

Received: 24 November 2017 Accepted: 21 December 2018 Published online: 12 February 2019

OPEN A biophysical model explains the spontaneous bursting behavior in the developing retina

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During early development, waves of activity propagate across the retina and play a key role in the proper wiring of the early visual system. During a particular phase of the retina development (stage II) these waves are triggered by a transient network of neurons, called Starburst Amacrine Cells (SACs), showing a bursting activity which disappears upon further maturation. The underlying mechanisms of the spontaneous bursting and the transient excitability of immature SACs are not completely clear yet. While several models have attempted to reproduce retinal waves, none of them is able to mimic the rhythmic autonomous bursting of individual SACs and reveal how these cells change their intrinsic properties during development. Here, we introduce a mathematical model, grounded on biophysics, which enables us to reproduce the bursting activity of SACs and to propose a plausible, generic and robust, mechanism that generates it. The core parameters controlling repetitive firing are fast depolarizing V-gated calcium channels and hyperpolarizing V-gated potassium channels. The quiescent phase of bursting is controlled by a slow after hyperpolarization (sAHP), mediated by calciumdependent potassium channels. Based on a bifurcation analysis we show how biophysical parameters, regulating calcium and potassium activity, control the spontaneously occurring fast oscillatory activity followed by long refractory periods in individual SACs. We make a testable experimental prediction on the role of voltage-dependent potassium channels on the excitability properties of SACs and on the evolution of this excitability along development. We also propose an explanation on how SACs can exhibit a large variability in their bursting periods, as observed experimentally within a SACs network as well as across different species, yet based on a simple, unique, mechanism. As we discuss, these observations at the cellular level have a deep impact on the retinal waves description.

Retinal waves, observed in many vertebrate species - chicks¹, ferrets², mice³, turtles⁴, macaques⁵ etc. are spontaneous bursts of activity propagating in the developing retina and playing a fundamental role in shaping the visual system and retinal circuitry. They emerge due to the conjunction of intrinsic single-cell properties (excitability and long refractoriness) and network interactions⁶. In the developing retina, waves evolve during three consecutive stages, mainly characterized by different types of synaptic transmission; gap junctions (stage I), acetylcholine (stage II) and glutamate (stage III)⁷ and are mediated by transient networks of specific cell types. An important type of such cells are the Starburst Amacrine Cells (SACs), which are involved in direction selectivity in the mature retina^{8,9} thanks to their special asymmetric synaptic distribution of excitatory and inhibitory synapses. However, during early development, these cells have a different role; they are responsible for eliciting stage II retinal waves via their spontaneous bursting behavior induced by autonomous intrinsic cellular mechanisms⁶. This activity disappears completely upon maturation⁶. During stage II, SACs constitute a transient network of autonomous bursters which, through cholinergic coupling, achieve opportunistic local synchrony. Under conditions yet to be identified, this local area of synchronized bursters can give rise to propagating waves. Although stage II retinal waves are quite thoroughly studied experimentally¹⁻³ as well as theoretically by several modeling approaches 10-13 (see Methods for a short review of these models and a comparison to ours) there are still important questions not yet addressed by existing studies. So far, previous models do not allow to understand how biophysical parameters, like conductances, impact the activity of SACs during this stage. This question is important

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as some of these parameters evolve during development, or can be controlled pharmacologically, with drastic effects on individual SACs bursting^{6,14}. Finally, and although stage II retinal waves are ubiquitous, experiments reveal a vast variability in the bursting period of SACs across species. Are SACs fundamentally different from one species to another, or is there a common mechanism of bursting under which the period is quite sensitive to physiological parameters variations?

In the spirit of previous works¹⁵⁻¹⁷ using dynamical systems and bifurcation theory to characterize neural systems, we develop a thorough modeling of the SACs bursting which accurately reproduces the experimental observations made in the seminal papers by Zheng *et al.*^{6,14}, several of them for the first time. In addition to new theoretical predictions, we also propose a simple scenario for the SACs bursting mechanism and its evolution during development, that could be tested experimentally. Bursting is thoroughly studied in the neuronal modeling literature¹⁸, as well as the possible bifurcations scenarios generating it. Through our analysis, we identify which parameters are essential to control the transient bursting of immature SACs occurring naturally along development or pharmacologically. As parameters variation is restricted to small ranges by biophysical constraints, our hypothesis is that, during development, SACs are close to a bifurcation point, consequently making them sensitive to small physiological parameters variations.

More precisely, we propose to describe and explain the experimental observations of Zheng et al.^{6,14} on individual immature SACs dynamics, in the following theoretical framework; SACs burst because, during stage II, they are close to a (saddle-node) bifurcation, driving the cell from a rest state to fast oscillations. The bifurcation is triggered by small variations of the membrane potential, due to noise or other cellular effect. After this bifurcation, SACs stay in the regime of fast repetitive firing, triggering the activation of slow calcium-gated potassium channels generating a slow hyperpolarization current (sAHP). This current eventually drives the cell to a homoclinic bifurcation. As a consequence, cells return back to the quiescent state allowing a long after hyperpolarization phase. This bifurcation analysis reveals a mechanism for SACs rhythmic bursting, interpretable with respect to biophysics and the experiments of Zheng et al.^{6,14} on individual immature SACs dynamics. In addition, we identify the fast voltage-gated potassium channels as a possible candidate for the fast hyperpolarizing channels involved in bursting (to our best knowledge not yet been experimentally determined). Based on 19,20 we propose they could be of Kv_3 type. This leads us to a conjecture on the role of the Kv_3 potassium conductance variation which may act on the transient intrinsic properties of SACs excitability during development. Furthermore, the closeness to a bifurcation has strong consequences on the interburst intervals of SACs. It is naturally associated with a high variability which can explain the wide range of interburst periods observed within a network²¹, as well as across species¹⁰. As we show, a small external current (~5 pA) strongly impacts the interburst intervals of SACs. This implies, that, in a network of cholinergically coupled SACs, the bursting period has a wide intrinsic variability, resulting from simple bifurcation arguments, without requiring exogenous mechanisms. In the discussion section, we briefly argue how a set of such inhomogeneous bursters may achieve synchrony and wave generations without any additional mechanism than the existing cholinergic interaction. The general discussion on how cholinergic interaction between nearby autonomous bursters generates wave-like, population-level behavior propagating in a similar way as is observed experimentally will be addressed in a forthcoming paper (see^{22–24}).

The paper is organized as follows. The section 'Results' introduces the model definition, the bifurcation analysis, and the main conclusions resulting from this analysis, including experimental predictions. The section 'Discussion' is mainly dedicated to the biological interpretation of our theoretical results, as well as their impact on the description of retinal waves. Finally, in Methods, we provide the technical aspects of our work including the biophysical derivation of the model's equations, as well as mathematical computations used in the paper. We also include in this section a short review on the comparison of existing models.

Results

During retinal development, Starburst Amacrine Cells (SACs), which are cells forming a lattice right above the ganglion cells layer, are found to exhibit spontaneous intrinsic rhythmic bursting activity, which disappears upon maturation. In the following, we derive a biophysical set of equations, describing the bursting activity of immature SACs, following experimental observations regarding the ionic currents involved in their bursting. We calibrate the equations parameters in order to reproduce the bursting characteristics found in 6,14. The analysis of our model leads to our central hypothesis, that the rest state of SACs lies near a bifurcation point, which means that small variations of parameters could lead to abrupt changes in the dynamics of the cell. This hypothesis, which is novel for immature SACs, jointly explains A) the biophysical mechanism of bursting in the developing SACs, B) accounts for the vast variability of bursting characteristics of SACs across species and C) predicts a possible mechanism on how SACs lose their excitability upon maturation.

Let us now start by deriving our model along with the biological justifications for our equations.

A biophysical model for bursting immature SACs. In order to address the mechanism of bursting and how the physiological parameters of SACs change during development we need to model their underlying cellular mechanisms. In the case of immature SACs, the two key biophysical mechanisms associated with the emergence of spontaneous bursting during early development are⁶:

- fast repetitive bursts of spikes mainly controlled by fast voltage-gated channels;
- prolonged After HyperPolarizations (AHPs) modulating fast oscillations, controlled by Ca^{2+} -gated K^+ channels.

In order to account for (i) and (ii), we model immature SACs activity with a conductance based model of the Morris-Lecar type²⁵, an electronic-circuit-equivalent equation for the local membrane potential, with additional non-linear currents. Our model involves 5 variables, whose evolution is controlled by a set of non-linear

Activated Ca-gated K+ channel R R R R Gating mechanism All 4 subunits R of the channel are bound to activate the channel with the channel subunit Saturated Calmodulin molecule (CaM) binds to each channel subunit 4 Calcium ions bind to each saturated Calmodulin complex (CaM)

Figure 1. Schematic representation of the modeling of the gating mechanism of Ca^{2+} -gated K^+ channels. The correspondence between the channel's activation steps and the modeling state variables R, S, C is also indicated.

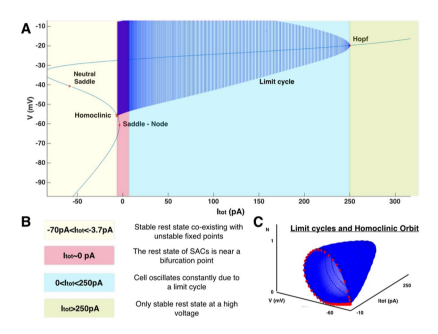


Figure 2. (A) Bifurcation diagram of the fast dynamics (N, V) (Eq. (19) in Methods) when the constant current I_{tot} is varied in a wide range. Each deep blue curve corresponds to the projection of a trajectory of the fast subsystem onto the V axis, for a fixed value of I_{tot} . (B) Interpretation of all the different dynamical regimes of the bifurcation diagram. (C) Same as in A, except that the various trajectories are now drawn in the (N, V) plane and not only projected onto the V axis. Blue. Evolution of the limit cycles when I_{tot} varies. Red. Homoclinic orbit. Bursting stops by an homoclinisation of the limit cycle.

differential equations with 3 timescales: (1) fast variables ($\sim 10 \, ms$) V(t), the local membrane potential, and, N(t), the gating variable for fast voltage-gated K^+ channels; (2) medium variable ($\sim 2 \, s$) C(t), the intracellular Ca^{2+} concentration; (3) slow variables ($\sim 10 \, s$) R(t) and S(t), the gating variables for slow Ca^{2+} -gated K^+ channels. All parameters values and the auxiliary functions involved are found in Methods.

The membrane voltage V(t) obeys:

$$C_{m}\frac{dV}{dt} = I_{L}(V) + I_{C}(V) + I_{K}(V, N) + \underbrace{I_{sAHP}(V, R) + I_{ext}}_{I_{tot}} + \sigma \xi_{t}, \tag{1}$$

where C_m is the membrane capacitance, $I_L = -g_L(V - V_L)$ is the leak current, with g_L , leak conductance and V_L , leak reversal potential.

The fast repetitive firing (see (i)) during the active phase of bursting generally results from the competition between a depolarizing and a hyperpolarizing current. Experiments in have specified voltage-gated Ca^{2+} channels as the depolarizing component of bursting in SACs. However, the ionic channels related to the fast hyperpolarizing component of SACs bursting have not yet been identified experimentally. In this work, we propose that fast voltage-gated K^+ channels play this role (see also Discussion for further justification). Therefore, the terms I_C and I_K , respectively corresponding to Ca^{2+} and K^+ currents, are generating the fast oscillations in our model.

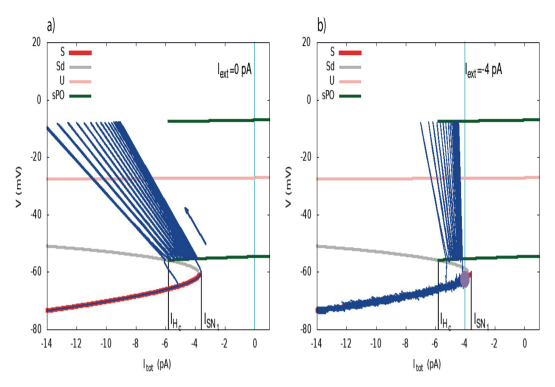


Figure 3. The two scenarios of triggering bursting. (a) Dynamically driven bursting, $I_{ext}=0$. The plot shows a representation of bursting in the plane (I_{top} V) (zoom of the bifurcation diagram of Fig. 2, near I_{SN}). Red lines correspond to stable fixed points, gray and pink lines to unstable fixed points, green lines are the extremal values of voltage fast oscillations. Upon this bifurcation diagram, obtained from the reduced model (Morris-Lecar with $I_{SAHP}=I_{tot}=$ constant), we superimpose the trajectory of the full system in the plane (I_{SAHP} , V) (deep blue). The blue arrow indicates the direction of the flow. In the fast oscillations regime V is varying periodically, with a fast period, inducing a fast variation of the term $V-V_K$ in the sAHP current, explaining the diagonal motion. Despite these fast oscillations one sees that the bifurcation is driven by the slow motion on the branches obtained from the bifurcation analysis assuming a constant current. The conductance $g_{SAHP}R^4$ is varying slowly, with the time scale of R, explaining the slow left-wise shift of the trajectory until the homoclinic bifurcation point is reached. The cyan vertical line corresponds to the value of $I_{ext}=0$. (b) Noise driven bursting. The dynamics has now a stable fixed point (purple circle). An additional current, $I_{ext}=-4$ pA (vertical cyan line) is now present. Bursting cannot occur without an exogenous excitation, here noise. The amount of noise (controlled by σ) necessary to induce bursting depends on the distance between the fixed point (controlled by I_{ext}) and the bifurcation point I_{SN} .

Note that have shown that voltage-gated Na^+ channels do not participate in the bursting mechanism of immature SACs (bursting activity of SACs was not altered upon tetrodoxin -TTX- application), thus, dynamics of Na^+ channels will not be considered in our modeling (see also Discussion).

The voltage-gated Ca^{2+} current is described by a Morris-Lecar model²⁵ where:

$$I_C(V) = -g_C M_{\infty}(V)(V - V_C). \tag{2}$$

 $g_C M_\infty(V)$ is the voltage dependent conductance of the Ca^{2+} channel (see Eq. (21) in Methods). The fast voltage-gated K^+ channel is modeled as:

$$I_K(V, N) = -g_K N(V - V_K).$$
 (3)

where the evolution of the fast voltage-gated K^+ channel gating variable N(t) is given by:

$$\tau_N \frac{dN}{dt} = \Lambda(V)(N_{\infty}(V) - N),\tag{4}$$

 $\Lambda(V)$ and $N_{\infty}(V)$ are given by Eqs (21) and (22) in Methods.

The form of equations (3) and (4) is based on²⁵. However, the timescale of the variable N is much faster ($\sim ms$) in our model that in the original Morris-Lecar model ($\sim s$), calibrated in order to capture the frequency of the fast repetitive firing of SACs, which is around 20 Hz⁶ (see also Bifurcation Analysis).

The long refractoriness in-between consecutive bursts is controlled by a slow After Hyper-Polarization (sAHP) K^+ current, I_{sAHP} . It was observed by 6 that I_{sAHP} is mediated by Ca^{2+} -gated K^+ channels, and that it resembles the sAHP observed by Abel $et\ al.^{26}$, generated by specific channels called SK. Following these tracks we propose a modeling of "SK"-like channel (as named in 6) based on 12 for the structure of the equations and 27 for the calcium

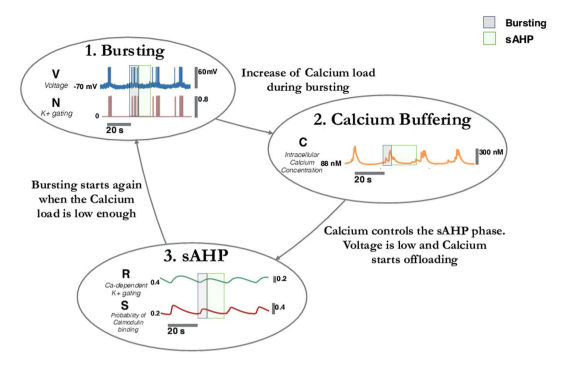


Figure 4. Biophysical mechanism for bursting in immature SACs. (1) Fast spiking occurs due to the competition between voltage gated Ca^{2+} (excitatory current) and K^+ channels (inhibitory current). (2) Calcium load increases during the rapid firing phase while the voltage is in a high voltage state. This leads to a slow increase of sAHP. When sAHP is large enough there is a sharp decay of the voltage (bifurcation) and the hyperpolarisation phase of the cell starts. As voltage is low, the calcium load starts to slowly decrease. (3) During the slow offloading calcium stage we observe a refractory phase. The decrease of calcium concentration induces a slow decay of the sAHP. When calcium is small enough rapid firing starts again via a new bifurcation.

dynamics. The mechanism of the opening of Ca^{2+} -gated ionic channels channels is analyzed in detail in the "Methods" section. In order to simplify the cascade of chemical reactions taking place while opening the sAHP channels, we approximate the channel dynamics by reducing the process into two discrete steps: a) Four ions of Ca^{2+} bind to a second messenger protein called calmodulin, forming a saturated calmodulin complex, CaM; b) CaM binds to each of the four intracellular subunits of the channel to open it (see Fig. 1). This process is mapped to our model through three variables: 1) the variable C which models the intracellular calcium concentration and mainly controls the gating variables of the sAHP channels, 2) the variable S which models the fraction of the saturated calmodulin and 3) the variable R which models the fraction of bounded terminals. This gating mechanism is sketched in Fig. 1.

The sAHP current takes the form:

$$I_{sAHP}(V, R) = -g_{sAHP}R^4(V - V_K),$$
 (5)

where g_{sAHP} is the maximum sAHP conductance. Indeed, 4 bound terminals are needed to open a Ca^{2+} -gated K^+ channel, thus the corresponding conductance is $g_{sAHD}R^4$, involving a fourth order nonlinearity. We model the gating mechanism of the Ca^{2+} gated K^+ channels as follows. The gating variable R(t) obeys:

$$\tau_R \frac{dR}{dt} = \alpha_R S(1 - R) - R,\tag{6}$$

the fraction of saturated calmodulin concentration S(t) obeys:

$$\tau_S \frac{dS}{dt} = \alpha_S C^4 (1 - S) - S,\tag{7}$$

and the intracellular calcium concentration C(t) obeys:

$$\tau_C \frac{dC}{dt} = -\frac{\alpha_C}{H_X} C + C_0 + \delta_C I_C(V). \tag{8}$$

The derivation of the equations (6), (7) and (8) is fully justified in Methods.

Finally, ξ_t is a white noise whose amplitude is constant with time and controlled by σ and I_{ext} is an external current. For subsequent analysis we introduce the current $I_{tot} = I_{sAHP} + I_{ext}$.

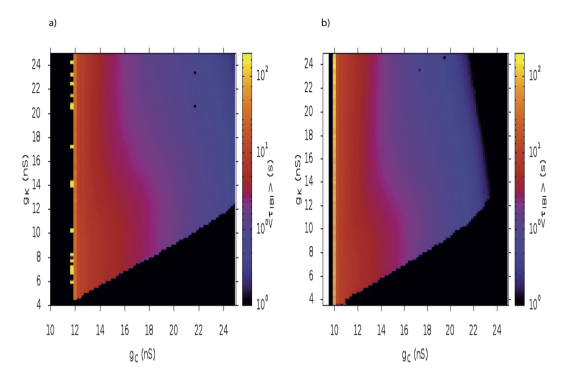


Figure 5. Heat map of the interburst period as a function of g_C and g_K . We sample g_C g_K on a grid with resolution 0.25 nS. For each point, we generate 20 trajectories of duration 2000 s and count the number of bursts in each trajectory. A period of high calcium activity has to last at least 1 second to be considered as a burst. The heat map show the average value of τ_{IBI} in color log scale for a fixed amount of noise $\sigma = 4 \,\mathrm{pA} \,ms^{1/2}$. (a) $I_{ext} = -4 \,\mathrm{pA}$ (noise driven bursting); (b) $I_{ext} = 0 \,\mathrm{pA}$ (dynamically driven bursting).

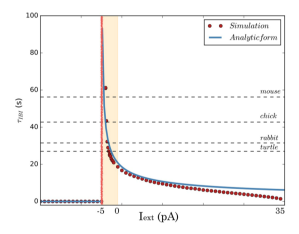


Figure 6. The behavior of the bursting period near a saddle-node bifurcation could explain interburst variability. *Red*: Computing the dependence of τ_{IBI} upon the variation of I_{ext} with our model. *Blue*: A 2 parameters fit of the simulated data with the curve $\frac{K}{2\sqrt{I_{ext}-I_c}}$ for $I_{ext} \in [-20, 40]$ pA. Note that this fit does not hold true away from the asymptote. (see Methods). *Red*: Values of experimentally measured interburst intervals for different species mapped to our results (see text).

Dynamical changes with respect to parameters variations. In our model, the variable V, N evolve with a fast time scale of the order of a few milliseconds under the influence of the current I_{sAHP} whose conductance is slow, being driven by slow variables C, S, R (time scale - several seconds). As a consequence, the main material of this paper relies on a thorough analysis of the bifurcation structure in the fast Morris-Lecar dynamics. There already exist remarkable numerical bifurcation analyses of the Morris-Lecar model (see for example the very detailed work of which has however many parameters: changing their range of variations can dramatically impact the dynamics. Our study on SACs bursting considers quite different range of parameters than the studies we know about (more related to the original paper investigating the barnacle giant muscle 25). For example , investigates the effect of a positive external current whereas we consider a negative external current, with quite different effects on

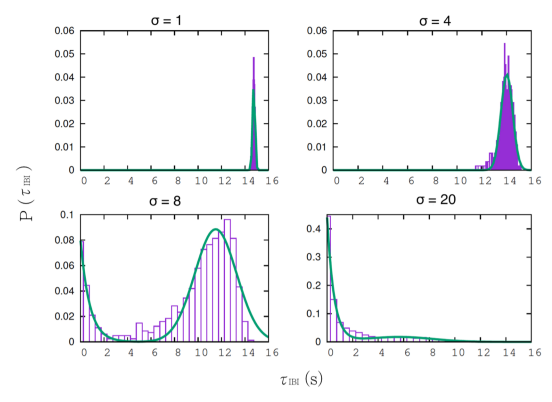


Figure 7. Histogram of bursting periods for different noise levels. We show the distribution of interburst interval distribution τ_{IBI} with different levels of noise. Top left: $\sigma = 1 \, pA \, ms^{1/2}$. Top right: $\sigma = 4 \, pA \, ms^{1/2}$. Bottom left: $\sigma = 8 \, pA \, ms^{1/2}$. Bottom right: $\sigma = 20 \, pA \, ms^{1/2}$. Green curves correspond to fit either by a Gaussian (top), or a linear combination of a Gaussian and a decaying exponential (bottom).

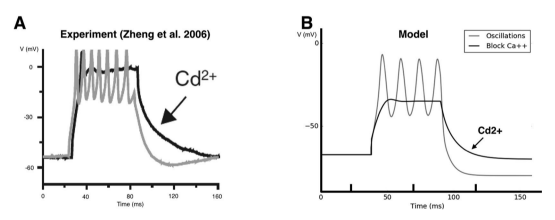


Figure 8. Modeling the cellular mechanisms generating bursting activity in immature SACs. (**A**) Experimental investigation of the cellular mechanisms involved in bursting activity by 6 . *Gray*: Fast sub-threshold oscillations and subsequent AHP generated by a short pulse of current of amplitude 150 pA and duration 60 ms. *Black*: Blocking of all Ca^{2+} related channels by Cd^{2+} . No oscillations are exhibited. (**B**) Simulating the experimental conditions of 6 *Gray*: Reproducing fast sub-threshold oscillations and subsequent AHP emulating the experiment of 6 . Application of a current step pulse of 150 pA for 60 ms. *Black*: Oscillations disappear when Ca^{2+} related conductances are set to zero.

dynamics. Other parameters such as V_1 , V_2 , V_3 , V_4 , V_L , g_K , g_{Ca} have very different ranges in our case (e.g. $V_3 = -1$ mV in 25 , $V_3 \in [0, 25]$ mV in 28 , $V_3 = -25$ mV in our case). Finally, the characteristic rate for opening potassium channels (variable N) is of order $\frac{1}{15}s^{-1}$ in Morris-Lecar's paper, and of order $0.1s^{-1}$ in 28 . That is the variable N is slow. In contrast, our variable N, mimicking fast potassium channels is very fast $\left(\frac{1}{\tau_N} = 200 \, s^{-1}\right)$. This results in a very fast contraction in the N direction near the rest state visible e.g. in the shape of stable and unstable manifolds (see Figs 9A,B, 11c,d, 12B,C and 13B,E,G). As a consequence, our bifurcations study is new and, in addition to explain bursting in SACs, brings original results in the analysis of the Morris-Lecar model dynamics.

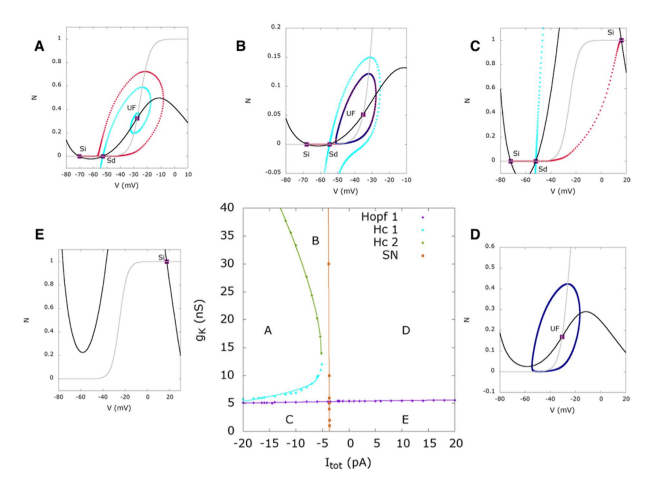


Figure 9. Bifurcation diagram as a function of I_{tot} , g_K . At the bottom center, the bifurcation map is plotted. Points are sampling points obtained by a direct inspection of the phase portrait whereas continuous lines correspond to fit. "Hc" means "homoclinic bifurcation" and "SN" stands for "saddle-node". The capital letters (A-E) inside correspond to the typical phase portraits surrounding the bifurcation map. In these phase portraits, continuous lines correspond to nullclines (N_V , in black, is the V-nullcline whereas N_N , in gray, is the N-nullcline). 'Si' stands for 'Sink, 'Sd' for 'Saddle, 'SF' for 'Stable Focus' and 'UF' for 'Unstable Focus.' Sink and stable focus correspond to stable rest state: a small perturbation about this state decay exponentially fast. On the opposite, saddle and unstable focus are unstable. For saddles we show the stable and unstable directions (black lines) as well as the stable (cyan) and unstable (red) manifolds W_s , W_u . 'SPo' means 'Stable Periodic Orbit'. It corresponds to fast oscillations, plotted in dark blue. In region (A), there is a low voltage stable state. In region (B), a stable state with low voltage coexists with a limit cycle (fast oscillations) separated by an unstable state. When the cell is in the low voltage state, a large enough perturbation (e.g. noise or other cells action) leads it to fast oscillations. In region (C), two stable states, one with low voltage and one with high voltage, coexist separated by an unstable point. In region (D), the cell only exhibits a fast oscillations regime, reached whatever the initial condition. Finally, in region (E), there is a stable state with high voltage. When varying parameters, continuous change occurs in the interior of a region (e.g. the period of oscillations continuously varies), whereas crossing the bifurcation lines leads to abrupt changes.

We distinguished two classes of parameters. The first one consists of modeling parameters constraining the Morris-Lecar dynamics, namely V_1 , V_2 , V_3 , V_4 (see Methods for the corresponding equations). We verified the robustness of the scenario described in the paper when varying these parameters. Especially, V_3 is the half activation potential of the fast potassium channels, playing a central role in SACs excitability as shown later in the text. The second class is the set of parameters constrained by biophysics such as conductances, reversal potentials, capacitance,... Most of them have been fixed based on the biophysical literature and are not considered to vary. Here, we choose to study the variation of 4 of these parameters: I_{tot} , because it controls bursting, and g_K , g_C and V_3 because, as we show later, their variations allows us to reproduce experimental facts, that have not been reproduced before by any model. The most striking consequence is a conjecture on the role of the potassium conductance during development.

Bifurcation analysis. Bursting is an alternation between a rest state and repetitive firing, often modulated by slow voltage or Ca^{2+} dependent processes¹⁸. In order to address which are the biophysical mechanisms for a) triggering, and, b) sustaining bursting in immature SACs, we will perform bifurcations analysis. This is a powerful mathematical tool providing a compact, geometric representation, of dynamical systems behavior when

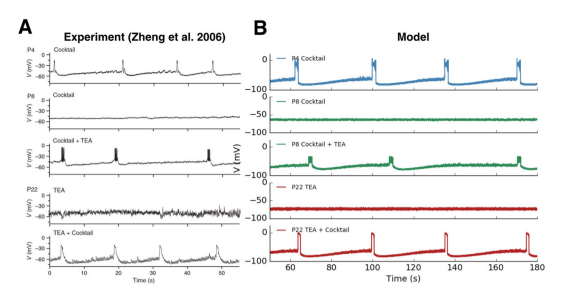


Figure 10. (**A**) Pharmacological manipulations made by⁶ in order to restore bursting activity in rabbit SACs at different ages. (**B**) Varying the potassium conductance g_K and the half-activation potential V_3 changes the excitability of SACs in our model. *Blue*: Modeling the bursting activity of isolated P4 SAC, gK = 8 nS, $V_3 = -16$ mV. *Green top*: Modeling P8 isolated SACs where we see no bursting activity, gK = 10 nS, $V_3 = -34$ mV. *Green bottom*: Decreasing gK = 8 nS, $V_3 = -34$ mV we restore oscillations. *Red top*: Modeling P22 coupled SAC where there is no bursting activity upon treatment with TEA. An inhibitory current $I_{ext} = -10$ pA is applied to coarsely mimic the inhibitory input to SAC at this stage of development. To account for the TEA application we take decrease the conductance gK = 4.5 nS ($V_3 = -35$ mV). *Red bottom*: Restoration of bursting at P22 by removing all inhibitory synaptic connections, $I_{ext} = 0$ pA. Removing inhibition from mature amacrine circuitry, along with blocking a subfamily of K^+ channels with TEA, is enough to re-initiate bursting activity. The values proposed here for V_3 , g_K are only indicative, as the same behavior is observed in a wide range of parameters (see the bifurcation diagram in Fig. 11).

parameters are modified. While, in general, a continuous variation of a parameter induces a continuous variation of the dynamical system's solutions, the variation of parameters around bifurcation points induces, in contrast, strong and abrupt changes in the dynamics.

We first perform a bifurcation analysis of the fast V, N dynamics in the presence of the current I_{top} constant in time, and used as a bifurcation parameter. The bifurcation diagram is represented in Fig. 2A (drawn using MATCONT²⁹), where we have explored a wide range of variation of I_{tot} (-70 to +310 pA). For the analysis, we consider an example with the potassium $g_K = 10$ nS, and calcium conductance $g_C = 12$ nS (see Robustness with respect to parameters variations). We reveal a rich dynamical profile upon the variation of an external current where for $I_{tot} < -3.7$ pA, stable rest state co-exists with unstable fixed points, for $-3.7 < I_{tot} < 250$ pA, cell oscillates constantly due to a limit cycle and for $I_{tot} > 250$ pA only a stable rest state at a high voltage exists (see Fig. 2B). We refer the reader to e.g.³⁰ for the dynamical systems terminology used in the next paragraphs.

In more detail, when $I_{tot} \lesssim -5.8 \, \mathrm{pA}$, there is a stable rest state with a membrane potential $V_{rest} \in [-70, -60] \, \mathrm{mV}$, coexisting with two unstable fixed points, a saddle and an unstable focus (see Fig. 9A for the phase portrait of this case). When $I_{tot} \simeq -5.8 \, \mathrm{pA} = I_{H_c}$, there is an homoclinic bifurcation giving rise to a stable limit cycle. In the range $I_{tot} \in [I_{H_c}, I_{SN_1}]$, where $I_{SN_1} \simeq -3.7 \, \mathrm{pA}$, the stable rest state coexists then with a stable limit cycle, corresponding to fast oscillations, and an unstable fixed point. When $I_{tot} = I_{SN_1} \simeq -3.7 \, \mathrm{pA}$ the stable rest state coalesces with the middle unstable branch and both disappear with a saddle-node bifurcation (SN_1) . For $I_{tot} > I_{SN_1}$ the dynamics has only one attractor, the limit cycle, corresponding to fast oscillations. This cycle eventually disappears by a Hopf bifurcation at $I_{tot} = 250 \, \mathrm{pA}$ (Fig. 2). However, this value of external current is quite beyond the range of plausible values during the bursting activity of SACs.

We observe that the rest state of SACs, corresponding to $I_{tot} = 0$ pA, lies near a saddle-node bifurcation point within a narrow region of a few pA, which is our main emerging hypothesis. This leads to important consequences for SACs described in the following.

Bifurcations analysis reveals two possible drives for triggering bursting. The bifurcations analysis for the fast subsystem shown in Fig. 2, reveals all possible dynamical regimes upon the variation of a constant external current within a wide interval of our choice. Thanks to the timescales separation in our model (fast, medium, slow), we make the approximation that the bifurcation diagram in Fig. 2A, does not change significantly when we consider the full model case. However, in the full model I_{tot} is not constant, but time-dependent, driven by sAHP. By superimposing the solution of the full model and the bifurcation diagram of the fast subsystem, we reveal how the trajectory driven by the slow variations of I_{sAHP} matches the bifurcation diagram, in the plane (I_{tot} V). See Fig. 3 showing a zoom of the bifurcation diagram shown in Fig. 2A, to our regime of interest (stable branch: red, neutral

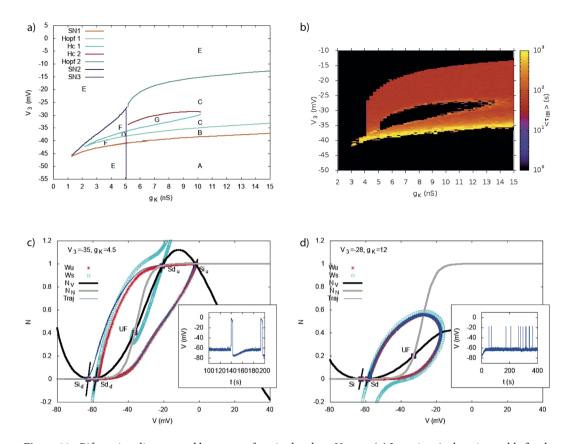


Figure 11. Bifurcation diagram and heat map of τ_{IBI} in the plane $V_3 - g_K$. (a) In region A, there is a stable fixed point with low voltage. In B, 2 stable fixed points coexist with a saddle-point. In C a stable fixed point with low voltage coincides with a stable periodic orbit giving rise to fast oscillations. In region D there are 4 fixed points (2 stable, 2 unstable) and a limit cycle. In region E there is a stable fixed point with low voltage and a stable fixed point whose position depends on g_K and V_3 . In region F there are five fixed points, 2 stables. Region G contains 2 stable fixed point (a sink and a focus) separated by an unstable fixed point. Bursting takes place in regions C and D. (b) Heat map of the interburst interval τ_{IBI} . Same representation as Fig. 5. We sampled the plane g_{K} , V_3 with a step of 0.25 nS on the g_K axis and 1 mV on the V_3 axis, explaining the irregular shape of the border. (c.d) There is a discrepancy between the heat map and the bifurcation diagram: (i) The bifurcation map does not predict bursting for $g_K \in [4, 5]$ nS, $V_3 \in [-40, -25]$ mV whereas the heat map shows calcium peaks lasting more than 1 s in this region. This is due to the particular structure of the phase portrait of the Morris-Lecar model in this region and the effect of noise, as shown in (c). This phase portrait has two saddles (Sd) down (Sd_d) and up (Sd_u) , and two sinks (Si), down (Si_d) and up (Si_u) . The unstable manifold (W_u) in red) of Sd_d connects to Si_u , on the right, and to Si_d , on the left; The unstable manifold of Sd_u connects to Si_u , on the right, and to Si_d , on the left. Thanks to noise, dynamics can leave Si_d and reach Si_u where it stays until sAHP rising drive it back to Si_u . This is illustrated by the blue trajectory (complete system with noise) which follows the unstable manifolds of the Morris Lecar model. The inset shows the trajectory of the voltage for the complete system in the presence of noise. (ii) (d) While the bifurcation diagram predicts bursting for $V_3 \sim -28 \,\mathrm{mV}$, $g_K \sim 12 \,\mathrm{nS}$, the heat map does not show bursting in this region (black). This is again due to noise. When the cell is on low rest state (Si) noise drives it to the limit cycle, but after a few oscillations, noise drive the cell back to the rest state Sd (see inset for the voltage trajectory). This effect depends on the distance between the two fixed points Si, Sd, fixed by the model parameters, and on the noise intensity σ (here 4 pA ms $\frac{1}{2}$).

saddles: gray, $unstable\ branch$: pink, green traces showing the maximum and minimum of V along the limit cycle), with the superimposed trajectories of the full system (blue curves).

Analyzing how the slow variations of I_{sAHP} generate bursting, we actually exhibit two possible drives of triggering bursting:

(i) Dynamical drive, induced by the mere dynamics of neurons, without external influence. In our model, the joint fast dynamics of Ca^{2+} and K^+ channels generates fast oscillations while the slow AHP, mediated by Ca^{2+} gated K^+ channels, modulates slow oscillations⁶. The conjunction of these two mechanisms generates bursting. Note that in this case, $I_{ext} = 0$ in Eq. (1), meaning there is no external current (Fig. 3 left). In this scenario, cells burst periodically, with a frequency controlled by the characteristic times τ_R , τ_S of variables R and S respectively. From a dynamical systems perspective, there are two important bifurcations associated with bursting: a) bifurcation of the rest state that leads repetitive firing and b) bifurcation of a spiking attractor that

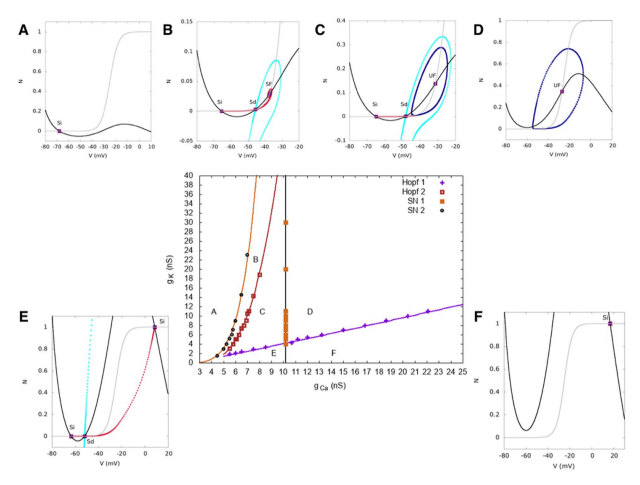


Figure 12. Bifurcation diagram as a function of g_C , g_K in the dynamically driven bursting regime. Same representation as Fig. 9 and same comments. Note the correspondence with the heat map Fig. 5.

- leads to a rest state¹⁸. There are two possible types of bifurcations linked to such type of behavior (1) Homoclinic saddle-node bifurcation and (2) Hopf bifurcation. We have therefore shown that (1) holds in our case. The first bifurcation associated with the breaking of the rest state is the saddle-node bifurcation. The "spiking attractor", here the limit cycle, is created by an homoclinisation. According to the classification of bursting made by Izhikevich in¹⁸, this corresponds to a "square-wave" point-cycle planar burster.
- (ii) Noise-drive, induced by an external excitation, typically noise. In order to move to this regime, it suffices to add an external constant current $I_{ext} < I_{SN_1} = -3.7 \, pA$ to Eq. (1). Then, the deterministic dynamics converges towards a stable fixed point, the cell ends up in a steady rest state and cannot switch to rapid firing. In this case, the bifurcation diagram (Fig. 3 right) has now two fixed points: V_{rest} , which is stable (purple point), and V_u which is unstable (intersection of the gray branch and the vertical blue line). However, in the presence of noise, sufficiently close to the bifurcation point (depending on the noise amplitude), the random fluctuations around the rest state leads eventually the cell into the rapid firing regime (Fig. 3 right). Note that when I_{ext} is sufficiently large in absolute value the total current I_{tot} is smaller than I_{SN_1} whatever the (negative) value of I_{sAHP} .

Taken together, since noise is always present in neurons, distinguishing those two scenarios for triggering bursting is not easy experimentally but still possible (see Discussion). Maybe both scenarios are possible in the same network of SACs, since it would correspond to a slight variability in the intrinsic parameters which control the rest state of each SAC, placing it on the right or left of the bifurcation within a narrow region ($\sim 10\,pA$ according to our model). We are actually able to capture both the dynamically and the noise driven bursting regimes in a single, unifying model. In the presence of noise the dynamically driven bursting regime still exists but the period has now random fluctuations. In fact, the noise smooths the transition between those 2 regimes. Also, what our analysis reveals is that, in the noise driven regime, bursting will occur only if the noise has a sufficiently large amplitude compared to the distance between the rest state and the bifurcation point. On the other hand, in the dynamically driven regime, bursting will occur only if the rest state is sufficiently close to the bifurcation in order for the amplitude of the negative sAHP current to be sufficient to compensate the depolarizing current (I_C) and enter the hyperpolarization phase. Otherwise, if the rest state is further on right from the bifurcation, SACs would exhibit only a repetitive firing regime without hyperpolarization. In other words, in both regimes, the cell has to be close enough to the bifurcation point to observe bursting.

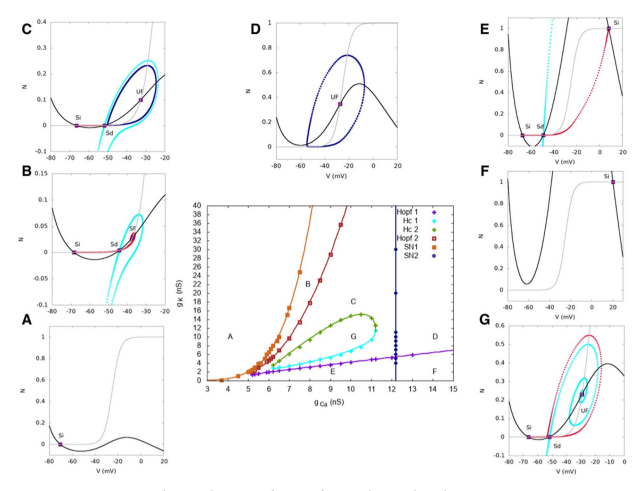


Figure 13. Bifurcation diagram as a function of g_C , g_K in the noise driven bursting regime. Same representation as Fig. 9. In region (**A**), there is a unique stable rest states with low voltage. In region (**B**) two stable rest state with low voltage coexist separated by an unstable point; hence dynamics is bistable. In region (**C**), a stable rest state with low voltage coexists with a limit cycle (fast oscillations) separated by an unstable state. When the cell is in the low voltage rest state, a large enough perturbation leads it to fast oscillations. In region (**D**) the cell only exhibits a fast oscillations regime, reached whatever the initial condition. In region (**E**) two stable rest states, one with low voltage and one with high voltage, coexist separated by an unstable point. Region (**F**) has only one high voltage stable rest state. Finally, in region (**G**), there is a low voltage rest stable state.

Bifurcations analysis reveals a biophysical mechanism for SACs sustained bursting. Our analysis reveals also a possible biophysical mechanism to explain how bursting activity is sustained periodically in immature SACs. Once fast repetitive firing is elicited in SACs, either through i) dynamical, or ii) noise drive, the biophysical cyclic mechanism which produces bursting is the same. From the bifurcation analysis we see that once the cell crosses the saddle-node SN_1 , it is firing rapidly (limit cycle). Being in a high voltage state, Ca^{2+} loads, leading to a slow rise of sAHP conductance. This corresponds, in the bifurcation diagram, to a motion towards negative current values, with a slow variation due to the sAHP conductance driven by the slow variable R, and fast oscillations due the fast variable $V - V_K$ appearing in the sAHP current $I_{SAHP} = -g_{SAHP}R^4(V - V_K)$. As revealed by a bifurcation analysis where the sAHP conductance is the bifurcation parameter (not shown), and as shown in Fig. 3, these fast oscillations have no effect on the bifurcation, what matters is the slow variation. The increase of sAHP conductance continues until the cell reaches the homoclinic bifurcation where rapid firing stop. $I_{tot} = I_{sAHP}$ is very negative now, leading to the hyperpolarization phase. Then, because voltage is low, $[Ca^{2+}]$ decreases and I_{sAHP} drops down. Eventually, SAC crosses the SN_1 bifurcation point and starts firing again either due to dynamics in one case or noise. This is illustrated in Fig. 3. The deep blue traces are the trajectory of a burst in the plane $(I_{tot} = I_{sAHP}, V)$, for the full system. One sees in particular the fast oscillations coming from the fast variable $V - V_K$.

Nevertheless, for *both* proposed scenarios the biophysical mechanism for sustaining bursting is the same. Although the conclusion drawn on the model is entirely based on a mathematical analysis via bifurcation theory, we have sketched the main mechanism of SACs bursting in Fig. 4, for readers not familiar with bifurcation theory. Note that Fig. 4 is only qualitative. In contrast, bifurcation analysis gives us more insights on bursting mechanism, where actually two regimes are possible, and quantitative results, especially on the parameters determining the transitions inducing bursting and the range of these parameters. To our best knowledge this is the first model able to provide a mechanism to account for the repetitive firing of SACs, which is the condition under which calcium controls the after-hyperpolarization phase.

Explaining the wide range of interburst intervals (IBI) across species by a unique mechanism.

Spontaneous bursting activity in immature SACs has been consistently observed across various species i.e. mice, rabbits, chicks, turtles, macaques etc., although, the measured mean interburst interval (τ_{IBI}) strongly varies $^{1-4}$. This experimental observation raises the question whether there exist several bursting mechanisms changing from species, explaining these variations in interburst. Bifurcations theory provides another explanation though. As we showed, in our model, bursting involves a saddle-node bifurcation. It is also known from³¹, that interburst is known to be sensitive to parameters variations near this type of bifurcation. In Fig. 5, we illustrate this effect by numerically computing the heat map of the mean interburst intervals of SACs, while varying the parameters g_K and g_{\odot} for both bursting scenarios (noise and dynamic driven). We show that the IBI increases when approaching the Saddle-Node line. Note that we define a burst numerically by a thresholding on Ca^{2+} : a burst occurs when this concentration exceeds 150 nM during a time larger than 1 s. There is a "critical" region (yellow zone) where interburst increases (up to several minutes). This is nicely explained by a bifurcation analysis (see section Robustness with respect to parameters variations). The irregular shape of this region - alternation between yellow and black regions - is due to our numerical procedure. We generate 20 trajectories of duration 2000 s and count the number of bursts in each trajectory. In our simulation, when going close to the critical region we observe that this sampling is not sufficient; we have rare bursts giving poor statistics, and, in some cases (black regions in-between vellow ones), we observe no burst in the sample.

To illustrate the sensitivity of τ_{IBI} to a parameter variation, we now vary the external current I_{ext} (Fig. 6). This indeed induces strong variations in τ_{IBI} compatible with the variations observed across species, which increases monotonously and sharply as I_{ext} decreases, following a hyperbola whose form has been derived analytically (see Methods and Fig. 6, blue trace). As shown in the paper³¹, hyperbola is one of the 2 possibilities, corresponding to the case where the noise amplitude is small compared to the excitability. We also observe a sharp transition from a bursting to a non-bursting regime, where $\tau_{IBI} = 0$, which corresponds to the loss of SACs excitability.

Based on the shape of our theoretical curve, we show that τ_{IBI} exhibits a *strong asymptotic behavior* around a very narrow regime of $I_{ext} = \{ [-5, 0] \text{pA}$. The analytic form of the function $\tau_{IBI} = f(I_{ext})$ is the following (see Methods for detailed derivation):

$$\tau_{IBI} = \begin{cases} 0, & I_{ext} \le I_c; \\ \frac{K}{\sqrt{I_{ext} - I_c}}, & I_{ext} > I_c; \end{cases}$$
(9)

(where K = 0.657 s $pA^{\frac{1}{2}}$ and $I_c = -5$ pA, for $\sigma = 4$ pA $ms^{1/2}$, $g_K = 10$ nS, $g_C = 12$ nS). The values K, I_c are obtained from fit. Note that for $I_{ext} \le I_c$, τ_{IBI} is not defined because neurons do not burst anymore.

This result suggests that, depending on slight variations of physiological conditions, a SAC could exhibit very variable bursting periods, providing an explanation how different species exhibit variability in the interburst interval of immature SACs. In real SACs, the inhibitory current I_{ext} could correspond to a change in the intrinsic inhibitory ionic currents of individual SAC or to external inhibitory inputs from other layers of the retina. It could also mimic changes in synaptic inhibition (e.g. $GABA_A$ maturation, changes in $[Cl]_{in}$). These factors could drive the disappearance of bursting activity and subsequently the loss of SACs excitability. On the other hand, an increase of I_{ext} leads to a gradual decrease of τ_{IBI} towards zero, meaning that in this regime of parameters, SACs tend to burst repeatedly, without a refractory period. This scenario is not observed under normal physiological conditions, but could be tested experimentally by varying I_{ext} as an externally applied current. Finally, in the experimental paper³², it is found that the bursting period increases upon maturation (experiment in P1–P2 rabbit SAC). In light of the present model-driven analysis, this could be linked to a change of intrinsic properties of SACs during development.

To compare with experimental results, we show in the same figure τ_{IBI} for different species as found in the literature (Fig. 6). We have not been able to find τ_{IBI} for turtles and chicks. We found the mean inter-wave intervals τ_{IWI} instead. In order to extrapolate to τ_{IBI} in these cases, we used a common constant scaling factor of 3, based on the ratio between τ_{IBI} and τ_{IWI} for rabbits and mice found in and respectively. Besides this narrow regime a decrease of I_{ext} results in a sharp disappearance of bursting activity, which means that bursting activity stops after a limit value.

To conclude, our study suggests that bursting activity in immature SACs could share a common mechanism across species. The same mechanism could also induce interburst variability within a network SACs, thus with cells bursting with a different period. Such variability has been implemented in 21 by varying the decay time of the sAHP from a Gaussian distribution. In our case, the variability arises without adding an extra mechanism and is induced by the hyperbolic behavior observed near a saddle-node bifurcation upon a tiny variation of a single parameter, I_{exp} interpreted as an effective depolarizing/hyperpolarizing current. Taken together, the vast variability of SACs bursting features accross different species, could be explained simply by our hypothesis that the rest state of SACs lie within a narrow regime around a saddle-node bifurcation point.

Characterizing the effect of noise on the probability distribution of bursts. In 6, recordings show that the bursting periods of SACs are not regular and have a certain probability distribution. We obtain this effect by adding the noise ξ to the dynamics of the voltage V (see Eq. (1)). Note that noise is always present in real neurons due e.g. to fluctuations in ionic currents from the random opening of ionic channels. What we want to illustrate here is the role played by the intensity of this noise, controlled by the parameter σ . For small σ , dynamics has fluctuations around the deterministic trajectory with little effects during the bursting phase. In contrast, low additive noise during the slow (after-hyperpolarization) dynamics, is enough to accelerate the start of a burst. For higher values of σ the bursting period of cells decreases dramatically. As shown in Fig. 7 the presence of noise has also a drastic impact on the shape of the interbursts intervals distribution.

Experimentally, maybe due to the lack of sufficiently large samples in⁶, the exact shape of the distribution of the interburst intervals is not sharply defined. Therefore, the comparison with our theoretical results is difficult. However, the experimental distribution obtained by these authors is definitely not exponential. As we observe an exponential distribution for large enough σ values, Fig. 7, this remark provides us with an upper bound on the level of noise which should be no greater than $8 pA m s^{1/2}$. Note that the left-wise peak in the bimodal distribution ($\sigma \sim 8 pA m s^{1/2}$), corresponds to short bursts. Similar bursts have been observed in the spontaneous activity of retinal ganglion cells during waves, in turtle and in mouse, see e.g. control in Figs 3 and 7 in⁴ (E. Sernagor private communication).

The role of the fast potassium conductance in bursting activity. We now address in further detail the potential role of the fast voltage-gated potassium channels used in our model to produce bursting (see also the work of E. Marder and collaborators, in a different context³³). Zheng $et\ al.^6$ propose that the ionic channels mainly involved in the bursting activity of SACs during early development are voltage-gated Ca^{2+} channels. In this work the hyperpolarizing current involved is not characterized and these authors don't mention potassium channels before the end where they perform experiments with TEA (see below for more details). We proposed above fast voltage-gated K^+ channels as a source of fast inhibition necessary for the active phase of bursting of SACs. We now justify this claim based on several experiments made by Zheng $et\ al.$, interpreted in the context of our bifurcation analysis.

We first show (Fig. 8) how our model accurately reproduces a key experiment of 6 . Here, the authors artificially control the triggering of fast oscillations by applying a short current pulse ($I_{ext} = 150 \, pA$ for $60 \, ms$) to individual immature SACs. Also, upon the pharmacological application of Cd^{2+} , which blocks all Ca^{2+} related channels (voltage-gated Ca^{2+} and sAHP), they show that no oscillatory activity is triggered upon stimulation, but only a raise in the plateau of the level of the voltage. The corresponding figure of their experiment in the paper 6 has been reproduced in Fig. 8A (with the kind authorization of the authors). As shown in Fig. 8B, we are able to reproduce reliably this result using the fast K^+ channels described in Eq. (3). Particularly, we are able to simulate the emergence of fast oscillations during a short current pulse, with an AHP phase after the end of the pulse (see gray curve in Fig. 8B). We emulate as well the disappearance of the oscillations observed by these authors, upon blocking all Ca^{2+} related channels (voltage-gated Ca^{2+} and sAHP), by setting, in our model, the corresponding conductances to zero (see black curve in Fig. 8B).

This is interpreted as follows from the bifurcation diagram Fig. 9, in the plane I_{tot} , g_K . The simulation has been done in the noise driven bursting regime where the cell, initially in its rest state, is in region A. A pulse of current $I_{ext} = 150 \,\mathrm{pA}$ drives the cell in region D where it fast spikes. Removing the current pulse drives back the cell in region A. Here we observe a small sAHP hyperpolarizing the cell. In order to observe the same behavior in the dynamically driven bursting regime, one needs to clamp the voltage so as to maintain the cell in the rest state.

These observations support therefore our proposition that the hyperpolarizing component of the fast oscillations observed in immature SACs are driven by fast voltage-gated K^+ channels.

Exploring the role of the potassium conductance in the loss of SACs excitability upon maturation.

SACs in the retina of vertebrates lose their ability to spontaneously burst once they reach a certain stage of development different for each species. This transient excitability is a key process in the developing retina and the shaping of the visual system. It is not yet clear which physiological properties of SACs change upon maturation causing the abrupt change from autonomous bursting to rest state. So far, to our best knowledge, no experiment has studied in detail the biophysical properties of immature SACs at the level of each ionic channels involved during this window of development. However, there are some indirect experimental implications which help us extrapolate a possible scenario on how SACs change their properties upon maturation. In²⁰ Ozaita *et al.* show that mature SACs not only stop bursting but also cannot be depolarized beyond $-20 \,\mathrm{mV}$. A specific type of inhibitory voltage gated K^+ channels, Kv_3 , is responsible for this property, providing an electric shunt to SACs somas. In the contrary, in the developing retina, SACs are autonomous bursters, depolarised beyond mature SACs ($\sim -5 \,\mathrm{mV}$)⁶. Therefore, upon maturation, the characteristics of voltage gated K^+ channels evolve, leading to a drastic change in SACs activity. In our model, these observations can be reproduced by the variation of two parameters; (i) the conductance g_K and (ii) the half-activation potential V_3 of the voltage gated K^+ channels.

Here a point of caution is necessary. The modeling chosen here, in the same spirit as Morris-Lecar or Hodgkin-Huxley model, characterizes the behavior of a small piece of a cellular membrane, large enough to contain many ionic channels so that the description in terms of activity variables (N, R, S) is correct. As a consequence, parameters such as g_K , V_3 are phenomenological, depending on membrane characteristics (e.g. the density of ionic channels of a given type), and on channels characteristics. When we vary a parameter such as V_3 , it means that we mathematically change the threshold of the term mimicking fast-K channels in the model. The biophysical interpretation of this variation is more difficult. The easiest interpretation is that it corresponds to a change of threshold response of the corresponding channels, but other explanations could be also possible. Assume for example that there are two types of fast K^+ channels (e.g. kV3.1 and kV3.2) on the membrane, with different densities and reactivity. Then, a mere change of their density during development would impact the value of the phenomenological parameters g_K , V_3 of our model.

With our model, we are able to reproduce an experiment by Zheng *et al.*⁶. The results of their experiment are shown in Fig. 10, left column (with the kind authorization of the authors), whereas the model results are presented in Fig. 10, right column. Our results are based on a bifurcation analysis in the plane $V_3 - g_K$, which reveals a wide region of parameters where bursting takes place. In Fig. 11 we show, on the top left, the bifurcation diagram in the plane V_3 , g_K and on the top right, a heat map, similar to Fig. 5). This map makes easier the interpretation of Zheng *et al.* experiment which mainly addresses the following questions:

- How does SACs autonomous bursting stop? Zheng et al. consider first isolated SACs (all synaptic connections are inactivated with a pharmacological cocktail). While SACs are bursting spontaneously (i.e. without the influence of the other cells) at P4 (first row, left in Fig. 10), this spontaneous activity disappears at P6 (second row, left). We can easily reproduce this observation (Fig. 10, first and second row, right) by moving from region C (fast oscillations) in the bifurcation diagram of Fig. 11 to another region. For example, a simple transition to region A is obtained by decreasing the half-activation potential V_3 of the fast K^+ channels, leading to the saturation of SACs depolarization below $-20\,mV$ observed in mature SACs. A concomitant variation of g_K is also possible leading to stop bursting as well. To illustrate this scenario, we set $g_K = 10\,\text{nS}$ and $V_3 = -34\,\text{mV}$ as an indicative example (see Fig. 10).
- How can bursting be restored in further mature SACs? Zheng et al.⁶ show that bursting activity can be re-initiated pharmacologically in P8 (stage III) and P22 retinas with chemical agents upon i) blocking all synaptic connections (gap junctions, cholinergic, gabaergic and glutamatergic synapses), especially inhibition by applying an antagonist cocktail and ii) decreasing the conductance of the fast inhibitory K^+ channels, by applying tetraethylammonium (TEA). This suggests that upon maturation, bursting is in fact suppressed by alterations in the intrinsic properties of individual SACs. Particularly, in the same preparation (P8 isolated SACs), the authors apply a voltage-gated K^+ channel blocker tetraethylammonium (TEA), which results in restoring bursting activity in isolated mature SACs (Fig. 10, third row, left). A potential type of potassium channels involved is the TEA -sensitive K^+ of the $K\nu_3$ family. Therefore, this experiment emphasizes once again the potential role of fast potassium channels and especially the Kv_3 family. We mimic the blocking of these channels in our model by decreasing the conductance g_K . When g_K is small enough, we reproduce the bursting restoration (Fig. 10, third row, right). This example corresponds, in Fig. 11, to a motion from $(g_K, V_3) = (10 \text{ nS}, -34 \text{ mV})$ (region B) to $(g_K, V_3) = (8 \text{ nS}, -34 \text{ mV})$ (region C). Note that the chosen value $V_3 = -34 \,\mathrm{mV}$ allows us to fix the maximal depolarization to $-20 \,\mathrm{mV}$, the value observed by Ozaita et al.²⁰ for further mature SACs. Remark also that the lower branch of the heat map Fig. 11 (region between G and B in the bifurcation map) corresponds to interburst intervals of the order of 8–16 min, so according to SACs recording not to a biophysically plausible bursting regime.
- How can bursting be restored in adult SACs? Upon further maturation, Zheng et al. show that the restoration of bursting in P22 SACs (late stage III-before eye-opening) depends on the interplay between the change of the intrinsic properties of K^+ channels strong inhibition (TEA application) and blocking the strong inhibitory (gabaergic) input induced by other amacrine cells at this phase (Fig. 10, fourth and fifth row, left). Neither of these conditions suffice alone for the bursting restoration. To model the gabaergic inhibitory input to coupled SACs at P22, we add an inhibitory current $I_{ext} = -10 \,\mathrm{pA}$ whereas the effect of TEA is modeled by decreasing the g_K conductance to 4.5 nS. This conductance is not set to 0 because TEA blocks only one subtype of Kv_3 channels while there exist several of them (see below) in the developing retina. The whole operation has the effect of suppressing bursting (red trace top) and corresponds moving from region B to region A in Fig. 9. Bursting re-initiation is obtained by setting back the current I_{ext} to zero.

Along these lines, we have been able to reproduce the effects of the pharmacological manipulation performed by Zheng *et al.* at the three separate ages; *P*4, *P*8 and *P*22 in Fig. 10.

Taken together, during development, SACs exhibit a transient bursting activity which is lost upon maturation. In the context of our bifurcation analysis, this process is interpreted as follows: immature SACs rest state lies close to a bifurcation point in order to facilitate the bursting activity which is essential for the generation of retinal waves and the shaping of the visual system. Upon maturation and once this functionality is no further useful for the retinal circuitry, we predict that SACs rest state is naturally driven towards the stable branch (further left in Fig. 2A), away from the bifurcation point, where bursting generation is no longer possible (we remind that noise could drive the cell to cross the bifurcation but, the further away, the less biophysically plausible this scenario is regarding the noise amplitude). In our model, this essentially corresponds to adding a hyperpolarizing (negative) current as an input. For identifying the source of such current, we propose a conjecture on the possible role of the $K\nu_3$ potassium channels during development, based on our analysis, which could reveal a mechanism of how SACs lose their excitability.

A conjecture on the role of the Kv₃ channels in the loss of SACs excitability upon maturation.

We predict that bursting occurs in immature SACs, during a certain age window while fast potassium channels are under-expressed. Upon development, we make the hypothesis that potassium channels increase their expression and spontaneous bursting stops when the hyperpolarizing K^+ dominate the dynamics over the depolarizing Ca^{2+} channels. Also, we suggest that upon maturation, the half activation potential V_3 decreases in order to provide a saturation in mature SACs activity as shown in²⁰. This theoretical prediction could be experimentally tested, by following the expression of the fast K⁺ channels in SACs during development and showing a potential increase of their activity along maturation (see discussion section). Moreover, we propose that the fast voltage gated K^+ channels involved is the $K\nu_3$ family. More specifically, two subtypes of fast voltage gated K^+ channels have been identified 19,20 , both belonging to the Kv_3 family: i) I_K delayed rectifier currents, sensitive to TEA (tetraethylammonium) emitted by subunits Kv_3 .1 and Kv_3 .2 and ii) I_A A-type currents, sensitive to 4-amino-pyridine (4AP), emitted by subunits Kv_3 , 3 and Kv_3 , 4^{19} which are not sensitive to TEA. The apparition and evolution of such channels in SACs during development has not been studied yet. However, in the experiment of⁵, the application of TEA in *immature* SACs (rabbit ~ P8) was crucial to the re-initiation of bursting activity. This finding implies that, during development TEA sensitive K^+ channels are already expressed in SACs. On this basis, we propose that the specific type of K^+ channels responsible for the spontaneous bursting activity during development and eventually its loss upon maturation, is the I_K rectifier currents of the Kv_3 .1 and Kv_3 .2 subtype.

To conclude, we propose that, during early development (<P6), Kv_3 channels are under-expressed, allowing a competition between inhibition (K^+) and excitation (Ca^{2+}), leading SACs to burst (see Fig. 10, blue trace). Upon maturation, the expression of these channels could evolve increasingly, leading to stronger inhibition, dominating fully the competition of inhibitory/excitatory channels, by suppressing oscillations completely. Therefore, we suggest that the evolution of the expression of fast Kv_3 channels could be part of the transient process that leads to a complete loss of excitability of mature SAC. Our theoretical results indicate that the level of the expression of Kv_3 channels could increase gradually upon maturation, which essentially means that the conductance of these channels would increase as well as other physiological properties of the channels such as kinetics parameters like the half-activation potential (fixing the characteristic activation sigmoid of the channel). This type of experiment could elucidate the exact role of Kv_3 channels in the intrinsic properties of the excitability of SACs.

Robustness with respect to parameters variations. We now demonstrate the robustness of this analysis to variations of our main parameters. We already showed the effect of varying I_{tot} and g_K in Fig. 9. As bursting is a competition between calcium and potassium dynamics, we have also checked the structure of bifurcations when varying the calcium conductance g_C and the potassium conductance g_K around the values we have fixed in our model. We have used different software (XPPAUT³⁴, MATCONT²⁹, PyCont³⁵) and we were not able to get all bifurcations with them (especially homoclinic bifurcations). So we obtained the bifurcation diagram by a direct inspection of the phase portraits with respect to parameters variations. The points shown in the bifurcation maps are sampling points whereas continuous lines correspond to fit. We note that Hopf and Saddle Node bifurcations were obtained identically using the bifurcation analysis software quoted above.

The resulting bifurcation diagrams, in the two regime of bursting, are drawn in Fig. 12 (dynamically driven bursting) and 13 (noise driven bursting). In the noise driven bursting regime there is a region (G in the Fig. 13), delimited by two homoclinic bifurcations lines, where the cell is bistable without fast oscillations. Note that this region does not exist in the dynamically driven bursting regime, hence Figs 12 and 13 have a different *x* axis range. (The goal is to enlarge region G, which does not appear in Fig. 6).

Discussion

In this paper we have proposed a model for spontaneous bursting of SACs, one of the key ingredients for the initiation of stage II retinal waves. To our best knowledge, this is the first model able to characterize SACs dynamics in great detail and to show that biophysical properties of SACs are sufficient to explain intrinsic bursting and refractoriness in these cells. The model can reproduce spontaneously occurring fast oscillatory activity followed by long refractory periods in individual SACs. It is explained by bifurcations arising purely from intrinsic excitable membrane properties. SACs can spontaneously switch to strong oscillatory activity followed by refractoriness, reflecting what has been reported experimentally in immature SACs during the period of retinal waves. The bursting frequency is determined by a Ca-activated K conductance responsible for the sAHP. This led us to propose several conjectures and possible experiments, directly inspired from the model analysis.

We would like now to develop other aspects, not considered in the main text.

Biophysical simplifications. Although our model is able to explain fairly well and reproduce several aspects in the physiology of bursting SACs, as well as to propose new experimental predictions it is yet simplified with respect to the real cellular processes in SACs, lacking potentially certain biological mechanisms with important consequences. Especially, there are several important ion channels (e.g. I_{Na})³⁶ and transporters in immature SACs that are not considered in our mathematical description. These currents make changes in the intracellular ion concentrations (ex Cl^- concentration³⁷) and make changes in the reversal potentials (e.g. V_L) of the ion channel currents. A change in V_L results in a change of the effective current $g_L V_L$ in eq. (1) and therefore can be absorbed in I_{ext} . Here, we justify further our approximations:

- The role of sodium channels. In⁶, it has been shown that voltage-gated Na^+ channels do not contribute to the bursting activity of immature SACs, since upon the application of tetrodoxin (TTX), their activity remained unaltered. Note that the paper³⁶ (10 years anterior to⁶) mention the role of Na^+ but it seems only responsible for the initial peak of voltage arising when the burst starts. This point is confirmed in⁶, see Fig. 4 of this paper. We have therefore decided to remove Na^+ channels from our modeling. Eliminating Na^+ channels allows us to consider a more simplified model than Hodgkin-Huxley, yet a canonical model in computational neuroscience, the Morris-Lecar model.
- On calcium dynamics. In^{6,12}, the authors showed that the long refractory period of SACs is controlled by Ca^{2+} -gated K^+ channels, probably SK-like, as those observed in pyramidal cells in hippocampal neurons studied in²⁶. Based on these experimental observations, we have proposed a model of sAHP dynamics. Although our modeling is widely inspired from¹², we get deeper in the biophysical description of Calcium dynamics, based on the seminal work by M. Graubner and collaborators^{27,38} (see Methods). Additionally, the parameters were tuned to match the SK channels experimental characterization made by Abel *et al.*²⁶. Yet, we have simplified the dynamics of NCX exchangers and we neglected the ionic pumps currents. These simplifications are detailed in Methods.

The Interburst variation and the interpretation of I_{ext} . Spontaneous bursting activity in immature SAC has been observed consistently across species i.e. mice, rabbits, chicks etc., but with different characteristic bursting periods. In Fig. 6, we show the relationship of the average bursting period τ_{IBI} with respect to the value of the tunable current I_{ext} whose effect could be translated on how an effective inhibition affects the characteristics of the bursting activity and even stops it completely. The source of such an inhibition could be linked either with intrinsic cell transient properties or inhibitory inputs from other layers of the retina upon maturation. Although it is unlikely

that changes in intrinsic properties of SACs would occur over the short period of stage II, such changes could be observed across species. As the interburst intervals (measured in retinal ganglion cells) increase and then decrease (in mouse) during Stage II waves (see³) the effective inhibition current I_{ext} is more likely arising from changes in synaptic inhibition (e.g. GABA A maturation or changes in the inner chloride concentration).

How identical cells can display variability in behaviour. One of the key result of this study is that SACs display bursting because they are close to a bifurcation point. As a consequence, the characteristics of bursting (IBI, amplitude) vary strongly upon tiny variations of physiological parameters such as g_K , g_C , I_{ext} , V_3 ... and noise can induce huge fluctuations in their bursting period. In a network of SACs those parameters may fluctuate. Especially, the presence of acetylcholine inducing a cholinergic current impacts the bursting period as was experimentally observed by Zheng et al. in^{6,14} and confirmed by our model^{22,23}. Therefore, a network of identical coupled SACs is expected to display a huge variability: SACs are not identical bursters. This variability has been invoked and modeled in a paper by Ford and Feller²¹ where they show how it impacts the retinal waves dynamics and provide more realistic results than an homogeneous model. In their paper, inhomogeneity is imposed ad hoc (Gaussian distribution of IBIs) and does not evolve in time. In contrast we propose here that (i) inhomogeneity purely results from the biophysical mechanisms leading SACs to burst; (ii) this heterogeneity evolves in space and time, depending on the collective dynamics (e.g. acetylcholine concentration varies in space and time and induces a variability in a given cell bursting period). In particular, one could have two populations of cells: one population acting as dynamically driven bursters with a well defined bursting period, and one population acting as noise driven bursters with a large variation of the period. Those populations would act in a different way on the collective dynamics of waves (see next section).

How inhomogeneous bursters can produce retinal waves? To our best knowledge, this model is the first able to characterize SACs dynamics in great detail and to show that biophysical properties of SACs are sufficient to explain intrinsic bursting and refractoriness in these cells. Yet, this work was concerned with individual neuron dynamics a crucial activity for the emergence of a propagating wave activity. A next step therefore consists of studying a network of coupled SACs. The crucial point is to understand how a network of coupled inhomogeneous bursters can achieve the third ingredient necessary to initiate retinal wave: synchrony. In the retina, at stage II, SACs are coupled via cholinergic synapses, whose strength evolve during development⁶. Modeling the cholinergic coupling, we observed indeed that SACs are able to achieve local synchrony and generate waves for sufficiently large value of the cholinergic conductance, compatible with experiments⁶. This is further developed in a forthcoming paper but we give here the main ideas. Local synchrony results from the cholinergic coupling via the classical mechanism of non linear synchronization between inhomogeneous oscillators, called "Arnold' tongues"39. For a sufficiently large cholinergic conductance, compatible with experiments, this synchrony leads to a wave propagation. This propagation can be described, in our model, via a transport equation allowing to compute the wave characteristics (speed, size, duration) as well as the effect of sAHP on dynamics. Because of its slow time scale, quite slower than the characteristic time of the wave propagation, sAHP somewhat imprints the medium where the wave has to propagates leading to transient spatial structures observed in Ford et al. 40. The full analysis of these aspects will be done in a forthcoming paper (see^{22–24}).

Pharmacological control. Although this model has many parameters, bifurcation analysis allows us to highlight several of them, g_C , g_K , V_3 and I_{exp} controlling important aspects of dynamics, in direct links with experiments. Varying those parameters have a deep impact on SACs dynamics as shown e.g. in Fig. 10. This is possible via a pharmacological control (e.g. TEA to vary K conductance, Cd to vary Ca conductance, synaptic input changing I_{ext} ...). It would be interesting to confirm experimentally our bifurcation diagram with such experiments.

In addition, this study opens the possibility that mature SACs are still potential bursters. Then, by a suitable pharmacological treatment, guided by the bifurcation analysis, they could start to burst in mature retinas. Would this restore a wave activity as recently observed in mature mice retinas⁴¹. This deserves however further investigations. Especially, SACs are not connected to each other in the adult retina (cholinergic synapses disappear at the end of stage II⁶). In adults SACs make synapses with bipolar cells axons, with retinal ganglion cells, and with some other amacrines types (inhibitory) providing several potential coupling mechanism to propagate waves.

Proposing new experiments towards the validation of our theoretical predictions. *How to validate experimentally a bifurcation diagram for SACs dynamics?* It would be very elucidating to conduct experiments in order to calibrate accurately SACs dynamical repertoire with respect to our bifurcation analysis shown in Fig. 2A. It would require to apply several consequent constant current pulses to an isolated immature SAC, within a wide range, and create an "experimental bifurcation diagram", showing the observed behaviors of the cell upon the variation of a constant current. Such an experiment, would shed light also on understanding the two possible drives for triggering SACs bursting which is a result of our modeling. Also, it would help validate our hypothesis about SACs being close to a bifurcation during development.

How to validate experimentally the possible role of K^+ channels on the loss of SACs excitability? One of the predictions that emerged from our model is that the Kv_3 channels, that are expressed in the SACs at adult stage²⁰, may not be expressed at stage II. To test this hypothesis, immunolabeling experiments are needed in the mouse retina, using an antibody labeling the Kv channels, both in adult and immature retina in order to compare and deduct whether the localization of Kv channels is weaker or inexistent in the immature retina (P3–P10 for mice). Along with immunochemistry experiments, actual measurements of the K^+ currents at different stages of development (we thank one of the reviewers for this comment) would elucidate their possible transient expression during development and their role in the lose of SACs excitability.

Methods

Deriving sAHP dynamics. To our best knowledge, the ionic channels types involved in sAHP for immature stage II SAC are not precisely known. However, Zheng *et al.* argue in that these channels could share characteristics with SK channels, thoroughly studied for pyramidal neurons by Abel *et al.* in 26 . On this basis we modeled SK channels dynamics. SK channels have four subunits associating to form a tetramer. The SK channel gating mechanism is controlled by intracellular calcium levels. The precise mechanism is: (i) calcium binds to the protein calmodulin forming the complex CaM where 4 ions of Ca^{2+} are fixed to calmodulin; (ii) CaM binds to a SK channel terminal to open it; (iii) 4 terminals must be open to let the SK channel open. We now model these different steps.

Saturated calmodulin production. The set of kinetic equations leading to CaM formation is widely described in M. Graupner's work 27,38 . This is a cascade of equations that we summarize in one kinetic equation, from free calmodulin, M, to the saturated one, CaM.

Let us call $k_{ass}(M^{-4}s^{-1})$ and $k_{diss}(s^{-1})$ respectively association and dissociation constants of calmodulin. Set $K_d^4 = \frac{k_{diss}}{k_{ass}}$. If we call $M_0 = [M] + [CaM]$ the total calmodulin concentration, and $S = \frac{[CaM]}{M_0}$ the fraction of saturated calmodulin, we have $\frac{[M]}{M_0} = 1 - S$ and we obtain a kinetic equation:

$$\frac{dS}{dt} = k_{ass}C^4(1-S) - k_{diss}S.$$

where C is the intracellular calcium concentration. Setting $au_S = \frac{1}{k_{diss}}$ and $au_S = \frac{k_{ass}}{k_{dis}} = \frac{1}{K_d^4}$ we arrive at:

$$\tau_S \frac{dS}{dt} = \alpha_S C^4 (1 - S) - S. \tag{10}$$

Note that equation (7) has a similar form to the one proposed by Hennig *et al.* in ¹². However, in their model our term C^4 is replaced by $\frac{C^4}{K_d^4 + C^4}$.

Binding of calmodulin to SK terminals. This corresponds to a reaction:

$$F + CaM \underset{P_{BF}}{\overset{P_{FB}}{\rightleftharpoons}} B$$
,

where F is the density of free terminals, B the density of bounded terminals, $P_{FB}(P_{BF})$ the transition rate from free to bound (bound to free).

Calling *R* the fraction of bounded terminals, $\tau_R = \frac{1}{P_{DE}}$, $\alpha_R = \frac{P_{FB}}{P_{DE}}$ we end up with a kinetic equation:

$$\tau_R \frac{dR}{dt} = \alpha_R S(1 - R) - R = \alpha_R S - (1 + \alpha_R S)R. \tag{11}$$

Finally, since R is the probability that a terminal is open and since 4 terminals must be open to let the SK channel open, the sAHP conductance is $g_{sAHP}R^4$.

Note that equation (6), is similar to Eq. (3) in $^{12}\left(\tau_R\frac{dR}{dt} = (\alpha C + S)(1 - R) - R\right)$, with a remarkable difference: in our model there is no direct dependence on calcium concentration whereas the term αC in 12 corresponds to a direct binding of Ca^{2+} to a terminal. Note that, taking the quite large value of the parameter α (~2400) considered by these authors, their equation is essentially equivalent to $\tau_R\frac{dR}{dt} = \alpha C(1 - R) - R = \alpha C - (1 + \alpha C)R$ with a steady state $R = \frac{\alpha C}{1 + \alpha C}$ very close to 1 whenever αC is quite larger than 1. In this case, S plays essentially no role.

Calcium concentration. Both variables R and S are driven by intracellular Ca^{2+} concentration dynamics, given by:

$$\tau_C \frac{dC}{dt} = -\frac{\alpha_C}{H_X} C + C_0 + \delta_C I_C(V)$$
(12)

Equation (12) is a linear approximation of a more complex equation ((15) below). This equation is similar to Hennig *et al.* (Eq. (5)) with two significant differences: (1) We have added a rest concentration C_0 avoiding unphysical situations where C can become negative; (2) the value of parameters are different.

Calcium concentration dynamics. The calcium current crossing a membrane section results from the opening of gates in ionic channels. Following³⁸ the equation for *Ca* concentration is (adapted with our notations):

$$\frac{dC}{dt} = \frac{G}{n_{cd}F} \left[\frac{I_C}{S} - J_X(C) - J_p(C) + L \right] \frac{1}{1 + \frac{dCa_{bound}}{dC}}.$$
(13)

Here $G = \frac{S}{V} = 6 \mu m^{-1} = 6 \times 10^5 dm^{-1}$ is the surface to volume ratio that accounts for the localization of the channels at the surface of the membrane, $n_{Ca} = 2$ is the calcium valence and $F = 96500 \ C \ mol^{-1}$ is the Faraday number. $\frac{dCa_{bound}}{dC}$ corresponds to a quasi-steady-state approximation for the calcium buffering where the bound calcium concentration (on calmodulin) Ca_{bound} is adapted instantaneously to the free calcium concentration C at each time. Since we have no way to estimate $\frac{dCa_{bound}}{dC}$ we shall consider it is a constant and set $\frac{1}{1 + \frac{dCa_{bound}}{dC}} \equiv K_{bound}$.

To alleviate notations we set:

$$r = \frac{GK_{bound}}{n_{Ca}F}. (14)$$

The first term in Eq. (13), $r\frac{I_C}{\varsigma}$ corresponds to an increase of internal Ca^{2+} concentration upon calcium influx (current $I_C(V)$) generated by spikes or, in experiments, by voltage clamp. As $V_{Ca} = 50 \,\text{mV}$ this current is positive unless $V > V_{Ca}$.

The second term is $-rJ_X(C)$, where

$$J_X(C) = \rho_X I_X \frac{C}{H_X + C},$$

is the efflux current density through sodium-calcium exchanger (NCX). It corresponds to an outward current through NCX exchangers, contributing to restoring the initial Ca^{2+} concentration. Here, ρ_X is the surface density of the NCX membrane proteins. We take $\rho_X=100~\mu m^{-2}=10^{12}~dm^{-2}$ (from²⁷). $I_X=4.8\times10^{-19}~Cms^{-1}=4.8\times10^{-16}~Cs^{-1}$ is the maximum ionic flux through a single NCX channel. This corresponds to 3+ charges (1.5 Ca^{2+}) per ms; $H_X=1.8~\mu M=1.8\times10^{-6}~M$ is the half activation concentration³⁸.

Also, in (13), $I_p(C)$ is the current density of Ca^{2+} pumps. We shall neglect this term from now on. Finally, L is the leakage surface current density representing the residual conductivity of the plasma membrane. We have not been able to find its value in the literature.

To summarize Eq. (13) becomes:

$$\frac{dC}{dt} = r \left[\frac{I_C}{S} - \rho_X I_X \frac{C}{H_X + C} + L \right]. \tag{15}$$

We also assume that NCX current term is approximated by a linear term. This is valid if one assumes that calcium concentration $C \ll H_X$ which is the case in our simulations.

In order to match (15) with the form (12) ((8) in the text), we set:

$$\frac{\alpha_{Ca}}{\tau_{Ca}} = r\rho_X I_X; \tag{16}$$

$$\frac{\delta_{Ca}}{\tau_{Ca}} = \frac{r}{S};\tag{17}$$

$$\frac{C_0}{\tau_{Ca}} = rL. \tag{18}$$

How does the average interburst depend on the parameter I_{ext} . In this section we analyze how the interburst interval τ_{IBI} depends on the current I_{ext} . In particular, we justify Eq. (9).

As explained in the text there are two distinct regimes where the interburst interval τ_{IBI} is constrained by different factors. In the noise driven bursting regime ($I_{ext} < I_{SN_i}$), the stochastic dynamics around the rest state is described, with a good approximation, by a Ornstein-Uhlenbeck process (upon neglecting the non linearities coming from g_C and g_K , which are small in this range of membrane potential). Therefore, fluctuations of V are Gaussian with a mean V_{rest} and a variance proportional to σ , the noise intensity. Thus, the probability to cross the bifurcation threshold and enter into the bursting regime can be easily computed. This is a sigmoid with a slope proportional to $\frac{1}{\sigma}$. As a consequence, bursting can take place when $I_{ext} < I_{SN_1}$, because of noise, but the probability to burst decreases rapidly when I_{ext} decreases, after a threshold value $I_C \equiv I_C(\sigma)$ depending on σ . Formally, the bifurcation value I_{SN_1} is shifted to I_C explaining this term in Eq. (9). For $I_{ext} < I_C$ the probability to induce bursting by noise is so small that we consider it vanishes. Here τ_{IBI} is not defined anymore but we set it to zero in Eq. (9) by convention.

In the dynamically driven bursting regime $(I_{ext} > I_{SN_1})$, τ_{IBI} is constrained by two times: (i) the time t_1 to get in a neighborhood of the bifurcation point V_{SN_1} in Fig. 3 where bursting starts (essentially determined by the sAHP) and (ii) the time t_2 to leave this neighborhood. When I_{ext} is quite larger than I_{SN_1} , τ_{IBI} is largely dominated by t_1 . Here, τ_{IBI} decreases slowly as I_{ext} increases. Indeed, the higher I_{ext} the more neurons are prone to bursting. When I_{ext} becomes too large, I_{sAHP} is not enough to stop the oscillatory phase and $\tau_{IBI} = 0$.

 I_{ext} becomes too large, I_{sAHP} is not enough to stop the oscillatory phase and $\tau_{IBI} = 0$. Close to I_{SN_1} , we can use the normal form of the saddle-node bifurcation for the variable V: $C\frac{dV}{dt} = -(I_{SN_1} - I_{ext}) + \alpha(V - V_s)^2$ where V_s is the rest state at the bifurcation point and α depends on the other parameters of the model. In this approximation, there are two branches of fixed points, existing only when $I_{ext} < I_{SN_1}$, approximated by $V_{\pm} \sim V_s \pm \sqrt{\alpha(I_{SN_1} - I_{ext})}$. The linear stability of the stable branch is given by the linearized equation $\frac{dx}{dt} = -2x\sqrt{\alpha(I_{SN_1} - I_{ext})}$ where x is a small perturbation around the fixed point. This equation has a characteristic time $\frac{1}{2\sqrt{\alpha(I_{SN_1} - I_{ext})}}$ which diverges as $I_{ext} \to I_{SN_1}$ from below. When $I_{ext} \gtrsim I_{SN_1}$ the normal

form still holds but there is no fixed point anymore. Nevertheless, around the point where bursting starts the flow has a very small amplitude. As a consequence, the time to leave a neighborhood of this point is also of order $\frac{1}{2\sqrt{\alpha(I_{SN_1}-I_{ext})}}$. In this case τ_{IBI} is largely dominated by the time t_2 to leave the neighborhood of the transition point

Physical quantity	Units
Time	ms
Potential	mV
Capacitance	pF
Current	pA
Conductance	nS
Concentrations	nM

Table 1. Units of physical quantities used in the paper.

where bursting starts. In the absence of noise, we have thus $\tau_{IBI} = \frac{K}{\sqrt{I_{ext} - I_{SN_1}}}$ for $I_{ext} > I_{SN_1}$, where $K = \frac{1}{2\sqrt{\alpha}}$. In the presence of noise the threshold value I_{SN_1} is shifted to I_c as explained above. This fully justifies Eq. (9).

Rescaled equations and Multi-time scale analysis. The dynamical system (1), (4), (6), (7), (8) has 3 characteristic time scales: fast variables V, N (of order ms); medium C (of order s); slow R, S (of order 10 s), fixed by the characteristic times given in the Table 1. In order to make explicit these time-scales separation we set $\widetilde{g}_X = \frac{g_X}{g_L}$ for conductances (X = C, K, sAHP); $\tau_L = \frac{C_m}{g_L}$, $\widetilde{I}_{ext} = \frac{I_{ext}}{g_L}$; $t_f = \frac{t}{\tau_L}$; $\widetilde{\tau}_X = \frac{\tau_X}{\tau_L}$, where X = N, C, R, S. This gives:

$$\begin{cases} \frac{dV}{dt_f} &= -(V - V_L) - \tilde{g}_C M_\infty(V)(V - V_C) - \tilde{g}_K N(V - V_K) - \tilde{g}_{sAHP} R^4(V - V_K) \\ &+ \tilde{I}_{ext} \end{cases}$$

$$\tilde{\tau}_N \frac{dN}{dt_f} &= \Lambda(V)(N_\infty(V) - N)$$

$$\tilde{\tau}_C \frac{dC}{dt_f} &= -\frac{H_X}{\alpha_C} C + C_0 - \delta_C g_C M_\infty(V)(V - V_C)$$

$$\tilde{\tau}_S \frac{dS}{dt_f} &= \alpha_S (1 - S) C^4 - S$$

$$\tilde{\tau}_R \frac{dR}{dt_f} &= \alpha_R S(1 - R) - R$$

Note that, with our parameters values, $\tau_L = 11$ ms, $\frac{1}{\tilde{\tau}_C} \sim 5.5 \times 10^{-3}, \frac{1}{\tilde{\tau}_S} = \frac{1}{\tilde{\tau}_R} \sim 2.5 \times 10^{-4}$. Therefore, on the fast time scale, one can use the approximation $\frac{1}{\tilde{\tau}_X} = 0$, X = C, S, R, so that the variables C, S, R are constant. So, fast dynamics reduces to a Morris-Lecar model (here with a fast variable N) in the presence of an additional current $I_{tot} = I_{sAHP} + I_{ext}$:

$$\begin{cases} C_{m} \frac{dV}{dt} = -g_{L}(V - V_{L}) - g_{C} M_{\infty}(V)(V - V_{C}) - g_{K} N(V - V_{K}) + I_{tot}; \\ \frac{dN}{dt} = \Lambda(V)(N_{\infty}(V) - N). \end{cases}$$
(19)

Parameters values and auxiliary functions. *Units.* In all the paper, physical quantities are expressed in the units displayed in Table 1. Having integrated over all the surface of the membrane we omit the surface units.

Calibrating parameters from experiments. All parameters values are calibrated with respect to biophysics, found in the literature or fitted from experimental curves in 14,26 and 6 . Morris-lecar tuning parameters V_1 , V_2 , V_3 , V_4 were calibrated (see Fig. 8), so as to reproduce the experiment of 6 (Fig. 4a), where the authors investigate the ionic mechanisms of the fast oscillations. Note that the bursting regime is robust to (small) variations of these parameters (results not shown). We tuned the sAHP parameters taking into account the analogy with SK channels studied in 26 (fit not shown). Also, we note that the intensity of sAHP observed by Abel *et al.* in pyramidal neurons (of order 150 pA) is quite bigger than in stage II SAC. In our model, this means a lower sAHP conductance g_{sAHP} ($g_{sAHP} = 2$ nS).

Auxiliary functions. The dimensionless auxiliary functions involved in the dynamical equations appearing in the model definition are:

$$M_{\infty}(V) = \frac{1}{2} \left[1 + \tanh\left(\frac{V - V_1}{V_2}\right) \right], \tag{20}$$

Parameter	Physical value
C_m	22 pF
g _L	2 nS
g_C	[3, 20] nS
g_K	[1, 20] nS
g_{sAHP}	2 nS
V_L	-70mV
V_C	50 mV
V_K	-90 mV
V_1	-20mV
V_2	20 mV
V_3	-25mV
V_4	7 mV
$ au_N$	5 ms
$ au_R$	8300 ms
$ au_{S}$	8300 ms
$ au_C$	2000 ms
δ_C	$10.503nMpA^{-1}$
$\alpha_{\scriptscriptstyle S}$	$\frac{1}{200^4}nM^{-4}$
α_C	4865 nM
α_R	4.25
H_X	1800 nM
C_0	88 nM

Table 2. Range of values for the parameters used in the paper.

$$\Lambda(V) = \cosh\left(\frac{V - V_3}{2V_4}\right),\tag{21}$$

$$N_{\infty}(V) = \frac{1}{2} \left[1 + \tanh \left(\frac{V - V_3}{V_4} \right) \right], \tag{22}$$

Parameters. The parameters used in the model are displayed in Table 2.

Comparison with existing models. In this section we shortly revisit models of SACs activity in the stage II and compare them to our model, (see^{11-13,40,42,43}, for a review see¹⁰). We would first like to remark that these existing models are devoted to describe wave activity and do not focus on thoroughly describing individual SACs dynamics. Especially, none of the models we know describe the biophysical mechanisms of SACs bursting activity and the role played by biophysical parameters. Instead their focus was more on having a relatively simple description of the cell activity with a minimal set of tunable parameters (a notable example is Butts *et al.* model⁴³ which has two free parameters governing the waves properties). Finally, our model is the first one, able to characterize SACs dynamics in great detail and to show that biophysical properties of SACs are sufficient to explain intrinsic bursting and refractoriness in these cells.

The closest model to ours has been proposed by M. Hennig and collaborators (refered to as "Hennig model" see also the extension by Ford and Feller and the recent paper of Xu *et al.* 44). Actually, our model has been widely inspired by this work with several notable differences. As exposed in the Methods section our biophysical analysis of sAHP dynamics leads to equations and parameters values departing from Hennig model. Additionally, Hennig model does not consider a fast potassium dynamics and there is no fast oscillation. The mechanism that mimics SACs bursting is a switch from low membrane potential level to high one. This switch is determined by an exogenous shot noise i.e. a voltage dependent rate modulated Poisson process with a slow decay. This activity is maintained long enough so that sAHP can be activated, enabling the cell to return to rest. In our model shot noise is not necessary to trigger activity. Instead, a cell can spontaneously switch to the bursting state, where it stays until the sAHP produced by its activity leads it back to the rest state. By "spontaneous" we mean literally "happening or done in a sudden way, without any planning or without being forced/without premeditation". In our model this sudden switch is a bifurcation, induced by the mere cells dynamics. The presence of a fast (Brownian) noise facilitates this transition, but, there is no need for a shot noise. The cell stays in the bursting state by its mere dynamics, even when it is isolated.

A similar modeling to Hennig holds in Lansdell *et al.* model¹³. It is ruled as well by an excitable Morris-Lecar model with a slow potassium variable linked to sAHP. There is no fast potassium and here too cells do not burst. As in Hennig model, the cell activity is triggered by a random excitatory current and maintained by network dynamics.

Although Hennig or Lansdell model are based on differential equations with many parameters, none of these authors made a bifurcation analysis of their model. We did it and we found that there is no bifurcation in a neighborhood of the parameters value they choose. In this sense, our model is in strong contrast with previous studies. This is precisely because cells are close to a bifurcation point that they are able to exhibit the wide repertoire of dynamics we have presented in close agreement with experimental findings.

References

- 1. Sernagor, E., Eglen, S. J. & O'Donovan, M. J. Differential effects of acetylcholine and glutamate blockade on the spatiotemporal dynamics of retinal waves. *J. Neurosci.* 20, RC56 (2000).
- 2. Feller, M. B., Butts, D. A., Aaron, H. L., Rokhsar, D. S. & Shatz, C. J. Dynamic processes shape spatiotemporal properties of retinal waves. *Neuron.* 19, 293–306 (1997).
- 3. Maccione *et al.* Following the ontogeny of retinal waves: pan-retinal recordings of population dynamics in the neonatal mouse. *J. Physiol.* **485**, 1545–1563 (2014).
- Sernagor, E. & Grzywacz, N. Spontaneous activity in developing turtle retinal ganglion cells: Pharmacological studies. J Neurosci. 19, 3874–3887 (1999).
- 5. Warland, D. K., Huberman, A. D. & Chalupa, L. M. Dynamics of spontaneous activity in the fetal macaque retina during development of retinogeniculate pathways. *J. Neurosci.* 26, 5190–5197 (2006).
- Zheng, J., Lee, S. & Zhou, Z. J. A transient network of intrinsically bursting starburst cells underlies the generation of retinal waves. Nat. Neurosci. 9, 363–371 (2006).
- 7. Sernagor, E. & Hennig, M. Retinal Waves: Underlying Cellular Mechanisms and Theoretical Considerations in *Cellular Migration and Formation of Neuronal Connections*. (ed. Rubenstein, J. L. R. & Rakic, P.) 909–920 (Academic Press, Oxford, 2013).
- 8. Fried, S. I., Münch, T. A. & Werblin, F. S. Mechanisms and circuitry underlying directional selectivity in the retina. *Nature.* 420, 411–414 (2002).
- 9. Yoshida, K. *et al.* A key role of starburst amacrine cells in originating retinal directional selectivity and optokinetic eye movement. *Neuron* **30**, 771–780 (2001).
- Godfrey, K. & Eglen, S. Theoretical models of spontaneous activity generation and propagation in the developing retina. Molecular BioSystems. 5, 1527–1535 (2009).
- 11. Godfrey, K. & Swindale, N. B. Retinal wave behavior through activity-dependent refractory periods. *PLoS Comp. Bio.* 3, e245 (2007).
- 12. Hennig, M. H., Adams, C., Willshaw, D. & Sernagor, E. Early-stage waves in the retinal network emerge close to a critical state transition between local and global functional connectivity. *J. Neurosci.* 29, 1077–1086 (2009).
- 13. Lansdell, B., Ford, K. & Kutz, J. N. A reaction-diffusion model of cholinergic retinal waves. PLoS Comp. Bio. 10, 1-14 (2014).
- 14. Zheng, J., Lee, S. & Zhou, Z. J. A developmental switch in the excitability and function of the starburst network in the mammalian retina. *Neuron* 44, 851–864 (2004).
- 15. Choi, H. et al. Intrinsic bursting of AII amacrine cells underlies oscillations in the rd1 mouse retina. J. Neurophysiol. 19, 1491–1504 (2014).
- Han, J. M., Tanimura, A., Kirk, V. & Sneyd, J. A mathematical model of calcium dynamics in HSY cells. PLoS Comp Bio. 13, e1005275 (2017).
- 17. Park, C. & Rubin, J. E. Cooperation of intrinsic bursting and calcium oscillations underlying activity patterns of model pre-Botzinger complex neurons. *J. Comp. Neurosci.* **34**, 345–66 (2013).
- 18. Izhikevich, E. M. Dynamical Systems in Neuroscience: The Geometry of Excitability and Bursting. (The MIT Press, Cambridge, MA, 2007).
- 19. Kaneda, M., Ito, K., Morishima, Y., Shigematsu, Y. & Shimoda, Y. Characterization of voltage-gated ionic channels in cholinergic amacrine cells in the mouse retina. *J Neurophysiol* **97**, 4225–4234 (2007).
- Ozaita, A. et al. A unique role for Kv₃ voltage-gated potassium channels in starburst amacrine cell signaling in mouse retina. J. Neurosci. 33, 7335–7343 (2004).
- 21. Ford, K. J. & Feller, M. B. Assembly and disassembly of a retinal cholinergic network. Vis. Neurosci. 29, 61-71 (2012).
- 22. Karvouniari, D., Gil, L., Marre, O., Picaud, S. & Cessac, B. Modeling the spontaneous emergence of stage II retinal waves. *Areadne Conference, Santorini, Greece* (2016).
- 23. Karvouniari, D., Gil, L., Marre, O., Picaud, S. & Cessac, B. Following stage II retinal waves during development with a biophysical model. *Bernstein Conference, Gottingen, Germany* (2017).
- 24. Karvouniari, D. Retinal waves: theory, numerics and experiments. PhD thesis (University of Côte d'Azur, March 2018).
- 25. Morris, C. & Lecar, H. Voltage oscillations in the barnacle giant muscle fiber. Biophys. J. 35, 193-213 (1981).
- 26. Abel, H. J., Lee, J. C. F., Callaway, J. C. & Foehring, R. C. Relationships between intracellular neocortical pyramidal neurons calcium and afterhyperpolarizations in neocortical pyramidal neurons. *J. Neurophysiol.* **91**, 324–335 (2004).
- 27. Graupner, M., Erler, F. & Meyer-Hermann, M. A theory of plasma membrane calcium pump stimulation and activity. *J. biol. phys.* 31, 183–206 (2005).
- Tsumoto, K., Kitajima, H., Yoshinaga, T., Aihara, K. & Kawakami, H. Bifurcations in Morris-Lecar neuron model. Neurocomputing 69, 293–316 (2006).
- Dhooge, A., Govaerts, W. & Kuznetsov, Y. A. MATCONT: A MATLAB Package for Numerical Bifurcation Analysis of ODEs. Acm Transactions on Mathematical Software 29, 141–164 (2003).
- 30. Guckenheimer, J. & Holmes, Ph. Non linear oscillations, dynamical systems, and bifurcation of vector fields. (Springer-Verlag, 1983).
- 31. Iwata, M. & Shinichi, S. Theoretical analysis for critical fluctuations of relaxation trajectory near a saddle-node bifurcation. *Phys. Rev. E.* 82, 011127 (2010).
- 32. Zhou, Z. J. Direct Participation of Starburst Amacrine Cells in Spontaneous Rhythmic Activities in the Developing Mammalian Retina. J. Neurosci. 18, 4155–4165 (1998).
- 33. Marder, E. & Goaillard, J. M. Variability, compensation and homeostasis in neuron and network function. *Nat. Rev. Neurosci.* 7, 563–574 (2006).
- Ermentrout, B. Simulating, Analyzing, and Animating Dynamical Systems: A Guide to XPPAUT for Researchers and Students. ASME. Appl. Mech. Rev. 56 (2003).
- 35. Clewley, R. Hybrid models and biological model reduction with PYDstool. *PLoS Comp. Bio.* **8**, 1–8 (2012).
- 36. Zhou, Z. J. & Fain, G. L. Starburst amacrine cells change from spiking to nonspiking neurons during retinal development. *PNAS* **93**, 8057–8062 (1996).
- 37. Zhang, L. L., Pathak, H. R., Coulter, D. A., Freed, M. A. & Vardi, N. Shift of Intracellular Chloride Concentration in Ganglion and Amacrine Cells of Developing Mouse Retina. *J. Neurophysiol.* **95**, 2404–2416 (2006).
- 38. Graupner, M. A theory of Plasma Membrane calcium pump function and its consequences for presynaptic calcium dynamics. PhD thesis, TU Dresden, Dresden (2003).

- 39. Arnold, V. Geometrical methods in the theory of ordinary differential equations. (Springer, New York, NY, 1988).
- Ford, K. J., Félix, A. L. & Feller, M. B. Cellular Mechanisms Underlying Spatiotemporal Features of Cholinergic Retinal Waves. J. Neurosci. 32, 850–863 (2012).
- 41. Kolomiets et al. Retinal waves can be reinitiated in the adult retina. Institut de la Vision preprint (2016).
- 42. Burgi, P. Y. & Grzywacz, N. M. Model for the Pharmacological Basis of Spontaneous Synchronous Activity in Developing Retinas. *J. Neurosci.* 14, 7426–7439 (1994).
- Butts, A., Feller, M. B., Shatz, C. J. & Rokhsar, D. S. Retinal Waves are governed by collective network properties. J. Neurosci. 19, 3580–3593 (1999).
- 44. Xu, H. P. et al. Retinal wave patterns are governed by mutual excitation among starburst amacrine cells and drive the refinement and maintenance of visual circuits. J. Neurosci. 36, 3871–3886 (2016).

Acknowledgements

D.K. was supported by a doctoral fellowship from Ecole Doctorale des Sciences et Technologies de l'Information et de la Communication de Nice-Sophia-Antipolis (EDSTIC). E.O. was supported by a doctoral fellowship from Ecole des Neurosciences de Paris Ile-de-France. This work also benefited from the support of the axis MTC-NSC of the University Côte d'Azur and the Doeblin federation. We warmly acknowledge Matthias Hennig and Evelyne Sernagor for their invaluable help. We thank Pr. Z. Jimmy Zhou for his authorization to reproduce Figs 8A and 10A. We thank the reviewers for their helpful criticism.

Author Contributions

D.K. contributed to the model definition-analysis and to the writing of the paper, she made numerical simulations and bifurcations studies. L.G. contributed to the model definition and to the writing of the paper. E.O. contributed to the biological interpretation of the model. O.M. contributed to the model definition and to its biological interpretation. S.P. contributed to the model definition and to its biological interpretation. B.C. contributed to the model definition and analysis and to the writing of the paper, he made numerical simulations and mathematical developments. He supervised the research and paper redaction.

Additional Information

Competing Interests: The authors declare no competing interests.

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