LP reduced oscillation frequency and lengthened the burst duration, consistent with our model (Fig. 8). LP, however, also induced slow (~1 Hz) oscillations in AIIIs that appeared to involve Ca channels (Fig. 9). The frequency and waveforms of bursts underlying these slow oscillations resembled Ca channel-dependent regenerative activity observed previously in isolated ONCB cells (Ma and Pan 2003) and in whole retinas when inhibition was blocked (Yee et al. 2012). We speculate that the enhanced burst duration in the presence of LP may allow activity in AIIIs to trigger such regenerative events in ONCB cells; in turn, the regenerative events in ONCB cells modulate the AII voltage.

To illustrate the feasibility of such a scenario, we extended our passive ONCB cell model to include a simple model of FitzHugh-Nagumo type (Fitzhugh 1961) to allow for regenerative events; active ONCB cells were coupled to model AIIIs (Choi 2014). In this simulation, decreasing the maximal M-type K conductance to mimic LP application generated a sudden transition from short bursts to much longer depolarizing events upon which spiking was superimposed (Fig. 13). We did not, however, assess whether this model captured all aspects of the experimentally observed transition to the slow oscillations induced by LP; while this is an interesting question, we considered it to be beyond the scope of the current work.

Clinical implications. A recent study demonstrates that MFA blocks the characteristic oscillatory activity of retinas of rd10 mice, which have a less severe retinal degeneration phenotype than rd1 mice, and improves signal transmission from surviving photoreceptors to ganglion cells (Toychiev et al. 2013). This finding is difficult to interpret, however, since gap junctions mediate interactions among many different classes of retinal neurons. Thus blocking all gap junctions may not be the most advantageous therapeutic intervention.

Our study indicates that MFA application eliminates oscillations not by uncoupling the AII network but by hyperpolarizing intrinsically active AIIIs. As a result, manipulating the resting membrane potential of AIIIs in some other manner might be an effective and benign way to restore signaling in a deteriorating retina. Introducing light sensitivity into bipolar cells (Lagali et al. 2008; Doroudchi et al. 2011) or stimulating them with retinal prostheses (Stingl et al. 2013) are possibilities that may be sufficient to reduce hyperpolarization of AIIIs and keep them from bursting.

In recent optogenetic experiments, introducing channel rhodopsin-2 expression in ON bipolar cells via electroporation (Lagali et al. 2008) or viral vectors (Doroudchi et al. 2011) has been successful in restoring light-induced responses both physiologically and behaviorally. Therefore, it will be interesting to examine how effective such light sensitivity is in suppressing the rd1 oscillations. To predict how much depolarization in ONCB cells is needed to suppress the oscillations, depolarization of the ONCB cell was simulated via current injections in our model of a passive ONCB cell electrically coupled to an AII cell with the coupling strength of 750 pS (Fig. 14). In our model, a current injection of 20 pA, which resulted in an ~20-mV depolarization in the coupled ONCB, suppressed the oscillatory activities in the AII and induced tonic spiking. Although variability in coupling strengths and number of coupled ONCB cells prevents us from predicting the precise amount of depolarization needed, this simulation predicts that depolarization of ~20 mV in ONCB cells is sufficient to eliminate the oscillations in the coupled AIIIs. Whether current methods of introducing light sensitivity in ON bipolar cells can bring the rd1 retina to the nonbursting regime would be an interesting topic for future work.

ACKNOWLEDGMENTS

We thank Rosemary Braun for many helpful discussions and Jiji Jiang for technical support.

GRANTS

This study was supported by National Eye Institute Grants EY-017836 (to J. H. Singer) and EY-021372 (to W. L. Kath, H. Riecke, and J. H. Singer). J. H. Singer was a Research to Prevent Blindness Special Scholar for Retinitis Pigmentosa.

DISCLOSURES

No conflicts of interest, financial or otherwise, are declared by the author(s).

AUTHOR CONTRIBUTIONS


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J Neurophysiol • doi:10.1152/jn.00437.2014 • www.jn.org


