Counting ion channels on cell membranes is of fundamental importance for the study of channel biophysics. Channel counting has thus far been tackled by classical approaches, such as radioactive labeling of ion channels with blockers, gating current measurements, and nonstationary noise analysis. Here, we develop a counting method based on patch-clamp fluorometry (PCF), which enables simultaneous electrical and optical recordings, and apply it to EGFP-tagged, hyperpolarization-activated and cyclic nucleotide–regulated (HCN) channels. We use a well-characterized and homologous cyclic nucleotide–gated (CNG) channel to establish the relationship between macroscopic fluorescence intensity and the total number of channels. Subsequently, based on our estimate of the total number of HCN channels, we determine the single-channel conductance of HCN1 and HCN2 to be 0.46 and 1.71 pS, respectively. Such a small conductance would present a technical challenge for traditional electrophysiology. This PCF-based technique provides an alternative method for counting particles on cell membranes, which could be applied to biophysical studies of other membrane proteins.

INTRODUCTION

The number of particles on cell membranes is a fundamental quantity required for biophysical studies of membrane proteins, especially ion channels and transporters (Hille, 2001). This information can be used to derive important biophysical properties such as single molecule conductance, open probability, ionic selectivity, and stoichiometry of ligand binding or subunit assembly. Long before the era of single molecule biophysics, elegant strategies were developed to count ion channels on cell membranes, including the use of radiolabeled neurotoxins to specifically label certain ion channels (saxitoxin and tetrodotoxin for Na channels and α-bungarotoxin for acetylcholine receptors; Moore et al., 1967; Salpeter and Loring, 1985), the measurement of macroscopic gating currents for voltage-gated channels divided by the number of gating charges per channel (Armstrong and Bezanilla, 1974, 1977), and the application of fluctuation analysis to ensembles of macroscopic current recordings (nonstationary noise analysis [NSNA]; DeFelice, 1981; Sigworth, 1984; Alvarez et al., 2002). These classical techniques—dating back to the 60s—elegantly bridged macroscopic observations with molecular properties that had been difficult to approach, such as the density of channels on cell membranes and single-channel conductance. Later on, the single-channel recording technique made it possible to directly detect transitions between open and closed channels and to derive the total number of channels from single-channel conductance and open probability measurements (Neher and Sakmann, 1976; Sigworth and Neher, 1980). However, single-channel recording is not suitable for ion channels with an extremely small conductance or flickering openings, nor for most transporters. Further, a condition of NSNA is that the membrane patch must sustain repetitive stimulations up to 100 times. Moreover, particular gating properties, such as cooperative opening or closing, could complicate the interpretation of noise analysis. Therefore, alternative strategies that help delineate macroscopic measurements and shed light on molecular properties are still useful.

In the potassium channel superfamily, cyclic nucleotide–gated (CNG) and hyperpolarization-activated and cyclic nucleotide–regulated (HCN) channels share similar architecture. Each subunit within the tetrameric assembly contains six transmembrane α-helices and a cyclic nucleotide–binding domain in the C terminus (Jan and Jan, 1990; Zagotta and Siegelbaum, 1996; Kaupp and Seifert, 2002; Biel et al., 2009). Intracellular cyclic nucleotides, including cAMP and cGMP, directly bind to and activate CNG and HCN channels. Single-channel recordings of CNG channels, first reported in the 80s, provided important mechanistic insights into their biophysical and physiological properties, such as...
the density of CNG channels on photoreceptor cell membranes (Matthews and Watanabe, 1988; Goulding et al., 1992, 1994; Ruiz and Karpen, 1997; Li and Lester, 1999). However, because of the extremely small single-channel conductance of HCN channels, studies to directly address single-channel properties have been rare (DiFrancesco, 1986; Biel et al., 2009). Among the four vertebrate HCN subtypes, HCN1 to HCN4, only HCN2 has been subjected to single-channel electrophysiology, and the conductance was determined to be <2 pS (Dekker and Yellen, 2006; Thon et al., 2013). The NSNA approach has also been applied to native and heterologously expressed HCN channels, yielding useful information (Table S1; Kole et al., 2006; Flynn et al., 2007; Barrow and Wu, 2009).

Here, we develop a new method to measure the number of channels on cell membranes, based on the patch-clamp fluorometry (PCF) technique. PCF combines simultaneous electrical recording of ionic currents and fluorescence intensity from a membrane patch held within a glass recording pipette (Zheng and Zagotta, 2003; Kusch and Zifarelli, 2014). It has been an effective tool in the study of channel biophysics, with topics ranging from the calmodulin-dependent regulation of CNG channels to ligand-dependent gating mechanisms in CNG and HCN channels (Zheng and Zagotta, 2000; Kusch et al., 2010; Wu et al., 2011, 2012). We apply this method to estimate the number of HCN1, HCN2, and spHCN channels on membrane patches and then study their single-channel conductance and ionic selectivity. Our results not only provide new information on basic biophysical properties of HCN channels but also highlight the application of PCF as an alternative channel counting method for the study of membrane proteins with ionic conductance.

MATERIALS AND METHODS

Construction of EGFP-tagged CNG and HCN fusion proteins

For the chimeric CNG channel (ROONS; provided by S. Siegelbaum, Columbia University, New York, NY), the DNA sequence encoding EGFP was added to the C terminus through the cut sites of SalI and HindIII (New England Biolabs, Inc.). Five amino acids (VDAGA) were added between CNG (...STQD) and EGFP (MVSK...). Another five amino acids (SGLRS) were added between EGFP (...DELYK) and the stop codon. For the spHCN channel (provided by B. Kaupp and R. Seifert, Abteilung Molekulare Neurosensorik, Center of Advanced European Studies and Research, Bonn, Germany), the EGFP was inserted between I690 and P691. An alanine and a glycine residue were added to the N and C termini of EGFP, respectively. For the mouse HCN1 channel, the EGFP was inserted between E609 and I610, with an alanine and a glycine added to the N and C termini of EGFP, respectively. For the mouse HCN2 channel, the EGFP was inserted after A602 through the BsmI cut site. An extra BsmI cut site was engineered to the C terminus of EGFP, with three extra amino acids (ENA) added after EGFP.

Functional expression in Xenopus laevis oocytes and electrophysiological characterization

cDNAs encoding HCN and CNG channels related to this study were cloned into the pGHI9 vector and linearized by NheI (CNG and spHCN) and SphI (HCN). mMESSAGE machine (Ambion) was used for cRNA synthesis. 40–50 ng cRNA was injected into each oocyte at stage VI. After 1–2 d of incubation at 18°C, injected oocytes were selected for recording. For patch-clamp recording, the electrode solution (extracellular) and bath solution (intracellular) were symmetrical and contained 110 mM KCl, 2 mM NaCl, 10 mM HEPES, and 1 mM EDTA (pH 7.4 adjusted by KOH). All experiments were performed at room temperature.

PCF

The PCF setup was constructed based on a BX50WI microscope (Olympus) equipped with a 60× water immersion lens (LUMPlanFL, N.A. 1.0). A 473-nm diode-pumped solid-state (DPSS) laser (Ultralasers Inc.) was used as the excitation light source. The following filter set was used for collecting the EGFP fluorescence signal: exciter, D480/30; dichroic mirror, DC505LP; emitter, D510LP (Chroma Technology Corp.). Optical signals were detected by a 16-bit EMCCD camera (Cascade 1K; Photometrics). An 18-bit data acquisition board (PCI-6289; National Instruments) was used for analogue and digital I/O. WinWCP was used for data acquisition. The laser light source, the CCD camera exposure, and the amplifier for patch-clamp recording were synchronized by TTL signals.

Macroscopic current traces were collected in the presence of a saturating concentration of cAMP and with a hyperpolarizing voltage step that, based on the Boltzmann equation, was enough to produce maximal channel opening. Fluorescence signals were collected with four increasing exposure times: 25, 50, 100, and 200 ms. No obvious bleach of the EGFP molecules was detected (Fig. S8). The optical signals were collected within the linear range of the CCD camera. The ImageJ program (National Institutes of Health) was used to analyze the fluorescence images (Schneider et al., 2012). To specifically measure the fluorescence signal near the excised membrane, a region of interest (ROI) was selected around the arc of membrane patch. ΔF was defined as mean fluorescence intensity in the ROI with background fluorescence subtracted. The background fluorescence was collected by moving the tip of the recording pipette out of the view field. The following linear equation was used to fit the F-I relationship:
where $A$ and $B$ represent the intercept and slope, respectively. OriginPro 8.1 was used for curve fitting. A linear equation was used to fit the F-I results. The Pearson correlation coefficient, of which the value of 1 indicates perfect positive correlation and 0 indicates no correlation, is listed in figure legends. The confidence interval bands (95%) are plotted in green.

$I$ is defined by the following equation:

$$I = N \cdot P_0 \cdot i = N \cdot P_0 \cdot V \cdot \gamma,$$  \hspace{1cm} (2)

where $N$ is the total number of channels, $P_0$ is the open probability, $i$ represents single-channel current, $V$ is the membrane voltage minus the reversal potential, and $\gamma$ represents single-channel conductance. Thus, the F-N relationship can be expanded to

$$F = A + B \cdot I,$$  \hspace{1cm} (1)

Because CNG and HCN channels share similar topology, we assume the slope of the F-N relationship is independent of the channel type. Thus, the conductance of HCN channel could be derived based on the following equation:

$$B_{CNG} \cdot P_{CNG} \cdot V_{CNG} \cdot \gamma_{CNG} = B_{HCN} \cdot P_{HCN} \cdot V_{HCN} \cdot \gamma_{HCN},$$  \hspace{1cm} (4)

where $i$ represents the single-channel current, $N$ represents the total number of channels, and $k$ is the background offset (Kole et al., 2006; Barrow and Wu, 2009). The polynomial fit function with two orders from OriginPro 8.1 was used for curve fitting. Adjusted $R^2$ values are provided in each figure legend. For HCN channels that show significant current rundown, using the difference between neighboring traces is essential to obtain reasonably good results. Separately, we calculated the current variance based on the difference from the averaged current trace but obtained extremely high values of variance and unrealistic single-channel conductance (Fig. S7).

Online supplemental material
A document supplementary to the main manuscript provides the information on previous published results for HCN channel conductance (Table S1), calibration of the optical system using fluorescent plates (Fig. S1), voltage-dependent channel activation curves for mHCN2 and spHCN channels (Figs. S2 and S3), NSNA results for mHCN2, spHCN-EGFP, and spHCN channels (Figs. S4, S5, and S6), and a test of a traditional method to calculate the current variance in NSNA (Fig. S7). Fig. S8 shows the selection of ROI and photobleaching of EGFP molecules. Online supplemental material is available at http://www.jgp.org/cgi/content/full/jgp.201511559/DC1.

RESULTS
Establishing the relationship between macroscopic fluorescence intensity $F$ and the total number of channels $N$ on a membrane patch
We began by establishing the relationship between the number of channels and the macroscopic fluorescence intensity from a well-characterized chimeric CNG channel (ROONS) for which the total number of channels can be determined purely by electrophysiology recordings at single-channel and macroscopic levels. ROONS was constructed based on the bovine CNGA1 channel, of which the N terminus and the pore domain of CNGA1 were replaced by the corresponding parts from catfish CNGA2 to improve the gating efficacy (Goulding et al., 1994; Tibbs et al., 1997). We first started from single-channel recordings of ROONS-EGFP, with a subsaturating concentration of cGMP (3.5 µM) applied to the intracellular side (Fig. 1A, top). Then we applied a high concentration of cGMP (35 µM) and determined the open probability to be 94.9% (Fig. 1A, top). Histogram analysis revealed that the single-channel conductance of

$$\sigma^2(t) = \frac{2}{M-1} \sum_{m=1}^{M} (y_m(t) - Y(t))^2.$$  \hspace{1cm} (7)

Step 4. Fit the data points of variance — macroscopic current amplitude by the following parabolic function:

$$\sigma^2(I) = i \cdot I - \frac{I^2}{N} + k,$$  \hspace{1cm} (8)

where $i$ represents the single-channel current, $N$ represents the total number of channels, and $k$ is the background offset (Kole et al., 2006; Barrow and Wu, 2009). The polynomial fit function with two orders from OriginPro 8.1 was used for curve fitting. Adjusted $R^2$ values are provided in each figure legend. For HCN channels that show significant current rundown, using the difference between neighboring traces is essential to obtain reasonably good results. Separately, we calculated the current variance based on the difference from the averaged current trace but obtained extremely high values of variance and unrealistic single-channel conductance (Fig. S7).
ROONS-EGFP is 53.6 ± 7.1 pS (n = 7), close to the original characterization of this channel (Fig. 1B; Goulding et al., 1993). Thus, based on the macroscopic current recorded under 35 µM cGMP, we could directly calculate the total number of ROONS-GFP channels on the membrane patch by dividing the macroscopic current amplitude by the open probability and the single-channel current (Fig. 1, C [left] and E [top and bottom x axes]; N = I/[i × Po], where I is the macroscopic current, i is the single-channel current, and Po is the open probability). Thus, electrophysiology recordings can provide a reliable estimation of the number of ROONS channels (N) on the cell membrane, which will be used to calibrate the corresponding fluorescence intensity (F).

To establish the F-N relationship, we applied PCF to collect the macroscopic current trace and the corresponding fluorescence image for each membrane patch (Fig. 1C, right). To ensure the optical recording system functions within the linear range, we sequentially collected four fluorescence images by increasing the exposure durations. The Cascade 1K is a 16-bit camera so that the pixel intensity ranges from 0 to 65,535. (right) Total fluorescence intensity from the ROI surrounding the patch of membrane patch (green) and the background (blue), which was separately recorded after moving the recording pipette out of the view. (E) Cross-plots of fluorescence intensity versus current amplitude. Two x axes show the macroscopic current (bottom) and the corresponding number of channels (top). The amplitude of single-channel current, 4.29 pA, was used in the conversion. Linear fit statistics: Pearson correlation coefficient, 0.823. Green curves show the confidence bands (95%).
mHCN2 channels were chosen based on the corresponding Boltzmann fit of the channel activation curve (Figs. S2 and S3). If we assume the open probabilities for these channels to be 100%, our analysis revealed that the single-channel conductance for spHCN-EGFP, mHCN1-EGFP, and mHCN2-EGFP channel are 0.88, 0.46, and 1.46 pS, respectively. These values represent a lower estimation because applying NSNA to these channels suggests the maximal open probabilities are <100%.

Applying NSNA to macroscopic currents to estimate the single-channel conductance and the total number of channels

To corroborate the aforementioned results based on PCF, we switched to NSNA, a classical approach for estimating the single-channel conductance and the total number of channels that is based on fluctuation analysis of macroscopic currents. Because of the technical difficulty in collecting an ensemble (ideally 100) of stable HCN current traces with extremely negative potentials, to our best knowledge, only four publications in the literature had applied NSNA to HCN channels: two for heterologously expressed WT spHCN or mHCN2 channels and two for native neuronal HCN currents (Johnson and Zagotta, 2005; Kole et al., 2006; Flynn et al., 2007; Barrow and Wu, 2009). Here we started from mHCN2-EGFP and mHCN2 channels and then extended the study to spHCN-EGFP and spHCN channels. We did not apply NSNA to mHCN1 channel because of the technical difficulty in obtaining stable current traces. For channel activation, we chose the hyperpolarizing voltage step of −130 mV (mHCN2 channels) or −90 mV (spHCN channels) in the presence of saturating concentrations of cAMP (3 µM for mHCN1-EGFP and mHCN2-EGFP; 30 µM for spHCN-EGFP; Fig. 2 A, left). The corresponding fluorescence images were recorded using the same protocol as for ROONS-EGFP (Fig. 2 A, right). For the patches with a high density of channels, histogram analysis of pixel intensities did reveal saturated pixels (exposure duration: 100 and 200 ms; Fig. 2 B). In those cases, images collected with shorter exposure durations were used.

After collecting electrical and optical recordings from membrane patches of a diverse range of channel densities, we plotted the current amplitudes versus fluorescence intensity and fitted the results with a linear relationship (Fig. 2 C; mHCN2-EGFP). For mHCN1 and spHCN channels, the macroscopic current trace and the corresponding images are shown in Fig. 3 (A and B, respectively). Linear fits of the F-I plots are shown in Fig. 3 (C and D), which we used in the final analysis (see Materials and methods; Eq. 4). To ensure channel opening reach steady-state, the hyperpolarizing voltage steps for activating spHCN, mHCN1, and mHCN2 channels were chosen based on the corresponding Boltzmann fit of the channel activation curve (Figs. S2 and S3). If we assume the open probabilities for these channels to be 100%, our analysis revealed that the single-channel conductance for spHCN-EGFP, mHCN1-EGFP, and mHCN2-EGFP channel are 0.88, 0.46, and 1.46 pS, respectively. These values represent a lower estimation because applying NSNA to these channels suggests the maximal open probabilities are <100%.

Applying the relationship between fluorescence intensity and channel number to count HCN channels

Because CNG and HCN channels are homologous and share similar folding and assembly, the relationship between fluorescence intensity and channel number, which was established based on ROONS-EGFP data, can be applied to count HCN channels. We constructed chimeric spHCN, mHCN1, and mHCN2 channels by inserting EGFP to the C-terminal end of the cyclic nucleotide-binding domain, comparable with the construction of ROONS-EGFP. Macroscopic currents were recorded in the presence of saturating concentrations of cAMP (3 µM for mHCN1-EGFP and mHCN2-EGFP; 30 µM for spHCN-EGFP; Fig. 2 A, left). The corresponding fluorescence images were recorded using the same protocol as for ROONS-EGFP (Fig. 2 A, right). For the patches with a high density of channels, histogram analysis of pixel intensities did reveal saturated pixels (exposure duration: 100 and 200 ms; Fig. 2 B). In those cases, images collected with shorter exposure durations were used.

After collecting electrical and optical recordings from membrane patches of a diverse range of channel densities, we plotted the current amplitudes versus fluorescence intensity and fitted the results with a linear relationship (Fig. 2 C; mHCN2-EGFP). For mHCN1 and spHCN channels, the macroscopic current trace and the corresponding images are shown in Fig. 3 (A and B, respectively). Linear fits of the F-I plots are shown in Fig. 3 (C and D), which we used in the final analysis (see Materials and methods; Eq. 4). To ensure
0.21 pS; n = 3; activation) channels are larger than the corresponding values based on PCF. We also adjusted the PCF results by taking into account the maximal open probability obtained by NSNA and obtained the value of 1.12 and 1.70 pS for spHCN-EGFP and mHCN2-EGFP, respectively (Table 1).

Determining the ion selectivity based on the slope of fluorescence-current relationship
Next we expanded the aforementioned PCF-based method to measure the ion selectivity of the mHCN2 channel. We focused on three charge carriers: K⁺, Na⁺, and NH₄⁺. Determining the relative permeability of different ions is a classical topic that has been addressed mostly by single-channel recordings, if applicable, or more popularly the measurement of reversal potential based on the Goldman–Hodgkin–Katz (GHK) equation (Goldman, 1943; Hodgkin and Katz, 1949). The GHK/reversal potential method is reliable in most cases; however, for ammonium (NH₄⁺), it is complicated because the NH₄ gradient across the membrane would lead to significant changes in local pH, which in turn affects...
the function of many channels including the HCN channel (Boron, 2004; Musa-Aziz et al., 2009).

Because the macroscopic fluorescence intensity of the membrane patch reflects the total number of channels on the membrane, the slope of the F-I relationship should directly reflect the permeability of the channel to that particular charge carrier. In turn, the ratio of the slope of F-I curves can be used to determine the relative selective permeability among different charge carriers. For each charge carrier, we used symmetrical solutions on both sides of the membrane. As in the aforementioned experiments, we collected the macroscopic current trace together with the fluorescence image from membrane patches that express a diverse range of number of channels (Fig. 5, A and B). We started from P_K/P_Na and used it as the positive control because the P_K/P_Na has been well documented to be around 3:1 for HCN channels (in the presence of at least 5 mM K+; Biel et al., 2009). Also, it is known that in the absence of K+ ions, the permeability of Na+ ions through the HCN channel is very low (Frase et al., 1992; Ludwig et al., 1998; Biel et al., 2009). Indeed, we calculated the ratio of the slopes

Figure 3. Cross-plots of fluorescence intensity versus current amplitude for mHCN1-EGFP and spHCN-EGFP channels. (A, left) Macroscopic current of the mHCN1-EGFP channel recorded with 3 µM cAMP in the bath. A voltage step from 0 to −120 mV was used for channel activation. Tail current was recorded at −40 mV. (B, left) Macroscopic current of the spHCN-EGFP channel recorded with 30 µM cAMP in the bath. A voltage step from 0 to −120 mV was used for channel activation. Tail current was recorded at 40 mV. (A and B, right) The corresponding brightfield and fluorescence images. (C) Cross-plots of fluorescence intensity versus current amplitude for the mHCN1-EGFP channel. Linear fit statistics: Pearson correlation coefficient, 0.973. (D) Cross-plots of fluorescence intensity versus current amplitude for the spHCN-EGFP channel. Linear fit statistics: Pearson correlation coefficient, 0.859. Green curves show the confidence bands (95%).
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(0.0230, 110/5 mM Na⁺/K⁺ versus 0.0067, 110/2 mM K⁺/Na⁺) and obtained the value of 3.43, which is in agreement with the previous results in the presence of K⁺ (Biel et al., 2009). However, in the absence of K⁺, the ratio of Pᵦ/Pₙa was as high as 101, indicating a high selective permeability for K⁺ and very low permeability for Na⁺ for HCN channels (Fig. 5, C and D). Finally, we tested the relative permeability for NH₄⁺. Based on the

Table 1. Summary of the results on HCN channel conductance

<table>
<thead>
<tr>
<th>HCN channels</th>
<th>Experimental method</th>
<th>Results (pS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>mHCN1-EGFP</td>
<td>PCF</td>
<td>0.46 ± 0.07</td>
</tr>
<tr>
<td>mHCN2-EGFP</td>
<td>PCF</td>
<td>1.46 ± 0.26</td>
</tr>
<tr>
<td>spHCN-EGFP</td>
<td>PCF</td>
<td>0.88 ± 0.22</td>
</tr>
<tr>
<td>mHCN2-EGFP</td>
<td>NSNA</td>
<td>2.70 ± 0.21 (n = 3) 2.29 ± 0.61 (n = 3)</td>
</tr>
<tr>
<td>mHCN2</td>
<td>NSNA</td>
<td>2.37 ± 0.24 (n = 4) 2.27 ± 0.28 (n = 4)</td>
</tr>
<tr>
<td>spHCN-EGFP</td>
<td>NSNA</td>
<td>1.03 ± 0.11 (n = 3) 0.99 ± 0.27 (n = 3)</td>
</tr>
<tr>
<td>spHCN</td>
<td>NSNA</td>
<td>0.76 ± 0.16 (n = 4) 0.49 ± 0.10 (n = 4)</td>
</tr>
</tbody>
</table>

SD of the PCF results was calculated using the following equation based on Eq. 4:

$$Δγ_{HCN} = γ_{HCN} \left( \frac{ΔB_{CNG}}{B_{CNG}} \right)^2 + \left( \frac{ΔP_{CNG}}{P_{CNG}} \right)^2 + \left( \frac{ΔY_{CNG}}{Y_{CNG}} \right)^2 + \left( \frac{ΔB_{HCN}}{B_{HCN}} \right)^2 + \left( \frac{ΔP_{HCN}}{P_{HCN}} \right)^2$$

where $ΔB$ and $ΔP$ represent the SD of the F-I slope and the open probability, respectively. Because the maximal open probability and the single-channel conductance of the CNG channel were directly derived from electrophysiology recordings, their contributions to the error should be minimal compared with the contributions by the slopes of the F-I plots and by the maximal open probability of HCN channels based on NSNA.
ratio of slopes, the $P_{K^+}/P_{NH_4^+}$ was determined to be 13.1 (without $K^+$) or 12.1 (with 5 mM $K^+$; Fig. 5, E and F).

**DISCUSSION**

Counting the number of particles on cell membranes is a classic topic for the study of not only ion channels but also transporters and other types of membrane proteins. It is not a trivial task and historically has been addressed by carefully designed experiments including radioactive labeling of channels/receptors (Moore et al., 1967), gating current and gating charge measurements (Armstrong and Bezanilla, 1974, 1977), and noise analysis like NSNA (DeFelice, 1981; Sigworth, 1984).
Here, based on the PCF technique, we developed a relatively straightforward method to count the number of channels by correlating the macroscopic current amplitude with the fluorescence intensity recorded from membrane patches. We used a well-characterized CNG channel to calibrate the F-N relationship and used this method to determine the single-channel conductance and relative permeability of HCN channels. Our results are consistent with previous studies using electrophysiology methods, which confirms the effectiveness of this PCF-based method (Dekker and Yellen, 2006; Kole et al., 2006; Flynn et al., 2007; Barrow and Wu, 2009; Bendord et al., 2012). Moreover, we obtained new information such as the single-channel conductance of mHCN1 channel, which is about threefold smaller than that of mHCN2. Such a small value (0.46 pS) would present a great challenge for traditional electrophysiology-based methods. Finally, we further expanded this approach and confirmed it as a straightforward and effective method for determining the relative permeability for different charge carriers.

The single-channel conductance of the mHCN2 channel has been tackled by two previous studies using high-quality single-channel recordings, which reported the value of 1.46 and 1.67 pS, respectively (Dekker and Yellen, 2006; Thon et al., 2013). Here our PCF-based method yielded a value of 1.71 pS for mHCN2 channel (after taking into account of the open probability determined by NSNA), which is close to the values determined by single-channel recording and thus confirmed the effectiveness of this method. Noticeably, if maximal open probability is assumed to be 100%, as for mHCN1 channel, the value of single-channel conductance provided by the PCF method represents a lower estimate. Another possibility of error is the small portion of channels that reach the cell membrane but are not fluorescent or do not function, mostly because of the misfolded or immature GFP or channel molecules (Ullrich and Isacoff, 2007). These optically or functionally “silent” channel undetected in PCF might explain the observation that some F-I fittings did not pass through the zero point.

Although two previous publications provide similar values for the single-channel conductance (1.46 or 1.67 pS), they were distinct in their interpretations of gating properties especially the cooperativity among different channels. In the study by Dekker and Yellen (2006), the mHCN2 channel was expressed in HEK293 cells, and a prominent cooperativity was observed during channel opening. The study by Thon et al. (2013) used the Xenopus oocyte expression system but failed to detect any cooperativity. Interestingly, our noise analysis results showed that the single-channel conductance of the same channel to be around 2.3 pS, very close to a previous publication using the similar method (Johnson and Zagotta, 2005) and significantly higher (at least 35%) than the values yielded by single-channel recording or PCF. Therefore, it is possible that mHCN2 shows some cooperativity under certain conditions, such as being expressed in mammalian cells or at high expression level (up to 20,000 per macropatch in our experiments).

Surprisingly, we discovered that the mHCN1 channel, with only a few residues different in the pore region from mHCN2, has a very small single-channel conductance, only about a third of that of mHCN2 if we assume the maximal open probabilities are comparable between the two channels. Physiologically, this extremely small conductance could play important roles in fine-tuning the resting membrane potential and membrane resistance, especially in the spatially restricted region like the neuronal dendrites, the presynaptic terminal, or the axonal initial segment (Kole et al., 2006; Huang and Trussell, 2014; Harnett et al., 2015; Ko et al., 2016).

If applicable, NSNA is still a robust method and can give results comparable with single-channel recording or PCF. Membrane patches expressing mHCN2 and spHCN channels but not the mHCN1 channel can sustain repetitive stimulations up to 100 times. Our NSNA results showed that for mHCN2, the single-channel conductance is $2.37 \pm 0.24$ pS ($n = 4$; activation), which is in close range but significantly larger than the value by other methods ($\sim 1.6$ pS by single-channel recordings), possibly because of cooperative openings at single-channel level. Still, NSNA should be considered as a robust method for estimating the number of channels and the single-channel conductance because in many cases collecting an ensemble of macroscopic current traces is relatively straightforward compared with recording extremely small single-channel currents or simultaneously recording the fluorescence signals from membrane patches. The good match between the NSNA-based determination of the distribution profile of HCN channels along the pyramidal cell dendrites with the immunostaining results showcases the strength of this traditional fluctuation-based counting method (Kole et al., 2006; Barrow and Wu, 2009).

For determining the selective permeability among different charge carriers, the PCF-based approach is even simpler and does not require the manipulation of ionic compositions. The ratio of the fluorescence-current relationship is directly correlated with the selective permeability. Our results of $PK/PNa$ are highly consistent with the electrophysiology results: in the absence of $K^+$, the HCN channel has a very small conductance for $Na^+$; however, in the presence of a small amount of $K^+$, the permeability for $Na^+$ could reach $\sim 30\%$ of that for $K^+$. Moreover, we extended this approach to $PK/PNH_4^+$ and obtained a value around 12–13, which is similar to an earlier study of native HCN current (14.3) but very different from a recent study of heterologously expressed HCN2 channel (1.95; Woll-
It is possible that in the later study applying high concentrations of NH₄⁺ to the extracellular solution leads to acidification of intracellular environment, which in turn affects the function of HCN channels.

Currently, our approach relies on the measurement of macroscopic fluorescence intensity and a separate calibration curve based on an EGFP-tagged homologous channel of which the total number can be purely derived based on electrophysiology recordings. Because we used the same settings during the optical recording, factors including efficiencies of excitation and collection of emitted fluorescent light can be cancelled out from the calculation. Moreover, the shape of the membrane patch and its orientation against the lens focal plane would affect the fluorescence reading, but both factors should equally affect EGFP-tagged CNG and HCN channels so that they could also be cancelled out. In practice, we noticed that it was difficult to consistently adjust the focal plane to the horizontal equator of the membrane dome, which should be a source of variation among patches of membrane but could be overcome by further increasing the number of data points. Finally, this PCF-based counting method could be improved by incorporating measurements of single fluorescent particles and thus bypassing the requirement of a well-characterized homologous channel for calibration purpose, which requires more careful experiment design and result interpretation but ought to broaden the application of this approach to other membrane proteins.

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REFERENCES


Figure S1. **Calibration of the optical detection system.** An autofluorescent plastic slide (#92001 by Chroma Tech. Corp.) was used to calibrate the optical detection system on a daily basis. (A) Fluorescence images were collected using the same procedure as the images of membrane patch, with the same 60× water immersion lens and settings for the EMCCD camera. (B) Fluorescence intensities within the central region where membrane patches were positioned. Each of the nine different positions was measured by an area as indicated by the dashed box in A. (C) For each day, six images of the fluorescent plate were collected at different positions before and after the experiments. The mean value of those images was used to calibrate the experimental results (fluorescence intensities of membrane patches) of that day. (D) Certain variations were observed across different days.
Figure S2. Voltage-dependent channel activation curves for mHCN2 and mHCN2/EGFP channels. (A) For mHCN2 channel, a series of hyperpolarizing voltage steps from −90 mV (without cAMP, top) or −70 mV (with cAMP, middle), at the interval of −10 mV, were used for channel activation. Tail currents were measured at −40 mV. Tail current amplitudes were normalized by the maximal tail current amplitude measured in the absence of cAMP. The averaged V_{1/2} values are −112 ± 4 mV (n = 7) with cAMP and −128 ± 7 mV (n = 7) without cAMP. (B) For mHCN2/EGFP, the averaged V_{1/2} values are −115 ± 8 mV (n = 5) with cAMP and −129 ± 9 mV (n = 9) without cAMP. (C) Normalized channel activation curves of mHCN2 channels. (D) Normalized channel activation curves of mHCN2-EGFP channels.
Figure S3. **Voltage-dependent channel activation curves for spHCN and spHCN-EGFP channels.** (A) For spHCN channel, a series of hyperpolarizing voltage steps from −20 mV (without and with cAMP), at the interval of −10 mV, were used for channel activation. Tail currents were measured at 40 mV. Tail current amplitudes were normalized by the maximal tail current amplitude measured in the absence of cAMP. The averaged V1/2 values are −67 ± 5 mV (n = 3) with cAMP and −67 ± 3 mV (n = 3) without cAMP. (B) For spHCN/EGFP, the averaged V1/2 values are −62 ± 7 mV (n = 3) with cAMP and −63 ± 7 mV (n = 3) without cAMP. (C) Normalized channel activation curves of mHCN2 channels. (D) Normalized channel activation curves of mHCN2-EGFP channels.
Figure S4. **NSNA of an ensemble of macroscopic mHCN2 currents.** (A, top) Voltage protocol used for channel activation and deactivation. (bottom) 6 representative traces from 100 repeatedly collected traces. (B) Current variance over the complete time course of a single episode. (C) Current variance versus mean current amplitude. Red, parabola fit of the macroscopic current part. Green, parabola fit of the tail current part. (D) Normal residual after curve fit for macroscopic current (corresponding to the red trace in C). Results: $i, -0.245$ pA ($-130$ mV); $\gamma, 1.89$ pS; $N, 7,319$; $Po, 82.5\%$. Adjusted $R^2, 0.891$. (E) Normal residual after curve fit for tail current (corresponding to the green trace in C). Results: $i, -0.0593$ pA ($-40$ mV); $\gamma, 1.48$ pS; $N, 7,707$; $Po, 82.4\%$. Adjusted $R^2, 0.937$. 
Figure S5. **NSNA of an ensemble of macroscopic spHCN-EGFP currents.** (A, top) Voltage protocol used for channel activation and deactivation. (bottom) 6 representative traces from 100 repeatedly collected traces. (B) Current variance over the complete time course of a single episode. (C) Current variance versus mean current amplitude. Red, parabola fit of the macroscopic current part. Green, parabola fit of the tail current part. (D) Normal residual after curve fit for macroscopic current (corresponding to the red trace in C). Results: $i$, $-0.111 \, \text{pA} \ (90 \, \text{mV})$; $\gamma$, 1.23 pS; N, 182,574; Po, 79.8%. Adjusted $R^2$, 0.940. (E) Normal residual after curve fit for tail current (corresponding to the green trace in C). Results: $i$, 0.061 pA (40 mV); $\gamma$, 1.53 pS; N, 178,014; Po, 68%. Adjusted $R^2$, 0.992.
Figure S6. **NSNA of an ensemble of macroscopic spHCN currents.** (A, top) Voltage protocol used for channel activation and deactivation. (bottom) 6 representative traces from 100 repeatedly collected traces. (B) Current variance over the complete time course of a single episode. (C) Current variance versus mean current amplitude. Red, parabola fit of the macroscopic current part. Green, parabola fit of the tail current part. (D) Normal residual after curve fit for macroscopic current (corresponding to the red trace in C). Results: $i, -0.451 \text{ pA} (-90 \text{ mV}); \gamma, 0.50 \text{ pS}; N, 33,980; Po, 74.4\%$. Adjusted $R^2$, 0.925. (E) Normal residual after curve fit for tail current (corresponding to the green trace in C). Results: $i, 0.011 \text{ pA} (40 \text{ mV}); \gamma, 0.27 \text{ pS}; N, 118,962; Po, 51.0\%$. Adjusted $R^2$, 0.947.
Figure S7. **NSNA of mHCN2-EGFP: Using the mean current as reference in the calculation of variance.** (A) 6 representative traces from 100 repeatedly collected traces. The same set of data as used in Fig. 4. (B) Current variance was calculated using the following equation instead of Eq. 7 in the main text:

\[
\sigma^2(t) = \frac{1}{M-1} \sum_{m=1}^{M} (i_m(t) - I(t))^2,
\]

where M is the total number of current traces, m is the index of current traces, \(i_m(t)\) is the raw current value at time \(t\), and \(I(t)\) represents averaged current trace. (C) Current variance versus mean current amplitude. Red, parabola fit of the macroscopic current part. Green, parabola fit of the tail current part. (D) Normal residual after curve fit for macroscopic current (corresponding to the red trace in C). Notice that to make the curve fit converge only a fraction of the data can be used. Results: \(i, -75.66\) pA (\(-130\) mV); \(\gamma, 582\) pS; \(N, 52\); Po, 50.0%. (E) Normal residual after curve fit for tail current (corresponding to the green trace in C). Results: \(i, -27.0\) pA (\(-40\) mV); \(\gamma, 674\) pS; \(N, 23\); Po, 76.3%.
Figure S8. **Selection of ROI and photobleaching of EGFP molecules.** (A) Selection of ROI from the fluorescence image of a membrane patch expressing spHCN-EGFP. (left) Brightfield image. (right) Fluorescence image. Yellow lines represent manually selected region for analysis. (B) Photobleaching of EGFP molecules. The same membrane patch was exposed to two sets of 25-50-100-200 (ms) laser pulses. The fluorescence intensity of the second laser pulse set was normalized by the values of the first set of laser pulses.

<table>
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<tr>
<th>Table S1. Summary of published results for HCN channel conductance</th>
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<tr>
<td><strong>HCN channels</strong></td>
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<tr>
<td>Native; HCN2 or HCN4?</td>
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<tr>
<td>Native; HCN1-like</td>
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**References**


