In Silico Determination of the Effect of Multi-Target Drugs on Calcium Dynamics Signaling Network Underlying Sea Urchin Spermatozoa Motility

Jesús Espinal-Enríquez1,2,3, Alberto Darszon4, Adán Guerrero5, Gustavo Martínez-Mekler1,2,6*

1 Instituto de Ciencias Físicas, Universidad Nacional Autónoma de México, Cuernavaca, Morelos, México, 2 Centro de Ciencias de la Complejidad, Ciudad Universitaria, México, México, 3 Instituto Nacional de Medicina Genómica, Arenal Tepeyan, Tlalpan, México, 4 Departamento de Genética del Desarrollo y Fisiología Molecular, Instituto de Biotecnología, Universidad Nacional Autónoma de México, Cuernavaca, Morelos, México, 5 Laboratorio Nacional de Microscopía Avanzada, Instituto de Biotecnología, Universidad Nacional Autónoma de México, Cuernavaca, Morelos, México, 6 Centro Internacional de Ciencias, Cuernavaca, Morelos, México

Abstract

The motility of spermatozoa of both *Lytechinus pictus* and *Strongylocentrotus purpuratus* sea urchin species is modulated by the egg-derived decapeptide speract via an oscillatory [Ca$^{2+}$]-dependent signaling pathway. Comprehension of this pathway is hence directly related to the understanding of regulated sperm swimming. Niflumic acid (NFA), a nonsteroidal anti-inflammatory drug alters several ion channels. Though unspecific, NFA profoundly affects how sea urchin sperm respond to speract, increasing the [Ca$^{2+}$]i oscillation period, amplitude, peak and average level values of the responses in immobilized and swimming cells. A previous logical network model we developed for the [Ca$^{2+}$]i dynamics of speract signaling cascade in sea urchin sperm allows integrated dissection of individual and multiple actions of NFA. Among the channels affected by NFA are: hyperpolarization-activated and cyclic nucleotide gated Na$^{+}$ channels (HCN), [Ca$^{2+}$]-dependent Cl$^{-}$ channels (CaCC) and [Ca$^{2+}$]-dependent K$^{+}$ channels (CaKC), all present in the sea urchin genome. Here, using our model we investigated the effect of blocking in silico HCN and CaCC channels suggested by experiments. Regarding CaKC channels, arguments can be provided for either their blockage or activation by NFA. Our study yielded two scenarios compliant with experimental observations: i) under CaKC inhibition, this [Ca$^{2+}$]-dependent K$^{+}$ channel should be different from the Slo1 channel and ii) under activation of the CaKC channel, another [Ca$^{2+}$]i channel not considered previously in the network is required, such as the pH-dependent CatSper channel. Additionally, our findings predict cause-effect relations resulting from a selective inhibition of those channels. Knowledge of these relations may be of consequence for a variety of electrophysiological studies and have an impact on drug related investigations. Our study contributes to a better grasp of the network dynamics and suggests further experimental work.

Introduction

Fertilization is an important process in life. Reproductive success is attained by means of different strategies that increase the probability of gamete encounter. Several species, including sea urchins, produce spermatozoa with swimming patterns regulated by egg secretions. *Strongylocentrotus purpuratus* and *Lytechinus pictus* sea urchin spermatozoa swimming is modulated by speract, a decapeptide contained in the outer coating of the egg which diffuses in the sea [1,2]. When these sperm detect speract by means of receptors along the flagellum, an intracellular signaling pathway that regulates fluctuations of the intracellular Ca$^{2+}$ concentration ([Ca$^{2+}$]i) is triggered. The pathway involves production of cyclic nucleotides and alterations in the ionic permeability of the plasma membrane (Fig 1A). These biochemical changes are timely translated to enhance the encounter of the sperm with the egg. Sea urchin spermatozoa swimming close to a surface describe circles, a very convenient trait to do imaging [3–5]. In the presence of speract gradients a cross-correlation of [Ca$^{2+}$]i oscillations and path curvatures takes place [3,5–7]. Increases in the [Ca$^{2+}$]i, are associated with sharp turning events (high path curvature) that are interspersed with periods of straighter swimming episodes (low path curvature). This swimming pattern is common to a wide variety of organisms with external fertilization [3,4,8–16].

Niflumic acid (NFA), a nonsteroidal anti-inflammatory drug, is able to block or modify several ion channels. Its lack of specificity, usually disadvantageous, turns out to be key to the profound effects it generates on how sea urchin sperm respond to speract. Immobilized *S. purpuratus* sperm exposed to NFA respond to speract with [Ca$^{2+}$]i fluctuations that are larger, longer and with increased time intervals between them than control speract
Network Multi-Target Drug Analysis for Sea Urchin Sperm Motility

Figure 1. Speract-activated \([\text{Ca}^{2+}]\) signaling pathway network model. A) Upper part: Schematic representation of the components of the signaling pathway triggered by speract in the sperm flagellum. Arrows traversing the membrane show ion fluxes. Arrows within the cell are indicative of causal relations. B) Bottom part: Signaling pathway operation diagram, black arrows correspond to activation, red lines to deactivation and yellow arrows can be activating or inhibitory depending on the relative state of the pathway elements being interconnected. Once speract binds to its receptor the several feedback loops are triggered according to the nature of the links involved. The concatenation of these loops leads to oscillatory stages of the whole pathway. The color code identifies corresponding upper and lower part components. Current models propose that the binding of speract to its receptor promotes the synthesis of cGMP that activates \(K^+\) selective and cyclic nucleotide-gated channels (KCNG) leading to membrane potential (V) hyperpolarization [3,4,7–11,18]. This V change first induces an intracellular pH increase via a \(\text{Na}^+/\text{H}^+\) exchanger (NHE) activation, [18,51,52], stimulates hyperpolarization-activated and cyclic nucleotide-gated channels (HCN) [53–55], removes the inactivation of voltage-gated \(\text{Ca}^{2+}\) channels HVA and LVA [18,56] (CaV), and facilitates \(\text{Ca}^{2+}\) extrusion by \(\text{Na}^+/\text{Ca}^{2+}\) exchangers (NCE) [51,52]. The opening of HCN and the influx of \(\text{Na}^+\) contribute to V depolarization, and concomitant increases in \([\text{Ca}^{2+}]_i\) and \([\text{Na}^+]_i\), further depolarize V. It has been proposed that the \([\text{Ca}^{2+}]_i\) increases could lead to the opening of \(\text{Ca}^{2+}\)-regulated \(\text{Cl}^-\) channels (CaCC) and/or \(\text{Ca}^{2+}\)-regulated \(K^+\) channels (CaKC), which would then contribute to hyperpolarize the V again, removing inactivation from CaV channels and opening HCN channels, [3,4,18]. It is thought that this series of events is then cyclically repeated generating a sequence of V-dependent turns. B) Network model of the signaling pathway. The network can be envisaged as a circuit where each element represents an element of the pathways and links, either in the form of arrows or lines, correspond to connections determined in the bottom part of (A). The activating or inhibitory nature of the yellow lines depends on the value of voltage (V). Yellow nodes represent binary nodes (0,1), and the four brown nodes are ternary nodes that can take values 0, 1 and 2. Changes in the node states are determined by the connected nodes by means of a regulatory function (or truth table). As an illustration we present the case of the cGMP shown at the bottom left of (B). The first 3 columns in this table contain all the possible activation states of the cGMP regulators: GC, which is an activator; PDE, an inhibitor and cGMP (cGMP) is a self-regulator; the fourth column shows the values for the function that correspond to each combination of the regulators. Additional nomenclature note: Speract receptor (SR); guanylate cyclase (GC); unknown blockers for \(\text{Ca}^{2+}\) channels HVA and LVA; dCa, dCl, dNa, dK are abbreviations for permeability changes in \(\text{Ca}^{2+}\), \(\text{Cl}^-\), \(\text{Na}^+\) and \(K^+\), respectively. doi:10.1371/journal.pone.0104451.g001

responses [17] (Fig 2). Furthermore, in S. purpuratus swimming sperm these alterations on flagellar \([\text{Ca}^{2+}]\), dynamics caused by NFA increase the speract-induced flagellar asymmetry resulting in more pronounced and prolonged sharp turns [4]. In L. pictus swimming sperm NFA inhibits chemotaxis (also in Arbacia Punctulata [15]), without altering their ability to detect the chemoattractant gradient [8].

It is because of the findings mentioned above that it is very important and interesting to unravel how NFA achieves these responses. There are not many tools that efficiently allow analysis of multiple variables regulating a transduction path; certainly a network model of a signaling pathway is one. In this regard using this approach to examine how NFA may achieve its remarkable effects on the speract signaling cascade seems warranted. This Boolean network model has already shown its ability to uncover systemic behaviors. There are not many tools that efficiently allow analysis of multiple variables regulating a transduction path; certainly a network model of a signaling pathway is one. In this regard using this approach to examine how NFA may achieve its remarkable effects on the speract signaling cascade seems warranted. This Boolean network model has already shown its ability to uncover systemic behaviors.
four individual-cell quantities measured previously by Wood et al., in [4], namely, average $[Ca^{2+}]_i$ level, amplitude, peak and frequency. We use them to compare the model predictions between oscillations in the wild type (WT) untreated network and in the NFA treated case, considering the two scenarios, depending on whether NFA inhibits or activates $CaKC$ channels.

### Methods and Models

#### The logical network

The logical signaling network corresponding to the SASP, first introduced in [18], consists of nodes interlinked according to Fig 1A, representing the principal components involved in the signaling cascade: ion channel activities, intracellular ion and molecular concentrations and the membrane potential amongst others. The network is shown in Fig 1B and the nomenclature is explained in its figure caption. To analyze the dynamics of the network, we implemented a discrete formulation that is a generalization of the Boolean approach and that has proven to be revealing for the gene regulation dynamics of many systems [38–42], as well as other cell signaling networks [43]. In this approach, the dynamical state of the network consists of a set of $N$ discrete variables $\sigma_1, \sigma_2, \ldots, \sigma_n$, each representing the state of a node. For this particular network, most of the variables take on two values, 0 and 1, depending on whether the corresponding element is absent or present, closed or open, inactive or active, etc. However, an accurate description of the dynamical processes in the network required four nodes to be represented by three-state variables: the membrane potential (hyperpolarized 0, resting 1, and depolarized 2); the low and high threshold voltage-gated $Ca^{2+}$ channels (inactive 0, closed 1, and open 2); and the intracellular $[Ca^{2+}]$ concentration $[Ca^{2+}]_i$ (basal 0, tonic 1 and supratonic 2). The state of each node $\sigma_n$ is determined by its set of regulators (which are some other nodes that also belong to the network). Let us denote as $\sigma_{n1}, \sigma_{n2}, \ldots, \sigma_{nk}$ the $k$ regulators of $\sigma_n$. Then, at each time step the value of $\sigma_n$ is given by

$$\sigma_n(t+1) = F_n \left( \sigma_{n1}(t), \sigma_{n2}(t), \ldots, \sigma_{nk}(t) \right),$$

where $F_n$ is a regulatory function constructed by taking into account the activating/inhibiting nature of the regulators. Each node has its own regulatory function. For the construction of these regulatory functions, we have made use of extensive biological knowledge, mainly of an electrophysiological nature, available to us in the literature and in our own laboratory. An illustration of such regulatory functions is given in Fig 1 for the cGMP node. For the case of $[Ca^{2+}]$, which is the main concern in our study, we have that it is regulated by 6 nodes (see origin of incoming arrows and red lines of dCa in Fig 1B). LVA, HVA and cAMP-dependent $Ca^{2+}$ channels are activators, i.e., their activation at time $t$, will favor and increase in $[Ca^{2+}]$ at the subsequent time $t+1$. There are also two inhibitory nodes: the $Ca^{2+}$ pump and the $Na^+/Ca^{2+}$ exchanger which when activated at time $t$ will favor a decrease in $[Ca^{2+}]$ at time $t+1$. Finally the sixth node is the $[Ca^{2+}]$ node itself that acts as an inactivator, hence favoring a decrease in its value at

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**Figure 2. Niflumic acid increases the $[Ca^{2+}]_i$ mean, maximum peak, amplitude and interval between successive $[Ca^{2+}]$ fluctuations.** Experimental determination of the $Ca^{2+}$-sensitive colorant Fluo-4 fluorescence in the flagellum, averaged along its length, of sperm exposed to 100 nM caged speract, in the absence of NFA (black trace) and the presence of 10 µM of NFA (red trace). Concentration is measured in florescence intensity (F.I) units and time in seconds according to the scale shown in the inset on the upper right part of the figure. The figure is a modified version of the equivalent shown in [4].

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the following time step. The balance among all the above inputs determined by physiological considerations is expressed in the form of a truth table with 432 rows, which constitutes its regulatory function. In the supplementary material S1 the regulatory functions for all the networks nodes are shown explicitly. With this model we can therefore observe in silico the effect of altering certain elements relevant to the pathway. In this paper we consider the case of NFA-sensitive channels: HCN, CaKC or GaCC. In order to test the effect of NFA on the network evolution, we proceeded with the deletion or activation of these channels according to the two scenarios mentioned above, individually, by pairs (HCN and GaCC, HCN and CaKC, CaCC and CaKC) and all three simultaneously. Within our model, node elimination takes place by assigning a zero value to all the regulatory entries in which it participates during the whole network evolution. Hence, each channel will be closed, even if its regulators are activated, thus simulating the blocking effect of NFA. Activation is put into practice by keeping the affected channel permanently open with a value of one. We calculated the modeled speract-triggered \([Ca^{2+}]\) fluctuations averaging the \([Ca^{2+}]\) node value over 10° different initial conditions of the entire network (with all nodes present) and compared the result with equivalent calculations done on the treated networks. We repeatedly checked that after 10° initial conditions the averaged \([Ca^{2+}]\) values essentially coincided with the complete initial condition calculations. With the averaged intracellular \([Ca^{2+}]\) ion concentration, determined from the model dynamics, which we shall denote by \([\langle Ca^{2+}\rangle]\), we have a representation with a finer resolution, more adequate for comparison with continuous experimental measurements.

**Steady state analysis**

For this type of network (finite number of nodes which take a finite number of values), all initial conditions lead to a periodic behavior where the network configuration is replicated after a certain number of steps. The time required to reach this condition is known as the transient time; it is important to mention that, independently of the initial condition, it is shorter than 45 steps. The attractor is reached after 22 iterations and the number of iterations between the repeated configurations is the period. These periodic solutions are the attractors of the network dynamics. For the wild type (WT) network (with all nodes present), 88.9% of all speract activated initial conditions lead to a period-4 attractor, while the remaining 11.1% converge to a period-8 attractor [18]. The total amount of initial conditions which reach a particular attractor is called the basin of attraction. In order to understand the differences between the averaged \([Ca^{2+}]\) concentration \([\langle Ca^{2+}\rangle]\), steady-state behavior of NFA-treated network and the WT, we calculate modifications of the network attractors after alteration of all combinations of NFA-sensitive channels, paying attention to the ensuing number of attractors and their associated periodicity. We make the comparisons between the characterization of the periodic behaviors of \([Ca^{2+}]\) and the period of the network attractors. This is done by means of a Fourier spectrum analysis of \([Ca^{2+}]\), from which we can determine differences in the temporal behavior of WT and NFA-treated network dynamics. We should point out that we have explored the effect of small perturbations on the regulatory functions, such as modifications in the outcome of any row in the truth tables of all the nodes. This holds particularly well for \([Ca^{2+}]\) and is a measure of the high robustness of the dynamics we are implementing and hence of the reliability of our results.

**Results**

**Individual Channel Results**

In the first column of Fig 3, we show the \([Ca^{2+}]\) fluctuations after a transient of 150 time steps, determined by the model as a function of time, for the original network and for the network with the channel specified in the figure blocked (A, D and G) and activated (J). We should point out that an extensive statistical study shows that steady state conditions are attained with less than 150 iterations. Note the changes in the peak, mean and amplitude values, comparing the colored lines against the WT (black) curve. For a more quantitative comparison, in the second column (B, E, H and K), we show the bar plot of the average over 50 time steps of the amplitude and mean of the series shown to the left. In the third column (C, F, I and L) we focus on the temporal characterization of the series by means of the Fourier power spectrum of the \([Ca^{2+}]\) fluctuations. For each row, the channel altered is the same. Looking at the right column, the lines indicate the amplitude of the dominant frequencies in the Fourier decomposition of the \([Ca^{2+}]\) fluctuations depicted on the left column. The magnitude of each line indicates the weight of an underlying periodic contribution to the \([Ca^{2+}]\) time series of a given Fourier decomposition mode. For the WT case, in Fig 3C, the first black line to the left, at the frequency 1/8 is the fundamental period-8 Fourier mode and reflects the contribution to the \([Ca^{2+}]\) of the period-8 attractor obtained from the network dynamics. The third line at 3/8 is its harmonic. The middle line at frequency 1/4 is the result of the averaging procedure of the \([Ca^{2+}]\) node network dynamics period-4 attractor together with the second harmonic of the period-8 attractor. We shall say that the \([Ca^{2+}]\) time series has a richer temporal behavior if its Fourier decomposition is more complex. Blockage of the HCN channel in our network model produces a more elaborate temporal behavior, which is registered in the Fourier power spectrum by the appearance of multiple lines that correspond to different modes in the Fourier decomposition. An opposite effect is observed with the blockage of the CaCC channel, where the Fourier decomposition consists of only one red line at the frequency value of 1/4 that corresponds to a unique period-4 attractor of the \([Ca^{2+}]\) dynamics. For the case of CaKC blockage the Fourier spectrum remains unchanged with respect to the WT; this is consistent with the persistence in the blocked network of the WT attractors. Finally, activation of the CaKC channel produces 3 peaks at 1/7, 2/7 and 3/7 frequency values which are related to a period-7 attractor obtained by the network dynamics.

**Double Channel Results**

With regard to the temporal behavior, a similar analysis to the one performed for Fig 3, reaffirms the trends found from the individual channel alterations (Fig 4), namely: i) HCN blockage leads to a more complex Fourier Decomposition, consistent with a more elaborate behavior of the \([Ca^{2+}]\) series shown on the left. ii) CaCC blockage reduces the Fourier components leading to a simpler periodic behavior. iii) CaKC activation enhances the effect of the concomitant deletion of either HCN or CaCC.

**Triple Channel Results**

With the joint alteration of the HCN-CaCC-CaKC channels the differences of the effect of blocking or activating the CaKC channel can be appreciated in Figs. 5 and 6 respectively. Fig 5A is a segment of the \([Ca^{2+}]\) time series once steady-state conditions have been reached. Fig 5B is the Fourier spectrum calculated over
1000 steady-state time steps. Besides the coincidence of the WT period 4 and 8 peaks with their harmonics as before, a new feature that arises in the dynamics of the blocked system is the occurrence of a period-9 peak and one of its harmonics. This is an indication of a richer temporal behavior. If we perform a running average of the $\frac{1}{2}Ca^{2+}$ series over a window of 4 steps, the smaller fluctuations are dampened and the behavior of an envelope at larger scales becomes evident. This is shown in Fig 5C where a recurrent module of 72 time steps comes to light. Note that 72 is the minimum common multiple of the period 8 and 9 Fourier components, this is a manifestation of a period locking phenomena at the level of the $\frac{1}{2}Ca^{2+}$ network attractors. If the running average is performed now over an 8 step window, then the WT fluctuations are completely leveled out, as well as the period-8 component of the blocked case network, allowing for the period-9 component to surface (see Fig 5D). When activation of CaKC channel is considered, we encounter the behaviors shown in Fig 6 which are determined as in Fig 5. In Fig 6A the emergence of a 3 step module absent in the WT is noticeable. This new feature is reflected in the period 3 Fourier component shown in Fig 6B. When a 4 step average is performed, the WT is strongly smoothened and a period 8 structure emerges while the structure of NFA treated case remains similar (see Fig 6C). In Fig 6D we show that though the WT is completely flatten to its over-all

Figure 3. $[Ca^{2+}]$ dynamics comparison between WT network and blocking an individual NFA-sensitive channel. The $[Ca^{2+}]$ steady state fluctuations calculated form averaging over 100,000 initial conditions. For each initial condition, $[Ca^{2+}]$ will take value 0 if it is in the basal state, 1 if it corresponds to a tonic state and 2 if it is supratonic. After averaging it will take values within the range [0–2] with a resolution of $10^5$. Concentration units are arbitrary, they comply with the above restrictions and are set for comparative purposes, time is measured by simulation steps and frequency refers to the fraction of a cycle covered by a simulation step, i.e cycles per simulation step. The values of $[Ca^{2+}]$ determined by the logical signaling network model, in the above mentioned A.U. (Arbitrary Units), are shown in black for the wild type case and in colors for the network with deletion of the node indicated in the figure: In the first column note A) for HCN, the loss of regularity; D) For CaCC, the increase in amplitude, and G) for CaKC on scenario 1, a higher average and peak values. J) CaKC on scenario 2 (activating CaKC), the decrease in peak, average and amplitude values. The central column (B, E, H and K) is the comparison between amplitude and mean of the $[Ca^{2+}]$ dynamics between the WT curve and those with altered channels. The colors are the same as in the first column. Right column (C, F, I and L): Fourier Power spectra obtained by the curves on the left.

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average, under an 8 step running average, when the effect of NFA is taken into account a module 3 structure is preserved. For the converse case of averaging over 3 steps, it is the treated case that gets flattened while the 8 module consisting of two alternating 4 modules structure of the WT survives.

Overall Picture

Given our in silico results, we can conclude from our data, the following: Blockage of the HCN channel affects mainly the temporal behavior of the $[\text{Ca}^{2+}]$ dynamics. This became evident in figures 3 and 4, where changes in the temporal structure of the $[\text{Ca}^{2+}]$ curve as well as the number of Fourier components can be observed. Another visible effect is the reduction of the $[\text{Ca}^{2+}]$ amplitude in all cases where HCN participates. When CaCC is blocked, the main effects are the increase of the amplitude in the $[\text{Ca}^{2+}]$ curve and a recovery of the regularity. Fourier components are reduced in all cases where CaCC is altered. Regarding scenario 1, in which CaKC channel is inhibited, the principal effect is an elevation in the $[\text{Ca}^{2+}]$ mean and maximum peak. In scenario 2, after activating the CaKC channel, the overall effect is quite similar to the one observed by HCN elimination: reduction in $[\text{Ca}^{2+}]$ amplitude, peak and mean but, an increase in Fourier modes. Though all of these effects are reproduced under combinations of the above channel alterations, it is possible to establish the dominance of one channel alteration over another. For example, with regard to the fluctuation amplitude we have that the effect of deleting HCN overrides CaCC blockage, however, there is a recovery of regularity, situation in which the effect on temporal behavior of CaCC blockage overrides HCN elimination (Fig 4A).

The same temporal observation is obtained by looking at the

Figure 4. Effect on the Calcium dynamics of altering the different NFA-sensitive channels taken by pairs. Average over 100,000 initial conditions of the steady-state $[\text{Ca}^{2+}]$ dynamics for the WT network is shown in black as in Figure 2. The columns are also distributed as in figure 2. A), B) and C) Blocking HCN and CaCC channels case (green line); D), E) and F) Blocking HCN and CaKC (dark green); G), H) and I) Blocking CaCC-CaKC (turquoise); J), K) and L) HCN blocked and CaKC activated; M), N) and O) CaCC blocked and CaKC activated. Notice that cases with HCN blocked produce non-regular $[\text{Ca}^{2+}]$ dynamics, opposite to the regularity generated by the CaCC blockage. Elimination of CaKC (scenario 1 in text) generates an elevation in $[\text{Ca}^{2+}]$ concentration compared with blockage of CaCC or HCN. Activation of CaKC (scenario 2) generates a decrease in $[\text{Ca}^{2+}]$, and peak, but the temporal behavior is more elaborate.

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Fourier power spectrum for the case of the joint blockage of $\text{HCN}$ and activation of $\text{CaKC}$. Both alterations result in an increase of the Fourier modes (Fig 4K) not only with regard to the WT but also to their action taken separately (Figs. 3C and 3L). This last result has a physiological explanation, because the effect of blockage of a channel which allows a cation entrance ($\text{HCN}$) or the activation of a channel which allows the cation extrusion ($\text{CaKC}$) will have a similar result in the regulation of membrane potential: shorter depolarizations.

Finally, the effect of altering the three channels together is analyzed. Here, we encounter the two scenarios mentioned previously, depending on whether $\text{CaKC}$ channel is inhibited (Scenario 1, Fig 5) or activated (Scenario 2, Fig 6).

Overall our main findings are the following:

With respect to scenario 1, our analysis shows that only when NFA blocks simultaneously the three channels under consideration, we recover the experimental observations of [4] with regard to WT conditions for the model $\left[\text{Ca}^{2+}\right]_i$ time steps, namely:

a. The mean taken over the time increases (Fig 5A, C and D).

b. The amplitude grows (Fig 5D).

c. The peak values are higher (Figs 5A, C and D).

d. The time interval between oscillations increases (Fig 5B).

In scenario 2 (Fig 6), the effect observed is a decrease in amplitude, period, and maximum peak, a trend opposite to the experiments. This behavior suggests the need of another $\left[\text{Ca}^{2+}\right]_i$ channel in the signaling pathway. An integrated presentation of our findings is condensed in Table 1. The red arrows correspond to those cases in which $\text{CaKC}$ channel is activated (scenario 2).

Table 1 clearly shows that within our modeling, experimental results from [4] can only be recovered under the joint NFA
deletion of the three HCN, CaCC and CaKC channels. The first three columns of Table 1, state the cause-effect connections of individual channel deletions: the blockage of the CaCC channel increases [Ca\(^{2+}\)] amplitude fluctuation, the blockage of the CaKC increases the [Ca\(^{2+}\)] peak and mean values, and the blockage of the HCN channel confers a more elaborate temporal behavior.

### Discussion

**Summary**

We developed a novel analysis of the effect of the multi-target drug niflumic acid, based on a logical network model of the speract-activated signaling pathway. We selectively modified the functionality of three ionic channels which are sensitive to NFA:

### Table 1. Effect of altering the NFA-sensitive channels in the Mathematical Model.

| Channels | HCN | CaCC | CaKC | HCN-CaCC | HCN-CaKC | CaCC-CaKC | All  |
|----------|-----|------|------|----------|----------|-----------|------|
| Mean     | ↓   | ↓    | ↑    | =        | =        | ↑         | ↑    |
| Amplitude| ↓   | ↑    | ↓    | ↓        | ↑        | ↓         | ↑    |
| Peak     | ↓   | =    | ↑    | ↓        | ↑        | ↑         | ↑    |
| Fourier Mode | ↑ | ↓ | = | ↑ | ↑ | ↑ | ↑ |
| Diversity | ↑ | ↓ | = | ↑ | ↑ | = | ↑ |

Compilation of results obtained by the logical network model altering one, two or the three NFA-sensitive channels in sea urchin spermatozoa. The non-bolded arrows correspond to scenario 1, in which the CaKC channel is inhibited. The bolded arrows indicate the alterations corresponding to scenario 2, with activation of CaKC channels.

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CaCC, HCN and CaKC. We calculated $[Ca^{2+}]$ mean, amplitude, maximum peak and periodicity and determined the conditions under which accordance with previous experimental findings are attained. Other approaches have been done in terms of alterations of NFA in sea urchin spermatozoa [8,44]. Although NFA is known to affect these three channels, we addressed the question of the physiological effect that NFA may exert on the $Ca_{i/C138}^{2+}$ channel in the speract activated signaling pathway (SASP) [18], and it is known that the Slo1 channel is activated by NFA [45,46], the above line of reasoning could hold if we consider the participation of a different $Ca_{i/C138}^{2+}$ channel, such as $SK_{Ca}$ or another $K^{+}$ channel in the pathway. Given the results of the last column of table 1, these findings support this alternative, hinted by the experimental discrepancy mentioned in the introduction.

In the case of scenario 2, after the first $KCNG$-dependent hyperpolarization, the activation of the $Ca_{i/C138}^{2+}$ channel will produce subsequent higher hyperpolarization values since this $Ca_{i/C138}^{2+}$ channel remains open. This in turn causes shorter lived depolarizations of a smaller magnitude, situation that leads to a premature closing of $Ca_{i/C138}^{2+}$, and it is known that the $Slo$1 channel is activated by NFA [45,46], the above line of reasoning could hold if we consider the participation of a different $Ca_{i/C138}^{2+}$ channel, such as $SK_{Ca}$ or another $K^{+}$ channel in the pathway. Given the results of the last column of table 1, these findings support this alternative, hinted by the experimental discrepancy mentioned in the introduction.

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Physiological Context

The physiological effect that NFA may exert on the $Ca_{i/C138}^{2+}$-activated $K^{+}$ channel $Ca_{i/C138}^{2+}$ in the speract-activated signaling pathway (SASP), either activating or inhibiting, can be explained in both cases (Fig 7):

i) When $Ca_{i/C138}^{2+}$ is blocked, changes in membrane potential are attained, the depolarization is more pronounced owing to the reduction of $[K^{+}]$ influx (Fig 7A, violet line), hence, the voltage-dependent $[Ca^{2+}]$ channels ($Ca_{i/C138}^{2+}$) will remain open for a longer period; consequently, $Ca_{i/C138}^{2+}$ blockage (scenario 1) produces a bigger $[Ca^{2+}]$ curve (Fig 7B, violet line). Since there is pharmacological evidence of the presence of Slo1-type $Ca_{i/C138}^{2+}$-activated $K^{+}$ channel in the SASP [18], and it is known that the Slo1 channel is activated by NFA [45,46], the above line of reasoning could hold if we consider the participation of a different $Ca_{i/C138}^{2+}$ channel, such as $SK_{Ca}$ or another $K^{+}$ channel in the pathway. Given the results of the last column of table 1, these findings support this alternative, hinted by the experimental discrepancy mentioned in the introduction.

ii) In the case of scenario 2, after the first $KCNG$-dependent hyperpolarization, the activation of the $Ca_{i/C138}^{2+}$ channel will produce subsequent higher hyperpolarization values since this $Ca_{i/C138}^{2+}$ channel remains open. This in turn causes shorter lived depolarizations of a smaller magnitude, situation that leads to a premature closing of $Ca_{i/C138}^{2+}$, and it is known that the $Slo$1 channel is activated by NFA [45,46], the above line of reasoning could hold if we consider the participation of a different $Ca_{i/C138}^{2+}$ channel, such as $SK_{Ca}$ or another $K^{+}$ channel in the pathway. Given the results of the last column of table 1, these findings support this alternative, hinted by the experimental discrepancy mentioned in the introduction.

Figure 7. Schematic representation of the effect of deletion or activation of $Ca_{i/C138}^{2+}$ channels in the speract activated signaling pathway. A) the membrane potential dynamics. B), the $[Ca^{2+}]$ dynamics. For (A) the membrane potential is depicted in black for the WT network after speract activation. Hyperpolarization (lower than $-40$ mV) due to a $[K^{+}]$ efflux via the KCNG channel, and the consequent opening of the voltage-dependent $HCN$ channel which in turn depolarizes the sperm flagellum are shown. Depolarization (higher than $-40$ mV) opens $LVA$ and $HVA$ $[Ca^{2+}]$ channels. The increase in $[Ca^{2+}]$ enhances depolarization. This last entrance of $[Ca^{2+}]$ opens the $CaCC$ and $CaKC$ channels with the corresponding influx of $[Cl^{-}]$, and efflux of $[K^{+}]$, that hyperpolarize the membrane potential. All these steps are cyclically repeated causing the $[Ca^{2+}]$ oscillation pattern depicted in (B). If the effect of the drug is an inhibition of $Ca_{i/C138}^{2+}$ channels, this would cause a bigger depolarization, because a decrease in the $[K^{+}]$ efflux. This is shown in the purple curve, notice that the amplitude is bigger than in the WT case. If the effect on $Ca_{i/C138}^{2+}$ is an activation instead, a bigger hyperpolarization is produced, due to the increase in the efflux of $[K^{+}]$. Depolarization takes less time and is smaller than in the WT case for the same reason (blue curve). For (B), the WT $[Ca^{2+}]$ dynamics are again depicted in black. Inhibition $Ca_{i/C138}^{2+}$ channels (purple curve), reduces the extrusion of $[K^{+}]$, diminishing hyperpolarization hence delaying the closure of $Ca_{i/C138}^{2+}$ channels. This will cause the elevation of $[Ca^{2+}]$, as well as the time between $[Ca^{2+}]$ peaks (the period). Overall, the activation of $Ca_{i/C138}^{2+}$ channels (blue curve), produces a shorter $[Ca^{2+}]$ oscillations due to the faster hyperpolarization, which in turn closes the $Ca_{i/C138}^{2+}$ channels avoiding a big elevation of $[Ca^{2+}]$.

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Our investigation is closely related to the temporal behavior of the \([\text{Ca}^{2+}]_i\) fluctuations. A proper understanding of temporal properties of the dynamics is fundamental for the comprehension of the link between internal biochemical oscillations that regulate molecular motors in the flagellum and the oscillatory space exploration of gradients of sperm activating peptides present in sea urchin swimming. Timing issues related to this interchange appear to be of importance for fertilization. In particular our study of the effect of NFA alterations on time-related characteristics may provide important issues on this link. Experimental analysis motivated by this work on the effect of NFA in the chemotactic response triggered by speract is provided in [8]. Finally, we would like to point out that the approach we have developed here can be used as a platform for the study of other biochemical signaling regulatory networks.

Supporting Information

Material S1  Regulatory tables for the Network Dynamics of the SASP. Each one of the 21 text files which compose this compressed file, contains the name of the element (the file name), the name of its regulators, the whole combination of the dynamical states of those regulators (at time \(t\)) as well as the state of the regulated node (at time \(t + 1\)), according to its logical rule (for more information, see Methods and Model section and figure 1).

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Author Contributions

Conceived and designed the experiments: JEE AD AG GMM. Performed the experiments: JEE. Analyzed the data: JEE AD GMM. Wrote the paper: JEE AD GMM. Designed the software used in analysis: JEE. Designed the figures: JEE GMM.
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