Conformational Flip of Nonactivated HCN2 Channel Subunits Evoked by Cyclic Nucleotides

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ABSTRACT Hyperpolarization-activated cyclic nucleotide-modulated (HCN) channels are tetrameric proteins that evoke electrical rhythmicity in specialized neurons and cardiomyocytes. The channels are activated by hyperpolarizing voltage but are also receptors for the intracellular ligand adenosine-3',5'-cyclic monophosphate (cAMP) that enhances activation but is unable to activate the channels alone. Using fcAMP, a fluorescent derivative of cAMP, we analyzed the effect of ligand binding on HCN2 channels not preactivated by voltage. We identified a conformational flip of the channel as an intermediate state following the ligand binding and quantified it kinetically. Globally fitting the time courses of ligand binding and unbinding revealed modest cooperativity among the subunits in the conformational flip. The intensity of this cooperativity, however, was only moderate compared to channels preactivated by hyperpolarizing voltage. These data provide kinetic information about conformational changes proceeding in nonactivated HCN2 channels when cAMP binds. Moreover, our approach bears potential for analyzing the function of any other membrane receptor if a potent fluorescent ligand is available.

INTRODUCTION

The activity of hyperpolarization-activated cyclic nucleotide-modulated (HCN) pacemaker channels (1–3) evokes electrical rhythmicity in various types of brain neurons (4–11) and specialized heart cells (12–15). The channels are primarily activated by hyperpolarization of the membrane voltage (16,17). In addition to this, the second messenger adenosine-3',5'-cyclic monophosphate (cAMP), produced upon sympathetic stimulation, can bind to the channels and further enhance activation (18–22), resulting in an acceleration of the electrical rhythm (7–10).

Structurally, HCN channels are tetramers. In mammals, four closely related genes, HCN1–HCN4, encode four homolog subunits (17,23) in which each contains a cyclic-nucleotide binding domain (CNBD) in its C-terminus (24). When expressed heterologously, all four isoforms can form functional homotetrameric channels (2,3,5,13,25).

The mechanism underlying the activation of HCN channels by voltage is poorly understood. The usage of voltage-clamp fluorometry in related spHCN channels provided the surprising result that the activation of only two of the available four subunits suffices to fully open the channels (26). When activating HCN channels in the presence of cAMP, the activation time course becomes accelerated and steady-state activation is shifted to less negative voltages, indicative of an intimate coupling between voltage- and ligand-induced activation. Experiments with channels formed by concatenated subunits containing a variable number of disabled binding sites suggest that in channels preactivated by voltage, all four subunits are involved in the generation of maximum activation (27). To quantify the dual gating of HCN channels, a cubic model has been proposed in which the voltage-independent closed-open transition is intimately coupled to a voltage step and a cAMP binding step (24,28). However, the degree of simplification in such a cubic model is a priori high because it ignores the tetrameric structure of the channels.

For ligand-induced activation of HCN2 channels preactivated by hyperpolarizing voltage, we recently substantiated a Markovian model that contains four binding steps and closed-open isomerizations from each of the five available closed states (29). Notably, the equilibrium association constants for the four binding steps, i.e., the microscopic affinity, reveals a cooperativity sequence of “positive-negative-positive” for the second, third, and fourth binding step, respectively (29). This cooperativity sequence differs fundamentally from the exclusively positive cooperativity for the four oxygen molecules binding to hemoglobin, probably the best-studied allosteric protein of all (for review, see Perutz et al. (30)). Consequently, Monod-Wyman-Changeux (MWC) models (31), with a constant allosteric factor used to describe the oxygen binding to hemoglobin, are invalid to describe the ligand-induced gating in HCN2 channels. Our previous analysis also showed that in HCN2 channels, preactivated by voltage, channel conformations with four, two, or zero ligands bound are more stable compared to channel conformations with either three ligands or one.

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ligand bound (29,32). This detailed insight into the gating of HCN2 channels was enabled by using confocal patch-clamp fluorometry (33) and jumps of a fluorescent cAMP (fcAMP) (29,34).

Our previous analyses also showed that not only HCN2 channels preactivated by a hyperpolarizing voltage pulse bind fcAMP, but also nonactivated channels at the voltage of −30 mV (34) and that channel activation enhances the overall affinity at the binding sites of the HCN2 channels by a factor of three, which was affirmed later by an alternative approach (35). However, the mechanism of binding in nonactivated channels has not been elucidated so far. In particular it is unanswered whether in nonactivated HCN2 channels the ligands bind independently or in a cooperative manner by interacting the subunits. It is also unknown how high the rates of binding and unbinding to the subunits are and how they differ from the values of a preactivated channel, and whether or not there are rate-limiting conformational changes associated with the ligand binding.

Herein we applied confocal patch-clamp fluorometry with jumps of the fcAMP concentration to nonactivated HCN2 channels. Analyzing time courses of ligand binding and unbinding and the steady-state concentration-binding relationship by Markovian models allowed us to identify a conformational flip (36,37) of the channel after the ligand binding and to quantify the kinetics of this flip. Furthermore, we quantified the interaction of the subunits, which was less pronounced compared to channels preactivated by hyperpolarizing voltage.

MATERIALS AND METHODS

Oocyte preparation and cRNA injection

Oocytes were obtained surgically under anesthesia (0.3% 3-aminobenzoic acid ethyl ester) from adult females of Xenopus laevis as described in Thon et al. (38).

Confocal patch-clamp fluorometry

Experiments were performed in inside-out macropatches of Xenopus laevis oocytes expressing homotrameric HCN2 channels of the mouse (NM_008226) (34). All measurements were started 3.5 min after patch excision to minimize run down phenomena. The bath solution contained 100 mM KCl, 10 mM EGTA, and 10 mM HEPES (pH 7.2). The pipette solution contained 120 mM KCl, 10 mM HEPES, and 1.0 mM CaCl2 (pH 7.2). Jumps of the ligand concentration (from zero to either 0.075, 0.25, 0.75, 2.5, or 7.5 μM, and back to zero) were performed by a double-barreled #glass pipette mounted on a piezo-driven device (39). The recording rate of our images was 10 Hz for recording the time courses and either 8.3 or 10 Hz for determining the steady-state values. The solution exchange at the pipette tip was completed within 1 ms. At the very membrane patch inside the pipette, the solution exchange is slowed by additional diffusion within the confined volume and effects of cytosolic cell material sticking at the patch. In the time courses analyzed herein, the solution exchange was shorter than 100 ms, the duration for one frame. Current recording was performed with the ISO3 hard- and software (MFK, Niedernhausen, Germany; sampling rate 200 Hz, 4-pole Bessel filter set to 2 kHz).

The fluorescence intensity in the patch was measured by patch-clamp fluorometry (40) combined with confocal microscopy. The method has been described in detail previously (33). As fluorescent ligand we used 8-DY547-AET-cAMP (fcAMP) (29,34). Recording was performed with the confocal microscope LSM 710 (Zeiss, Jena, Germany). All fluorescence signals were normalized with respect to the maximum fluorescence, $F_{\text{max}}$, determined at the voltage of −130 mV in the presence of the saturating fcAMP concentration of 15 μM.

To exclude that fcAMP binds to hypothetical additional binding sites outside the CNBD, eventually distorting our data, we performed control experiments with HCN2 channels in which a point mutation in the CNBD (R591E) generates a decrease of the apparent affinity for cAMP by more than three orders of magnitude (19,27). Functional expression of HCN2(R591E) channels was tested by voltage steps to −130 mV. With the saturating fcAMP concentration of 15 μM we obtained at −30 mV 0.9 ± 0.8 a.u. (n = 5) for patches from oocytes injected with HCN2(R591E) and 5.0 ± 1.2 a.u. (n = 4) for patches from water-injected oocytes. Both values are much smaller than the mean fluorescence of 34.7 ± 5.4 a.u. (n = 8) obtained for HCN2 channels. These experiments rule out any binding of fcAMP to additional binding sites outside the CNBD.

Data analysis

Steady-state concentration-binding relationships were fitted by

$$F/F_{\text{max}} = 1/\left(1 + (BC_{50}/[\text{fcAMP}])^H\right),$$

where $F$ is the actual fluorescence, $F_{\text{max}}$ the maximum fluorescence at −130 mV and 15 μM fcAMP, $BC_{50}$ is the fcAMP concentration generating the half-maximum binding and $H$ is the Hill coefficient. The time courses of ligand binding and unbinding were fitted according to

$$F/F_{\text{max}} = A[1 - \exp(-F/F_{\text{max}})]$$

and

$$F/F_{\text{max}} = A \exp(-t/\tau_a),$$

where $\tau_b$ and $\tau_a$ are the time constants of binding and unbinding, respectively. The value $A$ is a scaling factor and $t$ is the time. Data are shown as mean ± SE. Fits of equations to data points were performed with the software Origin8 (Northampton, MA).

Fitting Markovian models

The rate and/or equilibrium constants for Markovian models were obtained by globally fitting normalized averaged time courses of fcAMP binding and unbinding, following concentration jumps, and the respective steady-state values of binding. When building averaged time courses, each individual trace was considered to contribute with the same weight, i.e., the different number of channels in the patch was not considered. To determine the rate constants for a given model, the averaged and normalized time courses of binding/unbinding at 0.075, 0.25, 0.75, 2.5, and 7.5 μM fcAMP were subjected to a global fit together with the steady-state values at the same concentrations and one additional concentration of 15 μM, using a modified Levenberg-Marquardt algorithm (41,42). The approach is essentially the same as that reported earlier (29,33).

In brief: the goodness of the fit was judged by determining the χ² value from the fitted curves,

$$\chi^2 = g \left[ \sum_{j=1}^{n_m} \sum_{i=1}^{n_c} \frac{(F(t_j,x_i) - F_c(t_j,x_i))^2}{\sigma_F(t_j,x_i)^2} + \sum_{i=1}^{n_c} \frac{(B(t_i,x_i) - B_c(t_i))^2}{\sigma_B(t_i)^2} \right],$$

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where \( F_d(t,x) \) are mean fluorescence values, measured at time \( t_j (n_d = 60) \) and concentration \( x_i (n_i = 5) \) and normalized to the maximum value at the end of the activation interval. \( F_d(t,x) \) are the corresponding data points calculated by the fit. The square of the deviations at time \( t_j \) and concentration \( x_i \) was weighted by the reciprocal values of the observed variance \( \sigma_r^2 \) (\( t_j,x_i \)) calculated from the set of the normalized individual traces. \( B_{i}(x_i) \) are the steady-state values of the normalized fluorescence intensity (degree of binding) at the concentration \( x_i (n_i = 6) \), and \( B_{i}(x_i) \) are the respective calculated data points. Also the squared deviations of these steady-state values were weighted by the variance \( \sigma_{r_{st}}^2 (x_i) \). Because in the time courses the number of points to be fitted is \( n_d \)-times bigger than those in the steady-state relationships, we employed the factor \( g = n_d/(n_d n_i) = 0.02 \) to give the time courses and the steady-state relationships the same weight in the fit.

To calculate the data points of the time courses for the fit, the differential equation

\[
dp(t, x)/dt = p(t, x)Q(x)
\]

was solved with the eigenvalue method. Here \( Q(x) \) is the \( Q \)-matrix depending on the concentration \( x \) and \( p(t, x) \) is the row-vector of the probabilities to be in one of the states of the model \( (43) \). It should be noted that for a given concentration profile equation, Eq. 4 has to be solved for each pulse separately.

The steady-state values \( p(\infty, x_i) \) for each concentration were calculated by setting Eq. 4 to zero under consideration that the sum of all components must be 1. On the other hand, the sum of the components, weighted with their fraction of ligands, is the binding degree, and can be written as

\[
B_{i}(x_i) = p(\infty, x_i)u_{Bi}^{T},
\]

where \( u_{Bi}^{T} \) is the column vector of the fraction of ligands for each model state. In a similar way, we get the normalized fluorescence values of the time courses,

\[
F_{c}(t_j, x_i) = \alpha_{i} p(t_j, x_i)u_{Bi}^{T},
\]

where \( \alpha_{i} \) is an additional factor to normalize the time course to the maximum value at the end of the activation period according to the experimental data \( F_{d}(t_j, x_i) \).

To keep computation time within a reasonable limit, only \( n_d = 60 \) data points of each time course were chosen (each 30 points for activation and deactivation, respectively). This number of data points was sufficient to fully describe the time courses on the one hand and allowed us to perform global fits in reasonable time on the other.

The \( \chi^2 \) value computed this way was minimized in the fit procedure. The reduced global \( \chi^2 \) value was calculated by dividing \( \chi^2 \) by the degrees of freedom of the fit

\[
\chi^2 = \frac{1}{2n_d - m} \chi^2,
\]

where \( m \) equals the number of parameters. If only normally distributed random errors contribute to the scattering of the data and if the model fits to the data, then the reduced \( \chi^2 \) value should be close to unity. For the best model found in this study, the reduced \( \chi^2 \) value was -2.8.

**RESULTS**

**Ligand binding and unbinding in closed HCN2 channels**

To gain insight into the action of the subunits in homotetrameric HCN2 channels, we analyzed the relative steady-state binding of the ligand as well as the kinetics of ligand binding and unbinding \( (F/F_{\text{max}}) \) by using the fluorescently labeled ligand fcAMP and confocal patch-clamp fluorometry \( (29,34) \). In channels preactivated by a hyperpolarizing pulse to \(-130 \text{ mV}, \) a jump from 0 to 7.5 \( \mu \text{M fcAMP}, \) a nearly saturating fcAMP concentration, further enhances activation \( (\text{Fig. 1 A}) \). The time course of fcAMP binding is characterized by an initial rapid phase that is followed by a slow phase, resembling the time course of current activation. At this concentration, we previously assumed that all four binding sites are occupied \( (29) \). At \(-30 \text{ mV}, \) a jump to 7.5 \( \mu \text{M fcAMP} \) generated less relative binding at the steady state than at \(-130 \text{ mV} \) and, as expected, a current was not activated at this subthreshold voltage \( (\text{Fig. 1 B}) \). Comparison of normalized time courses reveals that the characteristic slow phases of binding and unbinding at \(-130 \text{ mV} \) are missing at \(-30 \text{ mV} \) \( (\text{Fig. 1 C}), \) supporting earlier results, which show that this slow phase is kinetically related to channel activation \( (29) \).

To explore the processes underlying binding and unbinding at \(-30 \text{ mV} \) more systematically, we performed respective experiments at the five ligand concentrations 0.075, 0.25, 0.75, 2.5, and 7.5 \( \mu \text{M fcAMP} \) and averaged between five and seven traces at each concentration \( (\text{Fig. 2 A}) \). Fit of the steady-state concentration-binding relationship with a Hill function \( (\text{Eq. 1}) \) showed a rightward shift for nonactivated channels at \(-30 \text{ mV} (BC_{50} = 2.46 \text{ \mu M fcAMP}) \) compared to activated channels at \(-130 \text{ mV} (BC_{50} = 0.61 \text{ \mu M fcAMP}) \) \( (\text{Fig. 2 B}) \). Notably, at \(-30 \text{ mV} \) the Hill coefficient \( H \) of 1.28 still exceeded unity although it was slightly below the value of 1.56 determined at \(-130 \text{ mV} \). This suggests that also in nonactivated HCN2 channels there is cooperativity among the subunits but with lesser intensity than in voltage-activated channels. A Hill coefficient,
steady-state binding at the five ligand concentrations to a
respective C-C model to the data (Fig. 3; model 1 in Table 1).

This model proved to be inadequate as judged by eye and
the large value of 15.86 for the reduced \( \chi^2 \) (\( \chi^2 \)), which
ideally approximates unity. When adding a conformational
change (flip) to a flipped state (C*) following the ligand
binding (36), the global fit improved (Fig. 4; model 2 in
Table 1). Nevertheless, the fit remained inadequate for
part of the time courses and, in particular, for the
steady-state concentration-binding relationship at the higher
concentrations.

We therefore repeated this approach with a model
including two coupled binding steps with a flip from both
the mono- and the biliganded state (model 3 in Table 1).
Despite a doubling of the free parameters from 4 to 8, the
result was that \( \chi^2 \) did not improve; it even deteriorated.

Extending this analysis to models with three binding and
respective flip steps led to a notable improvement of \( \chi^2 \).
In total 14 such models with between 6 and 9 parameters were
tested, including various assumptions for equity or differ-
ence of binding and flip steps as well as for a flip of the
whole channel or of the individual subunits (Table S1 in the
Supporting Material). It turned out that 11 models
(models 3-1 to 3-11) with a flip of the whole channel pro-
duced lower \( \chi^2 \) values than the three tested models with a
flip of the individual subunits (models 3-12 to 3-14),

however, only provides a guess for the minimum number of
subunits involved in a cooperative process but does not pro-
vide any further mechanistic insight.

We next quantified the time courses of ligand binding and
unbinding at the five fcAMP concentrations by fits with singe
exponentials, yielding the time constants \( \tau_b \) and \( \tau_u \),
respectively (Eqs. 2a and 2b). When comparing \( \tau_b = 0.31 \text{ s} \) at
0.075 \( \mu \text{M} \) fcAMP with \( \tau_u = 0.19 \text{ s} \) at 0.75 \( \mu \text{M} \)
fcAMP, i.e., a 10-fold different fcAMP concentration, it in-
creases only by a factor of 1.6 (Fig. 2 C). This shows that the
ligand binding is not rate-limited by the binding process
alone but either by the unbinding process or subsequent
conformational changes or both. Fig. 2 C also shows that
the unbinding depends on the ligand concentration only a
little, suggesting that in contrast to channels preactivated
by voltage (29), nonactivated channels do not trap any li-
gands, at least up to the fcAMP concentration of 7.5 \( \mu \text{M} \).

The subunits do not operate independently

To gain further insight into the kinetics of the confor-
mational changes in nonactivated channels, we subjected the
averaged time courses of ligand binding and unbinding,
shown in Fig. 2, A and C, together with the values of
steady-state binding at the five ligand concentrations to a
global fit analysis with Markovian models. Assuming as
the most simple case that in channels not preactivated by
voltage all subunits bind and unbind a ligand independently
from each other with first-order kinetics, we fitted a respec-
tive C-C model to the data (Fig. 3; model 1 in Table 1).
although the latter models belonged to those with the highest number of free parameters. This result supports the idea of a concerted flip of the subunits. Model 4 in Table 1 is the best of the models in Table S2 with respect to the value of \( \chi^2 \). Notably, despite the fact that the number of free parameters was only six, and thus by two less than in model 3, \( \chi^2 \) was markedly smaller. This result suggests that the binding of more than two ligands is of particular relevance for eliciting cooperative effects between the subunits.

In the closed channel, four subunits also cooperate

Because HCN channels are tetramers, it was plausible to also test models with four sequential binding and respective flip steps. In total, 51 such models with between 6 and 11 parameters were tested, again including various assumptions for equity or difference of binding and flip steps as well as for a flip of the whole channel or of the individual subunits (Table S3). Also in the case of four binding steps, a big number of models with a flip of the whole channel produced lower \( \chi^2 \) values (models 4-1 to 4-17) than models with flipping individual subunits (models 4-18, 4-35, 4-46, 4-48). Among those, the models 4-35, 4-46, and 4-48 contained more free parameters than the best 17 models with a concerted flip. But the best way to demonstrate the superiority of models with a concerted flip over models with independently flipping subunits is to compare two models with the same assumptions for the cooperativity and number of free parameters but with a concerted flip on the one hand and independently flipping subunits on the other. This is demonstrated for our favorite model (model 5 in Table 1) and model 4-18 (Table S3). The favorite model 5 (4-1) with a concerted flip produced a significantly lower \( \chi^2 \) value than model 4-18. As an additional control, a model without flip (model 4-42) proved to be poor, although all four rate constants for binding and unbinding were free parameters. Together, these results show that there is a conformational flip and that it is caused by one concerted step of the four subunits but not by independently operating subunits.

A further result was that six models with four binding steps indeed produced still lower \( \chi^2 \) values than the best model with only three sequential binding steps, despite the fact that the number of free parameters was similar or even equal. The best model with respect to \( \chi^2 \) (model 5 in Table 1) produced a \( \chi^2 \) of only 2.81 compared to 3.25 obtained for the best model with three binding steps (model 4 in Table 1). The fit with model 5 is shown in Fig. 5, A–C. The parameters are provided by Table 2. Because the number of parameters was 6 in both fits, our data directly suggest that liganding of all four subunits is involved in the concerted conformational change of the channel. These results led us to the conclusion that there is cooperativity of the four subunits already in a closed HCN2 channel.

Effect of voltage on the microscopic affinity

Knowledge of the equilibrium association constants \( K_{A1}, K_{A4} \) for the closed nonactivated channel according to
model 5 (Table 2) allowed us to relate these constants to those determined for the channel preactivated by a voltage pulse to $-130$ mV as reported earlier (29) and, thus, to determine the effect of voltage-induced activation on the microscopic binding affinity of the four binding steps (Fig. 6).

While $K_{A1}$ and $K_{A4}$ are nearly unaffected by voltage, there is a strong and opposite effect of hyperpolarization-induced activation on $K_{A2}$ and $K_{A3}$: $K_{A2}$ is by more than an order of magnitude increased whereas $K_{A3}$ is by more than one order of magnitude decreased. Hence, the high energy barrier associated with the third binding step (29) is only inferred by voltage-induced channel activation.

**DISCUSSION**

In this study we measured steady-state and time-dependent ligand binding to functional but closed HCN2 channels and we present evidence that the ligand binding evokes a conformational flip in the channel. We demonstrate that four sequential binding reactions are involved. For the best model with four ligand binding steps and one flip from each ligated state (model 5 in Table 1), we quantified the microscopic affinities and related them to those in channels preactivated by hyperpolarizing voltage as described earlier (29). In addition to this, we determined rate constants for all reactions. Hence, our data provide kinetic information about the molecular mechanism proceeding in nonactivated HCN2 channels when fcAMP binds.

The most prominent effect of channel activation by voltage on the microscopic binding affinity was to increase $K_{A2}$ and to decrease $K_{A3}$. In other words, the binding of the second ligand is facilitated and that of the third ligand is impeded, generating with respect to the microscopic binding affinity the complex cooperativity pattern “positive-negative-positive” (29,44). In contrast, at $-30$ mV the respective cooperativity pattern is “negative-no-positive” (Fig. 6). Hence, there is an essential effect of channel activation on the affinity of the binding sites involved in the binding of the second and third ligand. These results further specify earlier results on HCN2 channels showing a reciprocal relationship between channel activation and the binding affinity (34). It should be noted that in the fit of our favorite model 5 (Table 1) we set the rate constants underlying $K_{A2}$ and $K_{A3}$ equal. Despite this simplifying and somewhat arbitrary...

**TABLE 2 Rate and equilibrium constants for model 5**

| Rate Constant | Dimension | Mean       | Err% |
|---------------|-----------|------------|------|
| $a$           | $M^{-1} \cdot s^{-1}$ | $2.00 \times 10^6$ | 15   |
| $b$           | $s^{-1}$ | $4.05 \times 10^9$ | 16   |
| $c$           | $s^{-1}$ | $1.18 \times 10^3$ | 20   |
| $d$           | $s^{-1}$ | $6.87 \times 10^{-1}$ | 18   |
| $e$           | $s^{-1}$ | $3.51 \times 10^{-1}$ | 22   |
| $f$           | $s^{-1}$ | $7.11 \times 10^{-2}$ | 42   |

| Equilibrium Constant | Dimension | Mean       | Err% |
|----------------------|-----------|------------|------|
| $K_{A1}$             | $M^{-1}$ | $4.94 \times 10^5$ | 8    |
| $K_{A2}$             | $M^{-1}$ | $1.69 \times 10^5$ | 9    |
| $K_{A3}$             | $M^{-1}$ | $1.69 \times 10^5$ | 9    |
| $K_{A4}$             | $M^{-1}$ | $2.91 \times 10^6$ | 18   |
| $E$                  |           | $2.03 \times 10^{-1}$ | 21   |

The rate constants $a$–$f$ are specified in Fig. 4A. Err% is the relative error (mean ± SE divided by mean in %). $K_{A1}$ through $K_{A4}$ denote the equilibrium association constants of the four binding steps. $E$ is the flip equilibrium constant given by $eff$. The errors of the equilibrium constants are calculated using the covariance matrix of the fitted parameters.
assumption, the effect of activation on these two equilibrium association constants (Fig. 6) is viewed to be robust because in channels preactivated by voltage, $K_{A2}$ and $K_{A3}$ differed enormously by three orders of magnitude and no fit could be obtained there when equating $K_{A2}$ and $K_{A3}$. Furthermore, the errors obtained from the covariance matrix under both conditions are reasonably small (Fig. 6) and thus confirm our interpretation.

Concerning the flip following the ligand binding described herein, our results suggest that it is a concerted flip of all subunits but not of the liganded subunits only. This suggests that a nonactivated channel acts as whole tetramer (24,45) already when the first ligand binds and that this action becomes promoted in proportion to the number of further bound ligands. It is presently not clear whether this action involves only the tetrameric CNBD or also parts of the transmembrane portion of the channel. The question may arise why we did not detect the flipped state in our previous analysis on channels preactivated by a hyperpolarizing voltage pulse to $\sim 130$ mV (29). We assume that under those conditions the reciprocal feedback from the activated channel pore to the flip of the nonactivating channel considered herein prevented the possibility to identify the flip.

How can the conformational flip be attributed to structural changes in the channels? Recent work in isolated CNBDs provided a guess of the initial conformational changes following the binding of cAMP to the binding site, in which the movement of the C-helix to occlude the cyclic nucleotide in the binding pocket plays a key role (46–48). Other structures have also been proposed to participate, including a folding of the P-helix within the PBC element, a translational movement of the B-helix, and a folding and movement of the F-helix (47). These changes presumably lead to an elimination of steric clashes evoked by the CNBD on the tetrameric C-linker that finally leads to the pore opening (48), at least in channel preactivated by voltage. It is plausible that all these arrangements are included in the conformational flip of nonactivated channels studied herein. However, considerable caution is needed when directly relating the conformational flip to these structural results because our functional data were obtained from whole channels whereas the structural data are from isolated CNBDs only. Hence, in the isolated CNBD the energetic feedback of the transmembrane channel domain on the CNBD (34) is completely missing. As outlined above, it is also possible that rearrangements in the transmembrane domain contribute to the conformational flip studied herein. Moreover, the role of the N-terminus is completely elusive in this context. Hence, the conformational flip studied herein provides information about a global conformational change of the nonactivated channel.

Another result of this study is remarkable: Despite the superiority of models with four binding steps and one concerted flip (model 5 in Table 1; Fig. 5), a model with independently operating subunits, i.e., one binding step and one flip (model 2 in Table 1; Fig. 4), already produced a significantly better description than the C-C model (model 1 in Table 1; Fig. 3). This result supports the notion that, in a nonactivated channel, the flip evoked by the ligand binding to a subunit predominantly affects the binding affinity of this particular subunit (49) and that the interaction between the subunits is of lesser effect.

If relating the results of our modeling to established kinetic models, the concerted (allosteric) conformational flip in the type of models favored herein matches one of the basic assumptions of the MWC-model (31) but does not support the necessity to use more complex models of the Koshland-Nemethy-Filmer type (50). Nevertheless, the model favored herein differs from the MWC model by the nonmonotonous change of the affinity of the four binding reactions.

We finally like to emphasize that the measurements presented in this report are based on measuring the ligand binding and unbinding, and that these data were interpreted by hidden Markov models to substantiate a kinetic scheme describing the binding and gating of the closed channel. This approach is basically opposite to common approaches in electrophysiology, studying channel activation and substantiating hidden Markov models describing ligand binding and the activation gating (51–54). A major aspect of our approach is that it reports other properties of channel gating than those associated with activation. Hence, this approach is applicable to other nonactivated ligand gated channels as well and, most notable, to any other membrane receptors, as, e.g., the huge number of G-protein coupled receptors (55). Moreover, if a readout of conformational changes is included in the experiments, e.g., by evaluating the kinetics of Förster resonance energy transfer (56), the constraints in the fit will grow significantly, thereby enabling kinetic analyses with much greater detail than herein.

SUPPORTING MATERIAL

Three tables are available at http://www.biophysj.org/biophysj/supplemental/S0006-3495(15)01008-5.

AUTHOR CONTRIBUTIONS

S.T. performed the experiments, analyzed the data, and designed the figures; E.S. performed the global fits; J.K. performed experiments and contributed to the data analysis; and K.B. designed the study and wrote the article.

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