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Authors
Zohar, Assaf
Dekel, Noa
Rubinsky, Boris
et al.

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New Mechanism for Voltage Induced Charge Movement Revealed in GPCRs - Theory and Experiments

Assaf Zohar¹, Noa Dekel¹, Boris Rubinsky², Hanna Parnas¹*

¹ Department of Neurobiology, Hebrew University, Jerusalem, Israel, ² School of Computer Science and Engineering, Center for Bioengineering in the Service of Humanity and Society, Hebrew University, Jerusalem, Israel

Abstract
Depolarization induced charge movement associated currents, analogous to gating currents in channels, were recently demonstrated in G-protein coupled receptors (GPCRs), and were found to affect the receptor's Agonist binding Affinity, hence denoted AA-currents. Here we study, employing a combined theoretical-experimental approach, the properties of the AA-currents using the m2-muscarinic receptor (m2R) as a case study. We found that the AA-currents are characterized by a "bump", a distinct rise followed by a slow decline, which appears both in the On and the Off responses. The cumulative features implied a directional behavior of the AA-currents. This forced us to abandon the classical chemical reaction type of models and develop instead a model that includes anisotropic processes, thus producing directionality. This model fitted well the experimental data. Our main findings are that the AA-currents include two components. One is extremely fast, ~ 0.2 ms, at all voltages. The other is slow, 2 – 3 ms at all voltages. Surprisingly, the slow component includes a process which strongly depends on voltage and can be as fast as 0.3 ms at +40 mV. The reason that it does not affect the overall time constant of the slow component is that it carries very little charge. The two fast processes are suitable candidates to link between charge movement and agonist binding affinity under physiological conditions.

Introduction

Voltage gated channels were shown to exhibit charge movement associated currents, gating currents (GCs), already more than 30 years ago [1]. Since then, an overwhelming amount of experimental data was accumulated [2–6]. This data supplemented by mathematical models [7–10] indicates that voltage induced reorientation of electric charge within the channel protein produces a conformational change in the protein which leads to channel opening.

G-protein coupled receptors (GPCRs) are the largest family of proteins in the living cell and they mediate most signal transduction processes; the first step being binding of an agonist. Although being transmembrane proteins, they were not considered to be able to sense changes in membrane potential. Recently, however, it was found that several GPCRs exhibit voltage sensitivity where voltage modulates their agonist binding affinity [11,12]. Even more dramatic was the finding that GPCRs, like channels, display depolarization induced charge movement associated currents. Furthermore, a tight correlation was found between the charge that moves and the fraction of receptors that undergo a change in binding affinity at any membrane potential [13]. Because in GPCRs, the charge movement associated currents lead to alteration of the Agonist binding Affinity we denote these currents AA-currents.

Aiming at unraveling the mechanism that underlies the voltage induced change in agonist binding affinity we developed here a combined theoretical-experimental approach to study the properties of the putative voltage sensor(s), taking the m2-muscarinic receptor (m2R), a prototypical GPCR, as a cases study. We found that the AA-currents include two components. One is in the tenth of millisecond range and the other is slow, 2 – 3 ms. However, the slow component includes a fast constituent which is also in the tenth of millisecond range and carries very little charge. The two fast constituents are suitable candidates to link between charge movement and agonist binding affinity under physiological conditions.

Materials and Methods

Preparation of cRNA and Oocytes
cDNA plasmid of m2R was linearized and transcribed in vitro as described [14]. Xenopus oocytes were isolated and incubated in NDE solution composed of ND96 (in mM: 96 NaCl, 2 KCl, 1 CaCl₂, 1 MgCl₂, 5 HEPES-NaOH pH 7.5) with the addition of 2.5 mM Na⁺ pyruvate, 100 units/ml penicillin, and 100 μg/ml streptomycin. A day after their isolation, the oocytes were injected with 50 nL of the m2R cRNA, at 100 ng/μL. The oocytes were maintained at 18°C in NDE for 3 – 6 days before currents measurements.

Cut-Open Oocyte Voltage Clamp Setup
AA-currents recordings were performed with a CA-1B amplifier (Dagan, Minneapolis), as described [13,15]. Voltage commands were generated by using PCLAMP8 software (Axon Instruments, Union...
City, CA), a personal computer and a DigiData 1322A interface (Axon Instruments). Recording were performed at room temperature, unless specified otherwise. Data was sampled at 50kHz and filtered at 5kHz. Linear leak and capacitive currents were compensated by analog circuitry and online subtraction employing the pulse(P)/8mV protocol from a holding potential of +40mV
[16]. That is, administrating 1/8 of the test pulse amplitude from a holding potential where no charge movement was observed (+40mV). Then, subtracting the currents obtained from 8 such pulses from the test pulse current. The external solution contained in mM: 115 N-methyl-D-glucamine (NMDG)-methanesulphonate (NMG-MES), 2 CaCl$_2$, 20 HEPES, pH 7.5. The internal solution was similar but did not contain CaCl$_2$ and contained 2mM EGTA.

Statistical Evaluation
Significance was checked by Student’s two-tailed or one-tailed t test. Results are given as mean ± SD.

Numerical Simulations and Parameters Estimation
All the numerical operations were done using Matlab 7. The equations of the kinetic models were implemented in Matlab and were solved using stiff differential equations variable order method. Fitting eqs. 13 and 14 to the experimental AA-currents recordings was done using the curve fitting toolbox. Parameters optimization was done using multidimensional unconstrained nonlinear minimization method.

Results
Characteristic Features of the AA-Currents
Using the cut-open oocyte voltage clamp setup [15], we measured AA-currents in m2R expressing Xenopus oocytes. As seen, for the experimental protocol exhibited in Figure 1A (administrating 40ms depolarizing pulses of various amplitudes from a holding potential of −120mV), denoted standard protocol), the m2R AA-currents exhibit the following characteristic features: (i) The predominant feature is the “bump”, a complex behavior of a rather fast rise followed by a slow decay which appears both in the On and the Off responses. When existing it always follows the initial fast decay. (ii) In the On responses, the bump is clear at −70mV, less apparent at −40mV and completely disappears at more positive potentials. (iii) In the Off responses, the bump always appears irrespective of the level of the depolarizing pulse. Furthermore, the normalized responses overlap (Figure 1B). For comparison, the GCs in channels (Figure 1C) show a fundamental different behavior. Specifically, the bump does not characterize the GCs, it rather seems as an odd phenomenon appearing only at very specific conditions; in the Off response of −8mV (Figure 1C). Another important difference is that, in contrast to the AA-currents, the Off responses do not overlap. Rather, the behavior of the Off responses depend on the level of the depolarizing pulse. Because the bump is a predominant feature of the AA-currents, we designed experiments to check whether the bump is not an artifact but rather an intrinsic feature of the AA-currents. The AA-currents seen in Figure 1A were obtained by subtracting the linear capacitive currents from the total currents [15]. It is, however, possible that residual AA-currents still occur at the voltages used to assess the linear capacitive currents, hence an artificial bump is seen. To check for this possibility, we employed the P/8mV subtraction protocol from three holding potentials (+20, +30 and +40mV), all at the saturated region of the curve that describes the dependency of the charge that moves on depolarization (see later). Figure 2A shows that the kinetics of the AA-currents (normalized each to the peak amplitude of its fast component) is similar in the

Figure 1. Characteristic features of AA-currents and GCs. (A) AA-currents recorded from m2R expressing oocytes following 40ms depolarizing pulses to the indicated potentials from a holding potential of −120mV, notice the different scales. The arrows in (A) indicate the bumps observed in the AA-currents. Upper panel - the experimental protocol. Symmetric capacitive currents were subtracted by using pulses of P/8mV from a holding potential of +40mV. (B) Superposition of the results in (A) where each graph is normalized to the peak amplitude of its fast component. (C) recordings of GCs from oocytes expressing the Shaker K$^+$ channel, taken with permission from Bezanilla et al. [7], notice the different scales. doi:10.1371/journal.pone.0008752.g001
three subtraction holding potentials. In particular, the time constants of the fast decay, the slow decay and the rise of the bump obtained by fitting the AA-currents to a three exponential function, are not much affected by the subtraction holding potential (Figure 2B–D). Employing even more positive potentials for subtraction is problematic as it leads to opening of intrinsic voltage-dependent $K^+$ channels [17,18]. Using holding potentials that are more negative than $-120 mV$ are also problematic because of hyperpolarization-activated $Cl^-$ channels [18,19]. The quantitative features of the bump are not affected even if the only ions existing in the recording system, i.e., $Ca^{2+}$ and $Cl^-$, are replaced by $Ba$–$Acetate$ (Figure S1).

To further challenge our conclusion that the observed bump is an intrinsic feature of the AA-currents, we examined the dependency of the AA-currents kinetics on temperature. We focused on the Off responses where the bump always occurs. We expect that the $Q_{10}$ of the various time constants of the AA-currents, including that of the bump, will be in the range previously found for GCs in channels, i.e., $1.2 < Q_{10} < 4.5$ [20]. Figure 3A shows representative AA-currents recorded at $10^\circ$C and $20^\circ$C. As seen, the Off bump flattened at $10^\circ$C compared with $20^\circ$C. Average of 36 recordings showed that the bump amplitude was reduced by 1.5 fold ($p<0.01$) and the average time to peak of the bump was prolonged from $3.7 \pm 0.2 ms$ to $4.4 \pm 0.5 ms$ ($p<0.01$) when the temperature was reduced to $10^\circ$C. The various time constants of the Off responses (Figure 3B–D) were calculated by fitting the currents to a three exponential function. The time constants of the fast decay, the slow decay and the rise of the bump were increased by 1.3, 1.6 and 2.2 fold respectively when the temperature was reduced to $10^\circ$C ($p<0.01$).

Guide Lines in Developing a Mathematical Model for the AA-Currents

To obtain a bump, a minimum of two sequential transitions is required [21]. Accordingly, a minimal model that will produce a bump will be,

$$
\text{On response} \quad B \xrightleftharpoons[k_{+1}(V)]{k_{-1}(V)} C \xrightleftharpoons[k_{+2}(V)]{k_{-2}(V)} D
$$

$$
\text{Off response}
$$

For the model in scheme 1 to actually produce a bump two additional requirements must be met. (a) the parameters should guarantee that the 2nd transition ($C \Rightarrow D$ in the On response and $C \Rightarrow B$ in the Off response) will produce most of the measured AA-currents. For further details and means to achieve (a) see Text S1, Eqn S6. (b) Before administration of the depolarizing pulse the receptors need to populate mainly state $B$ while at the end of the depolarizing pulse the receptors need to populate mainly state $D$. This guarantees a delay in populating state $C$.

In order to account for the fast component seen in Figure 1A, an additional transition needs to be added either sequentially (scheme 2) or in parallel (scheme 3).
Charge Movement in GPCRs

that the fast component occurs in parallel to the slow one. Regarding the slow component, it must involve two transitions occurring in sequence in order to produce a bump. But, because the sequential part of the previous schemes does not guarantee a consistent depolarization independent bump in the Off responses the two transitions must occur in parallel rather than sequentially. Reconciliation of the two contradicting requirements; i.e., the two transitions occurring in parallel and in sequence, is achieved as follows. The slow component is now composed of two parallel transitions. The first transition, $R_1 \rightarrow R_2$, is faster than the second transition $C \rightarrow D$. The second transition is coupled to the first one by its voltage dependent rate constants being also dependent on the receptor configuration, $R_1$ or $R_2$. Such a model generates directionality.

The final model of the AA-currents is presented in scheme 5 and Eqn 6.

To ensure that the transition $C \rightarrow D$ (On response) will indeed follow the transition $R_1 \rightarrow R_2$, the rate constant for the transition $C \rightarrow D$ must be small when the receptor is in $R_1$ and large when in $R_2$. Similarly, to ensure that the transition $D \rightarrow C$ (Off response) will indeed follow the transition $R_2 \rightarrow R_1$ the rate constant for the transition $D \rightarrow C$ must be small when the receptor is in $R_2$ and large when in $R_1$. Accordingly,

$$\text{If the receptor is at state } R_1 \quad C \xrightleftharpoons{k_3(V)} D \quad (6a)$$

$$\text{If the receptor is at state } R_2 \quad C \xrightleftharpoons{k_4(V)} D \quad (6b)$$

To obtain $dC/dt$ we need to provide a quantitative description for $k_+(R, V)$ and $k_-(R, V)$. Following scheme 5 we notice that the receptor can be in one of four complex states; $R_1$ and $C$, $R_1$ and $D$, $R_2$ and C and $R_2$ and D. Thus, the fraction of receptors that will transit from state $C$ to $D$ with the rate constant $k_3(V)$ (Eqn 6a) will be the fraction of receptors that are in the complex state of $R_1$ and $C$. Because the $R$ states are independent of state $C(D)$
whether the receptor is in state $A$ transitions $B$ or $D$. To transform from fractions to amount of total receptors, the fraction of receptors at the complex states needs to be multiplied by $R_T$. Accordingly,

$$\frac{dC}{dt} = R_T \times \left( \frac{R_1}{R_T} + \frac{C}{R_T} \times k_3(V) \times \frac{R_1}{R_T} + D \right)$$

$$+ R_T \times \left( \frac{R_2}{R_T} + \frac{C}{R_T} \times k_4(V) \times \frac{R_2}{R_T} + D \right)$$

Rearrangement of Eqn 7 provides,

$$\frac{dC}{dt} = -\left( \frac{k_3(V)}{R_T} + \frac{k_4(V)}{R_T} \right) \times C$$

$$+ \left( k_{-3}(V) \times \frac{R_1}{R_T} + k_{-4}(V) \times \frac{R_2}{R_T} \right) \times D$$

Recalling that the transitions $R_1 \Rightarrow R_2$ do not depend on whether the receptor is in state $C$ or $D$, we obtain,

$$\frac{dR_1}{dt} = -k_3(V) \times R_1 + k_{-3}(V) \times R_2$$

For completion of the model equations, Eqn 10 describes the fast component,

$$\frac{dA}{dt} = -k_3(V) \times A + k_{-3}(V) \times B$$

The model differential equations obey the following conservation law,

$$R_T = A + B = R_1 + R_2 = C + D$$

Thus, the AA-currents model is given by Eqn 8–11. And the total AA-currents $I$ are given by the sum of the effective charges (the valence of the charge times the fraction of the electric field it traverse) carried by transitions $A \Rightarrow B$, $R_1 \Rightarrow R_2$ and $C \Rightarrow D$. Thus,

$$I(t) = -q_1 \frac{dA}{dt} - q_2 \frac{dR_1}{dt} - q_3 \frac{dC}{dt}$$

Where $q_1$, $q_2$ and $q_3$ are the effective charges carried by transitions $A \Rightarrow B$, $R_1 \Rightarrow R_2$ and $C \Rightarrow D$ respectively.

Parameters Estimation

To obtain an initial estimate of the various rate constants we aimed at achieving an analytical solution for the AA-currents. This became possible by simplifying scheme 5. Based on Eqn 6 we neglected $k_3(V)$ and $k_{-4}(V)$. Furthermore, because $k_{-3}(V)$ decreases with depolarization and increases with hyperpolarization while the opposite occurs for $k_4(V)$ we neglected $k_{-3}(V)$ and $k_4(V)$ during the On and Off responses respectively. The analytical solution is given by (for further details see Text S1, Eqn S17–S20),

$$I = \frac{X(V) \exp(-\lambda_1(V)t) + Y(V) \exp(-\lambda_3(V)t)}{\text{transitions } A \Rightarrow B}$$

$$+ \frac{Z(V) \times (\exp(-\lambda_2(V)t) - \exp(-\lambda_3(V)t))}{\text{transitions } C \Rightarrow D}$$

Where $X(V)$ is composed of the effective charge carried by transitions $A \Rightarrow B$ and of the rate constants of these transitions. $Y(V)$ is composed of the effective charge carried by transitions $R_1 \Rightarrow R_2$ and of the rate constants of these transitions. $Z(V)$ is composed of the effective charge carried by transitions $C \Rightarrow D$ and of the rate constants of these transitions. $R_1 \Rightarrow R_2$ and $C \Rightarrow D$. The exponents $\lambda_1(V)$, $\lambda_2(V)$ and $\lambda_3(V)$ are defined as the sums $k_1(V) + k_{-1}(V)$, $k_{-3}(V) + k_4(V)$ and $k_2(V) + k_{-2}(V)$ respectively (for further details see Text S1, Eqn S19).

The solution of Eqn 13 showed good agreement with the experimental results of Figure 1A (Figure S4A). It is satisfying to note that the parameters estimated from Eqn 13 corroborate the requirement of condition (a) above, necessary for a bump to be produced, that the contribution of transitions $R_1 \Rightarrow R_2$ to the AA-currents is negligible, i.e., it is a “hidden” transition (see Figure S4B). Neglecting this charge, Eqn 13 is further simplified to become,

$$I = \frac{X(V) \exp(-\lambda_1(V)t)}{\text{Fast component}}$$

$$+ \frac{Z(V) \times (\exp(-\lambda_2(V)t) - \exp(-\lambda_3(V)t))}{\text{slow component}}$$

The initial set of parameters was obtained by fitting Eqn 14 to the experimental results. The rate constants were estimated from the corresponding exponents in Eqn 14, i.e., $\lambda_1(V)$, $\lambda_2(V)$ and $\lambda_3(V)$, employing exponential voltage dependency as described in Bezanilla et al. [7]. The effective charges carried by transitions $A \Rightarrow B$ and $C \Rightarrow D$ were estimated from $X(V)$ and $Z(V)$ respectively using the already estimated rate constants.

Finally, the full model (scheme 5 and Eqn 6) was solved numerically and the initial values were adjusted (for most parameters no more than 15% change was required) such as to obtain best fit to the experimental results. Specifically, we used unconstrained nonlinear optimization method using the initial parameters estimation as an initial value. The final values of the parameters are given in Table S1.

Use of the Model to Characterize the Properties of the AA-Currents

We begin by contrasting the model and the estimated parameters (Table S1) with experimental results employing the standard protocol. For both, normalized average results with SD are provided. The normalized average of the simulation results was obtained as follows. Parameters were estimated for each of five individual oocytes. Then, five simulations of the AA-currents corresponding to the five sets of parameters were conducted. Finally each simulation was normalized to the peak amplitude of the fast component of its corresponding experimental result. This was done to contrast both the simulations kinetics and amplitude with the experimental currents. Then, as for the experiments, the
average of the simulation results was found. Figure 4 shows that the simulations (gray lines) and the experiments (black lines) match very well both in the On and the Off responses. Also seen that the model successfully accounts for the characteristic features of the AA-currents (i–iii above).

To further characterize the behavior of the AA-currents we need to estimate the time-constants of each component and the dependency of the charge that moves by each component on voltage \(Q-V\). A straightforward approach to evaluate the experimental time-constants would be by fitting the results to a double exponential decay function [7]. This procedure is not efficient in the present case due to the existence of a prominent bump. Because Eqn 14, a sum of three exponents, was shown to faithfully describe the experimental results (Figure S4C), we evaluated the time-constants of the fast component and that of the two constituents of the slow component from this equation. Figure 5A shows that the time-constant of the fast component (filled circles) around \(0.2\) ms and depends only weakly on voltage. The time-constant of the slow constituent of the slow component (filled diamonds) exhibits a parabolic dependency on voltage with values ranging between 2 to 3 ms. The surprising behavior concerns the time-constant of the other constituent of the slow component (filled triangles); it exhibits strong voltage dependency ranging from 3 ms at \(-100 mV\) to around 0.3 ms at \(+40 mV\).

The time-constants of the various components can be also evaluated from the model. Here, the time-constant of the fast component corresponds to transitions \(A\Rightarrow B\). The time-constant of the slow constituent of the slow component corresponds to transitions \(C\Rightarrow D\) and the fast constituent of the slow component corresponds to transitions \(R_1\Rightarrow R_2\). It can be seen that the time-constants evaluated from the model (Figure 5A, open symbols) match well the experimental ones. Because transitions \(R_1\Rightarrow R_2\) carry insignificant fraction of the charge, these transitions cannot be detected experimentally. Incorporating transitions \(R_1\Rightarrow R_2\) into the model was necessary for generation of the directionality required to account for the bump observed in the On and the Off responses. It is thus important to examine whether the characteristic properties of the transitions \(R_1\Rightarrow R_2\) (insignificant charge and strong voltage dependency) are indeed essential. We show in Figure S5 that if these transitions carry charge larger than 20% of the charge carried by transitions \(C\Rightarrow D\) or if they show weak voltage dependency the model fails to show a bump.

To evaluate how much charge is carried by each component we show in Figure 3B their respective \(Q-V\) curves. We see that as for the kinetics, the model also describes well the steady-state features of the AA-currents. The total charge (integral of the total current) shows a characteristic sigmoid shape. Also seen (extracted from Eqn 14) is that the slow component carries most of the charge (70%). The \(Q-V\) curve of the fast component, which is controlled by the transition \(A\Rightarrow B\), lacks the characteristic initial plateau (Figure 5C, squares), implying that a significant fraction of \(A\) shifted to \(B\) already at voltages lower than \(-120 mV\), which could imply that this transition is physiologically irrelevant. Indeed, when \(k_1\) and \(k_{-1}\) depend exponentially on voltage (Figure 5C, black lines) we find that already at \(-120 mV\) more than 60% of the receptors are at state \(B\) (Figure 5C, gray lines). Due to the physiological importance of this issue and because the estimation of the rate constants was performed with the constraint of exponential voltage dependency [7], we re-estimated the parameters but now relaxing this constraint. The parameters obtained from this fit are summarized in Table S2. Surprisingly, although no constraints were employed, the rate constants \(k_1\) and \(k_{-1}\) exhibit a sigmoid voltage dependency (Figure 5D, black lines) while the rest of the rate constants maintained the exponential voltage dependency.

Figure 4. Comparing simulation and experimental results employing the standard protocol (40 ms pulse duration, \(-120 mV\) holding potential). Average ± SD (n=5) currents at indicated depolarizing pulses, model, gray lines, experiments, black lines. The experimental currents are normalized, each to the peak amplitude of its fast component and the simulations are normalized, each to the peak amplitude of the corresponding experimental fast component (see text for details).

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Figure 5. Comparing average (n=5) simulation and experimental results employing the standard protocol. (A) Time constants of the various transitions, \(A\Rightarrow B\) (circles), \(R_1\Rightarrow R_2\) (triangles) and \(C\Rightarrow D\) (diamonds). Here and below, open symbols correspond to the model and filled symbols to the experiments. (B) \(Q-V\) curves of the total charge (diamonds), the fast component (squares) and the slow component (triangles). (C) and (D) Average occupancies of the fast component states \(A\) and \(B\) (gray lines), the values of \(k_1(V)\) and \(k_{-1}(V)\) (black lines) and the simulated normalized charge carried by the fast component (squares). In (C) the rate constants were estimated under the constraint of exponential dependency on voltage [7], while in (D) the rate constants were estimated after relaxing this constraint (Table S1 and Table S2 respectively). doi:10.1371/journal.pone.0008752.g005
Figure S6 depicts a comparison between the experiments and the simulations employing unconstraint parameters. It is seen that the fit to experimental results is even better than the very good fit obtained with the constraint parameters. However, in contrast to the behavior with the constraint parameters, the transitions $A \rightarrow B$ exhibit significant voltage dependency under the physiological range of the action potential (Figure 5D, gray lines). Specifically, at membrane potential of $-120\, mV$ more than 90% of the receptors are at state $A$ while at membrane potential of $+40\, mV$ more than 90% of the receptors are at state $B$. Furthermore, under these conditions the $Q-V$ curve of the fast component exhibits the characteristic initial plateau (Figure 5D, squares).

Further Experimental Validation of Key Assumptions of the Model

So far we have shown that the model fits well various aspects of the experimental results. We, nevertheless, wish to further test the model by examining whether its key assumptions can be validated experimentally. To do so, we first examine how the model generates the characteristic features of the AA-currents.

What generates the bumps in the On responses? Recalling Eqn 6, it is the transition $C \rightarrow D$ employed with the rate constant $k_4(V)$ (receptors that are at state $C$) which is expected to form the bump during the On response. Indeed, the occupancy of the receptors that are at state $C$ and $R_2$ (denoted $C \times R_2$) shows a bump like behavior (Figure 6, left panel). Comparing the time to peak of $C \times R_2$ to that of the bump at the same voltage (Figure 4, $-70\, mV$) we find that they match. The rising phase of the bump is thus controlled by the transitions $R_1 \Rightarrow R_2$.

Why is the bump in the On response detected only at low depolarizations? Examining the time course of the fast transition $A \rightarrow B$ at $+40\, mV$ (Figure 6, lower panel) and at $-70\, mV$ (Figure 6, upper panel), we find that the two are very similar. In contrast the transition $R_1 \rightarrow R_2$ is much faster at $+40\, mV$ than at $-70\, mV$ (0.3 and 2 ms respectively). Therefore, although a bump is formed both at $-70\, mV$ and at $+40\, mV$, it will be detected only at $-70\, mV$ because at $+40\, mV$ it merges with the fast transition $A \rightarrow B$.

Figure 6. How the model generates the AA-currents features. Occupancies of relevant states of the model during the On (left column) and the Off (right column) responses employing the standard protocol. The simulation results were obtained using the unconstraint parameters (Table S2).

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Why is the bump in the Off response always observed irrespective of the depolarizing pulse? In the Off response it is the occupancy of $D \times R_1$ which correlates with the bump [Figure 6, right panel]. As seen, the time to peak of the bump, which is governed by the transition $R_2 \rightarrow R_1$, is very similar at both $-70\, mV$ and $+40\, mV$. This is because the slow time constant of the transition $R_2 \rightarrow R_1$ corresponds to $-120\, mV$ (the holding potential). At the same time, the fast component, whose rate constant also corresponds to $-120\, mV$, remains fast. Consequently, the bump is distinct from the fast component at all voltages.

To test the above conclusion experimentally, we employed a “reverse” protocol. That is, we now administered pulses which are below $-80\, mV$ from a holding potential of $+40\, mV$. We expect to see a bump during the On response for all these pulses. Upon return to the holding potential of $+40\, mV$ (now the Off response) we expect a double exponential decay. Figure 7A shows that these predictions are fully met. The $Q-V$ curve of the reverse protocol is given in Figure 7B. It is seen that the model predictions (gray lines) match extremely well both the kinetics (Figure 7A) and the steady-state (Figure 7B) results. Since the model parameters were estimated from the standard protocol, the excellent match between the model and the reverse protocol results provides further support to the model.

Providing support to the notion that transitions $R_1 \Rightarrow R_2$ govern the bump, we now test the notion that the rate constants of these transitions depend strongly on voltage. To do so, we will repeat the experiment of Figure 1A but with a brief pulse instead of the standard 40 ms pulse and will focus on the Off response. The rational underlying such experiments is as follows. The Off response will be measured because the bump during the Off response depends on the fraction of receptors that populate states $D$ and $R_2$ at the end of the On response. We recall that the time constant of the transitions $R_1 \Rightarrow R_2$ is $\sim 3\, ms$ at low depolarizations, but it is $\sim 1\, ms$ at high depolarizations. Thus, with a 3 ms pulse we expect to see almost no bump at low depolarizations and a prominent bump at high ones (Figure 8A, left panel). Figure 8A, right panel, shows that these predictions are fully met by the experimental results. As expected, following a pulse of 1 ms (shorter than the time constant of transitions $C \Rightarrow D$) no bump is formed at any voltage. Figure 8B depicts the $Q-V$ curves of the various pulse durations normalized to that obtained following 40 ms pulse. As seen, the model also accounts for the steady-state behavior of the short pulses.

Figure 7. Kinetics of the AA-currents and $Q-V$ curves employing the reverse protocol. (A) Kinetics of the experimental and simulation results. The currents are normalized as describe in Figure 4. Here and in B, experiments, black lines and simulations gray lines. (B) $Q-V$ curves of the total charge. Experimental and simulation results are presented as mean $\pm$ SD ($n=5$). The simulation results were obtained using the unconstraint parameters (Table S2).

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Furthermore, because following the 1ms pulse only the fast component occurred, this result further supports the conclusion that the fast component carries only 30% of the charge (Figure 5B).

Discussion

We examined several models to their ability to account for the main characteristic features of the AA-currents. We found that only a non-linear model with rate constants that guarantee directionality can match the experimental results.

Conventional modeling of chemical reactions employ the Curie theorem which states that forces of one tensorial order cannot couple with fluxes of a different order. In the Curie theorem sense the affinity of a chemical reaction which is a scalar cannot couple with a vector such as the transition of a carrier from one side of the membrane to the other. In that sense a chemical reaction cannot drive a directional process, it can only cause activation or deactivation of a carrier. This concept is also closely tied to the concept of thermodynamic equilibrium. Conventional chemical reactions represent states of thermodynamic equilibrium. Therefore, conventional chemical reaction formulations of equilibrium are not rigorously correct when describing a vectorial process.

In our study we have encountered a situation in which there is a directional flux of electrical charges, i.e., a vectorial process that is not in a state of thermodynamic equilibrium. The bump pattern which we have observed (Figure 1A) involves a process that is clearly not in thermodynamic equilibrium and has directionality. Indeed attempts to employ conventional chemical reaction equilibrium formulations to describe the process failed. Therefore we used a new constitutive type of formulation in which we imposed a vectorial behavior to the process. It has been proposed [23] that anisotropy can act as a universal feature of vectorial couplings, the Curie theorem notwithstanding. We adopted this concept in Eqn 6. We showed that the constitutional anisotropic representation could indeed represent the experimentally observed process. Since the only fundamental thermodynamic criteria that constitute relations must satisfy are the laws of conservation, we have checked conservation of mass and have shown that the new formulation satisfies conservation of mass (Eqn 11).

Which of the AA-currents components could be responsible for the observed [11–13] voltage induced agonist binding affinity change? We previously showed [13], regarding the m2R, a tight correlation between the dependence of the charge that moves on voltage \(Q-V\) and the dependence of the fraction of receptors in low affinity state on voltage \(R^L-V\), suggesting that it is the charge that moves that drives the change in the GPCR’s binding affinity. Under physiological conditions the voltage induced change in binding affinity is expected to be produced by the brief (~1ms) action potential. Indeed this was shown to be the case for release of acetylcholine [24] and glutamate [25] from nerve terminals. Therefore, the two fast components, one from the fast voltage sensor and the hidden one from the slow voltage sensor, exhibit an appropriate time constant (faster than 1ms). The fast voltage sensor is a natural candidate. This is because we showed here that this fast component contributes 30% of the total charge that moves. Hence, it could play a major role in relaying charge movement to changes in conformation of the receptor and as a result change in binding affinity. The situation is different regarding the fast component of the slow voltage sensor (the transitions \(R_1 \rightarrow R_2\)), which does not display significant charge movement. Therefore, at first sight, this component is not expected to play a role in relaying charge movement to changes in conformation of the receptor. The question is then whether this fast component, whose rate constants strongly depend on voltage, can nevertheless be responsible for change in binding affinity. We argue that this could well be the case. It has been suggested regarding class A GPCRs (to which the m2R belongs) that a network of salt bridges forms an ionic lock that is disrupted during receptor activation [26–30]. It is thus quite possible that the transitions \(R_1 \rightarrow R_2\) represent a voltage dependent break of a salt bridge, hence the strong voltage dependency of the rate constants. This break, in turn, could cause a chain of conformational changes in the receptor resulting in changes in agonist binding affinity.

Our data is too preliminary to suggest a biophysical mechanism for the charge movement in GPCRs. However, it is possible that the main difference in the kinetics of GCs in channels and AA-currents in GPCRs stems from the lack of voltage dependent break of a salt bridge in the former.

Supporting Information

Text S1 The supporting information text and equations. Found at: doi:10.1371/journal.pone.0008752.s001 (0.11 MB PDF)

Figure S1 The effect of the ions present in the external solution on the kinetics of the AA-currents. On and Off currents elicited in
the m2R expressing oocytes following 40ms depolarizing pulses to the indicated potentials from −120mV. Standard external solution (black lines, see Methods) and when 2mM of CaCl2 was replaced by 2mM of Ba-Acetate (gray lines). The graphs are normalized, each to the peak amplitude of its fast component and are presented as mean ± SD (n = 4–9).

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Figure S2 AA-currents predicted by the sequential (scheme 7) and the parallel (scheme 8) models employing the standard protocol to the indicated potentials, (A) and (B), respectively. The Off AA-currents predicted by the slow component of the parallel model (scheme 8) are depicted in the inset of (B). The graphs are normalized, each to the peak amplitude of its fast component. The pulse protocol is presented on top.

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The transitions A expressing oocytes employing the standard protocol (black lines) superimposed with the three exponential fitting function, Eq. 20 (gray lines). (B) The total charge (circles) and the charge carried by the transitions A→B (triangles), R1→R2 (diamonds) and Ca-D (squares). (C) AA-currents recordings from m2R expressing oocytes employing the standard protocol (black lines) superimposed with the three exponential fitting function, Eq. 21 (gray lines). (D) Separate plot of the fast and the slow components, solutions were obtained from the same equation (Eq. 21) that was used to fit the experimental results seen in (C). Black lines represent the fast component (\( Z_e \exp(-\lambda_1 t) \)) and gray lines represent the slow component (\( Z_s \exp(-\lambda_2 t) - \exp(-\lambda_3 t) \)).

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Figure S4 Fitting of Eq. 20 and 21 to the experimental AA-currents recordings. (A) AA-currents recordings from m2R expressing oocytes employing the standard protocol (black lines) superimposed with the three exponential fitting function, Eq. 20 (gray lines). (B) The total charge (circles) and the charge carried by the transitions A→B (triangles), R1→R2 (diamonds) and Ca-D (squares). (C) AA-currents recordings from m2R expressing oocytes employing the standard protocol (black lines) superimposed with the three exponential fitting function, Eq. 21 (gray lines). (D) Separate plot of the fast and the slow components, solutions were obtained from the same equation (Eq. 21) that was used to fit the experimental results seen in (C). Black lines represent the fast component (\( Z_e \exp(-\lambda_1 t) \)) and gray lines represent the slow component (\( Z_s \exp(-\lambda_2 t) - \exp(-\lambda_3 t) \)).

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Figure S6 Comparing average (n = 5) simulation and experimental results employing the standard protocol. (A) Kinetics of AA-currents, experiments, black lines, simulations, gray lines. The currents are normalized as describe in Fig. 4 (see text). (B) Time constants of transitions A→B (circles), R1→R2 (triangles) and Ca→D (diamonds). In all, open symbols correspond to simulations while filled symbols correspond to experiments. (C) Q–V curves of the total charge and the fast and slow components. Diamonds, total Q–V, triangles, Q–V of the slow component and squares, Q–V of the fast component. In all, open symbols correspond to the model and filled symbols to experiments. The parameters were estimated after relaxing the constraint of exponential dependency on membrane potential (Table S2).

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Table S1 List of the parameters and the standard deviations estimated for the AA-currents model (text, scheme 4).

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Table S2 List of the unconstraint parameters and the standard deviations estimated for the AA-currents model (text, scheme 4).

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Author Contributions

Conceived and designed the experiments: AZ ND HP. Performed the experiments: ND. Analyzed the data: AZ. Wrote the paper: AZ ND BR HP. Supervised the theoretical and experimental work: HP. Developed and simulated the model: AZ. Suggested the experiments that were done to test the model: AZ. Supervised the theoretical work: BR. In particular, contributed to the anisotropic representation of GCs in GPCRs: BR.

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Figure S3 AA-currents predicted by the cyclic model (scheme 9). Computed AA-currents employing 40ms depolarizing pulse to −70mV from a holding potential of −120mV. The model was assigned with parameters that satisfy the conditions in Eqs. 15 and 16 and microscopic reversibility. Inset - the initial phase of the Off response.

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Figure S5 Examining the behavior of transitions R1→R2. (A) Predicted AA-currents following 40ms depolarizing pulse to −70mV from holding potential of −120mV. The three simulations differ in the value that was assigned to the effective charge carried by the transition R1→R2. The values were 0, 20 and 40% of the effective charge carried by the transition C→D (blue, green and red lines respectively). Here and below, the graphs are normalized each to the peak amplitude of its fast component. (B)
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